# **Thermo-Calc Documentation Set**

**Thermo-Calc Version 2018b** 





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# Contents

Installation Guide 1
Installing Thermo-Calc 2
System Requirements 3
Platforms Tested 3
Key to Using this Guide 4
Installation Terminology5
License and Installation Types 8
License and Installation Types9
About License Keys and Files9
Standalone Installations (SUNLL)10
Multi-User Network Installations (NWL)10
Consolidated Network Installations .10
Distributed Network Installations11
License Installations on a Server Network12
Preparing for Installation13
First Steps of Installation14
Default Directory Locations14
Thermo-Calc Default Directories 15
Subnet Information for Network Licenses18
Mac OS Installation Notes18
Linux Installation Notes
Installing Libgfortran19
openSUSE Administration (sudo) Users20

Opening Thermo-Calc in Linux20
Standalone Installations21
Typical Standalone Installations22
Request your license file from Thermo-Calc Software
Download your Thermo-Calc installation software22
Run the applicable installation file for your operating system22
Select the installation type and upload the license file23
Install the software and start using Thermo-Calc23
Add New Components to an Existing Installation24
Custom Standalone Installation 24
Complete Steps 1 to 4 of a typical standalone installation25
Select the installation type and what components to install25
Set the installation directory path, license file location and shortcuts
Install the software and start using Thermo-Calc27
Network Client Installations
Typical Network Client Installations 29
Install the licensing software 29
Complete Steps 1 to 3 of a typical standalone installation29
Choose the installation type and 29

enter the computer server details
Install the software and components29
Set Up the License Manager on Windows
Installation Requirements
Installing the License Manager on Windows31
Install the Sentinel RMS License Manager software 31
Choose the installation directory31
Choose the setup type and install the software31
Copy the license files to the destination directory32
Reboot or restart the license manager
Set Up the License Server on Linux .33
Installation Requirements
Installing the License Server on Linux
Stop any older versions of the License Server
Write a script to start the License Server
Confirm the HOSTNAME variable and open the UDP port35
Start the License Server35
Thin Client Installations35

Determining the Current License File Directory Path51
From Thermo-Calc51
Specifying a New Server Name or IP Number54
MATLAB <sup>®</sup> Installation Directory55
Firewalls55
Client/Server Mismatch56
Compare computer names on the client computer and server 56
Try these options to correct the issue
Mac OS Installations: Running the License Initialization Script57
Linux Installations: Running the License Initialization Script58
Uninstalling Thermo-Calc60
Uninstalling Thermo-Calc61
Uninstalling Standalone Versions of TC-PRISMA62
Quick Installation Guides64
Windows <sup>®</sup> Standalone Quick Install Guide65
Request a License File65
Install the Software67
Download and run your Thermo-Calc installation software67
Select the installation mode – typical or custom67
Customize your installation 68

Mac OS <sup>®</sup> Standalone Quick Install
Guide69
Request a License File70
Install the Software71
Download and install XQuartz71
Download and run your Thermo-Calc installation software71
Select the installation mode 72
TC-Python Quick Install Guide73
Install a Python Distribution73
Install Anaconda74
Install Thermo-Calc and the TC- Python SDK74
Install TC-Python75
Install an IDE (Integrated Development Environment)76
Open the IDE and Run a TC- Python Example76
Open the TC-Python Project in PyCharm76
Fixing Potential Issues with the Environment77
Updating to a Newer Version77
Mac OS: Setting Environment Variables78
Thermo-Calc User Guide 1
Introduction to Thermo-Calc 2
The Thermo-Calc Software
Add-on Modules 3

Typographical Conventions 3
Graphical Mode vs Console Mode 6
About the File Types and Formats 8
Graphical Mode 8
Console Mode 8
Opening Graphical Mode Project Files
How to Open any Project File 8
How to Open an Example Project
Opening Console Mode Macro Files
Opening a Macro From the Main Menu10
Opening a Macro Using a Command11
Drag and Drop a Macro File into the Console Window11
Examples, Manuals and Materials Folders
About the PDF Documentation Folder14
About the Examples17
Graphical Mode Examples 17
Console Mode Examples17
Examples by Product17
About the Material Files18
Austenitic Stainless Steel19
Duplex Stainless Steel19

High Speed Steel19
High Temperature Steel20
HSLA Steel20
Low Carbon Steel
Maraging Steel20
Martensitic Stainless Steel20
Tool Steel20
Training Steel20
Help Resources21
'Online' Help: No Internet Required 21
Context Help (Graphical Mode Only)21
Console Mode Help22
The Thermo-Calc Modules and Activities22
Displaying Thermo-Calc License Information25
Using This Guide27
The Graphical Mode Interface29
The GUI Layout30
Graphical Mode Windows 30
Project Activities and the Tree Structure31
Activity Successor and Predecessor Nodes31
Creating Activities and Successors32
Graphical Mode Workflow34
Main Menu, Toolbar and Keyboard Shortcuts35

Saving and Opening Project Files 38
Project Activity Nodes
Project and Activity Menus41
My Project Node41
Activity Node42
Organizing the Windows on the Desktop44
Moving Nodes in the Tree and Using the Grid45
Zooming In and Out46
Resetting a Plot to the Default View47
Node Status Markers47
Rotating 3D Plots48
Graphical Mode Projects49
The Quick Start Wizard50
Creating a Project with the Quick Start Wizard50
Creating a Project from a Template52
Cloning Activities and Trees53
Performing Projects, Trees and Activities55
About the Property Models in Thermo-Calc56
Models56
Calculation Types56
Property Model Development Framework57
Examples and Video Tutorials57
Using the Scheduler58

The Event Log	59
Graphical Mode Activities	61
System Definer	62
Configuration Settings	62
Creating a System Definer Activity	70
Property Model Calculator	70
Property Model Calculator Workflow	71
Defining the Property Model Calculator	71
Configuration Settings	73
About the Uncertainty Calculations	75
Phase Transition Model	80
Driving Force Model	80
Interfacial Energy Model	81
Estimation of Coherent Interfacial Energy	81
Coarsening Model	82
Particle Coarsening	82
Equilibrium Model	83
Equilibrium Calculator	83
Conditions Settings	84
Calculating with a Fixed Phase	.86
Options Settings	87
Single equilibrium and propert grid	-
Property Diagram	90

Phase Diagram90
Calculating and Plotting Functions91
Diffusion Calculator92
Available Options in Graphical Mode93
Diffusion Simulation Template .93
Using the Template93
Diffusion Calculator94
Demo Database Packages for the Examples Collection94
Demonstration (Demo) Mode94
Diffusion Simulation Template95
Diffusion Calculator95
Configuration Settings95
Configuration Settings95
Configuration Settings95 General97
Configuration Settings95 General97 Composition Unit97
Configuration Settings95 General97 Composition Unit97 Length Unit97
Configuration Settings95 General97 Composition Unit97 Length Unit97 Geometry97
Configuration Settings95 General97 Composition Unit97 Length Unit97 Geometry97 Options Settings97 Diffusion Calculator Plot
Configuration Settings
Configuration Settings
Configuration Settings

About the Homogenization
Functions103
Precipitation Calculator104
Available Options with the Precipitation Module104
Precipitation Simulation Template104
Precipitation Calculator 105
Demo Database Packages for the Examples Collection105
Demonstration (Demo) Mode .106
Configuration Settings
Show or Hide Details 107
Composition Unit107
Matrix Phase107
Precipitate Phase109
Calculation Type112
Numerical Parameters114
Particle Size Distribution (PSD)115
Accessing the 'Pre-existing size distribution' settings115
Defining the Preexisting size distribution
Plot Renderer117
Plot Settings117
Non-Isothermal Simulations120
Plot Renderer
Plot Renderer for the Add-On 122

Modules
Plot Toolbar122
Configuration Settings123
Examples of Calculation Type Differences123
Settings124
Plot Types126
Property Grid and Grid Calculations126
Uncertainty Calculations131
Plotting Options132
About Legend Styles135
Legend and Label Styles 136
Legend Styles136
Label Styles136
Plot Labels137
Plot Labels: Entering a Mixture of Plain Text and Equations138
Plot Labels: Rotation Angle and Show Anchor139
Moving Plot Labels139
Changing the Plot Properties140
Saving Plots140
Plotting Several Calculations in One Diagram141
Adding Axes to a Diagram142
Merging Plots into One Diagram144

Results Window Status Marker 146
Table Renderer146
Configuration Settings148
Setting Number Format and Number of Decimal Digits148
Saving Tabulated Data148
Setting Background Colours 149
Experimental File Reader149
Binary Calculator150
Configuration Settings150
Ternary Calculator152
Configuration Settings152
Calculation Types154
Scheil Calculator154
Configuration Settings154
Advanced Options and Settings155
The Console Mode Interface158
The Console Mode Layout
Main Menu Commands 161
Keyboard Shortcuts162
Create New Console or Result Tabs162
Navigate Between Tabs on Windows162
Pause, Resume or Stop Output to the Console Window
Console Window163

Editing the Default Console
Directory164
Console Mode Results Window166
Event Log Window166
Console Mode Workflow167
Performing Calculations in Console Mode168
Using the Command Line Prompt .168
Moving Between Modules and Submodules169
Entering Command Names or Abbreviations169
Specifying Parameters170
Wild Card Characters171
Command History173
Macro, Workspace and Experimental Data Files173
Log Files173
Macro Files174
Workspace Files177
Increasing the Buffer Size178
Defining a System in Console Mode178
Equilibrium Calculations181
About Equilibrium Calculations182
Setting Conditions for Equilibrium Calculations182
Calculating an Equilibrium184
Calculating an Equilibrium with a 184

About Phase Diagrams ......192

Fixed Phase

Calculating and Plotting a Phase Calculating a Quasi-Binary Phase Calculating a Quasi-Ternary Phase About the Scheil Module in Thermo-Calc ......197 Scheil Module Frequently Asked Simulating a Scheil-Solidification .. 199 Scheil Module State Variables ..... 201 T0 Temperature Simulations ...... 203 T0 Temperature Simulations ......204

204
206
207
207
210
211
211
212
214
215
216
219
221
223
224
225
226
228
229
230
230
232
233

About the Database Files	234
Encrypted vs Unencrypted Database Files	234
Database Initialization File	234
User Database Files	234
Database Installation Folder	.235
About Cached Database Files	235
Editing the Database Initialization File or Unencrypted Database File	236
Best Practice Recommendations	236
Database Checker	237
Changing Global Settings	238
General	.239
Graphical Mode - Default Units	241
Graphical Mode: Activities	.242
Global Settings: System Definer	242
Database name	243
Creating a Database Package	.244
Clearing a Database Package Message	245
Global Settings: Equilibrium Calculator	.246
Global Settings: Diffusion Calculator	.246
Global Settings: Precipitation Calculator	.246
Global Settings: Scheil Calculator	.247

Global Settings: Tables247
Global Settings: Graphical and Console Mode Plotting247
Titles, Legends and Labels248
Fonts and Background Colors 249
Lines and Grids249
Data Points and Anchors
Plot Area Size251
Global Settings: Console Mode Default Appearance251
Technical Notes
Technical Note: Pipe or Redirect Output Data from Console Mode . 254
Windows254
Linux254
Mac254
Mac254 Thermo-Calc Property Model
Mac254 Thermo-Calc Property Model
Mac254 Thermo-Calc Property Model Development Framework
Mac
Mac
Mac

General Commands	3
ВАСК	4
EXIT	4
GOTO_MODULE	4
HELP (? and ??)	4
HELP	5
INFORMATION	5
MACRO_FILE_OPEN	5
SET_INTERACTIVE	8
BINARY_DIAGRAM Commands	9
BINARY_DIAGRAM Module1	0
Database Commands1	1
AMEND_SELECTION1	2
APPEND_DATABASE1	2
DATABASE_INFORMATION14	4
DEFINE_ELEMENTS14	4
DEFINE_SPECIES	4
DEFINE_SYSTEM1	5
GET_DATA10	6
LIST_DATABASE10	6
LIST_SYSTEM	7
NEW_DIRECTORY_FILE1	7
REJECT	8
RESTORE	0
SET_AUTO_APPEND_DATABASE2	1
SWITCH_DATABASE2	2

REINITIATE_MODULE	52
RESTORE_ALL_WEIGHTS	52
SAVE_WORKSPACES	. 52
SELECT_EQUILIBRIUM	53
SET_ALL_START_VALUES	. 53
SET_ALTERNATE_CONDITION	. 55
SET_CONDITION	.58
SET_NUMERICAL_LIMITS	.60
SET_REFERENCE_STATE	62
SET_START_CONSTITUTION	64
SET_START_VALUE	.64
SET_WEIGHT	65
SHOW_VALUE	.66
STORE_ALL_WEIGHTS	.67
TABLE_HEAD, TABLE_VALUES and TABLE_END	.67
TRANSFER_START_VALUES	.69
Experimental Data Files	.70
Commands in the Experimental Data Files (*.POP, *.DOP)	. 71
TABLE_HEAD, TABLE_VALUES and TABLE_END	.73
COMMENT	.75
FLUSH_BUFFER	.75
Gibbs Energy System (GES) Commands	76
ADD_COMMENT	. 78
AMEND_ELEMENT_DATA	. 78

AMEND_PARAMETER	79
Solution Models in the GIBBS Module	82
AMEND_PHASE_DESCRIPTION .	89
AMEND_SYMBOL	93
CHANGE_STATUS	96
ENTER_ELEMENT	96
About the ENTER_PARAMETER Command	97
Defining the Parameter	98
ENTER_PARAMETER	101
About the ENTER_PHASE Command	103
ENTER_PHASE	106
ENTER_SPECIES	108
ENTER_SYMBOL	109
LIST_CONSTITUENT	114
LIST_DATA	115
LIST_PARAMETER	116
LIST_PHASE_DATA	117
LIST_STATUS	117
LIST_SYMBOL	120
READ_GES_WORKSPACE	121
REINITIATE	121
SAVE_GES_WORKSPACE	122
SET_R_AND_P_NORM	122
Optimizing with the PARROT Module	124

AMEND_PARAMETER	.126
AMEND_SYMBOL	.128
COMPILE_EXPERIMENTS	.131
CONTINUE_OPTIMIZATION	.132
CREATE_NEW_STORE_FILE	.133
EDIT_EXPERIMENTS	.133
About the ENTER_PARAMETER Command	. 133
Defining the Parameter	. 134
ENTER_PARAMETER	. 137
LIST_ALL_VARIABLES	139
LIST_CONDITIONS	. 140
LIST_PARAMETER	141
LIST_PHASE_DATA (from GIBBS) .	. 142
LIST_RESULT	. 142
LIST_STORE_FILE	147
LIST_SYMBOL_IN_GES	.147
OPTIMIZE_VARIABLES	. 148
READ_PARROT_WORKSPACES	148
RECOVER_VARIABLES	149
REINITIATE	149
RESCALE_VARIABLES	149
SAVE_PARROT_WORKSPACES	. 149
SET_ALTERNATE_MODE	. 150
SET_EXTERNAL_PARAMETER	. 150
SET_FIX_VARIABLE	151
SET_OPTIMIZING_CONDITION	152

LIST_STATUS	. 200
LIST_SYMBOLS	201
LOAD_INITIAL_EQUILIBRIUM	201
MAP	201
POST	. 203
READ_WORKSPACES	203
RECOVER_START_VALUES	204
REINITIATE_MODULE	204
SAVE_WORKSPACES	204
SELECT_EQUILIBRIUM	206
SET_ALL_START_VALUES	206
SET_AXIS_VARIABLE	208
SET_CONDITION	. 209
SET_INPUT_AMOUNTS	212
SET_NUMERICAL_LIMITS	. 212
SET_REFERENCE_STATE	214
SET_START_CONSTITUTION	216
SET_START_VALUE	. 216
SHOW_VALUE	. 217
STEP_WITH_OPTIONS	217
TABULATE	. 230
Plotting and Visualization with the POST Module	231
Using the POST Module	. 233
ADD_LABEL_TEXT	234
APPEND_EXPERIMENTAL_DATA .	236
CHANGE_LEGEND	237

DUMP_DIAGRAM	.237
ENTER_SYMBOL	.238
FIND_LINE	. 241
LIST_DATA_TABLE	. 241
LIST_PLOT_SETTINGS	. 242
LIST_SYMBOLS	.242
MAKE_EXPERIMENTAL_DATAFILE	243
MODIFY_LABEL_TEXT	.243
PLOT_DIAGRAM	.244
PRINT_DIAGRAM	.244
QUICK_EXPERIMENTAL_PLOT	. 245
REINITIATE_PLOT_SETTINGS	. 245
RESTORE_PHASE_IN_PLOT	. 245
SELECT_PLOT	. 246
SET_AXIS_LENGTH	.246
SET_AXIS_PLOT_STATUS	. 246
SET_AXIS_TEXT_STATUS	.247
SET_AXIS_TYPE	. 247
SET_COLOR	. 247
SET_CORNER_TEXT	.249
SET_DIAGRAM_AXIS	.249
SET_DIAGRAM_TYPE	.252
SET_FONT	.252
SET_INTERACTIVE_MODE	253
SET_LABEL_CURVE_OPTION	. 253
SET_PLOT_FORMAT	.254

SET_PLOT_OPTIONS	255
SET_PLOT_SIZE	. 257
SET_PREFIX_SCALING	. 257
SET_RASTER_STATUS	. 257
SET_REFERENCE_STATE	258
SET_SCALING_STATUS	. 260
SET_TIC_TYPE	260
SET_TIELINE_STATUS	261
SET_TITLE	261
SET_TRUE_MANUAL_SCALING	261
SUSPEND_PHASE_IN_PLOT	. 262
TABULATE	. 262
POTENTIAL_DIAGRAM Module	. 263
POTENTIAL_DIAGRAM	264
POURBAIX_DIAGRAM Commands	.265
POURBAIX_DIAGRAM Commands	266
Getting started with the Pourbaix module	
What is a Pourbaix diagram?	. 266
What database should I use?	266
Choosing a custom aqueous solution database	. 266
Copy and Rename the POLY3 file	268
Enforce a Pause	. 268
Pourbaix Main Options	. 269
POURBAIX_DIAGRAM Commands for Options 1 to 4	.269

Option 1: Start a completely new Pourbaix diagram	
calculation	269
Chemical system (elements)	270
Initial bulk composition?	270
Option 2: Open an existing file and plot other property diagrams	275
Option 3: Open an existing file and make another Pourbaix calculation	277
Option 4: Open an Existing file and make another Stepping calculation	279
REACTOR_SIMULATOR_3 Commands	282
CREATE_DIVIDERS	283
CREATE_PIPES	283
CREATE_STAGE_BOXES	284
EXECUTE_POLY3_COMMAND?	285
LIST_RECORDS	286
READ_WORKSPACE	286
SAVE_WORKSPACE	286
START_SIMULATION	287
SCHEIL Module Commands	290
BCC_TO_FCC_TRANSFORMATION	291
EVALUATE_SEGREGATION_ PROFILE	291
GLOBAL_MINIMIZATION	292
LIQUID_PHASE_NAME	292

SAVE_FILE_NAME	292
SET_INTERACTIVE	292
SET_NUMERICAL_LIMITS	293
START_WIZARD	294
TEMPERATURE_STEP	295
TERMINATION_CRITERIA	295
System Utilities Commands	296
ABOUT	297
CLOSE_FILE	297
DISPLAY_LICENSE_INFO	297
HP_CALCULATOR	297
OPEN_FILE	298
SET_COMMAND_UNITS	298
SET_ECHO	299
SET_GES_VERSION	299
SET_INTERACTIVE_MODE	299
SET_LOG_FILE	300
SET_PLOT_ENVIRONMENT	300
SET_TC_OPTIONS	301
STOP_ON_ERROR	302
TABULATION_REACTION	
Commands	
ENTER_FUNCTION	
ENTER_REACTION	
LIST_SUBSTANCES	
SET_ENERGY_UNIT	
SET_PLOT_FORMAT	309

SWITCH_DATABASE	. 310
TABULATE	311
TABULATE_DERIVATIVES	.311
TABULATE_REACTION	. 316
TABULATE_SUBSTANCE	319
TERNARY_DIAGRAM Commands	.325
TERNARY_DIAGRAM Module	326
Thermodynamic Variables and Units	.327
Thermodynamic Variables	.328
Common Thermodynamic Variables	.328
How to Read the Tables	. 329
Intensive Variables	329
Extensive Variables	.330
Special Quantities	.335
The u-Fraction Variable	335
Suffixes	.336
Normalizing Suffixes	336
Derived Variables and Partial Derivatives	. 342
Defining New Derived Variables and Functions	. 344
Derived Variables for Aqueous Solutions	. 345
Defining New Derived Variables for Aqueous Solutions	.349
Thermodynamic Variables and Wildcard Characters	. 349

Units of State Variables and	
Derived Variables	.350
User-Specified Units	.350
Examples of User-Defined Units	. 351
For intensive variables of a defined system	.351
For extensive variables of a defined system	.352
For intensive variables of a system component	.353
For extensive variables of a system component	.354
For intensive variables of a species in a phase	.357
For extensive variables of a species in a phase	.358
Graphical Mode Examples Guid	e . 1
About the Graphical Mode Examples	2
Watch the Video Tutorials	2
Opening a Project File	3
Thermo-Calc Graphical Mode Examples	4
T_01: Calculating a Single-Point Equilibrium	4
T_02: Stepping in Temperature in the Fe-C System	6
T_03: Fe-C Phase Diagrams	8
T_04: Fe-Cr-C Ternary Phase Diagram at 1000 K	10

T_05: Stable and the Metastable Fe-C Phase Diagrams1	3
T_06: Serially Coupled Equilibrium Calculators1	4
Table Renderer Results1	4
T_07: User-Defined Functions1	8
T_08: Scheil and Equilibrium Solidification2	0
T_09: Carbide Driving Force Heat Map2	3
T_10: Phase Transition2	3
T_11: Coarsening and Interfacial Energy2	6
Plot Results2	6
T_12: Driving Force and Interfacial Energy2	9
Diffusion Calculator Examples 3	2
Diffusion Calculator Examples 3 D_01 Homogenization of a Binary Fe-Ni Alloy	
D_01 Homogenization of a Binary	2
D_01 Homogenization of a Binary Fe-Ni Alloy3 D_02 Ferrite(bcc)/Austenite(fcc) Transformation in a Binary Fe-C	2
D_01 Homogenization of a Binary Fe-Ni Alloy	2 4 7
D_01 Homogenization of a Binary Fe-Ni Alloy	2 4 7 8
D_01 Homogenization of a Binary Fe-Ni Alloy	2 4 7 8

Precipitation Calculator Examples .	.52
P_01: Isothermal Precipitation of AI3Sc	52
Plot Results	.53
P_02: Stable and Metastable Carbides - Isothermal	54
Plot Results	.55
P_03: Stable and Metastable Carbides - TTT Diagram	56
Plot Results	.58
P_04: Precipitation of Iron Carbon Cementite	.59
Plot Results	.60
P_05: Precipitation of γ' in Ni Superalloys - Isothermal	.61
Plot Results	.62
P_06: Precipitation of γ' in Ni Superalloys - Non-isothermal	.64
Plot Results	.66
P_07: Cooling Rate (CCT) Diagram of Ni-Al-Cr γ-γ'	. 68
Plot and Table Results	.70
P_08: Precipitation of Cu-Ti CU4TI1 with Assumptions of Sphere and Needle Morphologies	
Plot Results	.73
P_09: Precipitation of Al-Sc AL3SC with Assumption of Sphere and Cuboid Morphologies	76
Plot Results	

P_10: Initial Particle Size	
Distribution of Fe-Cr-C	. 81
Plot Results	84
Volume Fraction with Initial Particle Size Distribution	.84
Volume Fraction with No Initial Particle Size Distribution	85
P_11: Interfacial Energy Function .	85
Plot Results	87
Mean Radius	.87
Thermo-Calc Console Mode Examples Guide	1
About the Thermo-Calc Console Mode Examples	2
Examples of Binary Systems	4
Examples of Ternary Systems	9
General Examples Using the Steel Database	. 13
Examples of Property and Potentia Diagrams	
Property Diagrams	.15
Potential Diagram	15
Examples of Multicomponent Systems	16
Examples Using Functions and Variables	. 18
Examples of the Gibbs Energy System (GES) Module	. 20
Notes related to the example .	. 22

Amending the binary excess model22
Amending the ternary extrapolation model23
Examples of Scheil- Gulliver Calculations24
Example using the PARROT and EDIT_EXPERIMENT Modules26
Example of the Tabulate Reaction (TAB) Module
Example of the Reactor Module 29
Examples with Specialized Databases
Pourbaix Module Examples
The PAQ2 Database33
Additional Examples in this Series34
Step 1: Single-Point Calculations for H2O-NaCl system34
Step 2: Single-Point Calculations for Fe-X (X = Cr- Ni-Co) system34
Step 3: Single-Point Calculations for Fe-Cr-Ni-Co + H2O-NaCl system34
Step 4: Pourbaix Diagram Mapping for Fe-Cr-Ni-Co + H2O-NaCl system - define the variables34
Pourbaix Module Examples TCEX40A to 40E
TCEX 40A
TCEX 40B

TCEX 40C
TCEX 40D
TCEX 40E
Data Optimization User Guide 1
Introduction 2
Introduction to Data Optimization 3
Data Optimization 3
The Least-Squares Method4
The CALPHAD Approach5
Working with Data Optimization in Thermo-Calc
Data Optimization in Thermo-Calc 7
Data Optimization Workflow7
The PARROT Module
The EDIT_EXPERIMENTS Module 8
Data Optimization in Other Modules
Optimization File Types
Optimization Workspaces
Method for Optimization and Simulation11
Creating the POP-File
Experimental Information13
POP-File Syntax15
Entering Experimental Information 16
Creating a New Equilibrium17
Setting Equilibrium Conditions and its Uncertainty17

Setting Reference States18
Entering the Experimental Value and its Uncertainty19
Entering Many Equilibria as a Table19
Setting Site Fraction Start Values21
Commenting About an Equilibrium21
Simultaneous Use of Binary and Ternary Experiments
Using Stability Conditions22
Grouping Equilibria with Labels23
Dividing the POP-File into Several Blocks23
The Setup File
Creating the Setup File26
Defining the System26
Creating Additional Composition Sets of a Phase27
Entering the Optimizing Variables27
Entering Optimizing Variables in Parameter Specifications27
Entering Interaction Parameters28
Initializing Optimizing Variables in PARROT29
Creating the Workspace File30
Compiling the POP-file
Returning Control to the Console31

Optimizing in PARROT	. 32
Optimizing the System in PARROT	33
Changes that Require POP-file Recompilation	.33
Optimization Workflow	. 33
Ensuring Computation of all Equilibria	.34
Setting Weights	. 35
Optimizing and Evaluating Results	. 36
The Critical Set of Experiments	. 37
Continuing the Optimization and Resetting Variables	. 37
Updating the Set of Optimizing Variables	. 39
Reducing the Number of Optimizing Variables	. 40
Plotting Intermediate Results	.40
Superimposing Experimental Data from EXP-files	.41
Finalizing the Optimization	.42
Finishing the Optimization	43
Rounding off Optimizing Variable Values	.43
Updating the Database File	. 43
Creating a Database File	44
Updating the Setup File and the POP-file	.44
Alternate Mode	.45
About Alternate Mode	46

	Optimizing in Alternate Mode	.47
	Preparing the POP-file for Alternate Mode	47
	Examples of SET_ALTERNATE_ CONDITION	47
	Entering Other Experiments in the POP with Alternate Mode	48
Т	roubleshooting	50
	Useful Guidelines	51
	Excluding and Including the Correct Equilibria	
	Conflicting Data	52
	xperiments and Experimental Data	53
	Phase Diagram Data	54
	Data for Individual Compounds	55
	Single Phase Mixing Enthalpies or Partial Enthalpies	.57
	Enthalpies of Formation and Enthalpies of Reactions	57
	Chemical Potentials via EMF Measurements	58
	Driving Force for Metastable Phases	.58
	Pressure-Temperature-Volume EOS Data	
	Crystal Structure, Lattice Parameters and Site-Occupancy	59
	Magnetism and Other Atomistic Properties	60
	Data About Systems of Different	61

Orders .....

Thermodynamic Models62
Gas63
Compound Energy Formalism63
Substitutional Liquid64
Ionic Two-Sublattice Liquid Model64
Models for Ordered Phases65
4SL Model for FCC, HCP and BCC66
2SL Model for FCC, HCP and BCC67
Magnetic Ordering67
Molar Volumes and High Pressure .68
Excess Models69
Pourbaix Diagrams1
Introduction to Pourbaix Diagrams . 2
About Pourbaix Diagrams in Thermo-Calc 3
Pourbaix Diagrams 3
Working with Pourbaix Diagrams 5
Key Concepts 6
Effective Interaction Ratio6
Solubility7
Basic Settings and Definitions7
Thermodynamic Data Requirements10
Required Thermodynamic Data11
Including or Excluding the Gas Phase11

Examples of Pourbaix Diagrams for Fe1	6
Pourbaix Diagrams with Gas Phase Excluded1	7
Pourbaix Diagrams with Gas Phase Included1	7
Other Examples of Pourbaix Diagrams	9
Variations of Pourbaix Diagrams 2	0
Pourbaix Diagrams for Complex Alloys2	2
Pourbaix Diagram References2	4
DATAPLOT User Guide	1
Introduction to the DATAPLOT User Guide	2
Using this Guide	3
About the DATAPLOT Graphical Language	3
DATAPLOT Examples	3
Plotting Experimental Data Files	4
Graphical Mode	4
Console Mode	4
Using DATAPLOT Graphical Language	6
DATAPLOT File Structure	7
DATAPLOT Language Syntax	7
Coordinate Systems	7
Graphical Operation Codes	8
Tables or Blocks	9

Drawing a Polygon	10
Drawing an Analytical Function	10
Painting of an Enclosed Area	10
Writing a Text	10
Plotting a Symbol	11
Other Commands	11
General	11
ATTRIBUTE	11
CLIP	12
COLOR	12
INCLUDE	12
LINETYPE	12
Interactive Plotting	12
Formatting DIGLAB Symbols in LaTeX Documents	12
LaTeX Formatting Codes	14
LTEXT vs LaTeX Commands for Labels	14
PROLOGUE Commands	15
PROLOG	16
DIAGRAM_TYPE	16
TIC_TYPE	16
TITLE	16
XLENGTH	17
XSCALE	17
XTEXT	17
ХТҮРЕ	

YLENGTH1	.7
YSCALE1	.8
YTEXT1	.8
YTYPE1	.8
DATASET Commands1	9
ATTRIBUTE2	0
BLOCK2	0
BLOCKEND2	1
CHARSIZE2	1
CLIP2	1
COLOR2	2
DATAPOINT2	3
DATASET2	3
DRAWLINE2	4
FONT 2	4
Font Numbers2	4
FUNCTION2	5
GLOBALSIZE2	5
INCLUDE2	5
LINETYPE2	6
Line Types2	6
PAINT2	6
PCFUNCTION2	7
STRING2	7
SYMBOLSIZE2	9
TEXT2	9

Examples of DATAPLOT Files and Outputs
Example 1 – Draw Lines and Symbols31
Output Example31
Macro Text31
Example 2 – Draw Polygons and Symbols
Output Example32
Macro Text33
Example 3 – Using Strings and Various Line Types34
Output Example34
Macro Text34
Example 4 – Draw Curves Defined by Functions36
Output Example37
Macro Text37
Example 5 – Use Included Files for Predefined Symbols
Output Example
Macro Text39
Example 6 – Triangular Diagrams for Ternary Systems40
Output Example40
Macro Text41
Example 7 – Color Codes46
Output Example47
Macro Text47

Example 8 - DIGLIB Symbols5	1
Output Example52	2
Macro Text52	2
Example 9 - DIGLIB Fonts and Lines 59	9
Output Example59	9
Macro Text59	9
Database Manager User Guide 2	1
Introduction to the Database Manager Guide	2
Thermodynamic and Kinetic/Mobility Databases	3
Initialization of the DATA Module 3	3
Initiation Parameter Examples	4
The Database Definition File and Keywords	6
Database Definition File Syntax	8
ELEMENT	9
SPECIES10	0
PHASE1	1
Examples of Recommended Suffixes1	1
GIBBS Phase-type Codes12	2
Additional clarification13	3
CONSTITUENT10	6
ADD_CONSTITUENT1	7
COMPOUND_PHASE18	8
ALLOTROPIC_PHASE18	8
TEMPERATURE_LIMITS19	9

DEFINE_SYSTEM_DEFAULT1	.9
DEFAULT_COMMAND1	9
DATABASE_INFORMATION2	0
TYPE_DEFINITION2	1
FTP_FILE2	4
FUNCTION2	5
PARAMETER2	7
GIBBS Parameter Name2	9
OPTIONS3	4
TABLE	5
ASSESSED_SYSTEMS3	5
REFERENCE_FILE3	8
LIST_OF_REFERENCE	9
ADD_REFERENCE4	0
CASE and ENDCASE4	1
VERSION_DATA4	2
Working with the Mobility Databases4	3
About the Diffusion Module (DICTRA) and the Databases4	4
Extensions to Database Definition File Syntax4	5
PARAMETER4	5
DIFFUSION4	6
ZEROVOLUME_SPECIES4	7
Database Definition File Examples4	8
Example 1: A Steel Database4	.9

Example 2: A Custom Database for the Sb-Sn Binary System	50
Example 3: A Public Database for the Fe-Cr-C Ternary System	53
The Database Checker6	68
About the Database Checker6	69
Database Checker Revision History 6	59

# **Installation Guide**

Windows<sup>®</sup>, Mac OS<sup>®</sup> or Linux Operating Systems

**Thermo-Calc Version 2018b** 





## **Installing Thermo-Calc**

### In this section:

System Requirements	3
Platforms Tested	3
Key to Using this Guide	4
Installation Terminology	. 5

### **System Requirements**

A typical installation installs Thermo-Calc and the latest version of the databases.

Additional add-on modules include:

- Diffusion Module (DICTRA)
- Precipitation Module (TC-PRISMA)

You can also install these Software Development Kits (SDKs):

- TQ-Interface (Windows and Linux)
- TC-API (Windows and Linux)
- TC-Toolbox for MATLAB<sup>®</sup> (Windows only)
- TC-Python (all OS)



System requirements and program availability are available on the web: <a href="http://www.thermocalc.com/products-services/software/system-requirements/">http://www.thermocalc.com/products-services/software/system-requirements/</a>

Also see our roadmap for platform support http://www.thermocalc.com/products-services/software/systemrequirements/platformroadmap/

### **Platforms Tested**

Thermo-Calc and its components are tested on these platforms:

Windows	Мас	Linux <sup>1, 2</sup>
Windows 7 64-bit Windows 10 64-bit	Mac OS X Sierra 10.13.4	<ul><li>Ubuntu 16.04 (Xenial)</li><li>CentOS 7.0</li><li>OpenSUSE 42.2</li></ul>

<sup>1</sup> Thermo-Calc Software cannot guarantee full functionality if the software is installed on other Linux distributions.

<sup>2</sup> If you are installing on CentOS 7, it is recommended you install gtk2 before you run the installation script.

### Key to Using this Guide

This installation guide is for all Thermo-Calc installations running on Windows, Mac OS or Linux operating systems. It is also applicable to demo (trial) installations as well as previously installed versions of Thermo-Calc (the new version is installed alongside the old version).

This guide includes information about:

- The types of licenses and installations available.
- Installing the Thermo-Calc software and its components on stand-alone computers, servers and client computers.
- Installing the license management software for network installations.
- Retrieving and installing license files.
- Troubleshooting to solve common installation problems.

Name	Description
Information	Provides important information and indicates that more detail is located in the linked or named topic.
Note	The information can be of use to you. It is recommended that you read the text or follow the link.
Operating system	The operating system is indicated when the information is specific to a <i>Windows, Mac OS</i> or <i>Linux</i> operating system.
License file	The license file, which contains the license keys and is installed on either a standalone computer or a network server.
A Thermo-Calc	The Thermo-Calc software and its components (including databases) to be installed.

### ICONS IN THIS GUIDE

#### **TYPOGRAPHICAL CONVENTIONS**

Convention	Definition
Forward arrow →	The forward arrow symbol $\rightarrow$ instructs you to select a series of menu items in a specific order. For example, <b>Tools</b> $\rightarrow$ <b>Options</b> is equivalent to: From the <b>Tools</b> menu, select <b>Options</b> .
<b>Boldface</b> font	A <b>boldface</b> font indicates that the given word(s) are shown that way in on a toolbar button or as a menu selection. For example, if you are told to select a menu item in a particular order, such as <b>Tools</b> $\rightarrow$ <b>Options</b> , or to click <b>Save</b> .
Italic font	An <i>italic</i> font indicates the introduction of important terminology. Expect to find an explanation in the same paragraph or elsewhere in the guide.
code and code bold	A code font indicates you are to make a keyboard entry. It also shows a programming code or code examples. The code bold font highlights the entry.

# Installation Terminology

Term	Definition
client computer	A computer without a license file of its own but connected through the network to a computer with the license information. The client computer receives a license token from the computer where the license management software is installed.
consolidated network	A type of network installation where the Thermo-Calc software and a Thermo-Calc licensing software are installed on a single server computer. This allows all computers in the network to access both the software and a license on the server via a client installation.
distributed network	A type of network installation where the Thermo-Calc software is installed on one server computer in a network and the Thermo-Calc licensing software is installed on a different computer. After a thin client installation, all computers in the network can access both the software and the network license key via the server.
host ID	The MAC/Ethernet Address on the computer that a license is locked to.

Term	Definition
license file	The file (called <i>lservrc</i> ) that contains all the purchased license keys. It is is issued by Thermo-Calc Software, usually by email.
license key	The information contained in the license file that represents the products purchased and which is then authorised for installation.
License Manager (Windows)	The Windows program used for license management: it distributes license tokens as specified in the network license file issued by Thermo-Calc Software.
License Server (Linux)	The Linux program used for license management: it distributes license tokens as specified in the network license file issued by Thermo-Calc Software.
license server	The use of the term <i>license server</i> (lower case letters) is reserved for use for scripts and option files (for example, in an <u>"Unattended Network</u> <u>Client Installations" on page 39</u> ). For the purposes of the step-by-step instructions it is not used to refer to the type of network installation that runs the license management program. Also see <i>License Manager</i> (Windows) and <i>License Server</i> (Linux).
licensing software	The Thermo-Calc license management software. When working in Windows it is called <i>License Manager</i> (full name is Sentinel RMS License Manager by SafeNet) and on Linux it is called <i>License Server</i> (full name is Sentinel License Sever by Safenet).
Network License (NWL)	The license type used with the licensing software. The NWL can only be used on one specific computer that hosts the licensing software. The Network License uses information about IP addresses and IP ranges to determine which client computers receive license tokens. The software may be used on the number of sites you have paid for and by the number of persons you have paid for on any number of network connected computers
standalone computer	A computer which uses a SUNLL license type.

Term	Definition
Single-User- Node Locked- License (SUNLL)	A unique license file for one designated computer at one designated site. It cannot be copied or moved outside the site. If the software is installed on a portable computer, it can be temporarily used off-site. The license can be used without network access (to the License Server).
thin client	An installation where the software is installed on a file server but is run from a client computer. The licensing software may be installed on the file server (consolidated network installation) or on a different computer (distributed network installation).
XQuartz	An open-source X Windows System, which is required to run Thermo- Calc on a Mac.

## License and Installation Types

#### In this section:

License and Installation Types	9
About License Keys and Files	9
Standalone Installations (SUNLL)	10
Multi-User Network Installations (NWL)	10
Consolidated Network Installations	10
Distributed Network Installations	11
License Installations on a Server Network	12

## **License and Installation Types**

Installation procedures are based on the license type, the installation type and the operating system.



**Demo (trial) installations**: Only a standalone installation is available and a license file is automatically installed.

License type	Installation Type	Windows	Mac OS	Linux
SUNLL	Standalone	Yes	Yes	Yes
	Consolidated network	Yes	No	Yes
NWL	Distributed network	Yes	No	Yes
	License installations on a server network	Yes	Yes <sup>1</sup>	Yes

<sup>1</sup> For Mac OS, the licensing software is installed on a Windows or Linux computer server.

www.thermocalc.com/purchase/licensing-options/

### **About License Keys and Files**



**Demo (trial) installations**: A license file is automatically installed with the software. You do not need to request it.

To run Thermo-Calc and its components you need a valid license key for each product. The license keys are included in the license file emailed to you by Thermo-Calc Software.

You can copy the license file to the applicable computer(s) before or after installing the software—the license file itself is not required during the installation process. However, it is required to make calculations in Thermo-Calc.

<u>"Request a License File from Thermo-Calc Software" on page 42</u>

### **Standalone Installations (SUNLL)**

The standalone installation is used with the Single-User Node-Locked License (SUNLL). The

Thermo-Calc software A and a Thermo-Calc license file are installed on a standalone computer. You can do a typical (all operating systems) or a custom installation (Windows and Linux only).



To get started, go to <u>"First Steps of Installation" on page 14</u> and then <u>"Standalone</u> Installations" on page 21.

### **Multi-User Network Installations (NWL)**

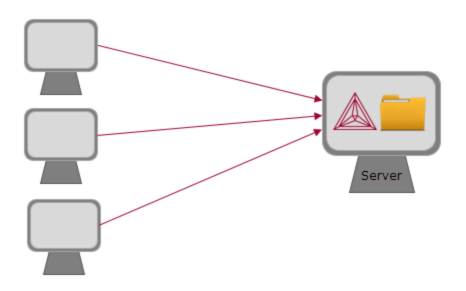
There are different types of multi-user network installations: consolidated, distributed and where the licensing software is installed on a computer network. There is also a subtype of installation called thin client, which is a step done with the consolidated and distributed network installations.

-	-
	Ξ

Thin client installations: A thin client installation is where the Thermo-Calc software is installed on a computer server but is run from a client computer. The licensing software can be installed on the computer server (a consolidated network) or on a different computer (a distributed network). This type of installation is available for Windows and Linux operating systems. The client computer must have the Thermo-Calc software installed. This type of installation mainly sets environmental variables and paths on the client computer.

### **Consolidated Network Installations**

The consolidated network installation is used with a Network License (NWL) and it is available for Windows and Linux operating systems.



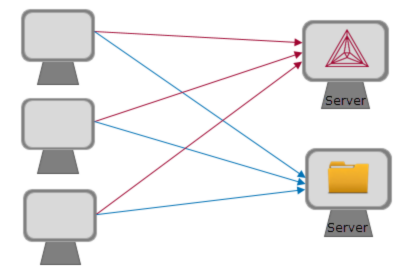
The Thermo-Calc software 🚵 and the license management program are installed on the

same server where the license file is also copied. A thin client installation allows all computers (the clients) in the server network to access both the software and the licensing on the same computer server.

To get started, go to "First Steps of Installation" on page 14 and then "Network Client Installations" on page 28.

# **Distributed Network Installations**

The distributed network installation is used with a Network License (NWL) and is available for Windows and Linux operating systems.



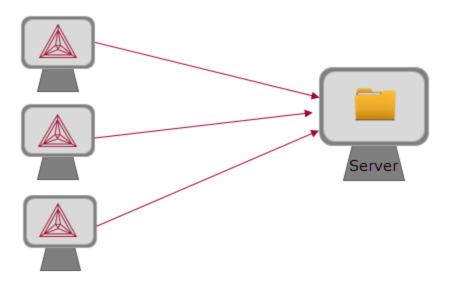
The Thermo-Calc software 📥 is installed on one computer server in a network and the

licensing software is installed on a different computer server where the license file is also copied. A thin client installation allows all computers (the clients) in the server network to access both the software and the licensing on the two computer servers.

To get started, go to <u>"First Steps of Installation" on page 14</u> and then <u>"Network Client</u> Installations" on page 28.

## **License Installations on a Server Network**

When a license file is installed on a server network, it is used with a Network License (NWL). It is available for all platforms. However, for Mac OS the server containing the licensing software must be set up on Windows or Linux.



The licensing software is installed on a computer server where the license file is also copied. The Thermo-Calc software is installed on every computer in the network that needs it.

To get started, go to <u>"First Steps of Installation" on page 14</u> and then <u>"Network Client</u> <u>Installations" on page 28</u>.

# **Preparing for Installation**

# In this section:

First Steps of Installation	14
Default Directory Locations	14
Thermo-Calc Default Directories	15
Subnet Information for Network Licenses	18
Mac OS Installation Notes	18
Linux Installation Notes	19
Installing Libgfortran	19
openSUSE Administration (sudo) Users	20
Opening Thermo-Calc in Linux	20

# **First Steps of Installation**

This section reviews some of the things you need to do before installing the Thermo-Calc software.

- Learn about the default directories where the software and license files are installed.
- For network installations, collect subnet information.
- For Mac, install XQuartz and make sure the security preferences are set correctly.
- For Linux, install libgfortran 4.4 (or higher).
- For a Linux Cent OS 7 installation, also install gtk2 before you run the installation script.

yum install gtk2



**Should I get my license file before or after software installation?** The license file itself is not required during the installation process (and not at all for demo installations). However, it is required to make calculations in Thermo-Calc. The license file has details about your computer(s) that in some cases cannot be determined accurately until *after* installation (especially for a network installation). For most users it is recommended you request the file before you start installing.

# **Default Directory Locations**

The default directories where the software and its components are installed, including where the license file is stored, vary by platform and user type.

If you want to select specific components to install or if you want to install the software in a non-default installation directory, then do a custom installation (Windows and Linux only). In the table, *<user>* stands for the user name and *<version>* for the version of Thermo-Calc, for example 2018b.

# **Thermo-Calc Default Directories**

# WINDOWS

OS	User type	Default directory
	Normal user	Users\< <i>user</i> >\Thermo-Calc\< <i>version&gt;</i> Users\< <i>user</i> >\Documents\Thermo- Calc\< <i>version&gt;</i> My documents
	Administrator	Program Files\Thermo-Calc\< <i>version&gt;</i> Users\Public\Public Documents\Thermo- Calc\< <i>version&gt;</i>
Windows	Custom installation (for all users)	Users\Public\Public Documents\Thermo-Calc\ <version></version>
	License software installations on a server network (the License Manager software)	C:\Program Files (x86)\Common Files\SafeNet Sentinel\Sentinel RMS License Manager\WinNT\

## MAC

OS	User type	Default directory
Mac	Administrator (user name and password required)	In the <b>Applications</b> folder: Thermo-Calc-< <i>version</i> >.app, Thermo-Calc-< <i>version</i> >-Demo.app, and, uninstall-Thermo-Calc- < <i>version</i> >.app Examples and manuals in /Users/Shared/Thermo- Calc/< <i>version</i> > To go to this folder, in Finder, from the <b>Go</b> main menu select <b>Go to folder</b> . Enter the above file path and click <b>Go</b> .

#### LINUX

OS	User type	Default directory
	Non root user	home/ <user>/Thermo-Calc/<version></version></user>
	Root user	usr/local/Thermo-Calc/ <version></version>
Linux	License software installations on a server network (the License Server software)	If you are doing a network installation then these files are available as a tar-file from the Thermo-Calc Software web site. In the downloaded tar-file, there is a <b>restart_lserv</b> script file, which can be used as a template for creating a script that starts up the License Server.

### DATABASES

Depending on the type of license and operating system, the encrypted Thermo-Calc databases and the database initialization file are located in a subfolder to the Thermo-Calc installation. For example, in a Windows standalone installation, you can find these in a folder called **data** (file path is *C*:\*Program Files*\*Thermo-Calc*\*<version>*\*data*).

OS (C:) $\rightarrow$ Program Files $\rightarrow$	Thermo-Calc	→ <version> &gt; data</version>
Name	Date modi	Туре
ALDEMO	2017-09-2	File folder
AQS2	2017-09-2	File folder
CUDEMO	2017-09-2	File folder
FEDAT	2017-09-2	File folder
FEDEMO	2017-09-2	File folder

### **TC-TOOLBOX FOR MATLAB**

Examples for the TC-Toolbox are placed under MATLAB in the same documents folder as the Thermo-Calc files (My Documents or Public Documents):

```
..\Documents\MATLAB\Thermo-Calc-Toolbox-X\Examples
```

Where X is the installed version number of the toolbox.

For most installations the examples are available in the MATLAB window when the software is opened.

### DATABASE CHECKER PROGRAM

The Thermo-Calc software package includes a program to check that the syntax of unencrypted database files is correct. The program applies the syntax rules set out in the *Database Manager User Guide* and reports errors and issues warnings. This program is intended for advanced users who develop and manage databases.

Thermo-Calc accepts deviations from these syntax rules. This means that a database can work even if the Database Checker reports errors and warnings. For example, an error is reported if an abbreviated phase name is found, but phase name abbreviations are accepted by Thermo-Calc and its add-on modules.

The executable **DatabaseChecker** file is found in the Thermo-Calc home directory. The program can also be launched by selecting **Tools**-> **Database Checker** from the main menu.

### **CUSTOM PROPERTY MODELS**

All property models (whether these are predefined or custom models) must be located in a subdirectory of the property model directory. You can go to **Tools** → **Options** and the **Model directory** field displays the default directory.

Predefined models included with the Property Model Calculator are installed in the locations below based on your operating system. If you create your own models, this is the default directory where these are saved. You can also use one of these as a template to create your own property model.

Operating system	Property Model installation directory	
Windows	C:\Users\Public\Public Documents\Thermo-Calc\ 2018b \PropertyModels	
Linux	/home/user/Thermo-Calc/2018b/PropertyModels	

Operating system	Property Model installation directory
Mac	/Users/Shared/Thermo-Calc/2018b/PropertyModels

# **Subnet Information for Network Licenses**

For network licenses, the subnet information is also required. Contact your company's network administrator for help as required. Additional requirements are included elsewhere for network installations.

- Subnets should be in the format aaa.bbb.ccc\*\*\* (for example: 192.168.29.0/24 or 192.168.29.\*
- A maximum of six subnets are permitted.
- An alternative way to get this information is to run the command:
- Windows: ipconfig /all
- Mac / Linux: ifconfig

# **Mac OS Installation Notes**

## **RESTARTING AFTER AN INSTALLATION**

In some types of installations (e.g. a network installation), you may have to restart your Mac to use Thermo-Calc for the first time.

## INSTALLING XQUARTZ

The XQuartz open-source X Windows system is required to run Thermo-Calc on Mac OS X but it is not required to install Thermo-Calc. Before installing Thermo-Calc, you may also need to ensure that the computer security permissions are set.

- 1. In a web browser go to <a href="https://www.macupdate.com">https://www.macupdate.com</a> or to download it directly from XQuartz, <a href="http://xquartz.macosforge.org/landing/">http://xquartz.macosforge.org/landing/</a>.
- 2. In the Search Mac Apps field, enter XQuartz.
- 3. Click the **XQuartz** search result. For example, based on the recent version, the name is XQuartz-2.7.7.dmg.
- 4. Click Download.

- 5. Double-click the **XQuartz.pkg** file. Follow the instructions.
- 6. Click **OK** when the message about logging out displays and click **Close** when the installation is successful.

#### **ENABLING THE SECURITY SETTINGS**

- 1. From the Apple main menu, select System Preferences.
- 2. Click Security and Privacy.
- 3. Click **General**. At the bottom of the **Security & Privacy** window, click the lock icon to make changes. Enter the administrator password as required.
- 4. If you have already tried to launch the Thermo-Calc installer, you can click **Open Anyway** to proceed. Otherwise, under **Allow apps downloaded from:** click **Anywhere** and then **Allow from Anywhere**.
- 5. Continue with the applicable installation type (standalone or network).



Remember to reset your security after installing Thermo-Calc.

# **Linux Installation Notes**

The following are details you need to know regarding your Linux installation as well as how to open Thermo-Calc.

### **Installing Libgfortran**

To install and run Thermo-Calc and its components, a runtime library (libgfortran 4.4 or higher) for GNU Fortran applications is required.

#### HOW TO INSTALL THE RUNTIME LIBRARY

Linux distribution	Notes	
openSUSE 42.2		On a standard installation, you do not need to install this runtime library. You may need GCC if it is not already installed on your computer.

Linux distribution	Notes
Ubuntu 16.04 (Xenial)	<pre>Open a terminal and enter the following: &gt; sudo apt-get -y update &gt; sudo apt-get -y install gfortran</pre>
CentOS 7.0	<pre>Open a terminal and enter the following: &gt; sudo yum -y update &gt; sudo yum -y install libgfortran</pre>

# openSUSE Administration (sudo) Users

Note the following if you are a sudo user (administrator) in openSUSE installing Thermo-Calc using this command:

sudo ./linux-installation-x64-6.0.0.17347.run

When this command is used, you will do a text-based installation instead of having a graphical interface to guide you. The information is the same, just without the user interface.

## **Opening Thermo-Calc in Linux**

To open Thermo-Calc double-click the **Thermo-Calc-2018b** application or from the Thermo-Calc installation directory enter \$ ./Thermo-Calc-2018b.sh in the terminal.

# **Standalone Installations**

You can either do a typical or a custom standalone installation. The license type is called a Single-User Node-Locked License (SUNLL).

### In this section:

Typical Standalone Installations	22
Add New Components to an Existing Installation	24
Custom Standalone Installation	. 24

# **Typical Standalone Installations**

The following describes a typical standalone installation. These steps assume you have completed the section "First Steps of Installation" on page 14.

# Request your license file from Thermo-Calc Software



**Demo (trial) installations**: Skip this step. A license file is automatically installed with the software. You do not need to request it.



**Updates to Thermo-Calc**: Customers who subscribe to the annual maintenance plan can skip this step if you have received your license file in an email from Thermo-Calc Software AB. Save the license file to your computer to use during software installation. The license file name must be the same as the existing file name to ensure there are no errors during the update.

Follow the instructions for your operating system in the section <u>"Request a License File from</u> Thermo-Calc Software" on page 42.

# Download your Thermo-Calc installation software

- *Web downloads*: For web links sent in an email from Thermo-Calc Software, go to the URL and log in with the information provided to download the software.
- DVD installations: Insert the DVD and follow the instructions.

## Run the applicable installation file for your operating system

- 1. Navigate to the folder for your operating system and click the link to download the software package, then go to the folder where you downloaded the software:
- Windows: Double-click the .exe file to launch the Setup program.
- Mac: Unzip the file and follow the instructions.
- *Linux*: To start the Setup Wizard in a Terminal window, enter these commands in the directory where the installation file is saved:

```
> chmod +x linux-installation-x64-2018b.run
```

> ./linux-installation-x64-2018b.run



If you are an openSUSE root user, see <u>"Linux Installation Notes" on page 19</u> for some information about your installation.

- 2. On the Setup-Thermo-Calc window click Next.
- 3. On the License Agreement window window click to accept the license agreement.
- 4. Click Next.

=	=		=	
-	=	-		
-	-	_	1	,
_	_	-1	1	

:

**Demo (trial) installations**: Your installation starts and is done when the **Completing the Thermo-Calc Demo Setup Wizard** opens. Click **Finish** and reboot your computer to start using Thermo-Calc.

#### Select the installation type and upload the license file

1. In the Select installation mode window click Typical Standalone → Install Thermo-Calc (incorporating Diffusion Module (DICTRA) and Precipitation Module (TC-PRISMA).

🛦 Setup	↔	-	
Select Installation Mode			
Typical Standalone <ul> <li>Install Thermo-Calc (incorporating Diffusion and Precipitation modules)</li> </ul>			
Choose a license file C:\Users\amanda\Downloads\\servrc			
Typical Network Client			
O Install Thermo-Calc (incorporating Diffusion and Precipitation modules)			
Enter the name or IP number of the licensing server			
O Custom Choose which features to install			
See the Installation Guide for explanation of the different choices: Installation Guide			

2. In the **Choose a license file** field enter the path to your license file (called **lservrc**) or click the file button related to a location on the computer (for example, where you saved it). Click **Next**.

To prevent errors, leave this field blank if you did not get a license file yet.

Install the software and start using Thermo-Calc

- 1. In the Ready to Install window click Next.
- 2. Click **Finish** and start using Thermo-Calc. For Linux users, log out and log back in to finish the installation and start using Thermo-Calc.



Mac OS installations and if you have previous Thermo-Calc installations: When you launch Thermo-Calc with the new or updated version, the splash screen hides the **Import User Settings** dialog box. In order to proceed you need to click the splash screen and then click **Yes** or **No** in the dialog box.





If you did not get your license file see <u>"Request a License File from Thermo-Calc</u> Software" on page 42.

# Add New Components to an Existing Installation

If you want to add new components to an existing Thermo-Calc installation (for example, you purchased a license for TC-API), run the executable (Windows) or binary (Linux) installation file again as in Step 3 of <u>"Typical Standalone Installations" on page 22</u>. This is not available for Mac.

- 1. Double-click the executable Thermo-Calc installation file. Click Next.
- 2. Click to accept the license agreement. Click Next.
- 3. Select the additional component(s) to install. Click Next.
- 4. Click Next to start the installation, which adds the new component(s).
- Use the **Uninstall** program to *remove* components.
- "Uninstalling Thermo-Calc" on page 61

# **Custom Standalone Installation**

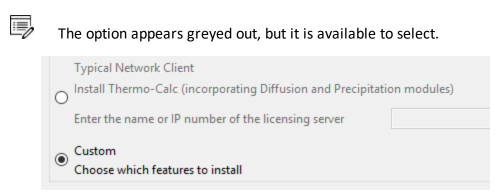
This topic describes how to do a custom installation on a standalone computer for either a Windows or Linux operating system. These steps assume you have completed the section "First Steps of Installation" on page 14.

### Complete Steps 1 to 4 of a typical standalone installation

Follow the instructions for <u>"Typical Standalone Installations" on page 22</u> to get your license file and to download and run the software.

Select the installation type and what components to install

1. On the Select installation mode window, click Custom. Click Next.



2. Set up the Desktop shortcuts.

For *Windows*, in the **Specify Configuration** window, the default is **Current user only**, meaning the installation is only available for the person logged in and installing the software. Select **All users** to allow anyone using this computer to access Thermo-Calc. By default, the **Create Shortcuts on Desktop** check box is selected. Click **Next**.

For *Linux*, and if you are installing as a non-root user, by default, the **Create Shortcuts on Desktop** check box is selected. Click **Forward**.

 In the Select Components window, click to select what you want to install. Click Next.



Additional licenses are required to run TC-Python, TC-API, TQ-Interface, TC-Toolbox for MATLAB (Windows only) and some of the databases. Some elements are included with Precipitation Module (TC-PRISMA) but a separate license is required for full functionality.

4	Setup		↔	-		×
	Select Components					
	Select the components you want to install; clear the components you do not	want to install. Click Nex	t when yo	ou are read	y to cor	ntinue.
	☑ Thermo-Calc	Click on a compone	nt to get a	a detailed d	lescript	ion
	✓ Databases					
	TC-Python					
	TC-API					
	✓ TQ-Interface					
	TC-Toolbox for MATLAB (R)					

#### Set the installation directory path, license file location and shortcuts

 In the Installation Directory window, either accept the default path or enter a path to where you want to install the software (see <u>"Default Directory Locations" on</u> page 14). Click the file button Page to navigate to another directory. Click Next.

🛕 Setup

#### Installation Directory

Please specify the directory where Thermo-Calc will be installed.					
Installation Directory	C:\Program Files\Thermo-Calc\ <mark><version></version></mark>	<b>1</b> 2			

For the Windows TC-Toolbox for MATLAB<sup>®</sup> component, and if you get a **MATLAB not** found message, enter the MATLAB installation directory and click **Next**.

- 2. In the **Choose Setup Type** window, click **Standalone**.
- 3. In the **Choose a license file** field enter the path to your license file (called **lservrc**) or click the file button 🗈 to navigate to a location on the computer. Click **Next**.

To prevent errors, leave this field blank if you do not have a license file yet.

📥 Setup						
Choose Setup Type						
Select one of the following setup types. See installation guide for detailed description of the choices.						
Standalone Choose a license	file	C:\Users\amanda\Downloads\Iservrc	P			
Network Client C Enter name or IP	number c	of the licensing server				

### Install the software and start using Thermo-Calc

- 1. In the Ready to Install window click Next.
- 2. When the installation is complete, click **Finish** and start using Thermo-Calc. For Linux users, log out and log back in to finish the installation and start using Thermo-Calc.

-	_	_

Mac OS installations and if you have previous Thermo-Calc installations: When you launch Thermo-Calc with the new or updated version, the splash screen hides the **Import User Settings** dialog box. In order to proceed you need to click the splash screen and then click **Yes** or **No** in the dialog box.

Import user	settings X
	Do you want to import user settings from
	Yes No



If you did not get your license file see <u>"Request a License File from Thermo-Calc</u> Software" on page 42.

# **Network Client Installations**

This installation type is available for Windows and Linux. It is applicable to the consolidated and distributed network types of installations as well as for the licensing installation on a server network.

- Do a *Typical Network Client* installation if you only want to install the Thermo-Calc program (including Diffusion Module (DICTRA)) and Precipitation Module (TC-PRISMA)) and databases.
- Do a *Custom* installation if you want to select specific components to install or if you want to install the software in a non-default installation directory.

### In this section:

Typical Network Client Installations	29
Set Up the License Manager on Windows	30
Installation Requirements	30
Set Up the License Server on Linux	33
Installation Requirements	33
Thin Client Installations	35
Custom Network Client Installations	37
Unattended Network Client Installations	39

# **Typical Network Client Installations**

The following outlines how you do a typical network client installation.

### Install the licensing software

Follow the instructions for your operating system:

- "Set Up the License Manager on Windows" on the next page, or
- "Set Up the License Server on Linux" on page 33

### Complete Steps 1 to 3 of a typical standalone installation

Follow the instructions for <u>"Typical Standalone Installations" on page 22</u> to download and run the software. It is up to you whether you get the computer information for the license files before or after installing the software.

### Choose the installation type and enter the computer server details

1. Click **Typical Network Client** installation to install Thermo-Calc incorporating Diffusion Module (DICTRA) and Precipitation Module (TC-PRISMA).



The option appears greyed out, but it is available to select.

2. In the field, enter the name or IP number of the computer where the licensing software is installed. Click **Next**.

۲	Typical Network Client Install Thermo-Calc (incorporating Diffusion and Precipitation modules) Enter the name or IP number of the licensing server
0	Custom Choose which features to install
	ne Installation Guide for explanation of the different choices: lation Guide

## Install the software and components

- 1. In the Ready to Install window click Next.
- 2. When the **Completing the Thermo-Calc Setup Wizard** window opens, click **Finish**.
- 3. For *Linux* users, log out and log back in to start using Thermo-Calc.

<sup>4.</sup> Mac OS installations and if you have previous Thermo-Calc installations:

When you launch Thermo-Calc with the new or updated version, the splash screen hides the **Import User Settings** dialog box. In order to proceed you need to click the splash screen and then click **Yes** or **No** in the dialog box.





If you did not get your license file see <u>"Request a License File from</u> <u>Thermo-Calc Software" on page 42</u>.

# Set Up the License Manager on Windows



**License Manager vs License Server**: Both terms refer to the licensing software program installed on a network computer. License Manger is for Windows and License Server is for Linux. See <u>"Installation Terminology" on page 5</u> for more information.

# **Installation Requirements**

This procedure explains how to install the License Manager on a computer that acts as a license server in your network. It is a Sentinel RMS License Manager Windows service and it supplies the client computers in your network with license tokens when it runs the Thermo-Calc software. A network license can only be used on one specific computer that hosts the License Manager. The network license determines which client computers receive license tokens using their IP addresses and IP ranges.

- For *Mac* installations, the server containing the license file must be set up on a Windows or Linux computer.
- For testing purposes, and in order to confirm the installation is working correctly, install the Thermo-Calc software on the same computer as the Sentinel RMS License Manager. Thermo-Calc can always be uninstalled after testing if required.
- Administrator rights are required.

## Installing the License Manager on Windows

Install the Sentinel RMS License Manager software

- 1. Follow the instructions to download your Thermo-Calc installation software.
- 2. Open the directory for Windows and right-click the file **Sentinel RMS License** Manager 8.5.1.msi.
- 3. Select **Save target as** and choose a suitable directory so that you can easily find the file later.
- 4. Right-click the downloaded file and choose Install. Click Next.

### Choose the installation directory

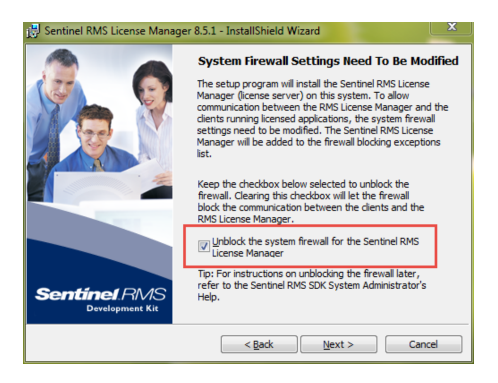
- 1. Click to accept the license agreement. Click Next.
- 2. Enter your User Name and Organization and select to install the License Manager for All users. Click Next.
- 3. In the **Destination Folder** window click **Next** to accept the suggested destination directory or **Change** to select another directory. Click **Next**.

#### Choose the setup type and install the software

- 1. In the Setup Type window, select Complete. Click Next.
- 2. Select the **Unblock the system firewall for the Sentinel RMS License Manager** check box.



Selecting the check box is required to runThermo-Calc.



3. Click Next and then Install.

8

4. Click Finish to exit the InstallShield Wizard.

#### Copy the license files to the destination directory

Copy the network license file to the destination directory. By default, it is installed in the following directory with the name **lservrc**.

```
C:\Program Files (x86)\Common Files\SafeNet Sentinel\Sentinel RMS License Manager\WinNT\
```

The name of the **license file** must be in lower case letters (**lservrc**). This is especially important if both Windows and Linux operating systems are part of your network; Linux commands and operations are case-sensitive.

If the default file path does not work, you can set the environmental variable **LSERVOPTS** to select the license file:

- a. In the Windows **Control Panel**, go to **System**  $\rightarrow$  **Advanced system settings**.
- b. In System Properties select Advanced → Environment Variables → New.
- c. Enter the **Variable name** LSERVOPTS to specify the parameters with which the License Manager is started.

d. To specify the path to the license file in the **Variable value** field enter the following and include the complete license file path and name.

-s <license file path and name>

Reboot or restart the license manager

# Set Up the License Server on Linux



**License Manager vs License Server**: Both terms refer to the licensing software program installed on a network computer. License Manger is for Windows and License Server is for Linux. See <u>"Installation Terminology" on page 5</u> for more information.

#### **Installation Requirements**

躑

There are a variety of things needed to prepare for a Linux License Server installation. The following points are described in this section.

- You need to stop older versions of the License Server before running the new one. To stop the license server you run **Isrvdown**.
- Root access to your computer is required or you need to be on the sudoers list.
- To run the License Server on a Linux 64-bit computer, you need to install 32-bit library support. For example, on CentOS this can be done by running the command:

>yum install glib2.i686

• You need a valid **Iservrc** network license file supplied by Thermo-Calc Software and the actual License Server—a file(program) called **Iserv**.

The name of the **license file** must be in lower case letters (**lservrc**) because Linux commands and operations are case-sensitive.

- The license file need to be read and writable by the license server(license manager)
- If you have a license server; Upgrade your license server if the new one is a higher version than the present one. Clients work with newer license servers(managers) but not necessarily with ones older than client.

#### **OPENSUSE ADMINISTRATION (SUDO) USERS**

Note the following if you are a sudo user (administrator) in openSUSE installing Thermo-Calc using this command:

sudo ./linux-installation-x64-6.0.0.17347.run

When this command is used, you will do a text-based installation instead of having a graphical interface to guide you. The information is the same, just without the user interface.

#### Installing the License Server on Linux

Stop any older versions of the License Server



Skip this step if this is a new installation.

1. To stop an old version of the License Server that is already running, go to the directory where the old file is found, and enter the commands:

```
>chmod +x <lsrvdown>
>./lsrvdown <host name>
```

2. Edit existing scripts that start the older version of the License Server so that they start the newer version instead.

For more information about the **Iservdown utility**, go to <u>http://sentinelrms.safenet-</u> inc.com/RMSDocumentation/SysAdmin/Content/Isrvdown.htm

#### Write a script to start the License Server

The following is an excerpt from the template script for starting the License Server found in the restart\_lserv file.

Replace the paths for LICENSE\_SERVER\_DIR, LICENSE\_FILE\_DIR and LOG\_DIR to match your system. Save it as a script:

```
# set the directory where the executables are located
LICENSE_SERVER_DIR=/usr/local/Thermo-Calc/2018b/Licensing-files
# set the directory of the license file lservrc
LICENSE_FILE_DIR=/usr/local/Thermo-Calc/2018b/Licensing-files
# set the directory where you wish to save the log-files
LOG_DIR=/tmp
# stop the running server
# (omit this line if calling this script at startup)
$LICENSE SERVER DIR/lsrvdown $HOSTNAME
```

```
# start the license server
$LICENSE_SERVER_DIR/lserv -s $LICENSE_FILE_DIR/lservrc -l $LOG_DIR/lserv_use.log -f
$LOG_DIR/lserv_err.log
```

Confirm the HOSTNAME variable and open the UDP port

1. Before starting the License Server make sure that the value of the \$HOSTNAME variable is correct. This value can be shown by typing the > nslookup \$HOSTNAME command in the terminal window:

The value should be the same IP address that is shown when you use the ifconfig command.

Open the UDP port 5093 in your firewall. In case you have installed SELinux you
might also need to disable it by setting SELINUX=permissive in the
/etc/selinux/config file.

#### Start the License Server

1. To start the License Server, run the script created as root. Enter these commands in the console:

```
> chmod +x <name_of_script_file>
```

```
> ./<name_of_script_file>
```

 If you want the License Server to start when you boot the system, put a copy of the start script under /etc/rc.d/init.d and start the script from the file /etc/rc.d/rc.local.

# **Thin Client Installations**

This topic describes how you install a thin Thermo-Calc client to run Thermo-Calc from a file server. It is applicable to consolidated and distributed network installations for either a Windows or Linux operating system.

#### **INSTALLATION REQUIREMENTS**

- You need access to the directory where Thermo-Calc is installed on the file server.
- You need the IP number or name of the computer where the license file is installed.
- For Linux you may have to disable your firewall. If SELinux is installed, try disabling it by setting SELINUX=permissive in /etc/selinux/config.

#### **INSTALLING A THIN CLIENT ON WINDOWS**

- 1. Map the file server share as a network drive so that it gets an assigned drive letter (e.g. E:).
- 2. Open the **\Thermo-Calc\**<*version*>\ directory on the file server, where *version* is the current version of the software.
- 3. Open the directory Thin-client. Double-click the client-install-windows.exe file.
- 4. Click to accept the terms of the license agreement. Click Next.
- Enter the name or IP number of where the license file and the License Manager software is installed. Click Next.
- 6. In the Specify Configuration window, the default is Current user only, meaning the installation is only available for the person logged in and installing the software. Select All users to allow anyone using this computer to access Thermo-Calc. By default the Create Shortcuts on Desktop check box is selected. Click Next.
- 7. In the Ready to Install window, click Next.
- 8. Click **Finish**. To start Thermo-Calc find the program on the server and start it from there.

#### INSTALLING A THIN CLIENT ON LINUX

1. Add the following to the smb.conf file, where *<user>* is your user name:

```
[public]
path=/home/<user>/Thermo_Calc/2018b
public=yes
writable=yes
comment=smb share
guest OK=yes
force user=<user>
```

 From the client, mount the directory on the server computer that contains the Thermo-Calc server installation (by default, this directory is called /Thermo-Calc/2018b). How the directory is mounted depends on the Linux distribution being used as well as how the Thermo-Calc directory on the server computer is shared. Use the Linux command mount. For example:

```
> mkdir public
> sudo mount -t cifs //ip_number_server_computer/public ./public -o
user=<user>
```

- 3. Open the mounted directory on your computer, and run client-install-linuxx64.run or client-install-linux-x86.run in the /Thin-client directory.
- 4. If you do this from the terminal window, use the following to start the installation (assuming that the file server is a 64-bit computer):

> ./client-install-linux-x64.run

- 5. Click Forward and click to select the license agreement. Click Forward.
- 6. Enter the name or IP number of the License Server. Click Forward.
- 7. If you are installing as a non-root user, by default, the **Create Shortcuts on Desktop** check box is selected. Click **Forward**.
- 8. In the Ready to Install window click Forward.
- 9. Click Finish. Log out and log back in to start using Thermo-Calc.

# **Custom Network Client Installations**

This section describes how to do a custom network client installation for Windows and Linux operating systems. The requirements are the same as for a typical client installation.

#### Install the licensing software

Follow the instructions for your operating system:

- "Set Up the License Manager on Windows" on page 30, or
- "Set Up the License Server on Linux" on page 33

#### Complete Steps 1 to 4 of a typical standalone installation

Follow the instructions for <u>"Typical Standalone Installations" on page 22</u> to download and run the software. It is up to you whether you get the computer information for the license files before or after installing the software.

### Choose the installation type and what to install

1. In the Select installation mode window click Custom. Click Next.



The option appears greyed out, but it is available to select.

С	Typical Network Client Install Thermo-Calc (incorporating Diffusion and Precipitation modules)
	Enter the name or IP number of the licensing server
۲	Custom Choose which features to install

2. In the Select Components window, choose what you want to install. Click Next.

#### Set the installation directory path to the license file

1. In the **Installation Directory** window, enter the path to an installation directory (see "Default Directory Locations" on page 14). Click **Next**.

🛕 Setup	↔ – □ ×
Select Components	
Select the components you want to install; clear the components you do	not want to install. Click Next when you are ready to continue. Click on a component to get a detailed description
<ul> <li>✓ Databases</li> <li>✓ TC-Python</li> </ul>	
✓ TC-API	
☐ TQ-Interface	
TC-Toolbox for MATLAB (R)	

For the Windows TC-Toolbox for MATLAB component and if you get a **MATLAB not found** message, enter the MATLAB installation directory and click **Next**.

2. In the **Choose Setup Type** window, click **Network Client** and enter the name or IP number of the computer where the licensing software is installed. Click **Next**.

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## Specify the configuration and create shortcuts

For *Windows*, in the **Specify Configuration** window, the default is **Current user only**, meaning the installation is only available for the person logged in and installing the software. Select **All users** to allow anyone using this computer to access Thermo-Calc. By default, the **Create Shortcuts on Desktop** check box is selected. Click **Next**.

For *Linux*, and if you are installing as a non-root user, by default, the **Create Shortcuts on Desktop** check box is selected. Click **Forward**.

### Install the software and start using Thermo-Calc

- 1. In the Ready to Install window click Next.
- 2. When the **Completing the Thermo-Calc Setup Wizard** window opens, click **Finish**.
- 3. Log out and log back in and start using Thermo-Calc.



Mac OS installations and if you have previous Thermo-Calc installations: When you launch Thermo-Calc with the new or updated version, the splash screen hides the **Import User Settings** dialog box. In order to proceed you need to click the splash screen and then click **Yes** or **No** in the dialog box.



If you have not got the license file, go to <u>"Request a License File from Thermo-Calc Software" on page 42</u>.

# **Unattended Network Client Installations**

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**Terminology note**: The use of the term *license server* (lower case letters) is applicable to the option files as described in this section. For the purposes of the step-by-step instructions it is not used to refer to the type of network installation that runs the licensing software.

Sometimes it is convenient to install Thermo-Calc network clients without providing user input after the installation is initiated (an unattended network client installation). This is available for Windows and Linux operating systems.



Make sure you know the name or IP number of the computer where the licensing software and the license file are to be installed.

#### Create an option file in a text editor to enter the options

The following is an example of an option file for a custom network client installation, where <*name/IP number>* is the name or IP number of the computer where the license file is installed and <*MATLAB directory>* is the path to the MATLAB directory (when applicable):

```
installation_mode=custom
setup_type=client
enable-components=thermo,databases,tq,tcapi,toolbox
config=all
icons=1
license_server=<name/IP number>
matlab_folder=<MATLAB directory>
```

Below is a shorter example, specifying that a typical Thermo-Calc network client should be installed:

```
installation_mode=typical_client
license_server=<name/IP number>
```

### Go to the directory where the installation file is stored



Thermo-Calc can only be installed in unattended mode from the command line prompt.

Start the installation in unattended mode using the option file created with this command:

- For Windows: <installationfile>.exe mode unattended --optionfile <path\optionfile>
- For Linux: <installationfile>.run --mode unattended --optionfile <path/optionfile>

To see other options when you start the Thermo-Calc installation program from the command line prompt, run this command:

- For Windows: <installationfile>.exe --help
- For Linux: <installationfile>.run --help

# **Request License Files**

# In this section:

Request a License File from Thermo-Calc Software	42
Getting License Information Using Thermo-Calc	42
Requesting the License File for Windows	43
Requesting the License File for Mac OS	45
Requesting the License File for Linux	46

# **Request a License File from Thermo-Calc Software**

The following explains how to retrieve the necessary diagnostic and network adaptor information about your computer and then send an email to Thermo-Calc Software to get your license file(s) containing the license key(s).

To set up the correct licensing (excluding demo installations where a license file is not required):

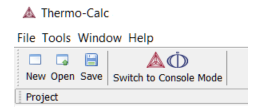
- 1. Make a note of your company or organisation's *site number*. You can find the site number on an invoice or any correspondence from Thermo-Calc Software.
- 2. Retrieve your computer diagnostic and network adapter information.
- 3. Email Thermo-Calc Software with this information.
- 4. A Thermo-Calc Software representative emails you the license file. Save the license file **lservrc** (there is no file suffix) to a location where you can easily access it later.
- http://www.thermocalc.com/support/request-support/

# **Getting License Information Using Thermo-Calc**

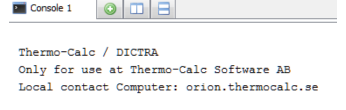
If you have already installed Thermo-Calc, you can view the license information in the GUI. Select Help  $\rightarrow$  Show License Info.

In Console Mode you can run the SYS module command DISPLAY\_LICENSE\_INFO.

- 1. Open Thermo-Calc.
- 2. Click Switch to Console Mode.



3. At the sys: prompt, enter DISPLAY\_LICENSE\_INFO (all caps).



SYS:DISPLAY\_LICENSE\_INFO

4. You can output the information to a FILE or to the SCREEN.

```
SYS:DISPLAY_LICENSE_INFO
OUTPUT TO SCREEN OR FILE /SCREEN/:SCREEN
```

5. For both options, the information is displayed on screen. Copy the section containing the Ethernet and IP address. You can save it to a text file or paste it directly into an email. If you choose FILE as the output, then you can attach the file to an email. For Windows and Linux you can also run the diagnostic program located on the website.

# **Requesting the License File for Windows**

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**Demo (trial) installations**: Skip this section. A license file is automatically installed with the software. You do not need to request it.

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- **Upgrades to a new version of Thermo-Calc**: Skip this section if you are upgrading to a new version of Thermo-Calc and (and you have a maintenance plan). Your license is sent to you in an email from Thermo-Calc Software AB. Save it to your computer to use during software installation.
- To access the computer information you need to provide to get a license, start Thermo-Calc and select Help → Show License Info.

🛕 License Information		×
		^
License diagnostics progra	am	
build: created: Wed (	Oct 11 09:15:37 2018	
Log generated at: 09-11-20	018 09:15:39	
Safenet environment varial	bles	
LSHOST: NO-NET		
LSERVRC: C:\Program Fil	les\Thermo-Calc-academic\&version>\LSERVRC	
Local (client) info		
hostname:	amanda	
license version:	8.5.1.0017	
ethernet:	F0-D5-BF-4E-CA-D7	~
		ОК

- Copy the information from the License Information window. A lot of information may be shown below the initial screen, so make sure you scroll down to the end when you select and copy the window's contents. (Alternatively, you can copy the same information from the console by using the SYS module command DISPLAY\_ LICENSE\_INFO.)
- 3. Send an email to <a href="mailto:support@thermocalc.com">support@thermocalc.com</a>. Make sure to include the following in the subject line:

**License diagnostic information from <Name>** where Name is replaced with your **name**, the **organisation name** and the **Site Number**. *You can find the five-digit site number on an invoice or any correspondence from Thermo-Calc Software*.

te 12345 al Science et 190						
support@thermocalc.com						
License diagnostic information from Jane Smith, XYZ Company, Site 12345						
ments 📋 tclog.log (7.3 KB) 🗵						

- 4. A representative from Thermo-Calc Software replies to the email and attaches your license file(s).
- 5. Save the license file(s) to your computer or network.

# **Requesting the License File for Mac OS**

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**Demo (trial) installations**: Skip this section. A license file is automatically installed with the software. You do not need to request it.

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**Upgrades to a new version of Thermo-Calc**: Skip this section if you are upgrading to a new version of Thermo-Calc and (and you have a maintenance plan). Your license is sent to you in an email from Thermo-Calc Software AB. Save it to your computer to use during software installation.

- 1. From the Apple main menu, select System Preferences.
- 2. Click Network.
- 3. In the left column select **Ethernet** or **Built-in Ethernet** (do not select a WiFi connection as a local static MAC address is required).
- 4. Click Advanced → Hardware.

The **Network** window shows you the **MAC Address**. For example, the MAC address (the host ID) might be **3c:07:54:28:5f:72**.

000 <>	Network					Q Search	
<> Ethernet							
	TCP/IP			802.1X	Proxies	Hardware	
	MAC Address: 3c:07:54:28:5f:72						
	Configure:			Automatically			
		Sp	eed:			٢	
		Dup	olex:	half-duplex		٢	
		Ν	ITU:	Standard (1	500)	٥	
			V	AVB/EAV N	Node		

5. Send an email to <a href="mailto:support@thermocalc.com">support@thermocalc.com</a> and copy and paste the MAC address into the body of the email. Make sure to include the following in the subject line:

**License diagnostic information from <Name>** where Name is replaced with your **name**, the **organisation name** and the **Site Number**. *You can find the five-digit site number on an invoice or any correspondence from Thermo-Calc Software*.

Designated S XYZ Company Dpt. of Mater 900 Main Stro Stockholm 11 SWEDEN	rial Science Set
To:	support@thermocalc.com
Cc:	
License	diagnostic information from Jane Smith, XYZ Company, Site 12345
Add attach	ments tclog.log (7.3 KB) 🗵

- 6. A representative from Thermo-Calc Software replies to the email address provided and attaches your license file or files.
- 7. Save your license file to your computer or network.

# **Requesting the License File for Linux**

- 1. Go to Thermo-Calc's website.
- 2. From the Support menu select Diagnostic tools.
- 3. Under TCLOG→ Linux computers click the link tclog for Linux computers.
- 4. For the tclog.tar file, either Save File or Open with the default program.
- 5. Go to the **Downloads** directory. Extract the files using the mouse (for example, in Ubuntu right-click to extract the files) or use this command in the Terminal window:

> tar -xf tclog.tar

- 6. This creates a subdirectory called *tclog* in the Downloads directory.
- 7. In a Terminal window run the following command replacing *<tclog\_path>* with the path to tclog:

> sh /<tclog path>/script/getreg.sh

- 8. Double-click to open the **tclog** folder. Then either copy the output from the Terminal window or right-click and save the **tclog** file. Name it **tclog.log**.
- 9. Send an email to <a href="mailto:support@thermocalc.com">support@thermocalc.com</a> and attach the <a href="mailto:tclog.log">tclog.log</a> file. Make sure to include the following in the subject line:

**License diagnostic information from <Name>** where Name is replaced with your **name**, the **organisation name** and the **Site Number**. *You can find the five-digit site number on an invoice or any correspondence from Thermo-Calc Software*.

Designated XYZ Compar Dpt. of Mate 900 Main St Stockholm 1 SWEDEN	ny erial Science reet
To:	support@thermocalc.com
Cc:	
License	diagnostic information from Jane Smith, XYZ Company, Site 12345
Add attact	hments 🗋 tclog.log (7.3 KB) 🛞

- 10. A representative from Thermo-Calc Software replies to the email and attaches your license file(s).
- 11. Save the license file(s) to your computer or network.

# Troubleshooting

## In this section:

License Errors	. 49
Determining the Current License File Directory Path	51
Specifying a New Server Name or IP Number	. 54
MATLAB <sup>®</sup> Installation Directory	55
Firewalls	55
Client/Server Mismatch	56
Mac OS Installations: Running the License Initialization Script	. 57
Linux Installations: Running the License Initialization Script	58

# **License Errors**

lssue	Platform	Installation type	Action or link to more information
How do I add or update a standalone (SUNLL) license on a Windows computer where Thermo-Calc is already installed?	Windows	Standalone	"Determining the Current License File Directory Path" on page 51
How do I add or update a standalone (SUNLL) license on a Linux computer where Thermo-Calc is already installed?	Linux	Standalone	"Determining the Current License File Directory Path" on page 51
How do I add or update a network license on Windows?	Windows	Network	"Determining the Current License File Directory Path" on page 51
How do I add or update a network license on Linux?	Linux	Network	"Determining the Current License File Directory Path" on page 51
No license file found. Copy a valid license file to the installation directory after completing the installation.	All	All	You get this message if the wrong path or no license file is found at a path entered during the installation process. See <u>"Default</u> <u>Directory Locations" on page 14</u> and <u>"Determining the Current</u> <u>License File Directory Path" on</u> <u>page 51</u>
MATLAB not found (for the TC-Toolbox for MATLAB component)	Windows	All	<u>"MATLAB® Installation Directory "</u> on page 55

lssue	Platform	Installation type	Action or link to more information
Warning Environment variable "LSHOST" is not set. Software will not function properly.	All	Network	"Specifying a New Server Name or IP Number" on page 54
Problem running post- install step. Installation may not complete correctly. Unable to run post- installation script, run the license initialization script as root after the installation.	Linux and Mac	Standalone	<u>"Mac OS Installations: Running the</u> <u>License Initialization Script " on</u> <u>page 57 or "Linux Installations:</u> <u>Running the License Initialization</u> <u>Script" on page 58</u>
Firewall warnings related to license management software (License Manager and License Server)	Windows and Linux	Network	"Firewalls" on page 55
Client / Server naming mismatches	Windows and Linux	Network	"Client/Server Mismatch" on page 56
InstallScript engine missing	Windows	Network	You are not notified if the InstallScript scripting engine is not installed. Install the ISScript.msi file and then run the downloaded file from Thermo-Calc again as described in <u>"Set Up the License</u> <u>Manager on Windows" on page 30</u>

# **Determining the Current License File Directory Path**

This topic explains how to find out where your license file is installed, either by default or if a different directory was chosen for previously installed versions. For new installations, where you may not have had a license file when you installed Thermo-Calc (for example, you had a demo version), you can also use these instructions to determine the correct file path.

## From Thermo-Calc

If you have already installed Thermo-Calc, you can view the path to the license file in the GUI. Select Help  $\rightarrow$  Show License Info.

In Console Mode you can run the SYS module command DISPLAY\_LICENSE\_INFO.

### WINDOWS STANDALONE INSTALLATIONS

How do I add or update a standalone (SUNLL) license on a Windows computer where Thermo-Calc is already installed?

1. Unpack the .zip file that contains your standalone license and save the LSERVRC file on your computer. Find out where to place the new license file by opening a Command Prompt Window (cmd.exe) and type in the command line:

>set LSERVRC

2. Press Enter. The output from the command is the path to the license file (including the name of the license file). For example: if the output is

LSERVRC=C:\Program Files\Thermo-Calc\2018b\lservrc

then the new license file should be placed in C:\Program Files\Thermo-Calc\2018b\

### LINUX STANDALONE INSTALLATIONS

How do I add or update a standalone (SUNLL) license on a Linux computer where Thermo-Calc is already installed?

Find out where to place the new license file by opening a Terminal window and type in the command line:

```
echo $LSERVRC
```

### WINDOWS NETWORK INSTALLATIONS

How do I add or update a network license on Windows?

- 1. You need administrator rights to do the following.
- On *Windows 7*, click the Start button and in the search field type Services. Doubleclick the Services result.

Programs (3)	
les Component Services	
Services	
services.exe	

- On *Windows 8*, move the mouse over the right edge of the screen to access Search, click Settings and then enter and select View Local Services.
- 2. In the list of installed services, right-click **Sentinel RMS License Manager** and select **Properties** from the pop-up menu.
- 3. On the **General** tab under **Path to executable**, click the line containing the path to see where the file **Iservnt.exe** is located. The license file (Iservrc) is in the same directory.

Sentinel RMS Licen	se Manager Properties (Local Computer)
General Log On	Recovery Dependencies
Service name:	Sentinel RMS License Manager
Display name:	Sentinel RMS License Manager
Description:	Sentinel RMS License Manager
Path to executab	
	Net Sentinel/Sentinel RMS License Manager/WinNT/Iservnt.exe"
Startup type:	Automatic
Help me configu	re service startup options.
Service status:	Started
Start	Stop Pause Resume
You can specify here.	the start parameters that apply when you start the service from
Start parameters	:
	OK Cancel Apply

The default path for the license file is

```
C:\Program Files (x86)\Common Files\SafeNet Sentinel\Sentinel RMS
License Manager\WinNT\LSERVRC
```

- 4. Move your copy of the license file (lservrc) to this directory.
- 5. Restart the Sentinel RMS License Manager by:
- Restarting the computer, or
- Right-clicking Sentinel RMS License Manager and selecting Restart from the list of installed services.

### LINUX NETWORK INSTALLATIONS

How do I add or update a network license on Linux?

Replace your current license file with the new license file. The license file is located in the same directory as the license server.

# Specifying a New Server Name or IP Number

The Thermo-Calc Setup wizard does not warn you if the name or IP number entered to the server (with the license management software) is incorrect.

If the name or IP number is not set at all, or if it is set to an empty string, then when you start Thermo-Calc the following message displays: *Warning Environment variable "LSHOST"* is not set. Software will not function properly.

### WINDOWS

This section explains how to specify or change the name or IP number of the server where the License Manager is installed and for a consolidated network client installation.

To install a license for an existing Thermo-Calc program, follow these steps:

- 1. Close all running Thermo-Calc programs.
- 2. Open a command line window.
- In *Windows 8,* use Search (mouse-over the right edge of your screen) to find **cmd** and then double-click the Command line icon.
- In Windows 7, from the Start menu click Run and type cmd in the field.
- 3. Specify the name or IP number of the licensing software server. Type:

set LSHOST=<name/IP number>

To display the current name or IP number, type:

set LSHOST

### MAC

Permanently set the environmental variable LSHOST to specify a new licensing server name or IP number. Refer to the file /Applications/Thermo-Calc-2018b.app/Contents/plist.info.

In this file, you can manually set the TC18B\_HOME variable, which specifies the home directory of the Thermo-Calc program. See the plist.info file for an example.

### LINUX

You can set or change the name or IP number of the license server at a later time by setting

the environmental variable "LSHOST" to this name or number.

# **MATLAB®** Installation Directory

For Windows installations of the TC-Toolbox for MATLAB component, you may get an error message or window asking for the MATLAB installation directory path. If it is during the installation Setup process, enter the path in the field.

### **DETERMINING THE LICENSING SOFTWARE VERSION**

Sometimes you may need to know the version of License Manager (Windows) or License Server (Linux) software being used. This can be useful while troubleshooting network installations.

• On *Windows* open a command prompt and go to the directory where <code>lservnt.exe</code> is found. For example:

```
>C:\Program Files (x86)\Common Files\SafeNet Sentinel\Sentinel RMS
License Manager\WinNT
```

Type the command >lservnt.exe -help and the version is displayed.

• On *Linux* open a Terminal window and go to the directory where the <code>lserv script</code> is saved. For example:

>cd /usr/local/Thermo-Calc/<version>/Licensing-files

Type the command >./lserv -help and the version is displayed.

# **Firewalls**

## WINDOWS

For a Windows License Manager installation you may have to change the system firewall settings to allow the License Manager to communicate with the computers running the client applications. Go to the Windows Firewall settings window to see if there are any problems. If prompted, select Unblock the system firewall for the Sentinel RMS License Manager and click Next.

### LINUX

For a Linux License Server installation you need to open the UDP port 5093 in your firewall. In case you have installed SELinux you might also need to disable it by setting

SELINUX=permissive in the /etc/selinux/config file.

# **Client/Server Mismatch**

For some network installations there can be a naming mismatch that causes errors. The server (where the license management software is installed) gets information from its DNS servers that an IP address (aaa.bbb.ccc) belongs to a computer (Computer A). But the client computer identifies itself as having a different IP address (aaa.bbb.ccc.eee) for the same computer (Computer A).

Even if the IP address of the client computer belongs to a subnet range listed in the license file, there is a problem with accessing the license file because the computer name does not match the information.

You may need your organisation's network administrator to assist with this issue.

## How to Check

## *Compare computer names on the client computer and server*

- 1. On the client computer, start a command prompt **cmd.exe** and type <code>ipconfig /all</code>. Press <code>Enter</code>. A variety of data is displayed on screen. Write down the following information for later comparison:
- Host name
- Dns Suffix
- Current IPv4 Address
- 2. In the command window, type nslookup <client\_computer\_name>, where client\_ computer\_name is the actual name of the client. Press Enter.

This shows what information is saved on the DNS Servers used by the client computer. This information should match what is shown in the output from the first step.

3. On the server that holds the license management software, open a command prompt and run the command line: nslookup client\_computer\_name. The output lists the information for this server.

Does it match the information given by the command ipconfig /all on the client computer? If not try the following:

## Try these options to correct the issue

- 1. If the client computer has a dynamic (DHCP) IP address, either change it to a static one or make a reservation on the DHCP Server so that the computer always get the same IP address.
- 2. Update the records on the DNS Servers used by both the client computer and the server computer.

You can also hard code computer names and IP addresses in the local "address book" of the client and server:

- 1. On the client computer, make a backup copy of the file C:\Windows\System32\drivers\etc\hosts.
- 2. Open the original in a simple text editor.
- 3. Add a new line which contains the IP and name of the server, for example: 192.168.29.27 license\_server.thermocalc.se.
- 4. Save the changes.
- 5. On the server, make a backup copy of the file /etc/hosts.
- 6. Open the original in a simple text editor.
- 7. Add a new line which contains the IP and name of the client, for example: 192.168.29.47 Computer\_A.thermocalc.se.
- 8. Save the changes.
- 9. In the host file, there is an example of how the operating system (Windows or Linux) wants the line in the hosts file to look like.

# Mac OS Installations: Running the License Initialization Script

As part of the Standalone installation of Thermo-Calc, the Setup runs a license initialization script. If this part of the installation process did not complete properly for a standalone installation, you must manually run the license initialization script. You need the administrator username and password for these steps.

## HOW TO RUN THE SCRIPT

For this script to successfully initialize the license, the license file must be called lservrc and be located in the folder /Applications/Thermo-Calc-2018b.app/Contents/Resources or /Applications/Thermo-Calc-2018b-Demo.app/Contents/Resources for demo installations.

## "Default Directory Locations" on page 14

- 1. Open a terminal window. The **Terminal** application is in the */Applications/Utilities* folder.
- 2. At the prompt, enter this command:

```
$ sudo /Applications/Thermo-Calc-
2018b.app/Contents/Resources/license-initialization-MacOS-x86_
64.sh
```

For demo installations, the path should be /Applications/Thermo-Calc-2018b-Demo.app/Contents/Resources/ license-initialization-MacOS-x86\_64.sh.

If you are not already logged in as an administrator, you probably cannot run the sudo command. If this is the case, then you can use the su command in the terminal window to login as an administrator. Contact your local system administrator for help.

- 3. Enter the administrator password and press Enter.
- 4. When you restart your computer and run the license initialization script, you are ready to start using Thermo-Calc.



If you did not install a license file during this procedure, you can still start Thermo-Calc although you will get the **No license file found** message.

# Linux Installations: Running the License Initialization Script

As part of a Standalone installation of Thermo-Calc, the Setup runs a license initialization script. If you get a warning message during the standalone installation Unable to run post-installation script, run the license initialization script as root after the installation, then you need to manually run the script as root.

### **OPENSUSE ADMINISTRATION (SUDO) USERS**

Note the following if you are a sudo user (administrator) in openSUSE installing Thermo-Calc using this command:

```
sudo ./linux-installation-x64-6.0.0.17347.run
```

When this command is used, you will do a text-based installation instead of having a graphical interface to guide you. The information is the same, just without the user interface.

### HOW TO RUN THE SCRIPT

For this script to successfully initialize the license, the license file must be called **Iservrc** and be located in the directory where you have installed Thermo-Calc.

Default Directory Locations" on page 14

1. In the console, go to the Thermo-Calc installation directory.

The default installation directory is /home/<user>/Thermo-Calc/2018b if you are installing as a non-root user, where <user> is your user name. If you are installing as root, then this directory is /usr/local/Thermo-Calc/2018b.

2. Enter this command:

\$ sudo sh license-initialization-linux.sh

If you are not able to execute the sudo command because you are not on the sudoers list, you must log in as a root user with the command su, and then use sh to run the license initialization script. For example, on a 64-bit computer, enter the following:

```
$ sh license-initialization-linux.sh
```

3. Log out of Thermo-Calc and then log back in to save these changes.

# **Uninstalling Thermo-Calc**

## In this section:

Uninstalling Thermo-Calc	61
Uninstalling Standalone Versions of TC-PRISMA	62

# **Uninstalling Thermo-Calc**

## WINDOWS 10

- 1. From the **Control Panel** Desktop app, go to **Programs** → **Programs** and **Features** → **Uninstall or change a program**.
- 2. In the list of programs, locate the version of Thermo-Calc you want to uninstall. Click to highlight that row.
- 3. Click Uninstall/Change.
- 4. In the **Setup** window, **Uninstallation Mode** dialog box, choose to uninstall the **Entire application** or **Individual components**. Follow the instructions to finish.

## WINDOWS 7

- 1. From the **Start** menu, go to **All Programs** or **All Apps** → **Thermo-Calc**.
- 2. Click Uninstall-Thermo-Calc<version>.
- 3. In the **Uninstallation Mode** window, choose to uninstall the **Entire application** or **Individual components**. Follow the instructions to finish.

## MAC

Deleting the program from the Applications folder does not remove the program from your operating system. Run the **Uninstall-Thermo-Calc**<*version>* program located in the Applications folder. This is also applicable to the Demo version of the software.

- 1. Go to the **Applications** folder and double click the file that starts with **Uninstall-Thermo-Calc**<*version*>. Enter the administrator's user name and password.
- 2. Click Yes in answer to the question Do you want to uninstall Thermo-Calc and all of its modules? Click No to quit.
- 3. The Uninstalling Thermo-Calc window shows the progress. Click OK when prompted.

### LINUX

1. To start the uninstallation procedure, in the Thermo-Calc installation directory use the command

./uninstall-Thermo-Calc-2018b

For *Thin Client* installations, use the command /uninstall-Thermo-Calc-Client-2018b

2. To remove the whole application, including all its components, select **Entire application** and click **Forward**, or to remove specific components, select **Individual components** and click **Forward**.

If you intend to reinstall Thermo-Calc 2018b it is important to remove the environmental variable TC18B\_HOME by logging out and back into the Linux user account.

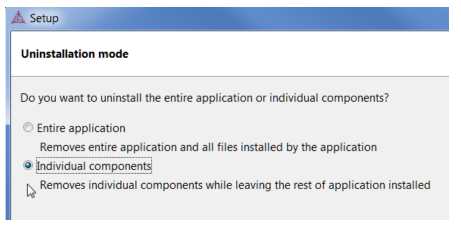
# **Uninstalling Standalone Versions of TC-PRISMA**

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As of Thermo-Calc version 2016a, TC-PRISMA is no longer a standalone program. It is integrated into the Thermo-Calc Graphical Mode and considered an add-on module called the Precipitation Module. It is also available for all platforms (Windows, Mac and Linux). If you have older versions of the TC-PRISMA software that you want to uninstall, follow these instructions.

- 1. Go to the Thermo-Calc uninstaller. For example, in Windows 7, go to Start → All Programs → Thermo-Calc or in Windows 10 go to Start → All Apps → Thermo-Calc.
- 2. Click uninstall-Thermo-Calc-2015b (or the version of TC-PRISMA you have).
  - Thermo-Calc
     TC-PRISMA-2015b
     Thermo-Calc-2015b
     uninstall-Thermo-Calc-2015b
- 3. In the Setup window, click Individual components. Click Next.



4. Select the **TC-PRISMA** check box.

Select Components To Uninstall	
select components to onnistan	
Select the components you want to uninst when you are ready to continue.	all;
Thermo-Calc	
TC-PRISMA	
Databases	

5. Click **Next** and then **OK** when the software component is uninstalled.

# **Quick Installation Guides**

# In this section:

Windows <sup>®</sup> Standalone Quick Install Guide	65
Request a License File	65
Install the Software	67
Mac OS <sup>®</sup> Standalone Quick Install Guide	69
Request a License File	70
Install the Software	71
TC-Python Quick Install Guide	73
Fixing Potential Issues with the Environment	77
Updating to a Newer Version	77
Mac OS: Setting Environment Variables	78

# Windows<sup>®</sup> Standalone Quick Install Guide

This quick guide helps you do a typical, custom or demo standalone installation. A *standalone* installation is used with the Single-User Node-Locked License (SUNLL), where the software and the license file are together on one computer.

### This guide is applicable to:

- Windows operating systems
- Standalone installation (SUNLL) -typical and custom types
- Demo installations during the installation you only need to do steps 2 and 3.
- Upgrading to a new standalone version of Thermo-Calc (maintenance plan only)



Also see the *Thermo-Calc Installation Guide* for detailed instructions for other operating systems and network installations.

## **Request a License File**

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**Demo (trial) installations**: Skip this section. A license file is automatically installed with the software. You do not need to request it.

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**Upgrades to a new version of Thermo-Calc**: Skip this section if you are upgrading to a new version of Thermo-Calc and (and you have a maintenance plan). Your license is sent to you in an email from Thermo-Calc Software AB. Save it to your computer to use during software installation.

5. To access the computer information you need to provide to get a license, start Thermo-Calc and select **Help** → **Show License Info**.

A License Information		Х
		^
License diagnostics pro	gram	
build: created: We	d Oct 11 09:15:37 2018	
Log generated at: 09-11	-2018 09:15:39	
Safenet environment var	iables	
LSHOST: NO-NET		
LSERVRC: C:\Program	Files\Thermo-Calc-academic\kversion>\LSERVRC	
Local (client) info		
hostname:	amanda	
license version:	8.5.1.0017	
ethernet:	F0-D5-BF-4E-CA-D7	~
		OK

- 6. Copy the information from the License Information window. A lot of information may be shown below the initial screen, so make sure you scroll down to the end when you select and copy the window's contents. (Alternatively, you can copy the same information from the console by using the SYS module command DISPLAY\_LICENSE\_INFO.)
- 7. Send an email to <a href="mailto:support@thermocalc.com">support@thermocalc.com</a>. Make sure to include the following in the subject line:

**License diagnostic information from <Name>** where Name is replaced with your **name**, the **organisation name** and the **Site Number**. *You can find the five-digit site number on an invoice or any correspondence from Thermo-Calc Software*.

Designated Sit XYZ Company Dpt. of Materia 900 Main Stree Stockholm 111 SWEDEN	cience
To:	support@thermocalc.com
Cc:	
License	gnostic information from Jane Smith, XYZ Company, Site 12345
Add attach	ents 📄 tolog.log (7.3 KB) 🗵

- 8. A representative from Thermo-Calc Software replies to the email and attaches your license file(s).
- 9. Save the license file(s) to your computer or network.

## **Install the Software**

#### Download and run your Thermo-Calc installation software

For web links sent in an email from Thermo-Calc Software:

- 1. Go to the URL and log in with the information provided. Then navigate to the Windows folder and click the link to download the software.
- 2. Double-click the .exe file to launch the Setup program.
- 3. On the Setup-Thermo-Calc window click Next.
- 4. On the License Agreement window click to accept the license agreement. Click Next.
  - **Demo (trial) installations**: Your installation starts and is done when the **Completing the Thermo-Calc Demo Setup Wizard** opens. Click **Finish** and start using Thermo-Calc.

#### Select the installation mode – typical or custom

Setup			↔	-	
Select Installation Mode					
Typical Standalone Install Thermo-Calc (inc Choose a license file	orporating Diffusion and Precipitation modu	ıles)			
Typical Network Client					
O Install Thermo-Calc (inc	orporating Diffusion and Precipitation modu	iles)			
Enter the name or IP nu	mber of the licensing server				
Custom Choose which features t	o install				

1. In the **Select installation mode** window click **Typical Standalone** or **Custom** to choose specific features to install.



The **Custom** option appears greyed out, but it is available to select.

2. For a Typical Standalone installation, in the Choose a license file field click the file

button <sup>[1]</sup> to navigate to where you saved the license file emailed to you from Thermo-Calc Software. The license file is called **lservrc**. Click **Next**.

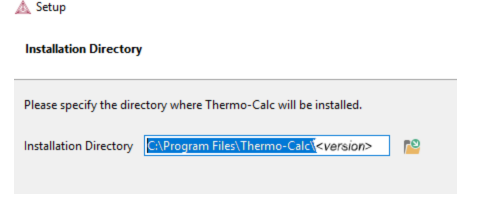
DEMO **Typical Standalone installations**: Your installation starts and is done when the **Completing the Thermo-Calc Demo Setup Wizard** opens. Click **Finish** and reboot your computer to start using Thermo-Calc.

### Customize your installation

1. In the **Select Components** window, click to select what you want to install. Click **Next**.

🛕 Setup	↔ – □ ×
Select Components	
Select the components you want to install; clear the components you do not	want to install. Click Next when you are ready to continue.
Thermo-Calc	Click on a component to get a detailed description
✓ Databases	
✓ TC-Python	
TC-API	
✓ TQ-Interface	
TC-Toolbox for MATLAB (R)	

2. In the **Installation Directory** window, either accept the default, enter a file path or click the file button 🖺 to navigate to another directory.



3. In the Choose Setup Type window it defaults to Standalone. In the Choose a license

**file** field you can accept the default file path or click the file button  $\square$  to navigate to, where you saved the license file emailed to you from Thermo-Calc Software (called **lservrc**). Click **Next**.

A	Setup		
Cł	noose Setup Type		
	ect one of the following se e installation guide for deta	tup types. iled description of the choices.	
۲	Standalone Choose a license file	C:\Users\amanda\Downloads\Iservrd	ø
0	Network Client Enter name or IP number o	f the licensing server	

4. In the Specify Configuration window, the default is Current user only, meaning the installation is only available for the person logged in and installing the software. Select All users to allow anyone using this computer to access Thermo-Calc. By default, the Create Shortcuts on Desktop check box is selected. Click Next.

📩 Setup
Specify Configuration
Current user only
○ All users
☑ Create Shortcuts on Desktop

5. In the Ready to Install window click Next. Click Finish and start using Thermo-Calc.

# Mac OS<sup>®</sup> Standalone Quick Install Guide

This quick guide helps you do a typical or demo standalone installation. A *standalone* installation is used with the Single-User Node-Locked License (SUNLL), where the software and the license file are together on one computer.

#### This guide is applicable to:

- Mac operating systems
- A typical Standalone installation (SUNLL)

- Demo installations during the installation process you only need to do steps 1 and 2.
- Upgrading to a new standalone version of Thermo-Calc (maintenance plan only)



Also see the *Thermo-Calc Installation Guide* for detailed instructions for other operating systems and network installations.

### **Request a License File**

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**Demo (trial) installations**: Skip this section. A license file is automatically installed with the software. You do not need to request it.

 -	_	-	
 -	_		

**Upgrades to a new version of Thermo-Calc**: Skip this section if you are upgrading to a new version of Thermo-Calc and (and you have a maintenance plan). Your license is sent to you in an email from Thermo-Calc Software AB. Save it to your computer to use during software installation.

- 1. From the Apple main menu, select System Preferences.
- 2. Click Network.
- 3. In the left column select **Ethernet** or **Built-in Ethernet** (do not select a WiFi connection as a local static MAC address is required).
- 4. Click Advanced → Hardware.

The **Network** window shows you the **MAC Address**. For example, the MAC address (the host ID) might be **3c:07:54:28:5f:72**.

$\circ \circ \circ < >$				Network			Q Search	
<> Ethernet								
	TCP/IP	DNS	WINS	802.1X	Proxies	Hardware		
	N	1AC Add	ress: 3c	:07:54:28:5	if:72			
		Config	gure: 🛛	Automatical	ly	٥		
		Sp	eed:			٢		
		Du	plex:	nalf-duplex		٥		
		N	NTU:	Standard (1	500)	٥		
			V	AVB/EAV N	Node			

5. Send an email to <a href="mailto:support@thermocalc.com">support@thermocalc.com</a> and copy and paste the MAC address into the body of the email. Make sure to include the following in the subject line: **License diagnostic information from <Name>** where Name is replaced with your **name**, the **organisation name** and the **Site Number**. *You can find the five-digit site number on an invoice or any correspondence from Thermo-Calc Software*.

Designated Sitt XYZ Company Dpt. of Materia 900 Main Stree Stockholm 111 SWEDEN	al Science
To:	support@thermocalc.com
Cc:	
License di	agnostic information from Jane Smith, XYZ Company, Site 12345
Add attachm	nents 📋 tclog.log (7.3 KB) 🗵

- 6. A representative from Thermo-Calc Software replies to the email address provided and attaches your license file or files.
- 7. Save your license file to your computer or network.

## **Install the Software**

### Download and install XQuartz

The XQuartz open-source X Windows system is required to run Thermo-Calc on Mac OS X.

- 1. In a web browser go to <a href="https://www.macupdate.com">https://www.macupdate.com</a> or to download it directly from XQuartz, <a href="http://xquartz.macosforge.org/landing/">http://xquartz.macosforge.org/landing/</a>.
- 2. In the Search Mac Apps field, enter XQuartz.
- 3. Click the **XQuartz** search result. For example, based on the recent version, the name is XQuartz-2.7.7.dmg.
- 4. Click Download.
- 5. Double-click the **XQuartz.pkg** file. Follow the instructions.
- 6. Click **OK** when the message about logging out displays and click **Close** when the installation is successful.

### Download and run your Thermo-Calc installation software

For web links sent in an email from Thermo-Calc Software:

1. Go to the URL and log in with the information provided. Then navigate to the Mac folder and click the link to download the software.

2. Unzip the file.

- 3. On the Setup-Thermo-Calc window click Next.
- 4. On the License Agreement window click to accept the license agreement. Click Next.

**Demo (trial) installations**: Your installation starts and is done when the **Completing the Thermo-Calc Demo Setup Wizard** opens. Click **Finish** and reboot your computer to start using Thermo-Calc.

### Select the installation mode

1. In the Select installation mode window click Typical Standalone.

🛕 Setup	↔	-	
Select Installation Mode			
Typical Standalone           Install Thermo-Calc (incorporating Diffusion and Precipitation modules)           Choose a license file         C:\Users\amanda\Downloads\ksevrc			
Typical Network Client			
O Install Thermo-Calc (incorporating Diffusion and Precipitation modules)			
Enter the name or IP number of the licensing server			
O Custom Choose which features to install			
See the Installation Guide for explanation of the different choices: Installation Guide			

2. In the **Choose a license file** field click the file button 🖸 to navigate to where you saved the license file emailed to you from Thermo-Calc Software. The license file is called **lservrc**. Click **Next**.

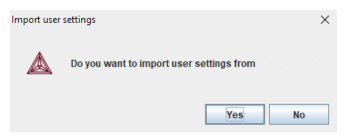


To prevent errors, leave this field blank if you did not get a license file yet.

- 3. The installation starts and is done when the **Completing the Thermo-Calc Setup Wizard** window opens.
- 4. Click Finish and start using Thermo-Calc.



Mac OS installations and if you have previous Thermo-Calc installations: When you launch Thermo-Calc with the new or updated version, the splash screen hides the **Import User Settings** dialog box. In order to proceed you need to click the splash screen and then click **Yes** or **No** in the dialog box.



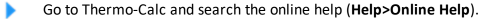
# **TC-Python Quick Install Guide**

This quick guide helps you do a TC-Python API installation.

There is an HTML help file you can open in your browser, plus a PDF of the same content, and several examples available with your Thermo-Calc installation.

Once you have installed TC-Python, go to Thermo-Calc and from the menu select **Help** → **Manuals Folder** and go to:

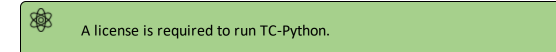
- TC-Python, then double-click index.html (or any .html file).
- Software Development Kits (SDKs) for the PDF.





For an installation video plus more information, go to http://www.thermocalc.com/products-services/software/tc-python/

This quick guide helps you to get a working TC-Python API installation.



## Install a Python Distribution



If you already have a Python distribution installation, version 3.5 or higher, skip this step.

These instructions are based on using the Anaconda platform for the Python distribution. Install version 3.5 or higher to be able to work with TC-Python, although it is recommended that you use the most recent version.

#### Install Anaconda

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- 1. Navigate to the Anaconda website: https://www.anaconda.com/download/.
- 2. Click to choose your OS (operating system) and then click **Download**. Follow the instructions. It is recommended you keep all the defaults.

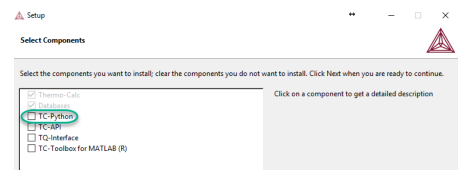
### Install Thermo-Calc and the TC-Python SDK

TC-Python is available starting with Thermo-Calc version 2018a.

- 1. Install Thermo-Calc and choose a Custom installation.
  - See "Custom Standalone Installation" in the Thermo-Calc Installation Guide.

If you have already installed Thermo-Calc, you need to find the installation file (e.g. Windows \*.exe, Mac \*.zip and Linux \*.run) to relaunch the installer and then continue with the next steps.

2. On the Select Components window, click to select the TC-Python check box.



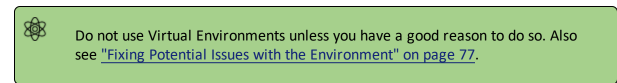
- 3. On the Install TC-Python window, click Next.
- 4. When the installation is complete, the TC-Python folder opens and includes the \*.whl file needed for the next step. There is also an **Examples** folder with Python files you can use in the IDE to understand and work with TC-Python.

The installation location for this API is the same as for other SDKs and based on the OS. For details, see "Default Directory Locations" in the *Thermo-Calc Installation Guide*.

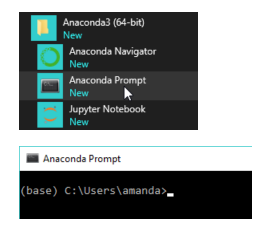
## Install TC-Python

Updating to a Newer Version

On Windows, it is recommended you use the Anaconda distribution prompt , especially if you have other Python installations.



1. Open the command line. For example, in Anaconda on a Windows OS, go to **Start** → **Anaconda** → **Anaconda** Prompt.



2. At the command line, enter the following. Make sure there are no spaces at the end of the string or in the folder name or it will not run:

```
pip install <path to the TC-Python folder>/TC_Python-
<version>-py3-none-any.whl
```

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	_	_	
-	-	-	-

For example, on a Windows OS Standalone custom installation, when you install for all users, the path to the TC-Python folder is C:\Users\Public\Documents\Thermo-Calc\2018b\SDK\TC-Python\



- Details for Mac and Linux installations are described in "Default Directory Locations" in the *Thermo-Calc Installation Guide*. On Linux typically *pip3* is used.
- 3. Press <Enter>. There is a confirmation that TC-Python is installed when the process is completed.

## Install an IDE (Integrated Development Environment)

Any editor can be used to write the Python code, but an IDE is recommended, e.g. PyCharm. These instructions are based on the use of PyCharm.

Use of an IDE will give you access to the IntelliSense, which is of great help when you use the API as it will give you the available methods on the objects you are working with.

- 1. Navigate to the PyCharm website: https://www.jetbrains.com/pycharm/download.
- 2. Click to choose your OS and then click **Download**. You can use the **Community** version of Pycharm.
- 3. Follow the instructions. It is recommended you keep all the defaults.

For Mac installations, you also need to set some environment variables as described below in Mac OS: Setting Environment Variables .

## Open the IDE and Run a TC-Python Example

After you complete all the software installations and set up the pip install path, you are ready to open the IDE to start working with TC-Python.

It is recommended that you open one or more of the included examples to both check that the installation has worked and to start familiarizing yourself with the code.

**Open the TC-Python Project in PyCharm** 



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When you first open the TC-Python project and examples, it can take a few moments for the Pycharm IDE to index before some of the options are available.

1. Open PyCharm and then choose **File**  $\rightarrow$  **Open**. The first time you open the project you will need to navigate to the path of the TC-Python installation as done in Step 4.



For example, on a Windows OS Standalone custom installation, when you install for all users, the path to the TC-Python folder is C:\Users\Public\Documents\Thermo-Calc\2018b\SDK\TC-Python\



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Details for Mac and Linux installations are described in "Default Directory Locations" in the *Thermo-Calc Installation Guide*.

- 2. Click the Examples folder and then click OK.
- 3. From any subfolder:
  - Double-click to open an example file to examine the code.
  - Right-click an example and choose Run .

## **Fixing Potential Issues with the Environment**

In most cases you should run TC-Python within your **global** Python 3 interpreter and not use Virtual Environments unless you have a good reason to do so. If there are problems with the interpreter settings, you can resolve them in the Settings window:

- 1. Go the menu File  $\rightarrow$  Settings.
- 2. Navigate in the tree to <*Project.YourProjectName*> and choose **Project Interpreter**.
- 3. Click on the settings symbol close to the **Project Interpreter** drop-down menu and choose **Add**.
- 4. Now choose System Interpreter and add your existing Python 3 interpreter.
- 5. Select your added interpreter and confirm.
  - If you are not following the recommended approach and create a new project (File → New Project), you need to consider that by default the options to choose the interpreter are hidden within the Create Project window. Therefore, click Project Interpreter: New Virtual Environment and in most cases choose your System Interpreter instead of the default New Virtual Environment.

If you really need to use a Virtual Environment, it is recommended you review the hints provided in the Best Practices section in the full TC-Python HTML help or PDF.

## **Updating to a Newer Version**

When updating to a newer version of Thermo-Calc, you always need to also install the latest version of TC-Python. It is not sufficient to run the installer of Thermo-Calc. The procedure is generally identical to <u>Step 3</u>:

```
pip install <path to the TC-Python folder>/TC_Python-<version>-
py3-none-any.whl
```

In case of problems you may wish to uninstall the previous version of TC-Python in advance:

```
pip uninstall TC-Python
pip install <path to the TC-Python folder>/TC_Python-<version>-
py3-none-any.whl
```

However, that should normally not be required. Note that on Linux usually pip3 is used.

You can check the currently installed version of TC-Python by running:

pip show TC-Python

### **Mac OS: Setting Environment Variables**

In order to use TC-Python on Mac you need to set some environment variables.

```
TC18B_HOME=/Applications/Thermo-Calc-2018b.app/Contents/Resources
```

If you use a license server:

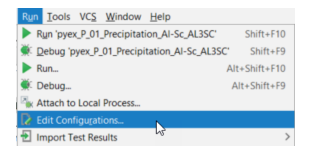
LSHOST=<name-of-the-license-server>

If you have a node-locked license:

```
LSHOST= NO-NET
LSERVRC=/Applications/Thermo-Calc-2018b.app/Contents/Resources/lservrc
```

In PyCharm, you can add environment variables in the configurations.

Select  $Run \rightarrow Edit$  Configurations to open the Run/Debug Configurations window. Enter the environment variable(s) bu clicking the button to the right of the Environment Variables text field.



# **Thermo-Calc User Guide**

Thermo-Calc Version 2018b





# **Introduction to Thermo-Calc**

This guide is an introduction to working with both *Graphical Mode* and *Console Mode* in Thermo-Calc.

## In this section:

The Thermo-Calc Software	
Add-on Modules	3
Typographical Conventions	3
Graphical Mode vs Console Mode	
About the File Types and Formats	8
Graphical Mode	8
Console Mode	8
Examples, Manuals and Materials Folders	12
Help Resources	21
'Online' Help: No Internet Required	21
Context Help (Graphical Mode Only)	21
Console Mode Help	22
The Thermo-Calc Modules and Activities	22
Displaying Thermo-Calc License Information	25
Using This Guide	27

# **The Thermo-Calc Software**

The Thermo-Calc software is a sophisticated database and programming interface package used to perform thermodynamic calculations. It can calculate complex homogeneous and heterogeneous phase equilibria, and then plot the results as property diagrams and phase diagrams.

The Thermo-Calc software (also referred to as the *Thermo-Calc program* or *the software* to distinguish it from the company name) fully supports stoichiometric and non-ideal solution models and databases. These models and databases can be used to make calculations on a large variety of materials such as steels, alloys, slags, salts, ceramics, solders, polymers, subcritical aqueous solutions, supercritical electrolyte solutions, non-ideal gases and hydrothermal fluids or organic substances. The calculations take into account a wide range of temperature, pressure and compositions conditions.

# **Add-on Modules**

The Thermo-Calc program also includes add-on modules:



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The Diffusion Module (DICTRA) and DIC\_PARROT is used to simulate diffusion controlled transformations in multicomponent systems.

The Precipitation Module (TC-PRISMA) is used to simulate diffusion controlled multi-particle precipitation process in multicomponent and multi-phase alloy systems.



Use online help for information about how to use Diffusion Module (DICTRA) and Precipitation Module (TC-PRISMA).

# **Typographical Conventions**

The following typographical conventions are used throughout the documentation, both online and in the PDF formats.

## Техт

Convention	Definition
Forward arrow symbol →	The forward arrow symbol $\rightarrow$ instructs you to select a series of menu items in a specific order. For example, <b>Tools</b> $\rightarrow$ <b>Options</b> is equivalent to: From the <b>Tools</b> menu, select <b>Options</b> .

Convention	Definition
Boldface font	A <b>boldface</b> font indicates that the given word(s) are shown that way on a toolbar button or as a menu selection. For example, if you are told to select a menu item in a particular order, such as <b>Tools</b> ->Options, or to click <b>Save</b> .
Italic font	An <i>italic</i> font in the body of the text indicates the introduction of important terminology. Expect to find an explanation in the same paragraph or elsewhere in the guide.
Command	For features in Thermo-Calc that use Console Mode (i.e. the command line), this font and all capital letters indicates that this is a Console Mode COMMAND. Examples of how you can use a command are written with code font. For example:
	Use DEFINE_ELEMENTS followed by a list of the elements that you want in your system. (To list the elements that are available in your current database, use LIST_DATABASE and choose Elements).
<u>HELP</u>	Text in <u>blue and underline</u> and a page number is a link to another topic in the current or referenced guide. Command names are often also topics. Clicking the link takes you to more detail about a particular command or subject in the PDF or documentation set.
<enter></enter>	Text with <angle brackets=""> indicates a keyboard entry. Usually to press <enter> (or Return) or to use a series of keys such as <ctrl +="" s="">.</ctrl></enter></angle>
code <b>and code</b> <b>bold</b>	A code font shows a programming code or code example. The code bold font highlights the entry.
click vs press	In general, you <i>click</i> with the mouse to perform an action on the screen (e.g. click <b>Save</b> ) and you <i>press</i> keys on a keyboard to enter a set of commands (e.g. press Ctrl+S).

Convention	Definition
	When working in Console Mode, you can use keyboard shortcuts. Sometimes a window opens where you have the option to <b>Save</b> , <b>Cancel</b> or <b>Open</b> a file, for example. In these cases the instructions might say <i>click</i> Save, whereas you would need to <i>press</i> the applicable keys to perform the action.

## ICONS

Convention	Definition
Important	Provides important information. It is recommended that you read the text or follow the link.
Note	The information can be of use to you. It is recommended that you read the text or follow the link to more information.
Also see	<ul> <li>Go to the link or guide to see more general information about the topic being discussed. Usually includes both text and a link.</li> <li>The small blue arrow is used when there is one (sometimes two) cross reference and no text describing the link. Often used in settings pages to link you to accompanying theory or extra detail.</li> </ul>
Тір	This is general information that can be of use but is not required knowledge.

Convention	Definition
Examples	Go to the example collection to learn more.
Console Mode	This note relates specifically to something in Console Mode.
Graphical Mode	This note relates specifically to something in Graphical Mode.
Diffusion Module	This indicates that the information relates to the add- on Diffusion (DICTRA) Module.
Precipitation Module	This indicates that the information relates to the add- on Precipitation Module.

# **Graphical Mode vs Console Mode**

There are two interfaces available in Thermo-Calc: *Graphical Mode* with a graphical user interface (GUI) and *Console Mode*, which uses a command line interface. Thermo-Calc and the Diffusion Module (DICTRA) are available in both modes, and the Precipitation Module (TC-PRISMA) is only available in Graphical Mode.



In Graphical Mode calculations are set up, carried out, and visualized as part of a *project*. The steps in the project are performed with *activities*. There are templates and a Wizard available to guide you through the process of defining the project.

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In Console Mode you work with *modules*, which are managed using commands typed at a prompt. Some modules, called *response-driven modules*, ask you a series of questions that typically take you through the process of defining your system, setting calculation conditions, performing calculations and plotting the results.

## SWITCHING BETWEEN MODES

The first time you open Thermo-Calc, it defaults to Graphical Mode. For any future instance

it defaults to the last mode used.

Along the top of the GUI is the **GRAPHICAL MODE TOOLBAR** where you switch between modes.

F	ile T	ools	Wind	ow Help
				۵
	New	Open	Save	Switch to Console Mode

In Console Mode, except for the **SWITCH TO GRAPHICAL MODE** button, the toolbar is not used.

New Open Save Switch to Graphical Mode



See <u>"Main Menu, Toolbar and Keyboard Shortcuts" on page 35</u> in the *Thermo-Calc User Guide* for information about the **Tools**, **Window** and **Help** menus, which are available in both modes.

## **FILE EXTENSIONS**

The file extensions are different based on the mode you are in (Graphical vs. Console). In Console Mode the extensions are also based on whether you are working with Thermo-Calc or the Diffusion Module (DICTRA).



In Graphical Mode, the Thermo-Calc project files have the extension \*.tcu. This includes the examples for the add-on Diffusion Module (DICTRA) and Precipitation Module (TC-PRISMA).



In Console Mode, Thermo-Calc macro files have a \*.TCM extension and the Diffusion Module (DICTRA) file extension is \*.DCM. In addition to macro files in Console Mode the are also *Log* files and *Workspace* files, which also differ.

## CAN I WORK WITH BOTH MODES AT THE SAME TIME?

The two modes can be run simultaneously, but there is no communication between them. What you do in Graphical Mode does not affect the state of the Console Mode session and vice versa. One exception is the plot settings. See <u>"Global Settings: Graphical and Console</u> Mode Plotting" on page 247 in the *Thermo-Calc User Guide*.

## DATA OPTIMIZATION AND THERMODYNAMIC / KINETIC ASSESSMENTS



Although many calculations can be done in either mode, data optimization and



thermodynamic or kinetic assessments are only available in Console Mode.

# **About the File Types and Formats**

The file formats (and the file extensions / suffixes) are based on whether you are working in <u>Graphical or Console Mode</u>, and in Console Mode, whether you are working with Thermo-Calc and/or the Diffusion Module (DICTRA).

## **Graphical Mode**

In Graphical Mode you work with one file type, a *project* file, which has a \*.tcu file extension. This is applicable to Thermo-Calc and the add-on modules.

## **Console Mode**

In Console Mode there are file *types*—macro (including log files) and workspace—and *extensions* —TCM, POLY3, GES5, PAR, POP, and EXP. For Thermo-Calc Console Mode the file types are described in <u>"Macro, Workspace and Experimental Data Files" on page 173</u> in this guide.



For DICTRA Console Mode see *File Formats* as described in the *Diffusion Module* (*DICTRA*) User Guide.

## **Opening Graphical Mode Project Files**

"Opening Console Mode Macro Files" on the next page

## How to Open any Project File

There are different ways to open project files.

## IN THERMO-CALC:

- From the main menu select File → Open Project.
- On the toolbar, click the **Open** button or press <Ctrl+O> on the keyboard. Then navigate to the file on your computer and click **Open**.

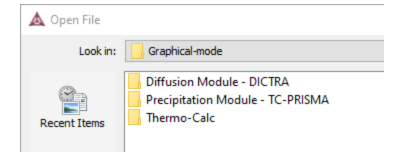
## **ON YOUR COMPUTER:**

• Navigate to the project file and double-click to open it. This launches a new instance of Thermo-Calc.

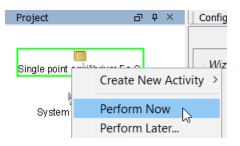
## How to Open an Example Project

To open an example project file for Thermo-Calc and the add-on Diffusion Module (DICTRA) and/or Precipitation Module (TC-PRISMA):

- 1. Open Thermo-Calc.
- 2. Confirm you are in Graphical Mode. If not, click the **Switch to Graphical Mode** button.
- 3. From the main menu, select File  $\rightarrow$  or Help  $\rightarrow$  Examples Files.
- 4. The examples are divided into folders by product (the add-on Diffusion Module (DICTRA) and/or Precipitation Module (TC-PRISMA) and Thermo-Calc). Navigate to the folder and project file you want and double-click to open.



5. The Graphical Mode examples are stored without calculated results so you will want to run the example calculation and generate a plot or table. To do this, click the first node in the tree at the top of the **Project** window. Then right-click and select **Perform now**.



## **Opening Console Mode Macro Files**

"Opening Graphical Mode Project Files" on the previous page



The macro files (with the extension \*.TCM) include comments, which you can either run in Thermo-Calc or open and read in a text editor.

You can open the macro files (\*.TCM or \*.DCM files) in different ways.

- From the main menu,
- Using a command,

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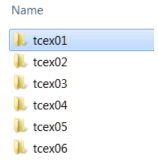
- By dragging and dropping the file into the Console window, or
- Navigate to the file location on your computer and double-click to open it. If you already have Thermo-Calc open, this opens a new instance of the software.
- You can also read the commands listed in the macro file using a text editor.

If you only open and read the macro file in a text editor, you do not see the output that Thermo-Calc gives in response to the commands stored in the macro file.

## **Opening a Macro From the Main Menu**

To open an example macro file for Thermo-Calc:

- 1. Open Thermo-Calc.
- 2. Confirm you are in Console Mode. If not, click the Switch to Console Mode button.
- 3. From the main menu, select File → or Help → Examples Files. The Console Mode folder opens.
- 4. Open the **Thermo-Calc** folder to see the available examples contained in separate folders.



▲ For the Diffusion Module (DICTRA) examples (\*.DCM macro files), open the Diffusion Module - DICTRA folder.

- 5. In the folder, double-click the \*.TCM file or click **Open**.
- 6. Follow the prompts in the Console window i.e. keep pressing <Enter> to run the macro and produce a plot in the **Console Results** window.

## **Opening a Macro Using a Command**

At the SYS prompt, type MACRO\_FILE\_OPEN and press <Enter>. In the file dialogue window that opens, navigate to and select the macro file to open.

## Drag and Drop a Macro File into the Console Window

For example, on your computer, navigate to the examples folder. Drag the file from its location into the Thermo-Calc Console window. When you release (drop) the file, the macro starts running. Follow the prompts in the Console window i.e. keep pressing <Enter> to run the macro and produce a plot in the **Console Results** window.

Console	
Console 1 💿 🔲 😑	
Thermo-Calc / DIO Only for use at Local contact reference and w SYS: → Move ↓ texQ2	
File Home Share View	
Pin to Quick Copy access Copy Paste	Move to • X Delete
Cipboard	Organize
← → ✓ ↑ 🔪 « Examples > Cons	sole-mode > tcex02
Name ^	Date r
tcex02.TCM	3/9/20
	õ <sup>a</sup> 0 ×
Console 1 💿 🔲 🚍	
Thermo-Calc / DICTRA Only for use at Server: thoth.ad.thermocalc.s Local contact Subnets: Clients not locked to	



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Drag and drop files into the Console window to open or run a macro, for example. File types that can be opened this way are macro files (\*.TCM and \*.DCM), database files (\*.TDB), and workspace files (\*.POLY3, \*.GES5,\*.PAR, and \*.DIC). Note that \*.DCM and \*.DIC file extensions are applicable to the Diffusion Module (DICTRA).

# **Examples, Manuals and Materials Folders**

There are several ways to access the examples, manuals (containing PDF documentation) and materials data folders.

	Main Menu	System Definer	From the Windows Start Menu	On your computer	Website
Manuals (PDF documentation, DATAPLOT EXP files, and Property Model Framework Tutorial Examples)	x	-	x	x	x
Examples (project and macro files)	x	-	x	x	x
Materials	-	x	x	x	x

## FROM THE MAIN MENU

Once your Thermo-Calc software is installed you can open the Examples and PDF documentation (Manuals) from the main menu:

- To open the Manuals folder select Help → Manuals Folder. Note for Linux openSUSE users: This link is not functional. To access the folder, navigate to the default location based on your installation type
- To open the Examples based on whether you are in Console Mode or Graphical Mode, select File → Examples Files or Help → Examples Files.

## FROM THE SYSTEM DEFINER (MATERIALS ONLY)

When you are working with the <u>"System Definer" on page 62</u> you can open the default **Materials** folder included with your installation.

At the bottom of the window under **Material**, click **Load material** to navigate to a material file.

Material Material name Amount N Fe	e: Mass percent V 99.99	
С	0.01	
	Copen Look in: Recent Items Desktop Documents	<ul> <li>Materials</li> <li>Aluminum Alloys</li> <li>Cast Iron</li> <li>My Materials</li> <li>Nickel Alloys</li> <li>Solder Alloys</li> <li>Steel</li> <li>Titanium Alloys</li> </ul>
	Network	File name:     Open       Files of type:     Material file     Cancel
	material	

## START MENU (WINDOWS ONLY)

On Windows, once Thermo-Calc is installed, you can locate the Examples and Materials folders, plus all the Manuals using the shortcuts located in the Start menu. Go to Start  $\rightarrow$  All **Programs** or All Apps  $\rightarrow$  Thermo-Calc and click Examples, Manuals, or Materials as required to open the applicable folder.

### **ON YOUR COMPUTER (ALL OPERATING SYSTEMS)**

The PDFs, example files and materials xml files are included with your installation in a folder

location based on the operating system.

In the table, *<user>* stands for the username and *<version>* for the version of Thermo-Calc, for example 2018b.

OS	User type	Default directory		
Windows	Normal user	Users\< <i>user</i> >\Thermo-Calc\< <i>version&gt;</i> Users\< <i>user</i> >\Documents\Thermo- Calc\< <i>version&gt;</i> My documents		
	Administrator	Program Files\Thermo-Calc\< <i>version&gt;</i> Users\Public\Documents\Thermo- Calc\< <i>version&gt;</i> Public documents		
Mac	Administrator (user name and password required)	Examples and manuals in /Users/Shared/Thermo- Calc/< <i>version&gt;</i> To go to this folder, in Finder, from the <b>Go</b> main menu select <b>Go to folder</b> . Enter the above file path and click <b>Go</b> .		
Linux	Non root user	home/ <user>/Thermo-Calc/<version></version></user>		
LIIIUX	Root user	usr/local/Thermo-Calc/ <version></version>		

## WEBSITE

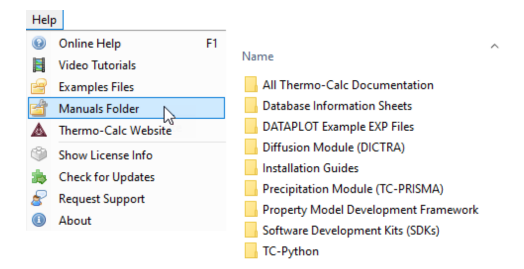
You can download the PDF documentation and examples from the website.

- 1. Go to http://www.thermocalc.com/.
- From the menu, select Support → Documentation and navigate to the applicable section.

## About the PDF Documentation Folder

There is PDF documentation available as part of your installation. This is the same information as that contained in the online help, just in a different format. The contents of each folder are described below.

Examples, Manuals and Materials Folders" on page 12



### **All Thermo-Calc Documentation**

Includes all the manuals needed to work with Thermo-Calc including the installation guides, Thermo-Calc User Guide, Console Mode Command Reference, Graphical Mode Examples Guide, Data Optimization User Guide, Database Manager User Guide, and the DATAPLOT User Guide. It also includes a copy of the Property Model Development Framework Guide.

#### **Database Information Sheets**

The extended information database information sheets are included in the online help system and in this sub-folder. You can also download these from the Thermo-Calc website.

#### **DATAPLOT Example EXP Files**

The examples are available as supplementary information as described in the *DATAPLOT User Guide*, which you can find in either the online help, Thermo-Calc Documentation Set or Diffusion Module (DICTRA) Documentation Set.

#### **Diffusion Module (DICTRA)**

This add-on module is used for accurate simulation of diffusion controlled reactions in multicomponent alloy systems, which is based on numerical solution of the multicomponent diffusion equations. The documentation set includes Diffusion Module (DICTRA)-specific manuals plus the Thermo-Calc User Guide, Console Mode Command Reference, Data Optimization User Guide, Database Manager User Guide, and the DATAPLOT User Guide.

The following guides are also available separately in this folder as well included in the documentation set:

- Diffusion Module Quick Guide Console Mode
- Diffusion Module Quick Guide Graphical Mode

### Installation Guides

Includes the quick install guides for Standalone Windows and Mac installations as well as the full installation guide for all platforms and installation types.

### Precipitation Module (TC-PRISMA)

This add-on module treats concurrent nucleation, growth/dissolution and coarsening under arbitrary heat treatment conditions in multicomponent and multi-phase systems using Langer-Schwartz theory and the Kampmann-Wagner numerical approach.

### **Property Model Development Framework**

The PDF guide contains details about setting up a development environment so you can build custom property models in Thermo-Calc.

The folder also includes tutorial Python files you can copy and use in your own custom property models. The examples are available as supplementary information as described in the *Property Model Development Framework Guide*.

### Software Development Kits (SDKs)

- **TQ-Interface Programmer Guide**: The TQ-Interface is available as a way to make Thermo-Calc an integral part of application programs such as those for process simulation, microstructure evolution modeling and materials property prediction.
- TC Toolbox for MATLAB<sup>®</sup> Programmer Guide: Use this to enable communication between Thermo-Calc and MATLAB to be able to retrieve thermodynamic and kinetic quantities.
- **TC-API Programmer Guide**: This is the Thermo-Calc API that can be used by programmers to set up your own applications to work with Thermo-Calc.

## **TC-Python API Reference Documentation**

TC-Python is an SDK and you can search and use the special HTML file located in this folder. Double-click the Index.html file to open it.

## About the Examples

To learn more about how to use the software you can open and run the example projects (Graphical Mode) and macro files (Console Mode).

## Graphical Mode Examples

Graphical Mode examples are available for Thermo-Calc and the add-on Diffusion Module (DICTRA) and Precipitation Module (TC-PRISMA). These are in the format of project files (\*.tcu).

A brief description of all the examples, and how to access them, is in the *Graphical Mode Examples Guide*, which you can find in this documentation set or by searching the online help.

## **Console Mode Examples**

Console Mode examples are available for Thermo-Calc and the Diffusion Module (DICTRA). The *macro* files are in the format of \*.TCM and .DCM, respectively.

There are other file types when you are working in Console Mode. See <u>"Thermo-Calc Console Mode File Types" on page 2</u> in the *Console Mode Command Reference*, and *About the File Formats* in the *Diffusion Module Quick Start Guide*.

## **Examples by Product**

## THERMO-CALC EXAMPLES

A brief description of the Thermo-Calc examples, and how to access them, is in <u>"About the Thermo-Calc Console Mode Examples</u>" in the *Thermo-Calc Console Mode Examples Guide*.

## **DIFFUSION MODULE (DICTRA) EXAMPLES**

A brief description of the Diffusion Module examples (both Graphical Mode and Console Mode), and how to access them, is in the *Diffusion Module (DICTRA) Examples Guide*.

## **PRECIPITATION MODULE (TC-PRISMA)**

A brief description of the Precipitation Module examples (only in Graphical Mode), and how to access these, is in <u>"Precipitation Calculator Examples" on page 52</u> in the *Graphical Mode* 

## Examples Guide.

## **TC-PYTHON EXAMPLES**

A brief description of the TC-Python SDK examples, and how to access these, is available in the Help.

## TQ-INTERFACE AND TC-TOOLBOX FOR MATLAB EXAMPLES

There are examples for the TQ-Interface and TC-Toolbox for MATLAB available as described in <u>"Examples, Manuals and Materials Folders" on page 12</u>. You can also find information about these in the respective user guides or by searching the online help.

## About the Material Files

## "Examples, Manuals and Materials Folders" on page 12

When you are working with the <u>"System Definer" on page 62</u> you can open the default **Materials** folder included with your installation. The files are in .xml format and the data contained are from some common alloys. You can also save and create your own materials files from the System Definer, or copy the xml files to the Materials folder.

## **ALUMINUM ALLOYS**

- UNS\_A95005
- UNS\_A96053

## CAST IRON ALLOYS

- UNS\_F10007
- UNS\_F33101

## NICKEL ALLOYS

- UNS\_N06600
- UNS\_N6600
- UNS\_N06625
- UNS\_N06690
- UNS\_N07001

- UNS\_N07718
- UNS\_N08800
- UNS\_N08825

#### **SOLDER ALLOYS**

- ASTM\_25A
- ASTM\_96TS
- UNS\_L50134
- UNS\_L54321

## **STEEL ALLOYS**

## Austenitic Stainless Steel

- ASTM\_304\_(UNS\_S30400)
- ASTM\_304L\_(UNS\_S30403)
- ASTM\_304LN\_(UNS\_S30453)
- ASTM\_316\_(UNS\_S31600)
- ASTM\_316L\_(UNS\_S31603)
- ASTM\_316LN\_(UNS\_S31653)
- ASTM\_317L\_(UNS\_S31703)
- UNS\_J92710
- UNS\_\$30908

### **Duplex Stainless Steel**

- UNS\_S31803\_A
- UNS\_S32750\_A
- UNS\_S32750\_B

### High Speed Steel

• ASTM\_M10\_A5

## High Temperature Steel

• UNS\_\$31008

## HSLA Steel

• UNS\_K12000

### Low Carbon Steel

• UNS\_H61180

### Maraging Steel

• UNS\_K92810

### Martensitic Stainless Steel

- ASTM\_420
- UNS\_J82090
- UNS\_J91639
- UNS\_S44004

### Tool Steel

- UNS\_T30106
- UNS\_T30109
- UNS\_T51620

## Training Steel

- Ferritic
- M7\_Steel

### **TITANIUM ALLOYS**

- UNS\_R54520
- UNS\_R56260

## **My Materials**

This is where you can add your own material files to be able to load it to the System Definer. When you are in the System Definer Configuration window, you define your material then click **Save material as**. You can save the profile to this folder or create a folder with a new name.

# **Help Resources**

## 'Online' Help: No Internet Required

To access online help in a browser, open Thermo-Calc and press <F1> on the keyboard, or select Help  $\rightarrow$  Online Help.

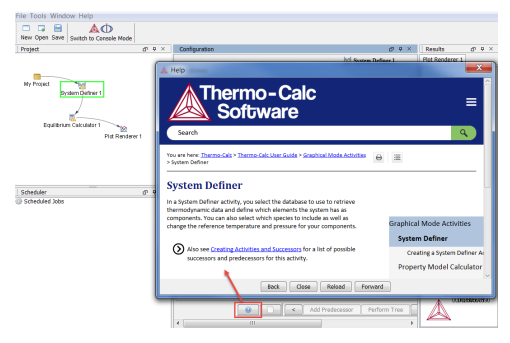
The content opens in a browser but uses local content so you don't need an Internet connection except for links to external websites.

# **Context Help (Graphical Mode Only)**



When you are in Graphical Mode, you can access feature help (also called *topic-sensitive* or *context help*) for the activity nodes in the tree.

- 1. In the Project window, click a node. For example, **System Definer**.
- 2. In the lower left corner of the **Configuration** window, click the help button <sup>1</sup>
- 3. The **Help** window opens to the relevant topic.



The window that opens has the same content as the help you access in the browser. There are these extra navigation buttons in this view.

• Use the buttons on the bottom of the window, Back Close Reload Forward, to navigate **Back** and **Forward** (these are only active once you have started using the help to load pages and create a history), to **Close** the window, and **Reload** the original content.

## **Console Mode Help**



Console Mode is for Thermo-Calc and the Diffusion Module (DICTRA).

In Console Mode at the command line prompt, you can access help in these ways:

- For a list of all the available commands in the current module, at the prompt type a question mark (?) and press <Enter>.
- For a description of a specific command, type Help followed by the name of the command. You can only get online help about a command related to the current module you are in.
- For general system information type Information. Specify the subject or type ? and the available subjects are listed. This subject list is specific to the current module.

# **The Thermo-Calc Modules and Activities**

Although many calculations can be done in either Graphical or Console Mode, only data optimization and thermodynamic or kinetic assessments are available in Console Mode. Conversely, only the Property Model Calculator is available in Graphical Mode.



For the Console Mode only commands see <u>"The Console Mode Interface" on</u> page 158 and the *Thermo-Calc Console Mode Command Reference* included with this documentation set.

The table briefly describes the available modules. The specific modules, using the names in the table, are accessed through Console Mode. For Graphical Mode, the underlying principles, and the final calculations, are incorporated into the projects and activities. Some modules require additional licenses.

The first and second columns are the Console Mode module names. The third column indicates whether there is an equivalent activity available in Graphical mode.

Full Module Name in Console Mode	Short Name	Graphical Mode Activity	Primary Functions
BINARY_ DIAGRAM_ EASY	BIN	<u>"Binary</u> <u>Calculator"</u> on page 150	Calculate binary phase diagrams. Access to specific databases designed for BIN, such as TCBIN, is required.
DATABASE_ RETRIEVAL	DATA	<u>"System</u> Definer" on page 62	Select appropriate databases, define chemical systems and retrieve the data required for calculations.
DICTRA_ MONITOR	DICTRA	"Diffusion Calculator" on page 92	Run simulations of diffusion controlled transformations in Thermo-Calc using the Diffusion Module (DICTRA). An additional license is required for this feature.
DIC_PARROT	DIC_ PARROT	Console Mode only	This is the data optimization module for the Diffusion Module (DICTRA). It allows you to assess experimental data and use such data to optimize calculations. An additional license is required for this feature.
EDIT_ EXPERIMENTS	ED_EXP	Console Mode only	Part of the PARROT submodule, use it to edit experimental data points to make reliable optimizations.
GIBBS_ ENERGY_ SYSTEM	<u>GIBBS</u>	Console Mode only	Handles thermodynamic models and quantities. Generally you do not need to use this module, but you can get information about models, and list, add or amend the thermodynamic parameters in use.
PARROT	PARROT	Console Mode only	A data optimization module to assess experimental data and use the data to optimize calculations.

Full Module Name in Console Mode	Short Name	Graphical Mode Activity	Primary Functions
not available	-	"Precipitation Calculator" on page 104	Set the conditions for, and perform, a precipitation calculation.
not available	-	"Property Model Calculator" on page 70	You can evaluate models simultaneously over a range of compositions and cross plot the results. You can then conduct an uncertainty analysis and plot the results as either a histogram or as a probability plot.
POLY-3	POLY	"Equilibrium Calculator" on page 83	Equilibrium calculation module. Specify conditions and make calculations.
POST_ PROCESSOR	<u>POST</u>	<u>"Plot</u> <u>Renderer" on</u> page 122	For post-processing and plotting of calculation results. Modify property and phase diagrams, add labels, change the diagram colours, etc. Submodules to POLY-3 and DICTRA module for example.
POTENTIAL_ DIAGRAM	<u>POTENTIAL</u>	"Ternary Calculator" on page 152	Calculate and plot gas potential diagrams in a ternary system. The potential of two gas species are used as the diagram axes.
POURBAIX_ DIAGRAM	<u>POURBAIX</u>	Console Mode only	Calculate and plot pH-Eh diagrams (Pourbaix diagrams) and property diagrams for heterogeneous interaction systems involving aqueous solutions.

Full Module Name in Console Mode	Short Name	Graphical Mode Activity	Primary Functions
REACTOR_ SIMULATOR_3	REACTOR	Console Mode only	Simulate chemical reaction processes in several feed-forward steady-state stages or in several dynamic stages.
SCHEIL_ SIMULATION	<u>SCHEIL</u>	<u>"Scheil</u> Calculator" on page 154	Simulate and plot Scheil-Gulliver solidification processes with no diffusion in the solid phases and with/without consideration of back diffusion of interstitial components (such as C, N, O, S).
SYSTEM_ UTILITIES	<u>SYS</u>	Console Mode only	In Console Mode, interact with the operating system and change environmental settings. Create and execute MACRO files.
TABULATE_ REACTION	TAB	Console Mode only	Tabulate properties of chemical reactions and substances (stoichiometric or solution phases).
TERNARY_ DIAGRAM	<u>TERN</u>	"Ternary Calculator" on page 152	Calculate ternary phase diagrams.

# **Displaying Thermo-Calc License Information**

You can start (and install) the software without a valid Thermo-Calc license but you cannot do any calculations. To show information about the available and installed licenses, from the main menu select Help  $\rightarrow$  Show License Info.

## Thermo-Calc Installation Guide

This is an example of part of a License Information window for a local installation.

A License Information

```
License diagnostics program
  build: 10636 created: Thu Feb 23 17:29:02 2017
Log generated at: 28-02-2017 12:19:56
Safenet environment variables
 LSHOST: NO-NET
 LSERVRC: C:\Program Files/Thermo-Calc/2017a/lservrc
Local (client) info
 hostname:
 license version:
 ethernet:
 ip address:
               #1
 ip address: #2
               #1
 dns name:
 dns name:
               #2
```

### **NETWORK LICENSES**

For a *network* license, under each license type with instances checked out, there is a list of who checked out each license.



In Graphical Mode, an instance is checked out when an Equilibrium Calculator activity is created, and checked back in when that Equilibrium Calculator is removed. For the add-on Diffusion Module (DICTRA) and/or Precipitation Module (TC-PRISMA) this also applies to the Diffusion Calculator and the Precipitation Calculator.



In Console Mode, an instance is checked out when you enter the POLY module and is checked back in when you exit the POLY module.

- The Features column is a list of the specific software, databases and API licenses purchases.
- The Version is an internal version number for the feature.
- The **number of licenses** column is how many instances of each license type available to be checked out simultaneously.
- The in use column shows how many license instances are checked out.

#### **CONSOLE MODE LICENSE RESTRICTIONS**

• If you start Thermo-Calc without a valid Thermo-Calc license you cannot leave the

SYS module.

- If all the licenses are checked out for a network client installation of Thermo-Calc, you may not be able to enter the POLY or DICTRA modules until a license is available.
- To enter the DICTRA and the DIC\_PARROT modules, you need a valid Diffusion Module (DICTRA) license key.
- If you enter and use either the DIC\_PARROT module or the PARROT module, then you cannot enter the other module in the same session. You must close down Thermo-Calc and launch the program again to enter the other data optimization module.

## **NETWORK COMPUTER LICENSE RESTRICTION**

If you have a network computer installation of Thermo-Calc, then you may not be able to do any calculations even if you have access to a valid network license file. This is because others who are part of your network installation may have checked out all allowed instances of the software.

If you are running a network installation of Thermo-Calc, you can see how many instances of the client licenses are currently checked out and how many of them are left. You can also see which client computers have checked out the licenses.

# **Using This Guide**

If you are a new user, Graphical Mode is a good way to learn how to work with Thermo-Calc. Depending on what you want to simulate, Console Mode can have more functionality but until you are comfortable with the concepts, it is recommended you start in Graphical Mode.



Additional documentation and training material is described in<u>"Help Resources"</u> on page 21.

## **GRAPHICAL MODE**

"The Graphical Mode Interface" on page 29 introduces you to the GUI layout and the workflow. In "Graphical Mode Projects" on page 49, it goes into more detail about how to set up and run your projects (with the Quick Start Wizard), using templates, scheduling jobs and creating activities. "Graphical Mode Activities" on page 61 specifically describes how to set up each type of available activity in the tree structure.

For both Graphical and Console mode, <u>"Changing Global Settings" on page 238</u> gives information about adjusting global user settings.

## **CONSOLE MODE**

In general, Console Mode has more functionality than Graphical Mode but the underlying concepts are the same. There are, however, command lines and extensive details applicable to this mode. This guide is only the beginning of the learning objectives. In particular, refer to the *Thermo-Calc Console Mode Command Reference* for details about all the commands available.

In this guide it is a basic introduction to the layout of the Command windows, as described in "The Console Mode Interface" on page 158.

Several topics describe and give an example of the calculation types:

- "Equilibrium Calculations" on page 181
- "Property Diagrams" on page 186
- <u>"Phase Diagrams" on page 191</u>
- "Scheil Simulations" on page 196
- <u>"T0 Temperature Simulations" on page 203</u>
- "Paraequilibrium" on page 206
- "Potential Diagrams" on page 210
- "Aqueous Solutions" on page 214
- "Console Mode TAB Module" on page 223

Then in <u>"Console Mode POST Module" on page 228</u> there is an overview of plotting.

For both Graphical and Console mode, <u>"Changing Global Settings" on page 238</u> gives information about adjusting global user settings.

# **The Graphical Mode Interface**

These topics introduce you to the concepts and terminology of Graphical Mode and describe the basic workflow.

## In this section:

_		
	The GUI Layout	30
	Project Activities and the Tree Structure	31
	Activity Successor and Predecessor Nodes	31
	Creating Activities and Successors	32
	Graphical Mode Workflow	34
	Main Menu, Toolbar and Keyboard Shortcuts	35
	Saving and Opening Project Files	38
	Project Activity Nodes	39
	Project and Activity Menus	41
	Organizing the Windows on the Desktop	44
	Moving Nodes in the Tree and Using the Grid	45
	Zooming In and Out	46
	Resetting a Plot to the Default View	47
	Node Status Markers	47
	Rotating 3D Plots	48

# **The GUI Layout**

Open Thermo-Calc to view the default GUI layout for Graphical Mode. You can easily reorganize the windows.



The windows are:

- 1. **Project**: Create, manipulate and navigate between the activities that make up a project.
- 2. **Configuration**: Shows the settings that can be configured for the currently selected activity.
- 3. **Results**: Shows the results of a calculation, either plotted as a diagram or displayed in table format. In this example, there is a Plot Renderer *tab* in the Results window.
- 4. **Scheduler**: Displays information about jobs, such as calculations, that are being performed or are scheduled to be performed. You can cancel scheduled jobs and if a job has failed, then you can view information about the error.
- 5. **Event Log**: By default this window is closed but it displays during calculation processes. You can always open it to view progress of calculations and to troubleshoot.

### Graphical Mode Windows

In this example, the Results window also has a Plot Renderer tab. You can also have a Table Renderer tab.

File Tools Window Help	
□ □ □ □ □ ▲ ① New Open Save Switch to Console Mode	2
Project 라무×	Configuration
Step in temperature FE-C	Plot Renderer 1       Save Diagram Show Triangular Show Grid Switch Axes Retain Labels       Beguilbrium Calculator 1
System Definer 1	Tie lines:     0 (m)       X Axis
Equilibrium Calculator 1	Avis variable: Temperature Kelvin Avis type: Linear Limits: 500.0 to 3000.0 step 250.0 V Automatic
Scheduler ar e × Scheduled Jobs	Aus variable: l'emperature Keivin Aus variable: L'inear Y Azis Aus variable: Amount of phase No normalizat Aus variable: Amount of phase O. 6- Aus variable: Amount of phase O. 6- L'imits: 0.0 to 1.0 step 0.1 V Automatic
12:12:01,718 INFO Phase Region from 12:12:01,718 INFO BCC_A2#1 FCC_A1#1	ding phase at 1.01117E+03 5

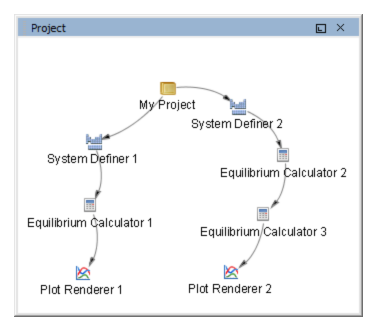
# **Project Activities and the Tree Structure**

In a *project*, a set of linked activities is called a *tree*. A result calculated within a tree is fed as input into the next *activity* in the tree. Consequently, if you have an Equilibrium Calculator with another Equilibrium Calculator as its *successor*, the successor takes the calculation results of the *predecessor* as the starting values for its calculation. When working with activities in the Project window these are also referred to as *nodes* (or *activity nodes*) in the tree structure.

## "Activity Successor and Predecessor Nodes" below

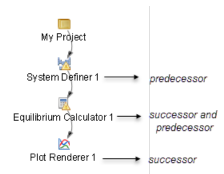
The example below shows a My Project node with two trees. In each tree, settings and calculation results are propagated downward until the calculation and the visualisation of the results are completed in the Plot Renderer nodes.

The system definitions, settings and calculation results of the two trees are independent of each other.



# **Activity Successor and Predecessor Nodes**

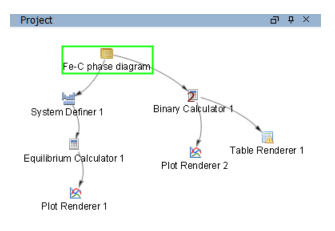
An activity node located below another activity node in a tree is referred to as that activity's *successor*. An activity located above another activity is called that activity's *predecessor*. A predecessor is performed before the predecessor's successors and its result is fed forward to any successor activities.



For example, to calculate and display a phase diagram, create a branch with three linked activities: A System Definer activity linked to an Equilibrium Calculator activity, which in turn is connected to a Plot Renderer activity.

See the example below that shows that Binary Calculator 1 is a predecessor to the successors, Plot Renderer 2 and Table Render 1.

You can determine the available successors by right-clicking a node in the Project window and choosing options from the menus.



# **Creating Activities and Successors**

You can create a new activity in these ways:

- In the **Project** window, right-click an activity and choose **Create New Activity (My Project** node only) or **Create New Successor** (all other nodes).
- Select the activity you want to create from the submenu.
- At the bottom of the **Configuration** window, click **Create New Activity** or **Create New Successor** and select the activity to create.

Activity	Possible Predecessors	Possible Successors
My Project	None; this is the first node of the tree structure	System Definer, Binary Calculator, Ternary Calculator, Experimental File Reader. Also choose templates from this level.
<u>"System Definer" on</u> page 62	My Project, System Definer	Property Model Calculator, Equilibrium Calculator, Precipitation Calculator, Scheil Calculator
"Property Model Calculator" on page 70	System Definer	Plot Renderer, Table Renderer
"Equilibrium Calculator" on page 83	System Definer, Equilibrium Calculator	Equilibrium Calculator, Plot Renderer, Table Renderer
"Precipitation Calculator" on page 104	System Definer	Plot Renderer, Table Renderer
"Diffusion Calculator" on page 92	System Definer	Plot Renderer, Table Renderer
<u>"Plot Renderer" on</u> page 122	Property Model Calculator, Equilibrium Calculator, Precipitation Calculator, Diffusion Calculator, Binary Calculator, Ternary Calculator, Scheil Calculator, Experimental File Reader	None

Activity	Possible Predecessors	Possible Successors
<u>"Table Renderer" on</u> page 146	Equilibrium Calculator, Property Model Calculator, Diffusion Calculator, Precipitation Calculator, Binary Calculator, Scheil Calculator	None
"Experimental File Reader" on page 149	My Project	Plot Renderer
"Binary Calculator" on page 150	My Project	Plot Renderer, Table Renderer
"Ternary Calculator" on page 152	My Project	Plot Renderer
"Scheil Calculator" on page 154	System Definer	Plot Renderer, Table Renderer

# **Graphical Mode Workflow**

You can set up a tree in the Project window (starting with the My Project node) and then perform all the activities at once, or create and perform one activity at a time.

The typical workflow is to:

- 1. *Define a System:* Create a <u>"System Definer" on page 62</u> activity (in the **Project** window) where you select a database and the elements to have as system components (in the **Configuration** window).
- Set up and run a calculation: Create an <u>"Equilibrium Calculator" on page 83</u> activity (a successor to the System Definer), where calculation conditions are set (temperature, pressure, etc.). This is where axis variables are set to create a property or phase diagram. These settings are also made in the Configuration window. You can also add a <u>"Property Model Calculator" on page 70</u>, <u>"Precipitation Calculator" on page 104</u>, <u>"Diffusion Calculator" on page 92</u> or <u>"Scheil Calculator" on page 154</u>.
- 3. *Visualize the results:* Create either a <u>"Plot Renderer" on page 122</u> or a <u>"Table</u> <u>Renderer" on page 146</u> activity in the **Project** window. When calculated, this creates

a plot or table in the **Results** window. The results are shown in a **Plot Renderer** or a **Table Renderer** tab.



You need to **Perform** a Plot Renderer or a Table Renderer activity to visualize the calculation results from an Equilibrium Calculator.

# Main Menu, Toolbar and Keyboard Shortcuts

F	File Tools Window Help			
*****				۵
	New	Open	Save	Switch to Console Mode

The **File** menu and toolbar are reserved for Graphical Mode but in Console Mode the **Tools**, **Window** and **Help** menus have the same options.



In Console Mode you use commands to access the other options, for example see <u>"Main Menu Commands" on page 161</u>.

## **GRAPHICAL MODE MENU AND TOOLBAR OPTIONS**

Option	Description	Action (s)
	ew Create a new project.	Click <b>New</b> on the toolbar
New		Select <b>File → New</b> <b>Project</b>
		Windows: Press <ctrl+n></ctrl+n>
		Mac: Press < ℜN>
	Open an existing project. See <u>"Saving and</u> Opening Project Files" on page 38.	Click <b>Open</b> on the toolbar
Open		Select <b>File → Open</b> Project
		Windows: Press <ctrl+o>.</ctrl+o>
		Mac: Press < 光O>

Option	Description	Action (s)
Save	Save a project. See <u>"Saving and Opening</u> Project Files" on page 38.	Click <b>Save</b> on the toolbar Select <b>File → Save</b> <b>Project</b> Windows: Press <ctrl+s>. Mac: Press &lt; ℋS&gt;</ctrl+s>
Save Project As	Save an existing project with a new name.	Select File → Save Project As Windows: Press <ctrl+shift+s> Mac: Press &lt; îHS&gt;</ctrl+shift+s>
Switch to Console Mode	Launch Console Mode and the command prompt.	Click the button to exit Graphical Mode and open Console Mode.
Switch to Graphical Mode	Launch Graphical Mode to work with the GUI version.	Click the button to exit Console Mode and open Graphical Mode.
Append Project	Combine two projects into one project file. See <u>"Saving and Opening Project Files" on</u> page 38.	Select File → Append Project Windows: Press <ctrl+shift+o> Mac: &lt; îૠO&gt;</ctrl+shift+o>
Exit	Exit the program.	In the upper right- corner of the program, click the X. Select <b>File → Exit</b> Windows: Press <ctrl+q>.</ctrl+q>

Option	Description	Action (s)
		Mac: <
Options	Change the global defaults for a variety of settings. See <u>"Changing Global Settings" on page 238</u> .	Select <b>Tools →</b> Options
Database Checker	Open the Database Checker, a program to check that the syntax of Thermo-Calc database files is correct. This is for advanced users who develop and manage databases.	Select <b>Tools →</b> Database Checker
Window menu	Highlight a specific window on the GUI	From the <b>Window</b> menu, select an option to refocus on that specific window.
Help	Open the online help system where you can browse or search all the Thermo-Calc documentation. This does not require an Internet connection. The content is stored locally.	Press <f1> Select <b>Help → Online</b> <b>Help</b></f1>
Video Tutorials	Open a link to the Thermo-Calc website where a variety of video tutorials are available.	Select <b>Help → Video</b> Tutorials
Examples Files	Open the examples folder based on whether you are in Console Mode or Graphical Mode.	Select File → or Help → Examples Files
Manuals Folder	Open the folder on your computer that has the PDF documentation and guides.	Select <b>Help →</b> Manuals Folder
Show License Info	Open the License Information window.	Select <b>Help → Show</b> License Info

Option	Description	Action (s)
	Also see <u>"Displaying Thermo-Calc</u> <u>License Information" on page 25</u> and the <i>Thermo-Calc Installation</i> <i>Guide</i>	
Check for update	Check the Thermo-Calc Software website for updates to the software.	Select Help → Check for Updates
Request Support	Open a link to the Support page on the Thermo-Calc website. There are instructions about how to submit your question to the email address below. http://www.thermocalc.com/support/request- support/ E-mail: support@thermocalc.com	Select <b>Help → Request</b> Support
About	Includes the version number of the installed software and information about third party software licenses.	Select <b>Help →About</b>

# **Saving and Opening Project Files**

Thermo-Calc Graphical Mode uses project files with the file name suffix .tcu. The add-on Diffusion Module (DICTRA) and/or Precipitation Module (TC-PRISMA) also use the same file format in Graphical Mode.



For information about file formats in Console Mode, see <u>"About the File Types</u> and Formats" on page 8.

To save your project and all its settings and results, on the toolbar click **Save**. Select to **Include calculated results in the project file** (the default) as required.

To open a Thermo-Calc project file, on the toolbar click **Open** and select the project file in the **Open file** window.

You can only have one project file open at a time. However, you can attach (*append*) the trees from additional project files to the topmost My Project node. To append an additional project file in this way, from the main menu select **File**  $\rightarrow$  **Append Project** and open a project file.

Saving Plots" on page 140 and Saving Tabulated Data for the <u>"Table Renderer" on</u> page 146

# **Project Activity Nodes**

The table below is a brief description of the project activity types. When working with activities in the Project window these are also referred to as *nodes*.

# ACTIVITY NODES

Activity type	Description
"System Definer" on page 62	Define a certain thermodynamic system and read it from file into memory.
"Property Model Calculator" on page 70	Choose one or more general models to predict and optimize material properties. You can evaluate models simultaneously over a range of compositions and cross plot the results. You can then conduct an uncertainty analysis and plot the results as either a histogram or as a probability plot.
"Equilibrium Calculator" on page 83	Set thermodynamic conditions and define axis variables when a series of equilibrium calculations are to be performed in one or more dimensions.
"Precipitation Calculator" on page 104	Set the conditions for, and perform, a precipitation calculation using the Precipitation Module (TC-PRISMA).
"Diffusion Calculator" on page 92	Using the Diffusion Module (DICTRA), this activity is where you set the geometry, add one or more regions, enter grids and phases in the regions, and then enter compositions of phases.

These are the project activity nodes available in Graphical Mode.

Activity type	Description	
"Plot Renderer" on page 122	Determine the layout of non-text based output.	
"Table Renderer" on page 146	Use it for text-based output.	
<u>"Binary Calculator"</u> on page 150	Use it for some calculations involving two components only. It is like a combination of System Definer and Equilibrium Calculator activities with adaptations to simplify the configuration of calculations on binary systems. To perform this activity, you need a database designed for the Binary Calculator, such as the TCBIN database.	
<u>"Ternary</u> <u>Calculator" on</u> page 152	Use it for some calculations involving three components. It is like a combination of System Definer and Equilibrium Calculator activities with adaptations to simplify the configuration of calculations on ternary systems. To get a reliable result you need a database that fully assesses binary and ternary systems.	
<u>"Scheil Calculator"</u> on page 154	Perform Scheil-Gulliver calculations (also known as Scheil calculations). A default Scheil calculation is used to estimate the solidification range of an alloy assuming that i) the liquid phase is homogeneous at all times and ii) the diffusivity is zero in the solid. However, it is possible to disregard the second assumption for selected components.	
"Experimental File Reader" on page 149	Read experimental data files (*.EXP). This type of file contains information specifying a plotted diagram, written in the DATAPLOT graphical language.	

# **Project and Activity Menus**

When working in the Project window you can right-click nodes to open menus with the available actions. For example, click the **My Project** node to open the menus. Most nodes in the tree have a common menu with differences due to placement in the tree and the activity type.

## MENUS

## My Project Node

My Project	Create New Activity >	System Definer
	Perform Now Perform Later	Binary Calculator Ternary Calculator Experimental File Reader
	Rename	Use Template
	Apply Auto Layout Snap to Grid Show Grid	
	Zoom In Zoom Out Reset Zoom	

## Activity Node

Create New Successor 🔅	Equilibrium Calculator
Perform Now Perform Later	Scheil Calculator Property Model Calculator Diffusion Calculator
Rename Remove	Precipitation Calculator
Clone Clone Tree	
Apply Auto Layout Snap to Grid Show Grid	
Zoom In Zoom Out Reset Zoom	

## MENU OPTIONS FOR ACTIVITIES

Option	Description	Action (s)
Create New Activity	Add System Definer, Binary Calculator, Ternary Calculator, and Experimental File Reader nodes. See <u>"Creating</u> <u>Activities and</u> <u>Successors" on</u> page 32.	Right-click <b>My Project</b> and from the <b>Create</b> <b>New Activity</b> menu choose an option.
Create New Activity → Use Template	Choose from the available templates to use the Wizard or add predefined nodes.	Right-click <b>My Project</b> and from the <b>Create</b> <b>New Activity → Use Templates</b> menu choose an option.

Option	Description	Action (s)
	See <u>"Creating a</u> Project from a Template" on page 52.	The templates are also available on the <b>Configuration</b> window, which you can access by clicking the <b>My Project</b> node, or from the <b>Window</b> menu.
Create New Successor	Add a successor to the selected node. See <u>"Creating</u> <u>Activities and</u> <u>Successors" on</u> <u>page 32</u> .	Right-click one of these nodes System Definer, Property Model Calculator, Equilibrium Calculator, Precipitation Calculator, Diffusion Calculator, Binary Calculator, Ternary Calculator or Experimental File Reader, and from the Create New Successor menu choose an option.
Add a Predecessor	Add a predecessor to the selected node.	Right-click a node to add a predecessor to it (when available).
Perform Now or Perform Later	Perform a calculation or create a plot or table immediately, or schedule it for a time in the future. See "Using the Scheduler" on page 58.	Right-click any node.
Rename	Change the name of the node.	Right-click any node.
Remove	Delete the selected node from the tree.	Right-click any node except My Project.
Clone	Duplicate the selected node and add it to the same tree. See <u>"Cloning</u> <u>Activities and Trees"</u> on page 53.	Right-click any node except My Project.

Option	Description	Action (s)		
Clone tree	Duplicate a tree and add it to My project.	Right-click any node except My Project.		
Apply Auto Layout	Apply an automatic layout to the nodes in the tree.	Right-click any node.		
Snap to Grid	Snap a node to align it with the grid. See "Moving Nodes in the Tree and Using the Grid" on the next page.	Right-click any node.		
Show Grid	Turn the grid on or off in the Project window.	Right-click any node.		
Zoom In, Zoom Out, Reset Zoom	Zoom in and out of the Project window or reset the zoom to the default.	Right-click any node.		

# **Organizing the Windows on the Desktop**

The windows on the Thermo-Calc desktop can be fixed or free-floating. By default, the windows are fixed and open.



An exception is the Event Log window, which is set to auto-hide by default.

## **AUTO-HIDING A WINDOW**

When a window is fixed it can also be set to auto-hide. Auto-hiding means that the window is minimized when you select another window. The minimized window is then shown either along the bottom or side of Thermo-Calc. A free-floating window is shown on top of the other windows and can be moved outside the frame of Thermo-Calc.

## RESETTING THE DESKTOP LAYOUT

To return to the standard desktop layout, from the main menu, select **Windows** → **Reset Windows**.

### FLOATING, HIDING AND MINIMIZING WINDOWS IN THE GUI

For each window you can rearrange, resize, minimize and close it. At the top right corner are the buttons listed in the table.

Button(s)	Description
J D	Click to toggle on/off free-floating
<del>Р</del> –	Click to toggle on/off auto-hide.
_	Click to minimize an open window that has auto-hide turned on.
_	The window automatically minimizes if another window is selected.
×	Click to close the window. You can open the window again from the Window main menu.

# Moving Nodes in the Tree and Using the Grid

In the Project window, you can work with the activity nodes in these ways.

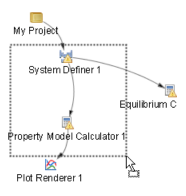
### SELECT AND MOVE INDIVIDUAL NODES

- Click an activity node to select it.
- To move activities, click and hold the activity node and move the cursor.

#### **SELECT SEVERAL NODES AT A TIME**

- To select several activity nodes, hold down <Ctrl> while clicking on each node you want to select.
- Click and hold the mouse button and draw a square around the activity nodes to select. Both the name and the icon that represents an activity must be within the

square.

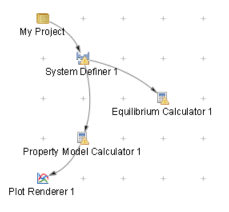


### SHOW GRID AND SNAP TO GRID

To show a grid of light-grey cross-hairs overlaid on a project, click **Show Grid** in the toolbar. Click the button again to turn the grid off. Or right-click in the **Project** window and select **Show Grid**.

Click **Snap to Grid** in the toolbar or right-click in the **Project** window and select **Snap to Grid** to automatically position each activity in the Project window at a cross-hairs point in the grid.

The Project window with the grid turned on and activity nodes moved from the default location. These nodes are snapped to the grid.



# **Zooming In and Out**

In the **Results** window (where the plots are):

- Use the mouse scroll wheel to zoom in and out.
- Click and drag the mouse from left to right to highlight an area of the plot to zoom into a specific point. Use the scroll wheel to continue zooming in or out.

### **Resetting a Plot to the Default View**

- For 2D plots, click and drag the mouse from right to left.
- For 2D and 3D plots, click **Perform Tree** to regenerate the plot in the **Results** window.
- "Rotating 3D Plots" on the next page

In the **Project** window (where the activity nodes are):

- Right-click and select Zoom In or Zoom Out.
- To go back to the default zoom, select Reset Zoom.

# **Node Status Markers**

Status markers indicate whether an activity is ready to be performed, is being performed, or has been performed. The markers are overlaid on the activity nodes in the **Project** window. The status markers are displayed on a **System Definer** node.

"Results Window Status Marker" on page 146

#### **STATUS MARKER DESCRIPTIONS**

Node Status Markers for the Project and Scheduler Windows

Window	Status marker	Description				
Project window	<b>`kal</b>	No status marker means the activity is ready to be performed (it has not been performed).				
	1	A yellow triangle means the activity cannot be run. The necessary configurations for the activity may not be set or other prerequisites are not met.				
	1	A green disc means the activity is performed.				
	0	A red circle with a dial (a clock face) means the activity is currently being performed.				

Window	Status marker	Description				
Scheduler window	8	No status marker means the activity (the job) is scheduled to be performed (the job has not started).				
		A yellow triangle means there an error happened during the activity job.				
	<u>}</u>	A green disc means the activity (the job) performed successfully.				
	·	A green right-facing arrow means the activity (the job) is being performed.				

# **Rotating 3D Plots**

Using an Equilibrium Calculator (property grid calculation) or Property Model Calculator (grid calculation) you can plot the diagram in 3D.

In the Results window after you create the 3D plot:

- Rotate the diagram automatically. Double-click in the **Results** window and the diagram rotates clockwise. Click once to stop the automatic rotation.
- Rotate the diagram manually. Click and hold the mouse and move it around the **Results** window.
- Use the mouse wheel to zoom in and out. To reset the plot to its default view click **Perform Tree** to regenerate the plot.
- "Equilibrium Calculator" on page 83
- "Property Model Calculator" on page 70
- "Plot Types" on page 126

# **Graphical Mode Projects**

These topics describes how you create, perform and link activities.

## In this section:

The Quick Start Wizard	50
Creating a Project with the Quick Start Wizard	. 50
Creating a Project from a Template	. 52
Cloning Activities and Trees	. 53
Performing Projects, Trees and Activities	55
About the Property Models in Thermo-Calc	56
Models	.56
Calculation Types	56
Property Model Development Framework	. 57
Examples and Video Tutorials	. 57
Using the Scheduler	58
The Event Log	. 59

# The Quick Start Wizard

Even before you understand how to work with Thermo-Calc projects, the Quick Start wizard helps you set up a project and run calculations in Graphical Mode. You can set up, calculate and visualize any of the following:

- Single point equilibrium
- Property diagram
- Phase diagram
- Scheil solidification simulation

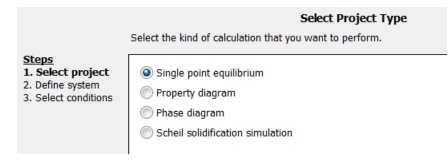
# Creating a Project with the Quick Start Wizard

To set up, perform and visualize a calculation using the Quick Start wizard:

1. In the **Configuration** window click **Quick Start**. If you cannot see the **Quick Start** button, first click **My Project** in the **Project** window.

File Tools Window Help	
□ □ □ □ ▲ Ď New Open Save Switch to Console Mode	Configuration
Project	
My Project	Wizards Quick Start

 In the Select Project Type window click to select the type of calculation to perform. Single point equilibrium, Property diagram, Phase diagram or Scheil solidification simulation. Click Next.



3.	In the	Define S	ystems	window,	choose	an	available	Database.
----	--------	----------	--------	---------	--------	----	-----------	-----------

🛕 Quick Start		$\times$
<u>Steps</u>	Define System You can either define the system by selecting elements from a database with thermodynamic information and/or kinetic data or by loading a pre-existing file that defines a material.	
<ol> <li>Select project</li> <li>Define system</li> <li>Select conditions</li> </ol>	Databases FEDEMO: Iron Demo Database v2.0 v	
	Element selection     C     C     C     Fe     Mn     Ni	
	O Material file	

- 4. Click to select Elements check boxes to include as system components. To add elements from an existing material file, click the Material file button to enter the file path to a material file or click the Select Material File button (to the right of the field) to navigate to a file on the computer.
- 5. Click Next.

The **Select Conditions** window has different options depending on the Project type and the elements selected.

In this example figure, using a **Single Point Equilibrium**, choose the **Composition unit** and under **Condition Definitions** specify the values of the state variables in the system.

<u>Steps</u>	Edit the conditions of compositions.		t Conditions Indent compositions can be edited by first clearing one of the other
<ol> <li>Select project</li> <li>Define system</li> <li>Select conditions</li> </ol>	Composition unit		
	Condition Definit	ions	
	Temperature	Kelvin 🔻	1000.0
	Composition	С	99.99
	Composition	Fe	0.01

For a **Property diagram** or **Phase diagram**, specify the **Axis Definitions** of the stepping or mapping operation. Choose minimum and maximum values for the variable(s) (select **Linear – min no. of steps**, **Linear – max step size** or **Logarithmic – scale factors**) and whether it is **Normal** or **Separate phases**.

For a **Scheil solidification simulation**, and where applicable, select **Fast diffuser** to account for back-diffusion of any fast-diffusing elements.

- 6. Click Finish.
- 7. Click to expand the **Event Log** window to see information about the progress of the calculation.

Event Log	
12:12:02,234 INFO	Global test at 5.20000E+02
12:12:02,249 INFO	Terminating at 500.000
12:12:02,252 INFO	*** Buffer saved on file: C:\Windows\TEMP\TH35C8~1\STATE_~1.POL
12:41:16,752 INFO	Opened project file: Ex_05_Fe-C_stable_and_metastable_phase_diagram.tcu
12:42:17,239 INFO	Opened project file: Ex_03_Fe-C_phase_diagram.tcu
14:11:35,181 INFO	Define component C
14:11:35,181 INFO	Define component Fe
14:11:35,211 INFO	Performing general equilibrium calculation

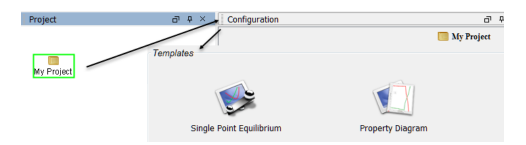
8. The diagram is plotted in a **Plot Renderer** tab in the **Results** window. If you have calculated a single-point equilibrium, then a **Table Renderer** tab shows information about the equilibrium.

Results				ē
Table Renderer 1				
System				
Moles	1.00000			
Mass	12.01194	[g]		
Temperature	1000.00000	[K]		
Total Gibbs Energy	-12659.53136	[J]		
Enthalpy	11784.43824	[J]		
Volume	5.39784E-6	[m3]		
Component	Mole Fraction	Mass Fraction	Activity	Potential
Fe	0.00002	0.00010	0.00619	-42278.5054
С	0.99998	0.99990	0.21816	-12658.8942
Stable Phases				
	Moles	Mass	Volume Fraction	
BCC_A2#1	0.00002	0.00120	0.00003	Composition

9. To save the project, click **Save** on the main toolbar. To save a diagram or table, rightclick the diagram or table and select **Save As**.

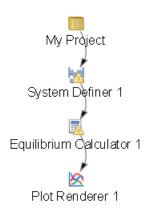
# **Creating a Project from a Template**

Use a *Template* to create all the activities for a certain type of calculation. In the **Project** window click the **My Project** node to display the templates in the **Configuration** window.

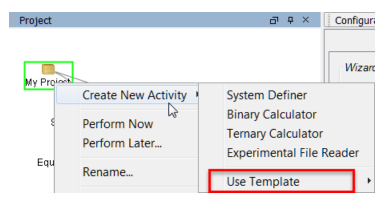


When you click a template, an activity tree is added to the Project window. You can then configure and perform the activities or the project.

For example, if you click **Property Diagram**, a tree is added to the **My Project** node with **System Definer**, **Equilibrium Calculator** and **Plot Renderer** nodes.



You can also select templates from the Project window. Right-click the **My Project** node and choose templates from the **Create New Activity**  $\rightarrow$  **Use Template** submenu.



# **Cloning Activities and Trees**

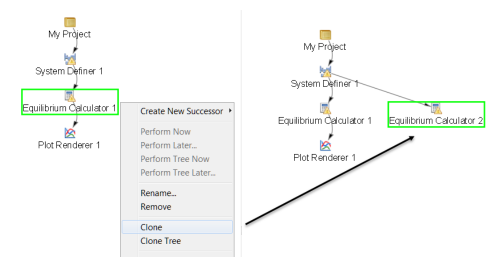
As described in <u>"Creating Activities and Successors" on page 32</u>, you can clone a single activity or the selected activity and all the activities that come after it (successors). The predecessor of the selected activity also is the predecessor of the clone.

Cloned activities are configured exactly like the activities that they were cloned from, and any results of calculations or plots are also cloned. To clone a selected activity, right-click the activity and select **Clone**.

For example, if you clone the Equilibrium Calculator in this project the result is an **Equilibrium Calculator 2** created as a successor to **System Definer 1**.

The following applies to clones.

- The new Equilibrium Calculator has the same settings for conditions, functions and options as its Equilibrium Calculator 1 clone.
- If the Equilibrium Calculator is already performed, then the calculation results are cloned.
- If the activity is a Plot Renderer or a Table Renderer, then the plot or table is cloned.
- If you want to clone all the successor activities that follow a selected activity, then right-click and select **Clone Tree** instead.



## CLONING AN EQUILIBRIUM CALCULATOR ACTIVITY NODE

### EQUILIBRIUM CALCULATOR CLONES AND SUCCESSORS

If you create an Equilibrium Calculator as a successor to another Equilibrium Calculator, then the successor inherits all the settings for conditions, functions and options from its predecessor. In this respect, it is similar to a clone.

The Equilibrium Calculator successor is different from a clone in that it does not inherit any calculated results and it is a successor to the Equilibrium Calculator instead of a successor to its predecessor.

The calculated result from an Equilibrium Calculator that is the predecessor to another Equilibrium Calculator is the starting value for the latter's calculation. The clone of an Equilibrium Calculator on the other hand, does not receive any data as input from the Equilibrium Calculator that it is cloned from.

# **Performing Projects, Trees and Activities**

You can trigger the performance of an activity (also called a *job*) and all the activities below it in the same tree in either the **Project** or **Configuration** window. In the Project window you can also perform a single activity without performing its successors or perform the whole project (all the activity trees).

In the **Configuration** window, click **Perform Tree** to perform the currently selected activity and all the activities below it. If there are no activities below the selected activity, then the button says **Perform** instead.



8

All activities that must be completed as prerequisites are automatically performed first. For example, if you Perform Tree for a Plot Renderer, then the predecessors, Equilibrium Calculator and System Definer, are automatically performed before the Plot Renderer.

In the Project window, you can do any of the following:

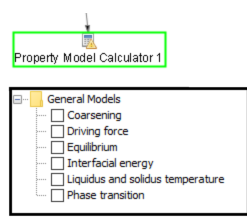
- Perform an activity: Right-click the activity and select **Perform Now**, or **Perform Later** and schedule a time to perform the activity
- Perform an activity and all activities below it in the tree: Right-click any node in the tree and select **Perform Tree Now**. Or select **Perform Tree Later** and schedule a time for all activities in the tree to be performed
- Perform a project: Right-click **My Project** and select **Perform Now** or **Perform Later** and schedule a time to perform all project activities in the tree



If there is an error during the performance of an activity, icons display status markers, which are described in <u>"Node Status Markers" on page 47</u>.

# **About the Property Models in Thermo-Calc**

## **Models**



There are these general models available when using the <u>"Property Model Calculator" on</u> page 70:

- **Coarsening**: Calculates the coarsening rate coefficient K (m<sup>3</sup>/s) of a spherical precipitate phase in a matrix phase.
- Driving force: Calculates the thermodynamic driving force for a phase.
- **Equilibrium**: Calculates the equilibrium for the given conditions. Optionally define additional Function definitions.
- Interfacial energy: Estimates the interfacial energy between a matrix phase and a precipitate phase using thermodynamic data from a CALPHAD database.
- Liquidus and solidus temperature: Use this to complete a common calculation. For example, with this model one can easily use uncertainty calculations, varying one or more conditions, and see how that affects the liquidus and solidus temperatures.
- **Phase transition**: Calculates the point when a new phase may form by varying set conditions. The model is useful to determine melting temperature, boiling temperature or solubility limits. It returns the phase transformation temperature, or composition, depending on the varied condition.

You can evaluate models simultaneously over a range of compositions and cross plot the results, then conduct an uncertainty analysis and plot the results as either a histogram or as a probability plot.

## **Calculation Types**

The Property Model Calculator includes these calculation types:

- Single: Calculates a single point.
- One axis: Varies a quantity on the X-axis.
- **Grid**: Evaluates two axis variables of the selected quantities in the specified range and number of steps. You can then select to create heat map, contour and 3D plots on the Plot Renderer.

"Plot Types" on page 126

- Min/Max: Evaluates the property model(s) for all variations of the selected quantities at the given limits. The Mean field is as defined under Condition Definitions for the respective quantity. The total minimum and maximum of the model(s) results are shown in the Event log.
- Uncertainty calculation: Evaluates the property model(s) where the values of the quantities are sampled from Gaussian distributions. The Mean field is as defined under Condition Definitions for the respective quantity. The result is visualized as a histogram or normal probability plot by adding a Plot Render activity.

"About the Uncertainty Calculations" on page 75 and "Plot Types" on page 126

### **Property Model Development Framework**

With the Property Model Calculator you can now develop your own models and integrate them into Thermo-Calc. In other words, you can customise the software to meet your modelling needs.

Models are developed using the programming language Python, and model development is possible without any additional installation using a simple editor, such as Notepad++. You can also use a Python Integrated Development Environment (IDE) to access advanced features such as debugging of the property models and autocompletion.

Models are automatically encrypted, for safe and secure file sharing.

Introduction to the Property Model Development Framework" on page 2

### **Examples and Video Tutorials**

#### WEBSITE

For an introduction to the Property Model Calculator, go to the video tutorial page on our <u>website</u> and under **Release and New Products**, click the arrow next to **Property Model Calculator**.

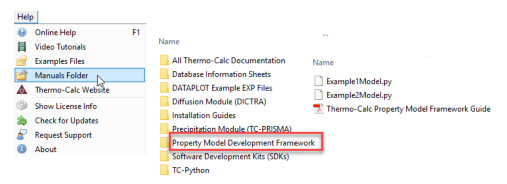
Then under **Examples - Graphical Mode**, click the arrows to open *T\_10*, *T\_11* and *T\_12*, which all use the Property Model Calculator.

#### ΥουΤυβε

You can also go to <u>YouTube</u> channel to review the available training videos. Subscribe to the channel to be updated when new videos are added. Also see the <u>video introduction</u> to the Property Model Calculator.

#### **PROPERTY MODEL DEVELOPMENT FRAMEWORK PDF AND EXAMPLES**

There is a PDF guide and tutorial examples included with your installation. In Thermo-Calc from the menu, select Help  $\rightarrow$  Manuals Folder. Then double-click to open the Property Model Development Framework folder. This guide is also available on our website.



# **Using the Scheduler**

The **Scheduler** window has activity information about current or scheduled jobs. A *job* is any activity you have selected to be performed. The Scheduler also indicates if there is an error during a job and when a job has stopped. If you have set an activity, tree or project to be performed at a later time then this is also listed.

#### ABOUT THE SCHEDULER JOBS

- Whenever you trigger the performance of a whole project, tree or individual activity, all the activities to be performed are shown under a job heading.
- When all the activities belonging to a job are completed the job and its activities disappear from the Scheduler window.
- If there is an error during the performance of an activity, icons for that job and activity remain in the Scheduler window until removed.
- <u>"Node Status Markers" on page 47</u>

Scheduler	L	×
Scheduled Jobs		
🚊 🕕 🕕 Job no 1		
Equilibrium Calculator 3 (127.0.0.1)		
i≟		
🖃 🥅 Stable and metastable FE-C phase diagram		
System Definer 1		
System Definer 2		
Equilibrium Calculator 2		
Equilibrium Calculator 1 (127.0.0.1)		

### MANAGING THE SCHEDULE

In the Scheduler window, you can cancel scheduled jobs, remove errors and show information about errors.

- To cancel all jobs: Right-click the **Scheduled Jobs** label or the cogwheel icon, and select **Cancel All Jobs**
- To remove (clear) all failed activities and jobs: Right-click the **Scheduled Jobs** label or the cogwheel icon, and select **Clear All Errors**
- To cancel a specific job: Right-click the job label (for example, **Job no 1**) and select **Cancel Job**
- To remove (clear) a specific failed job: Right-click the job label (for example, Job no 1) and select Cancel Job
- To open a window with error details: Right-click the label for the failed job (for example, **Job no 1**), and select **Show Error Log**

# **The Event Log**

The **Event Log** window is closed by default. Information about what Thermo-Calc is processing or doing is in blue text and error or warning messages are in red text.



```
EventLog 
V

14:50:34,751 INFO Cancelling activity: Equilibrium Calculator 1

14:50:36,266 WARN The subprocess completed with status code 2: the execution was
cancelled by the user

14:50:36,376 ERROR Gobbled: 14:50:36,376 ERROR Error when executing the activity:
Error when sending a remote event: Software caused connection abort: socket w
rite error

14:50:36,594 WARN The subprocess completed with status code 2: the exec
ution was cancelled by the user

15:36:25,495 INFO Activity was renamed to Appended System 1
```

- To open the Event Log window: Click the Event Log window once to expand it or select it from the Window → Event Log menu. Double-clicking the window maximizes it
- To specify the level of detail in the Event Log: From the main menu select Tools
   →Options. In the General tab set the Log level slide bar to anything between Debug (most detailed) and Error (least detailed)

# **Graphical Mode Activities**

These topics describe the activity types and has instructions for procedures tied to specific activities.

### In this section:

System Definer	62
Property Model Calculator	70
Property Model Calculator Workflow	
Equilibrium Calculator	
Diffusion Calculator	92
Precipitation Calculator	
Plot Renderer	
Plot Renderer for the Add-On Modules	
Plot Toolbar	
Table Renderer	146
Experimental File Reader	
Binary Calculator	150
Ternary Calculator	152
Scheil Calculator	

# **System Definer**

In a System Definer activity, you select the database to use to retrieve thermodynamic data and define which elements the system has as components. You can also select the species to include as well as change the reference temperature and pressure for your components.

The Configuration window has several tabs to set such things as:

- Elements
- Species
- Phases and Phase Constitution
- Components

There are also two tabs to list **Data Sources** and a **Description** for the selected database.

"Creating Activities and Successors" on page 32

### Configuration Settings

The System Definer Configuration window has many settings available, which are briefly described below.

igura	tion																		
										Systen	n Defi	ner: F	erritic	steel					
base	-										_								
O F	EDEMO	): Iron	n Demo I	Databa	se v2.(	0					$\sim$		Packa	ge:		•			$\sim$
ents	Specie	es Ph	nases ar	nd Pha	se Con	stitutior	n Con	ponen	ts Da	ta Sour	ces [	Descrip	tion						
								Г	Perio	dic Tab	le	Alph	abetic	List					
									ZE	VA							Â	Material	
																		Material na Ferritic	ime:
н	_									-		_			_	_	He	- Cinde	
Li	Be											В	С	Ν	0	F	Ne	Amount	Mass percent $\smallsetminus$
Na	Mg											AI	Si	Ρ	s	CI	Ar	Fe	80.5200000
к	Ca	Sc	ті	v	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr	Gr	17.0
Rb	Sr	Y	Zr	Nb	Мо	Тс	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Те	1	Xe	Ni	1.4
				-		_									-		_		
Cs	Ba	•	Hf	Та	W	Re	Os	lr	Pt	Au	Hg	TI	Pb	Bi	Po	At	Rn	с	0.08
Fr	Ra	**	Rf	Db	Sg	Bh	Hs	Mt	Ds	Rg	Cn	Uut	FI	Uup	Lv	Uus	Uuo	Mn	1.0
• 1 201	thanide	sories	La	Ce	Pr	Nd	Pm	Sm	Eu	Gd	ТЬ	Dy	Но	Er	Tm	Yb	Lu		ad material
Lan	unanioe :	selles	La	Ce		ING	-m		Eu	Ga	ID	Uy	но	Er	Im	TD	LU		e material as
** Ac	ctinide s	eries	Ac	Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr 🗸	Sav	e material dS
_																			
				😥 He	lp			<	Add P	redece	ssor	Pe	rform T	ree	Cre	ate Ne	w Succes	sor >	

CHOOSE DATABASE(S) OR A DATABASE PACKAGE

```
        Q = 102FE-5 Steels/Fe-Aloys v9.0
        V

        Package:
        Steels and Fe-aloys (TCFE9, MOBFE-6)
        V
```

At the top of the **Configuration** window you can choose to add one or more databases by clicking the **Add a database** button. You can also choose a predefined or custom database package from the **Package** list. Depending on your license, there are predefined database packages available to select.

You can also click **Remove this database** a button as required.

"Global Settings: System Definer" on page 242

If you have chosen a database package and then remove or change a database, a message displays *Selected databases and selected package may differ*. This means that the original package you selected, which added preset databases, does not match the databases you are using for the project. For information about how to clear the message, see <u>"Global Settings: System Definer" on</u> <u>page 242</u>.

There are several settings tabs available for the System Definer activity.

Elements Species Phases and Phase Constitution Components Data Sources Description

The following example uses the FEDEMO and MFEDEMO databases and Graphical Mode example T\_12.

Databases	
FEDEMO: Iron Demo Database v2.0	~
MFEDEMO: Fe-Alloys Mobility demo database v2.0	$\sim$

#### **ELEMENTS**

On the **Elements** tab, and based on the databases or database package selected, you can select and review elements using a **Periodic Table** or **Alphabetic List** format. Unavailable elements are greyed out on the Periodic Table view.

#### **PERIODIC TABLE**

In this example, using the demonstration databases FEDEMO and MFEDEMO, there are only

									ZE	VA							
н									J	-							He
Li	Be											в	С	Ν	0	F	Ne
Na	Mg											AI	Si	Р	s	CI	Ar
к	Са	Sc	ті	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr
Rb	Sr	Y	Zr	Nb	Мо	Тс	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Te	1	Xe
Cs	Ba	*	Hf	Та	W	Re	Os	Ir	Pt	Au	Hg	ті	Pb	Bi	Po	At	Rn
Fr	Ra	**	Rf	Db	Sg	Bh	Hs	Mt	Ds	Rg	Cn	Uut	FI	Uup	Lv	Uus	Uuo
* Lant	thanide :	series	La	Ce	Pr	Nd	Pm	Sm	Eu	Gd	Tb	Dy	Но	Er	Tm	Yb	Lu
Ad	ctinide s	eries	Ac	Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr

a select number of elements to choose from.

### **ALPHABETIC LIST**

Element	FEDEMO	MFEDEMO
1-	1	<b>v</b>
C Cr	1	<b>√</b>
	1	<b>√</b>
Fe	1	$\checkmark$
Mn	1	$\checkmark$
Ni	1	$\checkmark$
VA	1	$\checkmark$

### **MATERIAL COMPOSITION**

On the **Periodic Table** page, you can also define the material composition:

Material	
Material na	ime:
Ferritic	
Amount	Mass percent 💌
Fe	80.52
Ni	1.4
с	0.08
Mn	1.0
Cr	17.0

At the bottom of the window, click **Load material** to open a material file. The default folder opens to the **Materials** folder included with your installation.

"About the Material Files" on page 18 and "Examples, Manuals and Materials Folders" on page 12

Amount Mass percent V Fe 99.99 C 0.01 C 0.01 C 0.01 C 0.01 Cook in: Materials C Cast Iron Aluminum Alloys C Cast Iron Aluminum Alloys C Cast Iron My Materials Nickel Alloys S Solder Alloys S Solder Alloys S Steel This PC Network File name: Open Cancel	Material Material nam		
C 0.01	Amount	Mass percent $$	
▲ Open   Look in:   Materials   ▲ Aluminum Alloys   ■ Aluminum Alloys   ■ Cast Iron   ■ My Materials   ■ Nickel Alloys   ■ Solder Alloys   ■ Steel   ■ Titanium Alloys   ■ Steel   ■ Titanium Alloys   ■ Steel   ■ Titanium Alloys	Fe	99.99	
Look in: Materials Recent Items Desktop Documents This PC Haterials Aluminum Alloys Cast Iron My Materials Nickel Alloys Solder Alloys Solder Alloys Titanium Alloys File name: Open	с	0.01	
Recent Items   Desktop   Desktop   Documents   This PC   Vickel Alloys   Solder Alloys   Steel   Titanium Alloys   Solder Alloys   Steel   Titanium Alloys		🛦 Open	×
Recent Items   Desktop   Desktop   Documents   This PC   File name:		Look in:	] Materials 🗸 🤌 🗊 🖬 🗸
Desktop   Desktop   Solder Alloys   Steel   Titanium Alloys     This PC     File name:     Open		Recent Items	Cast Iron My Materials
Documents  This PC  File name:  Open		Desktop	Solder Alloys
File name: Open			Thankin Anoys
Natural .		This PC	
		Network	
	-		
Load material			

You can also click Save material as to save the material you defined, for example Ferritic.

"Global Settings: System Definer" on page 242



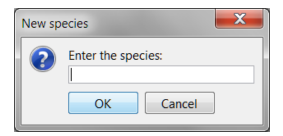
If you Load a material file and then change the composition, a message displays *The composition has changed*. This means that the original values/composition does not match the values you are using for the project. The message is cleared by either reloading the material file or editing the composition field to the original value.

### **S**PECIES

Elemer	nts Species Ph	ases and P	hase Consti
Specie	s Stoichiometry	FEDEMO	MFEDEMO
1-	-1	1	1
С	С	1	$\checkmark$
Cr	Cr	1	1
Fe	Fe	1	$\checkmark$
Mn	Mn	1	$\checkmark$
Ni	Ni	1	$\checkmark$
VA	VA	1	$\checkmark$
C2	C <sub>2</sub>	$\checkmark$	
C3	C <sub>3</sub>	$\checkmark$	
C4	C <sub>3</sub> C <sub>4</sub>	$\checkmark$	
C5	C <sub>5</sub>	$\checkmark$	
C60	C <sub>60</sub>	1	

On the **Species** tab for each database you can select and edit the **Species**.

At the bottom of the window, click **Add new species** and enter the name of species in the field.



PHASES AND PHASE CONSTITUTION

Status	Name	FEDEMO	MFEDEMO
Entered 🔹	BCC_A2	1	1
Entered 🔹	CEMENTITE	1	
Entered 🔹	CHI_A12	1	
Entered 🔹	DIAMOND_FCC	1	
Entered 🔹	FCC_A1	1	1
Entered 🔹	FCC_A1#2	1	1
Entered 🔹	GAS	1	
Entered 🔹	GRAPHITE	1	
Entered 🔹	HCP_A3	1	
Entered 🔹	KSI_CARBIDE	1	
Entered 🔹	LAVES_PHASE	1	
Entered 🔹	LIQUID	1	
Entered 🔹	M23C6	1	
Entered 🔹	M3C2	1	
Entered 🔹	M5C2	1	
Entered 🔹	M7C3	1	
Entered <	SIGMA	1	

At the top of the **Phases and Phase Constitution** tab for each database, you can review and edit the **Status (Entered** or **Dormant)** for the selected**Phases**. Select the **Check/uncheck all** check box to toggle between the options.

To review the **Phase constitution** for a specific phase, click it in the **Phase** list (for example **BCC\_A2**) and its details are displayed at the bottom of the window. Click **Edit** then click an element to select or reject a constituent. The last remaining constituent on a sublattice cannot be rejected.

FEDEMO MFEDEMO				
Phase BCC A2				
Phase constitution for current system Edit				
( Cr Fe Mn Ni ) <sub>1.0</sub> ( C VA ) <sub>3.0</sub>				
Phase description in database				
(Cr,Fe,Mn,Ni) <sub>1</sub> (C,VA) <sub>3</sub>				

### **COMPONENTS**

The Components tab is where you can review and edit the Component, Reference phase,

**Reference temperature**, and **Reference pressure**. Click to toggle the **SER** (Stable Element Reference) check box on and off. Then choose a **Reference phase** from the list and edit accordingly.

Elements Species Phases and Phase C	Constitution Components Data Sources	Description
Component Reference phase	Reference temperature Kelvin	Reference pressure Pascal
Fe	Current temperature 25.0	100000.0
Cr  Stable Element Refe	crence Current temperature 25.0	100000.0
Ni 🔹 🔽 SER	▼ Current temperature 25.0	100000.0
C   SER	▼ Current temperature 25.0	100000.0
Mn 🔻 🗸 SER	▼ Current temperature 25.0	100000.0

### **DATA SOURCES**

Data sources are not available until the activity has been performed. Click **Perform Tree** to display available data sources.

Elements	Species	Phases and Pha	se Constitution	Components	Data Sources	Description
FEDEMO	MFEDEM	0				
'B. mat	Uhrenius ( ter, Vol. 12	SGTE Data for Pu 1993-1994), Inte 2, pp. 121-127; N 1, Swerea KIMAB,	rnational journa Aolar volumes'	of refractory		d

### DESCRIPTION

The **Description** tab displays a description of the selected database(s). In this example, **MFEDEMO**.

Elements	Species	Phases and Phase Constitution	Components	Data Sources	Description
FEDEMO	MFEDEM	0			
	*** Ov	MFEDEMO - Database for Demo [Version 2.0: released March 2 vnship/Copyright: 2017 @ Thermo All rights reserved.	017]	********	^

## Creating a System Definer Activity

This topic describes how to define a system; specifically how to select one or more databases and define the elements in the system.

- "About the Database Files" on page 234
- "Global Settings: System Definer" on page 242
  - 1. Right-click **My Project** and from the menu select **Create New Activity**→ **System Definer**.
  - 2. In the **Project** window, click the <u>"System Definer" on page 62</u> activity node.
  - 3. In the **Configuration** window, from the **Databases** menu choose **Database(s)** or a **Database Package**.

### **OPTIONS:**

- Select <User database> to define your own.
- You can choose to add one or more databases by clicking the Add a database
   <sup>(3)</sup> button.
- From the **Package** list, choose a predefined database package to add multiple databases at a time. Depending on your license, there are predefined database packages available to select.
- 4. Select the **Elements** tab to choose the elements in the **Periodic Table** or use the **Alphabetic List** to choose elements for the system.
- 5. As required click the tabs to make edits: **Species**, **Phases and Phase Constitution**, and **Components**. The **Data Sources** and **Description** tabs provide information based on the database.
- 6. Click **Perform** to start the System Definer activity immediately and retrieve thermodynamic data about the system. Click the **Event Log** window to watch the progress of the job.

# **Property Model Calculator**

### About the Property Models in Thermo-Calc" on page 56

A Property Model Calculator allows you to predict and optimize material properties. A set of general models is available to use:

- "Phase Transition Model" on page 80
- "Driving Force Model" on page 80
- "Interfacial Energy Model" on page 81

- <u>"Coarsening Model" on page 82</u>
- "Equilibrium Model" on page 83
- Experienced users can also create custom property models. See the <u>"Introduction to the Property Model Development Framework" on page 2</u> for details.

You can evaluate models simultaneously over a range of compositions and cross plot the results. You can then conduct an uncertainty analysis and plot the results as either a histogram or as a probability plot.

## **Property Model Calculator Workflow**

When a project is first being designed, it is recommended you test it by first performing a **Single** calculation type. This verifies that the configuration of the model is valid, otherwise you will get a *Not a Number* result in <u>"Event Log Window" on page 166</u> indicating the calculation did not work correctly.

Once the Single calculation type performs successfully, you can experiment with other calculation types such as **Grid**, **Min/Max** or **Uncertainty**. Best practice for both Grid and Uncertainty is to start with a small number of steps (Grid) or samples (Uncertainty) to confirm that it works before increasing the number of steps or samples.

For a Grid calculation, you can additionally plot a *Heat map, Contour* or *3D* diagram. This setting is done on the "Plot Renderer" on page 122.

In general, the workflow for the different models differs. A recommended start point is to do a Property Diagram calculation to determine what phases you get and where the transitions are before setting up the Property Model Calculator.

- <u>"The Quick Start Wizard" on page 50 and "Creating a Project from a Template" on page 52</u>
- For examples using the Property Model Calculator, see examples T\_10, T\_11, and T\_12 in the *Graphical Mode Examples Guide*.



The **Configuration** window for a Property Model Calculator using the *Phase transition* model is shown and described in <u>"Configuration Settings" on page 73</u>.

## Defining the Property Model Calculator

1. From the left side of the **Configuration** window, click to select one or more of the check boxes under the **General Models** folder.

A tab with the same name as the property model opens in the right side of the window under *Condition Definitions*. When you select more than one model, a matching tab is added in the same order that the check box is selected.

Configuration				
Property Model Calculator 1				
General Models	Composition unit Mass percent v			
Driving force     Guillorium     Interfacial energy	Temperature Celsius 🗸			
Liquidus and solidus temperature	Composition Fe Composition Ni	80.52 1.4		
	Composition C	0.08		
	Composition Mn	1.0		
	Composition Cr	17.0		
	Phase transition Driving force			
	Configuration Description			
	Matrix phase: Any phase	~		
	Precipitate phase: BCC_A2	~		
<				
(i) Help (	Add Predecessor Perform Tree	Create New Successor >		
<		>		

- 2. Enter the settings described in <u>"Configuration Settings" on the next page</u> and for each *Calculation Type* (Single, Grid, Min/Max, and Uncertainty).
- 3. For each model on the **Configuration** tab, select or enter settings as required. Click the **Description** tab for more information about the model and refer to the individual sections for details.
  - "Phase Transition Model" on page 80
  - "Driving Force Model" on page 80
  - "Interfacial Energy Model" on page 81
  - "Coarsening Model" on page 82
  - "Equilibrium Model" on page 83

Configuration	<b>Settings</b>
---------------	-----------------

Configuration		
	Property Model Calculator 1	1
General Models Coarsening Coarsening General Models General	Composition unit Mass percent, Condition Definitions Temperature Celsius Composition Fe Composition Ni Composition C Composition Mn Composition Cr	700.0       80.52       1.4       0.08       1.0       17.0
	Phase transition Configuration Description	
	Matrix phase: Any phase Phase to form: SIGMA Condition to vary: Temperature Search direction: Negative	> > >
	Calculation Type O Single One axis	⊖ Grid ⊖ Min/Max
	Sampling of Data from Gaussian Distri	ributions
		Mean ∆ Min/Max
		700.0 0.0
	Composition Ni	1.4 0.4
	Composition C	0.08 0.02
	Composition Mn	1.0 0.3
	Composition Cr	17.0 2.0
	Sampling Parameters	
	Truncate distribution at	3 🜩 standard deviations
	Total number of samples	10 🛓
l l l l l l l l l l l l l l l l l l l	Add Predecessor Perform Tree	e Create New Successor >

#### **Composition unit**

Choose a unit: Mass percent, Mole percent, Mass fraction or Mole fraction.

#### **Condition Definitions**

Choose a unit (Kelvin, Celsius or Fahrenheit) and then enter a value for the Temperature and Composition.

The values entered in Condition Definitions are used as the start values for the different calculation types.

#### Define the property model

For each model chosen, select or enter settings as required. Click the **Description** tab for more information about the model and refer to the individual sections for details.

- "Phase Transition Model" on page 80
- "Driving Force Model" on page 80
- "Interfacial Energy Model" on page 81
- "Coarsening Model" on page 82
- "Equilibrium Model" on page 83

#### **Calculation Type**

Choose a calculation type: Single (no axes), One axis, Grid, Min/Max or Uncertainty.

- Select Single to calculate a single point.
- Select One axis to vary a quantity on the X-axis.
- Select a **Grid** to evaluate two axis variables of the selected quantities in the specified range and number of steps.
- The **Min/Max** calculation evaluates the property model(s) for all variations of the selected quantities at the given limits. The **Mean** field is as defined under Condition Definitions for the respective quantity. The total minimum and maximum of the model(s) results are shown in the Event log.
- The **Uncertainty** calculation evaluates the property model(s) where the values of the quantities are sampled from Gaussian distributions. The **Mean** field is as defined under Condition Definitions for the respective quantity. The result is visualized as a histogram or normal probability plot by adding a Plot Render activity.

#### Single calculation

For the **Single** calculation type no further settings are required. The results from this calculation are displayed in the **Event Log**.

#### **Axis Definition**

For **One axis**, select a **Quantity** to vary along the X-axis, for example, **Temperature** then

### enter a Min, Max and Number of steps.

#### Grid Definitions

For the **Grid** calculation type, define the two axes variables using the fields and menus: **Quantity**, **Min**, **Max** and **Number of steps**. The number of steps along with the minimum and maximum values for the axes define a grid. For each grid point the selected models are evaluated.



For a **Grid**, on the Plot Renderer, you can choose to plot a *Heat map*, *Contour* or *3D* diagram.

### Min/Max Definitions

For the **Min/Max** calculation type, click to select the applicable check boxes under **Quantity to enable or disable an axis**. The **Mean** field is as defined under Condition Definitions for the respective quantity.

Enter numerical values in the  $\Delta$  **Min/Max** fields to evaluate the change to the quantity. The highest and lowest value of each selected model results are shown in the Event log.

### Sampling of Data from Gaussian Distributions

For the **Uncertainty** calculation type, click to select the applicable check boxes under **Quantity to vary this quantity**. The **Mean** field is for the distribution function as defined under Condition Definitions for the respective quantity.  $\Delta$ Min/Max is the 95% confidence interval.

#### **Sampling Parameters**

For the **Uncertainty** calculation type, choose a number of **Standard deviations** where you want to **Truncate the distribution**. Choose the **Total number of samples**.

"About the Uncertainty Calculations" below

## About the Uncertainty Calculations

The following example uses the <u>"Phase Transition Model" on page 80</u> and an **Uncertainty Calculation Type** to describe how the sampling of data is performed. The example <u>"T\_10:</u> <u>Phase Transition" on page 23</u> in the *Graphical Mode Examples Guide* predicts the transition temperature to the unwanted brittle Sigma phase and shows how the temperature is influenced by changes to a steel alloy's composition.

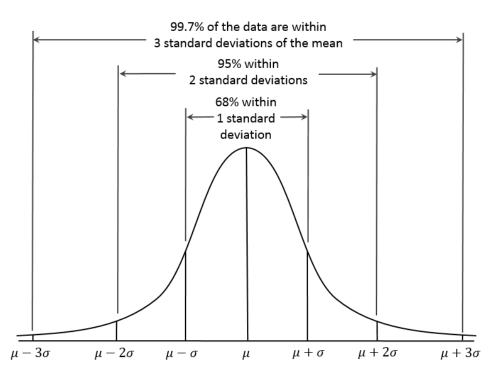
The **Uncertainty** calculation in Thermo-Calc uses the values (the mean) entered for quantities in the Condition Definitions section as the starting point to estimate, within a certain probability, the true variation of the quantity.

An example of a Gaussian distribution is shown next.

### **GAUSSIAN DISTRIBUTION, THE 3-SIGMA RULE**

- The *mean* is at the peak of the curve (labelled  $\mu$ ).
- The standard deviation ( $\sigma$ ) is a plus and minus range,  $\pm 1\sigma$ ,  $\pm 2\sigma$  or  $\pm 3\sigma$ , of the variance from the mean.

Image by Dan Kernler: https://en.wikipedia.org/wiki/Normal\_ distribution#/media/File:Empirical\_Rule.PNG.



As long as the number of events (the sample size) is very large, then the Gaussian distribution function can be used to describe physical events. A Gaussian distribution is assumed to describe how each quantity varies in an uncertainty calculation, as shown next.

#### **CONDITION DEFINITIONS AND SAMPLING SETTINGS**

In Thermo-Calc, the mean of the quantity's distribution is defined in the *Condition Definitions* section. The **Mean** is then used in the *Sampling of Data from Gaussian Distributions* section as the starting point for how much you want to deviate from this value.

The mean defined in Condition Definitions is used during the sampling.

Composition unit	Mass percent 💌			
Condition Definit	lions			
Temperature	Celsius	600.0		
Composition	Fe	80.52		
Composition	Ni	1.4		
Composition	с	0.08		
Composition	Mn	1.0		
Composition	Cr	17.0		
		Sampling of Data from	Gaussian Distri	butions
		Quantity	Mean	Δ Min/Max
		Temperature	600.0	0.0
		☑ Mass percent Ni	1.4	0.4
		☑ Mass percent C	0.08	0.02
		Mass percent Mn	1.0	0.3

The values entered in the  $\Delta$  **Min/Max** fields are the resulting width of the distribution corresponding to the 95% confidence interval (approximately  $\mu \pm 2\sigma$ ) shown in *Gaussian distribution, the 3-sigma rule*. This confidence interval means that approximately 95% of the sampled values fall within these limits, as shown next.

### SAMPLING PARAMETER SETTINGS

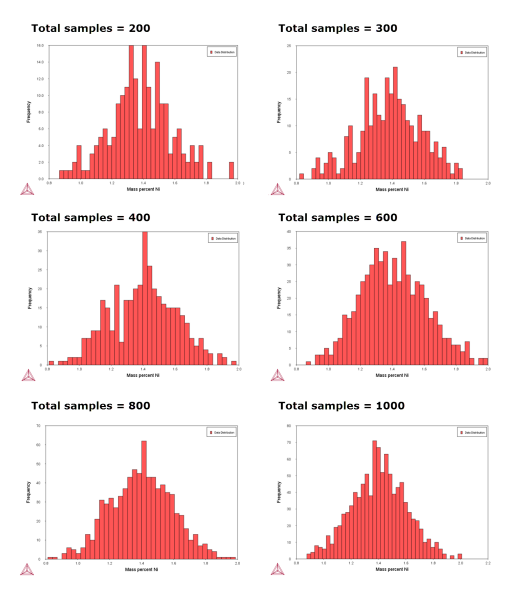
The default value for truncating the distribution is set at **3 standard deviations** in the *Sampling Parameters* section. This is also where you set the **Total number of samples**. The default is 200. In order to maintain a Gaussian distribution for the varied quantities, this number should not be too low.

Sampling Parameters	
Truncate distribution at	3 🔷 standard deviations
Total number of samples	200 🚔

You can experiment by varying the conditions and the number of samples to see the distribution results smooth out as is to be expected based on the Central Limit Theorem, as shown in the histogram plot. You can also plot the results as a normal probability.

#### **HISTOGRAM PLOT**

This series of histogram plots shows the uncertainty calculations sampling data from Gaussian distributions. On the **Plot Renderer** the variable is set to measure the **Composition** of **Ni**. It compares the Ni composition where the total sample size is increased. In the example, the initial sample size uses the default, 200, and then the sample size is increased to 300, 400, 600, 800, and 1000, showing how the plot follows the Gauss distribution.



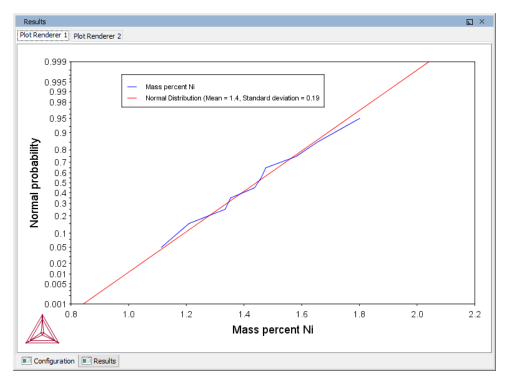
## NORMAL PROBABILITY PLOT

You can also plot the results as a Normal probability. Continuing with this example, on the

Property Model Calculator Configuration window the X-Axis variable is plotted as Composition and the Y-Axis variable is plotted as the Normal probability.

🖄 Property Mo	iel Calculator 1
Tie lines: (	Legend option: On V Legend style: None
X Axis	
Axis variable:	Mass percent Ni
Axis type:	Linear 🔻
Limits:	500.0 to 3000.0 step 250.0 V Automatic scaling
Y Axis	
Axis variable:	Normal probability
Axis type:	Probability 💌
Limits:	500.0 to 3000.0 step 250.0 V Automatic scaling
◎ ● 🔽	

This is a normal probability plot example of the uncertainty calculations sampling data from Gaussian distributions. It compares the Ni composition where the total sample size is 200. The sampled data (blue line) closely follows the ideal normal distribution (red line). Deviations are naturally larger at the tails ( $\approx 1\%$ Ni and  $\approx 2\%$ Ni) since the number of samples are fewer there compared to the center of the distribution.



# Phase Transition Model

By varying set conditions, the **Phase Transition** model calculates the point when a new phase may form. The model is useful to determine melting temperature, boiling temperature or solubility limits. It returns the phase transformation temperature, or composition, depending on the varied condition.



For an example see <u>"T\_10: Phase Transition" on page 23</u> in the *Graphical Mode Examples Guide*.

Define the:

- Matrix phase: The single phase region to start in. Select 'Any phase' to use the phases that take part of the initial equilibrium or choose a specific phase.
- **Phase to form:** For the new phase to be formed, select 'Any phase' or choose a specific phase.
- **Condition to vary:** Choose whether to vary a composition or temperature condition when calculating the phase transition.
- Search direction: For the composition or temperature condition being varied, choose the Search direction to be in a **Positive**, **Negative**, or **Positive and Negative** direction.
- Upper search limit and Lower search limit: These values also relate to the search direction, where an estimated change to the released condition implies where a new phase is expected.
- The start value (given in Condition Definitions) and phase transition point need to be within these limits, otherwise the model returns *Not a Number* as result. For example, a start temperature is 1100 K then the limits need to be outside

these, e.g. 500-1500.

# Driving Force Model

The **Driving Force** model calculates the thermodynamic driving force for a phase.

- Matrix phase: Choose a Matrix phase, the single phase region to start in. Select 'Any phase' to use the phases that take part of the initial equilibrium or choose a specific phase.
- Precipitate phase: Choose a Precipitate phase to calculate its driving force.

The calculation result is the driving force divided by RT.

For an example of a driving force model, see <u>"T\_12: Driving Force and</u> Interfacial Energy" on page 29 in the *Graphical Mode Examples Guide*.

# Interfacial Energy Model

Use the **Interfacial Energy** model to estimate the interfacial energy between a matrix phase and a precipitate phase using thermodynamic data from a CALPHAD database. The approximation model is based on Becker's bond energy approach and is also available in Precipitation Module (TC-PRISMA). In the actual calculation, the coefficients taken distinguish between only bcc and fcc structure for the (110) and (111) lattice plane, respectively.

For other planes and/or other phases, the estimate is approximate. In most situations, it is recommended to perform calibrations against some experimental data because the interfacial energy is dependent on so many factors that are ignored in the estimation, for example entropy, incoherency, orientation, and curvature, etc. Uncertainty factors must also be considered as thermodynamic data in CALPHAD databases are not always accurate.

Select the Matrix phase and Precipitate phase to return the interfacial energy [J/m<sup>2</sup>].

....

For an example, see <u>"T\_11: Coarsening and Interfacial Energy" on page 26</u> in the *Graphical Mode Examples Guide*.

# Estimation of Coherent Interfacial Energy

Interfacial energy is an important parameter used in precipitation simulations to calculate the rates of nucleation, growth/dissolution, and coarsening. The value of interfacial energy can vary dramatically (usually between 0.01 to 2.0 J/m<sup>2</sup>).

The extended Becker's model functions to estimate coherent interfacial energy by using thermodynamic data from existing CALPHAD thermodynamic databases:

$$\sigma_c = rac{n_s z_s}{N_A z_l} \Delta E_s$$

where  $\sigma_c$  is the coherent interfacial energy,  $n_s$  is the number of atoms per unit area at the interface,  $z_s$  is the number of cross bonds per atom at the interface,  $z_l$  is the coordination number of an atom within the bulk crystal lattice, and  $\Delta E_s$  is the energy of solution in a multicomponent system involving the two phases being considered [Becker, 1938]<sup>1</sup>.

<sup>&</sup>lt;sup>1</sup>. Becker, 1938. 'Die Keimbildung bei der Ausscheidung in metallischen Mischkristallen', Ann. Phys. 424, 128–140.

# **Coarsening Model**

The **Coarsening** model calculates the coarsening rate coefficient K (m<sup>3</sup>/s) of a spherical precipitate phase in a matrix phase.

- Matrix phase: Choose a Matrix phase.
- Precipitate phase: Choose a Precipitate phase to calculate its coarsening rate.

Returns the coarsening rate coefficient  $[m^3/s]$  for the precipitate phase and the interfacial energy  $[J/m^2]$  between matrix and the precipitate phase.



This model requires that the thermodynamic and kinetic databases used include descriptions of molar volume of the matrix and precipitates as well as mobilities for the matrix phase.



For an example of a coarsening model, see <u>"T\_11: Coarsening and Interfacial</u> <u>Energy" on page 26</u> in the *Graphical Mode Examples Guide*.

# Particle Coarsening

In the coarsening regime where the driving force is capillarity, the change of mean particle radius evolves according to the equation:

$$r^{-3} - r_0^{-3} = Kt$$

Following Morral and Purdy (1994)<sup>1.</sup>, the rate constant of precipitating  $\beta$  phase in a multicomponent  $\alpha$  phase is

$$K = rac{8 \sigma V_m^eta}{9} \Bigg[ igg[ c_i^eta - c_i^lpha igg]^T [M]^{-1} \Big[ c_k^eta - c_k^lpha \Big] \Bigg]^{-1}$$

The mobility matrix M is not the same as the mobilites that can be obtained directly from the Diffusion Module (DICTRA). The M matrix instead corresponds to the L" matrix as defined by Andersson and Ågren  $(1992)^2$ .

Using the u-fractions that is defined as

$$u_k = rac{x_k}{\sum\limits_{j \in S} x_j}$$

<sup>&</sup>lt;sup>1.</sup> Particle coarsening in binary and multicomponent alloys," *Scr. Metall. Mater.* 30, 905–908.

<sup>&</sup>lt;sup>2</sup>. Models for numerical treatment of multicomponent diffusion in simple phases. J. Appl. Phys. 72, 1350–1355.

where the summation only is performed over the substitutional elements give the final equation for the coarsening rate coefficient.

$$K=rac{8\sigma V_m^eta}{9}iggl[iggl[u_i^eta-u_i^lphaiggr]^T[L_{jk}^{\prime\primelpha}]^{-1}iggl[u_k^eta-u_k^lphaiggr]iggr]^{-1}$$

The interfacial energy  $\sigma$  is calculated using the extended Becker's model (that also is available as a separate Property Model Calculator model).

# Equilibrium Model

The Equilibrium model calculates the equilibrium for the given conditions. Optionally define additional **Function Definitions**. All quantities that normally can be obtained from an equilibrium calculation can also be obtained when plotting the results, as well as the defined functions.

# **Equilibrium Calculator**

An Equilibrium Calculator allows you to set the conditions for, and perform, a calculation. The Configuration window for an Equilibrium Calculator has these tabs:

- **Conditions:** Set the conditions for your calculation that define the stepping or mapping axis variables. This tab can be viewed in a simplified mode and in an advanced mode. See "Conditions Settings" on the next page.
- Functions: Define your own quantities and functions, which you then can calculate and plot. See "Calculating and Plotting Functions" on page 91.
- **Options:** Modify numerical settings that determine how equilibria are calculated, as well as how stepping and mapping calculations are performed. See <u>"Options</u> <u>Settings" on page 87</u>.
- The **Options** tab is where you can locally define specific Equilibrium Calculator settings. These settings can also be applied globally. To open the Options window: From the main menu select **Tools** → **Options** and click the **Activities** tab. Then click **Calculation**.

See <u>"Creating Activities and Successors" on page 32</u> for a list of possible successors and predecessors for this activity. An Equilibrium Calculator that is the successor of another Equilibrium Calculator inherits all the Configuration window settings of the predecessor.



For an example of how serially coupled Equilibrium Calculators can be used, see example T\_06 in the *Graphical Mode Examples Guide* 

Conditions Funct	ions Options			
Composition unit	Mass percent 🔻	•		
Condition Definit	ions			
Temperature	Celsius	▼ 1000.0		
Pressure	Pascal	▼ 100000.0		
System size	Mole	▼ 1.0		
Calculation Type	-	operty diagram	O Property	grid 💿 Phase diagram
Axis Definitions				
Quantity	Min	Max	Step division	Туре
Temperature	▼ 500.0	3000.0	50.0	Linear - min no. of steps 🔻
Temperature	▼ 500.0	3000.0	50.0	Linear - min no. of steps 🔻

## **Conditions Settings**



See examples <u>"T\_01: Calculating a Single-Point Equilibrium" on page 4</u>, <u>"T\_02:</u> <u>Stepping in Temperature in the Fe-C System" on page 6 and <u>"T\_05: Stable and</u> <u>the Metastable Fe-C Phase Diagrams" on page 13</u> in the *Graphical Mode Examples Guide*.</u>

Conditions and axis variables for the Equilibrium Calculator are set in the **Conditions** tab of the Configuration window. The tab can be viewed in two modes: **simplified** (the default) or **advanced**. The conditions to set depend on the type of calculation.

#### ABOUT ADVANCED MODE

Click **Switch to advanced mode** to add and remove conditions as well as set additional axis definitions. However, the number and types of conditions set must still conform to the Gibbs' phase rule. The **Number of missing conditions is** field, at the top of the tab shows how many conditions that you have left to set. If the number is negative, then that number of conditions need to be removed.

The following outlines what you can set on the **Conditions** tab in simplified mode.



Click Switch to advanced mode for more options.

#### **Composition unit**

Choose from Mass percent, Mole percent, Mass fraction or Mole fraction.

## **Condition Definitions**

Specify the values (and units) of the state variables: **Temperature**, **Pressure**, and **System** size.

## Calculation Type

Single equilibrium (no axes), Property diagram, Property grid or Phase diagram.



For the **Single** calculation type no further settings are required. The results from this calculation are displayed in the **Event Log**.

#### **Axis Definitions**

- Property diagram: Define the axis variable using these fields and menus: Quantity, Min, Max, Step division and Type.
- Property grid: Define the two axis variables using these fields and menus: Quantity, Min, Max and Number of steps.
- Phase diagram: Define the two axis variables using these fields and menus: Quantity, Min, Max, Step division and Type.
  - If the Equilibrium Calculator already has a Plot Renderer as a successor and you change the stepping/mapping quantities, then the quantities represented on the X- and Y-axis are automatically updated in the Plot Renderer.



For a **Property grid**, on the Plot Renderer, you can choose to plot a *Heat map*, *Contour* or *3D* diagram.

"Plot Types" on page 126

#### Number of steps (Property grid)

Set a fixed **Number of steps**. The number of steps along with the minimum and maximum values for the axes define a grid. For each grid point an equilibrium is calculated.

#### Type and Step division (Property diagram and Phase diagram)

Type options: Linear - min no. of steps, Linear - max step size and Logarithmic - scale factor.

- For Linear min no. of steps, the **Step division** value specifies a minimum number of steps that is calculated between the minimum and maximum value during the stepping/mapping operation.
- For Linear max step size, the **Step division** sets the maximum size of each step.For Logarithmic scale factor, **Step division** specifies the scale factor of a logarithmic function that distributes the steps from the initial equilibrium to the minimum and maximum values defined.

# Calculating with a Fixed Phase

This describes how to use the advanced mode to set a phase to be fixed at a certain amount. This allows you to, for example, find out at what temperature a material starts to melt. If you set the phase to be fixed to liquid phase at zero amount, and do not define the temperature state variable, then you can calculate at what temperature the material enters a state where the liquid phase is no longer zero (that is, when it starts to melt).

- For a fixed phase equilibrium calculation, it is recommended to serially couple the Equilibrium Calculators. This is where one Equilibrium Calculator activity is the successor to another Equilibrium Calculator activity that performs an ordinary equilibrium calculation, and then, for the condition you are interested in, you make a rough estimate of the value. This gives the fixed phase calculation better starting values.
  - 1. Right-click a node that can add an Equilibrium Calculator to your project.
  - 2. In the **Project** window, click the **Equilibrium Calculator** activity node.
  - 3. In the **Configuration** window, click **Switch to advanced mode**.
    - If the **Default calculation mode** is globally set to **Advanced** then by default the Configuration window is already in advanced mode.
- "Changing Global Settings" on page 238

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- 4. Under **Condition Definitions**, click the **Add quantity** <sup>(2)</sup> button.
- 5. Select **Fix phase** from the first menu.
- 6. Select the phase that you want to fix in the second menu, and set the amount of that phase you want to fix the phase to. For example, if you want to know when your material starts to melt, set the phase to **LIQUID** and the value to **0.0**:

Conditions Functions Options			
Number of missing conditions is -1		Switch to simplified mode	]
Condition Definitions		· · · · · · · · · · · · · · · · · · ·	~
ⓒ 🤤 🔽 Temperature 🗸 ∨	Kelvin v 1000.0		
③	LIQUID V 0.0		

Above the condition definitions, the **Number of missing conditions is** field is probably **-1**. This means that you must remove one condition.

7. To remove a condition, either click the remove Sutton, or click to clear the checkbox. For example, if you want to know at what temperature your material starts to melt, then you cannot have a fixed temperature, in which case the temperature condition is not selected:

Conditions Functions Options		
Number of missing conditions is 0		Switch to simplified mode
Condition Definitions		^
💿 😑 🔲 Temperature 🗸 🗸	Kelvin v 1000.0	
Image: Optimized state       Image: Optimized state <th>LIQUID V 0.0</th> <th></th>	LIQUID V 0.0	

The **Number of missing conditions is** field displays **0**. When you have set all your conditions, you can perform the Equilibrium Calculator (now or later).

## **Options Settings**

These settings are for the <u>"Equilibrium Calculator" on page 83</u> and are located on the **Configuration** window  $\rightarrow$  **Options** tab. You can change these locally for a specific Equilibrium Calculator or globally in the **Options** settings. To open the Options window: From the main menu select **Tools**  $\rightarrow$  **Options**. Click the **Activities** tab then click **Calculation**.

Default calculation mode: Simplified   Default	calculation type: Single equilibrium 🔻
Single Equilibrium and Property Grid	
<ul> <li>Global minimization</li> <li>Max grid points</li> <li>Coarse          <ul> <li>2,000</li> <li>2,000</li> <li>Approximate driving force for metastable phases</li> <li>Force positive definite phase Hessian</li> <li>Control step size</li> </ul> </li> </ul>	Max no of iterations:500 Image: 500 Image
Property Diagram           Image: Constraint of the second	Phase Diagram Global minimization
Every 10th eq 10	At node points
	Generate starting points automatically Use inside meshing points No of meshes along an axis: 3

#### Default calculation mode

Simplified (the default) or Advanced.

#### Default calculation type

Single equilibrium (the default), Property diagram, Phase diagram or Property grid.

#### Single equilibrium and property grid

#### **Global minimization**

The check box is selected by default to ensure that the most stable minimum under the specified conditions is computed.

#### Max no of iterations

By default, the program tries 500 iterations before it gives up. As some models give computation times of more than 1 CPU second/iteration, this number is also used to check the CPU time and the calculation stops if 500 CPU seconds/iterations are used.

#### Max grid points

Coarse (2000 grid points, the default), Medium (20,000 grid points), Fine (200,000 grid

points), or Custom (set your own number of grid points).

#### Required accuracy

The default is 1.0E-6. This is a relative accuracy, and the program requires that the relative difference in each variable must be lower than this value before it has converged. A larger value normally means fewer iterations but less accurate solutions. The value should be at least one order of magnitude larger than the machine precision.

#### Smallest fraction

The default is 1.0E-12. This is the value assigned to constituents that are unstable. It is normally only in the gas phase you can find such low fractions.

The default value for the smallest site-fractions is 1E-12 for all phases except for IDEAL phase with one sublattice site (such as the GAS mixture phase in many databases) for which the default value is always as 1E-30.



In Console Mode, the default of 1E-30 is true unless you are using the command SET\_NUMERICAL\_LIMITS to reset an even-lower value (e.g. 1E-45, that is naturally enforced to all the phases in the system.

#### Approximate driving force for metastable phases

The check box is selected by default, which means the metastable phases are included in all iterations. However, these may not have reached their most favourable composition and thus their driving forces may be only approximate.

If it is important that these driving forces are correct, click to clear the check box to force the calculation to converge for the metastable phases.

#### Force positive definite phase Hessian

The check box is selected by default and this setting determines how to reach the minimum of an equilibrium state.

**<u>ADVANCED\_OPTIONS</u>** on page 167 in the Console Mode Command Reference

#### **Control stepsize**

The check box is selected by default and this setting determines how to reach the minimum of an equilibrium state.

ADVANCED\_OPTIONS" on page 167 in the Console Mode Command Reference

## **Property Diagram**

"About the ADVANCED\_OPTIONS Command" on page 162 in the Console Mode Command Reference

#### **Global minimization**

The check box is selected by default to ensure that the most stable minimum under the specified conditions is computed.

#### Global test interval

Every 10th eq (the default), Always or Custom.

The default is the recommended global test interval i.e. every tenth step and at each phase change during calculations.

If you choose **Custom**, the integer number determines how often the Global Minimization should be used during step and mapping. Any other positive integer number, n, suggests it performs a Global Minimization test at every n-th step during calculations.

## Phase Diagram

<u>"About the ADVANCED\_OPTIONS Command" on page 162</u> in the Console Mode Command Reference

#### **Global minimization**

The check box is selected by default to ensure that the most stable minimum under the specified conditions is computed.

#### **Global test interval**

#### At node points (the default), Every 10th eq, Always or Custom.

The default is the recommended global test interval is used only at node points during the calculations.

If you choose **Every 10th eq**, this is every tenth step and at each phase change during calculations.

If you choose **Custom**, the integer number determines how often the Global Minimization should be used during step and mapping. Any other positive integer number, n, suggests it performs a Global Minimization test at every n-th step during calculations.

#### Generate starting points automatically

The check box is selected by default.

#### Use inside meshing points

Click to select the check box as required. The initial mesh consists of initial equilibrium points added only along the edge/border of the diagram defined by the axis-variables. If you select the check box, then the initial equilibrium points added are also added to the inside edge.

#### No of meshes along an axis

The default is 3, which represents how many intervals of initial equilibrium points are added along an edge in the diagram. For example, if set to 2 (two), initial equilibrium points are added at the beginning, center, and end of each axis-variable forming 2 intervals.

## **Calculating and Plotting Functions**

The **Functions** tab allows you to define your own functions that can be plotted in a Plot Renderer.



For an example of how you can use functions that you yourself define, see
 example <u>"T\_07: User-Defined Functions" on page 18</u> in the *Graphical Mode Examples Guide*.

- 1. Create an **Equilibrium Calculator** node and click it in the **Project** window.
- 2. In the **Configuration** window click the **Functions** tab.
- 3. Under **Quantity Definitions**, define **Q1**. Click the add <sup>(2)</sup> button to **Add a new quantity**. Then from the lists, choose the variable for each quantity to define. The quantities are called **Q1**, **Q2**, **Q3** and so on. In this example, the quantity **Q1** is defined as the **Amount of phase**, with **No normalization**, **LIQUID** in **Mole** units.

Configuration	리 무 ×
	Equilibrium Calculator 1
Conditions Functions Options	
Quantity Definitions	
Q1 = Amount of phase	▼ No normalization ▼ LIQUID ▼ Mole ▼
Function Definitions	
③	1-Q1
○ ○ ▼ f_solid =	1-NP(LIQUID)
	Id Predecessor Perform Tree Create New Successor >

4. Under **Function Definitions**, click the add So button to add a new function. In the left field edit the default name of each function and enter the function itself in the right

field after the equal sign. In this example, the quantity defined as Q1 is renamed to **fraction\_solid**, which gives as its output the fraction of solid phase (**1-Q1**).

You can also use Console Mode syntax when entering the function. For example, the function above can be entered as 1-NP(LIQUID).

- 5. To plot the defined function, right-click **Equilibrium Calculator** and choose **Create new successor**→**Plot Renderer**.
- 6. Click the **Plot Renderer** node. In the tab for the Equilibrium Calculator that contains the function, set an **Axis variable** to **Function**, then from the menu select the defined function. In this example the Y-axis is set to plot the value of the function **fraction\_solid**.

Configuration
🖄 Plot Renderer 2
Save Diagram Show Triangular Show Grid Switch Axes Retain Labels
Equilibrium Calculator 1
Tie lines:     0 ▲       Legend option:     Axis quantity       ▲     Legend style:
X Axis
Axis variable: Temperature
Axis type: Linear
Limits: 1500.0 to 1700.0 step 250.0 Automatic scaling
Y Axis
Axis variable: Function
Axis type: Linear
Limits: 0.0 to 1.0 step 0.1 Automatic scaling
Add Predecessor Perform Create New Successor >
۰ III ا

When run, the Plot Renderer plots the value of the function.

# **Diffusion Calculator**

The Diffusion Calculator is available with three elements if you do not have the additional Diffusion Module (DICTRA) license. With the add-on module you can use all available elements as per the licensed database selected.

A **Diffusion Calculator** allows you to set the conditions for, and perform, a diffusion calculation. The Configuration window for a Diffusion Calculator has these tabs:

- **Conditions:** Set the conditions for your calculation that define the geometry, regions, phases, grids, composition profiles and the thermal profile.
- **Options:** Modify the simulation conditions such as the choosing the default solver (automatic or homogenization) and the time integration method. Also define the time step controls and if you choose homogenization as the solver, define further settings such as the interpolation scheme and whether to use global minimization or not.

# Available Options in Graphical Mode

The Diffusion Module is an add-on module to the core Thermo-Calc software.



A separate license is required to perform calculations for more than three elements. Without it you are able to use the module in *Demo Mode*. This is not available in Console Mode. See <u>"Demonstration (Demo) Mode" on the next page</u> to learn more.

## **Diffusion Simulation Template**

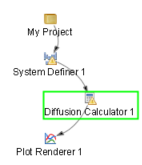
A **Diffusion Simulation** template is available to all Thermo-Calc users when in Graphical Mode.

If you are using the Diffusion Module in Demo Mode, see <u>"Demonstration</u> (Demo) Mode" on the next page for what is available to you. Even if you have a license, you may find yourself in Demo Mode if you have a network license and all the licenses are checked out.

## Using the Template

When you first open Thermo-Calc in Graphical Mode, on the **My Project** window, click the **Diffusion Simulation** button to add a *System Definer*, *Diffusion Calculator* and *Plot Renderer* to the **Project** tree.

Click the **Diffusion Simulation** icon in the Templates window to add **THESE NODES** to the Project.



"Creating a "Project from a Template" in the *Thermo-Calc User Guide* or search the Online Help.

# Diffusion Calculator

A Diffusion Calculator allows you to set the conditions for your calculation that define the geometry, regions, phases, grids, composition profiles and the thermal profile, plus additional options. Many of the settings are automatically set as you define your system. Details about this activity are described in <u>"Diffusion Calculator" on page 92</u> in the *Thermo-Calc User Guide*.

# Demo Database Packages for the Examples Collection

Both a thermodynamic and mobility database are required to run the Diffusion Module simulation. If you have a Diffusion Module (DICTRA) license you can run all the examples because the demonstration database packages are included with your installation.

Select the database packages from the **System Definer Configuration** window to run a simulation.



If you are in Demo Mode, see <u>"Demonstration (Demo) Mode" below</u> to learn more.

Configuration			6 F	φ×
😾 s	ystem Definer 1			
Databases				
③	Package:	Demo: Steels and Fe-alloys (FEDEMO, MFEDEMO)	$\sim$	
◎		 Steels and Fe-alloys (TCFE9, MOBFE4)	^	
Elements Species Phases and Phase Constitution Components Data Sour		Nickel-based superalloys (TCNI8, MOBNI4) Aluminium-based alloys (TCAL4, MOBAL3)		
Periodic Table Alph		Magnesium-based alloys (TCMG4, MOBMG1) Ultra-high-purity silicon (TCSI1, MOBSI1)		
		Demo: Steels and Fe-alloys (FEDEMO, MFEDEMO) Demo: Nickel-based superalloys (NIDEMO, MNIDEMO)	~	

# Demonstration (Demo) Mode

The Diffusion Module and some examples are available to all Thermo-Calc users but only for simulations with three elements. If you do not have a license for then you are in *Demo Mode* when using the Diffusion Calculator or Diffusion Simulation template.

## **Diffusion Simulation Template**

When you are in DEMO mode, in the **Templates** area this is indicated by the text under the logo.



# Diffusion Calculator

If you are experimenting with the Diffusion Calculator in Demo Mode, you may have access to a variety of databases based on your license. However, you can only define three elements for a demo simulation.

If you define more than three elements on the System Definer, when you go to the Diffusion Calculator, the **Perform** button is unavailable and the tooltip explains why.

1 Help	<	Add Predecessor	Perform Tree	Create New Successor	>	
			hr			-
			Diffusion o	alculations in Demo mode	limited	to three elements only



Even if you have more than three elements, the Plot Renderer **Perform** button is still available. However, if you click the button and try to run the simulation the Event Log displays an error message.

# **Configuration Settings**

The following Configuration window is from the *Graphical Mode Quick Start Guide*. It is using project example file *D\_07: Diffusion Carburization Multiphase*.

Configuration
🖄 Carburization multiphase
Conditions Options
Composition unit: Mass percent v
Geometry
Planar V
Austenite
Austenite
Left Interface Boundary
✓ Show left interface boundary
Snow left interface boundary
💿 🜑 Region
Name: Austenite
Width: 0.003 m 100 points and type Geometric v Ratio: 1.02 🜩
Q ⊆ Phase: M3C2
Composition profiles: Edit
Right Interface Boundary
Show right interface boundary
Left Boundary Condition
Mixed zero flux and activity v C Activity v 1.0 Closed system v
Tip: The reference state is defined in the System Definer Cr Zero flux v
Thermal Profile
Isothermal     Non-isothermal
Temperature: 1123.0 Kelvin V
Simulation time: 1000.0 Hours V
•
Image: Weight of the second

These are the settings available from the **Conditions** tab. To edit the defaults for some settings, go to **Tools**  $\rightarrow$  **Options**  $\rightarrow$  **Graphical Mode**  $\rightarrow$  **Diffusion**.

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The Diffusion Module (DICTRA) in Console Mode has commands to perform some of the settings also available on the Diffusion Calculator. Where possible the name of the command is indicated. All the commands are described in the *Diffusion Module (DICTRA) Command Reference*.

# General



See also these Console commands: ENTER\_COMPOSITIONS and ENTER\_ GEOMETRICAL\_EXPONENT

## **Composition Unit**

Choose from Mass percent, Mole percent, Mass fraction or Mole fraction.

### Length Unit

Choose **m**, **µm** or **nm**.

#### Geometry

Choose Planar, Spherical or Cylindrical. For Cylindrical, also enter a value for the First interface position .

With a cylindrical or spherical geometry, the system's zero coordinate (left boundary) is at the centre of the cylinder or sphere. The highest coordinate (right boundary) is defined by the cylinder or sphere radius.

## **Options Settings**

The following settings are available on the **Options** tab for a Diffusion Calculator.

You can edit the defaults for some of these settings by going to **Tools**  $\rightarrow$ **Options**  $\rightarrow$  **Graphical Mode**  $\rightarrow$  **Diffusion**. Also see <u>"Global Settings: Diffusion</u> <u>Calculator" on page 246</u>.



The Diffusion Module (DICTRA) in Console Mode has commands to perform some of the settings also available on the Diffusion Calculator. Where possible the name of the command is indicated. All the commands are described in the *Diffusion Module (DICTRA) Command Reference*.

# Diffusion Calculator Plot Renderer

The following is information about the settings available in both Simplified and Advanced mode for a Plot Renderer when it is a successor to a "Diffusion Calculator" on page 92.



If you are using Console Mode search for "Plot Condition and Independent Variable" in the *Diffusion Module Command Reference*.

# Simplified Mode

In this mode the most common axis variables can be selected. You do not have to think about setting the correct **Plot condition** and **Independent variable**. Use the Advanced mode to have more flexibility.

When you click the **Switch to advanced mode** button, this expands the variable options available from both the X-axis and Y-axis lists. Otherwise the axis variables default to the most logical combination.

#### X-AXIS VARIABLE

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Select the variable for the *x*-axis: **Distance** or **Time**.

- For Distance also choose a unit (m, μm, or nm) and what region to include. The list contains the regions defined on the Diffusion Calculator or you can choose All regions.
- For Time choose a unit (Seconds, Minutes, Hours, or Days).

#### **Y-AXIS VARIABLE**

Select the variable to plot on the *y*-axis. The available options are based on choices such as whether **Distance** or **Time** is chosen for the *x*-axis, whether isothermal or non-isothermal is selected, whether there is a homogenization solver, or if there is one or more regions.

#### **Distance along the x-axis**

If **Distance** is plotted along the x-axis, then by default the **Composition** is plotted along the *y*-axis.

- From the center list, choose the component to plot, or select All.
- From the next list, choose the unit to be **Mass percent**, **Mole percent**, **Mass** fraction, Mole fraction, or u-fraction.

#### Time along the x-axis (one region)

If **Time** is plotted along the x-axis, then choose the *y*-axis variable to plot plus the additional settings:

- Total phase fraction: By volume or By mass. Then select a phase from the list.
- Total fraction of element: By mole or By mass. Then select a component from the list.
- Total fraction of element in phase: By mole or By mass. Then select a component and a phase from the lists.
- For a non-isothermal thermal profile **Temperature** is available: **Kelvin**, **Celsius**, or **Farenheit**.

#### Distance along the x-axis (Homogenization)

If **Distance** is plotted along the x-axis and a Homogenization solver is being used, then choose **Composition** or **Phase fraction** to plot on the *y*-axis. Then select the applicable settings.

- Choose the component to plot, or select All and the unit Mass percent, Mole percent, Mass fraction, Mole fraction, or u-fraction.
- Choose By mole or By mass then select a component from the list.

#### Time along the x-axis (two or more regions)

If **Time** is plotted along the x-axis, then choose the *y*-axis variable to plot plus the additional settings:

- Total phase fraction: By volume or By mass. Then select a phase from the list.
- Total fraction of element: By mole or By mass. Then select a component from the list.
- Total fraction of element in phase: By mole or By mass. Then select a component and a phase from the lists.
- Composition at interface: Mass percent, Mole percent, Mass fraction, Mole fraction. Then choose a Region and whether it is Upper or Lower and a component.
- Position of interface: Choose a unit (m, µm, or nm) and the interface.
- Velocity of interface: Choose a unit (m, µm, or nm) and the interface.
- Width of region: Choose a unit (m, µm, or nm) and the interface.
- For a Non-isothermal profile, Temperature is available. Choose a unit Kelvin, Celsius or Fahrenheit.

## AXIS TYPE

Select the type of axis scale: Linear, Logarithmic, Logarithmic 10 or Inverse.

## LIMITS

Specify the range along the axis to show in the plot. In the fields, enter the minimum and maximum values of the axis variable. You can also determine the **step** size between the tick marks along each axis.

Select the Automatic scaling check box to allow the program to set the limits.

## ΤΙΜΕ



An alternative **Time** setting is available in Advanced mode.

When the *x*-axis variable is **Distance**, choose a **Time** unit (**Seconds**, **Minutes**, **Hours**, or **Days**). Enter a numerical value.

## ADD AND REMOVE AXES

Use the **Add an axis** and **Remove this axis** buttons to add additional X- and/or Y-axes to a plot or to merge plots (to overlay one plot with another in the same diagram).

## **Advanced Mode**

Click the **Switch to advanced mode** button to access more settings. These settings enable you to manually select any combination of axis variables, **Plot condition** and **Independent variable**. Examples of advanced plot options are activity of a component at an interface, diffusivities, and diffusion paths. Also user-defined functions can be used as plot variable where Console Mode syntax is used to define functions. In some cases, sensible default values are selected, but they can be changed according to your needs.

## PLOT AXIS SETTINGS IN ADVANCED MODE

- If any plot axis variable of the type **Integral** is selected, then the plot condition is set to **Integral** and the Independent variable is set to **Time**.
- If a plot axis variable is set to **Time** then the plot condition is set to **Time** and the Independent variable is also set to **Time**.
- If a plot axis variable is set to **Distance** then the plot condition is set to **Distance** or **Interface** and the Independent variable is set to **Distance**.

Plot condition	Independent Variable settings based on the Plot condition			
<b>Time</b> : Choose a unit ( <b>Seconds, Minutes,</b> <b>Hours,</b> or <b>Days</b> ). Enter a numerical value.	The default is <b>Distance</b> : Choose a <b>Global</b> or <b>Local</b> distance. A <b>Global</b> distance is counted from the rightmost interface of the system. A <b>Local</b> distance is counted from the lower interface of the Region selected. You can also choose <b>Time</b> , which requires no further settings.			
<b>Distance</b> : Choose a specific <b>Region</b> from the list or use the default <b>All regions</b> .	The default is <b>Time</b> and there are no further settings. You can also choose a <b>Global</b> or <b>LocalDistance</b> . A <b>Global</b> distance is counted from the rightmost interface of the system. A <b>Local</b> distance is counted from the lower interface of the Region selected.			
Interface: From the lists choose a <b>Region</b> and whether it is the <b>Upper</b> or <b>Lower</b> region.	The default is <b>Distance</b> : Choose a <b>Global</b> or <b>Local</b> distance. A <b>Global</b> distance is counted from the rightmost interface of the system. A <b>Local</b> distance is counted from the lower interface of the Region selected. You can also choose <b>Time</b> , which requires no further settings.			
Integral	The default is <b>Time</b> and there are no further settings. You can also choose a <b>Global</b> or <b>LocalDistance</b> . A <b>Global</b> distance is counted from the rightmost interface of the system. A <b>Local</b> distance is counted from the lower interface of the Region selected.			

# Pause, Resume and Cancel Diffusion Calculations

Diffusion calculations are often complex simulations that take some time to complete. Sometimes you may want or need to pause or resume a calculation, or make adjustments to your compositions and start again.

# **GRAPHICAL MODE**

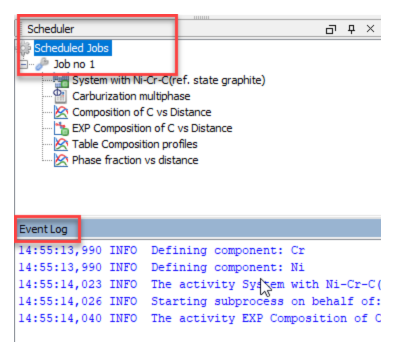
In Graphical mode, you pause and resume from the Diffusion Calculator **Configuration** window.



This is not available for the other calculators.

#### To Pause and Resume a Job

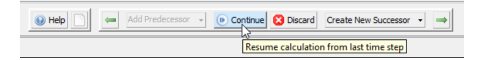
1. Run the job (i.e. click **Perform Tree**). In the **Scheduler**, you can see the job listed and in the **Event Log** you can follow the progress of the calculation.



- 2. In the **Project** window, click the **Diffusion Calculator** node.
- 3. In the **Configuration** window at the bottom, click **Pause**. If there are intermediate results available these will be listed in the **Event Log**.

😢 Help 📄	Add Predecessor 👻	U Pause	Create New Successor 👻	<b>a</b>
		Pa	use to display intermediat	e result if available

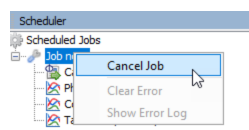
 When ready, click **Continue** to resume your calculations from the last time step or click **Discard** to discard the calculation (then click **Yes** or **No** on the window that opens).



#### To Cancel a Job

• In the Scheduler window, right-click the job you want to cancel and select Cancel

Job.



# CONSOLE MODE

In response to a command or a lengthy calculation, Thermo-Calc may display a lot of text in the Console window. Use these keyboard shortcuts to pause, resume or stop a process.

- To pause the text on printing press <Ctrl+S>.
- To resume the printing on screen, press <Ctrl+Q>.
- When Thermo-Calc is performing a stepping or mapping operation, the results are continuously printed in the Console window. To stop the calculation of the current region of the mapping and stop the output, press <Ctrl-A> (Windows) or <Ctrl-C> (Linux).

# About the Homogenization Functions

The geometrical interpretation of the Hashin-Shtrikman bounds are concentric spherical shells of each phase. For the general lower Hashin-Shtrikman bound the outermost shell consists of the phase with the most sluggish kinetics and vice versa for the general upper bound. The geometrical interpretation of the Hashin-Shtrikman bounds suggest further varieties of the bounds, viz. *Hashin-Shtrikman bound with prescribed matrix phase* and *Hashin-Shtrikman bound with majority phase as matrix phase*, where the outermost shell consist of a prescribed phase or the phase with highest local volume fraction, respectively.

The geometrical interpretation of the Wiener bounds are continuous layers of each phase either parallel with (upper bound) or orthogonal to (lower bound) the direction of diffusion.



The labyrinth factor functions (described below) are only available in Console Mode.

The labyrinth factor functions implies that all diffusion takes place in a single continuous matrix phase. The impeding effect on diffusion by phases dispersed in the matrix phase is taken into account by multiplying the flux with either the volume fraction (*Labyrinth factor f with prescribed matrix phase*), or the volume fraction squared (*Labyrinth factor f\*\*2 with prescribed matrix phase*), of the matrix phase.



8

The varieties with excluded phases (described below) are only available in Console Mode.

The varieties with excluded phases are useful in several respects. First, if a phase is modelled as having zero solubility for a component, the mobility of that component in that phase is undefined, which causes a (non-terminal) error. Setting a phase as excluded causes the mobility of all components in that phase to be set to zero. Second, often there are some major matrix solid solution phases and some minor precipitate phases. If the mobilities in the minor precipitate phases are zero the lower Hashin-Shtrikman bound is useless as it produces a kinetic coefficient of zero. However, using *General lower Hashin-Shtrikman bound with excluded phase(s)* the excluded phases are not considered when evaluating what phase has the most sluggish kinetics.

# **Precipitation Calculator**

The Precipitation Calculator is available with three elements if you do not have the additional Precipitation Module (TC-PRISMA) license. With the add-on module you can use all available elements.

A Precipitation Calculator allows you to set the conditions for, and perform, a precipitation calculation. The Configuration window for a Precipitation Calculator has these tabs where you define the settings:

- **Conditions**: Set the conditions for your calculation that define the Matrix and Precipitate phases. Choose the Calculation Type.
- **Options**: Modify Numerical Parameters that determine how the conditions are calculated. The Growth rate model can be set to Simplified or Advanced mode.

# Available Options with the Precipitation Module

The Precipitation Module, previously referred to as TC-PRISMA, is an add-on module to the core Thermo-Calc software. A separate license is required to perform calculations for more than three elements. Without it you are able to use the module in *Demo Mode*.

"Demonstration (Demo) Mode" on page 106

# **Precipitation Simulation Template**

A Precipitation Simulation template is available to all Thermo-Calc users.

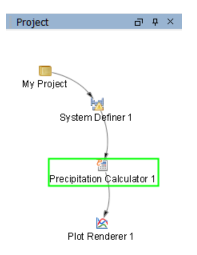


If you are accessing the Precipitation Module in Demo Mode, see <u>"Demonstration (Demo) Mode" on page 106</u> for what is available to you.

## USING THE TEMPLATE

When you first open Thermo-Calc in Graphical Mode, on the **My Project** window, click the **Precipitation Simulation** button to add a *System Definer*, *Precipitation Calculator* and *Plot Renderer* to the **Project** tree.

Click the **Precipitation Simulation** button in the templates section to add these nodes to the Project.



"Creating a "Project from a Template" in the *Thermo-Calc User Guide* or search the Online Help.

# **Precipitation Calculator**

A Precipitation Calculator allows you to set the conditions for, and perform, a precipitation calculation.

# Demo Database Packages for the Examples Collection

Both a thermodynamic and mobility database are required to run the Precipitation Module simulation. If you have a Precipitation Module (TC-PRISMA) license you can run all the examples as the demonstration database packages are included with your installation. Select the database packages from the **System Definer** $\rightarrow$ **Configuration** window to run a simulation.



If you are using the Precipitation Module in Demo Mode, see <u>"Demonstration</u> (Demo) Mode" on the next page to learn more.

Configuration			0 P
	System Definer 1		
Databases			
FEDEMO: Iron Demo Database v2.0	<ul> <li>Package:</li> </ul>	Demo: Steels and Fe-alloys (FEDEMO, MFEDEMO)	$\sim$
S S MFEDEMO: Fe-Alloys Mobility demo database v2.0	~	Steels and Fe-alloys (TCFE9, MOBFE4)	^
Elements Species Phases and Phase Constitution Components Data S	ources Description	Nickel-based superalloys (TCNI8, MOBNI4) Aluminium-based alloys (TCAL4, MOBAL3)	
Periodic Table	lphabetic List	Magnesium-based alloys (TCMG4, MOBMG1) Ultra-high-purity silicon (TCSI1, MOBSI1)	
		Demo: Steels and Fe-alloys (FEDEMO, MFEDEMO) Demo: Nickel-based superalloys (NIDEMO, MNIDEMO)	~

# Demonstration (Demo) Mode

The Precipitation Module, and some examples, are available to all Thermo-Calc users but only for simulations with three elements. If you do not have a license for the Precipitation Module then you are in *Demonstration Mode* when using the Precipitation Calculator or Precipitation Simulation template.

#### **PRECIPITATION SIMULATION TEMPLATE**

When you are in DEMO mode, in the **Templates** area this is indicated by the text under the logo.



#### **PRECIPITATION CALCULATOR**

If you are experimenting with the Precipitation Calculator in Demo Mode, you may have access to a variety of databases based on your license. However, you can only define three elements for a demo simulation.

If you define more than three elements on the System Definer, when you go to the Precipitation Calculator, the **Perform** button is unavailable and the tooltip explains why. In this case one of the chosen elements needs to be removed and then the Perform button is made available.

🔞 Help	<	Add Predecessor	Perform Isothermal S	imulation	Create New Successor	>	
				Precipitation simulations in Demo mode limited to three elements only.			

Even if you have more than three elements, the Plot Renderer or Table Renderer **Perform** button is still available. However, if you click the button and try to run the simulation the Event Log displays an error message.

## **Configuration Settings**

Below are details about the settings available from the Conditions and Options tabs.

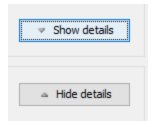
Configuration	L X
🔠 Precipitation Calculator 1	
Conditions Options	
Composition unit: Mass percent 🗸	^
Composition Fe 99.984	
Composition C 0.016	
Matrix Phase	
Phase: BCC_A2 v Show details	
O Precipitate Phase	- 1
Phase: CEMENTITE V A Hide details	
Nucleation sites: Dislocations V Calculate from matrix settings 5.87951601467259E20 m-3	
Interfacial energy: User-defined $\checkmark$ f(r,T) = 0.24 J/m <sup>2</sup>	
Growth rate model: Advanced 🗸	
Morphology: Sphere 🗸	
Transformation strain: Disregard $\checkmark$	
Molar volume: Database V 1.0E-5 m <sup>3</sup> /mol	
Phase boundary mobility: 10.0 m <sup>4</sup> /Js	
Phase energy addition: 0.0 J/mol	
Approximate driving force:	
Preexisting size distribution: Edit particle size distribution	
Calculation Type	
● Isothermal ○ Non-isothermal ○ TTT diagram ○ CCT Diagram	
Temperature: 102.0 Celsius 🗸	
Simulation time: 600000.0 Seconds 🗸	
	~
🕘 Help 📄 👄 Add Predecessor 🗸 📀 Perform Isothermal Simulation Create New Successor 🗸 👄	



You can edit the defaults for some of these settings by going to **Tools**  $\rightarrow$  **Options**  $\rightarrow$  **Graphical Mode**  $\rightarrow$  **Precipitation**.

## Show or Hide Details

Click **Show details** (found to the right of the section) to view some of the additional settings listed below.



## **Composition Unit**

Choose from Mass percent, Mole percent, Mass fraction or Mole fraction.

Matrix Phase

Phase

Only phases with kinetic data can be selected as the matrix phase. If the list is empty, go to the System Definer to make sure that both thermodynamic and kinetic databases are selected and defined.

Search the Help for "System Definer".

Choose a phase from the list. The list is based on the settings for the System Definer. When setting up your system, choose a matrix phase with kinetic data available in the database.

## **Elastic properties**

Choose the default **Disregard** to ignore the elastic properties. Otherwise choose **Isotropic** or **Cubic** and enter the elastic constants accordingly. Default elastic constants are given based on the major element of your alloy system. The elastic properties can affect nucleation rate, nucleation size, and particle shape.

#### Molar volume

Choose the default to take the value from the **Database** or **User defined** to enter another value. **Database** is available in the list when the molar volume for the phase is defined in the thermodynamic database.

#### Grain size

Enter a numerical value and choose a unit from the list. The default is 1.0E-4 m. The value entered for grain size changes the available nucleation sites when Grain boundaries, Grain edges, or Grain corners is selected along with "Calculate from matrix settings" in the Precipitate Phase options.

#### Grain aspect ratio

Enter a numerical value. The default is 1.0. The value entered for grain aspect ratio changes the available nucleation sites when Grain boundaries, Grain edges, or Grain corners is selected along with "Calculate from matrix settings" in the Precipitate Phase options.

#### **Dislocation density**

Enter a numerical value. The default is 5.0E12 m<sup>-3</sup>. The value entered for dislocation density changes the available nucleation sites when Dislocations is selected along with "Calculate from matrix settings" in the Precipitate Phase options.

#### Mobility enhancement prefactor

A parameter that multiplies to the mobility data from database. This value scales the

mobility by a constant amount. This can be useful, for example, when the material has a higher than normal vacancy concentration at the start of the precipitation simulation. (e.g. from a prior solutionizing and quenching treatment).

### Mobility enhancement activation energy

A value that adds to the activation energy of mobility data from database. This value scales the mobility by a temperature dependent amount. Similar usage as mobility enhancement prefactor.

## Precipitate Phase

#### Phase

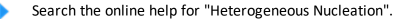


The phases available to choose have both thermodynamic and kinetic data. If the list is empty, go to the System Definer to make sure that both types of databases are selected and defined.

Search the Online Help for "System Definer".

Choose a phase from the list. The list is based on the System Definer settings .

#### **Nucleation sites**



Choose one of the following from the list.

- Bulk, Grain boundaries, Grain edges, Grain corners, or Dislocations.
- For Grain boundaries, Grain edges and Grain corners, enter the Wetting angle in addition to the matrix settings.

Click to select the **Calculate from matrix settings** check box if you want to calculate the number density of sites from the matrix grain size or dislocation density.

To enter a specific value for the number of **Nucleation sites**, deselect the check box.

#### **Interfacial energy**

Search the online help for "estimation of coherent interfacial energy".

Choose **Calculated** to use the estimated value and then enter a different **prefactor** value if you want to adjust the estimated value. You can also choose **User defined** to enter a value in J/m<sup>2</sup>.

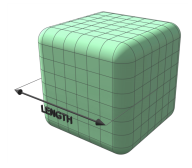
## Morphology

Search the online help for "Precipitation Morphology".

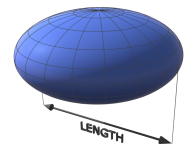
Choose the particle shape from the list: Sphere (default), Cuboid, Plate and Needle.

**Cuboid** is available only when **Cubic** is chosen in **Elastic properties** in the Matrix Phase section.

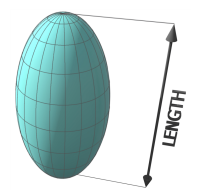
When **Plate** or **Needle** is chosen, a check box is provided to **Calculate aspect ratio from elastic energy**, which is available when **Isotropic** or **Cubic** is chosen in **Elastic properties** in the Matrix Phase section. Alternatively, a value can be entered in the **Grain aspect ratio** field to provide a constant aspect ratio.



*Figure 1: The cuboid shape is described by a supersphere. Cuboids have six faces, which form a convex polyhedron.* 



*Figure 2: The plate is described as oblate spheroid. Oblate spheroids have rotational symmetry around an axis from pole to pole.* 



*Figure 3: The needle shape is described as prolate spheroid. A prolate spheroid is a surface of revolution obtained by rotating an ellipse about its major axis.* 

#### **Transformation strain**

Search the online help for "Precipitation Morphology".

Choose the default, **Disregard**, to ignore the transformation strain. Otherwise choose **Calculated from molar volume** to obtain a purely dilatational strain, or **User defined** to enter the values.

These options are available only when **Isotropic** or **Cubic** is chosen in **Elastic properties** in Matrix Phase. In addition, if **Cuboid** is chosen as **Morphology**, only **Calculated from molar volume** is available.

#### Molar volume

Choose the default to take the value from the **Database** or **User defined** to enter another value. **Database** is available in the list when the molar volume for the phase is defined in the thermodynamic database.

#### Phase boundary mobility

A parameter that accounts for interface-controlled growth. Only effective if a very small, positive value is used. Use with caution due to a tentative treatment.

#### Phase energy addition

An energy value that adds to the Gibbs free energy from database.

#### Approximate driving force

Search the online help for "Nucleation Theory".

Select the check box to include this if simulations with several compositions sets of the same

phase create problems.

#### Preexisting size distribution

Select the check box to include this. Click **Edit particle size distribution** to open the **Preexisting Particle Size Distribution** window where you can edit the parameters and view a graph comparing the radius and number density for the selected component. See *Particle Size Distribution* for instructions.

See example P\_10 and P\_11 in the *Graphical Mode Examples Guide*.

## Calculation Type

## Isothermal

Use an **Isothermal** calculation type to do a precipitation simulation at constant temperature. Enter a **Temperature** and **Simulation time**.



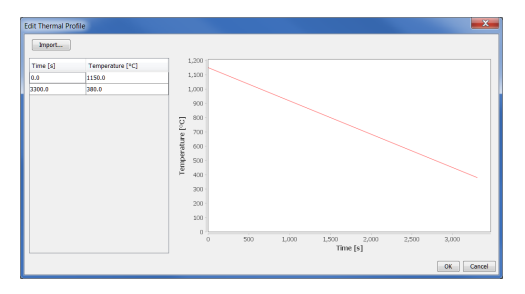
See examples P\_01, P\_02, P\_04, P\_05, P\_08 and P\_09 in the *Graphical Mode Examples Guide*.

#### Non-isothermal

Search the help for "Nucleation during a non-isothermal process".

For the **Non-isothermal** calculation type, select a **Temperature unit** and **Time unit** from the lists. Enter a value for the **Simulation time**.

Click **Thermal Profile**. Here the **Temperature** and **Time** coordinates of thermal profile points are entered. A minimum of two points is required. You can also click **Import** to add your own thermal profile from an Excel spreadsheet.



See example P\_06 in the *Graphical Mode Examples Guide*.

# TTT diagram

Use a **TTT-diagram** calculation type to do a precipitation simulation of the time-temperature-transformation (TTT).

Temperature: Enter Min, Max, and Step values for the and choose a Unit (Kelvin, Celsius, or Fahrenheit).

Max annealing time: Enter a numerical value and choose a Unit (Seconds, Minutes, Hours, or Days).

**Stop criteria**: Choose **Volume fraction of phase** or **% of equilibrium fraction** and then enter a numerical value in the field. For each temperature, the simulation stops when the stop criteria is fulfilled or if the maximum simulation time is reached, whichever happens first.



See example P\_03 in the *Graphical Mode Examples Guide*.

# **CCT diagram**

Use a **CCT-diagram** calculation type to do a precipitation simulation of the continuouscooling-transformation (CCT)

Temperature: Enter Min and Max values and choose a Unit (Kelvin, Celsius, or Fahrenheit).

**Cooling rate(s)**: Enter a range of values in the field, e.g. .01 .1 1 10 100. These values are equal to K/s, °C/s or °F/s per second based on the Unit selected above.

**Stop criteria**: Enter a numerical value for the **Volume fraction of phase**. For each temperature, the simulation stops when the stop criteria is fulfilled or if the maximum simulation time is reached, whichever happens first.



See example P\_07 in the *Graphical Mode Examples Guide*.

### Numerical Parameters

Search the help for "numerical method".



Click **Options** to go to these settings.

#### Max time step fraction

The maximum time step allowed for time integration as fraction of the simulation time. The default is 0.1.

### No. of grid points over one order of magnitude in radius

Default number of grid points for every order of magnitude in size space. The default is 150.0.

#### Max no. of grid points over one order of magnitude in radius

The maximum allowed number of grid points in size space. The default is 200.0.

### Min no. of grid points over one order of magnitude in radius

The minimum allowed number of grid points in size space. The default is 100.0.

#### Max relative volume fraction of subcritical particles allowed to dissolve in one time step

The portion of the volume fraction that can be ignored when determining the time step. The default is 0.01.

#### Max relative radius change

The maximum value allowed for relative radius change in one time step. The default is 0.01.

#### Relative radius change for avoiding class collision

Set a limit on the time step. The default is 0.5.

### Max overall volume change

This defines the maximum absolute (not ratio) change of the volume fraction allowed during one time step. The default is 0.001.

### Max relative change of nucleation rate in logarithmic scale

This parameter ensures accuracy for the evolution of effective nucleation rate. The default is 0.5.

### Max relative change of critical radius

Used to place a constraint on how fast the critical radium can vary, and thus put a limit on time step. The default is 0.1.

### Min radius for a nucleus to be considered as a particle

The cut-off lower limit of precipitate radius. The default is 5.0E-10 m.

### Max time step during heating stages

The upper limit of the time step that has been enforced in the heating stages. The default is 1.0 s.

### Maximum relative solute composition change at each time step

Set a limit on the time step by controlling solute depletion or saturation, especially at isothermal stage. The default is 0.01.

# Particle Size Distribution (PSD)

See example P\_10 and P\_11 in the *Graphical Mode Examples Guide*.

On the Precipitation Calculator you can set the pre-exisiting particle size distribution (PSD) parameters in the **Show details** section. The size distribution can be entered as a predefined distribution, by importing a file, or by manually entering information into a table.

Accessing the 'Pre-existing size distribution' settings

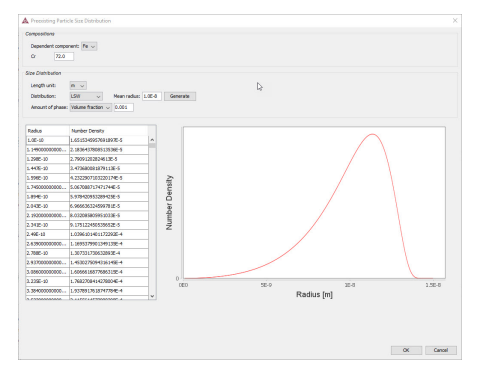
1. To the right of the Precipitate Phase settings fields, click Show details.

💿 🔵 Precipitate Ph	ase		
Phase:	CEMENTITE ~		L
Nucleation sites:	Bulk ~	Calculate from matrix settings 8.603059285714286E28 m <sup>-3</sup>	ł
Interfacial energy:	Calculated 🗸	with prefactor 1.0	

2. Click to select the Preexisting size distribution check box.

😳 🛑 Precipitate Phase			
Phase:	CEMENTITE	~	
Nucleation sites:	Bulk	~	Calc.
Intering:	Calculated		w <sup>all</sup> of
Ph. Je energy addi.	0.5	J/h	
Approximate driving force:			
Preexisting size distribution:	Edit pa	article size distribut	ion

3. Click Edit particle size distribution to open the window as in example P\_10.



#### Defining the Preexisting size distribution

- 1. Under **Compositions**, choose a **Dependent component** from the list and enter the composition for the other component.
- 2. Under Size Distribution define the following:
  - a. Choose a Length unit: m, µm, or nm.
  - b. Choose a **Distribution**: **LSW** (Lifshitz-Slyozov-Wagner), **Normal**, **Log normal** or **From file**. You can also manually enter numbers into the table instead of importing a file.

Radius [m]	Number Density	
	2	

- c. For LSW, Normal and Log normal, enter a Mean radius.
- d. For Normal and Log normal, enter a value for the standard deviation (Std).
- e. For **From File**, click **Import** and navigate to the file containing the required information and click **Open**. This file can be in .xls, .xlsx, .csv or .txt formats. The file should consist of two columns with values where the first column contains **radius** data and the second contains **number density** data.
- f. For all options, choose an **Amount of phase**: **Volume percent** or **Volume fraction** and then enter a number in the field.
- g. Once the **Distribution** is defined, click **Generate**.

### **Plot Renderer**

The following is information about the settings available for a Plot Renderer when it is a successor to a "Precipitation Calculator" on page 104.

"Plot Renderer" on page 122.

There are also specific settings related to non-isothermal simulations that are detailed in this topic.

#### Plot Settings

#### Legend option

Select whether the diagram's legend displays **On** or **Off**.

### Axis variable

Set the state variable you want plotted along the X-axis and the Y-axis.

Select the **Separate multimodal PSD** check box to enter settings for **Points**, **Excess kurtosis**, **Valley depth ratio** and **Minimum peak**.



See "Non-Isothermal Simulations" on page 120 for definitions.

#### Axis type

Select the type of axis scale: Linear, Logarithmic, Logarithmic 10 or Inverse.

### Limits

Specify the range along the axis to show in the plot. In the fields, enter the minimum and maximum values of the axis variable. You can also determine the **step** size between the tick marks along each axis.

Select the Automatic scaling check box to allow the program to set the limits.

### Unit (Time X Axis)

Choose a Unit : Seconds, Hours, or Days.

#### Add an axis and Remove this axis buttons

Use the Add an axis and Remove this axis a buttons to add additional X- and/or Y-axes to a plot or to merge plots (to overlay one plot with another in the same diagram). When merging plots, you define this variable as the Z-axis.

### **Axis Variables**

- Mean radius Spherical radius of average volume of all particles for a specific phase and nucleation type, regardless of their actual shapes.
- **Critical radius** Spherical radius of critical nuclei for a specific phase and nucleation type.
- Matrix composition Instantaneous compositions of the matrix phase.
- Number density Instantaneous number of particles per unit volume for a specific phase and nucleation type.
- **Size distribution** Number of particles varying with their sizes per unit volume per unit length, for a specific phase and nucleation type, at a specific time.
- Volume fraction Instantaneous volume fraction for a specific phase and nucleation type.
- Nucleation rate Instantaneous number of nuclei per unit volume per unit time for a specific phase and nucleation type.
- Normalized driving force Instantaneous nucleation driving force per unit mole of elements for a specific phase and nucleation type. It is normalized with RT and therefore, dimensionless.
- Mean cubic factor Average cubic factor of cuboid particles. Available only when cuboid is selected as the **Morphology**.

The following settings area available as indicated when **Needle**, **Plate** or **Cuboid** are selected as the **Morphology** in the *Precipitate Phase* section on the Precipitation Calculator.

- **Cubic factor distribution** Variation of cubic factor with particle size at a specific time. Available with a cuboid morphology.
- Mean aspect ratio Average aspect ratio of non-spherical particles. Available with a needle or plate morphology. Note that this is always larger than 1, where 1 = a sphere.
- **Mean particle length** Diameter of non-spherical particles along the longer axis. Available with a needle or plate morphology.
- Aspect ratio distribution Variation of aspect ratio with particle size at a specific time. Available with a needle or plate morphology.

# Non-Isothermal Simulations

When doing non-isothermal simulations it is common that particles grow in different generations. This results in multi-modal size distributions. To correctly estimate the properties of these different generations of particles you need to separate the peaks of multi-modal distributions.

### Separate multimodal PSD

When the **Separate multimodal PSD** check box is selected on a Plot Renderer activity for the Precipitation Calculator, the size distribution is evaluated at the given time steps and checked for multi-modal peaks. These are separated and used to calculate the specified property.

Configuration	
Plot Renderer (PSD Ni-8Al-8Cr)	
Save Diagram Show Triangular Show Grid Switch Axes Retain Labels	
Precipitation Ni-8AI-8Cr	
Legend option: On 💌	
Y Axes	
Axis variable:       Size distribution         Excess kurtosis:       25.0         Valley depth ratio:       0.05         Minimum peak:       100000.0	

It can be applied on the following plot properties:

- Mean radius
- Size distribution
- Number density
- Volume fraction

### Points

This field is available when **Mean radius**, **Number density** or **Volume fraction** is selected as the **Axis variable**. Since the evaluation of multi-modality at each time step is costly, you can specify how many evaluation **Points** to use. The points are geometrically distributed over the time scale where nucleated particles are found. The default is 40 points.

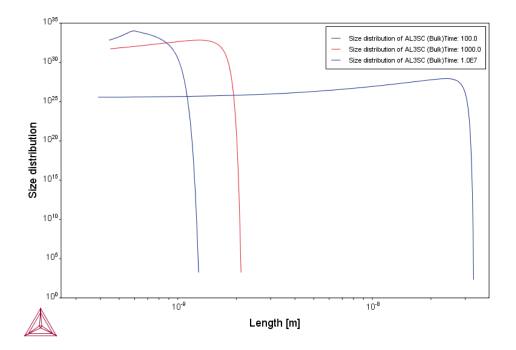
### Time

This field is available when **Size distribution** is selected as the **Axis variable**. Enter one or a series of numbers in the field, separated by a space.

For example, if you enter 100.0, 1000.0 and 1.0E7 in the field:

Y Axes				
•				
Axis variable:	Size distribution	▼ AL3SC (Bulk)	Time: 100.0 1000.0 1.0E7	

When you click **Perform** the times are plotted:



#### **Excess Kurtosis**

The **Excess kurtosis** number specifies the sensitivity when the program evaluates whether a curve is bimodal or not. The evaluation process calculates the excess kurtosis of the given size distribution and compares it with the input value. If the value is lower than the given value, the distribution is split. The excess kurtosis is a statistical measure that measures how *peaked* and *how heavy tail* a distribution has. For the definition used see http://en.wikipedia.org/wiki/Kurtosis. The default is 25.0.

#### Minimum Separation Limit (Valley Depth Ratio)

The **Minimum Separation Limit** is a rough method to prevent that noise or wiggles are interpreted as different distributions. If a local minima is encountered above this value the following peak is not interpreted as a new distribution. The valley depth ratio is the ratio of the minimum separation limit to the peak value. The default is 0.01.

#### **Minimum Peak**

Any separated peak below the entered **Minimum peak** value is removed. The default is 100000.0.

# **Plot Renderer**

The Plot Renderer is where you define the settings and output results in a diagram that displays in the Results window. The available Plot Renderer settings are based on the System Definer and calculator activity node predecessors (e.g. Equilibrium, Scheil, and Property Model Calculators). Setting availability is also determined by the calculation type.

Plotting Options" on page 132

# Plot Renderer for the Add-On Modules

If you have an add-on Diffusion Module (DICTRA) and/or Precipitation Module (TC-PRISMA), there can be additional functionality available on the Plot Renderer. See <u>"Diffusion</u> <u>Calculator Plot Renderer " on page 97</u> and <u>"Plot Renderer " on page 117</u>.

Setting/Button	Description
Save Diagram	Save the plot diagram.
Show Triangular	Display the diagram in triangular form, with the X-axis along the base and the Y-axis along the left side. Such a diagram is often useful if you want to plot the fractions of two components in a ternary diagram.
Show Grid	Overlay a grid on the diagram.
Switch Axes	Show the plotting X axis variable on the diagram's Y-axis, and the Y axis variable on the diagram's X-axis.
Retain Labels	Toggle the default to keep labels displaying on the plot. By default labels are retained.

# **Plot Toolbar**

# **Configuration Settings**

The following is an example of the **Configuration** window for a Plot Renderer that is the successor to a Property Model Calculator.

Configuration	
🖄 Plot Renderer 1	
Save Diagram Show Triangular Show Grid Switch Axes Retain Labels	
Property Model Calculator: Coarsening	
Tie lines: 0 → Plot type: Heat map ▼ Interpolate: Never ▼	
X Axis	
Axis variable: Mass percent Cr 🔹	
Axis type: Linear	
Limits: 500.0 to 3000.0 step 250.0 V Automatic scaling	
Linits. 500.0 to 5000.0 step 250.0 V Automatic scaling	
Y Axis	
Axis variable: Mass percent C 🔹	
Axis type: Linear 🔻	
Limits: 500.0 to 3000.0 step 250.0 V Automatic scaling	
Z Axis	
Axis variable: Coarsening rate coefficient	
Axis type: Linear	
Limits: 500.0 to 3000.0 step 250.0 V Automatic scaling	
Add Predecessor Perform Create New Successor >	

# **Examples of Calculation Type Differences**

 If you choose a *Property grid* (Equilibrium Calculator) or *Grid* (Property Model Calculator) type, select a **Plot type** and whether to **Interpolate** the data or not. With these calculation types you can choose to plot a *Heat map*, *Contour* or *3D* diagram. See "Plot Types" on page 126.

For a Property grid or Grid the X- and Y-axes always represent the quantities that define the calculation axes, but you need to set a variable for the Z-axis (the axis perpendicular to the plane defined by the X- and Y-axes).

- For a Property Model Calculator *Uncertainty* calculation type, you choose an **Axis** type: Frequency, Histogram or Normal probability.
- For other calculators (e.g. Equilibrium, Binary, and Scheil) there are different axis types available such as Linear, Logarithmic, Logarithmic 10, and Inverse.

# Settings

The options along the top of the window are available for most of the calculation types (e.g. phase diagrams, property diagrams, min/max, uncertainty and so forth).

## Tie lines

Select how many tie lines to be plotted in the diagram. Every  $n^{\text{th}}$  number of tie line is plotted, where n is the number set here. Consequently, the higher the number, the fewer the number of plotted tie lines in the diagram.



This setting is not available if the Plot Renderer is a successor to an Equilibrium Calculator where only one axis is defined and varied (that is, an Equilibrium Calculator that has performed a stepping calculation).

### Legend option

Select whether the diagram's legend displays the **Stable phases**, the **Axis quantity**, or choose **Off** for no legend.

### Legend style

You can select a Legend style for the **Stable phases** or **Axis quantity** selected for **Legend option**. Choose **None**, **All**, **Constitution description**, or **Ordering description**.

"About Legend Styles" on page 135

# Plot type (property grid or grid)



This option is available if the Plot Renderer is a successor to either an Equilibrium Calculator that is set up to calculate a *Property grid* or a Property Model Calculator that is calculating a *Grid*.

# "Plot Types" on page 126

- On a **Heat map** diagram, each Z-axis variable value in the grid is represented by a colour-coded pixel, where dark red represents the highest Z-axis variable value and dark blue represents the lowest Z-axis variable value.
- On a **Contour** diagram, Z-axis variable values are traced with curves in the same way that height values are traced with curves on a topographical map. The colors used originate from the different data series.
- On a **3D** diagram, the 3D surface is created by Delaunay tesselation of the calculated

data. The surface uses a cold to warm color map where red represents the highest Zaxis variable value and blue represents the lowest Z-axis variable value. The color map color legend can be turned on or off with the legend option.



#### Interpolate (property grid or grid)

From the **Interpolate** list, choose to interpolate data **Never** (the default), **Once**, **Twice** or **3 times**. This means points are put in between calculated points in the grid and interpolated instead of calculated in the calculation engine.

Choose how many points of data to interpolate on a grid or property grid to improve curve smoothness – **Never**, **20x20**, **50x50**, or **100x100**. Try different settings for your own set of data.

#### Label decimal digits

Choose the number of decimal digits to display in the plot label. The default is 2.

#### Label number format

Choose from Auto (the default), Decimal or Scientific.

### Axis variable

Set the state variable you want plotted along the X-axis and the Y-axis.



If the stepping/mapping variables are changed in the Equilibrium Calculator that precedes the Plot Renderer, then the variables to be plotted along the diagram's X- and Y-axis are automatically updated to the appropriate quantities.

#### Axis type

Select the type of axis scale: Linear, Logarithmic, Logarithmic 10 or Inverse.

#### Automatic scaling

The **Automatic scaling** check box is selected by default to allow the program to set the limits.

#### Limits

This setting is available when the **Automatic scaling** check box is cleared (the scaling is turned off).

Enter the lower and upper **Limits** of the variables to display on the axis. You can also determine the **step** size between the tick marks along each axis.

If the **Automatic scaling** is turned off (i.e. the check box is not selected) then it is important to enter a large enough number in the **step** field for the numerical values to display correctly on the axis. That is, if the step size is too small, the program ignores this setting and the axis labels will not be as expected.



8

For logarithmic axes, the step size is related to the exponent. For example, if the step entered is 0.1, then the exponent increases by one each time  $(10^{-9}, 10^{-8}, 10^{-7} \text{ and so forth})$ .

# Contour values and Use custom contour values (Z-axis, contour plots).

The **Use custom contour values** check box is available when **Contour** is selected as the **Plot type**. Select this check box to enter one or a series of numbers in the **Contour values** field. Numbers can only be separated by a space.

# Add an axis and Remove this axis buttons

Use the Add an axis <sup>(2)</sup> and Remove this axis <sup>(2)</sup> buttons to add additional X- and/or Y-axes to a plot or to merge plots (to overlay one plot with another in the same diagram). When merging plots, you define this variable as the Z-axis.

"Plotting Options" on page 132

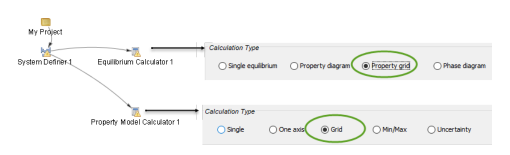
# Plot Types



The following examples use Graphical Mode.

# **Property Grid and Grid Calculations**

Using an **Equilibrium Calculator** (Property grid calculation) or **Property Model Calculator** (Grid calculation) you can choose to plot a *Heat map, Contour* or *3D* diagram.



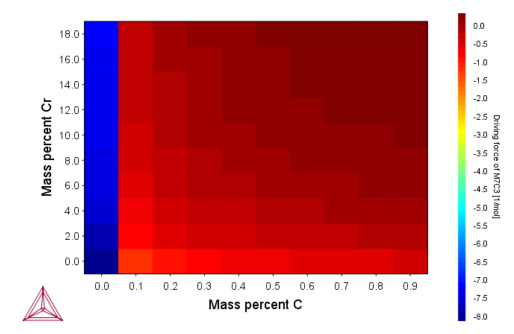
Equilibrium Calculator" on page 83 and "Property Model Calculator" on page 70

### HEAT MAP

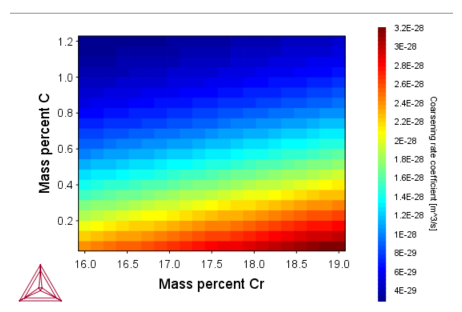
<u>"T\_09: Carbide Driving Force Heat Map" on page 23 and "T\_11: Coarsening and</u> Interfacial Energy" on page 26 in the *Graphical Mode Examples Guide*.

Each Z-axis variable value in the grid is represented by a colour-coded pixel, where dark red represents the highest Z-axis variable value and dark blue represents the lowest Z-axis variable value.





### Using a Property Model Calculator

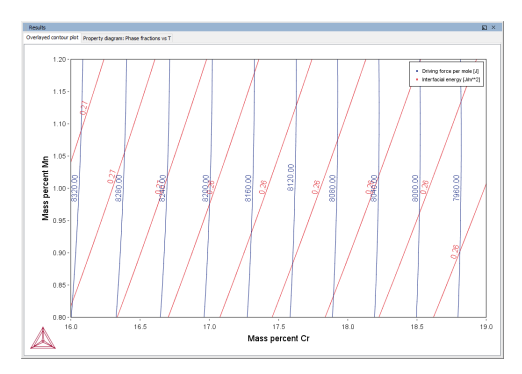


# CONTOUR

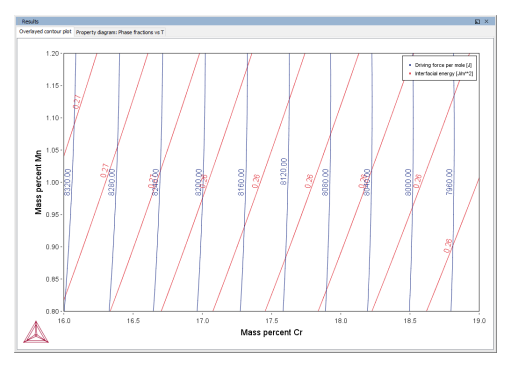


"T\_12: Driving Force and Interfacial Energy" on page 29 in the Graphical Mode Examples Guide.

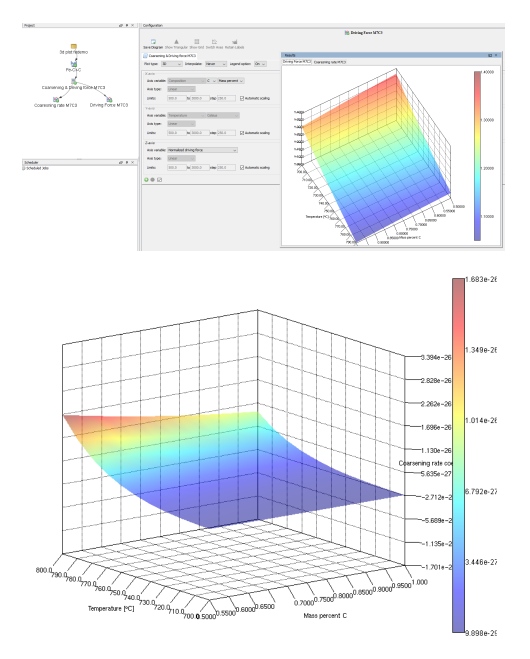
Z-axis variable values are traced with curves in the same way that height values are traced with curves on a topographical map. The colors used originate from the different data series.



Z-axis variable values are traced with curves in the same way that height values are traced with curves on a topographical map. The colors used originate from the different data series.



3D



"Rotating 3D Plots" on page 48

### Note for Linux OS Using VirtualBox

The following is applicable to users running Thermo-Calc under Linux in a virtual machine using VirtualBox.

Due to limitations in the OpenGL-drivers for VirtualBox, you will experience problems with 3D-plots unless the 3D-acceleration is disabled for the virtual machine being used.

To disable the 3D-acceleration:

- 1. Open the main VirtualBox application.
- 2. In the list, right-click on the virtual machine running Thermo-Calc.
- 3. Select Settings.
- 4. Navigate to **Display**.
- 5. On the Screen tab, click to clear the Enable 3D acceleration check box.



If you are using other virtualization software and experience problems with the 3D plot, a similar solution might apply.

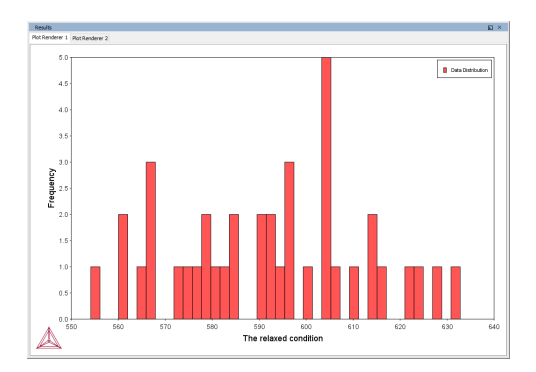
# **Uncertainty Calculations**

"T\_10: Phase Transition" on page 23 in the Graphical Mode Examples Guide

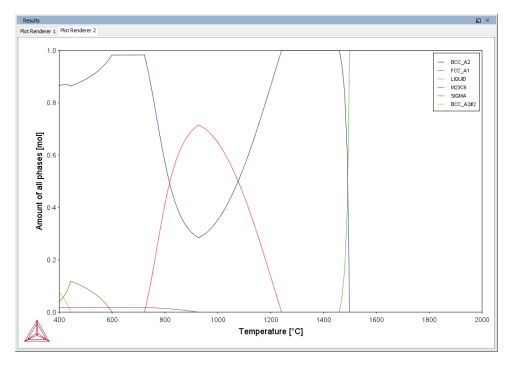
Using a **Property Model Calculator** (Uncertainty calculation) you can choose to create either a *histogram* or *probability* plot.

🥅 My Pròject				
HAL				
System Definer 1				
Prope	nty Model Calcula	tor 1		
Calculation Type				
◯ Single	One axis	⊖ Grid	◯ Min/Max	Uncertainty
Property N	Iodel Calculat	or" on pag	e 70	

HISTOGRAM



# PROBABILITY



# **Plotting Options**

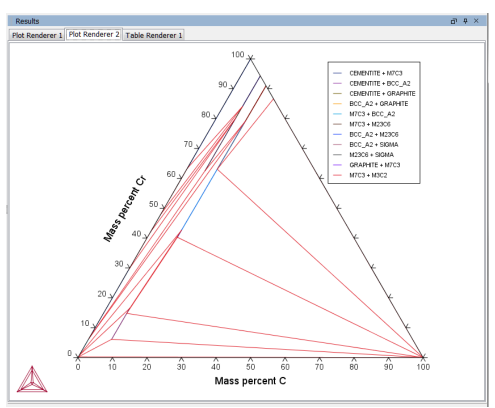
The <u>"Plot Renderer" on page 122</u> is a dynamic activity node with options such as being able to:

- add more than one Plot Renderer to its predecessor node (i.e. one of the calculator activities),
- add axes to the plots, and
- merge results in one plot.

### PLOTS CAN HAVE MORE THAN ONE PREDECESSOR

A Plot Renderer can have several predecessors (if you want to plot several calculations in the same diagram), in which case there are several tabs in the Configuration window, one for each predecessor.

Several Plot Renderer activities (as well as Table Renderer activities) can be shown in the Results window as different tabs. The figure is from Graphical Mode example T\_04.



# Plotting Several Calculations in One Diagram" on page 141

### ADD AXES TO PLOTS

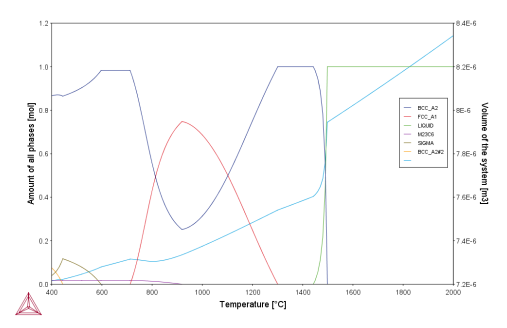
You can add additional X- and/or Y-axes to a plot to compare two or more variables at the

same time as in the figure.



"Adding Axes to a Diagram" on page 142

Two variables plotted on the Y-axis.



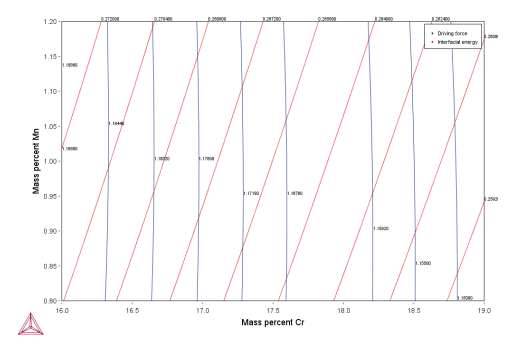
### MERGE RESULTS IN ONE PLOT

You can merge (or overlay) calculated results for several variables into one plot. This is similar to adding an axis, but instead of displaying different variables along the X- and/or Y- axes, the same X and Y variables are used to compare the third variable (defined in the Plot Renderer as the Z-axis).

As in the figure (from example T\_12 in Graphical Mode Examples), the driving force and interfacial energy results are combined into one diagram. The X and Y axis is the same for both (Mass percent Cr/Mass percent Mn).

"Merging Plots into One Diagram" on page 144

A contour plot showing both the driving force and interfacial energy calculation results comparing Mass percent Cr to Mass percent Mn.



# **About Legend Styles**

There are a variety of legend styles you can use to display detail about the phases. The same legend styles are available for both Graphical Mode and Console Mode plots.



For Console Mode, use the command CHANGE\_LEGEND.

In Graphical Mode the label styles are defined when using the <u>"Plot Renderer" on page 122</u>. You can select a **Legend style** for the **Stable phases** or **Axis quantity** selected for **Legend option**.

When there is a legend in the diagram you can append constitution and/or ordering descriptions to the phase name.

If constituent description is included in the legend, then for each equilibria along a line in the diagram, the largest constituent fractions are summed until they are equal or greater than 0.6. This is done separately for each sublattice. If the equilibria along the line gets different sets of constituents these are all appended to the list for that sublattice.

The ordering description is only printed for partitioned phases (phases that have a disordered contribution) that begins with at least two identical sublattices (sublattices where the set of constituents are identical). DISORD (disordered) is shown if site fractions are identical between identical constituents on all identical sub lattices (with an error margin) otherwise ORD (ordered) is shown in the legend.

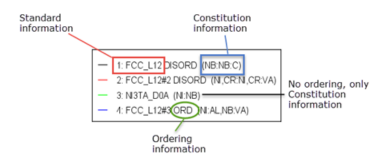
If ordering between equilibria in a line are different, then ORD + DISORD is shown in the legend.

# Legend and Label Styles

# Legend Styles

To show the differences between labels, the following uses the same example file that creates Figure 6 in <u>TCNi8 database extended information</u> document (found on the Thermo-Calc website).

When **All** is selected for this specific example the label includes the following:



Specifically:

- There are two disordered FCC phases (matrix and carbide), one ordered FCC (gamma prime), plus a phase NI3TA\_DOA (called Delta in Ni-alloys).
- ORDERING shows the different FCC phases.
- CONSTITUTION shows that FCC\_L12#1 is NbC, FCC\_L12#2 is matrix with Ni and Cr as major constituent, FCC\_L12#3 is Ni3Al with Nb addition and NI3TA\_D0A is Nb-rich (eg. Ni3Nb).

# Label Styles

These examples are for each label style choice - **None**, **All**, **Constitution** or **Ordering**. The instructions are for when you are setting the options using the Plot Renderer. In Console Mode, you choose the options (ALL, NONE, CONSTITUTION, or ORDERING) after the command CHANGE\_LEGEND.

### NONE

When **None** is selected the legend includes the standard information about the phases.

-	1: FCC_L12
-	2: FCC_L12#2
-	3: NI3TA_D0A
-	4: FCC_L12#3

# All

When you select **All** the **Constitution description** and the **Ordering description** (when there is ordering) are included.



— 3: NI3TA\_D0A (NI:NB)
 — 4: FCC\_L12#3 ORD (NI:AL,NB:VA)

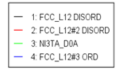
# **CONSTITUTION DESCRIPTION**

When you select **Constitution description**, this is added to the standard description.



## **ORDERING DESCRIPTION**

When you select **Ordering description** and when there is ordering, this is added to the standard description.



# **Plot Labels**

To add a label text at a certain point in a plot, right click the point, and select **Add Label**. For phase diagrams, the program suggests a label that you can keep or edit. The suggested label specifies the names of the stable phases at the coordinate where you clicked. For all other plot types, you enter your own text, which can either be Plain text or LaTeX formatted text.

🔺 Add Label	X
Label: Font: AaBbCc123 AαBβΓγ Font: AaBbCc123 AαBβΓγ LaTeX-formatted text	-
Rotation angle: 0.0	
Show details OK Cancel Reset Delete	

You can enter the label as **Plain text** or in **LaTeX-formatted text**. For **LaTeX-formatted text** enter or edit the text as required. For example, underscores (\_) are added before the numbers to make subscripts in the label.

4

For a list of supported LaTeX commands, see: http://www2.ph.ed.ac.uk/snuggletex/documentation/math-mode.html.

By default when using LaTeX, you are in math mode and Thermo-Calc automatically interprets the entered text as math. The above link uses the dollar sign (\$) in all the examples but this is NOT required. For example, if you enter \$E=mc^2\$ this is displayed including the \$ instead of the equation.

#### Plot Labels: Entering a Mixture of Plain Text and Equations

When you enter text using LaTaX formatted text it displays by default on the plot in italics and treats the text as an equation.

As in the following example, the top label does not display correctly because the label is a mixture of text (Mole-Fraction) and an equation  $(Al_2O_3)$ . In the bottom image, when the same information is added as Plain text to the plot it now does not include the subscripts.

[	$Mole$ - $FractionAl_2O_3$	
🔺 Edit La	bel	<b>—</b> X
Label:	Mole-Fraction Al_20_3	LaTeX-formatted text V
Mo	ole-fraction AI_20_3	
🛕 Edit Lab	el	
_	ole-fraction Al_20_3 aBbCc123 ΑαΒβΓγ	ັ ຊິ Plain text Modify

To add a mixture of plain text and equation text, you can exit math mode using the string \textrm followed by the text to display enclosed in curly brackets. For example \textrm {Mole-fraction }Al 20 3 results in the label as shown:

		Mole-fraction $Al_2O_3$		
ſ	🛕 Edit Lab	el		×
1	Label:	\textrm{Mole-fraction }Al_20_3	• III	LaTeX-formatted text



In this example, you need to add a space at the end of fraction in order to separate the text from the equation part of the label.



When working in Console Mode, you can also use the above windows on the Console Results window to add and edit labels. In addition, you can use the ADD\_ LABEL\_TEXT command described in the *Thermo-Calc Console Mode Command Reference*.

### Plot Labels: Rotation Angle and Show Anchor

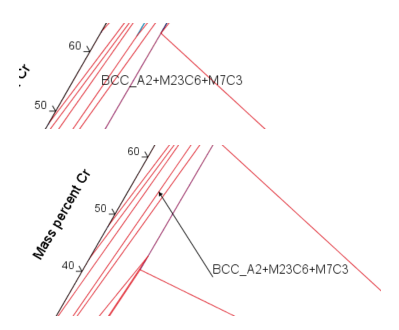
By default the **Show anchor** check box is selected, which includes an arrow from the label point to the coordinates.

If you do not want the label oriented horizontally in the Plot Renderer tab, then specify the number of degrees of rotation in the **Rotation angle** field.

Click **OK** to add the label to the plot. You can also move a label by clicking and dragging it. The default anchor arrow is added to indicate the point that belongs to the label.

### Moving Plot Labels

Drag and drop labels anywhere on the plot. The anchor arrow points to the point of origin.

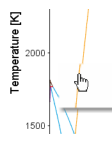


**Changing the Plot Properties** 

For global changes to the default settings plots see <u>"Global Settings: Graphical and Console</u> <u>Mode Plotting" on page 247</u>. The settings are the same but applied to all new plots.

To make local changes to the appearance of a specific plot, in the **Results** window right-click a plot and select **Properties**. You can also edit some properties for individual plot lines (the color, the line width and type, and whether data points are included).

In the **Results** window, hover the mouse over a plot line. When it is over a line that can clicked the crosshair cursor turns into a pointing finger.

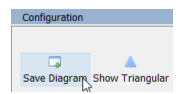


To prevent unintended edits, hold down <Ctrl> while moving the cursor around the plot to only display it as a crosshair.

# Saving Plots

8

1. To save a plot, at the top of the **Configuration** window click **Save Diagram**.



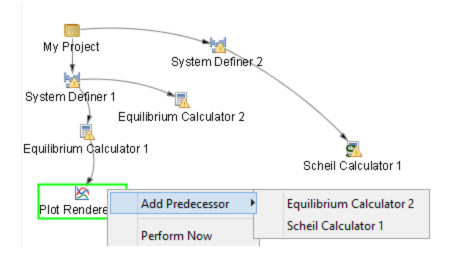
Or right-click the diagram in the Results window and select Save As.

2. In the **Save** window, navigate to where you want to save the plot, enter a **File name**, and choose the **File type** to save it in. Click **Save**.

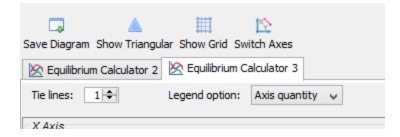
# Plotting Several Calculations in One Diagram

To plot several calculations in one diagram, you must have, in addition to the Plot Renderer's original predecessor, at least one available Plot Renderer predecessor in your project, for example, the additional predecessor can be an Equilibrium Calculator, Binary Calculator, Ternary Calculator, Scheil Calculator or Experimental File Reader.

- 1. In the Project window click the **Plot Renderer** node.
- 2. In the **Configuration** window click **Add Predecessor**. Or right-click the **Plot Renderer** and choose **Add Predecessor**.

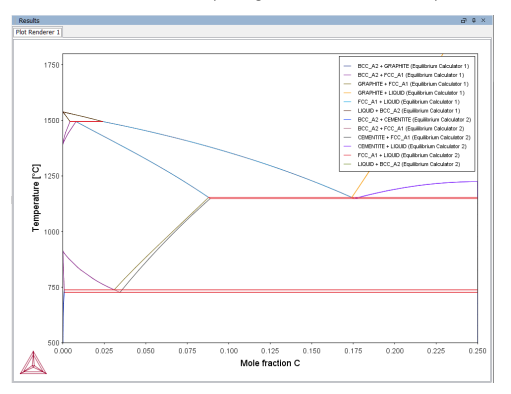


3. Select a predecessor. When added, the extra predecessor is linked to the Plot Renderer. On the Plot Renderer's Configuration window, the predecessors are shown as two tabs:



4. Configure how you want each calculation plotted on the tabs in the **Configuration** window, then click to **Perform** the Plot Renderer activity.

The calculations are plotted in the same diagram in a tab in the **Results** window. If any of the axis variables of the plots differ, then the label and scale of all calculations are shown parallel to each other. The following screen shot shows a diagram in which two calculations with different plotting axes variables have been plotted.



# Adding Axes to a Diagram

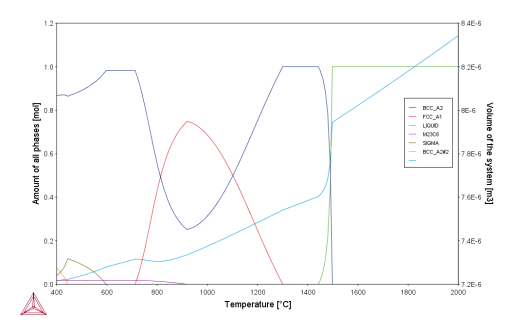
You can add additional X- and/or Y-axes to a plot to compare two or more variables at the same time.

- 1. In the **Project** window right-click the calculator node (e.g. an Equilibrium Calculator) and add a **Plot Renderer** node from the **Create New Successor** menu.
- 2. Click the **Plot Renderer** node.

- 3. In the **Configuration** window, define the plot. Available settings are based on the calculation type.
- 4. Click the **Add an axis** S button. A new settings section is added underneath.
- 5. From the *X* Axis and/or *Y* Axis sections define another **Axis variable** and choose an **Axis type**.
- 6. Click **Perform** to plot both variables on the selected axis, in this example, the Y-axis.

In this example **Amount of phase** and **Volume** are selected as the Y-axis variables and **Temperature** as the X-axis variable. The plot is an example of the output.

X AXIS			
Axis variable:	Temperature	~	Celsius 🗸
Axis type:	Linear 🗸		
Limits:	500.0 to 3000.0	step 250.0	Automatic scaling
Y Axis			
Axis variable:	Amount of phase	~	No normalization $\checkmark$ All phases $\checkmark$ Mole $\checkmark$
Axis type:	Linear 🗸		
Limits:	0.0 to 1.0	step 0.1	✓ Automatic scaling
0 😑 🗹			
X Axis			
Axis variable:	Temperature	$\sim$	Celsius 🗸
Axis type:	Linear 🗸		
Limits:	500.0 to 3000.0	step 250.0	Automatic scaling
Y Axis			
Axis variable:	Volume	~	No normalization V SER System V Cubic meter
Axis type:	Linear 🗸		
Limits:	0.0 to 1.0	step 0.1	Automatic scaling



# Merging Plots into One Diagram

You can merge (or overlay) calculated results for several variables into one diagram. The same X and Y variables are used to compare the third variable (defined in the Plot Renderer as the Z-axis).

You cannot overlay heat map or frequency plots.

The following uses example T\_12 in Graphical Mode Examples to describe the steps to overlay two plots: the driving force and interfacial energy results are combined into one diagram. The X and Y axis is the same for both (Mass percent Cr/Mass percent Mn).

- 1. In the **Project** window right-click the calculator node (e.g. Property Model Calculator) and add a **Plot Renderer** node from the **Create New Successor** menu.
- 2. Click the Plot Renderer node.
- 3. In the **Configuration** window, define the plot. Available settings are based on the calculation type.

### EXAMPLE

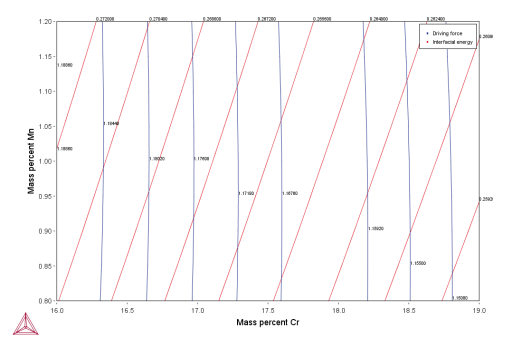
In this example the Property Model Calculator is set to a Grid calculation type.

Choose On for the Legend option and Contour as the Plot type.

0 🗢 🔽				
X Axis				
Axis variable:	Mass percent Mn		Ŧ	
Axis type:	Linear	Ŧ		
Limits:	0.0	to 100.0	step 10.0	Automatic scaling
Y Axis				
Axis variable:	Temperature	3	▼ Celsius	•
Axis type:	Linear	Ŧ		
Limits:	500.0	to 3000.0	step 250.0	Automatic scaling
Z Axis				
Axis variable:	Driving force			
Axis type:	Driving force Interfacial energy			
Limits:	0.0	to 1.0	step 0.1	Automatic scaling
0 🗢 🔽				

- 4. Click the **Add an axis** Dutton. A new axes settings section is added underneath.
- 5. From the Z Axis section choose the Axis variable. In this example, Interfacial energy.
- 6. Click **Perform** to plot both on the same diagram.

In this case, a contour plot showing both the driving force and interfacial energy calculation results comparing Mass percent Cr to Mass percent Mn.



# **Results Window Status Marker**

If you have plotted a diagram and then modify something in the tree that the Plot Renderer is associated with, then a notification —an exclamation mark enclosed in a yellow triangle—is shown in the Plot tab.

	"Node Status Markers" on page 47
Results Plot Rer	ndere 🔺 Plot Renderer 2

# **Table Renderer**

A Table Renderer visualizes calculation results in a table. Several Table Renderer activities (as well as Plot Renderer activities) can be shown in the Results window as different tabs.

You can only successfully tabulate the results of a single-equilibrium calculation or a stepping operation. The tabulated data from an equilibrium calculation is different from the tabulated data from a stepping calculation.

"Creating Activities and Successors" on page 32

# **RESULTS FROM AN EQUILIBRIUM CALCULATION**

The following tab shows the result of an equilibrium calculation (from example <u>"T\_01:</u> Calculating a Single-Point Equilibrium" on page 4 in the *Graphical Mode Examples Guide*).

Table Renderer 1				
Custom				
System	1 00000			
Moles	1.00000	r_1		
Mass	55.64392	[g]		
Temperature	1000.00000	[K]		
Total Gibbs Energy				
Enthalpy	24707.16903	[1]		
Volume	7.29746E-6	[m3]		
Component	Mole Fraction	Mass Fraction	Activity	Potentia/
Fe	0.99537	0.99900		-42277.77189
C	0.00463	0.00100		-12658.35450
C	0.00403	0.00100	0.21010	-12038.33430
Stable Phases				
	Moles	Mass	Volume Fraction	
BCC_A2#1	0.99609	55.59691	0.99711	Composition and constitution -
Composition				
Component	Mole Fraction	Mass Fraction		
Fe	0.99928	0.99984		
с	0.00072	0.00016		
Constitution	(Fe) <sub>1</sub> (C,VA) <sub>3</sub>			
	Constituent	Site Fraction		
Sublattice 1:				
	Fe	1.00000		
Sublattice 2:				
	VA	0.99976		
	С	0.00024		

For each stable phase listed, from the **Composition and constitution** menu you can select the information to view. In this example, the maximum amount of information is shown and includes details about both composition and constitution for **BCC\_A2#1**.

The default for the Phase description detail can be changed globally. To open the Options window: From the main menu select **Tools**  $\rightarrow$  **Options** and click the **Activities** tab. Then click **Tabulation**.

### **RESULTS FROM A STEPPING OPERATION**

The **Table Render 1** tab in the example shows the results of a stepping calculation (from example <u>"T\_02: Stepping in Temperature in the Fe-C System" on page 6</u> in the the *Graphical Mode Examples Guide*).

Results  Plot Renderer 1 Table Renderer 1				
Temperature [K]	Amount of BCC_A2 [mol]	Amount of LIQUID [mol]	Amount of FCC_A1 [mol]	Amount of GRAPHITE [mol]
1000.00000	0.99609			0.00391
1010.00000	0.99618			0.00382
1011.17278	0.99620			0.00380
1011.17278	0.99620		0.00000E0	0.00380
1011.17278	0.87364		0.12636	0.00000E0
1011.17278	0.87364		0.12636	
1020.00000	0.86046		0.13954	
1030.00000	0.84331		0.15669	
1040.00000	0.82356		0.17644	
1050.00000	0.80082		0.19918	
1060.00000	0.77395		0.22605	
1070.00000	0.74180		0.25820	

# **Configuration Settings**

In the **Configuration** window for the Table Renderer 1 node, set the level of detail for the Columns.

Table Renderer 1
Number format: Auto 🗸
V Kelvin V
✓ No normalization ✓ All phases ✓ Mole ✓

# Setting Number Format and Number of Decimal Digits

The table results can be presented in ordinary decimal number format or with normalized scientific notion. The **Number format** defaults to **Auto** to use the most appropriate format based on the tabulated data. Or choose **Scientific** or **Decimal** to increase or decrease the number of **Decimal digits** used in the table. The default is 5 digits.



The defaults for these settings can be changed globally. From the main menu select **Tools**  $\rightarrow$  **Options** and click the **Activities** tab. Then click **Tabulation**.

# Saving Tabulated Data

To save all the data in a table as a plain text file, in the **Configuration** window click **Save table** or in the **Results** window right-click in the table and select Save As.

To copy the data from a single cell of a table to the clipboard, right-click the cell and select **Copy**. To copy the whole table to the clipboard, select **Copy all**.



The data saved is what is shown in the **Table Renderer** tab. If data is calculated but not displayed it is not saved.

### Setting Background Colours

To change the background colours of the table, in the **Results** window right-click a Table Renderer tab, and select **Properties**. Modify as required.

Table Properties	
Primary display color:	Modify
Secondary display color:	Modify
	OK Cancel Reset

The defaults for these settings can be changed globally. To open the Options window: From the main menu select **Tools**  $\rightarrow$  **Options** and click the **Activities** tab. Then click **Tabulation**.

# **Experimental File Reader**

The Experimental File Reader activity can only be created from the top My Project node. This activity allows you to read an experimental data file (\*.EXP). This type of file has information specifying a plotted diagram, written in the DATAPLOT graphical language. From this activity node you can plot the EXP file or save a copy of the file to another location.

"Creating Activities and Successors" on page 32 and "Plotting Experimental Data Files" on page 4 in the DATPLOT User Guide included with this documentation set.

### PLOTTING AN EXPERIMENTAL DATA FILE

- 1. Right-click **My Project** and from the **Create New Activity** submenu choose **Experimental File Reader**.
- 2. In the **Configuration** window in the **EXP file** field, enter the name of the EXP file to plot. Or click the **Select Exp file** button to navigate to a location on the computer.
- 3. Right-click the **Experimental File Reader** node and from the **Create New Successor** submenu choose **Plot Renderer**.

If the source EXP file is changed, you can also click the **Reload the selected EXP file from disk** button to refresh the data. If you want to save or export the loaded file, click the **Save EXP file** button, then navigate to where you want to save it and give it a file name. 4. In the **Configuration** window click **Perform** or right-click the **Plot Renderer** node and select **Perform**.

# **Binary Calculator**

A Binary Calculator can be used for some calculations involving two components. You can think of this activity as a combination of a System Definer and an Equilibrium Calculator, but designed to simplify setting up and performing calculations on binary systems.

The Binary Calculator relies on some specifications that are not supported by all databases. You need a specially designed database for the Binary Calculator such as the TCBIN database.

For an example of how a Binary Calculator can be used in a project, see example <u>"T\_03: Fe-C Phase Diagrams" on page 8</u> in the *Graphical Mode Examples Guide*.

### **Configuration Settings**

The table lists the available settings to configure before performing the calculation and the image shows the **Configuration** window of a **Binary Calculator** activity.

"Creating Activities and Successors" on page 32

Config	uratio	n																		
									2	Binar	y Cal	culato	r 1							
Databa					_															
TCBIN	TCBIN																			
Elemer	Elements Phases and Phase Constitution Data Sources Description																			
																		1		
									ZE	VA										
Н		1								-				_			He			
Li	Be											В	С	Ν	0	F	Ne			
Na	Mg											AI	Si	Р	S	СІ	Ar			
K	Са	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr			
Rb	Sr	Y	Zr	Nb	Мо	Тс	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Те	H	Xe			
		 	Hf					$\square$	$\square$				Pb				님			
Cs	Ba			Та	W	Re	Os	lr	Pt	Au	Hg	TI		Bi	Po	At	Rn			
Fr	Ra	**	Rf	Db	Sg	Bh	Hs	Mt	Ds	Rg	Cn	Uut	FI	Uup	Lv	Uus	Uuo			
• Lan	thanide	series	La	Ce	Pr	Nd	Pm	Sm	Eu	Gd	Tb	Dy	Ho	Er	Tm	УЬ	Lu			
									$\square$								$\square$			
A	ctinide s	series	Ac	Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr			
Calcul	ation	Туре																		
0 P	hase	diagra	am																	
() G	ibbs e	energ	y curv	/es	Тетр	eratu	re Ke	lvin		-	1000	).0								
_							_			_	1000									
A	ctivity	curv	es		remp	eratu	re Ke	ivin		*	1000	.0								
() P	Phase fractions Mole fraction 0.0																			
																				٦l
	0					Add	Prede	cesso	r	Calo	ulate	Phas	e Diag	jram		Create	e New	Successor	>	

Setting	Description
Select database	Select a database that contains specifications dedicated to the Binary Calculator.
Elements (periodic table)	In the periodic table, select the two elements that make up the components of the system.
Phases and Phase Constitution	<ul> <li>This tab is the same as for a System Definer. See <i>Phases</i> and <i>Phase Constitution</i> for an example.</li> <li><u>"System Definer" on page 62</u></li> </ul>
Data Sources	This tab is the same as for a System Definer. See <i>Data Sources</i> for an example.

Setting	Description
Description	This tab is the same as for a System Definer. See <i>Description</i> for an example.
Calculation Type	
Phase diagram	Select to perform a mapping calculation. Click <b>Calculate</b> <b>Phase Diagram</b> to plot the results.
Gibbs energy curves	Select the <b>Temperature</b> and click <b>Calculate Gibbs Energy</b> <b>Curves</b> to calculate at a constant temperature over the whole composition range.
Activity curves	Select the <b>Temperature</b> and click <b>Calculate Activity</b> <b>Curves</b> to calculate at a constant temperature over the whole composition range.
Phase fractions	Select the <b>Mole fraction</b> and click <b>Calculate Phase</b> <b>Fractions</b> to calculate as a function of temperature at a constant composition.

# **Ternary Calculator**

A Ternary Calculator is used for some calculations involving three components. You can think of this activity as a combination of a System Definer and an Equilibrium Calculator, but designed to simplify setting up and performing calculations on ternary systems.



For an example of how a Ternary Calculator is used in a project, see example <u>"T\_04: Fe-Cr-C Ternary Phase Diagram at 1000 K" on page 10</u> in the *Graphical Mode Examples Guide*.

"Creating Activities and Successors" on page 32

### **Configuration Settings**

The table lists the available settings that must be configured before performing the calculation and the image shows the Configuration window of a Ternary Calculator activity.

onfigu	uratio	n																			L
									3	Terna	ry Ca	lculate	or 1								
ataba					_																
CFE8	3				•																
emen	its P	hases	and	Phase	e Con	stituti	on D	ata S	ource	s De	script	ion									
							_											1			
									ZE	VA											
Н										-							He				
									′								님				
Li	Be											В	С	N	<u> </u>	F	Ne				
Na	Mg											AI	Si	Р	S	СІ	Ar				
к	Са	So	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr				
Rb	Sr	Y	Zr	Nb	Мо	Тс	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Te	-	Xe				
Cs	Ва	*	Hf	Та	w	Re	Os	Ir	Pt	Au	Hg	ті	Pb	Ві	Po	At	Rn				
Fr	Ra	**	Rf	Db	Sg	Bh	Hs	Mt	Ds	Rg	Cn	Uut	FI	Uup	Lv	Uus	Uuo				
• Lani	thanide	series	La	Ce	Pr	Nd	Pm	Sm	Eu	Gd	ть	Dy	Ho	Er	Tm	Yb	Lu				
Ac	ctinide s	eries	Ac	Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr				
alcula	ation	Туре																			
I	sother	mal s	ection	n To	empe	rature	Kel	in		•	100	0.0									
_																					
M	lonov	ariant	lines																		
O Li	iquidu	is pro	jectio	n Te	empe	rature	Keh	/in			Mi	in 298	3.15		Max	3273	.15	In	terval	100.0	
					1						-									_1	
		0				<	Add	Prede	cesso	or [	Per	form <sup>·</sup>	Tree		Create	New	Succe	ssor	>		

#### Select database

Select a database that contains specifications dedicated to the Ternary Calculator.

#### Elements (periodic table)

In the periodic table, select the three elements that make up the components of the system.

#### **Phases and Phase Constitution**

This tab is the same as for a <u>"System Definer" on page 62</u>. See *Phases and Phase Constitution* for an example.

#### **Data Sources**

This tab is the same as for a <u>"System Definer" on page 62</u>. See *Data Sources* for an example.

#### Description

This tab is the same as for a <u>"System Definer" on page 62</u>. See *Description* for an example.

### **Calculation Types**

#### **Isothermal section**

Select the **Temperature** at which the ternary system is calculated for the whole composition range.

### **Monovariant lines**

The variation of the liquid compositions with temperature is calculated.

### Liquidus projection

Select the **Temperature** range (**Min** and **Max**) and an **Interval**. The projection liquid surface and the monovariant lines are calculated over the given temperature range.

# **Scheil Calculator**

The Scheil Calculator activity is used to perform simulations of *Scheil (Scheil-Gulliver) solidification processes*. By default, a Scheil calculation results in an estimation of the solidification range of an alloy. The calculation is based on the assumption that the liquid phase is homogeneous at all times and that the diffusivity is zero in the solid. However, it is possible to disregard the latter assumption for selected components.

By default, a Plot Renderer successor has the default plot axes **Mole fraction of solid** (Xaxis) and **Temperature Celsius** (Y-axis). When these default axes are used in the Plot Renderer, the results of normal equilibria calculations are automatically plotted in the same diagram as the Scheil calculations.

- "Creating Activities and Successors" on page 32 and "Scheil Simulations" on page 196 in the Console Mode section.
  - See example <u>"T\_08: Scheil and Equilibrium Solidification" on page 20</u> in the *Graphical Mode Examples Guide*.

### **Configuration Settings**

The main **Configuration** window of a **Scheil Calculator** activity is shown below and listed in the table. These settings must be configured before performing the calculation.

Configuration		
	😴 Scheil Calculator 1	
Start temperature:	2500.0	<ul> <li>Show advanced options</li> </ul>
Temperature step:	1.0	
Temperature unit:	Kelvin 🔻	
Composition unit:	Mass percent 🔹	
Composition C	99.99 Tast diffuser	
Composition Fe	0.01 Fast diffuser	
	< Add Predecessor Perform Tree Cr	eate New Successor >

#### Start temperature

Enter a **Start temperature** that is higher than the liquidus temperature of the alloy, in other words, the temperature at which the alloy is completely melted.

#### **Temperature step**

Enter a **Temperature step**. Decreasing the temperature step increases the accuracy, but the default value is usually adequate.

#### **Temperature unit**

Select the Temperature unit: Kelvin, Celsius or Fahrenheit.

#### **Composition unit**

Select the Composition unit: Mass percent, Mole percent, Mass fraction or Mole fraction.

#### Composition

This is based on what is entered in the "System Definer" on page 62.

#### Fast diffuser

Click to select the **Fast diffuser** check box to allow redistribution of this component in both the solid and liquid parts of the alloy.

#### **Advanced Options and Settings**

Additional advanced settings are available as shown below and in the table:

Configuration		
	Scheil Calculator 2	
Start temperature:	2500.0	<ul> <li>Hide advanced options</li> </ul>
Temperature step:	1.0	
Temperature unit:	Kelvin 🔻	
Composition unit:	Mass percent 🔻	
Global minimization		
$\square \text{ Allow BCC} \rightarrow \text{FCC}$		
Liquid phase:	LIQUID	
Terminate on:	Fraction of liquid 🔻 at: 0.01	
Composition Fe 9	9.99 🔲 Fast diffuser	
Composition C 0	.01 Fast diffuser	
<u> </u>	Add Predecessor Perform Tree	Create New Successor >

Click Show advanced options to open this configuration window.

#### **Global minimization**

By default the **Global minimization** check box is not selected. Click to select it to perform a global minimization test when an equilibrium is reached. This costs more computer time but the calculations are more robust.

#### Allow BCC→FCC

By default the **Allow BCC**  $\rightarrow$  **FCC** check box is not selected. Select the check box to allow transformations in the solidified part of the alloy caused by each of the components specified to be a *fast diffuser*. It is recommended that you only select this for steels.

#### Liquid phase

The default in the list is the Liquid phase already defined in the database.

#### **Terminate on**

Enter a value to terminate the calculation on the **Fraction of liquid phase** or at a specific **Temperature**.

#### Max no. of iterations

By default, the program tries 500 iterations before it gives up. As some models give computation times of more than 1 CPU second/iteration, this number is also used to check the CPU time and the calculation stops if 500 CPU seconds/iterations are used.

#### **Required accuracy**

The default is 1.0E-6. This is a relative accuracy, and the program requires that the relative difference in each variable must be lower than this value before it has converged. A larger value normally means fewer iterations but less accurate solutions. The value should be at least one order of magnitude larger than the machine precision.

#### **Smallest fraction**

The default is 1.0E-12. This is the value assigned to constituents that are unstable. It is normally only in the gas phase you can find such low fractions.

The default value for the smallest site-fractions is 1E-12 for all phases except for IDEAL phase with one sublattice site (such as the GAS mixture phase in many databases) for which the default value is always as 1E-30.



In Console Mode, the default of 1E-30 is true unless you are using the command SET\_NUMERICAL\_LIMITS to reset an even-lower value (e.g. 1E-45, that is naturally enforced to all the phases in the system.

#### Approximate driving force for metastable phases

The check box is selected by default, which means the metastable phases are included in all iterations. However, these may not have reached their most favourable composition and thus their driving forces may be only approximate.

If it is important that these driving forces are correct, click to clear the check box to force the calculation to converge for the metastable phases.

# The Console Mode Interface

The following is an introduction to the Console Mode's user interface and the command line prompt.



Also see <u>"The Graphical Mode Interface" on page 29</u> for information about that mode.

### In this section:

The Console Mode Layout	
Main Menu Commands	
Keyboard Shortcuts	
Create New Console or Result Tabs	
Navigate Between Tabs on Windows	
Pause, Resume or Stop Output to the Console Window	
Console Window	
Editing the Default Console Directory	164
Console Mode Results Window	
Event Log Window	
Console Mode Workflow	
Performing Calculations in Console Mode	
Using the Command Line Prompt	
Moving Between Modules and Submodules	
Entering Command Names or Abbreviations	
Specifying Parameters	
Wild Card Characters	171
Command History	173
Macro, Workspace and Experimental Data Files	
Log Files	173

Macro Files	174
Workspace Files	177
Increasing the Buffer Size	178
Defining a System in Console Mode	

# The Console Mode Layout

By default the first time you launch Thermo-Calc it opens in Graphical Mode. After this, the mode defaults to the most recent one. You can switch back and forth between modes. To open the Console Mode interface, click **Switch to Console Mode** on the toolbar.

### THE CONSOLE MODE USER INTERFACE

File Tools Window Help			
New Open Save Switch to Graphical Mode			
Console	9 t ×	Console Results	급 후 ×
Console 1 💿 🔲 🗄		🖄 Results Console 1	
Calculated 66 e	equilibria	Plot 1 🕥 💷 😑	
Phase region boundary 21 at: 6.8 LIQUID#1 ** BCC A2#1	76E-01 2.042E+03	EX 2015.07.27.10.42.33 PTERN: CR, FE P=1.01325E5, N=1	cample 4a
Calculated 67 e	equilibria		
Phase region boundary 22 at: 6.8 LIQUID#1 ** BCC A2#1	76E-01 2.042E+03	2000 -	
Calculated 35	equilibria	_ 1800	
Phase region boundary 23 at: 9.90 LIQUID#1 ** BCC A2#1	06E-01 2.178E+03		
	equilibria	H H 1400	
Phase region boundary 24 at: 9.90 LIQUID#1 ** BCC A2#1	06E-01 2.178E+03		
Calculated 15 e	equilibria	U 1000	
C:\Users\Public\Documents\Thermo-Calc .POLY3	·	800-	
CPU time for mapping	2 seconds	600	
Event Log			- a ×
09:10:59,431 INFO Initialization complete 10:42:21,815 INFO Dragging and dropping a		\2015a\Examples\Console-mode\toex04\toex04.TC	3

The currently selected window is called the *active* window. This has a darker title bar than the other windows. The windows are:

- 1. Console, with a Console 1 tab.
- 2. **Console Results**, with a **Results Console 1** tab as well as a sub-tab (**Plot 1**), which is what is displayed in the window in the figure.
- 3. Event Log, which is the active window in this example.

### **CONSOLE TABS**

By default, the tabs are side-by-side:

Console			
Console 1 ×	Console 2	Console 3	○ □ =

You can toggle to view the tabs top-to-bottom/stacked (click the  $\square$  button) and back to sideby-side (click the  $\square$  button) or the  $\blacksquare$  button.

#### **OPEN CONSOLE WINDOWS**

Open a specific window from the **Window** main menu, or select **Reset Windows** to return to the default window layout.

File Tools Wi	ndow Help
New Open	Project Scheduler
Console	Configuration
Console	Results
INTEGRAI	Console Console Results
Integrat	Event Log
POST:help	Reset Windows

# **Main Menu Commands**



In Console Mode you use commands to access these menu options. If there is an equivalent command, it is listed below.



The **File** menu and toolbar are reserved for Graphical Mode. The **Tools**, **Window** and **Help** menus are described in <u>"Main Menu, Toolbar and Keyboard</u> Shortcuts" on page 35 in the *Thermo-Calc User Guide*.

#### **OPEN FILES**

- In the SYS module: <u>"MACRO\_FILE\_OPEN"</u> on page 5
- In the SYS module: <u>"OPEN\_FILE" on page 298</u>
- "Opening Console Mode Macro Files" on page 9

Drag and drop files into the Console window to open or run a macro, for example. File types that can be opened this way are macro files (\*.TCM and \*.DCM), database files (\*.TDB), and workspace files (\*.POLY3, \*.GES5,\*.PAR, and \*.DIC). Note that \*.DCM and \*.DIC file extensions are applicable to the Diffusion Module (DICTRA).

### SAVE (WORKSPACE)

This command depends on the module you are in. Type ? at the prompt to look for a command such as "SAVE\_GES\_WORKSPACE" on page 122 (GIBBS module).

#### Εχιτ

In any module: EXIT

### CLOSE

In the SYS module: "CLOSE\_FILE" on page 297

### **DISPLAY LICENSE INFO**

In the SYS module: "DISPLAY\_LICENSE\_INFO" on page 297

### **Keyboard Shortcuts**

### **Create New Console or Result Tabs**

- Create a new Console tab: <CTRL + T>
- Create a new Result tab: <CTRL+SHIFT + T>

#### Navigate Between Tabs on Windows

Use keyboard shortcuts to move between the tabs available in the Console and Results windows

#### WINDOWS

Press CTRL + TAB to move forward sequentially through each open tab. When you
are focused on the Console window, you move forward simultaneously through the
Console Results tabs, e.g. If you are on Console 1 and press CTRL + TAB, you move
forward to Console 2 and Results Console 2 and so forth.

Console		_	0 4 ×	Console Results		
Console 1	Console 2 ×	Console 3		Results Console 1	🖄 Results Console 2	🖄 Results Console 3

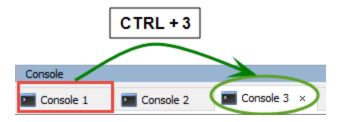
- Press CTRL + SHIFT + TAB to move back through the open tabs. This is useful if you
  have multiple tabs open, otherwise you can use the CTRL + TAB to cycle through the
  windows.
- Press CTRL + [tab number 1 to 9] to move directly to the numbered tab. For example,

if you are focused on Console 1 and want to move to Console 3, press CTRL+3 and the focus jumps to that tab.



The tab number corresponds to the order of the tabs, not necessarily the name. That is, if Console 4 is first, then it is what is highlighted when CTRL + 1 is used.

For **EXAMPLE**, press CTRL + 3 on the keyboard to move focus from Console 1 to Console 3.





When you are focused on the **Console Results** window and use any of the shortcuts this only moves you through the open tabs on this window.

#### LINUX AND MAC

The same **Windows** keyboard shortcuts are used but must be pressed twice to move forward and backward.

### Pause, Resume or Stop Output to the Console Window

In response to a command or a lengthy calculation, Thermo-Calc may display a lot of text in the Console window. Use these keyboard shortcuts to pause, resume or stop a process.

- To pause the text on printing press <Ctrl+S>.
- To resume the printing on screen, press <Ctrl+Q>.
- When Thermo-Calc is performing a stepping or mapping operation, the results are continuously printed in the Console window. To stop the calculation of the current region of the mapping and stop the output, press <Ctrl-A> (Windows) or <Ctrl-C> (Linux).

### **Console Window**

By default the Console window is on the left-hand side of the console. This is where you enter commands. The window can have several Console tabs.

To open and close Console tabs:

- To open (or add), click the S button, press <Ctrl+T> when the Console window is active, or right-click a tab header and select **New Console**.
- To close, click the button, press <Ctrl+W> or <Ctrl+F4>, or right-click the tab header and select Close.
- To close all tabs except one, right-click the header of the tab you want to keep and select **Close Other**.
- To close all tabs and open a new one, right-click the header of any tab and select **Close All**.

You can change the tab name, font and colours for an individual Console tab by right-clicking its header and selecting **Properties**.

Console Properties					
Console name:	Console 2				
Buffer size:	40,000 🚔				
System output font:	АаВЬСс123 АаВβГү				
Command prompt font:	AaBbCc123 ΑαΒβΓγ				
User input font:	AaBbCc123 ΑαΒβΓγ				
Background color:					
Default directory:	C:\Users\amanda\Documents				

# **Editing the Default Console Directory**

You can add a startup parameter to a shortcut when launching Thermo-Calc on any of the platforms. When you launch the revised shortcut, the file path is set to a user-defined default directory. An example of why you might want to have a custom directory path is if you use batch projects or if you have macro files for batch jobs. The custom directory file path can also be viewed from within Console Mode.

The shortcut is only valid throughout the existing session; it is not saved as a default. Each time you want to use this file path you must launch Thermo-Calc from the customized shortcut.

The following uses Windows for the example.

8

#### SETTING UP THE DEFAULT CONSOLE DIRECTORY

📩 Thermo-Calc-2017	<b>Open</b> Troubleshoot compatibility Open file location Send to	Þ
	Cut Copy Paste	
	Create shortcut Delete Rename	
-	Properties	

1. Right-click a shortcut to Thermo-Calc and click **Properties**.

- 2. In the **Thermo-Calc 2018b Properties** dialog, **Target** field, add the following to the end of the string: **–r** plus the *"path to the default directory"*. Include a space between the –r and path, and use quotation marks around the full file path. A simple example:
  - -r "C:\Thermo-Calc\_Macro\_Files"

Target type:	Application			
Target location:	2017b			
Target:	nermo-Calc.exe	" -r "C:\Thermo-Calc_Macro_Files"		
Start in:	"C:\Program Files\Thermo-Calc\2017b"			

- 3. Click Apply and OK.
- 4. Launch Thermo-Calc from the shortcut where you added the parameter. You can confirm that it is set correctly from within Thermo-Calc.

### VIEWING THE FILE PATH IN THERMO-CALC

- 1. Launch Thermo-Calc from the short cut where you added the parameter.
- 2. In Console Mode, right click the Console window and choose Properties.

Console		
🔚 Console 1		1
	Properties	
Thermo	Close 🗟	
Only f	Close All	Calc So
Local	Close Other	orion
	New Console	
SYS:		

3. In the **Console Properties** window the **Default directory** section displays the custom file path.

Console Properties				
Console name:	Console 1			
Buffer size:	40,000 🚔			
System output font:	АаВbCc123 АαВβГү			
Command prompt font:	AaBbCc123 ΑαΒβΓγ			
User input font:	AaBbCc123 ΑαΒβΓγ			
Background color:				
Default directory:	C:\Thermo-Calc Macro Files			
L				

# **Console Mode Results Window**

By default, the Console Results window is on the right-hand side of the screen. This is where plotted diagrams display. Each Console has its own Results tab in the Results window. All the plots and tables generated are presented in the Results tab.

- To open new **Plot** tabs press <Ctrl+Shift+T> or click the add <sup>O</sup> button.
- To open a new Plot tab for the currently active Console tab, press <Ctrl+Shift+T>. To close the Plot tab, press <Ctrl+Shift+W> or <Ctrl+Shift+F4>.

# **Event Log Window**

By default the Event Log window is collapsed (closed). It includes important information about the commands that run and are completed, including error messages. Click once to toggle it open and closed.

Event Log		
14:59:28,042 1	INFO	Loading database TCFE9
14:59:34,752 1	INFO	Loading database FEDEMO
14:59:39,933 1	INFO	Opened project file: T_01_Single-point_equilibrium.tcu
14:59:42,358 1	INFO	Defining component: C
14:59:42,359 1	INFO	Defining component: Fe
14:59:42,405 1	INFO	The activity System Definer 1 executed in 624 ms
14:59:42,408 1	INFO	Performing general equilibrium calculation
14:59:42,474 1	INFO	The activity Equilibrium Calculator 1 executed in 68 ms
14:59:42,602 1	INFO	The activity Table Renderer 1 executed in 127 ms

# **Console Mode Workflow**

Working with Thermo-Calc involves moving between different modules. The workflow differs depending on the type of calculation:

- If you want to perform a calculation using the POLY module, you typically first retrieve thermodynamic data in the DATA module, perform the calculation in POLY, and visualize the results in the POST module. It is possible to define a system directly in POLY using <u>"DEFINE\_MATERIAL" on page 189</u> and <u>"DEFINE\_DIAGRAM" on</u> page 187.
- If you want to calculate and plot a diagram using a response-driven module, such as POTENTIAL or SCHEIL, then you can go directly to that module. Response-driven modules prompt you to go through the steps to do the calculation and the required post-processing. These modules include BIN, TERN, POTENTIAL, POURBAIX, and SCHEIL. You typically end up in the POST module after having used a response-driven module. In the POST module you can modify the plots and save the diagram.
- If you want to tabulate a chemical substance, phase, or reaction, go directly to the TAB module.

The basic workflow is:

- 1. You are in the **SYS** module when you start Thermo-Calc in Console Mode.
- 2. In the **DATA** module define the system. Before performing a calculation, you must define the system and retrieve thermodynamic data. The data is retrieved from a database file (\*.TDB or \*.TDC).
- 3. The data needed to describe a system is then sent to the **GIBBS** module. This data is accessed by the **POLY** module when you instruct it to perform calculations.
- 4. In the **POLY** module, perform the calculations.
- In POLY, the conditions are set for an equilibrium calculation (temperature, pressure, system composition, etc.), then an equilibrium calculation is done.

- If you want to make a property or phase diagram, you set the conditions for a stepping or mapping operation and perform that operation.
- This generates data that can be used to plot a property diagram (if a stepping operation is performed) or a phase diagram (if a mapping operation is performed).
- 5. In the **POST** module plot and visualize your data.
- If a stepping or mapping operation is performed, then you can go to the POST module and plot a property or phase diagram.
- The diagram can be plotted quickly using default settings, but you can also modify which variables to plot of the diagram axes and change the appearance of the diagram.
- You can save the diagram either as a plain text file with data about all the coordinates or as an image file.

# **Performing Calculations in Console Mode**

Calculations can be performed either in the POLY module, or using some of the responsedriven modules designed to perform specific types of calculations.

- <u>"About Equilibrium Calculations" on page 182</u> describes generally how to calculate and plot *Property Diagrams* and *Phase Diagrams*. To calculate and plot a property diagram or a phase diagram, you must first calculate an equilibrium.
- "About Scheil Simulations" on page 197
- "T0 Temperature Simulations" on page 204
- "About Paraequilibrium" on page 207
- "About Potential Diagrams" on page 211
- "About Aqueous Solutions" on page 215
- Also search the help or see <u>"TABULATION\_REACTION Commands" on page 303</u> and <u>"Equilibrium Calculations with the POLY\_3 Module" on page 157</u> in the *Thermo-Calc Console Mode Command Reference*.

# **Using the Command Line Prompt**

The command line prompt in the Console window tells you which module you are currently in. When Thermo-Calc is first opened in Console Mode, you are in the SYS module, which is indicated with the command line prompt sys.

To use Console Mode, you type in commands at the command line prompt. The available commands depend on which module you are in. You can list the commands by typing ? at the prompt and then press <Enter>.

- "Moving Between Modules and Submodules" below
- "Entering Command Names or Abbreviations" below
- "Specifying Parameters" on the next page
- "Wild Card Characters" on page 171
- "Command History" on page 173
- Keyboard Shortcuts" on page 162

# **Moving Between Modules and Submodules**

To go to a specific module, you typically use GOTO\_MODULE (or just GOTO) followed by the name of the module. For example, to go to the DATA module, type GOTO DATA.

The exceptions are the submodules POST and ED-EXP:

- To go to the POST module, you must enter the POST command from within the POLY module or DICTRA module (if you have the add-on license).
- To go to <u>"Experimental Data Files" on page 70</u>, you must enter the ED-EXP command from within the PARROT module.

To go back to the previous module type Back. For example, if you are in the DATA module and entered it from the SYS module, when you type Back, you return to the SYS prompt. This command is also available to exit from the submodules POST and ED-EXP.



Without a valid Thermo-Calc license, you cannot leave the SYS module. To enter the DICTRA and DIC\_PARROT modules, you also need a valid Diffusion Module (DICTRA) license key.

### **Entering Command Names or Abbreviations**

The name of a command typically consists of terms linked with underscores, for example, "LIST\_EQUILIBRIUM" on page 198. Note the following:

- UPPER or lower case letters are OK.
- Hyphens (-) instead of underscores (\_) are OK, e.g. LIST-EQUILIBRIUM.

- You can abbreviate commands as long as it is unambiguous, e.g. when more than one command begins with LIST\_, you need to type the next letter or word to distinguish between the commands.
- Each word can be abbreviated e.g., L\_E for LIST\_EQUILIBRIUM.

#### **EXAMPLES**

These are examples of how you can abbreviate commands at the command line prompt.

Command name	Abbreviation
CALCULATE_EQUILIBRIUM	C-E
CALCULATE_ALL_EQUILIBRIA	C-A
LIST_EQUILIBRIUM	L-E
LIST_INITIAL_EQUILBRIUM	LI-I-E
LOAD_INITIAL_EQUILBRIUM	LO-I-E
LIST_PHASE_DATA CBCC	L-P-D CBC
LIST_PHASE_DATA CEMENTITE	L-P-D CEM
SET_ALL_START_VALUES	S-A-SS-AL
SET_AXIS_VARIABLE 1 X(FCC,FE) 0 0.89 0.025	S-A-V 1 X(F,FE) 0 .89 .025
SET_START_CONSTITUENT	S-S-C
SET_START_VALUE	S-S-V
SET_AXIS_PLOT_STATUS	S-A-P
SET_AXIS_TEXT_STATUS	S-A-T-SS-A-TE
SET_AXIS_TYPE	S-A-TY
SET_OPTIMIZING_CONDITION	S-O-C
SET_OPTIMIZING_VARIABLE	S-O-V
SET_OUTPUT_LEVEL	S-O-LS-OU

### **Specifying Parameters**

Running a command typically requires that several parameter values are specified, which can be done directly in the command line after the command name. For example, to set an axis variable for a stepping or mapping operation, enter:

SET\_AXIS\_VARIABLE 1 X(FCC,FE) 0 0.89 0.025

You can also use SET\_AXIS\_VARIABLE and press <Enter>. At the command line prompt, you then specify the parameters one at a time.

On all command lines, you can enter a question mark (?) for help. For some parameters, enter two question marks (??) for more detailed information.

#### **DEFAULT PARAMETERS VALUES**

When you are requested to specify a parameter, Thermo-Calc usually suggests an appropriate default value, which is displayed at the prompt between two slash signs (//). For example, this prompt means the default answer is TERMINAL:

```
OUTPUT FILE /TERMINAL/:
```

Press <Enter> to accept the default.

If you specify some parameters directly after the command name, then you can still choose to accept default values for some parameters.

To accept the default value for a parameter, type a comma (,) instead of a value for that parameter. For the comma to be interpreted correctly, type a blank space on each side of the comma. However, if you accept the default values for several parameters that follow each other, then the commas can be typed in without blank spaces separating them: (, , ,) is interpreted in the same way as (, , ,).

For some commands and parameters, Thermo-Calc treats the values previously assigned as default values if the command is used again. For example, if the first stepping or mapping axis variable is specified as:

```
SET AXIS VARIABLE 1 X(FCC,FE) 0 0.89 0.025
```

Then if you want to change a parameter value for this axis variable, type:

```
SET AXIS VARIABLE 1 , 0.1 ,,,
```

This changes the minimum value from 0 to 0.1 at which a stepping or mapping operation halts. The other parameter values remain the same. Consequently, what is typed is equivalent to:

SET\_AXIS\_VARIABLE 1 X(FCC,FE) 0.1 0.89 0.025

### Wild Card Characters

You can sometimes use the asterisk (\*) as a wild card character to, for example, refer to all components, phases, or species, when showing calculated properties in the POLY module, or when you set axis variables for plotting diagrams in the POST module.

When showing calculated properties in the POLY module, you can also use the dollar sign (\$) to refer to all stable phases.

For activity and/or chemical potential properties, the \* and \$ normally do not work properly as wild card characters.

### **EXAMPLES**

8

These are examples of the wild card characters you can use to search commands at the command line prompt.

Command	Action
SHOW_VALUE B(*)	Lists mass (gram) of all components in the system.
SET_AXIS_VAR Y B(*) *	Sets mass (gram) of all components in the system as Y-axis variable.
SHOW_VALUE MUR(*)	Lists chemical potentials of all components in the system.
SET_AXIS_VAR Y ACR(*) *	Sets activities of all components in the system as Y-axis variable.
SHOW_VALUE HM(*).T	Lists heat capacities of all phases.
SET_AXIS_VAR Y GM(*) *	Sets molar Gibbs free energies of all phases as Y-axis variable.
SHOW_VALUE TC(\$)	Lists curie temperature of all stable phases.
SHOW_VALUE W(*,*)	Lists mass fractions of all components in all phases.
SHOW_VALUE W(\$,*)	Lists mass fractions of all components in all stable phases.
SHOW_VALUE W(FCC,*)	Lists mass fractions of all components in the FCC phase.
SHOW_VALUE W(*,FE)	Lists mass fractions of the Fe component in all phases.
SHOW_VALUE W(\$,FE)	Lists mass fractions of the Fe component in all stable phases.
SHOW_VALUE Y(*,*)	Lists site fractions of all species in all sublattices of all phases.
SHOW_VALUE Y(\$,*)	Lists site fractions of all species in all sublattices of all stable phases.
SHOW_VALUE Y(*,*)	Lists site fractions of all species in all sublattices of all phases.

# **Command History**

- To scroll through the last twenty commands used, type two exclamation marks (!!) and press <Enter>.
- To repeat the last n commands, type ! followed by the number of previous commands to be executed again.
- To scroll through previous performed commands, use the Up ( $\uparrow$ ) or Down arrow ( $\downarrow$ ) keys.
- To read more information about the command history functionality in Thermo-Calc, type ! ? at the command line prompt.

# Macro, Workspace and Experimental Data Files



For DICTRA Console Mode see *File Formats* in the *Diffusion Module* (*DICTRA*) User Guide

In Console Mode, Thermo-Calc uses different kinds of files, including:

- <u>"Macro Files" on the next page</u> (including <u>"Log Files" below</u>) with the file extension \*.TCM.
- <u>"Workspace Files" on page 177</u> in POLY (\*.POLY3), GIBBS module (\*.GES5), and the PARROT module (\*.PAR and \*.POP).
- Experimental Data Files (\*.EXP)
  - "Commands in the Experimental Data Files (\*.POP, \*.DOP)" on page 71 in the Thermo-Calc Command Reference

Drag and drop files into the Console window to open or run a macro, for example. File types that can be opened this way are macro files (\*.TCM and \*.DCM), database files (\*.TDB), and workspace files (\*.POLY3, \*.GES5,\*.PAR, and \*.DIC). Note that \*.DCM and \*.DIC file extensions are applicable to the Diffusion Module (DICTRA).

# **Log Files**

Log files with the file extension \*.TCM are plain text files used to save a sequence of commands. Log files can be edited in a text editor.

- To start saving your input into such a file, use SET\_LOG\_FILE in the SYS module, followed by the name of the file that you want to save your command sequence to.
- If you want to save the output in the log file as well, use SET\_ECHO before the SET\_ LOG\_FILE command. Doing this is useful if you want use the log file later as a macro file.
- System Utilities Commands" on page 296 in the Thermo-Calc Console Mode Command Reference

# **Macro Files**

8

Macro files (with the file extension of either \*.TCM or \*.LOG) are plain text files used to save a sequence of commands that can be loaded and executed. Macro files can be edited in a text editor. When creating a command sequence, you add comments to the file by starting a line with @@.

Macro files include comments, which you can either run in Thermo-Calc or open and read in a text editor.

If you only open and read the macro file in a text editor, you do not see the output that Thermo-Calc gives in response to the commands stored in the macro file. It is recommended you run the macro to view all the details.

### **OPENING A CONSOLE MODE EXAMPLE MACRO FILE**

To open an example macro file for Thermo-Calc:

- 1. Open Thermo-Calc.
- 2. Confirm you are in Console Mode. If not, click the **Switch to Console Mode** button.
- 3. From the main menu, select File → or Help → Examples Files. The Console Mode folder opens.
- 4. Open the **Thermo-Calc** folder to see the available examples contained in separate folders.

Name	
👢 tcex01	
👢 tcex02	
👢 tcex03	
👢 tcex04	
👢 tcex05	
👢 tcex06	

▲ For the Diffusion Module (DICTRA) examples (\*.DCM macro files), open the Diffusion Module - DICTRA folder.

- 5. In the folder, double-click the \*.TCM file or click **Open**.
- 6. Follow the prompts in the Console window i.e. keep pressing <Enter> to run the macro and produce a plot in the **Console Results** window.

### MACRO FILE COMMANDS

Action	Command	Description and Comments
To load a macro file	In the SYS module, <u>"MACRO_FILE_OPEN"</u> <u>on page 5</u> followed by the name of the macro file.	Thermo-Calc starts the command sequence that the file contains.
Regain control of the console	"SET_INTERACTIVE" on page 292	End a macro file with this command to have control of the Console returned to you.
Add comments to a macro file	Start a line with @@, or enclose it between an @ (-line and a @)- line.	When lines are enclosed the software ignores them when running the macro.
Open a macro file	MACRO_FILE_OPEN	When creating the log file, you can make a macro file load up to five other macro files.

Action	Command	Description and Comments
Nest macro files inside of each other	MACRO_FILE_OPEN SET_INTERACTIVE	If a nested macro file ends with SET_ INTERACTIVE, then Thermo-Calc resumes with the higher-level macro file at the command immediately following the MACRO_FILE_OPEN command that loaded and executed macro file that has just been terminated. If a nested macro file doesn't end with SET_INTERACTIVE (but with an end-of-file character), then the console shuts down and the macro is stopped.

### MACRO FILE CONTROL CHARACTERS

The @? character allows you to make a macro interactive by allowing input. At the @? character, which is placed where a parameter value or argument is normally put, Thermo-Calc prompts to input the value of a parameter or argument. You can enter a string immediately following the @? character. This string is presented to the user when prompted to enter the parameter value or argument. The entered value is used by Thermo-Calc as input to the command in question. For example, you can request the user to specify the temperature range of a stepping calculation by entering the following in a macro file:

```
SET_AXIS_VARIABLE 1 T @?Low-temperature-limit: @?High-temperature-
limit:
```

You can also use up to nine variables in your macro file and prompt the user to enter values that can be assigned to these variables. Use the @#n character when you want to prompt the user to provide a value to the variable, where the n is a digit between 1 and 9. You can then use this value by with the ##n character. For example, you can request the user to provide the first element of a system by entering the following in the macro file:

```
@#3First-element?
```

You can then use this variable with the entering the character ##3 later in the macro file. For example, you can write:

```
SET_AXIS_VARIABLE 1 x(##3) 0 1,,,
```

Finally, there is the @ pause character. Thermo-Calc pauses and waits for input from the user when this pause character is read from a macro file. Inserting pause characters is useful if you want to allow the user to monitor what is happening when Thermo-Calc is running the macro.

### **RESPONSE-DRIVEN MODULES AND MACROS**

Macro files that are created while you use a *response-driven module* begin with the moduleentering command (for example, Goto\_Module Scheil) followed by a number of lines with responses to the module's questions and requests. The file terminates with the commands "<u>POST" on page 203</u> or <u>"SET\_INTERACTIVE" on page 292</u> (this command gives you back control of the Console). An empty line cannot be edited with any input rather than the default answer (to a specific question). Comment lines, commands to open other macro files, pause characters or input-controlling characters cannot be inserted between these empty lines (otherwise the response-driven module cannot be executed properly).

# **Workspace Files**

# For DICTRA Console Mode see *File Formats* in the *Diffusion Module* (*DICTRA*) User Guide

Workspaces files allow you to save all the data in your current workspace. A workspace contains all the data, specified settings and the result of any calculations performed. The saved data includes original and modified thermodynamic data, the last set of conditions and options, and the results of any simulations. You use the command SAVE\_WORKSPACES to create this file type, which is available in the POLY, GIBBS and PARROT modules.

When in Thermo-Calc Console Mode these are the workspace file extensions:

- POLY Module (\*.POLY3)
- GIBBS Module (\*.GES5)
- PARROT Module (\*.PAR and \*.POP)
- For DICTRA in Console Mode, the PARROT Module file extension is \*.DOP.

The workspace has all the settings specified and the result of any stepping or mapping operations performed after using SAVE\_WORKSPACES. Consequently, the command opens a workspace in which all data is saved which is generated after the SAVE\_WORKSPACE command is executed. The saved data includes original and modified thermodynamic data, the last set of conditions and options, and calculation results.

To load the data and calculation results of a workspace file, use the READ\_WORKSPACES command available in the POLY module. The file can be used for calculation in POLY, visualization in POST, or data manipulation in GIBBS. You are also given the option in the POURBAIX and SCHEIL modules to open a previously saved workspace file. This allows you to make new POURBAIX or SCHEIL calculations on the same chemical system as before but with different temperature, pressure, and composition conditions, or to plot new property diagrams based on the previous calculations.

Drag and drop files into the Console window to open or run a macro, for example. File types that can be opened this way are macro files (\*.TCM and \*.DCM), database files (\*.TDB), and workspace files (\*.POLY3, \*.GES5,\*.PAR, and \*.DIC). Note that \*.DCM and \*.DIC file extensions are applicable to the Diffusion Module (DICTRA).

### **Increasing the Buffer Size**

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Sometimes the output overflows the window text buffer.

- "Keyboard Shortcuts" on page 162
- 1. Right-click the Console tab header in the Console window (if this is the first Console tab, it is labeled **Console 1**).
- 2. Click Properties, and increase the Buffer size in the Console Properties window.

Console Properties			
Console name: Buffer size:	Console 1 40,000		
System output font:	AaBbCc123 ΑαΒβΓγ		
Command prompt font:	ΑαΒbCc123 ΑαΒβΓγ		
User input font:	ΑαΒbCc123 ΑαΒβΓγ		
Background color:			
Default directory:	C:\Users\amanda\Documents		

# Defining a System in Console Mode

*Defining a system* means to select the chemical components and to retrieve thermodynamic data about those components from an appropriate database in order to do the calculations. You use database(s) with thermodynamic data and then define what system elements as components. Once the system is defined, you retrieve the system's thermodynamic data from the database(s).

• If you use a *response-driven module* to perform your calculation, such as BIN or POURBAIX, then the module prompts you to select database(s) and define the system.

- When doing calculations in the POLY module, you typically have to manually select databases and define the system in the DATA module. However, it is also possible to define the system directly in POLY using <u>"DEFINE\_MATERIAL" on page 189</u> and <u>"DEFINE\_DIAGRAM" on page 187.</u>
- <u>"Database Commands" on page 11</u> in the Thermo-Calc Console Mode Command Reference

#### HOW TO DEFINE A SYSTEM IN CONSOLE MODE

- 1. Go to the DATA module. At the SYS prompt, use GOTO\_MODULE and enter Data.
- 2. Change the default database unless you directly specify the name of the database as a parameter to the command. Type SWITCH\_DATABASE and follow the prompts.

The second part of the command prompt indicates the current default database. For example, if the prompt is TDB TCFE8, then the current database is TCFE8.

3. Use DEFINE\_ELEMENTS followed by a list of the elements that you want in your system. (To list the elements that are available in your current database, use LIST\_DATABASE and choose Elements). For example, if you want to have Fe and C in your system, type:

DEFINE ELEMENTS Fe C

4. Use REJECT and choose Phases if you want to avoid retrieving any phases from the database. (To list the phases that are available in your current database, use LIST\_DATABASE and choose Phases. To list the phases that can form in the defined system, use LIST\_SYSTEM and choose Constituent.) For example, if you do not want the graphite phase to be retrieved, you type:

```
REJECT PHASES GRAPHITE
```

If the number of phases to include is much lower than the total number of phases, then it can be convenient to first Reject Phases \* and then restore the phases to include using Restore Phases.

5. Use GET\_DATA to search the database and send the thermodynamic data about your system to the GIBBS workspace.

At this point you can proceed to the POLY module (with GOTO\_MODULE POLY). However, you may want to add elements, phases and species to the GIBBS workspace from other databases. If so, then proceed to the next step.

6. Use APPEND\_DATABASE to select the (additional) database from which you want to

retrieve data. This command works exactly in the same way as the SWITCH\_DATABASE command, with the exception that it does not reinitialize the DATA module and the GIBBS workspace, but instead appends or replaces the data that has already been retrieved with new data from the additional database.

- Define your elements and specify whether to reject and restore any phases. Do this in exactly the same way when using the SWITCH\_DATABASE command (see steps 2 and 3).
- 8. Use GET\_DATA to search the database and add the thermodynamic data to the data that already exists in the GIBBS workspace.
- 9. Use APPEND\_DATABASE again if you want to add data from yet another database. When you have retrieved all the data you need, you can proceed to the POLY module.

# **Equilibrium Calculations**

### In this section:

About Equilibrium Calculations	.182
Setting Conditions for Equilibrium Calculations	. 182
Calculating an Equilibrium	184
Calculating an Equilibrium with a Fixed Phase	184
Calculating an Equilibrium with Suspended or Dormant Phases	.185

# **About Equilibrium Calculations**

An equilibrium describes what the composition of the end state of a system is, given a full specification of state variables such as temperature, pressure, initial composition, system size, etc. An equilibrium calculation is normally done in POLY-3 according to the *Global Minimization Technique*, which ensures that the most stable minimum under the specified conditions is computed.

For an equilibrium calculation to be performed, the state variables must all be set as conditions for the calculations. Such conditions include, for example, temperature, pressure, and system composition. When you calculate an equilibrium in the POLY module, you have to set these conditions manually.

# **Setting Conditions for Equilibrium Calculations**

Setting a condition normally involves giving a single state variable a specific value. For example, you can set the temperature to 1273.5 Kelvin (T=1273.5). Alternatively, setting a condition can involve giving a linear expression with more than one state variable a specific value. For example, you can set the mole fraction of the S component to be the same in the liquid and the pyrrohotite phases (X (LIQ, S) - X (PYRR, S) = 0).

```
POLY 3:1-e
OUTPUT TO SCREEN OR FILE /SCREEN/:
Options /VWCS/:
Output from POLY-3, equilibrium = 1, label A0 , database: TCFE7
Conditions:
T=1000, P=1E5, X(C)=5E-2, N=1
DEGREES OF FREEDOM 0
Temperature 1000.00 K ( 726.85 C), Pressure 1.000000E+05
Number of moles of components 1.00000E+00, Mass in grams 5.36552E+01
Total Gibbs energy -4.07975E+04, Enthalpy 2.41176E+04, Volume 7.21740E-06
Component
                     Moles
                                W-Fraction Activity Potential Ref.stat
                     5.0000E-02 1.1193E-02 2.1816E-01 -1.2659E+04 SER
С
                     9.5000E-01 9.8881E-01 6.1895E-03 -4.2279E+04 SER
FE
BCC A2
                         Status ENTERED Driving force 0.0000E+00
Moles 9.5069E-01, Mass 5.3063E+01, Volume fraction 9.6312E-01 Mass fractions:
FE 9.99844E-01 C 1.55514E-04
                          Status ENTERED Driving force 0.0000E+00
GRAPHITE
Moles 4.9313E-02, Mass 5.9230E-01, Volume fraction 3.6881E-02 Mass fractions:
   1.00000E+00 FE 0.00000E+00
```

The number of state variables that need to set is determined by the *Gibbs Phase Rule*. Typically, the state variables to give values to are:

- temperature (in K)
- pressure (in Pascal)
- system size in number of moles (in mole) or mass (in grams)
- the fraction of each component (in number of moles or mass)

If you fix the phase of the equilibrium and all but one of the state variables, you can discover the value of the other state variable at equilibrium.

It is possible to specify a set of conditions that does not have any equilibrium and the program detects this by failing to reach equilibrium during the calculation.

### STATE VARIABLES

The following table lists some of the available state variables you can use to set conditions:

State Variable	SET_CONDITION Parameter
temperature in the system (in K)	т
pressure in the system (in Pascal)	Ρ
system size (mole number in moles)	Ν
system size (mass in grams)	В
number of moles of a component in the system	N( <component>)</component>
mole fraction of a component in the system	X( <component>)</component>
mass fraction of a component in the system	W( <component>)</component>
activity of a component in the system	ACR( <component>)</component>
chemical potential of a component in the system	MUR( <component>)</component>
mole fraction of a component in a phase	X( <phase>,<component>)</component></phase>
mass fraction of a component in a phase	W( <phase>,<component>)</component></phase>
activity of a species referred to a phase at ambient temperature and pressure	ACR( <species>,<phase>)</phase></species>
chemical potential of a species referred to a phase at ambient temperature and pressure	MUR( <species>,<phase>)</phase></species>

State Variable	SET_CONDITION Parameter
enthalpy in the system (in J)	н
enthalpy of a phase (in J/mol)	HM( <phase>)</phase>

# **Calculating an Equilibrium**

This topic explains how to calculate an equilibrium in POLY. To calculate an equilibrium means to calculate the equilibrium composition of your system, given a full specification of conditions that reduce the degrees of freedom of the calculation to zero.

If you are running a network client installation of Thermo-Calc and cannot enter the POLY module, this may be because all the license instances on that the license server makes available have been checked out.

- "Equilibrium Calculations with the POLY\_3 Module" on page 157 in the Thermo-Calc Console Mode Command Reference
  - 1. Define your system.

"Defining a System in Console Mode" on page 178

- 2. Go to the POLY module. At the SYS prompt, type GOTO\_MODULE and enter POLY.
- 3. Use SET\_CONDITION followed by conditions and value assignments to set the conditions of your calculation. For example, to set temperature, pressure, initial composition and system size for a Fe-Cr-C system, you might enter the following:

SET CONDITION T=1200 P=1E5 W(CR)=0.18 W(C)=0.0013 N=1

This sets the temperature (T) to 1200 K, the pressure to 1 bar (100,000 Pascal), the mass fraction of Cr to 18 mole percent, the mass fraction of C to 0.13 weight percent, and the total amount of material to 1 mole. The fraction of Fe in the system is calculated from the fractions of Cr and C. You have to set the fraction of all the components in your system except one.

- 4. Use COMPUTE\_EQUILIBRIUM to run the calculation.
- 5. Use LIST\_EQUILIBRIUM to see the results of the calculation.

# **Calculating an Equilibrium with a Fixed Phase**

There are many ways to calculate equilibriums in Thermo-Calc. The following is one example.

You can calculate an equilibrium that has a certain amount of a certain stable phase. Use "CHANGE\_STATUS" on page 96 and choose Phase to specify the phase and the amount of that phase (in normalized mole number) that you want to set as fixed. For example, if you want to find out at what temperature a system starts to melt, enter the following:

CHANGE STATUS PHASE LIQUID=FIX 0

You must leave the state variable whose equilibrium value you are interested in unspecified. However, if you have already specified that state variable, you can make it unspecified again by using <u>"SET\_CONDITION" on page 209</u> and set that state variable to NONE. For example, if you have given temperature a value, you can type:

```
SET CONDITION T=NONE
```

The calculated equilibrium includes the value of the unspecified variable at which the equilibrium enters the phase that is fixed.



For an example where an equilibrium is calculated with a fixed phase, see Console Mode example tcex07. Go to <u>"About the Thermo-Calc Console Mode</u> Examples " on page 2 in the *Thermo-Calc Console Mode Examples Guide*.

# Calculating an Equilibrium with Suspended or Dormant Phases

You can calculate an equilibrium under the assumption that one or several phases are SUSPENDED or DORMANT using CHANGE\_STATUS and choose Phase. For example, to specify that all phases except one should be suspended, you can first suspend all phases and then enter a single phase in the following way:

CHANGE\_STATUS PHASE \*=SUSPENDED CHANGE STATUS PHASE FE LIQUID=ENTERED



For an example where this is done, as well as where the status of phases is set to be dormant, see <u>tcex10</u> in the *Thermo-Calc Console Mode Examples Guide*.

## **Property Diagrams**

About Property Diagrams	. 187
Calculating and Plotting a Property Diagram	. 187
Calculating a Property Diagram One Phase at a Time	. 189
Calculating Several Properties in the Same Diagram	. 190

### **About Property Diagrams**

When you calculate and plot a property diagram, there is only one independent state variable. Many different properties can be plotted as a function of this independent variable. For example, if the independent state variable is temperature, then the mole fractions of all phases can be plotted as a function of temperature. Or the composition of a specific phase may be plotted relative to temperature, or the activity of a component in the system as a whole or in a specific phase may be plotted relative to it.

A property diagram is plotted based on a series of equilibria that is computed while the value of one state variable is varied between a minimum and a maximum value. This variable is referred to as the stepping axis variable. In Thermo-Calc, first calculate one initial equilibrium, and then new equilibria are calculated at incremental steps in both directions on the stepping axis from the initial equilibrium. This continues until the stepping operation has covered the length of the axis between a minimum and a maximum value that is specified.



For an example of the calculation of a property diagram, see tcex08 in the *Thermo-Calc Console Mode Examples Guide*.

## **Calculating and Plotting a Property Diagram**

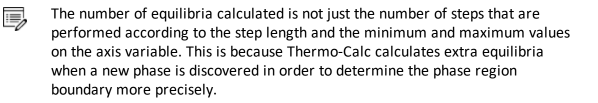
This topic explains how to calculate a property diagram in POLY. To calculate a property diagram means to calculate a series of equilibria while the value of the stepping axis variable varies between a minimum and a maximum value. With the exception of the stepping axis variable, all the state variables that you set when you calculate the initial equilibrium retain the values when the new equilibria are calculated.

- "Equilibrium Calculations with the POLY\_3 Module" on page 157 in the Console Mode Command Reference
  - 1. Calculate an initial equilibrium in the POLY module (see <u>"Equilibrium Calculations"</u> on page 181).
  - 2. Use SET\_AXIS\_VARIABLE to set the axis variable, the minimum and the maximum stepping variable values and the step length. The first parameter of SET\_AXIS\_ VARIABLE is the axis number. Since a property diagram only has one axis variable, this is set to 1. For example, if you want to set the axis variable to temperature, and calculate an equilibrium at every 50 K between a minimum temperature of 100 K and a maximum temperature of 2000 K, enter

SET\_AXIS\_VARIABLE 1 T 100 2000 50

- An axis variable must be a state variable that is set when you calculated the initial equilibrium. For example, if you set the fraction of a component in number of moles, then you cannot set the mass fraction of this component as an axis variable. Also, the minimum stepping variable value must be smaller than, and the maximum value larger than, the value that you set the state variable to when calculating the initial equilibrium.
  - 3. Use STEP\_WITH\_OPTIONS and choose Normal to perform the stepping operation.

Thermo-Calc lists the phase regions along the stepping axis, the phases contained in each region, and the number of equilibria calculated in that region. Each region is defined by the phases it contains.



```
POLY 3:s-a-v 1 T 1000 2000 100
POLY 3:step
Option? /NORMAL/:
No initial equilibrium, using default
 Step will start from axis value 1000.00
...OK
 Phase Region from 1000.00 for:
    BCC A2
    GRAPHITE
 Global check of adding phase at 1.01118E+03
 Calculated 4 equilibria
 Phase Region from 1011.18 for:
    BCC A2
    FCC A1
    GRAPHITE
 Calculated 2 equilibria
 Phase Region from 1011.18 for:
    FCC A1
    GRAPHITE
 Global test at 1.09000E+03 .... OK
 Global check of removing phase at 1.15727E+03
 Calculated
             17 equilibria
```

4. To plot the diagram, use POST to open the POST module and then use PLOT\_DIAGRAM. By default, the X-axis represents the stepping axis variable and Y-axis represents the sum of the mole fractions of all phases (NP(\*)).

### Calculating a Property Diagram One Phase at a Time

There are many ways to calculate property diagrams in Thermo-Calc. One example is where you can calculate a property diagram with a separate stepping operation being performed for each phase at a time in its default most stable composition (the major constitution). This is useful if you want to create a property diagram for a heterogeneous system with both ordered phases and their disordered pairs.

To calculate a property diagram one phase at a time, use <u>"STEP\_WITH\_OPTIONS" on</u> page 217 and when prompted choose <code>one\_phase\_at\_time</code> to perform the stepping operation.

### **Calculating Several Properties in the Same Diagram**

There are many ways to calculate property diagrams in Thermo-Calc. The following is another example. If you perform several stepping calculations after each other, the results of these calculations are all saved in your workspace file. This allows you to the following:

- Calculate (and then plot) missing parts of a specific property inside the first property diagram. These parts are calculated inside the range of the stepping variable with a different control condition.
- Calculate (and then plot) two or more sets of a specific property on the same property diagram for the same system. These can be calculated under different control conditions with stepping operations being performed across the same stepping axis variable range.
- Calculate phase boundary lines and then plot them in a corresponding phase diagram for the same system. This can be especially useful for some defined secondary phasetransformations. For example, if you want to find the phase boundary between BCC\_ A1 and BCC\_B2, or the equal-Gm for two specific phases, or the equal-fraction or equal-activity for two specific phases of a certain species, then this can be useful.

Unless you have opened a new workspace file previously using SAVE\_WORKSPACES, the results of the stepping calculations are saved in a RESULT.POLY3 file. To save the results of several stepping calculations in a file different from the workspace file that your results are currently saved to, then use SAVE\_WORKSPACES before you perform the first stepping calculation.



SAVE\_WORKSPACES overwrites and deletes the results of all previous stepping calculations.

## **Phase Diagrams**

About Phase Diagrams	192
Calculating and Plotting a Phase Diagram	192
Calculating a Quasi-Binary Phase Diagram	194
Calculating a Quasi-Ternary Phase Diagram	194

### **About Phase Diagrams**

Phase diagrams have two or more independent axis variables. Any state variable that has already been set can be used as the mapping variable for a mapping calculation and then as the axis variable for a phase diagram. From a mapping calculation, many types of phase diagrams can be plotted, with one of the mapped variables as one axis variable, and with other mapped variables or any varied property (state or derived variables) or entered symbol (variables, functions or table values) as the other axis variables.

All phase diagrams consist of zero phase fraction lines. There are two distinct types of phase diagrams: those with the tie-lines in the plane of the diagram and those where the tie-lines are not in the plane. The former includes binary phase diagrams and ternary isotherms. The latter includes more general isopleth diagrams with one or more fixed extensive variables (normally, this is a composition). The BIN and TERN modules calculate binary and ternary phase diagrams. The POTENTIAL and POURABIX modules calculate the more general isopleth diagrams.

The following topics describe how to calculate the types of phase diagrams in the "Equilibrium Calculations with the POLY\_3 Module" on page 157 module. In POLY, you can calculate phase diagram for systems with up to 40 components and with thousands of phases. You can combine activity conditions and fixed phase status and fraction conditions in any way. Thermo-Calc can calculate any arbitrary 2D section through composition space.



There is no guarantee that the conditions set will result in a calculation that reaches an equilibrium.



For an example of the calculation and plotting of a phase diagram, see Console Mode example tcex04 in the *Thermo-Calc Console Mode Examples Guide*.

## **Calculating and Plotting a Phase Diagram**

This topic describes how to calculate a phase diagram in the POLY module. The commands are described in <u>"Equilibrium Calculations with the POLY\_3 Module" on page 157</u> and <u>"Plotting and Visualization with the POST Module " on page 231.</u>

- 1. Calculate an initial equilibrium in the POLY module (see <u>"Equilibrium Calculations"</u> on page 181).
- 2. Use SET\_AXIS\_VARIABLE to set the first axis variable, its minimum and maximum mapping value and the length of each incremental step on this axis. The first parameter of SET\_AXIS\_VARIABLE is the axis number and is set to 1. For example, suppose you want the mass fraction of C as your first axis variable. You want it to vary between 0 and 0.1, with the length of each incremental step being no more than 0.002. You then enter:

```
SET AXIS VARIABLE 1 W(C) 0 .1 0.002
```

- An axis variable must be a state variable that is set when you calculated the initial equilibrium. For example, if you set the fraction of a component in number of moles, then you cannot set the mass fraction of this component as an axis variable.
  - 3. Use SET\_AXIS\_VARIABLE to set the second axis variable and specify its minimum, maximum, and step length values. The axis number is set to 2. For example, suppose you want temperature as your second axis variable and you want it to vary between 900 K and 1900 K, with an incremental step of 25 K. You then enter:

```
SET AXIS VARIABLE 2 T 900 1900 25
```

 If you want more than two axis variables, then use SET\_AXIS\_VARIABLE until you have set all the axis variables. You can set up to five axis variables. Axis variables 3, 4 and 5 must be set temperature, pressure or to the chemical potentials of components.

You may want to save your workspace with SAVE\_WORKSPACES before you perform the mapping calculation. This saves the axis variables you have set. However, it overwrites the results of any previous stepping or mapping calculations done.

5. Use MAP to perform the mapping calculation. The console lists the phase region boundaries and the phases contained on one side of each boundary.

```
Phase region boundary 30 at: 1.363E+03 2.012E-01
FCC_A1
** GRAPHITE
Calculated. 3 equilibria
Terminating at known equilibrium
Phase region boundary 31 at: 1.990E+03 2.526E-01
LIQUID
** GRAPHITE
Calculated. 13 equilibria
Terminating at known equilibrium
```

6. To plot the diagram, use POST to open the POST module and then use PLOT\_DIAGRAM. By default, the X-axis represents mapping axis variable 1 and the Y-axis represents mapping axis variable 2.

### **Calculating a Quasi-Binary Phase Diagram**

There are many ways to calculate phase diagrams in Thermo-Calc. The following is an example where a quasi-binary phase diagram is used for calculations on a ternary system in which one component has an activity or chemical potential that is fixed (although if you have fixed a phase and the phase composition varies, then the activity or chemical potential may also vary). The tie-lines in a quasi-binary diagram are in the diagram's plane of the diagram. This means that the calculation follows the lever rule as well as other rules.



For an example of a calculation of a quasi-binary phase diagram, see <u>tcex17</u> in the *Thermo-Calc Console Mode Examples Guide*.

When calculating a quasi-binary phase diagram it is necessary to set a condition on either activity or chemical potential of the third component. To specify a meaningful value it is recommended that you first change the component's reference state using SET\_REFERENCE\_STATE. For example, to calculate a phase diagram for a quasi-binary Fe-Cr-O system with fixed oxygen activity, you can enter the following:

SET\_REFERENCE\_STATE O GAS ,,

You must also use SET\_CONDITION to set the activity of the component whose activity or chemical potential you have fixed. Do this by assigning a value to the ACR state variable. For example, if the component is oxygen, you might enter

SET CONDITION ACR(O)=0.2

#### ALTERNATIVE WAY TO SET THE COMPONENT ACTIVITY

As an alternative to set the activity of the component, you can Use CHANGE\_STATUS to set a phase to fixed. For example, suppose you want to compute the iron-saturated section in "FeO"-SiO2. You then enter:

```
CHANGE-STATUS FCC=FIX 0
```

### **Calculating a Quasi-Ternary Phase Diagram**

There are many ways to calculate phase diagrams in Thermo-Calc. The following is an example of a quasi-ternary phase diagram used for calculations on a quaternary system where one component has a fixed activity or fixed chemical potential.

When calculating a quasi-ternary phase diagram it is necessary to set a condition on either activity or chemical potential of the fourth component. To specify a meaningful value it is recommended that you first change the component's reference state using SET\_REFERENCE\_STATE. For example, to calculate a phase diagram for a quasi-ternary Fe-Cr-Ni-C system with fixed carbon activity, you can enter the following:

 $\texttt{SET}\_\texttt{REFERENCE}\_\texttt{STATE}$  C GRAPH ,,

You must also use SET\_CONDITION to set the activity of the component whose activity or chemical potential you have fixed. Do this by assigning a value to the ACR state variable. For example, if the component is carbon, you might enter

```
SET CONDITION ACR(C)=0.002
```

### **Scheil Simulations**

About Scheil Simulations	. 197
About the Scheil Module in Thermo-Calc	. 197
Scheil Module Frequently Asked Questions	198
Simulating a Scheil-Solidification	199
Scheil Module State Variables	. 201

### **About Scheil Simulations**

Thermo-Calc is primarily a program for performing equilibrium calculations, but some nonequilibrium transformations or partial-equilibrium transformations can be simulated. One example of such a transformation is a Scheil-Gulliver solidification.

In a Scheil-Gulliver solidification, the diffusion in the solid phases is assumed to be so slow that it can be ignored, while the diffusion in the liquid phase is assumed to be very fast. With this approximation, the conditions at the liquid/solid interface can be described as a local equilibrium.

By making a stepping operation on the temperature variable (or enthalpy or amount liquid phase) with small decrementing steps, the new composition of the liquid can be determined. After each step, the amount of formed solid phase is removed and the overall composition is reset to the new liquid composition. In effect, the whole system is described as a non-equilibrium state regarding various parts of solidified phases at various solidification stages.

### About the Scheil Module in Thermo-Calc

You can simulate Scheil-Guilliver solidification processes with the SCHEIL module. When you enter the module, you are prompted to answer a series of questions about which database to use, what the major element is and in which amount. Thermo-Calc then displays a property diagram showing how the fraction of solid phase varies with temperature (in Celsius). You can then plot diagrams using other variables you are interested in. Do this by starting a new simulation, by opening an old file with the results of a previous simulation that you can plot differently. For example, you might be interested in plotting the fraction of remaining liquid against temperature, the fraction of each solid phase or the total of solid phases against temperature, the microsegregation in each solid phase, or the latent heat evolution against temperature.

It is possible to use the SCHEIL module to run a modified Scheil-Gulliver simulation that also takes back-diffusion of some fast-diffusing interstitial elements such as C or N into account, as well as BCC to FCC phase transformation phenomena.

#### SCHEIL MODULE EXAMPLES

See Thermo-Calc Console Mode examples tcex15 and tcex30 in the *Thermo-Calc Console Mode Examples Guide* to simulate non-equilibrium transformations without considering fast-diffusion elements.

See Thermo-Calc Console Mode example tcex48 in the *Thermo-Calc Console Mode Examples Guide* to simulate partial-equilibrium transformation with fastdiffusion elements taken into consideration. This example also shows how you can perform lever-rule calculations (full-equilibrium transformations).

#### SCHEIL AND THE DIFFUSION MODULE (DICTRA)

 $\bigcirc$ 

If you also have a license for the Diffusion Module (DICTRA), you can import a previously calculated Scheil segregation profile into the software using the command INPUT\_SCHEIL\_PROFILE. Search the online help or see the *Diffusion Module (DICTRA) Command Reference*.

### **Scheil Module Frequently Asked Questions**

#### HOW CAN I PLOT THE LATENT HEAT EVOLUTION AFTER A SCHEIL SIMULATION?

The total or apparent heat release during the solidification process consists of two parts: one is the so-called latent heat, i.e. heat due to the liquid -> solid phase transformation, and the other is the heat related to the specific heat of liquid and solid phases. To examine the heat evolution after a Scheil simulation, one can in Console Mode choose either NH or BH from the list of available plotting quantities to plot both (total or apparent) heat release and latent heat evolution against T, the temperature. This will result in a diagram with two curves, where the upper curve represents the latent heat evolution, whereas the lower curve represents the (total or apparent) heat release. By choosing NH the result is displayed as J/mol, whereas BH gives the result as J/gram. If one wants to plot just a single curve, one has to save the results to an .EXP file and re-plot after editing it.

# WHAT IS THE DIFFERENCE BETWEEN HEAT RELEASE AND LATENT HEAT EVOLUTION FOR A SCHEIL SOLIDIFICATION SIMULATION?

The total or apparent heat release during the solidification process consists of two parts: one is the so-called latent heat, i.e. heat due to the liquid -> solid phase transformation, and the other is the heat related to the specific heat of liquid and solid phases. To examine the heat evolution after a Scheil simulation, one can in Console Mode choose either NH or BH from the list of available plotting quantities to plot both (total or apparent) heat release and latent heat evolution against T, the temperature. This will result in a diagram with two curves, where the upper curve represents the latent heat evolution, whereas the lower curve represents the (total or apparent) heat release. By choosing NH the result is displayed as J/mol, whereas BH gives the result as J/gram. If one wants to plot just a single curve, one has to save the results to an .EXP file and re-plot after editing it.

#### WHY ARE THERE TWO CURVES WHEN PLOTTING HEAT EVOLUTION AFTER A SCHEIL SIMULATION?

The total or apparent heat release during the solidification process consists of two parts: one is the so-called latent heat, i.e. heat due to the liquid -> solid phase transformation, and the other is the heat related to the specific heat of liquid and solid phases. To examine the heat evolution after a Scheil simulation, one can in Console Mode choose either NH or BH from the list of available plotting quantities to plot both (total or apparent) heat release and latent heat evolution against T, the temperature. This will result in a diagram with two curves, where the upper curve represents the latent heat evolution, whereas the lower curve represents the (total or apparent) heat release. By choosing NH the result is displayed as J/mol, whereas BH gives the result as J/gram. If one wants to plot just a single curve, one has to save the results to an .EXP file and re-plot after editing it.

### Simulating a Scheil-Solidification

Simulating a Scheil-Gulliver solidification process involves calculating the liquid composition of a higher-order multicomponent system at each step of a cooling process, and resetting the liquid composition as the composition of the entire system (after having removed all amount of solid phase).

- SCHEIL Module Commands" on page 290 in the Console Mode Command Reference.
  - 1. Use GOTO\_MODULE SCHEIL to enter the SCHEIL module.
  - 2. Type START-WIZARD.
  - 3. Specify which Database to use.
  - 4. Specify the Major element or alloy in your system.

For example, for a steel/Fe-alloy, enter Fe, for an Al-based alloy, enter Al, and for a Ni-based super alloy, enter Ni.

- 5. Set whether to specify the Composition input in mass (weight) percent (the default) or in Mole percent. Enter Y for mass percent and N for mole percent.
- 6. Specify the name of the 1st alloying element. You can directly specify mass or mole percent after the name (for example, enter cr 5). If this is not specified, then you are prompted to enter it separately.
- 7. Specify the other alloying elements in the same way as you specify the first (for example,  $_{\rm C}$  \_1).
- 8. After you have specified your last alloying element, press <Enter> when requested to specify the next element. This ends the process of defining the bulk composition of the alloy system.

You can also specify all the alloying elements and their corresponding compositions on the same line when you are prompted to specify your first alloying element in Step 6. For example, you can enter Cr 5 Ni 1 Mo 0.5 C 0.01 N 0.02.

9. Specify the starting Temperature (C) in Celsius. The default is 2000.

This value should be sufficiently high so that the solidification simulation starts with the alloy system in the liquid single-phase region.

- 10. Decide which phases (if any) to reject at the Reject phase(s) prompt. Enter the name of the phases to reject or press <Enter> for NONE.
- 11. Decide whether to Restore phase(s). You may want to restore a phase that you rejected when you ran the simulation earlier, or you may want to restore a phase that is rejected by default in your database. Enter the name of the phases to restore or press <Enter> for NONE.
- 12. When prompted OK? type N to go back to Step 9 to reconsider which phases to reject or restore. Type Y to continue.

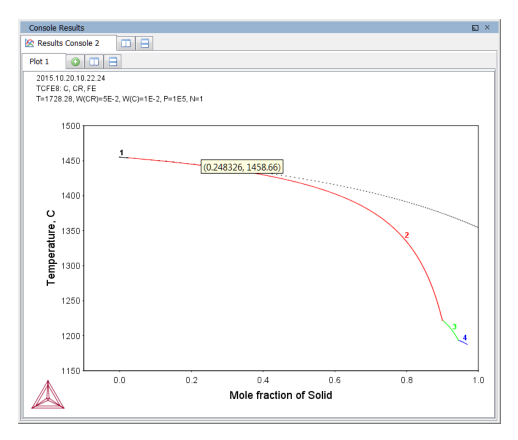
Thermodynamic data about the alloy system you defined is retrieved from the database.

13. At the Should any phase have a miscibility gap check prompt, press <Enter> for N or answer Y.

If you answer Y, you are prompted to specify the Phase with miscibility gap. Answer with a solution phase name(s) (in this example FCC is entered) as well as the Major constituents for sublattice (for each sublattice site) (for this example, Cr is entered for sublattice 1 and c for sublattice 2). In this example the Phase with miscibility gap is LIQUID PHASE NAME: LIQUID.

- 14. Press <Enter> to specify another phase.
- 15. Specify the names of any Fast-diffusing components. Press <Enter> for NONE or type all the names on the same line then press <Enter>. If you do specify any fast-diffusing components, then you are prompted to set whether BCC to FCC phase transformations should be simulated.

The Scheil-Gulliver solidification simulation runs, and the default diagram (Temperature versus Mole Fraction of Solid) is automatically plotted in the Console Results window.



### **Scheil Module State Variables**

Post processing is done the same way as POLY-3: specific variables for plotting in the Scheil Module are listed.

#### "Equilibrium Calculations with the POLY\_3 Module" on page 157

The variables can be plotted along the X- and Y-axes. In the table, *ph* is an abbreviation of *phase*, and *el* is an abbreviation of *element*. Use the abbreviations of the variables when you specify what the X- and Y-axes represent in a diagram.

Abbreviation	State variable	
т	Temperature in Celsius	
NL/BL	Mole/mass fraction of liquid	
NS/BS	Mole/mass fraction of all solid phases	
NS(ph)/BS(ph)	Mole/mass fraction of a solid phase	
W(ph,el)	Weight fraction of an element in a phase	

Abbreviation	State variable
X(ph,el)	Mole fraction of an element in a phase
Y(ph,el)	Site fraction of an element in a phase
NN(ph,el)	Distribution of an element in a phases
NH/BH	Heat release and Latent heat per mole/gram
CP/BCP	Apparent heat capacity per mole/gram
NV/NV(ph)	Molar volume of the system or a phase
DS/DS(ph)	Average density of the system or a phase
ВТ	Apparent volumetric CTE of the system

# **T0 Temperature Simulations**

T0 Temperature Simulations	204
Making a TO Temperature Simulation	204

### **T0 Temperature Simulations**

The T0-temperature is the temperature at which two phases have the same Gibbs energy for a certain composition in a multicomponent system. The  $T_0$ -temperature is located within the two-phase field between the phases and it is the theoretical limit for a transformation without diffusion. T0-temperatures are often of interest if you want to understand how diffusionless transformations work. The T0 temperature in a multicomponent system with a fixed composition is temperature at which the tangent lines of the Gibbs energies of the two target phases are the same.

If the composition of one or two components varies, the common Gibbs energy for the two phases in partial equilibrium of a diffusionless transformation becomes a plane or surface, and the T0 temperature becomes a line or plane.



See Console Mode examples tcex23 and tcex41 in the *Thermo-Calc Console Mode Examples Guide*.

### **Making a T0 Temperature Simulation**

This describes how to perform a T0 temperature simulation and apply a stepping calculation to its initial result.

Before you perform a TO temperature calculation, you must have defined your initial system and be in POLY. The initial overall composition must have appropriate settings for the desired TO temperature calculation for the two target phases.

- "Equilibrium Calculations with the POLY\_3 Module" on page 157 in the Thermo-Calc Console Mode Command Reference
  - 1. Use SET\_CONDITION to set the conditions of your calculation in the same way as when you calculate an ordinary equilibrium.
  - 2. Use COMPUTE\_EQUILIBRIUM to run the calculation. The calculation does not need to reach an equilibrium in which any of the target phases is stable.
  - 3. When you have calculated your initial equilibrium, use ADVANCED\_OPTIONS and choose T-zero.
  - 4. When prompted, specify the names of the two target phases for which the T0 temperature (where the phases' Gibbs energies are equal) is to be calculated.
  - 5. If the TO temperature between the two target phases is calculated, this temperature (in Kelvin) is shown in the Console window.
  - 6. If you want to do a stepping calculation based on this initial T0 temperature calculation, then use SET\_AXIS\_VARIABLE to specify which state variable to use as the

stepping variable. This is done the same way as when you calculate a normal property diagram.



When a T0 temperature simulation is run, you cannot set temperature as the stepping variable.

7. Use STEP\_WITH\_OPTIONS and choose T-zero to initiate the stepping calculation. You are prompted to specify the names of the target phases for which the T0 temperature is calculated.

During the STEP T-ZERO calculation procedure, the calculated TO values are presented next to the stepping variable.

For example, part of the output might look like this:

Phase Region from	1.000000E-01 for:
BCC_A2	
FCC_A1	
1.00000E-01	940.24
9.250000E-02	941.20
2.50000E-03	977.61
7.500000E-09	979.34

When the stepping calculation has finished, you can move to the POST module and plot the T0 line against the stepping composition variable or another varying axis value. Or you can impose the calculated T0 line onto a normal phase diagram.

## Paraequilibrium

About Paraequilibrium	
Calculating a Paraequilibrium	

### **About Paraequilibrium**

A paraequilibrium is a partial equilibrium where one interstitial component (such as carbon C and nitrogen N) in a multicomponent alloy can diffuse much faster than the other components (the substitutional elements, including the matrix element and alloying elements). The chemical potential for the interstitial component in two partially equilibrated phases is equal in such a case, but this is not true for the other components. In such a paraequilibrium state, it is possible to have a transformation that is partly without partitions, where a new phase of the mobile component can form with different content but where the slowly diffusing components retain their compositions.

A paraequilibrium calculation is useful when, for example, you want to study phase transformations in systems with large differences in the diffusivities of different elements. Transformations occurring under paraequilibrium states can be much more rapid than if full local equilibrium holds at the phase interface.



See Console Mode example tcex42 in the *Thermo-Calc Console Mode Examples Guide*.

## **Calculating a Paraequilibrium**

This topic describes both how to calculate an initial paraequilibrium with two target phases, and how to base a stepping calculation on this initial calculation.

Before you do any paraequilibrium calculations, you must have defined your initial alloy system and be in POLY. The initial overall composition must have appropriate settings for the desired paraequilibrium calculation for the two target phases.



These commands are described in the *Thermo-Calc Console Mode Command Reference* included with this documentation set.

- 1. Use SET\_CONDITION to set the conditions of your calculation just like when you calculate an ordinary equilibrium.
- 2. Use CHANGE\_STATUS to set the status of the chosen interstitial components to SPECIAL. For example, if C is an interstitial component, enter:

CHANGE\_STATUS COMPONENTS C=SPECIAL

This gives you a clear picture on u-fractions of the substitutional and interstitial components, which are different from the overall composition in the system. The SPECIAL status means that the specified components are not included in summations for mole or mass fractions. Therefore, all the composition variables plotted from paraequilibrium calculations are u-fraction related quantities.

- 3. Use COMPUTE\_EQUILIBRIUM to run the calculation.
- 4. When you have calculated your initial equilibrium, use ADVANCED\_OPTIONS Paraequilibrium to perform the paraequilibrium calculation.
- 5. Specify the names of the target phases of the paraequilibrium state. For example, FCC#1 BCC and FCC#2 M23C6.

Both phases must have similar interstitial/vacancy sublattices that the fastdiffusion interstitial components occupy. Both target phases should have phase constitution definitions that cover all the defined substitutional and interstitial components of the current alloy system.

6. Specify the names of one or more fast-diffusing components. These components must be located on the interstitial/vacancy sublattices in both of the chosen phases.



Interstitial components (for instance C and N combined) may have significantly different behaviours depending on how they are partitioned in different structured phases.

If the paraequilibrium between the two specified phases is successfully calculated, then a message is displayed, for example:

```
NP(FCC) = 0.3586 with U-fractions C = 2.71821E-02 N = 4.1548129E-03
NP(BCC) = 0.6414 with U-fractions C = 7.10061E-04 N = 2.3781027E-04
All other compositions are the same in both phases
```

The first two lines show the phase amounts expressed in mole-percent [NP(phase)] and the contents of the interstitial components C and N in each phase. These contents are expressed in u-fractions. The third line states that the compositions of the matrix component and all the remaining compositions (regarding substitutional components) are equal in the two target phases at the paraequilibrium state.

- If you want to do a stepping calculation based on this initial paraequilibrium, then use SET\_AXIS\_VARIABLE to specify which state variable to use as the stepping variable. This is done exactly in the same way as when you calculate a normal property diagram.
- 8. Use STEP\_WITH\_OPTIONS → Paraequilibrium to initiate the stepping calculation. You are prompted to specify the names of the target phases of the paraequilibrium states as well as the fast-diffusing components.

The stepping calculation is performed, and the different phase regions are listed with columns for the value of the stepping variable, the amounts of the two target phases, the u-fractions of interstitial(s) in each of the two phases, and the LNACR value(s) of the interstitial component(s).

```
This command calculates a paraequilibrium between two phases.
You must calculate an equilibrium with the overall composition first.
Name of first phase: fcc_a1
Name of second phase: bcc a2
Fast diffusing component: /C/: C
Fast diffusing component: /NONE/:
Output during stepping is:
axis value, phase amounts, u-fractions of interstitial(s) in phase 1 and 2,
and LNACR value(s) of interstitial(s)
Phase Region from 900.000 for:
   BCC A2
   FCC A1
  9.000000E+02 0.176 0.824 4.977501E-02 6.609390E-04 -2.303643E-01
  8.900000E+02 0.161 0.839 5.421135E-02 6.889794E-04 -1.689777E-02
  8.800000E+02 0.148 0.852 5.872152E-02 7.140931E-04 1.943989E-01
  8.700000E+02 0.137 0.863 6.329608E-02 7.362105E-04 4.040959E-01
  8.600000E+02 0.127 0.873 6.792661E-02 7.552871E-04 6.126989E-01
  8.500000E+02 0.119 0.881 7.260558E-02 7.713010E-04 8.206607E-01
  8.400000E+02 0.111 0.889 7.732629E-02 7.842505E-04 1.028392E+00
  8.300000E+02 0.105 0.895 8.208278E-02 7.941530E-04 1.236267E+00
  8.200000E+02 0.099 0.901 8.686971E-02 8.010431E-04 1.444633E+00
  8.100000E+02 0.093 0.907 9.168236E-02 8.049714E-04 1.653814E+00
  8.000000E+02 0.089 0.911 9.651648E-02 8.060032E-04 1.864114E+00
```



It is possible to now move to the POST module and plot some of the results of the calculation. For an example of a paraequilibrium calculation and how it can be plotted, see Console Mode example tcex23. Go to <u>"General Examples Using the Steel Database" on page 13</u> in the *Thermo-Calc Console Mode Examples Guide*.

## **Potential Diagrams**

About Potential Diagrams	.211
Calculating a Potential Diagram	211
Calculating a Potential Diagram with Different Pressure	212

### **About Potential Diagrams**

If you have a metal-oxide/sulphide-gas interaction system, you might want to calculate a socalled potential phase diagram given certain temperature and pressure. A potential diagram uses the activities of two major species in the gaseous mixture phase as mapping variables. The activities (that is, the fugacities at 1 bar) of these two species are typically set as the Xand Y-axis when the diagram is plotted. The phase relations between the gaseous mixture and various metal forms, metal-oxides, metal-sulphides, or other metal-bearing solids, are represented by different fields controlled by these two potential quantities.

This topic describes how to calculate such potential diagrams with the POTENTIAL module, and how to later modify the pressure of such a calculation in POLY. The calculation done by the POTENTIAL module always assumes a pressure of 1 bar.



"POTENTIAL\_DIAGRAM Module" on page 263 in the Console Mode Command Reference.

## **Calculating a Potential Diagram**

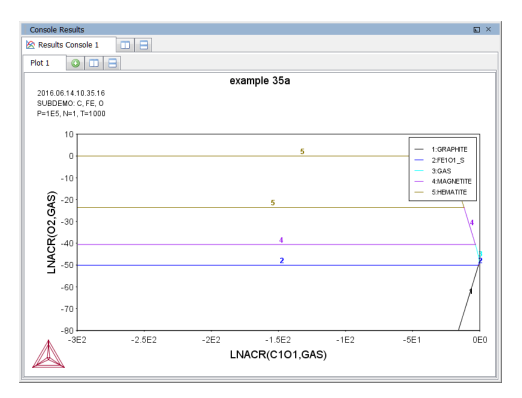
You need access to a substance or solution database that contains a gaseous mixture phase, metals (or alloys), and solids such as metal-oxides, sulphides, sulphates, carbonates, silicates, nitrides or nitrates (stoichiometric or solution phases).

POTENTIAL\_DIAGRAM Module" on page 263 in the Console Mode Command Reference

- 1. Use GOTO\_MODULE and enter POTENTIAL.
- 2. Specify which substance or solution database to use. Any substance or solution database can be used that has a gaseous mixture phase, metals (or alloys), and solids such as metalxides, sulphides, sulphates, carbonates, silicates, nitrides, and nitrates (stoichiometric or solution phases).
- 3. Specify the matrix element (a metal) and the two gaseous species that define you metal-gas interaction system. The names of these components must be available in the database you are using.
- 4. Set the temperature (in Kelvin). (The pressure is always 1 bar.)

A potential diagram with the activities of the two gaseous species as mapping variables is calculated. These mapping variables are also set as the axis variables in the plotted diagram.

The following is an example diagram (from example 35 in the *Console Mode Examples*).



A POLY3-file is also automatically saved in the current working directory with the default name POT.POLY3.

5. When the diagram is plotted, it opens the POST module where you can modify the plotted diagram in any way you like.

# Calculating a Potential Diagram with Different Pressure

It is possible to modify both the calculation and the plotted diagram that the POTENTIAL module generates in the POLY and POST modules. You can do this by opening a POLY3-file with the results from a POTENTIAL module calculation in either POLY or POST. You can modify the calculation in the same way that you can modify any ordinary mapping calculation or phase diagram. For example, this is useful for calculating a potential diagram at a pressure other than 1 bar.

"Equilibrium Calculations with the POLY\_3 Module" on page 157 in the Console Mode Command Reference

You must have calculated a potential diagram already, and have the saved POLY3-file available.

- 1. Use GOTO\_MODULE and enter POLY.
- 2. Use READ followed by the name of the POLY3-file that contains the results of the potential diagram calculation that you want to modify.
- 3. This opens the workspace file and loads the previous settings and results. The activities of the two gaseous species is now part of the conditions, and these conditions are set as the mapping variables.
- 4. Use SET\_CONDITION to specify the pressure under which you want the potential diagram to be calculated. For example, to set the pressure to 1000 bar, enter:

```
SET CONDITION P=1e8
```

5. Use COMPUTE\_EQUILIBRIUM to compute the initial equilibrium.

You may also want to use ADD\_INITIAL\_EQUILIBRIUM to calculate some additional equilibria that the mapping calculations are based on.

- 6. Use MAP to perform the mapping calculations.
- 7. Use GOTO\_MODULE and enter POST to set conditions for visualization and to plot the diagram.

# **Aqueous Solutions**

About Aqueous Solutions	215
Calculating a Pourbaix Diagram	216
Plotting More Pourbaix Diagrams	219
Stepping Calculations in an Aqueous Solution	221

### **About Aqueous Solutions**

Heterogeneous equilibrium calculations involving aqueous solutions are not different from other equilibrium calculations, but the setup of the equilibrium conditions and the reference states of the system is more complex. For example, the pH and Eh values of the aqueous solution phase are normally complex functions of bulk composition, temperature and pressure conditions.

For some simple systems, thermodynamic approximations of pH-Eh calculations can be used. But Thermo-Calc also has comprehensive thermodynamic models and databases for various non-ideal solution phases. These let you accurately calculate pH and Eh property diagrams (and many other types of property diagrams) in complex aqueous solutions in a variety of conditions.

A Pourbaix diagram is a pH-Eh phase diagram where the tie-lines are not in the plane. The chemical potentials or activities of two system components (H+ and EA) are used as mapping variables and the pH and Eh quantities are normally plotted on the X- and Y-axes of the diagram. You can use many other properties as axis variables in various property diagrams for the interaction system at a specific temperature and pressure. The phase relations between the aqueous solution and various metal forms, oxides, hydroxides, sulphides, sulphates, nitrates, silicates, carbonates and other solids, or gaseous mixtures, are represented as different fields defined by acidity and electric potential.

You can calculate Pourbaix diagrams with the Pourbaix module. The compositional definition and variations and the basic solvent and electron components H2O, H+1 and ZE are automatically determined by the aqueous solution phase in the Pourbaix module.



See Console Mode examples tcex40 and tcex53 in the *Thermo-Calc Console Mode Examples Guide*.

#### **KEY COMPONENTS**

These key components are defined as follows:

Component	Component status	Reference state	Ref-T(K)	Ref-P(Pa)
H2O	ENTERED	AQUEOUS	*	100000
H+1	ENTERED	SER		
ZE	ENTERED	REFERENCE_ELECTRODE	*	100000

Besides the aqueous solution phase, your system may also contain a gaseous mixture phase. Other neutral and charged gaseous species may also be constituents. For multicomponent systems there are normally also some interacting solid phases (stoichiometric or solution) as the so-called secondary phases. These are typically the matrix and/or precipitate phases in the alloys of interest.

Some databases have been specially developed for the Pourbaix module's diagram calculations. Such a database contains all necessary phases, i.e., an aqueous solution phase, a gaseous mixture phase, some alloy solid solution phases and some secondary solid phases. Databases developed for the Pourbaix module include *PAQ* (TCS Public Aqueous Solution Database) or *PAQS* (TGG Public Aqueous Solution Database).

If such a dedicated Pourbaix module is not used, then an aqueous solution database such as *TCAQ* (TC Aqueous Solution Database), *AQS* (TGG Aqueous Solution Database), or any user-specified aqueous databases can be used.

This database contains at least the AQUEOUS and REFERENCE\_ELECTRODE phases. The aqueous solution database needs to be appended with additional required data and phases, such as interacting alloy (solid) solution phases and secondary (solid and liquid) phases. Additional data is from one or more databases, such as SSUB/SSOL/TCFE or other substances/solutions databases.

### **Calculating a Pourbaix Diagram**

The POURBAIX module lets you construct Pourbaix diagrams and other types of property diagrams for heterogeneous interaction systems with aqueous solution phases.

When calculating a Pourbaix diagram, you need to have a database with an aqueous solution phase (that is, a phase with water as the dissolving solvent and with aqueous cation/anion and complex species as the dissolved solutes). Due to restrictions of aqueous solution models (SIT, HKF, Pitzer) used in the Thermo-Calc software, the aqueous solution database must be designed in the same Thermo-Calc database format as that used in the PAQ, PAQS, TCAQ and AQS databases for the aqueous solution phase.

Pourbaix diagram calculations are done in the advanced Pourbaix module.



<u>"POURBAIX\_DIAGRAM Commands" on page 265</u> in the *Console Mode Command Reference*.

#### HOW TO CALCUATE A POURBAIX DIAGRAM

1. At the  ${\tt sys}$  prompt, enter GOTO\_MODULE POURBAIX.

- 2. At the Need any instruction on the POURBAIX module? prompt, press <Enter> to skip or enter y to learn more.
- 3. Press <Enter> at the prompt Enforce a PAUSE after plotting when running a MACRO?.
- 4. Press <Enter> to accept the default 1 to start a completely new POURBAIX diagram calculation or enter 3 to Open an old file and make another POURBAIX calculation.

To choose 3, you must have already done a Pourbaix calculation and have it saved in a POLY3-file. The new calculation uses the same system definition as the previous calculation, but the other conditions, such as the bulk composition, temperature and pressure, can be modified in the next step.

5. Type Y or N to set whether to Consider the GAS phase (gaseous mixture phase) in the calculation.

The gaseous mixture phase should at least contain H2O, O2 and H2. If a GAS phase is not considered, then only the interactions between the aqueous solution phase and various solid phases (alloy phases and secondary phases) are calculated. Such a calculation may not accurately present all the heterogeneous interactions. Some secondary phases are therefore usually needed. Such phases exist in the PAQ or PAQS Public Aqueous Solution Databases. They can also be appended from the SSUB/SSOL/TCFE or other appropriate substances/solutions databases.

6. Type Y or N to set whether to Use single database (Y) or a multiple database (N).

If you choose to use a single database, at the Combined Database: /PAQ2/ prompt specify a database that is specially developed for the Pourbaix module's diagram calculations.

If you choose to use a multiple database, then you are prompted to specify an aqueous solution database and to append any additional required data and select necessary phases, such as various interacting alloy (solid) solution phases, and secondary (solid and liquid) phases, for example.

7. Specify all your solutes and the molality of each solute when prompted to do so. This can be done in terms of elements (such as Fe, Ni, Na, Cl, C or S) or arbitrary complex species (such as NaCl, CO2, CoCl3, Fe0.93S, NaSO4-1, or H2SO4). You can enter the element/species and the molarity on one line (for an example, NaCl 2.5).

The first letter of the element abbreviation must always be upper case. Any second letters must be lower case. When you have finished your specification, press <Enter> when asked to specify Next solute.

The definition of your system and its initial bulk composition is now presented.

8. Type Y to confirm the defined system and its bulk composition.

The thermodynamic data for your system is now retrieved from the database(s) chosen.

- 9. Answer the questions that you are prompted regarding various parameters of your calculation. You are asked to do any of the following:
- Rejectphase(s) **Of** Restore phases(s)
- Check for miscibility gaps on any phase (Should any phase have a miscibility gap check?)
- Enforce the Global Minimization Technique in calculations?
- Save all functions, variables and tables in POLY3 file?
- Set numerical limits on the calculation
- Confirm defined conditions?

S B

- Accept the default settings for two mapping variables
- 8. The Pourbaix calculation starts. Answer the question Any missing phase boundary you could possibly think of? Type Y or N to set whether you want to add starting points to the calculation. If you suspect that the calculation has missed some phase boundaries, this may help the program to detect them.
- 9. Answer questions about how the diagram is plotted and how to save the results of your calculation. You are asked to do any of the following:
- Change the pH/Eh steps for smoother curves?
- Zoom in on a specific region of the diagram
- Change the curve-label option for the diagram
- Add label-texts onto the Pourbaix diagram
- Change the subtitle of the diagram
- Change the axis-texts

- Further refine the diagram in the POST Module
- Print the diagram (save a Hard copy of the diagram)
- Save the X-Y coordinates of curve on a text file
- Modify the diagram
- Calculate and plot another Pourbaix diagram (Any more diagram?)

### **Plotting More Pourbaix Diagrams**

When you have calculated a Pourbaix diagram, you can plot additional diagrams based on the results of the same calculation, but with different axis variables than Ph and Eh.

The variables can be plotted along the X- and Y-axes. Use the abbreviations of the variables when you specify what the X- and Y-axes represent in a diagram. In the table, *AQsp* refers to the name of a specific aqueous species. The asterisk (\*) can be used as a wild card character for all such species. If neither *AQsp* nor \* is entered in parenthesis, then all species are searched.

You must have plotted a Pourbaix diagram to plot a diagram with other variables than Ph and Eh on the Y-axis and the X-axis. You can either plot such additional diagrams directly after having finished the plotting and saving/printing of a Poubaix Ph-Eh diagram or by choosing option 2 (Open an old file and plot other property diagrams) when you enter the POURBAIX module and opening a POLY3 workspace file.

Abbreviation	Variable
рН	Acidity
Eh	Electronic Potential (V)
Ah	Electronic Affinity (kJ)
Pe	Electronic Activity (log10ACRe)
IS	Ionic Strength
ТМ	Total Concentration
Aw	Activity of Water
Oc	Osmotic Coefficient
MF(AQsp)	Mole Fractions of Aqueous Species

#### **ABBREVIATIONS FOR AQUEOUS SOLUTION VARIABLES**

Abbreviation	Variable
ML(AQsp)	Molalities of Aqueous Species
AI(AQsp)	Activities of Aqueous Species
RC(AQsp)	Activity Coefficients of Aqueous Species

#### HOW TO PLOT AN AQUEOUS SOLUTION DIAGRAM

- 1. Type Y when prompted whether to plot any more diagrams, or select option 2 when you enter the POURBAIX module.
- 2. Specify which variable you want the X-axis to represent. Use the abbreviation specified in the preceding table. When you specify the name of an aqueous species, you can use the asterisk (\*) to select all species.
- 3. Specify which variable you want the Y-axis to represent. The diagram is then automatically plotted.
- 4. Answer questions about how the diagram is plotted and how to save it. You are asked to do any of the following:
  - a. change the axis-type (linear, logarithmic or inverse) on the X-axis or Y-axis
  - b. zoom in on a specific region of the diagram
  - c. change the curve-label option for the diagram
  - d. add label-texts onto the Pourbaix diagram
  - e. change the subtitle of the diagram
  - f. change the axis-texts
  - g. refine the diagram in the POST Module
  - h. print the diagram (save a hard copy)
  - i. save the X-Y coordinates of curve on a text file
  - j. modify the plotted diagram
  - k. calculate and plot another aqueous solution diagram

# **Stepping Calculations in an Aqueous Solution**

Performing the stepping calculation overwrites all the previous settings and stepping results on the POLY3-file. To avoid this, use <u>"SAVE\_WORKSPACE" on</u> page 286 to create a new workspace.

When you have calculated a Pourbaix diagram, you can use the saved results of the calculation (the POLY3-file), and make a property diagram based on stepping calculation.

Since the program normally sets the calculated initial equilibrium point as the starting point, the minimum value for the stepping variable must be specified as smaller or equal to the specific initial condition, and the maximum value as larger or equal to that condition. If pH is set as the stepping variable, the program automatically calculates the equilibrium constant in logarithmic (log10Ka) of the electrolysis reaction H2O (water) = H+ + OH- for the real aqueous solution under the current temperature-pressure-composition conditions, and thus this constant log10Ka value is considered as the highest pH in specifying the maximum value along stepping. Subsequently, confirm or modify an increment factor with a default value; for pressure and mole of a component, specify the step in logarithmic (log10P or log10N).

# **STATE VARIABLE ABBREVIATIONS**

8

State variable	Abbreviation
Temperature (K)	Т
Pressure (bar)	Р
Acidity	рН
Electronic potential (V)	Eh
Mole number of a dissolving component (such as Fe, Na, Cl, or NaCl) in 1 kg of water.	N(Comp) (for example, N(Fe))

You can use any of these state variables as a stepping variable.

# HOW TO DO A STEPPING CALCULATION

You must have plotted a Pourbaix diagram before making a stepping calculation on an aqueous solution.

- 1. Use GOTO\_MODULE POURBAIX to enter the POURBAIX module.
- 2. Type 4 to Open an old file and make another STEPPING calculation.
- 3. Choose which POLY3-file to load.
- 4. Specify the conditions of the calculation by answering the questions that follows.

The system definition is the same as in the previous Pourbaix calculation, but you can modify conditions such as bulk composition, temperature and pressure, and change some component definitions if necessary.

5. Specify which variable to use as the stepping variable (see the preceding table), its minimum and maximum value, and the length of each incremental step.

The settings and stepping results are written to the opened POLY3-file, and a first default property diagram is plotted.

# **Console Mode TAB Module**

# In this section:

Tabulation of Chemical Substances, Phases or Reactions	224
Tabulating a Reaction	.225
Tabulating a Substance or Solution Phase at Fixed Composition	.226

# Tabulation of Chemical Substances, Phases or Reactions

The TABULATION\_REACTION (TAB) module can be used to tabulate thermodynamic functions of any type of substance, stoichiometric phase or solution phase, or reaction. The module can tabulate thermodynamic functions of pure stoichiometric phases or solution phases at fixed composition, as well as various kinds of partial derivatives for a pure stoichiometric phase or solution phase at a given composition.

The TAB module begins by evaluating the most stable species of a substance in either a homogeneous state or in a reaction. Results are presented in a table or plotted as a graph.

# "Tabulating a Reaction" on the next page

A reaction tabulation presents you with the following properties, on rows representing different temperature levels:

- Heat capacity change / Delta-Cp (Joule/K) in column 2.
- Enthalpy change / Delta-H (Joule) in column 3.
- Entropy change / Delta-S (Joule/K) in column 4.
- Gibbs energy change / Delta-G (Joule) in column 5.

In the case of a tabulation of a substance or solution phase at a fixed composition, the properties presented are the heat capacity, enthalpy, entropy and Gibbs energy, rather than the degree of change in those properties.

Tabulating a Substance or Solution Phase at Fixed Composition" on page 226

You can add a 6th column to the table that shows the values of a property or user-defined function of your choice.

*****	*****	*****	*****	*****
т	Delta-Cp	Delta-H	Delta-S	Delta-G
(K)	(Joule/K)	(Joule)	(Joule/K)	(Joule)
******	*****	*****	*****	****
298.15	-4.44006E+01	-9.18800E+04	-1.98115E+02	-3.28120E+04
300.00	-4.43267E+01	-9.19621E+04	-1.98389E+02	-3.24452E+04
400.00	-3.92294E+01	-9.61533E+04	-2.10482E+02	-1.19604E+04
500.00	-3.34122E+01	-9.97861E+04	-2.18613E+02	9.52022E+03
600.00	-2.77768E+01	-1.02842E+05	-2.24200E+02	3.16779E+04
700.00	-2.26324E+01	-1.05358E+05	-2.28088E+02	5.43040E+04
800.00	-1.81080E+01	-1.07390E+05	-2.30808E+02	7.72568E+04
900.00	-1.41889E+01	-1.09000E+05	-2.32710E+02	1.00438E+05
1000.00	-1.08095 <mark>E+01</mark>	-1.10245E+05	-2.34025E+02	1.23779E+05
1100.00	-7.77802E+00	-1.11169E+05	-2.34908E+02	1.47229E+05
1200.00	-5.07556E+00	-1.11807E+05	-2.35464E+02	1.70750E+05
1300.00	-2.93467E+00	-1.12203E+05	-2.35782E+02	1.94314E+05
1400.00	-1.19414E+00	-1.12407E+05	-2.35934E+02	2.17901E+05
1500.00	2.55400E-01	-1.12452E+05	-2.35966E+02	2.41497E+05
1600.00	1.49022E+00	-1.12363E+05	-2.35909E+02	2.65091E+05
1700.00	2.56484E+00	-1.12159E+05	-2.35785E+02	2.88676E+05
1800.00	3.51909E+00	-1.11854E+05	-2.35611E+02	3.12246E+05
1900.00	4.38259E+00	-1.11458E+05	-2.35397E+02	3.35797E+05
2000.00	5.17775E+00	-1.10980E+05	-2.35152E+02	3.59325E+05

# **Tabulating a Reaction**

A tabulation of a given chemical reaction provides you with data about the rate of change of the standard tabulation properties at various levels of temperature. You can also add a column for a property of your choice to the table.

- <u>"TABULATION\_REACTION Commands" on page 303</u> in the Console Mode Command Reference
  - 1. Use GOTO\_MODULE TAB to enter the TAB module.
  - 2. If you want to add one thermodynamic functions to the tabulation calculation, which is present in a sixth column in the table, use ENTER\_FUNCTION.

You are prompted to enter the name of the column that is given at its head, and the function itself. The following state variables can be used to define the function: G, H, S, T, P, V and H298.

3. Use TABULATE\_REACTION followed by a formula that specifies of the reacting species and products to perform the tabulation calculation. The name of the elements must

be entered in upper case letters only. Terminate the chemical reaction with a semicolon. For example, you can enter:

TABULATE REACTION 3H2+N2=2N1H3;

Elements that are designated with a single letter must be followed by a stoichiometry factor, even if this factor is 1. co hence be interpreted as Cobalt rather than as carbon monoxide. Carbon monoxide is entered as c1o1 or o1c1.

4. Set the pressure, the range of temperature that you want the tabulation to cover, and change in temperature between each row in the table. You can also specify if you want to output the calculation results to an EXP-file and whether you want any graphical output, and if so, which column you want to be plotted against temperature.

The tabulation calculation is performed and you are given a table that shows the various properties at each temperature level within the range defined.

If you have chosen to receive graphical output, then the Console Results window presents a graph that plots the column of you chose (on the Y-axis) against temperature (on the X-axis). When this graph is plotted, you are now in the POST module, where you can modify the graph in various ways.

# Tabulating a Substance or Solution Phase at Fixed Composition

A tabulation of a substance or solution phase at fixed composition provides you with data about the standard tabulation properties at various levels of temperature for a given chemical reaction. You can also add one extra property column of your choice to the table.

It is important to have already defined the system and retrieved thermodynamic data from a database before using the TAB command.

# <u>"TABULATION\_REACTION Commands" on page 303</u> in the Console Mode Command Reference

- 1. Use GOTO\_MODULE TAB to enter the TAB module.
- Use SWITCH\_DATABASE to switch to a database that contains solution-based data, such as TCFE for example.



8

You cannot use the default SSUB database when tabulating solution phases at a fixed composition.

3. If you want to add one thermodynamic function to the tabulation calculation, which is present in a sixth column in the table, use ENTER\_FUNCTION.

You are prompted to specify the column header and the function itself. The following state variables can be used to define the function: G, H, S, T, P, V and H298.

- 4. Use TABULATE\_SUBSTANCE followed by the name of the substance or solution phase to perform the tabulation calculation.
- 5. Set the pressure, the range of temperature that you want the tabulation to cover, and change in temperature between each row in the table. You can also specify if you want to output the calculation results to an EXP-file and whether you want any graphical output, and if so, which column you want to be plotted against temperature.

The tabulation calculation is performed and a table shows the various properties at each temperature level within the defined range.

If you have chosen to receive graphical output, then the Console Results window presents a graph that plots the column you chose (on the Y-axis) against temperature (on the X-axis). ). When this graph is plotted, you end up in the POST module, where you can modify the graph in other ways.

# **Console Mode POST Module**

### In this section:

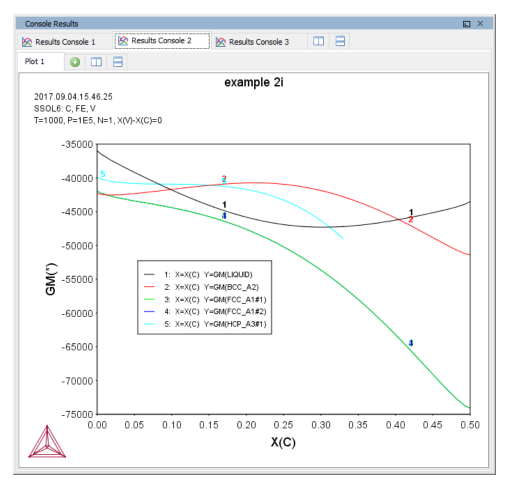
Console Mode Results	229
Plotting Diagrams	230
Modifying Diagrams	230
Saving Diagrams	232
Loading Saved Diagrams	233

# **Console Mode Results**

The results of a calculation in Thermo-Calc can be plotted by the post-processor module, also called the POST module. The diagram is presented in the Console Results window.

In the POST module, you can define and plot various kinds of diagrams based on a calculation. You can modify diagram type, which variables to plot on each axis and between which minimum and maximum values, append experimental data from an EXP-file, change labels and symbols in the diagram, zoom in on (and out from) a certain region in a diagram, etc.

If you calculate a property or phase diagram in POLY, then you have to open the POST module yourself. If your calculation was performed by a response-driven module such as SCHEIL or POURBAIX, then the plotting of a diagram based your calculation is part of the dialogue of questions and answers you have in that module (but under the hood, the response-driven module is actually utilizing the POST module to get the plotting done).



# **Plotting Diagrams**

Plotting a diagram is to graphically represent how several variables (typically two variables) co-vary. Each axis in a property diagram or a phase diagram represents the value of one variable. The default variables represented by the X- and Y-axes are the following:

- If you plot a diagram based on the results of a mapping operation (a phase diagram), then by default the X- and Y-axes represent the mapping axis variables 1 and 2 (set using <u>"SET\_AXIS\_VARIABLE" on page 208</u> in POLY).
- If you plot a diagram based on the results of a stepping operation (a property diagram), then by default, the X-axis represents the stepping variable (set using SET\_AXIS\_VARIABLE) and the Y-axis represents the sum of the mole fractions of all phases (that is, NP(\*)).
- "Equilibrium Calculations with the POLY\_3 Module" on page 157 in the Console Mode Command Reference

# HOW TO PLOT A DIAGRAM

To plot a diagram, you must have already made a stepping or mapping operation in POLY (or in some cases, made a calculation as part of some opening a response-driven module), and you must have entered the POST module.

The POST command is available inside POLY. Use POST to enter the POST module

- If you want to directly plot your diagram with the default variables represented by the X- and Y-axes, type Plot and press <Enter>. This plots a property or phase diagram with the default variables represented at each axis. If you do not want to plot your diagram with these default variables, go to the next step.
- 2. Use SET\_DIAGRAM\_AXIS to set which variables to plot on the X-axis and the Y-axis. For example, if you want to plot a phase diagram with pressure on the Y-axis and temperature on the X-axis, type:

```
SET_DIAGRAM_AXIS X T
SET_DIAGRAM_AXIS Y P
```

3. Use PLOT\_DIAGRAM to plot your diagram.

# **Modifying Diagrams**

Modifying a diagram amounts to changing the settings that determine how the POST module presents the calculated results and plots a new diagram. Suppose that you have already plotted a binary phase diagram with the mapping axis variables temperature and

mole fraction of one of your components (the Fe component, say) on the X- and Y-axes. If you then want to modify your diagram so that the Y-axis instead represents, say, the mass fraction of Fe rather than the mole fraction, then you use <u>"SET\_DIAGRAM\_AXIS" on page 249</u> to set diagram axis Y to  $_{W}(Fe)$ , and then use <u>"PLOT\_DIAGRAM" on page 244</u>.

There are many ways in which you can modify your diagram and how it is presented. Any modification made in the plot settings is taken into account the next time you use PLOT.



The following commands are described in the *Thermo-Calc Console Mode Command Reference*.

# **EXAMPLE COMMANDS TO USE**

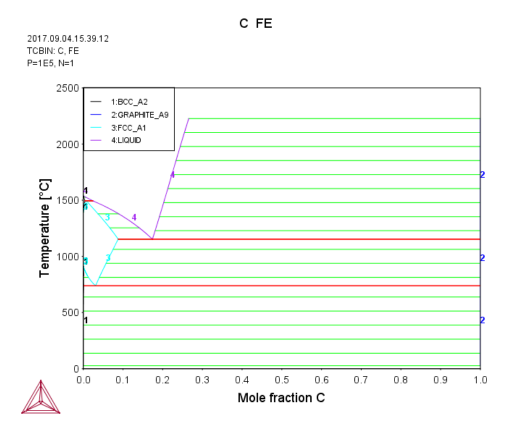
The following are some of the modifications you can make.

- Use <u>"SET\_TITLE" on page 261</u> to set add a title or to change an already existing title. The title is displayed above the plotted diagram in the Console Results window.
- Use <u>"ADD\_LABEL\_TEXT" on page 234</u> to add a label text that starts at a certain Xand Y-coordinate. If you have plotted a phase diagram, then you can let Thermo-Calc set the label text that specifies the names of the stable phases at that coordinate.
- Use <u>"SET\_AXIS\_TYPE" on page 247</u> to set whether the scale of an axis is linear, logarithmic or inverse.
- Use <u>"SET\_DIAGRAM\_TYPE" on page 252</u> to plot a triangular diagram with the X-axis on the triangle's base and the Y-axis on the triangle's left side. Such a diagram is often useful if you want to plot the fractions of two components in a ternary diagram.
- Use <u>"SET\_SCALING\_STATUS" on page 260</u> to modify the range of values that are displayed on an axis. In effect, you can use it to zoom in or out of the diagram. For example, to set the scale of the X-axis to range from 0 to 0 .3, type:

```
SET_SCALING_STATUS X N 0 0.3
```

The x specifies which axis to change the scaling on, the  $\mathbb{N}$  turns off Thermo-Calc's automatic scaling, and 0 and 0.3 specifies the minimum and maximum values on the scale. You can turn on automatic scaling again by setting the second parameter to  $\mathbb{Y}$ .

• Use <u>"SET\_LABEL\_CURVE\_OPTION" on page 253</u> to label each set of curves that have the same stable phases with a certain digit. A legend is also added, where you specify which phases each digit designates.



# LTEXT VS LATEX COMMANDS FOR LABELS

As of Thermo-Calc 2015a the LTEXT text formatting is replaced by LaTeX. For existing users who may have plots and files that use the LTEXT format, and especially when using Console Mode, the following examples show the command changes from LTEXT and its LaTeX equivalent:

Symbol or text format	LTEXT	LaTeX
Subscripts in an axis text	S-A-TEXT Y N Mole-fraction Al^DO2\$O^DO3\$	S-A-TEXT Y N \latex Mole- fraction Al_2O_3
Label with subscript	add .13 .15 n c-ZrO^DO2\$	add .13 .15 n \latex c-ZrO_2
Greek symbol	ADD .05 .50 N ^GRd\$	ADD .05 .50 N \latex \delta

# **Saving Diagrams**

When a diagram is plotted, there are several ways in which you can save it:

- Use DUMP\_DIAGRAM to save the diagram image to a PS, PDF, SVG, EMF, GIF, or PNG file. You are asked what format to save it in.
- To save the diagram image in a variety of image file formats, right click the diagram and select **Save Plot**.
- Use PRINT\_DIAGRAM to print a hard copy of the diagram.



Information about the underlying calculations that the diagram is based on is not saved in an EXP-file.

**DATAPLOT** User Guide included with this documentation set.

# **Loading Saved Diagrams**

If you have previously saved a diagram in an EXP-file, then you can load and superimpose the diagram on another diagram. Both the following command requires you to specify which prologue and dataset(s) that are loaded from an experimental data file.

Use <u>"APPEND\_EXPERIMENTAL\_DATA" on page 236</u> to plot selected data from an experimental data file (EXP-file). If you have set the axes of the diagram already, the labels on the diagram are not changed when you plot the additional data from the EXP-file, irrespectively of the data in that file.

Use <u>"QUICK\_EXPERIMENTAL\_PLOT" on page 245</u> to plot selected data from an experimental data file (EXP-file). If you have set the axes of the diagram already, these settings are overwritten with the axes settings from the EXP-file.

# About the Database Files

# **Encrypted vs Unencrypted Database Files**

Any licensed databases purchased from Thermo-Calc Software are encrypted in files with the extension . TDC; these files cannot be edited. These are referred to as *encrypted databases*.

*Unencrypted databases* have the file extension . **TDB** and can be edited. For example, PURE5 or PG35 (the free databases) or a user-defined database file. If you plan to edit any of these files.

Editing the Database Initialization File or Unencrypted Database File" on page 236

# **Database Initialization File**

In Windows, the database initialization file is called tc\_initd.tdb. In Linux and Mac it is called initd.tdb. The file contains a list of all the databases available from Thermo-Calc Software. Individual licenses are still required to use each database.

The file is used differently by the program depending on whether you are working with databases in Console Mode or Graphical Mode. For this reason, editing the file can be problematic if you don't understand the differences.

<u>"About Cached Database Files" on the next page and "Editing the Database</u> Initialization File or Unencrypted Database File" on page 236

# **User Database Files**

A user-defined database is considered to be an unencrypted database because you can make changes to it.

In Graphical Mode you can add user databases via the **Options**  $\rightarrow$  **Graphical Mode**  $\rightarrow$  **System definer** window. Once you have added the user database it is available from the database list in the System Definer Configuration window.

Global Settings: System Definer" on page 242 to add a user database.

In Console Mode, users may decide to add their own user-defined database to the database initiation file, which then makes it available in the Console window. If you plan to edit this file.

Editing the Database Initialization File or Unencrypted Database File" on page 236

# **Database Installation Folder**

Depending on the type of license and operating system, the encrypted Thermo-Calc databases and the database initialization file are located in a subfolder to the Thermo-Calc installation. For example, in a Windows standalone installation, you can find these in a folder called **data** (file path is *C*:\*Program Files*\*Thermo-Calc*\*<version>*\*data*).

OS (C:) > Program Files >	Thermo-Calc	> <version> &gt; data</version>
Name	Date modi	Туре
ALDEMO	2017-09-2	File folder
AQS2	2017-09-2	File folder
CUDEMO	2017-09-2	File folder
FEDAT	2017-09-2	File folder
FEDEMO	2017-09-2	File folder

# **About Cached Database Files**

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L				L	
E	-	-	-	Ł	
-	-	Т	-		

This is applicable to Graphical Mode.



Also see Global Settings <u>"General" on page 239</u> where you can set the default database directory path.

The first time you open a database in Graphical Mode it can take a little while for it to load, especially if it is one of the large databases such as TCFE or TCNI. However, the next time the database opens quickly because now the databases are cached, or stored, in Graphical Mode.

Think of a cache as an intermediate storage area. Once a database cache is established the file made easily accessible, even if you close the software program. It is the same concept as when you load a web page for the first time. The next time the page opens faster. This is also how it works for the databases in Thermo-Calc Graphical Mode.

Thermo-Calc caches the database files, both encrypted and unencrypted types, at two points –when you first open a database, sometimes called an *in-memory cache*, and then later as a *disk-based cache*.. The disk-based cache is what speeds up database loading and it is at this point that Thermo-Calc refers to the database initialization file for information. The disk-based cache uses a lookup key to find the abbreviated database name (e.g. TCFE or TCNI) contained in this initialization file.

It is important to understand the way databases are cached in Graphical Mode to ensure best practices for those who also use Console Mode, especially if the database initialization file is edited. If you plan to edit this file, see Editing a Database Initialization File or Unencrypted Database File for important information.

# Editing the Database Initialization File or Unencrypted Database File

Some advanced users may decide to edit the database initialization file, for example to reorder the list of encrypted databases displaying in the Console window or to add their own user-defined database for use in Console Mode. It is important to understand that changes to this file impact the database caching process used in Graphical Mode.

"About Cached Database Files" on the previous page

The following recommendations are also applicable if you edit an unencrypted database file, which includes user databases and free databases such as PURE5 or PG35.

# **Best Practice Recommendations**

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If you edit the database initialization file (tc.initd.tdb in Windows or initd.tdb in Mac and Linux), then:

• If a user database is added to the file, it is subject to the same caching as the encrypted databases. This means that further changes (to the user database) will not be detected when you are working with databases in the System Definer in Graphical Mode.

If you have a user database, which is subject to regular updates, then:

- If you make changes to a user database, you then need to clear the cache.
- It is recommended that you create a separate directory to store unencrypted database files.

If you edit an unencrypted database file included with the installation (such as the PURE5 and PG35 files):

• Do not edit the file in the installation folder. Instead make a local copy and load it into the application as a user database.

# **Database Checker**

The Thermo-Calc software package includes a program to check that the syntax of unencrypted database files is correct. The program applies the syntax rules set out in the *Database Manager User Guide* and reports errors and issues warnings. This program is intended for advanced users who develop and manage databases.



Thermo-Calc accepts deviations from these syntax rules. This means that a database can work even if the Database Checker reports errors and warnings. For example, an error is reported if an abbreviated phase name is found, but phase name abbreviations are accepted by Thermo-Calc and its add-on modules.

The executable **DatabaseChecker** file is found in the Thermo-Calc home directory. The program can also be launched by selecting **Tools → Database Checker** from the main menu.

Database Manager User Guide

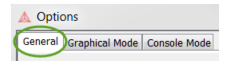
# **Changing Global Settings**

In the **Options** window (select **Tools**  $\rightarrow$  **Options**) you can globally set general settings for Graphical Mode as well as default settings for any new activities for both Graphical Mode and Console Mode.

# In this section:

General	239
Graphical Mode - Default Units	. 241
Graphical Mode: Activities	242
Global Settings: Graphical and Console Mode Plotting	247
Titles, Legends and Labels	. 248
Fonts and Background Colors	. 249
Lines and Grids	. 249
Data Points and Anchors	. 250
Plot Area Size	. 251
Global Settings: Console Mode Default Appearance	251

# General



These settings are for both Graphical and Console modes. To open this window, from the main menu select **Tools**  $\rightarrow$  **Options**.

# **Tooltips enabled**

Select whether to turn on tooltips information. By default the **Tooltips enabled** check box is selected. This displays a small text box when you hover the cursor above some buttons or other items.

# Localization

To change the GUI language from the **Localization** list choose **English** (the default), **Swedish**, **Spanish**, **German**, **Russian**, **Chinese**, **Japanese** or **French**.

### Look and feel

To change the **Look and feel** of the GUI layout, choose **Windows** (the default), **Metal**, **Nimbus**, **CDE/Motif** or **WindowsClassic**.

### **Database directory**

In the **Database directory** field, specify the path to the directory that contains the **data** directory. This is where the Thermo-Calc database directory called **data** is located. Do not specify the path to where the database files are found (that is, in the **data** directory).

# MODIFYING THE DATABASE DIRECTORY PATH (GRAPHICAL MODE ONLY)

**B** Important Note: Database Directory Path (Graphical Mode only)

When you open a database for the first time, it is stored in a cache (an intermediate storage area), that is maintained even if you close the program. This improves the speed of loading the next time you open that database. The cache is used for all databases that are listed in the database initialization file, typically all the standard databases that are included with Thermo-Calc.

If you modify your own user-defined database, or if you choose to edit an unencrypted database, it is recommended that you create a separate directory to store these database files. This way the most recent version of the user-defined or edited databases are loaded into the program when the System Definer is performed.



### **Model directory**

This is where the predefined and custom property models are saved and stored by default. Click **Modify** to open the default folder or to create a custom folder for your own models.

All property models (whether these are predefined or custom models) must be located in a subdirectory of the property model directory. For that reason it is recommended that you do not change the directory location.



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See <u>"Property Model Calculator" on page 70</u> for information about using the predefined models. Also see "Creating a New Property Model" in the *Property Model Development Framework Guide* for details about creating a custom model.

#### Log level

Select the level and type of information to display in the **Event Log** window using the **Log level** slide bar. Choose from a **Debug**, **Info**, **Warning** or **Error** level of detail.

### Check update interval

From the **Check update interval** list, choose not to check (**Don't check**, the default) or **On startup**. You can also manually check for updates – choose Help→Check for Updates.

#### **Experimental**

Select the Use GES6 in Graphical Mode check box to enable this functionality.

#### Reset to original settings

In the lower corner of the window, click the **Reset to original settings** button to restore the original settings. You may have to expand the window to view the button.

# **Graphical Mode - Default Units**

🛕 Opti	ons	
General	Graphical Mode	Console Mode
Activitie	s Default Units	

To open this window, from the main menu select **Tools**  $\rightarrow$  **Options**and click the **Default Units** tab.

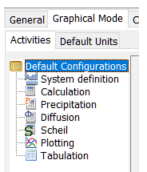
### **Reset to original settings**

In the lower corner of the window, click the **Reset to original settings** button to restore the original settings. You may have to expand the window to view the button.

Setting	Options
Temperature	Kelvin (the default), Celsius or Fahrenheit
Pressure	Pascal (the default), Atmospheres or Bar
Amount	Mole (the default), Gram, Kilogram or Pound
Composition	Mass percent (the default), Mole percent, Mass fraction or Mole fraction
Energy	Joule (the default), Calorie or Electron volt
Volume	Cubic meter (the default), Cubic decimeter or Cubic centimeter
Density	Kilogram per cubic meter (the default) or Gram per cubic meter
Entropy	Joules per Kelvin (the default), Calorie per Kelvin or Electron volt per Kelvin
Length	Meter (the default), Micrometer or Nanometer
Time	Seconds (the default), Hours or Days

# **Graphical Mode: Activities**

### A Options



To open this window, from the main menu select **Tools**  $\rightarrow$  **Options** and click the **Activities** tab.

There are **System definition**, **Calculation**, **Precipitation**, **Diffusion**, **Scheil**, **Plotting** and **Tabulation** settings windows.

For Plotting, the default settings are shared with the Graphical and Console Modes. Any changes you make also apply to the default settings on the **Console Mode**  $\rightarrow$  **Plotting** tab in the **Options** window (and vice versa).

"Global Settings: Graphical and Console Mode Plotting" on page 247

# **Global Settings: System Definer**

When you create a new System Definer, the default databases are taken from these defaults. The database selections for each System Definer can then be configured individually. Once created, the specific (local) settings for a System Definer are not affected if you later make changes to the global defaults.

To open this window, from the main menu select **Tools** → **Options** and click the **Activities** tab. Then click **System definition**.

General Graphical Mode	Console Mode	
Activities Default Units		
Default Configurations System definition Calculation Precipitation Diffusion Scheil Plotting Tabulation	Default database selection:       Database       TCFE9: Steels/Fe-Alloys v9.0         User Databases       Database name:        Add       Remove         Database name:        Add       Remove         Database Packages        Steels and Fe-alloys        Show details         Image: Imag	

# System Definer" on page 62

"Creating a Database Package" on the next page and "Clearing a Database Package Message" on page 245

### Default database selection

Choose to define a Database or a Database package.

If you choose **Database**, then select the default installed database to use globally for all System Definer nodes. For example, choose **TCFE9** or **TCNI8**.

If you choose **Database Package**, select an option from the list. These Database Packages are defined in *Database Packages*.

### User Databases

### Database name

Click the **Add** button to add a custom database. After you have added the user database it is available from the list to choose as the default.

Once a user database is added to the **Database name** list, you can also click the **Remove** button to delete it from your project.

#### **Database Packages**

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Based on the installed databases, there are also default Database Packages available.

Create and remove **Database Packages** by clicking the **Add a database package** button or **Remove this database package** button.

The addition or removal of a database package does not take effect until a new System Definer node is defined.

Click **Show details** to display the list of databases included with the package. Click **Hide details** to only display the name of the package.

"Creating a Database Package" on the next page

### **Reset to original settings**

In the lower corner of the window, click the **Reset to original settings** button to restore the original settings. You may have to expand the window to view the button.

#### Creating a Database Package

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- 1. From the main menu select **Tools**  $\rightarrow$  **Options**.
- 2. Click the Activities tab and click System definition.
- 3. Under Database Packages, click the Add a database package<sup>Q</sup> button.
- 4. Replace the default text with a name for the package, for example **My database package**.
- 5. Select a database from the **Choose database** list, for example, **TCFE9 Steels/Fe-Alloys**.
- 6. Click the Add a database S button and from the Choose database list, select another database, for example, TCCU2 Cu-Alloys
- 7. Click **OK.** The Database Package is now available to choose as the default for new System Definer activities.

Default database selection: Database Package 💌	Demo: Steels and Fe-alloys [FEDEMO, MFEDEMO] 🔻
User Databases	Demo: Steels and Fe-alloys [FEDEMO, MFEDEMO]
	My database package [TCFE8, TCAL4]
Database name:	▼ Add Remove
Database Packages	
Oemo: Steels and Fe-alloys	
💿 🤤 My database package	Show details

The package is available for new System Definer nodes defined after adding the package.

System Definer 2		
Package:		~
	Magnesium-based alloys (TCMG4, MOBMG1)	^
	Ultra-high-purity silicon (TCSI1, MOBSI1)	
	Solder alloys (TCSLD3, MOBSLD1)	
	Copper based alloys (TCCU2, MOBCU2)	
	Demo: Steels and Fe-alloys (FEDEMO, MFEDEMO)	
	Demo: Nickel-based superalloys (NIDEMO, MNIDEMO)	
Material	Demo: Aluminium-based alloys (ALDEMO, MALDEMO)	
Material name:	My database package (TCFE9, TCCU2)	۷

# "Clearing a Database Package Message" below and "Global Settings: System Definer" on page 242

# Clearing a Database Package Message

Sometimes when working on the System Definer Configuration window, you may add or delete Databases. If you originally added a Database Package, the addition or removal of a database means that the original package you selected does not now match the databases you are using. To clear the message *Selected databases and selected package may differ* and reset the databases you need to either:

- Change the newly added database back to the one that matches the database package, or
- Choose another package from the **Package** list, then select the package you want.

# For example, from the Package list you select Demo: Steels and Fe-alloys.

Package: Demo: Steels and Fe-alloys [FEDEMO, MFEDEMO] 🔻

This adds the FEDEMO and MFEDEMO databases.

Databas	es	
0 🔾	FEDEMO: Iron Demo Database v2.0	~
0 🔾	MFEDEMO: Fe-Alloys Mobility demo database v2.0	$\sim$

If you add or remove **Databases**, in this example **MFEDEMO** is removed, then this does not match the **Package** and the message displays:

🔚 System Definer 1				
Databases				
🔾 🜑 FEDEMO: Iron Demo Database 🔹	P	Package:	Demo: Steels and Fe-alloys (FEDEMO, MFEDEMO) 🔻	
			Selected databases and selected package may differ	

To clear the message, click the **Add a database** Subtton and from the **Choose database** list, select the database that belongs with the package, in this example, **MFEDEMO**. The message is cleared.

You can also choose another package from the list, which changes the databases, and then re-select the original package to reset the message.

"Creating a Database Package" on the previous page and "Global Settings: System Definer" on page 242

# Global Settings: Equilibrium Calculator

When you create a new Equilibrium Calculator, its initial settings are taken from these defaults. The settings of each Equilibrium Calculator can then be configured individually. Once created, the specific (local) settings for an Equilibrium Calculator are not affected if you later make changes to the global defaults.

To open this window, from the main menu select **Tools**  $\rightarrow$  **Options** and click the **Activities** tab. Then click **Calculation**.

# "Options Settings" on page 87



You can reset the defaults. In the lower corner of the window, click the **Reset to original settings** button to restore the original settings. You may have to expand the window to view the button.

# **Global Settings: Diffusion Calculator**

When you create a new Diffusion Calculator, its initial settings are taken from these defaults. The settings for each Diffusion Calculator can then be configured individually. Once created, the specific (local) settings are not affected if you later make changes to the global defaults.

To open this window, from the main menu select **Tools**  $\rightarrow$  **Options** and click the **Activities** tab. Then click **Diffusion**.

"Options Settings" on page 97

### **Reset to original settings**

In the lower corner of the window, click the **Reset to original settings** button to restore the original settings. You may have to expand the window to view the button.

# **Global Settings: Precipitation Calculator**

When you create a new Precipitation Calculator, some of its initial settings are taken from these defaults. The settings for each Precipitation Calculator can then be configured individually. Once created, the specific (local) settings for a Precipitation Calculator are not affected if you later make changes to the global defaults.

To open this window, from the main menu select **Tools**  $\rightarrow$  **Options** and click the **Activities** tab. Then click **Precipitation**.

"Configuration Settings" on page 106

### Reset to original settings

In the lower corner of the window, click the **Reset to original settings** button to restore the

original settings. You may have to expand the window to view the button.

# Global Settings: Scheil Calculator

When you create a new Scheil Calculator, its initial settings are taken from these defaults. The settings of each Scheil Calculator can then be configured individually. Once created, the specific (local) settings for a Scheil Calculator are not affected if you later make changes to the global defaults.

To open this window, from the main menu select **Tools**  $\rightarrow$  **Options** and click the **Activities** tab. Then click **Scheil**.

"Configuration Settings" on page 154

### **Reset to original settings**

In the lower corner of the window, click the **Reset to original settings** button to restore the original settings. You may have to expand the window to view the button.

### **Global Settings: Tables**

When you create a new Table Renderer, its initial settings are taken from these default settings. The settings of each Table Renderer can then be configured individually. Once created, the settings of a Table Renderer are not affected by changes in the default settings.

To open this window, from the main menu select **Tools**  $\rightarrow$  **Options** and click the **Activities** tab. Then click **Tabulation**.

"Configuration Settings" on page 148

### Reset to original settings

In the lower corner of the window, click the **Reset to original settings** button to restore the original settings. You may have to expand the window to view the button.

# **Global Settings: Graphical and Console Mode Plotting**



The default settings are shared with both modes. Any changes you make also apply to the default settings on the **Console Mode**  $\rightarrow$  **Plotting** tab in the **Options** window (and vice versa).

🛦 Opt	ions			
General	Graphical Mode	Console M	ode	
Activities	Default Units			
Ca	It Configurations ystem definition alculation recipitation ffusion	General	Х-ах	
	otting	Options		
T	Send	eral Graphi	ical Mode	Console Mode
	Defa	ault Appeara	ance Plott	ing
	Ger	ieral X-axi	s Y-axis	

To make local changes to the appearance of a specific plot, in the **Results** window right-click a plot and select **Properties**. In the **Plot Properties** window a variety of settings can be made.

You can also edit some properties for individual plot lines (the color, the line width and type, and whether data points are included). In the **Results** window, hover the mouse over a plot line. The crosshair cursor turns into a cursor resembling a pointing hand when it is over a line that can clicked. Alternatively, hold down the <Ctrl> key while you move the cursor around the plot to only display it as a crosshair and prevent unintended edits.

To globally configure the default settings for plotting, from the main menu select **Tools**  $\rightarrow$  **Options**. Click the **Graphical Mode**  $\rightarrow$  **Activities** tabs and then click the **Plotting** node.

When you create a new Plot Renderer, its initial settings are taken from these default settings. The settings can then be configured individually. Once created, the settings of a Plot Renderer are not affected by changes to the default global settings.

To configure the settings of an individual Plot Renderer rather than the default settings, right-click a plot in the **Results** window and select **Properties** from the menu.

Plot Renderer Settings for Console and Graphical Modes

These global settings are available for both Console and Graphical Modes.

#### Reset to original settings

In the lower corner of the window, click the **Reset to original settings** button to restore the original settings. You may have to expand the window to view the button.

# **Titles, Legends and Labels**

#### **Title format and Label format**

# Select Plain text (the default) or LaTeX formatted text.



For LaTeX formatted text you can use the LaTeX math text to enter text. See <u>"ADD\_LABEL\_TEXT" on page 234</u> and <u>"Plot Labels " on page 137</u> for more details.

### Show legend

Select the check box to include a legend on the plot. By default no legend is displayed.

### Legend phase caption style

### Choose All, Constitution description or Ordering description.



### **Retain labels**



Graphical Mode only.

By default the check box is selected. Plot labels are kept (retained) when plots are updated. It can be applied globally or locally to individual plots.

#### Show header



Console Mode only.

Click to select the **Show header** check box to display the basic details about the plot along the top. This includes the date and time the plot is generated, the database used and the properties.

# **Fonts and Background Colors**

Title font, Legend font, Label font, Header font

Click **Modify** to edit the Font Name, Style, Size and Color.

Legend background color, Background color, Canvas color, Border color,

Click **Modify** to choose a color or enter a specific color.

# **Lines and Grids**

Show grid

Select the check box to include a grid on the plot. By default no grid is displayed.

#### Show every Nth tieline



Tielines are only applicable to phase diagrams.

Choose an integer value ranging from 0 to 99.

### **Color option (plot lines)**

For the plot line colors, choose from the options in the list: Legacy, Printer friendly, JFree chart, Pastel, Medium dark, Bright dark, Vivid or Earth.

This option is also available for specific plot lines. To change the color of an individual plot line double-click it in the Results window and use the color palette to define it or enter a specific color.

#### **Border width**

Choose a plot border width between 1 (thin, the default) and 10 (thick).

#### Line width

For the plot line width, choose from the options in the list. These options are also available for specific plot lines. To change an individual plot line width double-click it in the Results window.

#### Line type

This option is only available for individual plot lines. To change an individual plot line type double-click it in the Results window and choose an option from the list.



# **Data Points and Anchors**

#### Show anchor

By default, an anchor between the label and the plot point is displayed.

#### Show data points

Click to select the check box to display the data points on the plot lines. This option is also

available for specific plot lines. To show data points on an individual plot line double-click it in the Results window and select the check box.

#### Data points marker size

Choose the data point marker size between 1 (small) and 10 (large). The default is 3.

#### Plot area size

The **Fit the plot area size to the enclosing window** option is selected by default. Excluding triangular plots, click Define the plot area size to enter a Plot area height and Plot area width.

### **Plot Area Size**

#### Plot area size

The Fit the plot area size to the enclosing window option is selected by default.

Click **Define the plot area size** to enter a **Plot area height** and **Plot area width**. The plot width is not applicable to triangular plots.

# **Global Settings: Console Mode Default Appearance**

To open this window, from the main menu select **Tools**  $\rightarrow$  **Options**. To change the name, buffer size, fonts and colours for a specific Console window, right-click the label for that particular Console and select **Properties**.

A Options		
General Graphical Mode	Console Mode	
Default Appearance Plot	ting	
Console name:	Console	
Buffer size:	40,000 🜩	
System output font:	AaBbCc123 AαBβΓγ	Modify
Command prompt font:	АаВЬСс123 АмВβГү	Modify
User input font:	ΑαΒbCc123 ΑαΒβΓγ	Modify
Background color:		Modify
Default directory:	C:\Users\amanda\Documents	Modify

#### **Console name**

The default is **Console** or enter another name to display.

### **Buffer size**

The default buffer size is 40,000.

<u>"Increasing the Buffer Size" on page 178</u> to adjust locally when in Console Mode.

### System output font

Click **Modify** to open the **Select Font** window and makes changes to the font type, style, size or color as required.

### User input font

Click **Modify** to open the **Select Font** window and makes changes to the font type, style, size or color as required.

### Command prompt font

Click **Modify** to open the **Select Font** window and makes changes to the font type, style, size or color as required.

### Background color

Click **Modify** to open the **Select Color** window to choose a background color for the Console Mode window where you enter commands.

### **Default directory**

Set the default directory file path to where the various file types are saved.

### Reset to original settings

In the lower corner of the window, click the **Reset to original settings** button to restore the original settings. You may have to expand the window to view the button.

# **Technical Notes**

### In this section:

<b>T</b>   ·   N   D'			254
Technical Note: Pipe or	Redirect Output Data fro	om Console Mode	

# Technical Note: Pipe or Redirect Output Data from Console Mode

Audience:	Advanced Programmers and Users
OS:	Windows, Linux, or Mac
Application:	For users who want to pipe or redirect data from Thermo-Calc running a macro in Console Mode.
Command examples:	<pre>Windows Thermo-Calc.bat macro.tcm &gt; results.log Linux Thermo-Calc.sh -E &lt; macro.tcm &gt; results.log Mac Thermo-Calc.command -E &lt; macro.tcm &gt; results.log</pre>
Command lin	e definitions
E	For Linux and Mac users, the Emacs style editing feature must be disabled (turned off) using the switch $-E$ in order for redirection and piping to work correctly with Thermo-Calc. For Windows this flag is not necessary.
<>	Redirection commands. The direction of each arrow indicates the flow of data. For example in ./Console.sh -E < my_commands.tcm > results.log, the contents of my_commands.tcm is used as input to the Console.sh script file and the resulting output is then written to results.log.

# Thermo-Calc Property Model Development Framework

Thermo-Calc Version 2018b



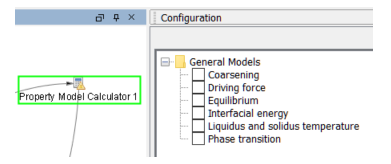


# Introduction to the Property Model Development Framework

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Only an overview is included here. Please use the standalone PDF or the Online Help for IDE and tutorial instructions. The PDF guide is available in the **Manuals** folder, accessible from the **Help** menu. There is also a subfolder containing the tutorial Python files.

The <u>"Property Model Calculator" on page 70</u> in Thermo-Calc Graphical Mode has predefined models available for immediate use and new models are being added on a regular basis. The *Property Model Development Framework* enables you to create custom property models that are then able to be included with the Property Model Calculator and ultimately used for your simulations.



When you use the *Property Model Development Framework API*, it extends the software to enable you to create property models. The models are written in the easy-to-learn programming language **Python** and the API provides seamless integration into Thermo-Calc. This includes access to thermodynamic and kinetic calculation routines as well as to the Thermo-Calc plotting system.

This API is like a plug in to Thermo-Calc. It is not an SDK. That is, you cannot use this API to build other applications to work with Thermo-Calc like you would use the *TC-API* or the *TQ-Interface*.

The API is based on *Jython*, which is a Python implementation running in the Java Virtual Machine. All Python libraries that are compatible to the Jython distribution can be used within the model (but not libraries based on native C-code).

Model development is possible without any additional installation by using a text editor (for example *Notepad++*). However, it is recommended you use a Python Integrated Development Environment (IDE), which provides you with advanced features such as debugging and auto-completion.

### PREDEFINED PROPERTY MODEL INSTALLATION LOCATION

8

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All property models (whether these are predefined or custom models) must be located in a subdirectory of the property model directory. You can go to Tools
 → Options and the Model directory field displays the default directory.

Predefined models included with the Property Model Calculator are installed in the locations below based on your operating system. If you create your own models, this is the default directory where these are saved. You can also use one of these as a template to create your own property model.

Operating system	Property Model installation directory				
Windows	C:\Users\Public\Public Documents\Thermo-Calc\ 2018b \PropertyModels				
Linux	/home/user/Thermo-Calc/2018b/PropertyModels				
Мас	/Users/Shared/Thermo-Calc/2018b/PropertyModels				

# **Property Model Tutorials: Python Files Available with Your Installation**

The tutorial Python files are available with your installation. These examples are also included with the PDF guide but placed in the installation folder to make it easier to copy the text and re-use it in your own Python files.

There is a PDF guide and tutorial examples included with your installation. In Thermo-Calc from the menu, select Help  $\rightarrow$  Manuals Folder. Then double-click to open the Property Model Development Framework folder. This guide is also available on our <u>website</u>.

TC-Python
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# Thermo-Calc Console Mode Command Reference

**Thermo-Calc Version 2018b** 





# **Introduction to the Command Reference**

### In this section:

About the Console Command Reference	2
Thermo-Calc Console Mode File Types	2

# **About the Console Command Reference**



This command reference is for Console Mode users of Thermo-Calc software. This guide describes the function and syntax of the commands available in the Thermo-Calc modules.



If you have the Diffusion Module (DICTRA) also see the *Diffusion Module* (*DICTRA*) *Command Reference* found in the Diffusion Module (DICTRA) Documentation Set PDF.

The commands available in all or most modules are described in <u>"General Commands" on</u> page 3.

The next topics are listed in alphabetical order based on the module names, then in each subtopic, the module commands.

### ABBREVIATIONS USED IN THIS GUIDE

- For the prompts,  ${\tt Y}$  stands for Yes and  ${\tt N}$  stands for No.
- TC is for Thermo-Calc. For example, default file names often start with tc.ini.

# **Thermo-Calc Console Mode File Types**

In Console Mode, Thermo-Calc uses different kinds of files, including:

- Macro Files (\*.TCM). These include log files created using the <u>"SET\_LOG\_FILE" on</u> page 300 command.
- Workspace Files in <u>POLY</u> (\*.POLY3), <u>GIBBS</u> module (\*.GES5), and the <u>PARROT</u> module (\*.PAR and \*.POP).
- Experimental Data Files (\*.EXP). See <u>"Commands in the Experimental Data Files</u> (\*.POP, \*.DOP)" on page 71.



For links to more information about this topic go to <u>"Macro, Workspace and</u> <u>Experimental Data Files" on page 173</u> in the *Thermo-Calc User Guide*.



If you are working in Console Mode*File Formats* in the *Diffusion Module* (*DICTRA*) User Guide.

# **General Commands**

### In this section:

BACK	. 4
EXIT	4
GOTO_MODULE	4
HELP (? and ??)	. 4
HELP	. 5
INFORMATION	. 5
MACRO_FILE_OPEN	. 5
SET_INTERACTIVE	. 8

# BACK

### "GOTO\_MODULE" below.

Syntax	BACK
	Switch control back to the most recent module. Going from the POST module (post-processor), BACK goes only to the TAB or POLY module (from where the POST module is entered).

# EXIT

Synta	EXIT
	Exit the program and return to the operating system. Use a save command before exiting otherwise all data and results are lost (in either the GIBBS, POLY, PARROT or REACTOR module).

# **GOTO\_MODULE**

Switch between modules. The name of the module must be typed. To get a list of available modules, press the <Enter> key.

Syntax	GOTO_MODULE
Prompt	Module name: <module name=""></module>
	If the <enter> key is pressed without typing a unique module name (or typing a ? mark) it prompts for the MODULE NAME and lists the available modules.</enter>

# HELP (? and ??)

Gives text help about the current prompt (whether this is the prompt of a module, or a prompt that requests you to specify the value of some parameter).



# HELP

Lists the available commands or explains a specified command. Specifying a unique command displays an explanation on screen. Typing a command abbreviation that is not unique lists all matching commands. Get the command information by typing a unique abbreviation or the complete command name.

Syntax	HELP
	Command: <command name=""/>
Prompt	<i>Command name</i> is the name of the command (one of the-module commands) to get help.
	If the <enter> key is pressed without typing a command name, then all the available commands are listed.</enter>

# **INFORMATION**

Get basic information about topics related to the module you are in.

Syntax	INFORMATION
	Which subject /Purpose/: <topic name=""></topic>
Prompt	Specify a subject (or its abbreviation as long as it is unique). Enter a question mark ? for a list of topics.

# MACRO\_FILE\_OPEN

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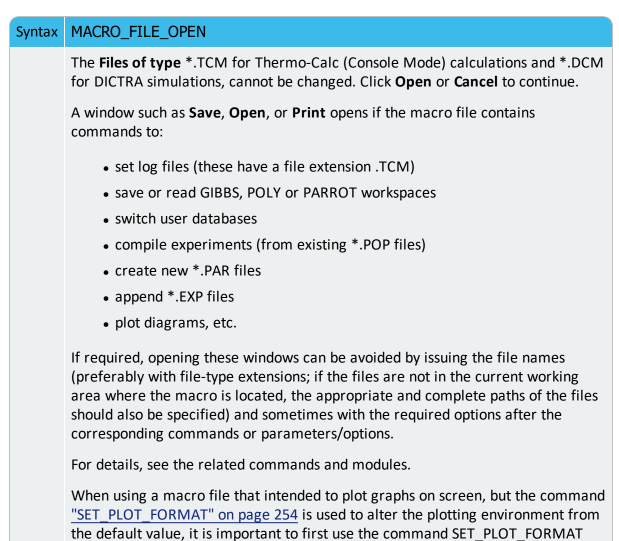
Some of the information in this topic is related to the add-on Diffusion Module (DICTRA).

The macro file functionality is a way to predefine sequences of Thermo-Calc commands stored in a macro file (a basic text file usually with the default extension \*.TCM or\*.DCM) and then executing all of the sequences using this command (preceded by the macro file name). This command can be used within various modules (i.e. SYS, POLY, PARROT, TAB and DICTRA\_Monitor).

### Syntax MACRO\_FILE\_OPEN

<Name of a macro file \*.tcm>

An **Open** window displays to specify the filename of the macro file (\*.TCM) with the macro command, so that the path (in the **Look in** field) and **File name** can be specified.



again to change back to the default value.

See example <u>tcex12</u> in the *Thermo-Calc Console Mode Examples Guide*.

This is useful when the same or similar calculations are made frequently with small changes [in terms of system definitions, data manipulations, conditions (for single points, stepping or mapping calculations), plotting settings, etc.). For example, use this feature when calculating phase/property diagrams during an assessment of thermodynamic data.

A macro file can be automatically generated in Console Mode if in the SYS module the <u>"SET\_UOG\_FILE" on page 300</u> command is used and a log file name (with a file extension \*.TCM) is given before any other SYS, DATA, TAB, GIBBS, POLY, POST, PARROT, or ED-EXP command or any special-module command (e.g. BIN, TERN, POTENTIAL, SCHEIL, POURBAIX, or REACTION). Such a log file generated in Console Mode is a text file. You can use a text

editor (such as Notepad++) to remove unnecessary command lines, modify some commands, settings and definitions, add some pause points, add comment lines that start with @@, and so forth. Then save it as a macro file with the standard file extension TCM.

Experienced users can also write/edit an appropriate macro file for calculations/simulations, using any basic textual editor outside the Thermo-Calc program.

All commands can be used in a macro file.



The file can be terminated with EXIT or interrupted with SET\_INTERACTIVE.

Within a macro file you can use comment-lines (for describing the problems and for explaining commands and inputs/outputs), which should always start with the @@ signs in the beginning of each comment-line. Such comment-lines help to document the macro file, while these are not considered as command lines and thus do not affect the proceeding of all the normal commands in modules of the Console Mode when the file is called by Thermo-Calc.

You can put multiple-line comment-blocks in a macro file between the @ (" and "@)" signs. The former sign indicates where the comment begins and the latter sign indicates where the comment ends. A comment-block begins from a line started with the *begin comment* sign @ (and ends with the *finish comment* sign @); all the lines written in between are ignored, and the line started with @) is also ignored.

Another use for a macro file is to allow you to interact at some specific points, using the "@?" sign starting a text line (note that the text describing the expected user-specification/input should be written as a continuous string without any empty space), for user's on-time specifications of arguments/parameters or inputs of parameter-values that are requested by a certain command. The macro temporally stops at the "@?" sign, prompt on screen the text given after "@?", and waits for the specified argument/parameter/value. Thermo-Calc then uses specified argument/parameter/value as the input(s) for the associated command. For example, you can input the values of lower and higher temperature limits for the second axis-variable as follows:

```
GO POLY-3
SET-AXIS-VAR 2 T @?Low-temperature-limit:
@?High-temperature-limit:
```

You can have macro-variables denoted by the signs of @#n (for definition) and ##n (for usage); and up to nine variables inside a single macro file. Such a macro-variable can be assigned with its desired value, as for example:

@#3First-element?

This writes the text (note that the text describing the expected user specification/input is written as a continuous string without any empty spaces) after the "@#3" sign as prompted on screen and then wait for a specification. The input is assigned to the macro-variable ##3, which is directly called in different parts within the current macro file.

For example, the content of the macro-variable ##3 is inserted in the command:

DEFINE-SYSTEM ##3

It is also useful in more complicated commands, such as:

SET AXIS VAR 1 x(##3) 0 1,,,

This command sets the mole fraction of macro variable 3 as axis 1.

A macro file can have any number of pauses at the @a signs, for the purpose of checking the details/results of executing certain commands when running the macro file. However, you can also prevent the Thermo-Calc (Console Mode) software from temporarily stopping at any pause by typing any character (except for the Y character for confirming a Yes answer to a command prompt) after specifying the name of a macro file.

A macro file can have a maximum of five nested levels, i.e. a macro file can call another macro file, and if one sub-level macro is terminated by the command it resumes at the next command in the previous macro. If it is terminated by end-of-file, the Thermo-Calc (Console Mode) software stops.

This feature is especially useful for alloy design that requires many (hundreds) of calculations/simulations on similar material system/processes (specified in many different but appropriately-documented macro files that are organized in up to five levels) during a certain period of time (e.g. in an evening), you can run the main macro (on the top level) at a certain time (e.g. before leaving the office) and afterwards (e.g. next morning) you can systematically and efficiently check/compare/analyse the results (saved as graphical files, and/or EXP, TXT, XLS... files).

By adding the SYS command <u>"SET\_ECHO" on page 299</u> at the beginning of a macro file (or in the primary macro file on the top level if any sub-level(s) of macro files are used), it is useful to automatically display the complete/detailed meaning of various commands in all the sequential operations in the software, all enforced according to the macro file(s).

### **SET\_INTERACTIVE**

### Syntax SET\_INTERACTIVE

Resets the input and output units to the initial values, i.e. keyboard and screen. Add this as the last command to the macro files.

# **BINARY\_DIAGRAM Commands**

In this	section:	
---------	----------	--

BINARY	DIAGRAM	Module	 	 	 	10

# **BINARY\_DIAGRAM Module**

The BINARY\_DIAGRAM module (short name, the BIN module) enables you to quickly calculate a simple binary phase diagram. Access to specific databases designed for BIN, such as TCBIN, is required.

To enter the module, at the SYS prompt type GOTO\_MODULE BINARY. There are no other commands for this module. Follow the prompts to plot a diagram.



Also see the Graphical Mode equivalent to this command, <u>"Binary Calculator"</u> on page 150 in the *Thermo-Calc User Guide*.

Syntax	BINARY_DIAGRAM
Prompts	Database: /TCBIN/
	First element
	Second element
	Phase diagram, Phase fraction (FE), G- OR A-curves (G/A): /Phase_ Diagram/

### **Database Commands**

.;:::

These commands are related to the DATABASE\_RETRIEVAL module, which is also known as the TDB module.

<u>See example tcex50</u> in the *Thermo-Calc Console Mode Examples Guide*.

Search the online help or see in the *Thermo-Calc User Guide* for step-by-step instructions to define a system.

### In this section:

AMEND_SELECTION	
APPEND_DATABASE	
DATABASE_INFORMATION	14
DEFINE_ELEMENTS	14
DEFINE_SPECIES	14
DEFINE_SYSTEM	15
GET_DATA	16
LIST_DATABASE	16
LIST_SYSTEM	
NEW_DIRECTORY_FILE	17
REJECT	
RESTORE	20
SET_AUTO_APPEND_DATABASE	21
SWITCH_DATABASE	22

<sup>&</sup>gt;\_

# **AMEND\_SELECTION**



This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

Use this command after defining the elements, species or the system (with the commands "DEFINE\_ELEMENTS" on page 14, "DEFINE\_SPECIES" on page 14, or "DEFINE\_SYSTEM" on page 15). The prompts allow changes to the predefined system. By answering Y or N, each of the selected elements, species or phases can be accepted or rejected (although this is not the case for the constituents or the entire system).

Syntax	AMEND_SELECTION
Prompt	Keep <name1> No/Quit/Yes/ Keep <name2> No/Quit/Yes/</name2></name1>
	Keyword Elements/Species/Phases name&
Options	Names of the pre-defined or pre-selected elements/species/phases

# APPEND\_DATABASE

>\_

This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

Append data from a file or additional database to the current set of data already read from another database. Data already read from another database and stored in the Gibbs energy system is kept in the GES5 workspace.

This command also enters all additional parameters (phase constituents, GO and interaction parameters, etc.) to already existing phases, and all existing parameters (phase constituents, excess model, GO and interaction parameters, etc.) in the phase are replaced with the values retrieved from the appending database.

The command is equivalent to the SWITCH\_DATABASE USER command sequence. It is also similar to <u>"SWITCH\_DATABASE" on page 22</u>, but does not reinitialize the DATA module and GIBBS workspace.

All the directly connected databases as predefined by the original database initiation file (TC\_INITD.TDB file in the /DATA/ area for Windows environments, or the initd.tdb file in the \data\ area for Linux), or by a user-specified database initiation file after using <u>"NEW\_DIRECTORY\_FILE" on page 17</u>, are listed after pressing <Enter> without giving any argument. You can supply your own database by giving the argument USER, the database name and the correct path if it is not located in the current working directory.

After the command is executed for the first time, the TDB\_XYZ: prompt (XYZ stands for the name of the primary switched database) is changed to the APP: prompt to go to more commands related to appending data.

Syntax	APPEND_DATABASE	
	Use one of these databases	
	PURE = SGTE pure element database	
Prompt	SSUB = SGTE substance database	
	User = User-defined database	
	Database name /XYZ/: <additional database="" name=""></additional>	
	Additional database name	
Options	The name of an existing database or a USER database definition file (***setup.TDB) that corresponds to the appending database.	
	If a USER database name or its path is not given on the same line of the APPEND_DATABASE command, or if the name or path is incomplete or incorrect, an <b>Open</b> window displays to open an appropriate database-setup. You can then open the USER database, or can cancel the open session; in the latter case, the program lists all predefined databases in the installation area, and you can select one of these or proceed with the USER option again for appending a desired database.	
	When this command is used in a macro (*.TCM) file, if the USER option is selected, the database setup file name (*setup.TDB) containing the setup definitions of the USER database, and its correct path, are required.	
Notes	In Linux, the file name of a USER database, or one of the predefined names, can be used under the prompt FILENAME: This is a valid file name for the USER database definition file (***setup.tdb) or a predefined database name, with the correct path. The default file name extension is <i>.tdb</i> .	
	This command can be called more than once if appending two or more databases to the primary switched database.	
	After this command, the commands to define a system (in terms of either elements or species), to reject/restore phases or species, and retrieve data must be repeated; however, the second keywords and parameter values might have to be different from the previous ones.	

# **DATABASE\_INFORMATION**

This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

### Syntax DATABASE\_INFORMATION

A short description of the current database is normally given by typing this command.

This can include information on covered systems, used models, valid ranges for parameters in temperature and composition, major applications, and so on.

### **DEFINE\_ELEMENTS**



This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

Define the system in terms of elements. All possible species that can be formed by the given elements are retrieved from the database. The names of elements must be separated with a space or comma. It is possible to use a wildcard \* after a common part of element names so that all the elements, which start with that common part and are available in the currently switched or appending database, are defined in the system. Up to 40 elements can be defined into a single system.



Also see <u>"Defining a System in Console Mode" on page 178</u> in the *Thermo-Calc User Guide* for step-by-step instructions to define a system with this command.

Syntax	DEFINE_ELEMENTS	
<b>D</b>	Element& <element1, element2,=""></element1,>	
Prompt	A list of elements to be defined into the system.	
	When appending database(s), this command, or <u>"DEFINE_SPECIES"</u> <u>below</u> or <u>"DEFINE_SYSTEM" on the next page</u> must be repeated with the same or similar elements as defined in the first switched database.	

# **DEFINE\_SPECIES**



This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

<sup>&</sup>gt;\_

Define the system in terms of species. Only those species given are retrieved. The different names of species must be separated with a space or comma. It is possible to use a wildcard \* after a common part of species names so that all the species, which start with that common part and are available in the currently switched or appending database, are defined in the system. Up to 1000 species can be defined in a single system.

Defining a System in Console Mode" on page 178 in the Thermo-Calc User Guide.

Syntax	DEFINE_SPECIES	
Duovoint	<pre>Species&amp; <species1, species2,=""></species1,></pre>	
Prompt	A list of species to be defined in the system.	
	When appending database(s), this command, <u>"DEFINE_ELEMENTS"</u> on the previous page or <u>"DEFINE_SYSTEM" below</u> must be repeated with the same or similar elements as defined in the first switched database.	

### **DEFINE\_SYSTEM**

L	>	
L	<u></u>	

This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

Define the system in terms of either elements (equivalent to <u>"DEFINE\_ELEMENTS" on the previous page</u>) or species (equivalent to <u>"DEFINE\_SPECIES" on the previous page</u>). Certain databases have a default value of the keyword (as either ELEMENTS or SPECIES) reflecting what is most appropriate when defining a system.

The different names of elements or species must be separated with a space or comma. It is possible to use a wildcard \* after a common part of elements or species names so that all the elements or species, which start with that common part and are available in the currently switched or appending database, are defined in the system.

When appending database(s), this command (or DEFINE\_ELEMENTS or DEFINE\_SPECIES) must be repeated with the same or similar elements as defined in the first switched database. Different databases might contain different elements and have different species definitions: avoid defining elements/species that are missing in the appending database(s). Otherwise, the program indicates these missing elements/species, and ignores them in subsequent steps. But additional elements/species, and additional phases not available in the first switched database can be defined and retrieved from the appending database(s). Up to 40 elements and 1000 species can be defined in a single system.



Also see <u>"Defining a System in Console Mode" on page 178</u> in the *Thermo-Calc User Guide* for step-by-step instructions to define a system with this command.

Syntax	DEFINE_SYSTEM
Prompt	Elements: <element1, element2,="">, or Species: <species1, species2,=""></species1,></element1,>
Options	Description
Species or Elements	Default keyword.
Element&	Specify a list of elements for the defining system.
Species&	Specify a list of species for the defining system.

# **GET\_DATA**



This command is for the DATABASE\_RETRIEVAL module (also called TDB module).



>\_

Only after executing this command can you go to any of the application programs such as GES, POLY or DICTRA and use the retrieved data. When appending database(s), this command must be repeated in order to obtain the additional system definitions, parameters and functions.

### Syntax GET\_DATA

Enter the defined system's elements, species, phases, and the connected parameters obtained from either the primary switched or additionally appending database(s) to the GIBBS and/or DICTRA workspace. This command is necessary for retrieval of all information concerning a defined system from the databank.

### LIST\_DATABASE

This is a DATABASE\_RETRIEVAL (TDB) command.

List all elements, species, phases or phase constituents in the database.

Syntax	LIST_DATABASE	
Options	Description	
Keyword	One of the keywords Elements, Species, Phases or Constituent must be used to indicate what to list	
Elements	All available elements, the reference state, atomic mass, H298-H0 and	

Syntax	LIST_DATABASE
	S298. Some elements have spaces in the column for the reference state. This implies that there are no parameters stored for this element.
Species	All available species together with the stoichiometric factors.
Phases	All available phases together with the number of sublattices and the number of sites in each sublattice.
Constituent	All available phases, the number of sublattices, the number of sites in each sublattice and the species dissolved in each phase. Species in different sublattices are separated with a colon (:). It is important to realize, for example, a phase can consist of Fe, Mo, V, and Cr, and its thermodynamic parameters can come from the binary systems Fe-Mo, Fe-V, Fe-Cr, and Mo-Cr. These data can give a relatively good description of the corners of the Fe-Cr-Mo system, but would most certainly give a bad one for the system Mo-Cr-V, due to the fact that the interaction parameters are, by default, set to zero, which originates from binary systems not included in the database.

# LIST\_SYSTEM

This is a DATABASE\_RETRIEVAL (TDB) command.

List all elements, species, phases or phase constituents in the defined system. It works only after a system is defined.

Syntax	LIST_SYSTEM
Keyword	One of the keywords Elements, Species, Phases or Constituent (as described in <u>"LIST_DATABASE" on the previous page</u> ) must be used to indicate what to list.

# **NEW\_DIRECTORY\_FILE**



>\_

This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

Open a new database initiation file (or called database directory file) generated by a local database manager or user, for accessing additional databases not predefined in the original database initiation file.

The original database initiation file is automatically copied by the installation script to the main database area, on a local computer for an independent installation locates or a connected server for a server installation.

- For Windows this is the TC\_INITD (or TC\_INITD.TDB) file in the \DATA\ area under the directory defined by the TCPATH parameter.
- For Linux, it is the initd.tdb file in /data/ area that is under the directory defined by the TC\_DATA parameter. If the Thermo-Calc (Console Mode) is run on a Linux platform, the new database initiation file must be located in the current working directory.

If there are too many databases at one installation or if there are some user-specified databases for a particular purpose, a local database manager can generate additional database initiation files, or each user can have their own initiation file.

The command switches the working initialization of database groups in the DATA module among the original and additional database initiation files. The Thermo-Calc (Console Mode) can use additional database initiation files that define accessing paths to database groups (databases located in different subdirectories under the directory defined by the TCPATH or TC\_DATA parameter). All directly accessible databases remain as in the same group, until this command is called or recalled.

Syntax	NEW_DIRECTORY_FILE
	<pre>File with database directory /tc_initd/: <database-initiation-file name=""></database-initiation-file></pre>
Prompt	Another database-initiation-file name
	The name of the next database initiation file (either additional or original) that is to be switched on as follows sessions.
	If the file name or its path is not given on the same line of the NEW_ DIRECTORY_FILE command, or if it is incomplete or incorrect, an <b>Open</b> window displays to specify the new database initiation file.

# **REJECT**



This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

Reject elements, species, phases or phase constituents that can form from the defined elements and species. Phases/species/constituents that are possible to form in the defined system are removed from the list of system phases/species/constituents (shown by the command <u>"LIST\_SYSTEM" on the previous page</u>). Phases/species/constituents that are not

included in the list cannot be entered without first being restored. The different names must be separated with a space or comma. You can use a wildcard \* after a common part of names so that all the elements/species/phases/constituents that start with that common part and are available in the currently switched or appending database, are rejected from the defined system.

This command can also reject a defined system, and thus reinitiate the entire DATA module memory and GES5 workspace.

Syntax	REJECT
Prompt 1	(Keyword = Elements or Species or Phases)
FIONDLT	<pre>If keyword: <name1, name2,=""></name1,></pre>
	(if keyword = Constituent)
Prompt 2	Phase: <phase name=""></phase>
	Sublattice number: <sublattice in="" number="" phase="" the=""></sublattice>
	Constituent: <constituent(s) in="" of="" phase="" sublattice="" the=""></constituent(s)>
Prompt 3	If keyword = SYSTEM the DATA module is reset to its initial state; GIBBS is reinitiated and data already entered to GES5 is lost.
Options	Description and additional information
Keyword	One of the keywords Elements, Species, Phases, Constituent or System must be used to indicate what is rejected.
Name&	Names of the pre-defined elements/species/phases/constituents/system
Elements	The given elements are rejected.
Species	The given species are rejected, making it impossible to form them from the defined elements.
Phases	The given phases are rejected, making it impossible to form them from the defined elements or species.
	The given constituent in one phase are rejected. Add the following when prompted:
	Phase: The name of the phase containing the constituent to be rejected.
Constituent	Sublattice number: The sublattice where the constituent enters (the first sublattice is 1). The question is omitted if there exists only one possible sublattice.
	Constituent: The name(s) of the constituent(s) to be rejected.

# RESTORE



This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

Restore already explicitly rejected elements, species, phases or constituents; it is the opposite of the command <u>"REJECT" on page 18</u> although it does not restore a completely rejected system.

Phases/species/constituents that are possible to form from the defined elements or species are entered to the list of system phases/species/constituents (shown by the command "LIST\_<u>SYSTEM" on page 17</u>). Phases/species/constituents that are not included on the list can now be added to the list. The different names must be separated with a space or comma. It is possible to use a wildcard \* after a common part of names so that all the elements/species/constituents, which start with that common part and are available in the currently switched or appending database, are restored in the defined system.

Syntax	RESTORE
Prompt1	(if keyword = ELEMENTS or SPECIES or PHASES)
	Keyword: <name1, name2,=""></name1,>
	(if keyword = CONSTITUENT)
Prompt2	Phase: <phase name=""></phase>
pt	Sublattice number: <sublattice in="" number="" phase="" the=""></sublattice>
	Constituent: <constituent(s) in="" of="" phase="" sublattice="" the=""></constituent(s)>
Options	Description and Information
Keyword	One of the keywords Elements, Species, Phases or Constituent must be used to indicate what is rejected.
Name&	Names of the pre-defined elements, species, phases, and constituents
Elements	The given elements are restored.
Species	The given species are restored and thus possible to form from the defined elements.
Phases	The given phases are restored and thus possible to form from the defined elements or species.
Constituent	The given constituent in one phase are restored.
	Add the following when prompted:

# Syntax RESTORE Phase: The name of the phase containing the constituent to be restored. Sublattice number: The sublattice where the constituent enters (the first sublattice is 1). Constituent: The name(s) of the constituent(s) to be restored.

# SET\_AUTO\_APPEND\_DATABASE



This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

Enforce an automatic action that appends thermodynamic data from a chosen database regarding the system to the data you already have about the system from the current database (that is, the default database or the database last chosen with <u>"SWITCH\_</u><u>DATABASE" on the next page</u> or <u>"APPEND\_DATABASE" on page 12</u>).



This is useful to simultaneously retrieve both thermodynamic and mobility data for a defined system when performing a DICTRA module simulation.

This command should be used before defining a system (with the commands <u>"DEFINE</u> <u>ELEMENTS" on page 14</u>, <u>"DEFINE\_SPECIES" on page 14</u>, or <u>"DEFINE\_SYSTEM" on page 15</u>) and retrieving the data with the <u>"GET\_DATA" on page 16</u> command from the primary switched database.

Syntax	SET_AUTO_APPEND_DATABASE
Prompt	Database name /XYZ/: <additional database="" name=""></additional>
	Additional database name
Option	The name of an existing database or a User database that corresponds to the automatically-appending database.

It works in a way that is similar to the APPEND\_DATABASE<DATABASE-NAME> command sequence, but only the phases that are also available in the primary switched database are appended. There is no possibility to manually list, reject and/or restore any phase that is available in the appending database. When retrieving data with the GET\_DATA command afterwards, all the phases that exist in the appending database but not in the primary switched database are automatically rejected. The prompt is kept as for the primary switched database, TDB\_XYZ: (where XYZ stands for the name of the primary switched database) until the execution of the GET\_DATA command.

Therefore, to selectively append more phases from a secondary database to a system that is defined and retrieved from a primary switched database, then the APPEND\_DATABASE command and sequential commands (such as DATABASE\_INFORMATION, DEFINE\_SYSTEM, DEFINE\_ELEMENT, DEFINE\_SPECIES, LIST\_SYSTEM, REJECT, RESTORE and GET\_DATA) should be used instead, before this command.

Although it is impossible to manually list, reject and restore any phase from the appending database, the DATA module automatically ignores all the phases that do not exist in both the primary switched database and the appending database, and append the data for the phases (that are also available in the primary switched database) from the appending database, as informed on screen when executing <u>"GET\_DATA" on page 16</u> afterwards.

# SWITCH\_DATABASE



This command is for the DATABASE\_RETRIEVAL module (also called TDB module).

Switch (or change) from the current database to a new one, and reinitialize the entire DATA module for defining a system and the GES5 workspace for storing retrieved data. All the directly connected databases as predefined by the original database initiation file, or by a user-specified database initiation file after using <u>"NEW\_DIRECTORY\_FILE" on page 17</u>, are listed by pressing the <Enter> key without giving any argument. You can also supply your own database by giving the argument USER and then the database name and its correct path if it is not located in the current working directory. The second part of the prompt TDB\_XYZ indicates the present database XYZ.

Syntax	SWITCH_DATABASE
Prompt	Database name /XYZ/:
	Use one of these databases:
	• PURE (SGTE pure element database)
	<ul> <li>SSUB (SGTE substance database)</li> </ul>
	• USER (user-defined database)
Options	Description and Information
New database name	The name of an existing database or a USER database definition file (***setup.TDB).
Database name	The new database is specified by typing the abbreviation given in the front of one of the available predefined databases. When switching/appending databases or additionally purchased databases you can add these to the

Syntax	SWITCH_DATABASE
	predefined database list in the database initiation file TC_INITD.TDB or initd.tdb of your installed software.
	When this command is used in a macro (*.TCM) file, if the USER option is selected, the database setup file name (*setup.TDB) that contains the setup definitions of the USER database, and its correct path, must be provided.
Filename	In Linux, the filename of a USER database, or one of the predefined names, can be used under this prompt. A valid filename for the USER database definition file (***setup.tdb) or a predefined database name, with the correct path. The default filename extension is .tdb.
Notes	If a USER database name or its path is not given on the same line of the SWITCH_DATABASE command, or if it is incomplete or incorrect, an Open window is displayed for the to-be-switched database.
	The USER database is used without reinitiating the Gibbs energy system, data from different databases can thus be combined. Therefore, the SWITCH_ DATABASE USER command sequence is equivalent to the APPEND_DATABASE USER command sequence. The result of a combination must be examined carefully, because differences in standard states, phase models and names can be disastrous. If the same parameters occur in several such switched or appended databases, the last one retrieved is used in the calculations. It is not advisable to use this method for large databases since these load slowly. After this command, those commands for defining system (in terms of either elements or species), rejecting/restoring phases or species, retrieving data, as well as for appending data from additional database(s), can be proceeded.

### **EDIT\_EXPERIMENTS Commands**

These commands are accessed through the PARROT module.

Some special commands are designed only for the ED\_EXP module and can also be used in \*.POP or \*.DOP files.

### In this section:

Running the EDIT_EXPERIMENTS Command	26
ADVANCED_OPTIONS	
COMPUTE_ALL_EQUILIBRIA	35
COMMENT	36
COMPUTE_ALL_EQUILIBRIA	
COMPUTE_EQUILIBRIUM	
CREATE_NEW_EQUILIBRIUM	38
DEFINE_COMPONENTS	
DELETE_SYMBOL	40
ENTER_SYMBOL	40
EVALUATE_FUNCTIONS	42
EXPERIMENT	43
EXPORT	44
FLUSH_BUFFER	45
GRAPHICS_PLOT	45
IMPORT	46
LABEL_DATA	46
LIST_ALL_EQUILIBRIA	47
LIST_CONDITIONS	47
LIST_EQUILIBRIUM	47
LIST_STATUS	48

LIST_SYMBOL	49
MAKE_POP_FILE	50
READ	51
READ_WORKSPACES	51
REINITIATE_MODULE	52
RESTORE_ALL_WEIGHTS	52
SAVE_WORKSPACES	52
SELECT_EQUILIBRIUM	53
SET_ALL_START_VALUES	53
SET_ALTERNATE_CONDITION	55
SET_CONDITION	58
SET_NUMERICAL_LIMITS	60
SET_REFERENCE_STATE	62
SET_START_CONSTITUTION	64
SET_START_VALUE	64
SET_WEIGHT	65
SHOW_VALUE	66
STORE_ALL_WEIGHTS	67
TABLE_HEAD, TABLE_VALUES and TABLE_END	67
TRANSFER_START_VALUES	69

# **Running the EDIT\_EXPERIMENTS Command**

To run the EDIT\_EXPERIMENTS (ED\_EXP) module:

- The first necessary command after entering the ED\_EXP module for the first time (with <u>"EDIT\_EXPERIMENTS" on page 133</u> in the PARROT module) should always be READ\_BLOCK in order to load the experimental data block from the current work \*PAR-file for editing. Also, the read command must also be used prior to any other ED\_EXP command if the module is reinitiated, or if no experimental data block has not previously been read from a work file compiled with a proper experimental data .POP/.DOP file, or if you want to change to another data block for editing.
- If any change is made in the ED\_EXP module remember to use the command <u>"SAVE</u> <u>WORKSPACE" on page 286</u> before going BACK to the PARROT module.



Although the TABLE\_HEAD, COMMENT and FLUSH\_BUFFER commands are visible in the ED\_EXP module, these cannot be used in the ED\_EXP module; these are only functional in an \*.POP or \*.DOP file.

### **ADVANCED\_OPTIONS**

Syntax ADVANCED\_OPTIONS

Prompt Which option? /Step\_and\_Map/: <Option>

"About the ADVANCED\_OPTIONS Command" on page 162.



LIST\_PHASE\_ADDITION has no additional prompt.

### STEP\_AND\_MAP (THE DEFAULT)

### Settings for MAP and STEP:

Global test interval /0/: <Integer number>

### Settings for MAP:

Automatically add initial equilibria /Y/: Number of mesh along an axis /3/: <Integer number> Use inside meshing points /N/: Global test interval /0/: <Integer number> The integer number determines how often the Global Minimization should be used during STEP\_WITH\_OPTIONS and MAP calculations. If it is set to 0 (zero), the recommended global test interval is used: i.e. every tenth step and at each phase change during STEP calculations, and only at node points during MAP calculations. Any other positive integer number, n, suggests it performs a Global Minimization test at every *n*-th step during STEP and MAP calculations. Of course, the Global Minimization test is always carried out at a phase change or a node point. The Thermo-Calc software is installed with 0 (zero) as the default value, but such a configuration for the default value can be changed by using SET\_TC\_OPTIONS in the SYS module.

Automatically add initial equilibria /Y/:

When <u>"ADD\_INITIAL\_EQUILIBRIUM" on page 160</u> is not used before MAP, a mesh of initial equilibrium points are added before the mapping itself takes place. This mesh is only available when two (2) axes are already defined. The default on start-up may be changed by SET\_TC\_OPTIONS in the SYS monitor.

Number of mesh along an axis /3/: <Integer number>

An integer for how many intervals of initial equilibrium points to be added along an edge in the diagram. For example, if set to 2 (two), initial equilibrium points are added at the beginning, center, and end of each axis-variable forming 2 intervals.

Use inside meshing points /N/:

If N the INITIAL\_EQUILIBRIUM mesh consists of initial equilibrium points added only along the edge/border of the diagram defined by the axis-variables. If Y, the initial equilibrium points added are also added to the inside edge.

### NONE

Take away any set advanced option in all subsequent single-point equilibrium and MAP/STEP calculations.

If there is any convergence problem in finding a stable solution, at any stage of an advanced option calculation conducted by an ADVANCED\_OPTIONS command-sequence, these messages display:

Convergence problems, increasing smallest site-fraction from 1.00E-30 to hardware precision 2.00E-14. You can restore using SET-NUMERICAL-LIMITS

The smallest site fraction in the current POLY3 workspace is automatically increased from the default value 1.00E-30 to the hardware-dependent precision (under Linux, as 2.00E-14). For other subsequent POLY-module calculation in the current TC run, you can use the POLY

command <u>"SET\_NUMERICAL\_LIMITS" on page 293</u> to restore or reset the smallest site fraction to the previous or another preferred value, as well as to reset other numerical limits.

### EQUILIBRIUM\_CALCULATION

The following prompts affect the ways how the POLY optimization does the Ordinary POLY Minimization to reach the minimum of an equilibrium state.

Settings for the minimization of an equilibria:

```
Force positive definite phase Hessian /Y/:
Control stepsize during minimization /Y/:
```

Choose Y or N to determine how to reach the minimum of an equilibrium state in a normal POLY optimization procedure. This is related to the special quantity, the phase stability function QF (phase), for all kinds of phases in an equilibrium state [A phase stability function for a phase is negative when the phase composition is inside a spinodal, and positive everywhere else].

If the composition of a solution phase is inside its spinodal, the enforcement of positive definite eigenvalues of a Hessian matrix (i.e. by answering Y) makes the step be taken towards the minima rather than the local maxima. Furthermore, if an eigenvalue (for a phase) of the Hessian matrix is near 0.0, the step-size is large and the answering Y sets a lower limit to this eigenvalue, and reduces the step-size and improves the convergence. The POLY command-sequence SHOW\_VALUE QF(phase) shows the size of the lowest eigenvalue of a phase in an equilibrium state.

By answering Y it might help the Ordinary POLY Minimization to converge, if there is such a problem, especially if QF(phase)=0 for phases.

By answering N the Ordinary POLY Minimization routine is applied.

For each phase in a defined system, the molar Gibbs energy of the phase is a function of the temperature-pressure condition and its composition:

$$G_m(X) = G_m(T,P;y_1,y_2,\ldots,y_n)$$

Compute a Hessian matrix as which describes the curvature of the Gibbs energy curve of this phase at the defined composition X:

$$\partial^2 G_m/\partial y_i\partial y_j$$

Diagonalize this matrix and call the Eigenvalues as e1,e2,...,en . The QF(phase) quantity for this phase is the lowest of these eigenvalues divided by the same values for a corresponding ideal (stoichiometric) phase:

```
QF(ph)=\min(\{e_1,e_2,\ldots,e_n\})/\min(\{se_1,se_2,\ldots,se_n\})
```

Here, se1,se2,...,sen are the eigenvalues for the ideal phase. Therefore, for an ideal (stoichiometric) phase, the QF(phase) should always be 1.0.

Control stepsize during minimization /Y/:

Choose Y or N to determine whether to introduce a control of step-size while reaching the minimum of an equilibrium state in a normal POLY optimization procedure. By answering Y it might help the POLY optimization to converge, especially if when some site fractions are less than 1E-4.

### **GLOBAL\_MINIMIZATION**

The following prompts affect how the Global Minimization is done. Settings for global minimization:

```
Use global minimization as much as possible /Y/:
Use global minimization for test only? /N/:
Maximum number of gridpoints per phase /2000/: <Integer number>
Use global minimization as much as possible /Y/:
```

Choose Y or N to decide if using the Global Minimization Technique from start of calculation or not. The default value on start-up may be changed by SET\_TC\_OPTIONS in the SYS module.

If Y the calculation is done when possible (depending on the condition settings), and a Global Minimization test is always performed when an equilibrium is reached. This costs more computer time but the calculations are more accurate.

If N the calculation is not used to reach the equilibrium state. Use global minimization for test only? determines if the Global Minimization test is made against the calculated equilibrium states obtained by the Ordinary POLY Minimization calculation.

Use global minimization for test only? /N/:

If Y a calculated equilibrium state obtained by the Ordinary POLY Minimization calculation is tested against the Global Minimization Technique after it is reached; and if found it as unstable, there is an error message.

If N the Global Minimization Technique is never tested for or done for all sequential calculations throughout the current TCC run. This implies that Global Minimization Technique is permanently turned off, and that consequently only the Ordinary POLY Minimization routine is used.

Maximum number of gridpoints per phase /2000/: <Integer number>

Specify the maximum number of grid points that are computed for each of the phases in the currently defined system, during the calculations enforced by the global minimization. More grid points give a higher robustness and take more computation time. The total number of computed grid points in all phases is limited to 2E6.

### IGNORE\_COMPOSI\_SET\_ORDER

```
Ignore composition set order /N/
```

### **NEW\_COMPOSITION\_SET**

The following prompts are to specify the additional composition set(s) to handle possible miscibility gap(s) of a certain solution phase:

Phase with miscibility gap: <Name of a phase>

Specify the name of the phase with miscibility gap.

New highest composition set number /2/: <#>

The default value is usually one higher than the current value. Each phase has initially one composition set. If a lower value is given, composition sets are deleted. You cannot take away the first composition set.

A message displays to show that you need to specify the composition for the composition set # (2,3,...).

Major constituent(s) for sublattice 1: /XX/: <YY>

The major constituents (YY) in each sublattice can be given. This may simplify giving start values when calculating the equilibrium as phases with miscibility gaps should have different major constituents for each composition set.

This prompt is repeated for each sublattice in the phase, sometimes even for all sublattices in the first composition set if such major constituents have not been specified.

### MAJOR\_CONSTITUENT

The following prompts are to specify the major constituent(s) on each of the sublattice sites

for a composition set of a certain solution phase:

Phase name: <name of a phase>

Specify the name of the solution phase with a new set of major constituents.

Composition set number /1/: <#>

The default value for the composition set number (#) is usually /1/ as the other composition sets are given major constituents when creating them. Each phase has initially one composition set.

Major constituent(s) for sublattice 1: /XX/: <YY>

The major constituents (YY) on each sublattice of a solution phase can be given. This may simplify giving start values when calculating the equilibrium as phases with miscibility gaps should have different major constituents for each composition set.

This question is repeated for each sublattice in the phase.

### PHASE\_ADDITION

The following prompts are to specify the additional energy term (always being a constant) of a given phase.

Phase name: <Name of a phase>

Specify the name of the (stoichiometric or solution) phase with the addition.

Addition to G per mole formula unit: <xxxxx>

The value (xxxxx) given is added to the Gibbs energy of the (stoichiometric or solution) phase. It can represent a nucleation barrier, surface tension, elastic energy or whatsoever.



It is not composition-, temperature- or pressure-dependent.

### PRESENT\_PHASE

Specify the name of the present phase a the name of the phase that should be present at all calculated equilibria:

Present phase: Phase name: <Name of a phase> Console Mode examples tcex45, tcex46 and tcex47 use this advanced option. See <u>"About the Thermo-Calc Console Mode Examples " on page 2</u> in the *Console Examples Guide*.

### **T-ZERO\_TEMPERATURE**

Name of first phase: <Phase A> Name of second phase: <Phase B>

The names of phases A and B must be given, for which the T0 temperature (where the Gibbs energies are equal) is to be calculated.

If the T0 temperature between the two specified phases is successfully calculated, a message displays, e.g.

The TO temperature is 840.82 K. Note: LIST-EQUILIBRIUM is not relevant

The first message shows the calculated T0 temperature between the two specified phases.

The second message indicates that after this option calculation with LIST\_EQUILIBRIUM is irrelevant and does not list the equilibrium for the system at the TO-temperature.

### PARAEQUILIBRIUM

```
Name of first phase: <Phase A>
Name of second phase: <Phase B>
```

The names of the two target phases A and B, between which the paraequilibrium state is to be calculated, must be entered subsequently or on the same (first) line at once then separated by an empty space, e.g. FCC#1 BCC or FCC#2 M23C6.



You need to understand what you are dealing with in terms of calculating a paraequilibrium state between the two specified phases.

Specifically, there are four distinguished cases to understand: (1) both chosen phases must have similar interstitial/vacancy sublattices where the fast-diffusion interstitial component (s) occupy; (2) the choice on the target phase pair must be reasonable for the defined system and specified initial overall composition; (3) both target phases should have phase constitution definitions that cover all the defined substitutional and interstitial components of the current alloy system; or (4) it is impossible to calculate the paraequilibrium state between the target phase pairs with given interstitial component(s) in the currently defined system.

Fast diffusing component: /c/: <interstitial component(s)>

```
Fast diffusing component: /none/: <interstitial component(s)>
```

The name(s) of the fast-diffusing component(s) (C as the default single component) must be given at the above prompts subsequently or at the same (first) prompt. It is possible to specify more than one interstitial component as fast diffusion species.

Note the following:

Such specified interstitial component(s) must be appropriately defined according to the phase constitution definitions of the two selected phases: these must be located on the interstitial/vacancy sublattices in both chosen phases;

If there is only one fast-diffusing component which is carbon, press the <Enter> key to accept the default input at the first prompt; if the single fast-diffusing component is another element (e.g. N), type its name at the first prompt;

If there are two or more fast-diffusing components (e.g. c and N), type their names at the above prompts subsequently or at the same (first) prompt (separated by an empty space, such as c N);

To finish the input of fast-diffusing elements, accept NONE at a repeated prompt, i.e. by pressing <Enter> key to start the paraequilibrium point calculation;

If NONE or a non-existing component name is typed at the first prompt, it means no back diffusion is to be considered, and the para-equilibrium calculation is thus canceled entirely.

If the paraequilibrium state between the two specified phases is successfully calculated, the messages displays e.g.

```
NP(FCC) = 0.3586 with U-fractions C = 2.71821E-02 N = 4.1548129E-03
NP(BCC) = 0.6414 with U-fractions C = 7.10061E-04 N = 2.3781027E-04
All other compositions the same in both phases
Note: LIST-EQUILIBRIUM is not relevant
```

The first and second lines list the phase amounts expressed in mole-percent [NP(phase)] and the contents of the interstitial components C and N in a specific phase expressed in the so-called u-fractions [u-f(phase,C) and u-f(phase,N)], for the phase A (in this case as FCC) and phase B (in this case as BCC), respectively. The third line states that the compositions of the matrix component and all the remaining compositions (regarding substitutional components) in both the target phase A and target phase B are the same at the current paraequilibrium state, while these are not shown on screen. The last line indicates that after this advanced-option calculation the LIST\_EQUILIBRIUM command is irrelevant and does not list the paraequilibrium state for the system at the current condition.

However, if the single-point calculation of the paraequilibrium state between the two specified phases has failed, these messages display:

```
*** Error 4 in ns01ad
*** Numerical error
```

This implies that the chosen target phase pair may be unreasonable for the defined alloy system or for the defined initial overall composition, or one or both phases may have inappropriate phase constitution definitions regarding the specified interstitial component (s). Then, you must either modify the settings of initial overall composition or specify the reasonable target phase pair with an appropriate choice of the fast diffusion interstitials in the defined alloy system.

#### STABILITY\_CHECK

```
Stability check on? /Y/:
Check also for unstable phases? /Y/:
```

If the automatic stability-check option is switched on choose to also check the stability for unstable phases. The default is Y and if an unstable phase is found to be located in a miscibility gas during a subsequent single-point equilibrium or MAP/STEP calculation, a warning message informs you to selectively make adjustments in the calculation settings [e.g. suspending the unstable phase, or using FORCED automatic start values for phase constituents by the S\_A\_S\_V F command-sequence, or creating other composition sets, etc.]. By answering N the stability check is enforced only to stable phases in the system.

#### TOGGLE\_ALTERNATE\_MODE

```
Set alternate toggle to DEFAULT, ALWAYS, or NEVER?
Set global toggle to DEFAULT, ALWAYS, or NEVER?
```

- DEFAULT: the experiment points are calculated according to the Alternate Technique depending on the SET\_ALTERNATE command.
- ALWAYS: the experiment points are always calculated according to the Alternate Technique even if the Alternate Mode is switched off in the PARROT module.
- NEVER: the experiment points are calculated as normal equilibria even if you SET\_ ALTERNATE in the PARROT module.

#### SHOW\_FOR\_T=

```
Temperature (K) /298.15/: <temperature_in_K>
```

Specify the new temperature condition (in K) under which the values of some specific state variable(s) or derived/partial variable(s) or defined symbol(s) for various thermodynamic properties of the entire system, of components, or of phases) in the currently-defined system (being in a *frozen* state) is shown on screen.

State Variable or symbol /VM/: <State variable or symbol name(s)>

Specify the name(s) of the desired state variable(s) or derived/partial variable(s) or defined symbol(s) for various thermodynamic properties of the entire system, of components, or of phases) in the currently-defined system. More than one state variable or symbol of interest can be simultaneously specified on the same line. For example, you can choose to show the values of VM or GM (i.e. molar volume or molar Gibbs energy of the entire system), or of VM(\*) or HM(\*).T (i.e. molar volumes or isobaric heat capacity of all phases), or of ACR(\*), DGM(\*) and LNACR(\*,FCC) [i.e. activities of all system components, driving forces for all phases, and activities (in logarithm) of all system components in the FCC solution phase], under the new temperature condition (being in a frozen state).

#### OUTPUT\_FILE\_FOR\_SHOW

A window opens with a default name for the file tc\_show.dat. Choose a location to save the file and click **Open**.

#### **COMPUTE\_ALL\_EQUILIBRIA**

#### Syntax COMPUTE\_ALL\_EQUILIBRIA

In ED\_EXP and PARROT modules, each experiment is treated as an individual equilibrium with some measured values. These are created with <u>"CREATE\_NEW\_EQUILIBRIUM" on page 185</u>, and stored in an experimental data \*.POP file and then compiled and saved in the POLY3 workspace of a PARROT work \*.PAR file.

With COMPUTE-ALL-EQUILIBRIA, all equilibria from the current to the last experimental points are calculated. If an equilibrium calculation fails, the calculation is stopped at that equilibrium. Equilibria with weight zero is skipped.

A current or present experimental point means the latest selected or calculated one. Once used the current point turns to the last point in the data block; in such a case you first use <u>"SELECT\_EQUILIBRIUM" on page 206</u> so that the current point is switched to a desired one.

For this command there is always a list output on screen, which consists of six columns for all experimental points available in the current data block. The first column is the *equilibrium identifier* (a number) assigned by this command, the second the *data label* assigned with <u>"LABEL\_DATA" on page 46</u>, the third the *number of iterations*, the fourth the

*current weight,* and the fifth the *current temperature*. In the sixth column the *fixed stable phases* are listed together with any *comment text* given after a <u>"COMMENT" on page 75</u> in the \*.POP or \*.DOP file.

If the weight is zero for an equilibrium columns 3-5 are replaced by the text *<unused>*. If the alternate mode is used for some experimental points, the listing is slightly different for such points. The 3-4 columns are displayed with \*alt\*, instead. If an alternate calculation is failed at one experimental point, the point is automatically assigned with a zero weight, and a warning message is shown above the data line (with all six columns).

#### **COMMENT**

This command is only used in the \*.POP/\*.DOP files to write brief comments or descriptive information on a newly created equilibrium (experimental) data point or a set of points given in a table.

Syntax	COMMENT
	A comment can have a maximum of about 60 characters written on the same line.
COM	PUTE_ALL_EQUILIBRIA

Syntax COMPUTE\_ALL\_EQUILIBRIA

In ED\_EXP and PARROT modules, each experiment is treated as an individual equilibrium with some measured values. These are created with <u>"CREATE\_NEW\_EQUILIBRIUM" on</u> <u>page 185</u>, and stored in an experimental data \*.POP file and then compiled and saved in the POLY3 workspace of a PARROT work \*.PAR file.

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*current weight*, and the fifth the *current temperature*. In the sixth column the *fixed stable phases* are listed together with any *comment text* given after a <u>"COMMENT" on page 75</u> in the \*.POP or \*.DOP file.

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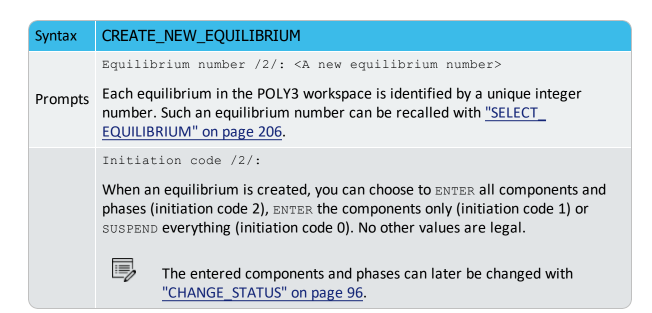
#### **COMPUTE\_EQUILIBRIUM**

The full equilibrium state is calculated for the given set of conditions. The Global Minimization Technique is by default enforced in this command (C\_E), while it can be disabled temporarily (for the current single-point equilibrium calculation) if using C\_E – or C\_E \* command-combination, or permanently (for all the sub sequential single-point calculations or stepping/mapping calculations within the current TCC run) if having decisively switched it off by a user (or possibly in some special modules) through changing the minimization option using the "ADVANCED\_OPTIONS" on page 167  $\rightarrow$  MINIMIZATION\_OPTION command-sequence.

Syntax	COMPUTE_EQUILIBRIUM
	The C_E command can be used, in order to enforce the ordinary POLY minimization routines in an equilibrium calculation; this is because of that the ordinary C_E command is now associated with the Global Minimization Technique, and only after the Global Minimization
	If technique is permanently switched off the C_E command makes no difference from the C_E – command-combination.
Prompt	Only certain types of equilibrium conditions [e.g. T, P, N, N( <component>), X (<component>), B, B(<component>), and W(<component>)] are fully supported in the Global Minimization mode (called <i>Direct Global Minimization</i>); and when other types of equilibrium conditions are used, after the initial POLY, a Global Minimization test and corrections are performed until the lowest minimum is found (called <i>Indirect Global Minimization</i>).</component></component></component></component>
	If there is any problem with convergence, you may try the C_E * command- combination. The character * enforces the command to use an advanced technique to obtain a complex equilibrium. However, after a successful C_E *calculation, you may repeat the C_E command and can check the status of phases, species or components (with <u>"LIST_STATUS" on page 200</u> $\rightarrow$ CPS command-sequence) and equilibrium conditions (with <u>"LIST_CONDITIONS" on page 198</u> ) and list out the calculation results (with <u>"LIST_EQUILIBRIUM" on</u>

#### Syntax COMPUTE\_EQUILIBRIUM page 198), because such actions may tell you how to further modify various settings for your current calculation. This command-combination is not that useful anymore, because the Global Minimization Technique that is always associated with the C E command is even more powerful and more precise in finding the most-stable equilibrium state in a complex heterogeneous interaction system; therefore, the C E \* command-combination is functional and can be used only after the Global Minimization mode has already been disabled temporarily or permanently. Some phases that are not stable in the current equilibrium state may not have their most favorable composition after this command, and thus their driving forces may not be correct. You can force the program to correctly calculate the driving forces of metastable phases, by giving repeated C E commands until the number of iterations (that is shown on screen after this command) is reduced to 2. Also see the POLY command "SET NUMERICAL LIMITS" on page 293 which can set the approximate driving force for metastable phases option on or off in all the subsequent POLY calculations within the current TCC run. If an equilibrium state for the defined system is not found, an error message is given. You can try repeating this command a few times, or change some of settings for the numerical limits, for starting variables and starting values, for starting constitutions of certain phases and for reference states of certain components, or to verify some of the defined conditions. CREATE\_NEW\_EQUILIBRIUM

# During data-assessments using the PARROT/ED\_EXP modules, you can, in the POLY module, create several equilibria with different sets of conditions and phases (but normally with the same set of components). By default, there is one equilibrium. To keep the set of conditions and phase for this equilibrium, create another one using this command, and use another set of conditions for that. Two equilibria may be useful to calculate easily the enthalpy difference between two states. In the PARROT module, the experimental information is stored as a sequence of equilibria.



# **DEFINE\_COMPONENTS**

Change the set of components. By default, the elements are used as components. The set of components can be important because some conditions are set using components, for example, the amounts, activities or chemical potentials.

For example, in the system Fe-Si-O, you can define FEO, FE2O3 and SIO2 as components, thus replacing the default FE, SI and O.

This implies a command <u>"REINITIATE\_MODULE" on page 204</u> and it should be given as the first command in the POLY module.

Syntax	DEFINE_COMPONENTS	
	Give all new components / existing components/: <new components=""></new>	
Prompt	The new components must be specified all on one line. These replace the existing component definitions.	
	The number of components cannot be changed with this command. Use <u>"CHANGE_STATUS" on page 96</u> instead.	
	To keep some existing components, it is recommended that you also enter these on the line. Otherwise, the added new components may not be defined correctly. This is especially important if some components are built out of several elements.	
	The new components do not have to be present as species.	

#### **DELETE\_SYMBOL**



This is a POLY3 module command.

Delete symbols (i.e. constants, variables, functions or tables. These symbols are entered with "ENTER\_SYMBOL" on page 109.

Syntax	DELETE_SYMBOL
	Name: <name a="" of="" symbol=""></name>
Prompt	Specify the name of the symbol to be deleted. Only one symbol can be deleted each time.

#### ENTER\_SYMBOL



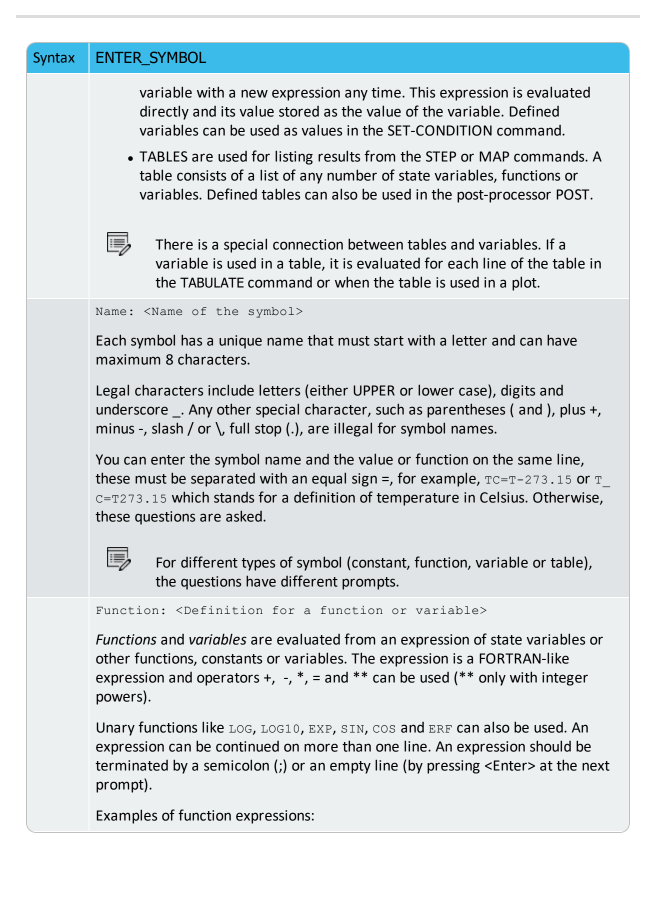
This command is specific to the ED-EXP module.



Also see <u>"ENTER\_SYMBOL" on page 109</u> for the GIBBS Module and <u>"ENTER\_SYMBOL" on page 238</u> for the POLY and POST Modules.

Symbols are a useful feature modules to define quantities that are convenient. Symbols can be constants, variables, functions or tables.

Syntax	ENTER_SYMBOL		
Prompt	Constant, Variable, Function or Table? /Function/: <keyword></keyword>		
	The Keyword can be specified as CONSTANT, VARIABLE, FUNCTION or TABLE.		
	<ul> <li>CONSTANTS can only be entered once and is a means of using a name for a numeric value. For example, the value of 1 atmosphere in Pascal can be denoted by P0 after the command ENTER CONSTANT P0=101325. Defined constants can be used as values in condition assignments, for example, SET-COND P=P0.</li> </ul>		
	<ul> <li>FUNCTIONS are expressions of state variables or other functions. These expressions are saved, and whenever a function value is requested all functions are evaluated. The reason for this is that they may depend on each other.</li> </ul>		
	<ul> <li>VARIABLES are similar to functions because they can also be expressions of state variables. However, contrary to functions, they are only evaluated when they are entered or if they are explicitly named in an <u>"EVALUATE_FUNCTIONS" on page 197</u> command. It is possible to enter a</li> </ul>		



Syntax	ENTER_SYMBOL	
	• GM(LIQUID): The Gibbs energy of liquid per mole component	
	• H.T/4.184: The heat capacity of the system in calories	
	• ACR(CR)/X(FCC,CR): The activity coefficient for Cr in FCC	
	• T-273.15: The temperature in Celsius	
	&: <continuation definition="" for="" of="" symbol="" the=""></continuation>	
The ampersand & means that you can continue to write the function on line if one line is not enough for the function. If you finish the function p <enter> again.</enter>		
	<pre>Value: <value a="" constant="" for=""> A constant can only be assigned a numeric value once.</value></pre>	
	Value or expression: <value a="" expression="" for="" of="" variable=""></value>	
	A <i>variable</i> can be assigned a numeric value or an expression. An expression is evaluated immediately and discarded. Only the numeric value is kept. This gives a possibility to save a value between calculations with different conditions because all state variables and functions are evaluated for the new conditions.	
	<pre>Variable(s): <variable(s) a="" in="" table=""></variable(s)></pre>	
	A <i>table</i> consists of a list of state variables or functions. One way to obtain results from a STEP command is through a table.	
	Example:	
	ENTER TABLE K=T,X(LIQ,C),X(LIQ,CR),ACR(C)	
	Which means that the table called K contains four columns, i.e. the temperature, the mole fractions of C and Cr in the LIQUID phase, and the activity of C.	
	To show the temperature in Celsius in a table, give the command ENTER FUNCTION $TC=T-273$ ; and then use the symbol $TC$ in the table.	
	& <continuation definition="" for="" of="" table="" the=""></continuation>	
	The ampersand $\&$ means that you can continue to write the table on the new line if one line is not enough for the table. If you finish the table press <enter> again.</enter>	

# **EVALUATE\_FUNCTIONS**

The value of one or more or all entered functions or variables are evaluated and listed.

# Syntax EVALUATE\_FUNCTIONS Name(s): <Name(s) of defined function(s)> The names of one or more entered functions or variables must be specified. By typing a wildcard \*, all functions and variables are evaluated. Image: Comparison of the synthesis of the synthe synthesis of the synthesis of the synthes

# **EXPERIMENT**

This command is used in original experimental data files (\*.POP or \*.DOP), but can also be given interactively inside the ED-EXP module, to change the value or uncertainty of an experiment or to add more experimental information.

An experiment usually consists of two parts with a colon (:) separating them:

- as a quantity relation, and
- as the uncertainty of the value for the quantity,

The quantity relation can be a normal POLY-module condition or an inequality (which is similar to condition but the relation between the quantity and given value is not in equality, i.e. < or >). The uncertainty can be expressed as an absolute value or relative to the value (x%).

An experiment that uses an inequality gives zero contribution to the sum of errors in the optimization procedure of PARROT if the value is on the right side. If the value is on the wrong side, the value after the colon determines how steeply the error increases with the value.

S	Syntax	EXPERIMENT	
given on the same line. The syntax of EXPERIMENT is similar to a CONDITION. Usually, a state var		The experiment must be typed after the command. Several experiments can be given on the same line.	
		The syntax of EXPERIMENT is similar to a CONDITION. Usually, a state variable set equal to a value with a given uncertainty is specified after the colon (:). For example, $T=1273.15:5$ .	
	<b>An inequality, &lt; or &gt;, can also be used. For example:</b> ACR(C)<-0.01:0.001, W(BCC,AG)>0.05:10%		

#### Additional Information About the Experment Command

A typical experiment added interactively specifies that a phase should not be stable in a certain experiment, because a phase may appear in a range of composition or of temperature where it has never been observed during the optimization. A phase is unstable if its driving force is negative, and you can add experimental information to enforce that. For example, you can suppress the HCP phase in an existing experimental point:

CHANGE-STATUS HCP=DORMANT EXPERIMENT DGM(HCP)<-.001:.0001



DGM is the POLY variable for the driving force that is already divided by RT, and it is dimensionless. Only dormant phases can have driving forces larger than zero, as this indicates that these would be stable if allowed to form. The experiment in the above case tries to modify the model parameters to make the HCP phase unstable.

More general examples of experiment definitions:

```
EXPERIMENT X(LIQ, PB)=.10:.01 X(FCC, PB)=0.02:.01
EXPERIMENT ACR(PB)=0.8:5%
EXPERIMENT T=700:5
```

The first experiment above describes a tie-line where the experimentally measured mole fraction of Pb is 0.1 in the liquid phase and 0.02 in the FCC phase, and its uncertainty is 0.01 for both measurements. The second experiment is that the activity of Pb should be 0.8 with an uncertainty of 5 percent.



The reference state of the component Pb must be set with the command <u>"SET\_REFERENCE\_STATE" on page 258</u>. The final one is that the temperature should be 700 K with an uncertainty of 5 degrees.

Experiments that are functions of states variable(s) must be described as the defined functions. For example:

```
ENTER_FUNCTION HTR=HM(LIQUID)-HM(FCC);
EXPERIMENT HTR=7000:1000
```

#### **EXPORT**



This command is for the ED-EXP module.

Transfer/export a calculated value from an equilibrium to an optimizing variable in the PARROT workspace. Thus the value can be used, for example, in the Gibbs energy description of a phase. Of course, this variable should not be optimized (i.e. it should be set as a fixed variable in PARROT).

Syntax	EXPORT		
	Function name: <function name="">#<n></n></function>		
Prompt	Specify the name of the function, the value of which should be transferred to a v variable. The number (n) of the v variable must be given after the function name, separated by hash character #.		
	The function name is UPPER/lowercase sensitive, and it should normally be given in UPPER case.		
	Example		
	<pre>Enter_function strngy=gm(fcc).x(cu);</pre>		
	Export strngy#6		
	This transfers the value of the partial derivative of the Gibbs energy of the FCC phase with respect to the mole fraction of Cu to the optimizing variable 6 (i.e. $v_6$ ).		

# **FLUSH\_BUFFER**

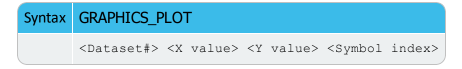
This command is only used in the \*.POP/\*.DOP files. It is needed if the number of experiments require more space than can be fitted into one buffer. When the PARROT program compiles experiments, it informs if any FLUSH commands are needed. The FLUSH\_BUFFER command terminates the current block, save it to the work file and starts a new block. With the READ command in the EDIT-EXP module, you can select the block to edit.

#### Syntax FLUSH\_BUFFER

After a FLUSH\_BUFFER command, the workspace is reinitiated and all functions or constants must be entered again in the \*.POP file. You can take advantage of this; for example, you can use the FLUSH command to have blocks with different components in one \*.POP file.

#### **GRAPHICS\_PLOT**

This command can help create a experimental data file from the \*.POP/\*.DOP file. Several data sets and different symbols are allowed.



#### **IMPORT**

Transfer/import the value of one of the optimizing variables to a constant. Normally, it is directly used inside an experimental data (\*.POP or \*.DOP) file. It is the inverse of <u>"EXPORT"</u> on page 44.

This is useful if several experiments are done at the same activity of a component, but this activity is not known. The activity should then be optimized and all equilibria with the same activity should use the same value. In this case the variable must be set as an optimizing variable, and an initial guess of the activity should be given. During the optimization, the PARROT program tries to find the activity that gives the best fit.



If an approximate value of the activity is known, it can be supplied as an experiment.



See <u>example tcex36</u> in the *Thermo-Calc Console Mode Examples Guide*.

Syntax	IMPORT		
	Constant name: <constant name="">#<n></n></constant>		
Prompt	The value of the $v$ variable must be assigned a symbolic constant. The number (n) of the $v$ variable must be given after the constant name, separated by hash character #.		
	The constant name is UPPER/lowercase sensitive, and it should normally be given in UPPER case.		
	Example		
	Enter_constant acu=0.1		
	Import acu#2		
	This transfers the value of the optimizing variable 2 (i.e. v2) to the constant ACU.		

#### LABEL\_DATA

Add a label to the experimental equilibrium point, either as a single point or several points given individually or in a table. The label is maximum four characters and must start with the letter A. Normally, it is directly used inside an experimental data (\*.POP or \*.DOP) file.

Several experimental equilibria can be given the same label, and the label can be used in the command <u>"SET\_WEIGHT" on page 65</u> to assign the same weight to all equilibria with the same label.

 Syntax
 LABEL\_DATA

 Label? /A1/: a<bcd>

 Prompt
 Specify a textual label (as maximum characters and must start with the letter A) for the current experimental equilibrium point(s).

# LIST\_ALL\_EQUILIBRIA

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		_

This is an EDIT\_EXPERIMENTS command.

#### Syntax LIST\_ALL\_EQUILIBRIA

List all the details (including equilibrium indicators, labels, iteration numbers, weight, and fixed phases or comments) of all equilibrium points from the current one to the last one on screen. You can always use <u>"SELECT\_EQUILIBRIUM" on page 206</u> to select a specific point as the current equilibrium point for such a list.

#### LIST\_CONDITIONS

~

This is a PARROT module command.

In the PARROT module, generate a list of the current values of optimization conditions, as well the current status of listing, either on screen or in a basic text file which can be opened and edited by a text editor.

Output to screen or file /Screen/: <File name>

Press <Enter> to output to screen or on the list file under a specific file name.

# LIST\_EQUILIBRIUM



Prompt

This is a POLY3 module command.

The result (always in SI units) from the last calculated equilibrium is listed on screen or in a textual file.



You can also execute this command if no calculation is made or if the calculation fails. It is your responsibility to interpret the results accordingly.

Syntax	LIST_EQUILIBRIUM
Dramat	Output to screen or file /Screen/: <file name=""></file>
Prompt	The name of the text file where the list of the calculation results shall be written.
	Options /VWCS/: <option(s)></option(s)>
	Select the output units and formats by optionally specifying a combination of these letters.
	• Fraction order: v means VALUE ORDER; A means ALPHABETICAL ORDER.
	• Fraction type: w means MASS FRACTION; x means MOLE FRACTION.
	<ul> <li>Composition: c means only COMPOSITION; N means CONSTITUTION and COMPOSITION.</li> </ul>
	<ul> <li>Phase: s means including only STABLE PHASES; P means including ALL NON-SUSPENDED PHASES.</li> </ul>
	Default options are vwcs. If the output fraction type should be in mole fraction (rather than mass fraction), then give vxcs or type x (implying that in this case the options V, C and S are accepted as the listing manners for fraction order, composition and phase).
	If accepting all the default options, or if accepting all the altered options that had already changed when using this command previously, you can type $L_E$ , , , or $LIST\_EQUILIBRIUM$ , , ,
	LIST_EQUILIBRIUM displays ORD (ordered) or DISORD (disordered) (when available). See <u>"CHANGE_LEGEND" on page 237</u> for details.

# LIST\_STATUS

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This command is for the POLY and ED-EXP modules.

List the status of components, species or phases.

Syntax	LIST_STATUS
	Specify what to list:
	Option /CPS/: <keyword(s)></keyword(s)>
Prompt	keyword = C or P or S, or any combination
	• c: list component status

Syntax	LIST_STATUS
	<ul> <li>P: list phase status</li> <li>s: list species status</li> <li>cs: list both components and species</li> </ul> Default is CPS.
	<ul> <li>The results depend on the key word specified in the options for <u>"CHANGE_STATUS" on page 96</u>, a table with the current status of phases or species or components, or the combinations, is shown:</li> <li>For components, the statuses and reference states are listed.</li> <li>For species, only the status is listed.</li> <li>For ENTERED and FIXED phases, the status, driving forces and equilibrium amounts are listed.</li> </ul>
	The metastable phases are listed in descending order of stability. To avoid long outputs only 10 metastable phases (in ENTERED status) are listed by lines, while all other less stable phases are merged onto one line. For DORMANT phases, their phase names and driving forces are listed. For SUSPENDED phases, only the phase names are listed.

#### LIST\_SYMBOL

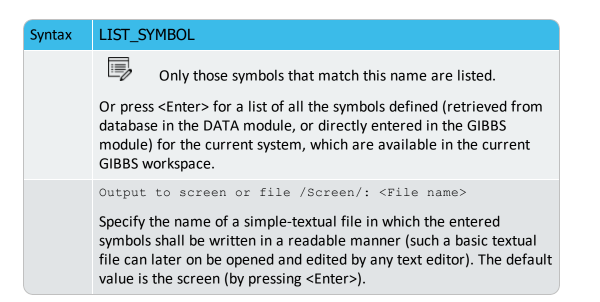
E	
I	>_

This is a GIBBS\_ENERGY\_SYSTEM (GES) command.

List the symbols available in the current GIBBS workspace, which are defined (retrieved from database in the DATA module, or directly entered in the GIBBS module) and used in TP-functions for the entered model parameters for various phases in the defined system.

For encrypted commercial databases you may not be able to use this command to list any retrieved symbol.

Syntax	LIST_SYMBOL
	Name: <symbol name=""></symbol>
Prompts	Specify a symbol name (either as the full name of a defined symbol, or as the first few common characters of several symbols which have names starting with such common characters).



# MAKE\_POP\_FILE

It is possible in the ED\_EXP module to make interactive changes and additions to the original experimental data file (the \*.POP or \*.DOP file).

To document or transfer the current or final experimental data to a \*.POP/\*.DOP file, you can use this command to write it to a new \*.POP/\*.DOP file or overwrite it on an old one.

Check this output carefully for errors. Tables in original \*.POP/\*.DOP files are written as individual experimental points. The output can be shown on screen or be saved as a new \*.POP file where to write or overwrite the edited experimental information in a simple text format.



If an existing \*.POP file name is specified, it is overwritten.



In the DICTRA module, a \*.DOP file is written or overwritten.

Syntax	MAKE_POP_FILE
	Output to screen or file /Screen/: <pop dop-file="" name=""></pop>
Prompt	Specify a name of the new *.POP/*.DOP file where to write or overwrite the edited experimental information in a simple text format.

# READ

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This command must be given each time the ED\_EXP module is entered, unless it is already given once in ED\_EXP and does not use <u>"LIST\_RESULT" on page 142</u> or <u>"OPTIMIZE\_VARIABLES" on page 148</u> in PARROT in between.

Syntax	READ
Prompt	Block number /#/: <n> The number of data block that should be edited must be given. If there is no command <u>"FLUSH_BUFFER" on page 75</u> in the original POP or DOP file, then there is only one data block with the number 1.</n>
	READ_WORKSPACE = READ This command (previously named READ_BLOCK) is equivalent to "READ_ <u>WORKSPACE" on page 286</u> in the POLY module, but it only reads the POLY3 workspace from the work file set by <u>"SET_STORE_FILE" on page 156</u> in the PARROT module.

#### **READ\_WORKSPACES**

This command is used in both the POLY and GES (GIBBS) modules.

The POLY3 and GIBBS workspaces and the calculated results from the MAP and STEP commands can be READ from a file where they are saved with <u>"SAVE\_WORKSPACES</u>" on the next page. Such an \*.POLY3 file is not printable.

Syntax	READ_WORKSPACES
Options	File name is the name of a saved POLY3-file where the POLY3 and GIBBS workspaces shall be read from must be specified. You do not need to type the extension if it is the default *.POLY3 (Windows) or *.poly3 (Linux), otherwise type the whole POLY-file name.
	A window opens so that the path (in the <b>Look in</b> field) and <b>File name</b> can be specified. The <b>Files of type</b> (i.e. POLY3) cannot be changed. Click <b>Open</b> to open the POLY3 and GIBBS workspaces from the saved *.POLY3 file.
	When reading back an original POLY3 workspace that has already been saved as an *.POLY3 file in the current Thermo-Calc (Console Mode) run or had been

#### Syntax READ\_WORKSPACES

read from an existing POLY3 file under the current work area, while some additional changes in the settings may be made but do not need to be kept in further steps in the current Thermo-Calc (Console Mode) run or any diagram is plotted in the POST module, you can type READ, , or READ\_WORKSPACE , ,

#### **REINITIATE\_MODULE**

Ensures that the whole POLY module (and thus the entire POLY workspace) is reinitiated to the state as it was when first entered.

#### Syntax REINITIATE\_MODULE

All the defined components, defined conditions, changed status, entered symbols, defined independent axis-variables, calculated starting equilibrium-points, added initial equilibrium points, stepped/mapped equilibrium data, and so forth, are removed completely. The saved file name is restored to the default.

#### **RESTORE\_ALL\_WEIGHTS**

Restore a specific weight-set (previously saved by <u>"STORE\_ALL\_WEIGHTS" on page 67</u>) and assign all the different experimental points in the current data block with the previously-set weights in the sequential assessments.

	Syntax	RESTORE_ALL_WEIGHTS
	Prompt	Set number (0 for list) /0/: <weight-set number=""></weight-set>
		The default value 0 is for a list on screen.

#### SAVE\_WORKSPACES



This command is for the ED-EXP module.

In the PARROT workspace this command saves the current status of the EDIT\_EXPERIMENT submodule before you switch back to the PARROT module. This is required when you want to save equilibria changes, including changes to conditions, experimental data, and/or computed equilibria results, as well as any newly added equilibria.

With this command all the changes made in the EDIT\_EXPERIMENT submodule are saved in the PARROT workspace. This means you do not need to execute <u>"SAVE\_PARROT\_</u> WORKSPACES" on page 149 in the PARROT module. However, when in the PARROT module, additional changes can only be saved with the SAVE\_ PARROT\_WORKSPACES command even if they have been made prior to executing SAVE\_ WORKSPACES in the EDIT\_EXPERIMENT module.

When you exit the EDIT\_EXPERIMENT module (or Thermo-Calc), the next time the file is opened it is in the same state as when you executed the SAVE\_WORKSPACES command. However, if you use <u>"OPTIMIZE\_VARIABLES" on page 148</u> the computed results are automatically updated.

Syntax	SAVE_WORKSPACES
Prompt	<file name=""> Sometimes good starting values ("SET_START_VALUE" on page 216) are required to successfully get or compute an equilibrium correctly. In this case, it is important to save the computed equilibrium in the PARROT workspace. It is important to remember that the computed results can be ruined by improper optimizations. For this reason, it is suggested you save the PAR file with a different name to back up the PARROT workspace.</file>

#### SELECT\_EQUILIBRIUM

If you create more than one initial equilibrium (during data-assessments using the PARROT/ED\_EXP modules), you can switch between them using this command.

Syntax	SELECT_EQUILIBRIUM
	Number /Next/: <choice equilibrium="" on=""></choice>
	Options: first, last, next, previous or present.
Prompt	Most commands affect only the PRESENT equilibrium. However, the commands <u>"REINITIATE_MODULE" on page 204</u> and <u>"DEFINE_COMPONENTS" on page 186</u> remove all the stored equilibria.

# **SET\_ALL\_START\_VALUES**

Set start values, e.g. if the calculation fails or if you have a miscibility gap or ordering. If temperature and pressure are not conditions, you are asked for values of them. Then for each phase prompt on if it should be stable and on its constitution.

Syntax	SET_ALL_START_VALUES
	T /XXXX/: <temperature in="" k=""></temperature>

Syntax	SET_ALL_START_VALUES
	If the temperature is not a condition, supply a guess of its final value (in K).
	P /100000/: <pressure in="" pa=""></pressure>
	If the pressure is not a condition supply a guess of its final value (in Pa).
	Automatic start values for phase constitutions? /N/: <y, f="" n="" or=""> <math display="inline"></math></y,>
	<ul> <li>For N, enter an initial amount of each entered phase and the major constituents or site fraction of each constituent.</li> </ul>
	<ul> <li>For Y, the program automatically sets the start values for phase constitutions in all possible phases.</li> </ul>
Prompts	<ul> <li>F is for force. In some cases the calculation fails because impossible conditions are set, e.g. W(C)=1.5 [W(C) is mass fraction and must thus be less than unity. The program tries to fulfill this condition by putting maximum amount of carbon in all phases, but can fail anyway. When you detect the error and set W(C) to 0.015, the calculation may still fail because it could start from the previous values. To get back to 'fresh' start values, choose F.</li> </ul>
	Should <phase> be stable /N/: <y 1="" 2="" n="" or=""></y></phase>
	A guess if this phase should be stable or not is requested. All entered phases are prompted for this question and next two.
	You cannot have more phases stable than you have components, but at least one (which dissolves all constituents) must be set as stable. For backward compatibility, this question can be answered by 1 (for Yes) or 0 (for No).
	The phase name may have a hash sign # followed by a digit, e.g. BCC_A2#2. For phases with miscibility gaps, there should be two phases with the same name but with different numbers after the hash sign.
	<pre>Major constituent(s): <name constituent(s)="" in="" major="" of="" phase="" the=""></name></pre>
	The constituent with the largest fraction in the phase should be specified. If there are more than one constituent with a large fraction, give them all on the same line. If the default major constituents should be used answer with an asterisk (*). By giving \$ the constitution is not changed. If there should be no major constituent give NONE and or if the major constituent(s) are improperly

#### Syntax SET\_ALL\_START\_VALUES

specified, you are asked for individual fractions in the phase.

Y(<phase>,<constituent>) /.XXXXXXXXX/: <.YYYYY>

The current value (.xxxxxxxxx) is default. You can accept the default by pressing <Enter> or give a new value (.YYYYY).

The phase name or constituent name may have a hash sign # followed by a digit, e.g. Y(BCC\_A2#2,FE), Y(BCC\_A2#2,C#2). For phases with miscibility gaps, there should be two phases with the same name but with different numbers after the hash sign. For phases with sublattices, the constituents in sublattice 2 or higher is also be suffixed with a hash sign # followed by a digit.

# **SET\_ALTERNATE\_CONDITION**

A special command used in the experimental data \*.POP/\*.DOP file but also possible in the ED\_EXP module. It is used only when you have specified that the alternate mode is set in the PARROT module.

The command syntax is almost the same as for the POLY command <u>"SET\_CONDITION" on</u> page 209 while the uncertainty should also be specified.

The alternate condition (including the normally POLY-module condition plus uncertainly; see below) must be given explicitly, but can be given on the same line or on separate lines with each one started with the command.

Syntax	SET_ALTERNATE_CONDITION
Prompt	State variable expression: <state expression="" linear="" name="" or="" variable=""></state>
	Give either a state variable or a linear expression of state variables.
Options	Description and Information
	Some of the state variables that can be used in conditions are:
	• $T$ : temperature in the system (in K)
State	<ul> <li>P: pressure in the system (in Pascal)</li> </ul>
variables	<ul> <li>N: system size (mole number in moles)</li> </ul>
	<ul> <li>B: system size (mass in grams)</li> </ul>
	• N( <component>): mole number of a component in the system</component>

Syntax	SET_ALTERNATE_CONDITION
	• X ( <component>): mole fraction of a component in the system</component>
	• W( <component>): weight fraction of a component in the system</component>
	• For the SET_CONDITION command this is ACR ( <component>): activity of a component in the system. For the SET-ALTERNATIVE_CONDITION command, this is ACR(<phase>,<component>) (activity of a component in a phase)</component></phase></component>
	<ul> <li>For the SET_CONDITION command this is MUR (<component>): chemical potential of a component in the system. For the SET-ALTERNATIVE_CONDITION command, this is MUR(<phase>,<component>) (chemical potential of a component in a phase)</component></phase></component></li> </ul>
	• X( <phase>,<component>): mole fraction of a component in a phase</component></phase>
	• W( <phase>, <component>): weight fraction of a component in a phase</component></phase>
	• ACR ( <species>, <phase>): activity of a species in a solution phase</phase></species>
	<ul> <li>MUR(<species>, <phase>): chemical potential of a species in a solution phase</phase></species></li> </ul>
	• H: enthalpy in the system (in J)
	• HM( <phase>): enthalpy of a phase (in J/mol)</phase>
	Factor: <a a="" continuation="" factor="" for="" or="" state="" the="" variable,=""></a>
Additional prompts	This question means that the previous question was not answered. The program is then expecting a single state variable or a complete state variable expression, or the numeric factor in an expression with only one state variable. In a state variable expression a state variable may be preceded by a constant factor. An example of this is:
	2*MUR(FE)+3*MUR(O)=-35000
	This means that it should be a condition that two times the chemical potential of FE plus three times the chemical potential of O should be -35000 J/mol.
	State variable: <a a="" continuation="" or="" specified="" state="" variable,=""></a>
	This question is prompted if a single state variable name has not given in either the prompt State variable expression or Factor, or a state variable expression is given but the expression is incomplete, for example, T- or 2*MUR(FE)+, for which the program is then expecting a continuation of the unfinished expression. You need to specify a state variable or a complete state variable expression, or complete the unfinished state variable expression. If a numeric factor is given before this prompt, only one state variable can be

Syntax	SET_ALTERNATE_CONDITION
	specified; otherwise, the program only takes the first state variable to complete the expression (i.e. the factor times the state variable).
	Value /X/: <a a="" constant="" numeric="" or="" value,="" variable=""></a>
	The value of the condition. This can be a numeric value, a constant or a variable. A suggestion is given as the default value. The special value NONE means that the condition is removed.
	Uncertainty /None/: <uncertainty condition="" of="" the=""></uncertainty>
	The uncertainty of the condition. This can be a numeric value, a constant or a variable. The default value NONE means that the uncertainty for the value specified above is zero. The uncertainty can either be expressed as an absolute value or relative to the value of the condition in percent (x%).
	There are more state variables that can be used in conditions. For more information, type an INFO STATE_VARIABLES command in the POLY module. A condition is normally a value of a single state variable with its value.
	FOR EXAMPLE:
	T=1273.15 P=1E5
	X(C)=.002
	W(CR)=1.5
	ACR (CR) =0.85
	X(FCC,C)=.001
<b>.</b>	H=-250000
Notes	HM (BCC) =-225000
	A condition can also be a value of a linear expression involving more than one state variable.
	FOR EXAMPLE:
	X(LIQ,S) - X(PYRR,S) = 0
	This means that it is a condition that the mole fraction of ${\rm s}$ should be the same in the LIQUID and PYRRHOTITE phases. In practice that should be the congruent melting point.
	After the equal sign only a numeric value is allowed.

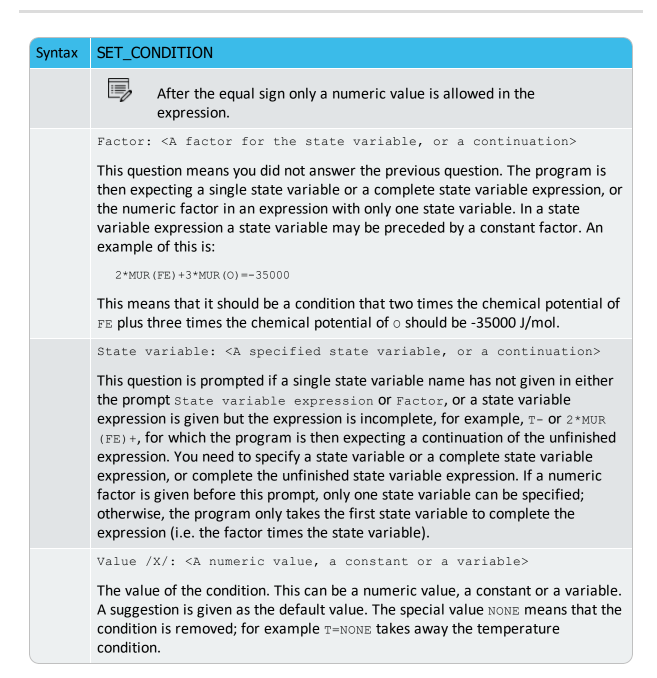
# **SET\_CONDITION**

Specify the equilibrium conditions for the calculation. All kinds of state variables, as well as most of the M/W/V/F-suffixed state variables (for normalization) and R-suffixed state variables (with respect to chosen reference states), can be used as conditions. Beside temperature and pressure conditions, a multicomponent system can have mixed types of conditions; this brings the extreme flexibility and powerfulness to the complex calculations with the Thermo-Calc software system. You should repeat this command for a multicomponent system, till the degree of freedom in the defined system becomes zero. When a FIXED phase status is used on a specific phase (use <u>"CHANGE\_STATUS" on page 177</u>), it is equivalent to one condition (implying that particular phase is stable in the defined system).

Syntax	SET_CONDITION
	Each condition must be given explicitly, but can be given on the same line (separated by a comma (,) or an empty space) or on separate lines with each one started with the command.
	EXAMPLE
	SET_COND T=1273, P=1E5, W(C)=.0015, X(LIQ,CR)=.22, ACR(N)=.2
	or
	SET_COND T=1273, P=1E5
	SET_COND $W(C) = .0015$ , $X(LIQ, CR) = .22$ ACR(N) = .2
	In this example, the temperature is set to 1273 K, the pressure to 1 bar (1E5 Pascal), the mass (weight) fraction of C to 0.0015 and the mole fraction of Cr to 0.22 and the activity of N to 0.2.
	<pre>State variable expression: <state linear<br="" name="" or="" variable="">expression&gt; or a linear expression of state variables.</state></pre> Some of the state variables that can be used in conditions are:
	• $T$ : temperature in the system (in K)
Prompt	<ul> <li>P: pressure in the system (in Pascal)</li> </ul>
Ρισπρι	• N: system size (mole number in moles)
	<ul> <li>B: system size (mass in grams)</li> </ul>
	• N( <component>): mole number of a component in the system</component>
	• X ( <component>): mole fraction of a component in the system</component>
	• W( <component>): weight fraction of a component in the system</component>

Syntax	SET_CONDITION
	• For the SET_CONDITION command this is ACR( <component>): activity of a component in the system. For the SET-ALTERNATIVE_CONDITION command, this is ACR(<phase>,<component>) (activity of a component in a phase)</component></phase></component>
	<ul> <li>For the SET_CONDITION command this is MUR (<component>): chemical potential of a component in the system. For the SET-ALTERNATIVE_CONDITION command, this is MUR(<phase>,<component>) (chemical potential of a component in a phase)</component></phase></component></li> </ul>
	• X( <phase>,<component>): mole fraction of a component in a phase</component></phase>
	• W( <phase>,<component>): weight fraction of a component in a phase</component></phase>
	• ACR ( <species>, <phase>): activity of a species in a solution phase</phase></species>
	<ul> <li>MUR(<species>, <phase>): chemical potential of a species in a solution phase</phase></species></li> </ul>
	• H: enthalpy in the system (in J)
	• HM( <phase>): enthalpy of a phase (in J/mol)</phase>
	There are many state variables that can be used in conditions. For more information, enter an INFO STATE_VARIABLES command. A condition is normally a value of a single state variable with its value.
	Example
	T=1273.15 P=1E5
	X(C)=.002
	W(CR)=0.5
	ACR (CR) =0.85
	X(FCC,C)=.001
	H=-250000
	HM(BCC)=-225000
	A condition can also be a value of a linear expression involving more than one state variable. For example,
	X(LIQ,S)-X(PYRR,S)=0
	This means that it is a condition that the mole fraction of ${\rm s}$ component should be the same in the LIQUID and PYRRHOTITE phases. In practice, that should be the

congruent melting point.



#### **SET\_NUMERICAL\_LIMITS**

Change the criteria for convergence. This is to speed up a calculation in a complex system.

If there is any convergence problem in finding a stable solution at any stage during a calculation procedure enforced by <u>"COMPUTE\_EQUILIBRIUM" on page 182</u>, <u>"STEP\_WITH\_</u> <u>OPTIONS" on page 217</u>, <u>"MAP" on page 201</u> or <u>"ADVANCED\_OPTIONS" on page 167</u>), this message displays:

Convergence problems, increasing smallest site-fraction from 1.00E-30 to hardware precision 2.00E-14. You can restore using SET-NUMERICAL-LIMITS

This implies that smallest site fraction in the current POLY3 workspace is automatically increased from the default value 1.00E-30 to the hardware-dependent precision (under Linux, as 2.00E-14). For other subsequent POLY-module calculation in Console Mode, you can use this command to restore or reset the smallest site fraction to the previous or another preferred value, as well as to reset other numerical limits.

Syntax	SET_NUMERICAL_LIMITS
	Maximum number of iterations /200/: <xxx></xxx>
Prompt	By default, the program tries 500 iterations before it gives up. As some models give computation times of more than 1 CPU second/iteration, this number is also used to check the CPU time and the calculation stops if 500 CPU seconds/iterations are used.
	Required accuracy /1E-6/: <yyy></yyy>
	This is a relative accuracy, and the program requires that the relative difference in each variable must be lower than this value before it has converged. A larger value normally means fewer iterations but less accurate solutions. The value should be at least one order of magnitude larger than the machine precision.
	Smallest fraction /1E-12/: <zzz></zzz>
	This is the value assigned to constituents that are unstable. It is normally only in the gas phase you can find such low fractions.
	The default value for the smallest site-fractions is $1E-12$ for all phases, except for IDEAL phase with one sublattice site (such as the GAS mixture phase in many databases) for which the default value is always as $1E-30$ (unless you have used the SET_NUMERICAL_LIMITS command to reset an even-lower value (e.g. $1E-45$ , that is naturally enforced to all the phases in the system).
	Approximate driving force calculation for metastable phases /Y/:
	Normally the POLY module only requires that the set of stable phases is in equilibrium in order to terminate the iterations. By default, the metastable phases are included in all iterations but these may not have reached their most favourable composition and thus their driving forces may be only approximate.
	If it is important that these driving forces are correct, choose N to force the calculation to converge for the metastable phases. This may require more iterations, and the STEP and MAP command may also terminate due to bad convergence in a metastable phase.

# **SET\_REFERENCE\_STATE**

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The reference state for a component is important when calculating activities, chemical potentials and enthalpies and is determined by the database being used. For each component the data must be referred to a selected phase, temperature and pressure, i.e. the reference state.

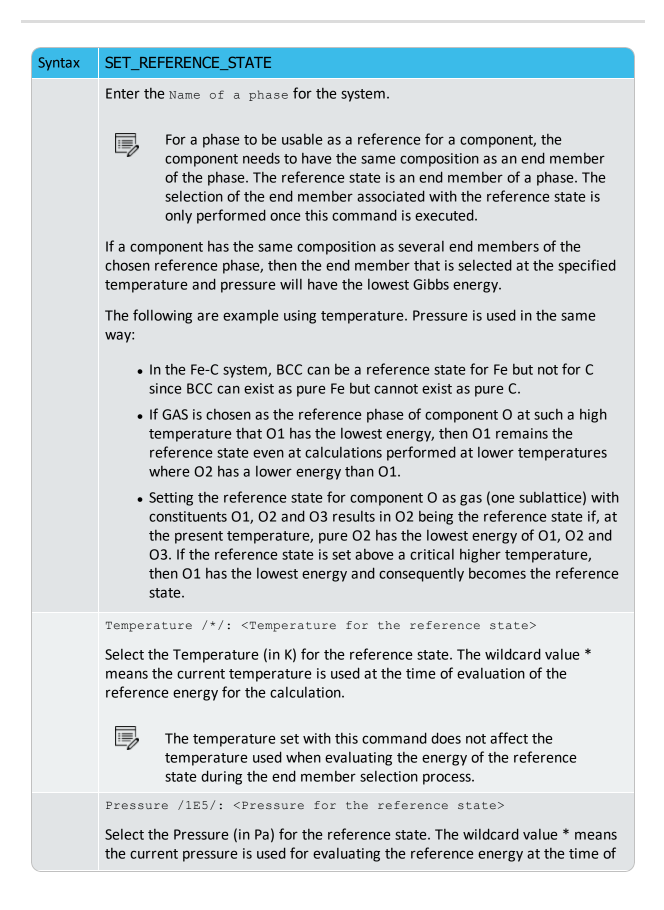
All data in all phases where this component dissolves must use the same reference state. However, different databases can use different reference states for the same element/component. It is important to be careful when combining data obtained from different databases.

By default, activities, chemical potentials and so forth are computed relative to the reference state used by the database. If the reference state in the database is not suitable for your purposes, use this command to set the reference state for a component using SER, i.e. the Stable Element Reference (which is usually set as default for a major component in alloys dominated by the component). In such cases, the temperature and pressure for the reference state is not needed.

In order to specify conditions in the specific reference state, you can append an R to the state variables. Thus, AC is the activity (of a system component or of a species in a phase) with respect to the default reference state, and ACR is the activity with respect to the selected reference state; MU is the chemical potential of a system component with respect to the default reference state, and MUR stands for the chemical potential with respect to the selected reference state. The AC and ACR variables, for both components in a system and species in a phase, can be specified in natural logarithm, e.g. LNAC(Fe), LNACR(C), LNAC (O2,GAS), LNACR(O2,GAS).

For the POST module, and after calculating with the <u>"STEP\_WITH\_OPTIONS" on page 217</u> or <u>"MAP" on page 201</u> commands, the reference state for a component can also be changed to plot various properties of the component in the entire system or in a specific phase. Then you can set the diagram axes as the chemical potential or activity quantities with the R suffix, i.e. MUR(comp), MUR(comp,ph), ACR(comp), ACR(comp,ph) or the common logarithms [e.g. LNACR(comp,ph)].

Syntax	SET_REFERENCE_STATE
Prompts	Component: <name component="" of="" the=""></name>
	The name of the component must be given.
	Reference phase: <name a="" as="" new="" of="" phase="" reference="" state="" the="" used=""></name>



Syntax	SET_REFERENCE_STATE
	calculation.
	EXAMPLES
	S-R-S Fe SER
	S-R-S Cr FCC * 100000
	S-R-S H2O AQUEOUS * 100000
	S-R-S ZE REF_ELECTRODE * 100000

# **SET\_START\_CONSTITUTION**

This command is similar to <u>"SET\_ALL\_START\_VALUES" on page 206</u>, but is used for an individual phase that may have a wrong constitution.

Syntax	SET_START_CONSTITUTION
Prompts	<pre>Phase name: <name a="" and="" constituent(s)="" major="" of="" phase,="" possible=""></name></pre>
	Specify the name of a phase for which the constitution shall be set.
	If there is a major constituent of the phase, you must specify this on the same line as the phase name. By giving an*, select the default major constituents. A \$ means keeping the same constitution, and NONE means to give individual site-fractions.
	Y( <phase>#<composition_set>,<constituent>#<sublattice>) /xxx/: <sf></sf></sublattice></constituent></composition_set></phase>
	The site fraction (SF) of the constituent shall be given. The default value $xxx$ is the last calculated one.

#### **SET\_START\_VALUE**

Set a start value of an individual variable.



This command is not often required as the automatic start values can be handled for most problems.

Syntax	SET_START_VALUE
Prompts	State variable: <name a="" of="" state="" variable=""></name>
	The state variable must be given.

Syntax SET\_START\_VALUE

Value

Specify the value to be assigned the state variable as start value.

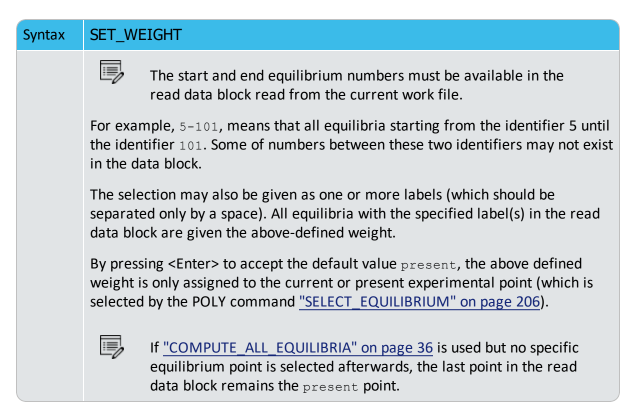
# **SET\_WEIGHT**

Each experimental value has an uncertainty associated with it, specified by the value after the colon (:) with the command <u>"EXPERIMENT" on page 43</u>. During an optimization the absolute difference between the experimental and calculated values gives a contribution to the sum of error.

With the SET\_WEIGHT command, you can change the scale of such a contribution (the uncertainty) for a single experimental point, or the contributions (all uncertainties) for a set of equilibria. The default weight is always unity. A value smaller than one means that the experiments should have less weight. A value larger than one that these should have higher weight. The special value zero means that the set of equilibria should be ignored in editing in ED\_EXP (as <u>"COMPUTE\_ALL\_EQUILIBRIA" on page 36</u> is given) and in <u>"OPTIMIZE\_VARIABLES" on page 148</u> in PARROT.

The weight may be needed to obtain a balance between different kinds of experiments. For example, if there are only five experimental values of the composition of a phase diagram but 500 experimental values of activities or enthalpies, then the five composition points may have to be given higher weight than unity, otherwise these are not well described by the optimizing procedure.

Syntax	SET_WEIGHT
	Value /1/: <weight value=""></weight>
Prompts	Specify a weight of the experiments in the specified equilibria (asked in the next prompt). The contribution to the sum of errors of these experiments are multiplied by this weight.
	The weight is squared, thus use 0.7 to make the error half (0.49) as big, and 1.4 to make it twice (1.96) as large.
	Equilibria (range) or label(s) /Present/: <selection></selection>
	Define the equilibria. These are available in the read data block from the current work file and are given the above-defined weight.
	The selection may be given as a range but the equilibrium numbers must then be separated only by a minus sign.



#### SHOW\_VALUE

Show the current value of any state variable, function or variable.

Syntax	SHOW_VALUE
Prompt	<pre>State variable or symbol: <name(s) (s)="" of="" or="" state="" symbol="" variable(s)=""></name(s)></pre>
	A single or several state variables or symbols (entered functions or variables) can be specified.
	The wildcard * can be used to denote all phases or all components.
	You can use the dollar-sign \$ to denote all stable phases. Thus, the command SHOW W(*,*)lists the mass fraction of all phases, whereas $SHOW \ W(\$, *)$ lists the mass fraction of all stable phases.
	If you SHOW a function, all functions are evaluated using the current values of the state variables. But if you SHOW a variable, it retains its value from the time it was ENTERED or last EVALUATED.
	Therefore to save a value from one calculation to another, ENTER it to a variable. This is frequently done in the PARROT module to transfer values

Syntax SHOW\_VALUE

8

between equilibria.

# STORE\_ALL\_WEIGHTS

Store a new weight-set that is the current situation of specified weights for all the different experimental points read from the currently-selected experimental data block in an assessment. This is useful when you want to try various weight-settings for some experimental points or for all the points, and then compare the assessment results from different weight-sets. Such a saved weight-set can be recalled and restored for all experimental data point with "RESTORE\_ALL\_WEIGHTS" on page 52 sequentially.

Syntax	STORE_ALL_WEIGHTS
	Command line: <weight-set name=""></weight-set>
Prompt	Give a weight-set name (comment line texts) for the current weight set for all experimental points.

#### TABLE\_HEAD, TABLE\_VALUES and TABLE\_END

These commands are used only in the \*.POP/\*.DOP files. It represents a convenient way to enter many experimental measurements of the same type in a table format.

These commands should always be used together and in a sequence, meaning that a table should always start with the TABLE-HEAD command, then follows the TABLE-VALUES command, and finish by the TABLE-END command. Between the two TABLEHEAD and TABLE-VALUES commands, there should exist some definition lines (e.g. on phase status, reference states, conditions, experiments, labels, etc.) for the experimental measurements of the current information type.

Syntax	TABLE_HEAD TABLE_VALUES TABLE_END
Prompt	
	After the TABLE-HEAD command, there must be an equilibrium description similar to a single experimental equilibrium but with some special notation. Then, there is always a TABLE-VALUES command, after which the actual data is

	TABLE_HEAD
Syntax	TABLE_VALUES
	TABLE_END
	given in columns. At the end of each table, there must be a TABLE-END command.
	The TABLE-HEAD command must be followed by a numeric value. This is used to generate unique numeric identifiers for all the equilibria in the table. The numeric identifier is incremented by one for each experimental (equilibrium) point in the table.
	Example
	An example of the use of a table for enthalpy measurements in the liquid is given here. Much more elaborate tables can be used
	\$ Enthalpies of mixing, Topor and Kleppa, Met Trans 1984 TABLE-HEAD 1
	CREATE-NEW 00 1
	CHANGE-STATUS PHASE LIQ=FIX 1
	SET-REFERENCE-STATE AU LIQ * 1E5
	SET-REFERENCE-STATE CU LIQ * 1E5
	SET-CONDITION P=1E5 T=1379 X(LIQ,AU)=01
	LABEL ALH
	EXPER HMR=02:5%
	TABLE-VALUES
	0.0115 -322
	0.0563 -1520
	0.8499 -2976
	0.9002 -2114
	TABLE-END
	The equilibrium description between TABLE-HEAD and TABLE-VALUES is similar as for a single experiment, except for these details:
	<ul> <li>@@ Used to automatically generate a series of identifying numbers instead of creating only one with the command <u>"CREATE_NEW_EQUILIBRIUM" on page 185</u>. The program automatically generates the identifying numbers starting after the number given after the TABLE-HEAD command (in this case it is 1).</li> </ul>

Syntax	TABLE_HEAD TABLE_VALUES TABLE_END
	• @1 values in the table are specified by an @ sign followed by a column number. In the above case, the mole fraction of Au in the liquid is in column 1 (see the line for SET-CONDITION). The columns may have other information than values, and you can use phase names or any text. If the text contains spaces or special characters, it must be surrounded by double quotes in the table, e.g. "ABC DEF&ghi".
	<ul> <li>For the syntax checker all commands describing the equilibrium must be in UPPER case after the TABLE-HEAD command.</li> </ul>
	• After the TABLE-VALUES command, there should be one line for each experimental measurement of the same type giving the values or texts that should be copied into the places of the condition or experiment defined by @1, t, etc. You can have columns that are not used but there must be exactly the same number of columns of each line, otherwise the syntax checker gives an error message in the PARROT module.

#### **TRANSFER\_START\_VALUES**

In the ED\_EXP module it is time consuming to calculate all equilibria when the optimization is sensitive to start values of the composition of the phases. Usually in one data block there are several experimental points with the same kind of equilibrium and each must have its start value set. When you manage to calculate equilibriums of such experimental points, this command is useful to transfer the site fractions from a successfully-calculated equilibria to the present experimental point.

Syntax	TRANSFER_START_VALUES
Prompt	From equilibrium /Previous/: <equilibrium number=""> The equilibrium number is the numeric identifier for the equilibrium from which the start values should be copied to the present experimental point. Previous is</equilibrium>
	the default if you press <enter>.</enter>

# **Experimental Data Files**

### In this section:

Commands in the Experimental Data Files (*.POP, *.DOP)	.71
TABLE_HEAD, TABLE_VALUES and TABLE_END	. 73
COMMENT	.75
FLUSH_BUFFER	.75

# Commands in the Experimental Data Files (\*.POP, \*.DOP)

In order to conduct an assessment for a system, the experimental data is described with a syntax, which is similar to the way that one calculates an equilibrium in the POLY module or you edit an experimental point in the ED\_EXP module. Similar to a Thermo-Calc macro file (\*.TCM), an experimental data file (i.e. the \*.POP or \*.DOP file) is a basic text file and can be opened and edited by any text editor. For this reason, an \*.POP or \*.DOP file is not hardware dependent.

A POP or DOP file consists of various commands from the POLY and ED\_EXP modules, as well some special commands which can only be used in such experimental data files. Various experimental information can be inputted in an \*.POP or \*.DOP file as different tables; the same type of experimental data are usually documented in the same table (see below).

An \*.POP or \*.DOP file is used in the PARROT module to provide experimental information for the optimization process, and is checked by a *syntax checker* when the PARROT command <u>"COMPILE\_EXPERIMENTS" on page 131</u> is proceeded.

Many POLY and ED\_EXP commands can be directly used in a POP or DOP file. But there are some differences with the set of POLY or ED\_EXP commands, and some specially designed commands as described in this topic can only be used in the \*.POP or \*.DOP files.

### FREQUENTLY USED COMMANDS

Most of the commands in the experimental data file are the same as in the POLY module. The most frequently used are listed below:

- "CREATE\_NEW\_EQUILIBRIUM" on page 185
- <u>"CHANGE\_STATUS" on page 177</u>
- "SET\_CONDITION" on page 209
- <u>"SET\_REFERENCE\_STATE"</u> on page 258
- <u>"ENTER\_SYMBOL" on page 238</u>
- <u>"SAVE\_WORKSPACES</u>" on page 52

As the last command in an \*.POP or \*.DOP file, you must always have the SAVE command.

### **OTHER COMMANDS**

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Other legal commands from the POLY module that are used less often are:

- "DEFINE\_COMPONENTS" on page 186
- <u>"EVALUATE\_FUNCTIONS" on page 197</u>
- "SET\_ALL\_START\_VALUES" on page 206
- "SET\_NUMERICAL\_LIMITS" on page 293
- <u>"SET\_START\_VALUE" on page 216</u>
- "ADVANCED\_OPTIONS" on page 167

The DEFINE-COMPONENTS command must be always used as the first command in an \*.POP or \*.DOP file, as it automatically reinitiates the whole workspace.

Most of the special commands for the ED\_EXP module are also often used in an \*.POP or \*.DOP file. For example:

- <u>"EXPERIMENT" on page 43</u>
- "EXPORT" on page 44

8

8\$B

- "IMPORT" on page 46
- "LABEL\_DATA" on page 46
- "SET\_ALTERNATE\_CONDITION" on page 55
- <u>"SET\_WEIGHT" on page 65</u>

The following special ED\_EXP commands are illegal and should not be used in any \*.POP or \*.DOP file:

- COMPUTE-ALL-EQUILIBRIA
- MAKE-POP-FILE
- READ (READ-BLOCK OR READ-WORKSPACE)
- TRANSFER-START-VALUES

The special commenting identifier, i.e. a dollar sign \$ which starts a line, is used to document comment lines at any position in an \*.POP or \*.DOP file. You may remember that this feature is universally the same as in almost all kinds of text files in Thermo-Calc

software, e.g. in \*.TDB, \*.DAT, \*.TCM, \*.POP/\*.DOP and \*.EXP files. Such a sign and all information after wards in the same comment line is skipped and ignored by the syntax checker.

# **TABLE\_HEAD, TABLE\_VALUES and TABLE\_END**

8

These commands are used only in the \*.POP/\*.DOP files. It represents a convenient way to enter many experimental measurements of the same type in a table format.

These commands should always be used together and in a sequence, meaning that a table should always start with the TABLE-HEAD command, then follows the TABLE-VALUES command, and finish by the TABLE-END command. Between the two TABLEHEAD and TABLE-VALUES commands, there should exist some definition lines (e.g. on phase status, reference states, conditions, experiments, labels, etc.) for the experimental measurements of the current information type.

	TABLE_HEAD
Syntax	TABLE_VALUES
	TABLE_END
Prompt	
	After the TABLE-HEAD command, there must be an equilibrium description similar to a single experimental equilibrium but with some special notation. Then, there is always a TABLE-VALUES command, after which the actual data is given in columns. At the end of each table, there must be a TABLE-END command.
	The TABLE-HEAD command must be followed by a numeric value. This is used to generate unique numeric identifiers for all the equilibria in the table. The numeric identifier is incremented by one for each experimental (equilibrium) point in the table.
	Example
	An example of the use of a table for enthalpy measurements in the liquid is given here. Much more elaborate tables can be used
	\$ Enthalpies of mixing, Topor and Kleppa, Met Trans 1984 TABLE-HEAD 1
	CREATE-NEW 00 1
	CHANGE-STATUS PHASE LIQ=FIX 1

	TABLE_HEAD
Syntax	TABLE_VALUES
	TABLE_END
	SET-REFERENCE-STATE AU LIQ * 1E5
	SET-REFERENCE-STATE CU LIQ * 1E5
	SET-CONDITION P=1E5 T=1379 X(LIQ,AU)=01
	LABEL ALH
	EXPER HMR=02:5%
	TABLE-VALUES
	0.0115 -322
	0.0563 -1520
	0.8499 -2976
	0.9002 -2114
	TABLE-END
	<ul> <li>The equilibrium description between TABLE-HEAD and TABLE-VALUES is similar as for a single experiment, except for these details:</li> <li>@@ Used to automatically generate a series of identifying numbers instead of creating only one with the command <u>"CREATE_NEW_EQUILIBRIUM" on page 185</u>. The program automatically generates the identifying numbers starting after the number given after the TABLE-HEAD command (in this case it is 1).</li> <li>@1 values in the table are specified by an @ sign followed by a column number. In the above case, the mole fraction of Au in the liquid is in column 1 (see the line for SET-CONDITION). The columns may have other information than values, and you can use phase names or any text. If the text contains spaces or special characters, it must be surrounded by</li> </ul>
	<ul> <li>double quotes in the table, e.g. "ABC DEF&amp;ghi".</li> <li>For the syntax checker all commands describing the equilibrium must be in UPPER case after the TABLE-HEAD command.</li> <li>After the TABLE-VALUES command, there should be one line for each experimental measurement of the same type giving the values or texts that should be copied into the places of the condition or experiment defined by @1, t, etc. You can have columns that are not used but there must be exactly the same number of columns of each line, otherwise the syntax checker gives an error message in the PARROT module.</li> </ul>

# **COMMENT**

This command is only used in the \*.POP/\*.DOP files to write brief comments or descriptive information on a newly created equilibrium (experimental) data point or a set of points given in a table.

Syntax	COMMENT
	A comment can have a maximum of about 60 characters written on the same line.

# **FLUSH\_BUFFER**

This command is only used in the \*.POP/\*.DOP files. It is needed if the number of experiments require more space than can be fitted into one buffer. When the PARROT program compiles experiments, it informs if any FLUSH commands are needed. The FLUSH\_BUFFER command terminates the current block, save it to the work file and starts a new block. With the READ command in the EDIT-EXP module, you can select the block to edit.

Syntax	FLUSH_BUFFER
	After a FLUSH_BUFFER command, the workspace is reinitiated and all functions or constants must be entered again in the *.POP file. You can take advantage of this; for example, you can use the FLUSH command to have blocks with different components in one *.POP file.

# Gibbs Energy System (GES) Commands

### In this section:

ADD_COMMENT	78
AMEND_ELEMENT_DATA	78
AMEND_PARAMETER	79
Solution Models in the GIBBS Module	82
AMEND_PHASE_DESCRIPTION	
AMEND_SYMBOL	
CHANGE_STATUS	
ENTER_ELEMENT	96
About the ENTER_PARAMETER Command	97
Defining the Parameter	
ENTER_PARAMETER	
About the ENTER_PHASE Command	
ENTER_PHASE	
ENTER_SPECIES	
ENTER_SYMBOL	
LIST_CONSTITUENT	
LIST_DATA	
LIST_PARAMETER	116
LIST_PHASE_DATA	117
LIST_STATUS	
LIST_SYMBOL	
READ_GES_WORKSPACE	
REINITIATE	121
SAVE_GES_WORKSPACE	122

SET_R_AND_P_NORM		12	22
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# **ADD\_COMMENT**

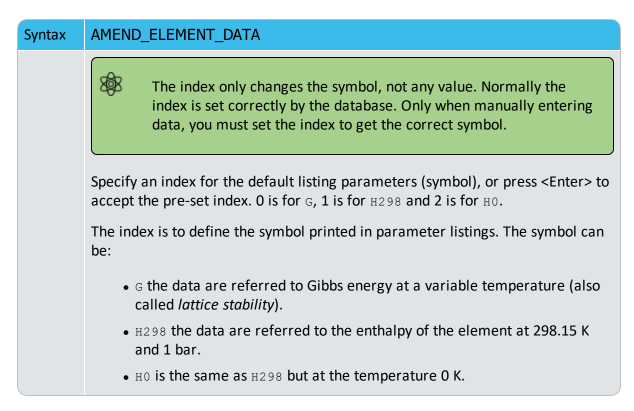
Syntax ADD\_COMMENT

Add a comment or make notes about parameters.

# AMEND\_ELEMENT\_DATA

The data for an element (in the SER, the Stable Element Reference, state) can be changed by this command. It should only be used for the elements that do not have any data in the database because the element data available in the database is set by the database-developer for the purpose of internal consistency. Except for the mass, the other values have no influence on the calculations.

Syntax	AMEND_ELEMENT_DATA
Deserves	Element name: <element name=""></element>
Prompts	Specify the name of the element for which you want to change the data.
	New stable element reference /ABCD/: <name of="" ser=""></name>
	Press <enter> to accept the default SER or specify a new SER for the element. Important: The default name should not be changed if this data is retrieved from a database. This name is used when parameters for a phase are listed and the database assumes that the stable element reference is the same as in the database. Only if the element's data have not been fetched from a database, e.g. entered manually, can you enter a new SER.</enter>
	New atomic mass /xx.xxxx/: <yyyyyyy></yyyyyyy>
	Press <enter> to accept the default atomic mass or specify a new value for the element. The atomic mass of the element is given in g/mol.</enter>
	New h(298.15)-h(0) /xxx.xxx/: <yyyyy></yyyyy>
	Press <enter> to accept the default H(298.15)-H(0) or specify a new value for the element. H(298.15)-H(0) is the enthalpy difference between 298.15 K and 0 K for the element in its SER state.</enter>
	New s(298.15) /xx.xxxx/: <yyyyy></yyyyy>
	Press <enter> to accept the default S(298.15) or specify a new value for the element. S(298.15) is the absolute value of the entropy at 298.15 K for the element in its SER state.</enter>
	Default element reference state symbol index /#/: <index></index>



# **AMEND\_PARAMETER**

>\_

This AMEND\_PARAMETER command is for the GES module. There is also a PARROT module command with the same name (<u>"AMEND\_PARAMETER" on page 126</u>).

Use this command to interactively change/amend the temperature-pressure function of a parameter. This is useful to correct typing errors because the old function is made available on the terminal for interactive editing. For most of these prompts, refer to <u>"ENTER</u> <u>PARAMETER" on page 137</u> for details.

Syntax	AMEND_PARAMETER
Prompts	Parameter: <parameter name=""></parameter>
	Specify a correct parameter name. If a parameter name is not acceptable or you only press <enter>, the error message displays:</enter>
	Error, please re-enter each part separately
	The program prompts for separate input for each part for a parameter name.
	Identifier (/X/): <g bm,="" l,="" or="" tc,="" v0="" va="" vb="" vc="" vk=""> <math display="inline"></math></g>

Syntax	AMEND_PARAMETER
	Specify one of these legal identifiers as described for ENTER_PARAMETER.
	Phase name (/ABCD/): <phase name=""></phase>
	Specify the phase name.
	Constituent (in sublattice # /ABC/): <species name=""></species>
	Specify the constituent name.
	<pre>Interacting constituent (in sublattice # /xyz/): <species name=""></species></pre>
	Specify the interacting constituent name: If there is no interacting constituent, press <enter>.</enter>
	<pre>Interacting constituent (in sublattice # /xyz/): <species name=""></species></pre>
	If there are more than one interacting constituents, specify them; otherwise press <enter>.</enter>
	Degree /#/: <degree></degree>
	Specify a numerical number as the degree of the phase parameter.
	After the parameter name is specified correctly, the program lists its current definition (either present in the database or defined by the ENTER_PARAMETER command), such as:
	L(PHASE2,AL,MG;1) =
	298.15 <t<2000.00: +5000<="" th=""></t<2000.00:>
	2000.00 <t<4500.00: +4500<="" th=""></t<4500.00:>
	4500.00 <t<6000.00: +4000<="" th=""></t<6000.00:>
	Then you are prompted to change the parameter definition, as shown below:
	Do you want to change the number of ranges /No/:
	Enter Y if you want to change the number of ranges for the chosen function, or change some of the temperature limits in the definition. Then retype both the low/high temperature limits and functions.
	If you do not want to change the number of ranges but want to change the function(s) in one or more ranges, press <enter> to accept the default N, then the whole definition of the chosen parameter in all ranges (if any) is listed on screen, such as:</enter>
	DIFFERENT FUNCTIONS IN THESE RANGES

```
298.15<T<2000.00
```

### Syntax AMEND\_PARAMETER

2000.00<T<4500.00

4500.00<T<6000.00

The prompt is given:

Do you want to change range limits /No/:

If there is more than one range, this question is prompted press <Enter>.

Range number (0 to exit) /0/: <Range number>

If the function of a parameter is different in two or more temperature ranges, you must specify the range of the function of which you want to amend. Or press <Enter> or type 0 to exit this command without making any changes.

```
Function:
```

The previous function is available for editing. The editing is performed within the general subroutine FOOLED, as described in <u>"AMEND\_SYMBOL" on</u> page 128. This routine prompts as follows:

1:+:>

The prompt consists of the current position in the string and the character at that position between colons, (::).

### COMMANDS

These commands can be given:

- Help: ?
- Move CP to last or first character: <+/-> A
- Delete characters from CP: <+-#characters> D
- Exit: E
- Find: <#occurrences> F<string>@
- Insert: I<string>@
- Move: <+-#positions> M
- Restore string: R
- Substitute: s<old>@<new>@
- Type string: T

Where CP denotes the current position in a string, # means number of, @ is a

# Syntax AMEND\_PARAMETER terminator of an input or search string. Image: When the string is typed the character at the current position is replaced by an underscore \_. To finish the editing of the current function, type E at the prompt.

# **Solution Models in the GIBBS Module**

The command <u>"AMEND\_PHASE\_DESCRIPTION</u>" on page 89 is used to specify/amend phase descriptions if a phase has a miscibility gap, uses a special excess energy model, or has a specific additional contribution to its Gibbs energy, for example.

This topic outlines the application of each option and includes important information about each command.

There are many solution models implemented in the GIBBS module (type INFO MODELS to see details about various thermodynamic models). However, these are switched on and handled differently via various GIBBS routines/commands:

- The first models the non-ideality as excess parts, i.e. by amending the phase's EXCESS\_MODEL and/or TERNARY\_EXTRAPOLAT models.
- The second models the non-ideality as ADDITIONAL parts, i.e. by amending the phase's other subjects in this command, e.g. MAGNETIC\_ORDERING for the Magnetic Ordering Model, DISORDERED\_PART for the CVM Approach in modelling chemical ordering/disordering phenomenon, QUASICHEM\_IONIC for using a quasi-chemical entropy term for an ionic two-sublattice liquid solution phase, QUASICHEM\_FACT00 or QUASICHEM\_IRSID for describing a substitutional liquid solution phase, DEBYE\_HUCKEL for the DHLL/SIT Model in describing a dilute aqueous solution phase, etc.
- The third implements the model entirely or partially into the GIBBS module and related database(s), such as for the electrostatic contribution in an AQUEOUS solution phase by the Complete Revised\_HKF Model, the Murngham Model, Birch-Murngham Model or Generalized PVT Model for high-pressure/temperature volume contribution in a solid or liquid phase, the SUPERFLUID Model for the non-ideal EOS and non-ideal mixing in a gaseous mixture.

### NEW\_CONSTITUENT

Use this if you want to add a new constituent to a phase. It is illegal to add new constituents to an ionic two-sublattice liquid phase.

### **RENAME\_PHASE**

Use this to change the names of some specific phases. For example, a phase called *FE3O4\_S* may be better named *Magnetite* to help identify it during the calculations and postprocessing. This is also a way to delete a phase by hiding it under a new name.

### SITE\_RATIOS

Use this to change the number of sites (i.e. the stoichiometric coefficients of various sublattices) in a sublattice phase.

### **COMPOSITION\_SETS**

Use this for solution phases that may have miscibility gap(s). However, this is less important today and often unnecessary to define additional composition set(s), since the implemented Global Minimization Technique can usually detect and add such composition set(s) in an automatic manner where it is really necessary during equilibrium calculations (of single-points, stepping or mapping).

### MAJOR\_CONSTITUENT

Use this to set major constituent(s) on each sublattice in each composition set for a solution phase. This is useful to make calculations converge faster because it may simplify giving start values when calculating the equilibrium as those phases with miscibility gaps should have different major constituents for each composition set. The databases often set major constituents for several phases automatically when data are retrieved.

### FRACTION\_LIMITS

Use this to limit the application range (in terms of mole-fractions of all the involved elements) of a particular solution phase. This is useful to avoid automatic creations (enforced by the Global Minimization Technique) of additional composition sets for some solution phases (such as AQUEOUS solutions or dilute Fe-based liquid mixtures) of which the used models [e.g. the SIT Model or the Complete Revised\_HKF Model for AQUEOUS solution, or the modified dilute solution parameters (plus a quadratic term, according to Hillert (1986) based on the SigworthElliot Model (Sigworth and Elliot, 1974) for Fe-rich liquid mixture] cannot be appropriately applied on a full scale.

Globally set the composition limits (in terms of mole fractions of various elements) in a specific solution phase, so that whenever the program finds a potential phase composition or a new composition set of possible miscibility gap(s) but that is out of this globally-set

composition range, the program automatically ignores such a phase composition in an equilibrium calculation. This can be done either permanently inside a database (by enforcing this option for amending the phase-description of the solution phase) or temporarily within the GIBBS module (by using this phase-description amendment option).

This is important for solution phases for which the thermodynamic models and assessed data for the phases are for specific composition ranges, for example, the FE\_LIQUID phase in the SLAG database is only applicable for Fe-rich liquid mixture, and the AQUEOUS phase in the TCAQ/PAQ and AQS/PAQS databases is only applicable for H2O-dominant aqueous solution phase.

Specify the low and high mole-fraction limits for each of the elements possibly available within the considered phase (in the currently defined system, or in a certain database). Do this for all the possible elements (defined in the phase) in a single AMEND\_PHASE\_ DESCRIPTION FRACTION\_LIMITS command-sequence, for example:

TYPE\_DEFINITION R GES AM\_PH\_DES FE\_LIQUID FRACTION\_LIMITS Fe 0 0.6 Ag 0 0.01 Al 0 0.05 Ca 0 0.05 Co 0 0.01 Cr 0 0.01 Cu 0 0.02 Mg 0 0.05 Mn 0 0.05 Mo 0 0.05 Nb 0 0.05 Ni 0 0.05 Pb 0 0.05 Si 0 0.10 Sn 0 0.02 Ti 0 0.05 U 0 0.01 V 0 0.02 W 0 0.02 Zr 0 0.03 B 0 0.01 C 0 0.01 H 0 0.01 N 0 0.01 O 0 0.01 P 0 0.01 S 0 0.01 !

### DISORDERED\_PART

This command is needed for the special treatment of chemically-ordered phases where the contributions from the disordered state are described by a phase without ordering sublattices (the disordered phase name must be specified).

Several checks are made that the ordered and disordered phases are compatible (sublattices, sites and constituents). A link is then created between the phases, and the disordered phase is hidden from application programs. The Gibbs energy for the ordered phase also includes the Gibbs energy for the disordered phase.

Phases which can have an order/disorder transformation have parameters split on two phases and are referred to as the *two phase* description in the GIBBS module. One of them has sublattices for chemical ordering, the other one represents the disordered state. Normally, the ordered BCC and FCC or HCP phases may have either 2 or 4 substitutional sublattice (plus one additional interstitial site), that are handled by the Two Substitutional-Sublattice Ordering Model or Four Substitutional-Sublattice Ordering Model, respectively. *Two phase* means that the *ordered* phase has parameters that describe the ordering. The *disordered* phase has all parameters for the reference state and those for describing the disordered phase. The ordered phase may occur only is some systems whereas the disordered phase may occur frequently, typical examples are the disordered FCC and BCC which may become ordered as L12 or B2 in certain systems. In order to treat multicomponent systems where some subsystems have ordering it would be necessary to transform all parameters of BCC into a B2 model.

The parameters describing the disordered phase are not changed but the Gibbs Energy system is informed that the Gibbs Energy for the two phases should be added.

The method used calculates the Gibbs Energy of a two phase model with two or four substitutional sublattices for ordering.

```
G_m = G_m^{ord}(y_i',y_i') + G_m^{dis}(x_i) - G_m^{ord}(y_i'=x_i,y_i''=x_i)
```



The mole fractions *x*i are calculated from the site fractions *y*i. The last term means that the contribution from Gmord in a disordered state (same site fraction, equal to the mole fraction, on both sublattices) is subtracted. The effect of this is that the parameters in the ordered phase have no contribution to the disordered state.

### MAGNETIC\_ORDERING

Change the magnetic ordering parameter for a certain phase with magnetic ordering contribution, in terms of its anti-ferromagnetic factor. By default this is -1 for BCC phase and -3 for all other phases (FCC, HCP, etc.). The fraction value of the total enthalpy (due to short-range ordering above the magnetic transition temperature) is by default 0.40 for BCC phase and 0.28 for all other phases (FCC, HCP, etc.).

### EXCESS\_MODEL

See <u>example tcex52</u> in the *Thermo-Calc Console Mode Examples Guide*.

Use this to change the default Excess Model (for interaction energies) from the default (REDLICH-KISTER\_MUGGIANU). The model handles the excess interaction energies in a solution phase is chosen from:

- REDLICH-KISTER\_MUGGIANU: for binary (R-K) & ternary (R-K-M) interactions
- REDLICH-KISTER\_KOHLER: for binary (R-K) & ternary (R-K-K) interactions

- FLORY-HUGGINS POLYMER MODEL: for interactions in a polymer mixture phase (F-H)
- MIXED-EXCESS-MODELS: (R-K default) for mixed binary excess model (R-K, Legendre & Polynom) of a substitutional solution phase
- HKF: for interaction in an AQUEOUS solution phase (HKF)
- PITZER: for interaction in an AQUEOUS solution phase (PIT)
- CENTRAL\_ATOM\_MODEL: for interaction in a liquid slag solution phase (C-A-M)

For extrapolations of excess energies from related binary systems to ternary or higher-order systems, the Redlich-Kister binary excess interaction parameters may be extrapolated with either a Muggianu extension (i.e. the default REDLICH-KISTER\_MUGGIANU model) and a Kohler extension (i.e. the alternative REDLICH-KISTER\_KOHLER model), where there is no ternary, quaternary or higher-order interaction parameter.



The REDLICHKISTER\_KOHLER model is implemented only for ternary systems.

The MIXED-EXCESS-MODELS option works only for a substitutional phase with no sublattice (such as the metallic LIQUID phase) and it can be used to invoke asymmetrical simple or Legendre polynomial as binary excess energy models in addition to the default symmetrical Redlich-Kister model for the chosen pair of constituents in a substitutional phase with no sublattice. Asymmetrical here means that the power series depend only on one of the constituents, for example with expansions based on the [1-2\*X(B)] term rather than [X(A)X (B)].



Binary interaction parameters for such a pair of constituents must be entered prior to turning on the non-default Legendre or Polynom models.

### TERNARY\_EXTRAPOLAT

See <u>example tcex52</u> in the *Thermo-Calc Console Mode Examples Guide*.

Use this to change the extrapolation method from the default REDLICH-KISTER\_MUGGIANU to another extrapolation model. This method extrapolates from binary to ternary (and higher-order) excess interaction parameters in a solution phase and is chosen from:

- MUGGIANU for Muggianu Extrapolation from R-K parameters
- TOOP-KOHLER for Toop-Kohler Extrapolation Model
- KOHLER-ALL for Kohler Extrapolation Model

- MUGGIANU\_RESTOR for Muggianu-Restor Extrapolation Model
- Only when all the relevant binary excess energies in the current solution phase are treated by the default Redlich-Kister Model (i.e. the Mixed-Excess-Model should have not been used), the MUGGIANU\_RESTOR method for ternary extrapolations is equivalent to the Redlich-Kister\_Muggianu Model, or the KOHLER-ALL method to the RedlichKister\_Kohler Model.

The default ternary extrapolation method MUGGIANU\_RESTOR applies to a solution phase as a whole, whatever the binary excess model(s) enforced to each of individual binary pairs in the phase. In case that all the binary pairs in the phase use the default binary excess model REDLICH-KISTER, then the ternary extrapolation method becomes the so-called Redlich-Kister\_Muggianu Model (in short as R-K-M), for extrapolations from binary parameters to ternary (and higher-order) excess energy terms; when no ternary L parameter is entered for that, such a default R-K-M Model is always used.

The KOHLER-ALL ternary extrapolation method can be turned on also for a solution phase as a whole, whatever the binary excess model(s) enforced to each of individual binary pairs in the phase. In case that all the binary pairs in the phase use the default binary excess model.

REDLICH-KISTER, then the ternary extrapolation method becomes the so-called Redlich-Kister\_Kohler Model (in short as R-K-K), for extrapolations from binary parameters to ternary (and higher-order) excess energy terms; when no ternary L parameter is entered for that, such an R-K-K Model is used.

The TOOP-KOHLER ternary extrapolation method is applied in a special way: for a specific ternary system (e.g. A-B-C) in a solution phase, specify which two constituents as the so-called Kohler constituents and the remaining constituent as the so-called Toop constituent on a given sublattice. If for the A-B-C ternary system you have specified A and B, or B and A, as the Kohler constituents (entered as the basis constituent and first interacting constituent) and C as the Toop constituent, only the A-B binary interaction parameters are used in accordance with the Kohler ternary extrapolation formula for A-B-C ternary interaction, while any other binary interaction parameters involving the Toop species C (i.e. of A-C and B-C binaries) are used in line with the Toop-Kohler ternary extrapolation formula (for the A-C-B and B-C-A ternary interactions).

### DEBYE\_HUCKEL

To use the DHLL (Debye-Hückel Limiting Law) model and SIT (Specific Ionic Interaction Theory) model for a dilute AQUEOUS solution phase, switch the DEBYE\_HUCKEL part on, and it removes previously set (if any) ADDITIONAL part on the non-ideality for the chosen AQUEOUS phase.

### **HKF\_ELECTROSTATIC**

To use the hypothetical electrostatic contribution for the chosen phase (it must be an aqueous solution phase using the Complete Revised\_HKF Model). It removes previously set (if any) ADDITIONAL part on the non-ideality for the chosen AQUEOUS solution phase.

### QUASICHEM\_IONIC

To use the Quasichemical-Ionic model for the chosen liquid phase, you must have already entered/retrieved a liquid phase as an ionic two-sublattice liquid solution phase (normally, that is the IONIC\_LIQ phase), and then use this amending option to switch QUASICHEM\_IONIC on. It creates a completely new liquid solution phase [namely the QUAS\_IONIC phase which uses the Quasichemical Model for the entropy, according to Mats Hillert (2001: *J of Alloys and Compounds*, 320, 161-176)], while the original IONIC\_LIQ phase remains in the system and is not changed by this option.

### QUASICHEM\_FACT00

To use the Quasichemical Model developed by Kongoli et al., you must have entered/retrieved a liquid phase using a normal substitutional liquid model with specified associates or species (on a single lattice site), and then turn on the option QUASICHEM\_ FACT00. This removes the previously-set (if any) ADDITIONAL part on the non-ideality for the chosen liquid phase.

See examples <u>tcex49</u> and <u>tcex50</u> in the *Thermo-Calc Console Mode Examples Guide*.

### QUASICHEM\_IRSID

To use the Kapoor-Frohberg-Gaye Quasichemical Cell Model (i.e. the Quasichemical Model developed by ISRID, France) for a liquid SLAG solution phase, you can use a normal liquid model with specified associates or species.

### **GLASS\_TRANSITION**

To use the special model for glass transition of a liquid phase. It removes previously set (if any) ADDITIONAL part on the non-ideality for the chosen liquid phase.

### **REMOVE\_ADDITION**

To remove all the selected ADDITIONAL part from the Gibbs energy description for the chosen phase. If preferred, set a specific phase-status bit for a phase, use the PHASE\_BITS option as long you know the restrict meaning of each part of a phase- status bits.

### DEFAULT\_STABLE

Set phases as default-stable, which helps you to have a better guess of which phases that should be stable at the first calculation.

# AMEND\_PHASE\_DESCRIPTION

Specify/amend phase descriptions if a phase has a miscibility gap, uses a special excess energy model, or has a specific additional contribution to its Gibbs energy, for example.



Also see <u>"Solution Models in the GIBBS Module" on page 82</u> for detailed information about this command and its options.

Syntax	AMEND_PHASE_DESCRIPTION		
	Phase name: <phase name=""></phase>		
	Specify the name of the phase.		
	Amend what /Composition_Set/: <subject></subject>		
Prompts	Several subjects for the phase can be amended but most often this command is used to enter two or more composition sets for a phase. If a phase has a miscibility gap it is necessary to have two composition sets, one for each possible composition that can be stable simultaneously.		
	Enter a question mark at the prompt to get a list of all possibly amended subjects for a phase.		
GLASS	There are no additional prompts for DEBYE_HUCKEL, HKF_ELECTROSTATIC, GLASS_TRANSITION, QUASICHEM_FACT00, QUASICHEM_ISRID, REMOVE_ ADDITION, and DEFAULT_STABLE.		
	New phase name /ABCD/: <phase name=""></phase>		
RENAME_PHASE	Give a new phase name for the chosen phase, or press <enter> to keep the default shown.</enter>		
	Sites in first sublattice /XX/ : <yy> Sites in second sublattice /XX/ : <yy></yy></yy>		
Site_ratios	Specify the site numbers for each of the prompted sublattices for the current phase. Press <enter> to accept the previous definitions.</enter>		
	Sublattice /#/: <sublattice number=""></sublattice>		
New	Specify the sublattice where the new constituents are located.		
constituent	Species: <species name=""></species>		
	Give a valid species name.		

Syntax	AMEND_PHASE_DESCRIPTION
	New highest set number /#/: <set n="" number=""></set>
	The default value (#) is usually one higher than the current value. All phases have initially one composition set. If a lower value (i.e. lower than the default one) is given, that specific composition sets are deleted.
	You cannot take away the first composition set.
Composition_	<pre>Major constituent(s) for sublattice #: /AB/: <major constituent(s)&gt;</major </pre>
sets	Specify the new major constituent(s) for the sublattice #, or press <enter> to accept the default which was automatically set according to the specified composition set of the phase.</enter>
	This prompt is repeated for all available sublattices in the chosen phase. The major constituents in each sublattice can be given. This may simplify giving start values when calculating the equilibrium as phases with miscibility gaps should have different major constituents for each composition set.
	Composition set /1/: <composition number="" set=""></composition>
	Give the composition set (digit number) for the chosen phase, or press <enter> if you want to set major constituents for the specified composition set.</enter>
	<pre>Major constituent(s) for sublattice #: /AB/: <major constituent(s)&gt;</major </pre>
Major_ constituent	Specify the new major constituent(s) for the sublattice #, or press <enter> to accept the default which is automatically set according to the specified composition set of the phase.</enter>
	This prompt is repeated for all available sublattices in the chosen phase for the specified composition set.
	The major constituents in each sublattice can be specified. This is useful in order to make calculations converge faster and more easily (because it may simplify giving start values when calculating the equilibrium as those phases with miscibility gaps should have different major constituents for each composition set). The databases often set major constituents for several phases automatically when data are retrieved.
Fraction_ limits	Element : <el1> Low fraction limit /0/ : <appropriate limit="" low=""></appropriate></el1>

Syntax	AMEND_PHASE_DESCRIPTION
	<pre>High fraction limit /1/ : <appropriate high="" limit=""> Element : <el2></el2></appropriate></pre>
	Low fraction limit /0/ : <appropriate limit="" low=""></appropriate>
	High fraction limit /1/ : <appropriate high="" limit=""></appropriate>
	Element : <eln></eln>
	Low fraction limit /0/ : <appropriate limit="" low=""></appropriate>
	High fraction limit /1/ : <appropriate high="" limit=""></appropriate>
	The antiferromagnetic factor /XX/: <yy></yy>
	Specify the anti-ferromagnetic factor for the chosen phase. This should be $-1$ for BCC phase and $-3$ for all other phases (FCC, HCP, etc.).
	Short range order fraction of the enthalpy /XX/: <yy></yy>
Magnetic_ ordering	The magnetic ordering is a second-order transformation and part of the enthalpy due to this transformation is due to short-range order. This value is the fraction of the total enthalpy that is due to short-range ordering above the magnetic transition temperature.
	The default value (xx) is 0.40 for BCC phase, and 0.28 for all other phases (FCC, HCP, etc.).
Disordered	Disordered phase name: <phase name=""></phase>
part –	Give the disordered phase name for which there is no ordering sublattice.
	Model name /ABCDEFG/: <model name=""></model>
	The default model is the pre-set model for the solution phase, normally the REDLICH-KISTER_MUGGIANU model, or choose another model for the phase to be amended:
	REDLICH-KISTER_MUGGIANU
	REDLICH-KISTER_KOHLER
Excess_model	FLORY-HUGGINS POLYMER MODEL
	<ul> <li>MIXED-EXCESS-MODELS (R-K default)</li> </ul>
	• HKF
	• PITZER
	CENTRAL_ATOM_MODEL
	The MIXED-EXCESS-MODELS option has sub-options:

Syntax	AMEND_PHASE_DESCRIPTION
	First (the independent) constituent: <constituent name=""></constituent>
	Second (the dependent) constituent: <constituent name=""></constituent>
	Specify the binary pair of constituents in the current substitutional solution phase (the first one as the so-called independent constituent, and the second one as the dependent constituent), for which you wish to change the binary excess model from the default REDLICH-KISTER model to another model (LEGENDRE or POLYNOM).
	You are prompted with the first sub-option (i.e. First (the independent) constituent after you have specified the desired binary Excess model type. To make further changes of binary excess model for other specific binary pairs in the current substitutional solution phase, press <enter> (implying that there are no more changes of binary excess model for all other possibly-remaining binary pairs that shall still use the default REDLICH-KISTER model).</enter>
	<pre>Excess model type: /Legendre/: <desired binary="" excess="" model="" type=""></desired></pre>
	For the currently-specified binary pair, choose one the legal binary excess models: LEGENDRE, POLYNOM or REDLICH-KISTER. If the default binary excess model REDLICH-KISTER is used for a certain binary pair, you can either specify the model name (or in short as R-K) or press <enter>. After this sub-option, it returns to the first sub-option <i>First (the independent)</i> <i>constituent:</i> either for further change(s) of binary excess model of any other binary pair(s) or for termination (by pressing <enter>) in the MIXED-EXCESS-MODELS option.</enter></enter>
	Extrapolation method: /Toop-Kohler/: <ternary model=""></ternary>
	Choose a ternary extrapolation model for the current solution phase:
	TOOP-KOHLER for Toop-Kohler Extrapolation Model
	KOHLER-ALL for Kohler Extrapolation Model
Ternary_ extrapolat	<ul> <li>MUGGIANU_RESTOR for Muggianu-Restor Extrapolation Model</li> </ul>
	The TOOP-KOHLER option further prompts:
	Constituent in sublattice #: <basis constituent="" name=""></basis>
	First interaction constituent: <interacting constituent="" name=""></interacting>
	Toop constituent: <toop constituent="" name=""></toop>

Syntax	AMEND_PHASE_DESCRIPTION
	Specify the so-called Kohler constituents (entered as the basis constituent and first interacting constituent) and Toop constituent on a given sublattice in the current solution phase.
Quasichem_ ionic	To use the Quasichemical Model for the chosen liquid phase, you must first enter it as an ionic liquid phase, and then use this amending option. It creates a completely new phase, the original IONIC_LIQUID phase remains in the system and is not changed.
	Number of bonds/atom /2/: <number atom="" bonds="" of="" per=""></number>
	Specify the Z value (number of bonds per atom).
	Correct the phase status bits. It is important to fully understand this command before using it:
	New status word (in hexadecimal) /04204000/: <new bits="" status=""></new>
	"LIST_STATUS" on page 117
Status_bits	Normally, these should never be changed in this way, because such phase status bits are automatically set on when all the phase descriptions (predefined in databases and amended in the GIBBS module) are defined properly. The only exceptions are for the Kapoor-Frohberg-Gaye cell model for liquid SLAG solution phase, which can be set only as 04208000; and for the complete Revised HKF AQUEOUS solution model, which can be set as 02084000.

# AMEND\_SYMBOL

This command is for the GES (GIBBS) and PARROT modules.

Interactively calculate the current values of a function or table (predefined by the switched database or previously entered with <u>"ENTER\_SYMBOL" on page 109</u>) at the current temperature and pressure condition, and also possible to modify the definitions of variables or functions.



>\_

The current values of the temperature and pressure used to calculate the functions or tables are the ones listed with <u>"LIST\_STATUS" on page 117</u>. There is no way to change the current temperature and pressure values interactively.

To modify the definitions of entered symbols (variable, functions or parameters), the performance of this command is slightly different one from another, as described below:

- For variables, the values can be changed.
- For functions the low and high temperature limits in various ranges (if any), as well the expressions, can be changed.
- For parameters the low and high temperature limits in various ranges (if any), as well the expressions, can be changed.

Syntax	AMEND_SYMBOL	
	Name: <symbol name=""></symbol>	
Prompts	Specify the name of an entered symbol.	
	For tables and functions (or parameters which are treated as functions after these are entered by the ENTER_SYMBOL command with the Parameter keyword, but not by the ENTER_PARAMETER command), after the symbol name is specified here, the program automatically calculates the values under the current temperature and pressure conditions, and lists the current values, such as:	
	FUNCTION VALUE 2.52500000E+01	
	TABLE VALUE 1.56000000E+02	
	For variables and functions (or parameters), there are additional prompts and depend on the symbol type.	
	For a variable, its current value is displayed, and you can change it to a new value:	
	Value /Current value/: <new value=""></new>	
	For a function (or a parameter entered as a symbol):	
	Do you want to change the number of ranges /No/:	
	Enter Y to change the number of ranges for the function, or change some of the temperature limits in the definition, then retype both the low/high temperature limits and functions (see all the remaining details in the ENTER_SYMBOL command.	
	If you do not want to change it, press <enter>. The definition of the chosen function in all ranges (if any) is listed, for example:</enter>	
	DIFFERENT FUNCTIONS IN THESE RANGES	
	298.15 <t<2000.00< td=""></t<2000.00<>	
	2000.00 <t<4500.00< td=""></t<4500.00<>	
	4500.00 <t<6000.00< td=""></t<6000.00<>	

# Syntax AMEND\_SYMBOL and this message displays: Do you want to change range limits /No/: If a function is different in two or more temperature ranges, you must specify the range of the function of which you want to amend. Or press <Enter> or type 0 to exit this command without making any change. Range number (0 to exit) /0/: <Range number> Function: The previous function is available for editing. The editing is performed within the general subroutine FOOLED. This routine prompts as follows: 1:+:> The prompt consists of the current position in the string and the character at that position between colons (::). **COMMANDS** These commands can be given: • Help: ? • Move CP to last or first character: <+/-> A • Delete characters from CP: <+-#characters> D • Exit: E • Find: <#occurrences> F<string>@ • Insert: I<string>@ • Move: <+-#positions> M • Restore string: R • Substitute: s<old>@<NEW>@ Type string: T Where CP denotes the current position in a string, # means number of, @ is a terminator of an input or search string.

When the string is typed the character at the current position is replaced by an underscore \_.

Syntax AMEND\_SYMBOL

To finish the editing of the current function, type E at the prompt.

# **CHANGE\_STATUS**

For the GIBBS\_ENERGY\_SYSTEM module, the status of an element or species or phase can be either ENTERED or SUSPENDED. The suspended status can be either implicit or explicit. The implicitly suspended status can be set e.g. for a species if any of the elements in its chemical formula is explicitly suspended. A species that is implicitly suspended becomes entered automatically if all its elements are set entered. After this command, a message shows which elements/species/phases are suspended or restored (entered). Consequently, the status bits for the specified elements or species or phases are changed, as listed with "LIST\_STATUS" on page 117.

Syntax	CHANGE_STATUS
Prompts	For Element, Species or Phase /Species/: <keyword></keyword>
	Specify the keyword (element, species or phase).
	Suspend /Y/:
	By default the status is changed from ENTERED to SUSPENDED.
	If an element is suspended all species with this element become implicitly SUSPENDED too. A phase may become implicitly suspended if all its constituents or all constituents in a sublattice are suspended.
	List of elements/species/phases: <name elements="" of="" or="" phases="" species=""></name>
	Specify the names or indices of those elements or species or phases that shall become suspended or active. For names, these should be separated by a space and terminated with a semicolon (;) or an empty line. For indices it is possible to give a range by separating two numbers by a hyphen (-). The list should be terminated by a semicolon (;). Example: $5 \ 1 \ 7-12 \ FE$ ;

# **ENTER\_ELEMENT**

Specify a system interactively. The program searches the currently switched or preset database for data for the given elements. The data for the elements in the database are the:

- mass in g/mol
- name of the Selected Element Reference State (SER) which normally is the stable phase for the element at 298.15 K

- enthalpy difference for the element in the SER state at 298.15 K and zero K
- absolute entropy for the element in the SER state at 298.15 K.

The two predefined elements, i.e. electrons and vacancies, have the chemical symbols /and va, respectively. Initially, these are suspended but can be entered either by this command or <u>"CHANGE\_STATUS" on the previous page</u>.

If an aqueous solution phase is involved, in the GIBBS module you enter a special aqueous electron called, ZE. This is specially designed for appropriately calculating the standard electric potential in the aqueous solution system.

Syntax	ENTER_ELEMENT	
	Element name: <element name=""></element>	
Prompt	Specify several elements on one line. The name of an element is its chemical symbol. The chemical symbols must be separated by spaces. Fictitious element names are legal but naturally no data are found in the database for them.	
	An element name (maximum 2 characters) can either have its first letter in upper and the second (if any) in lower case (i.e. Lower Case Mode) or both letters can be in upper or lower case (i.e. Upper Case Mode). The Upper or Lower Case Mode is selected by the command <u>"REINITIATE" on page 121</u> , which removes all data, and should be executed before any other command.	

# About the ENTER\_PARAMETER Command



For the PARROT Module, this command is the same and is described here. See <u>"ENTER\_PARAMETER" on page 137</u> for the command details.

In the descriptions of the standard thermochemical properties and special physical properties for a phase, there are a number of parameters which may depend on the temperature and pressure. The expressions for these parameters can be given in a free form as a sum of terms with powers of T and P and may also include the natural logarithm and exponential function. This type of expression is called *TP-functions*. Identical parameters (in terms of parameter-names) are stored only once in the GIBBS workspaces.

The composition-dependence of the Gibbs energy is described in the GIBBS module by the internal data structure, which is created when the phase is entered (see <u>"Gibbs Energy</u> <u>System (GES) Commands" on page 76</u>). The Gibbs energy of a phase is always referred to one formula unit of the phase, i.e. the amount derived from the number of sites (i.e. the stoichiometric coefficient) for each sublattice. If vacancy is a constituent of a sublattice, the amount of matter per formula unit of the phase may vary with composition.

### **Defining the Parameter**

A valid parameter should have the general form of:

```
<identifier>(<phase>, <constituent array>; <digit>) <xxx> <expression> <yyy>
<keyword Y or N> <zzz> !
```

The identifier must be followed by an opening parenthesis, a phase name, a comma and a constituent array. Optionally, the constituent array can be followed by a semicolon and a digit. The parameter name is terminated by a closing parenthesis. The parameter form is defined as:

- <identifier> is the parameter type;
- <phase> is the phase name (maximum 24 characters);
- <constituent array> is the specific constituent array in the phase;
- <digit> is the degree of composition-dependent interaction contribution (an integer number from 0 through 9), that is only for excess energy (L), Curie temperature (TC) and Bohr magneton number (BMAGN), as well as for volume-related parameters (V0 or VA or VB or VC or VK); if it is valued as zero, or if it is for the standard Gibbs energy (G) for which the degree is always zero, it can be omitted;
- <expression> is the mathematical relation to describe the parameter;
- <xxx> and <yyy> are the low and high temperature limits respectively for the applicable temperature range of the parameter expression;
- <keyword Y or N> is the indicator on if there is continuation for the parameter
  expression or not;
- <zzz> is the reference index/number for the assessment of this parameter;
- The exclamation point ! is used to indicate that the current parameter definition is ended.

### PARAMETER NAME

The GES parameter name has a general form of:

<identifier>(<phase>,<constituent array>;<digit>)

Examples of parameter names:

- G (GAS, C102): The Gibbs energy of formation of a CO2 molecule in gas.
- G(FCC, FE:VA): The Gibbs energy of formation of fcc Fe with interstitials.
- L(LIQ, Fe, Cr; 0: The regular solution parameter for Fe and Cr in liquid.

- L(LIQ, Fe, Cr; 1): The sub-regular solution parameter.
- TC (BCC, Fe:Va): The Curie temperature of bcc Fe.
- BMAGN (BCC, Fe:Va): The Bohr magneton number parameter of bcc Fe.

The parameter name consists of several parts. The first is a type-identifier and these can be used:

- G: Standard energy parameter (Gibbs energy of formation) or for interaction parameters;
- L: Excess energy parameter (Gibbs energy of interaction) always used for interaction parameters;
- TC: Curie temperature for magnetic ordering;
- BMAGN or BM: Bohr magneton number for magnetic ordering (or Born function  $\omega$ Pr,Tr for aqueous solute species).
- vo: Molar volume at 298.15 K and 1 bar (a numeric value only);
- VA: Integrated thermal expansivity;  $\int_{298.15}^T lpha(T) \mathrm{dT}$
- VB: Bulk modulus at 1 bar;
- vc: Isothermal compressibility;
- VK: High-pressure fitting parameter.

When necessary quantities as H (enthalpy), S (entropy), V (Volume), F (Helmholtz energy), etc., can be calculated from the Gibbs energy.

### PHASE NAME

Specifying the PHASE NAME in uppercase is recommended; however, if you prefer to write it as a mixture of uppercase and lowercase, it automatically converts all lowercase to uppercase, as the GIBBS module only recognises uppercase phase names. It is important that if a phase bears a legal phase-type (among G, A, Y, L, I, F and B) in its phase definition (already by the PHASE keyword; such as GAS:G, LIQUID:L,SLAG:L, IONIC-LIQ:Y, SPINEL:I, FCC\_L12:F, HCP\_D021:F, BCC\_B2:B, AQUEOUS:A), such a valid phase-type code should not be attached to the phase name when ENTER\_PARAMETER is executed.

### **CONSTITUENT ARRAY**

The constituent array consists of a list of constituent names. Interaction parameters have

two or more constituents from the same sublattice separated by a comma. If the phase has sublattices, at least one constituent in each sublattice must be specified. The constituents in different sublattices must be given in sublattice order and are separated by a colon.

After the component array, a sub-index digit can be specified after a semicolon. This digit must be in the range 0 to 9. The interpretation of the sub-index depends on the excess energy model used for the phase. If no semicolon and digit are given, the sub-index value is assumed to be as zero.

The excess energy parameters, e.g. the regular/subregular (binary) parameter or ternary parameters, are multiplied with two or more fractions of the constituents from the same sublattice of the solution phase. These additional constituents must be given as interacting constituents (as the following prompt).



Solution phases with sublattices can have interacting constituents in each sublattice.

### **INTERACTION PARAMETER**

An interaction parameter, which is used to describe the excess term of a quantity, must have two or more constituents that interact with each other on a specified sublattice site of the given phase. It is arbitrary which of these constituents is given as the first constituent and what is given as the interacting constituents. The software always sorts the constituents (in each sublattice) in alphabetical order when the parameter name is written as a prompt (for entering its parameter value) and when the parameter is listed (using the GIBBS commands "LIST\_PARAMETER" on page 141 or "LIST\_PHASE\_DATA (from GIBBS)" on page 142). This is important for all asymmetric interaction parameters where the sign of the interaction parameter must depend on the appearance order.

Use an asterisk\* to denote that the excess interaction parameter is independent of the constituents of a specific sublattice. For example,  $L(FCC_{L12}, AL, NI:*)$  means that the interaction parameter is for the binary interaction between constituents AL and NI on the first sublattice in the FCC\_L12 solution phase, while it is independent of all constituents on the second sublattice. An interaction parameter in the list of constituents is always added to the Gibbs energy and the asterisk \* is calculated with the term of  $[1\sum y(specified constituents)]$ , which implies that in an A-B binary system these L parameters are identical (but in higher-order systems, these are different):

- L(phase, A, B) is multiplied with X(A)\*X(B)
- L(phase, A, \*) is multiplied with X(A)\*(1-X(A))
- L(phase, B, \*) is multiplied with X(B)\*(1-X(B))

If you press <Enter> when you are asked for a parameter name or if you have improperly input the entire parameter name, you are asked for each of these items in the name.

# **ENTER\_PARAMETER**



This command is for both the GES and PARROT modules.

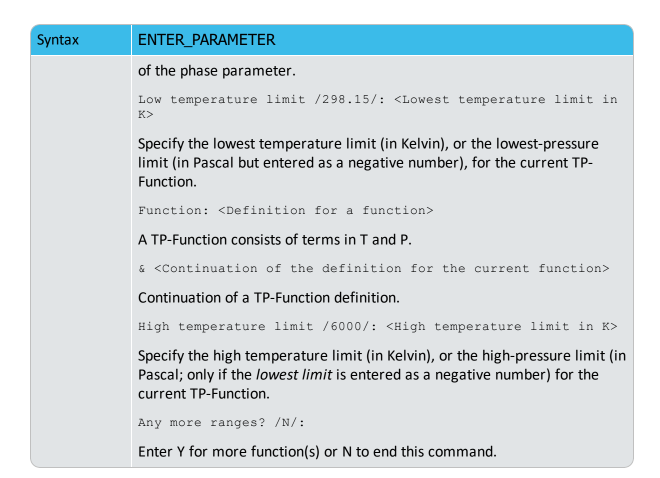
### "About the ENTER\_PARAMETER Command" on page 133

Use this command to enter TP-function(s) for a specific parameter for a phase interactively with this command. If there is already a parameter expression defined (for this parameter), that is deleted and replaced with newly entered one.

The entered TP-Function(s) for the parameter can be changed later with the command "AMEND\_PARAMETER" on page 79.

Syntax	ENTER_PARAMETER
Prompts	Parameter: <parameter name=""></parameter>
	As explained in <u>"About the ENTER_PARAMETER Command" on page 133</u> , specify a correct and complete Parameter Name, which should contain all the necessary parts of the general form:
	<identifier>(<phase>,<constituent array="">;<digit>)</digit></constituent></phase></identifier>
	If a parameter name is not acceptable or <enter>, is pressed, a message displays:</enter>
	*** Error, please re-enter each part separately
	and you are prompted for input for each required part for a parameter name.
Options	Description or Information
Identifier	Identifier (/X/): <g bm,="" l,="" or="" tc,="" v0="" va="" vb="" vc="" vk=""></g>
	If this command is used one or more times, the previous value on this prompt is set as default. Press <enter> for the same type identifier or specify a new type.</enter>
	Phase name (/ABCD/): <phase name=""></phase>
Phase name	Each parameter is valid for a specific phase only. The name of that phase must be supplied (maximum 24 characters). The name can be abbreviated.
Constituent	Constituent (in sublattice # /ABC/): <species name=""></species>

Syntax	ENTER_PARAMETER
	A parameter is identified by the constituents on a specified sublattice site of the given phase, the stoichiometric coefficients of which are multiplied with the parameter. The name of the constituent can be abbreviated.
	It is the species name, not the stoichiometric formula required here.
	If this command is used one or more times, the previous value on this prompt is set as default. Accept it by pressing <enter> if the constituent is the same, or specify a new species name.</enter>
	For phases with several sublattices, the program asks for one constituent in each sublattice.
Interacting constituent	<pre>Interacting constituent (in sublattice # /XYZ/): <species name=""></species></pre>
	If this command is used one or more times, the previous value on this prompt is set as default. Press <enter> to accept it if the constituent is the same, or specify a new species name.</enter>
	To cancel the default value of the interacting constituent type NONE or the name of another constituent.
	This question is repeated until all the interested interacting constituent(s) on a specific sublattice in the phase are specified, and finally an <enter> is enforced.</enter>
	Degree /#/: <degree></degree>
Degree	Degree is model-dependent. Specify an integer number (a value from 0 through 9) as the degree of composition-dependent interaction contribution for the phase parameter. This is valid for excess energy (L), Curie temperature (TC) and Bohr magneton number (BMAGN), as well as for volume-related parameters (V0, VA, VB, VC or VK).
	For binary interaction parameters, the degree is usually the power in the Redlich-Kister expression. For ternary interaction parameters, it is usually the Hillert ternary index.
	For a standard G parameter for a pure component (end-member) its degree should be always 0 and this prompt should not display.
Phase parameter	After this prompt, the program echoes on the screen the full TP-Function



# About the ENTER\_PHASE Command

With the command <u>"ENTER\_PHASE" on page 106</u>, the phase name, phase-type, sublattice number, and constituents or constituent array(s) for the phase are entered into the GIBBS workspaces.

### ABOUT PHASE NAMES

A thermochemical system must have at least one phase (which is a homogeneous part of the system with a uniform composition and structure). In the GIBBS module, any number of phases can be entered for a system and for each phase there must be a description of how its Gibbs energy varies with temperature, pressure and composition. A phase has a large amount of data connected to it, e.g. it starts with a phase name:

- It may be treated as a special phase-type;
- It may have structural information about sublattice(s), etc.,
- There must be a list of constituents (for a substitutional phase with no sublattice) or

of constituent arrays (for a sublattice phase);

- There may be basic information on what kind of EXCESS\_MODEL (polynomial expression) is used to describe the binary, ternary and/or higher-order interactions between constituents;
- There may be so-called Additional contributions to the Gibbs energy of the phase from special physical phenomena, e.g. magnetic ordering, hypothetical electrostatic interaction, and so forth;
- There must exist all the parameters required for the descriptions of thermochemical properties (i.e. G terms for standard Gibbs energies, and L terms for binary, ternary or higher-order interaction excess energies) and of some special physical properties (e.g. the Curie temperature TC and Bohr magneton number BMAGN (or BM) for magnetic ordering, V0-VA-VB-VC-VK parameters for volume contributions, Born functions for hypothetical electrostatic interaction in an aqueous solution phase) stored in connection with the phase.

### ABOUT PHASE TYPE CODE

A G phase (gaseous mixture) or an A phase (aqueous solution) is usually treated as a substitutional phase without sublattice, and that an L phase (ordinary liquid solution) is normally (but not always) modelled as a substitutional phase without sublattice, too.

The F and B phase-types are useful for ordered FCC (or HCP) and BCC solution phases handled by the so-called Four Substitutional-Sublattice Ordering Model, which always requires that the solution phase must have four sublattices for substitutional ordering and can additionally have an interstitial sublattice.

For ordered FCC or HCP phases, these four substitutional sublattices represent four corners of the regular tetrahedron on these lattices all of which are the nearest neighbours. A *Normal 4-Sublattice Model* requires that all the G parameters for each of end-members with the same elements but distributed on different sites be given separately. However, as these corners are identical lattice points, the phase-type option F means that the G parameters need be given only once. The possible permutations are handled automatically. To be more clarified: An A-B binary solution phase (with the element A locates on one sublattice site and B on three sublattice sites) treated by the Normal 4-Sublattice Model has to have 4 G parameters for 4 end-members, i.e. G(phase,A:B:B:B), G(phase,B:A:B:B), G(phase,B:B:A:B) and G(phase,B:B:A), because of that in the general case these G parameters can be different from each other. But for the FCC and HCP orderings, these are identical and thus all G parameters of such end-members need be given only once, and the possible

permutations are then automatically handled by the GIBBS module. This significantly simplifies the usage of this model (*Four Substitutional-Sublattice Ordering Model*) in multicomponent alloys.

There are restrictions on the excess parameters allowed with the phase-type option F. You can only have excess parameters of these types:

L(phase, A, B:\*:\*:\*;0...9) L(phase, A, B:C, D:\*:\*;0...9)

The asterisk \* means that the interaction parameter is independent on the constituents on its corresponding sublattice. No ternary interaction parameters (i.e. with three elements on one sublattice site) are allowed. The reason for this restriction is that it would be too complicated to handle all possible permutations. In the current *Four Substitutional Sublattice Ordering Model*, the binary interaction between A and B atoms is thus independent of the constituents on the other sublattices, where there are many other parameters to model the composition-dependence of the Gibbs energy (both in the ordered and disordered parts of the phase). The model for these ordered phases are always partitioned in a disordered part (with a single substitutional sublattice) and an ordered part (with four substitutional sublattices for ordering).

For ordered BCC phases, the phase-type option B means the same thing but the situation is more complicated, as the 4-sublattice ordering phase represents an irregular tetrahedron with two pairs of sites that are next nearest neighbours. Thus, for an A-B binary solution phase (with the element A locates on two sublattice site and B on other two sublattice sites) treated by the Normal 4-Sublattice Model, the end-member described by the G (phase,A:A:B:B) term has four nearest neighbour bonds between A and B atoms, whereas the end-member described by the G(phase,A:B:A:B) term has two nearest neighbour bonds between A and B atoms and two next nearest neighbour bonds



For detailed reference visit the Crystal Lattice Structure web page <u>http://www.nrl.navy.mil/mstd/branches/6390</u>.

The first end-member (described by the G(phase,A:A:B:B) term) represents B2- ordering and the second (described by the G(phase,A:B:A:B) term) stands for B32-ordering. There are two permutations of the G(phase,A:A:B:B) term and four permutations of the G(phase,A:B:A:B) term, automatically conducted in the Four Substitutional-Sublattice Ordering Model. And there are also two kinds of reciprocal interaction parameters, i.e.

```
L(phase,A,B:C,D:*:*;0...9) L(phase,A,B:*:C,D:*;0...9)
```

An advanced feature with the phase-type options F and B is that a composition set that represents the solution phase has a suffix (indicating that it is really as an ordered or disordered phase) that is automatically added to its phase name in some listings of

equilibrium calculations (when performing either single-point or stepping or mapping calculations, and when plotting the calculated property diagrams or phase diagrams). Such suffix indications can be:

- L12 or L10 for ordered FCC
- \_A1 for disordered FCC
- \_B2, \_B32, \_D03 or \_L21 for ordered BCC
- \_A2 for disordered BCC
- \_D019 and \_B19 for ordered HCP
- \_A3 for disordered HCP

Solution Phase	Suffix for Disordered Phase	Suffix for Ordered Phase
FCC PHASE	PHASE_A1	PHASE_L12
FCC PHASE		PHASE_L10
	PHASE_A2 PHASE_A3	PHASE_B2
		PHASE_B32
BCC PHASE		PHASE_D03
		PHASE_L21
HCP PHASE		PHASE_D019
nur Phase		PHASE_B19

# **ENTER\_PHASE**

See <u>"About the ENTER\_PHASE Command" on page 103</u> for details about the phase-type and other prompts.

With this command, the phase name, phase-type, sublattice number, and constituents or constituent array(s) for the phase are entered into the GIBBS workspaces. However, for the remaining parts of thermodynamic descriptions (i.e. thermochemical and physical parameters, excess model, and possibly additional contribution term) for the phase, the GIBBS commands <u>"ENTER\_PARAMETER" on page 137</u> and <u>"AMEND\_PHASE\_DESCRIPTION"</u> on page 89, should be used sequentially.

Normally, data for a system are retrieved from the preset database after the elements and species are specified. This is automatically done through <u>"GET\_DATA" on page 16</u>. In some cases, specify a phase interactively.

Syntax	ENTER_PHASE
Options	Description and Information
	Name of phase: <phase name=""></phase>
Phase name	The phase name is any string starting with a letter and containing only letters, digits or the underscore character. The phase name must be unique.
	Type code: <phase-type code=""></phase-type>
	The Phase-type code must be specified for the phase if it is not an ordinary solution phase. Press <enter> if the phase is ordinary.</enter>
Phase-type	
	Gaseous mixture phase.
G	There may only exist one gas phase in a system.
А	Aqueous solution phase.
L	Liquid solution phase but not ${\tt A}$ (aqueous) or ${\tt Y}$ (ionic liquid)
Ү	Ionic liquid solution phase (that is specially treated by the Ionic Two Sublattice Liquid Model).
I	A phase with charged species but not ${\tt G}$ (gaseous), ${\tt A}$ (aqueous) or ${\tt Y}$ (ionic liquid).
F	An ordered FCC or HCP solution phase with four substitutional sublattices (additionally, such a phase can also have an interstitial sublattice).
В	An ordered BCC solution phase with four substitutional sublattices (additionally, such a phase can also have an interstitial sublattice).
	Number of sublattices /1/: <sublattice number=""></sublattice>
Sublattice number	Phases with stoichiometric constraints usually have two or more sublattices. On each sublattice one or more species can be entered as constituents. The number of sublattices must not exceed 10. Substitutional phases with no sublattices are treated as phases with one sublattice.
	Number of sites on sublattice # /1/: <site number=""></site>
	For phases with sublattices, the ratio of the sites on each sublattice (donated with #) must be given. The program asks for values for all

Syntax	ENTER_PHASE
	sublattices (by repeating this question for each sublattice), but these may have a common factor extracted. It is recommended to use integer numbers as sites if possible. For substitutional phases with no sublattices, this question does not show up.
	Name of constituent: <constituent name(s)=""></constituent>
	For each sublattice specify at least one species as constituent. Several constituents can be given consequently (on the repeated prompt) or on the same line separated by a space. This question is repeated till pressing <enter> or giving a semi-colon (;), meaning that the list of constituents is terminated by (;) or an empty line.</enter>
	Will you add constituents later /N/:
Constituents	For Y add other constituents to the phase by specifying new components in e.g. an ENTER_PARAMETER command.
	If illegal constituents are used, you get a message e.g. when parameters are entered.
	Do you want a list of possible parameters $/N/$ :
	Enter Y for a list of all possible parameters up to the fifth interaction order for the entered phase.

# **ENTER\_SPECIES**

8\$B

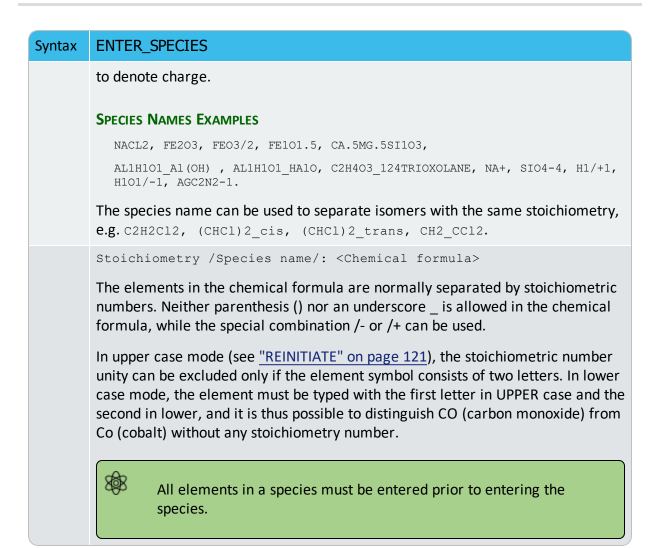
Specify a species from the already entered elements. The stoichiometry of the species is the chemical formula of the species. For each species, its name and chemical formula must be given.

All elements are automatically entered simultaneously as species. The constituents of a phase must be species, but there is no thermochemical data associated with the species except as constituents of a phase.

### Syntax ENTER\_SPECIES

Species name: <Species name>

Prompt Enter a unique species name. This is usually its chemical formula but it can be any string which starts with a letter and contains only letters or digits. Parenthesis () and an underscore \_ are permitted. The combination /- or /+ used



# **ENTER\_SYMBOL**

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This command is for the GES (GIBBS) module.

In the GIBBS module, a symbol can be used to represent a numeric quantity (e.g. a *variable*), a *function*, or a *table*. The entered symbol (with a symbol name of maximum eight characters) can be used later when parameters are entered, and this is a flexible way to manipulate thermodynamic functions. Symbols are especially useful if the same function or table is used in several thermochemical parameters. A *parameter* symbol is used to assign the values of phase parameters (which are defined with the command <u>"ENTER</u> <u>PARAMETER" on page 137</u>) to specific characterized symbols. If you want to define a constant quantity, enter it as a simple function with a constant value in all (temperature) ranges.

The symbols entered in the GIBBS module can also be listed and used in the <u>"Optimizing with</u> <u>the PARROT Module" on page 124</u> (for data optimization) where you can use the GIBBSentered symbols (variables, functions or parameters) to define parameters that shall be optimized.



Symbols entered in the GIBBS module are not the same symbols as defined in the POLY and POST modules.



Also see <u>"ENTER\_SYMBOL" on page 238</u> for the POLY and POST modules.



See <u>example 44</u> in the *Thermo-Calc Console Mode Examples Guide* for an example of using variables and functions.

Syntax	ENTER_SYMBOL	
	Variable, function, table or parameter? /Function/: <keyword></keyword>	
	The keyword can be a variable, function, table or parameter.	
	<ul> <li>Variables are similar to functions because these can also be expressions of state variables. It is possible to enter a variable with a new expression anytime.</li> </ul>	
	• Functions are expressions of state variables or other functions.	
Prompt	• Tables are used for listing results. A table consists of a list of any number of state variables, functions or variables.	
	<ul> <li>Parameter symbols are used to assign the values of phase parameters (which are defined by the command ENTER_PARAMETER) to specific characterized symbols. If the phase parameter is not defined yet, the parameter symbol is assigned zero or a symbol which is valued as zero. In this way it is easy to refer to entered phase parameters in further defining other phase parameters.</li> </ul>	
	Name: <name of="" symbol="" the=""></name>	
	Each symbol has a unique name that must start with a letter and can have maximum 8 characters.	
	The following prompts are based on the type of symbol.	
Variable	Value: <numeric a="" of="" value="" variable=""></numeric>	
symbol	Only a constant numeric value, not an expression, is accepted.	
Function <b>symbol</b>	Enter the lower-temperature limit, the expression, an upper-temperature limit and if there is another expression above this upper limit:	

Syntax	ENTER_SYMBOL
	Low temperature limit /298.15/: <lowest in="" k="" limit="" temperature=""></lowest>
	Specify the lowest-temperature limit, below which its (first) expression is not applicable. The default lowest limit of all kinds of data is 298.15 K.
	If a negative number is given as the lowest-temperature limit, it is assumed that there are breakpoints in pressure for this function; in such a case, it is interpreted as the lowest-pressure limit (in Pascal), and the other limits in the current function are also taken as pressure limit values (in Pascal).
	The temperature/pressure limits for the functions are checked during calculations. An indicator is set if the actual temperature/pressure condition is below the lowest temperature/pressure limit or above the highest temperature/pressure limit; and in such a case, an extrapolation is conducted using the TP-FUNCTIONS valid in the nearest temperature/pressure range.
	Function: <definition a="" for="" tp-function=""></definition>
	A TP-Function consists of terms in T and P. The expression is a FORTRAN-like expression and operators +, -, *, = and ** can be used (** only with integer powers). Unary-functions LN or LOG (both for natural logarithm) and EXP (for exponential) can also be used. An expression can be continued on more than one line. A PT-function must be terminated by a semicolon (;) or an empty line. No more than 78 characters can be written on a line. If this is not sufficient space on one line, press <enter> and continue on the next line. The program prompts with an ampersand <math>\&amp;</math> when a continuation is expected.</enter>
	A function can have several temperature ranges, and a different expression (i.e. a TP-Function) for each region; if required, a single function can have up to ten different temperature ranges (and thus have up to ten different TP-Functions).
	This question is repeated if the current function has more than one expression (TPFunction) in various temperature ranges [or pressure ranges; only if the <i>lowest limit</i> is entered as a negative number)].
	& <continuation current="" definition="" for="" of="" the="" tp-function=""></continuation>
	This is for a continuation of the definition for the current TP-Function. The ampersand & means that you can continue to write the function on the new line if one line is not enough for the current TP-Function. If you are finished the current TP-Function, press <enter> again.</enter>
	High temperature limit /6000/: <high in="" k="" limit="" temperature=""></high>

Syntax	ENTER_SYMBOL
	For a TP-Function, specify the high-temperature limit, above which its current expression is not applicable. The default high-temperature limit for all kinds of data is always 6000 K.
	This question is repeated if the function has more than one expression (TPFunction) in various temperature ranges.
	All the temperature limits for the presently-be-entered TP- Function MUST be in an increasing order; otherwise the whole function is given a single value of zero.
	If a negative number is given as the lowest-temperature limit, it is interpreted as the low-pressure limit (in Pascal), and the high-temperature limit is also taken as the high-pressure limit (in Pascal) for the current TP- Function.
	The high-temperature/pressure limit for the current applicable expression (TPFunction) is the low temperature/pressure limit for the next range if several ranges are needed to describe the temperature/pressure-dependence of this function.
	The temperature/pressure limits for the functions are checked during calculations. An indicator is set if the actual temperature/pressure condition is below the lowest temperature/pressure limit or above the highest temperature/pressure limit; and in such a case, an extrapolation is conducted using the TP-Function valid in the nearest temperature/pressure range.
	Any more ranges /N/: <y n="" or=""></y>
	Answer $\mathbb{N}$ to finish the definition of the current function. If $\mathbb{Y}$ the program asks for a new function valid above the last high temperature limit and below a new high temperature limit. The maximum number of temperature ranges is 10.
	Important note: Make sure that the function and its first derivatives are continuous across break points.
	Specify the low and high temperature limits, temperature step and tabled value at each specified temperature:
Table symbol	Low temperature limit /298.15/: <low in="" k="" limit="" temperature=""></low>
Symbol	The lowest temperature where the table has a value.
	High temperature limit /6000/: <high in="" k="" limit="" temperature=""></high>

```
ENTER SYMBOL
Syntax
            The highest temperature where the table has a value.
            Step in temperature /100/: <Temperature step>
            The step in temperature between each value in the table. This step must be
            constant for the whole table. If a table cannot be described with a single step
            it must be divided into several tables for different temperature ranges. The
            upper and lower limits are truncated to give reasonable values where the
            table values must be specified.
            Table value at XX /YY/: <Table value>
            The value of the table at the specified temperature (xx) must be given. The
            default value (yy) is the last value.
            Input the correct name of a phase parameter (including its identifier, phase
            name, constituent and interacting constituent(s) in various sublattices (if
            exist), and degree of the parameter. Most of the settings are the same as in
            the "ENTER PARAMETER" on page 137 command or the Function symbol
            prompts above.
            Parameter: <Parameter name>
            Specify a correct and complete parameter name, which should contain all
            the necessary parts of the general form:
            <Identifier>(<Phase>, <Constituent array>; <Digit>)
            If a parameter name is not acceptable or <Enter> is pressed, this error
            message displays:
Parameter
            *** Error, please re-enter each part separately
symbol
            The program prompts for separate input for each required part for a
            parameter name.
            Identifier (/X/): <G or L, or TC, or BM, or V0 or VA or VB or
            VC or VK>
            Specify one of these types of legal identifiers.
            Phase name (/abcd/): <Phase name>
            Specify the phase name.
            Constituent (in sublattice # /abc/): <Species name>
            Specify the constituent name on the specified sublattice site of the given
            phase.
```

Syntax	ENTER_SYMBOL
	<pre>Interacting constituent (in sublattice # /XYZ/): <species name=""></species></pre>
	Specify the interacting constituent name on the specified sublattice site of the given phase; if there is no interacting constituent, press <enter>.</enter>
	<pre>Interacting constituent (in sublattice # /XYZ/): <species name=""></species></pre>
	If there is more than one interacting constituent on the specified sublattice site of the given phase, specify them; otherwise press <enter>.</enter>
	Degree /#/: <degree></degree>
	Specify an integer number (a value from 0 through 9) as the degree for the phase parameter.
	Low temperature limit /298.15/: <lowest in="" k="" limit="" temperature=""></lowest>
	Specify the lowest-temperature limit (in Kelvin), or the lowest-pressure limit (in Pascal but entered as a negative number), for the current TP-Function.
	Function: <definition a="" for="" function=""></definition>
	Specify the desired TP-Function for the current temperature range (or current pressure range; only if the <i>lowest limit</i> is entered as a negative number).
	& <continuation current="" definition="" for="" of="" the="" tp-function=""></continuation>
	Specify the desired TP-Function continuation for the current temperature range (or current pressure range; only if the lowest limit is entered as a negative number).
	High temperature limit /6000/: <high in="" k="" limit="" temperature=""></high>
	Specify the high-temperature limit (in Kelvin), or the high-pressure limit (in Pascal; only if the lowest limit is entered as a negative number) for the current TP Function.
	Any more ranges /N/:

# LIST\_CONSTITUENT

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This is a GIBBS\_ENERGY\_SYSTEM (GES) command.

### Syntax LIST\_CONSTITUENT

List all the constitutions (as site-fractions) for all the phases in the defined system.



This is mainly for software managers. There are no commands in the GIBBS monitor to set the constitution.

# LIST\_DATA

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This is a GIBBS\_ENERGY\_SYSTEM (GES) command.

All data for the current system that is defined (retrieved from database in the DATA module, or directly entered in the GIBBS module) within the current GIBBS workspace is written in a readable manner on the output file. If no output file is specified, the data display on screen.

The output data consists of a list of all elements and their data followed by a list of all phases and the data associated with each phase. The thermochemical parameters listed for each phase are always in SI units.



For encrypted commercial databases you may not be able to use this command to list any retrieved data. However, it can be used to list some basic definitions (such as phase models and phase-constituents) retrieved from an encrypted database, as well as all the references related to original assessments (but not thermodynamic parameters) for the currently-defined system.

Syntax	LIST_DATA
	Output to screen or file /Screen/: <file name=""></file>
Prompts	Specify the name of a file in which the data shall be written in a readable manner. The default value is the screen (by pressing <enter>).</enter>
	Options?: <option(s)></option(s)>
	Choose one or several of these options for output:
	$\bullet{\ensuremath{\mathbb N}}$ the output is written as a user database format.
	<ul> <li>P the output is written as a macro file for future input. This is useful for creating *SETUP files for data assessments.</li> </ul>
	<ul> <li>s the symbols are suppressed.</li> </ul>
	<ul> <li>R the references for the parameters are listed (only for some databases in which references are available)</li> </ul>
	• ${\tt L}$ the output is written suitable for a LaTeX preprocessor.

# LIST\_PARAMETER



This command is for both the GIBBS and PARROT modules.

List the TP-function(s) of a specific parameter for a phase. You must supply the name of the phase parameter:

The parameter name: <Identifier>(<Phase>,<Constituent array>;<Digit>)



For encrypted commercial databases, you may not be able to use this command to list any retrieved parameter.

Syntax	LIST_PARAMETER
	Parameter: <parameter name=""></parameter>
	Specify a correct and complete parameter name, which should contain all the necessary parts of the general form: <identifier>(<phase>,<constituent array="">;<digit>)</digit></constituent></phase></identifier>
Prompts	See <u>"About the ENTER_PARAMETER Command" on page 133</u> to learn how to define the Parameter Name.
	If a parameter name is not acceptable or <enter> is pressed, an error message displays:</enter>
	*** Error, please re-enter each part separately
	The program prompts for separate input for each required part for a parameter name.
	Identifier (/X/): <g bm,="" l,="" or="" tc,="" va="" vb="" vc="" vk="" vo=""></g>
	Specify one of these types of legal identifiers.
	Phase name (/ABCD/): <phase name=""></phase>
	Specify the phase name.
	Constituent (in sublattice # /ABC/): <species name=""></species>
	Specify the constituent name on the specified sublattice site of the given phase.
	<pre>Interacting constituent (in sublattice # /XYZ/): <species name=""></species></pre>
	Specify the interacting constituent name on the specified sublattice site of the given phase; if there is no interacting constituent, press <enter>.</enter>

# Syntax LIST\_PARAMETER Interacting constituent (in sublattice # /XYZ/): <Species name> If there is more than one interacting constituent on the specified sublattice site of the given phase, specify them; otherwise press <Enter>. Degree /#/: <Degree> Specify a numerical number as the degree for the phase parameter.

# LIST\_PHASE\_DATA

This is a GIBBS\_ENERGY\_SYSTEM (GES) command.

All data for a specific phase are written in a readable manner on screen. The thermochemical parameters listed for the phase are always in SI units.



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>\_

For encrypted commercial databases you may not be able to use this command to list any phase data. However, this command can be used for listing some basic definitions (such as phase model and phase-constituents) for the specified phase, as well as all the references related to original assessments (but not thermodynamic parameters) for the currently-defined system.

Syntax	LIST_PHASE_DATA	
Prompt	Phase name: <phase name=""></phase>	
	Specify a phase name (if abbreviated, it should be unique).	

# LIST\_STATUS

This is a GIBBS\_ENERGY\_SYSTEM (GES) command.

# Syntax LIST\_STATUS For the GIBBS module, list the entered elements, phases and species with the status word. The command is included for system managers. Image: Comparison of the species of th

The bits are listed as hexadecimal, i.e. four bits are written as a hexadecimal number. Two hexadecimal numbers make a byte. In hexadecimal, 0 to 9 mean

### Syntax LIST\_STATUS

normal digits. A to F means values 10 to 15. The number E4000000 has thus bit 1, 2, 3 and 6 equal to one. The bits are numbered starting with 1 for the most significant (leftmost) bit.

### **ELEMENT STATUS WORD**

Set the element status word (Bit. Meaning) if:

- 1. Element cannot be deleted (only vacancy and electron)
- 2. Suspended (inclusive OR of bit 3 and 4)
- 3. Explicitly suspended
- 4. Implicitly suspended (cannot occur for an element)

### SPECIES STATUS WORD

Set species status word (Bit. Meaning) if:

- 1. Species record for an element (each element has a species record)
- 2. Suspended (inclusive OR of bit 3 and 4)
- 3. Explicitly suspended
- 4. Implicitly suspended (e.g. if a species element is suspended)
- 5. Charged (inclusive OR of bit 6 and 7)
- 6. Negative charge
- 7. Positive charge
- 8. Vacancy
- 9. Component (by default the elements are the components)

### PHASE STATUS WORD

Set phase status word (Bit. Meaning) if:

- 1. Ideal (no sublattices and no excess parameters)
- 2. Suspended (inclusive OR of bit 3 and 4)
- 3. Explicitly suspended

- 4. Implicitly suspended (e.g. if all constituents suspended)
- 5. Gas phase
- 6. Liquid phase
- 7. Solution phase
- 8. Only one constituent (in each sublattice)
- 9. Ions (inclusive OR of bit 10 and 12)
- 10. New ionic liquid model (charge balance handled internally)
- 11. Constituents cannot be added
- 12. External charge balance needed
- 13. Aqueous model
- 14. Charged species (ions)
- 15. Dilute entropy
- 16. Last calculation exceeded temperature range for any parameter
- 17. Kapoor-Frohberg-Gaye cell model
- 18. Turbo calculation used
- 19. Turbo calculation impossible
- 20. Turbo calculation illegal
- 21. Phase is not ideal
- 22. Current site fractions saved
- 23. This phase is the ordered part
- 24. This phase is the disordered part (bit 2 and 3 also set)
- 25. Shadow phase with diffusion data
- 26. Error in ionic model
- 27. Sometimes
- 28. CVM-SRO (short-range ordering) entropy expression
- 29. CVM initialization is made
- 30. Used to test quasi-chemical ordering model
- 31. Major constituents check even if no miscibility gap (ordering)
- 32. Hoch-Arpshofen model

### SYMBOL STATUS WORD

Set symbol status word (Bit meaning) if:

- 1. Constant
- 2. Variable
- 3. Function
- 4. Table
- 5. Value must not be amended
- 6. Undefined

# LIST\_SYMBOL

This is a GIBBS\_ENERGY\_SYSTEM (GES) command.

List the symbols available in the current GIBBS workspace, which are defined (retrieved from database in the DATA module, or directly entered in the GIBBS module) and used in TP-functions for the entered model parameters for various phases in the defined system.



>\_

For encrypted commercial databases you may not be able to use this command to list any retrieved symbol.

Syntax	LIST_SYMBOL
	Name: <symbol name=""></symbol>
	Specify a symbol name (either as the full name of a defined symbol, or as the first few common characters of several symbols which have names starting with such common characters).
Prompts	Only those symbols that match this name are listed.
	Or press <enter> for a list of all the symbols defined (retrieved from database in the DATA module, or directly entered in the GIBBS module) for the current system, which are available in the current GIBBS workspace.</enter>
	Output to screen or file /Screen/: <file name=""></file>
	Specify the name of a simple-textual file in which the entered symbols shall be written in a readable manner (such a basic textual file can later on be opened and edited by any text editor). The default value is the screen (by pressing <enter>).</enter>

# **READ\_GES\_WORKSPACE**

1	
	`
	/_

This command is specific to the GES (GIBBS) module.

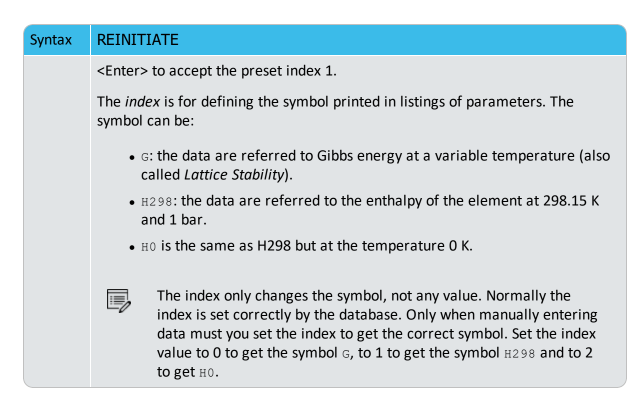
The data area saved onto a \*.GES5 file with <u>"SAVE\_GES\_WORKSPACE" on the next page</u> is read back into the GIBBS workspace.

Syntax	READ_C	GES_WORKSPACE
	Look in	<b>File name</b> and specify the working directory where to save the file in the field. Normally the default file-type in the <b>Files of type</b> field is the correct the GIBBS workspace format (i.e. *.GES5 file).
		The saved *.GES5 files are unique for each CPU type, and therefore a *.GES5 file saved on one CPU type cannot be read and used on another CPU type.

# REINITIATE

All data in the stored GIBBS workspace are erased, and all variables are initiated to their default values.

Syntax	REINITIATE
	Upper case only /Y/:
Prompts	Press <enter> to keep the default where the name of elements and species is in only UPPER case. Or choose N to have the elements with two letter names with the first letter in Upper case and the second in lower case.</enter>
	In upper case mode all input in lower case is automatically converted to upper case.
	Lower temperature limit /298.15/: <lowest in="" k="" t=""></lowest>
	This value is used as the lower temperature limit when data are entered interactively.
	Upper temperature limit /6000/: <highest in="" k="" t=""></highest>
	This value is used as the upper temperature limit when data are entered interactively.
	Default element reference state symbol index /1/: <1 or 2 or 3>
	Specify a proper index for the default listing parameters (symbol), or press



# SAVE\_GES\_WORKSPACE



This command is for the GES (GIBBS) module.

The data area used by the GIBBS workspace can be saved on a file (with a default extension GES5 under MS-Windows, or ges5 under Linux). The default file name is RESULT.GES5 or equal to the one used in a previous <u>"READ\_GES\_WORKSPACE" on the previous page</u> command.

Syntax	SAVE_GES_WORKSPACE
	Enter a <b>File name</b> and specify the working directory where to save the file in the <b>Save in</b> field. Normally the default file-type in the <b>Files of type</b> field is the correct
	one for the GIBBS workspace format (i.e. *.GES5 file).

The output is unformatted and cannot be printed.

# SET\_R\_AND\_P\_NORM

The value of the gas constant (R) is used to define the energy units that are used for input of data. On output, all data are in SI units. The value of the pressure 1 atm is used to interpret values of the pressure given by application programs.

Syntax	SET_R_AND_P_NORM
	Value of gas constant in your energy units /8.31451/: <new value=""></new>
Prompts	The default value of the gas constant is in SI units (i.e. 8.31451). If you change this value, it only affects the interpretation of energy data given interactively from the terminal, e.g. if the value is set to 1.98717 you can give values in Calories.
	Value of one bar in your presure units /100000/: <new value=""> The value of the pressure given in application programs is divided by this value and then multiplied with the value of one bar expressed in Pascal before it is</new>
	used inside the GIBBS module.

# **Optimizing with the PARROT Module**

### In this section:

AMEND_PARAMETER
AMEND_SYMBOL
COMPILE_EXPERIMENTS
CONTINUE_OPTIMIZATION
CREATE_NEW_STORE_FILE
EDIT_EXPERIMENTS
About the ENTER_PARAMETER Command
Defining the Parameter
ENTER_PARAMETER
LIST_ALL_VARIABLES
LIST_CONDITIONS
LIST_PARAMETER
LIST_PHASE_DATA (from GIBBS)
LIST_RESULT
LIST_STORE_FILE
LIST_SYMBOL_IN_GES
OPTIMIZE_VARIABLES
READ_PARROT_WORKSPACES
RECOVER_VARIABLES
REINITIATE
RESCALE_VARIABLES
SAVE_PARROT_WORKSPACES
SET_ALTERNATE_MODE
SET_EXTERNAL_PARAMETER

SET_FIX_VARIABLE	
SET_OPTIMIZING_CONDITION	
SET_OPTIMIZING_VARIABLE	
SET_OUTPUT_LEVELS	154
SET_SCALED_VARIABLE	
SET_STORE_FILE	156

# **AMEND\_PARAMETER**

>\_

This AMEND\_PARAMETER command is for the PARROT module. There is also a GES module command with the same name (<u>"AMEND\_PARAMETER" on page 79</u>).

Interactively modify the TP-function(s) for a specific parameter for a phase with this command. This is useful in order to correct typing errors because the old function is made available for interactive editing on the terminal.

Syntax	AMEND_PARAMETER
	Parameter: <parameter name=""></parameter>
Prompts	Specify a correct parameter name. If a parameter name is not acceptable or <enter> is pressed, the error message displays:</enter>
	*** Error, please re-enter each part separately
	The program prompts for separate input for each part for a parameter name.
	Identifier (/X/): <g bm,="" l,="" or="" tc,="" va="" vb="" vc="" vk="" vo=""></g>
	Specify one of these types of legal identifiers (see <u>"ENTER_PARAMETER" on</u> page 137).
	Phase name (ABCD/): <phase name=""></phase>
	Specify the phase name.
	<pre>Constituent (in sublattice # /ABC/): <species name=""></species></pre>
	Specify the constituent name.
	<pre>Interacting constituent (In sublattice # /XYZ/): <species name=""></species></pre>
	Specify the interacting constituent name; if there is no interacting constituent, press <enter>.</enter>
	<pre>Interacting constituent (in sublattice # /XYZ/): <species name=""></species></pre>
	If there is more than one interacting constituent, specify these; otherwise press <enter>.</enter>
	Degree /#/: <degree></degree>
	Specify a numerical number as the degree of the phase parameter.
	After the parameter name is specified correctly, the program lists its current definition (either preset in database or defined by the ENTER_PARAMETER

### Syntax AMEND\_PARAMETER

### command), such as:

```
L(PHASE2,AL,MG;1) =
298.15<T<2000.00: +5000
2000.00<T<4500.00: +4500
4500.00<T<6000.00: +4000
```

You are prompted to change the parameter definition:

```
Do you want to change the number of ranges /No/:
```

Enter Y to change the number of ranges for the chosen function, or change some of the temperature limits in the definition, then retype both the low/high temperature limits and functions

If you do not want to change the number of ranges but want to change the function(s) in one or more ranges, press <Enter> and the whole definition of the chosen parameter in all ranges (if any) is listed such as:

```
DIFFERENT FUNCTIONS IN THESE RANGES
298.15<T<2000.00
2000.00<T<4500.00
4500.00<T<6000.00
```

This message prompts:

Do you want to change range limits /No/:

If there is more than one range, this question is prompted. Press <Enter>.

Range number (0 to exit) /0/: <Range number>

If the function of a parameter is different in two or more temperature ranges, you must specify the range of the function of which you want to amend. Or press <Enter> or type 0 to exit this command without making any change.

```
Function
```

The previous function is available for editing. The editing is performed within the general subroutine FOOLED, as described in <u>"AMEND\_SYMBOL" on the next page</u>. This routine prompts as follows:

1:+:>

The prompt consists of the current position in the string and the character at that position between colons, (::).

Syntax	AMEND_PARAMETER
	COMMANDS
	These commands can be given:
	• Help: ?
	• Move CP to last or first character: <+/-> A
	• Delete characters from CP: <+-#characters> D
	• Exit: E
	• Find: <#occurrences> F <string>@</string>
	• Insert: I <string>0</string>
	• Move: <+-#positions> M
	Restore string: R
	• Substitute: s <old>@<new>@</new></old>
	• Type string: T
	Where CP denotes the current position in a string, # means number of, @ is a terminator of an input or search string.
	When the string is typed the character at the current position is replaced by an underscore
	To finish the editing of the current function, type ${\ensuremath{\mathbb E}}$ at the prompt.
	Range number (0 to exit) /0/: <range number=""></range>
	Give a range number to edit that function, or press <enter> or type 0 to exit this command.</enter>

# AMEND\_SYMBOL



This command is for the GES (GIBBS) and PARROT modules.

Interactively calculate the current values of a function or table (predefined by the switched database or previously entered with <u>"ENTER\_SYMBOL" on page 109</u>) at the current temperature and pressure condition, and also possible to modify the definitions of variables or functions.

The current values of the temperature and pressure used to calculate the functions or tables are the ones listed with <u>"LIST\_STATUS" on page 117</u>. There is no way to change the current temperature and pressure values interactively.

To modify the definitions of entered symbols (variable, functions or parameters), the performance of this command is slightly different one from another, as described below:

• For variables, the values can be changed.

- For functions the low and high temperature limits in various ranges (if any), as well the expressions, can be changed.
- For parameters the low and high temperature limits in various ranges (if any), as well the expressions, can be changed.

Syntax	AMEND_SYMBOL
	Name: <symbol name=""></symbol>
	Specify the name of an entered symbol.
Prompts	For tables and functions (or parameters which are treated as functions after these are entered by the ENTER_SYMBOL command with the Parameter keyword, but not by the ENTER_PARAMETER command), after the symbol name is specified here, the program automatically calculates the values under the current temperature and pressure conditions, and lists the current values, such as:
	FUNCTION VALUE 2.52500000E+01
	TABLE VALUE 1.5600000E+02
	For variables and functions (or parameters), there are additional prompts and depend on the symbol type.
	For a variable, its current value is displayed, and you can change it to a new value:
	Value /Current value/: <new value=""></new>
	For a function (or a parameter entered as a symbol):
	Do you want to change the number of ranges /No/:
	Enter Y to change the number of ranges for the function, or change some of the temperature limits in the definition, then retype both the low/high temperature limits and functions (see all the remaining details in the ENTER_SYMBOL command.
	If you do not want to change it, press <enter>. The definition of the chosen</enter>

### Syntax AMEND\_SYMBOL

function in all ranges (if any) is listed, for example:

DIFFERENT FUNCTIONS IN THESE RANGES

298.15<T<2000.00

2000.00<T<4500.00

4500.00<T<6000.00

and this message displays:

Do you want to change range limits /No/:

If a function is different in two or more temperature ranges, you must specify the range of the function of which you want to amend. Or press <Enter> or type 0 to exit this command without making any change.

Range number (0 to exit) /0/: <Range number>

Function:

The previous function is available for editing. The editing is performed within the general subroutine FOOLED. This routine prompts as follows:

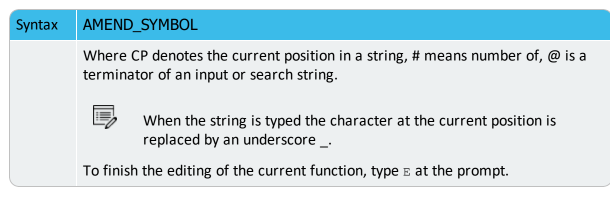
1:+:>

The prompt consists of the current position in the string and the character at that position between colons (::).

### COMMANDS

These commands can be given:

- Help: ?
- Move CP to last or first character: <+/-> A
- Delete characters from CP: <+-#characters> D
- Exit: E
- Find: <#occurrences> F<string>@
- Insert: I<string>@
- Move: <+-#positions> M
- Restore string: R
- Substitute: s<old>@<new>@
- Type string: T



# **COMPILE\_EXPERIMENTS**

The descriptions of the experimental equilibria saved on a textual \*.POP file, given as commands in the POLY and ED\_EXP syntax, is compiled by this command into the PARROT structured data, i.e. stored into the current work \*.PAR file which is created with <u>"CREATE\_NEW\_STORE\_FILE" on page 133</u>, set by <u>"SET\_STORE\_FILE" on page 156</u>, or opened with <u>"READ\_PARROT\_WORKSPACES" on page 148</u>, all done before COMPILE\_EXPERIMENTS.

This command also lists the compilation details on screen or onto a listing file (\*.TXT).



If a syntax error in the commands is detected, the compilation is terminated. You need to correct the error in the \*.POP file (using a textual editor) and recompile it with the \*.PAR work (store) file.

After a successful compilation, the program translates all the experimental data points in the \*.POP file into a graphical experimental data file (\*.EXP) in the DATAPLOT syntax, which can be further edited and used as a graphical experimental data file (\*.EXP) for the purposes of imposing onto calculated/plotted phase diagrams and/or property diagrams for comparisons during the assessment process.

**DATAPLOT** User Guide included with this documentation set.

### Syntax COMPILE\_EXPERIMENTS

A window displays to open the original experimental data file (\*.POP). Enter a \*.POP **File name** box, and specify the working directory in the **Look in** box. Normally the default file-type in the **Files of type** field is the correct one for the experimental data format (i.e. \*.POP file). Click **Open** or **Cancel** button as applicable.

Output to screen or file /Screen/: <\*.pop file name>

Prompts A list of the source code in the \*.POP file and error messages during compilation is written on screen (by pressing <Enter>) or on a list \*.TXT file under a specific file name which can later on be opened and edited by any basic

### Syntax COMPILE\_EXPERIMENTS

textual editor.

Initiate store file /Y/:

After a successful compilation, the data in the POLY3 workspace is stored as a new block of equilibria on the current work (store) file. By default the store file is initiated before compilation and only the compiled block is stored on the current work file (consequently, the \*.PAR file is updated).

A window displays to generate/store the graphical experimental data file (\*.EXP) in the DATAPLOT syntax. Enter a \*.EXP file name in the **File name** field and specify the working directory when opening the \*.EXP file in the **Look in** field. Normally the default file type in the **Files of type** field is the correct one for the graphical experimental data format (i.e. \*.EXP file). Click **Open** or **Cancel** as applicable.

To keep the current work file, enter N.

### **CONTINUE\_OPTIMIZATION**

This command continues the optimization using the same Hessian matrix.

In some cases it is not possible to attempt continuation. For example, if the optimization has already converged or if the optimizing parameters or the set of experimental equilibria used in the current optimization runs are changed. The program gives a warning message if you try to continue when you make these changes.

Syntax	CONTINUE_OPTIMIZATION
	Are you sure? /N/: <n or="" y=""></n>
Prompts	Prior to this question, a message displays to confirm if it is OK to continue:
	It is safe to CONTINUE only after TOO MANY ITERATIONS $% \left( \left( {{{\left( {{{\left( {{{}} \right)}} \right)}}}} \right)$ and no change in variables and experiments
	You can accept the default (N) in order to cancel this special attempt.
	Number of iterations /N/: <integral iteration="" number="" of=""></integral>
	Specify an integral number for the optimization iteration. The specified iteration number is shown as the default number. The PARROT program tries this exact number of different sets of values of the optimizing variables without stopping the optimization even it would have converged earlier or should give up earlier.

# **CREATE\_NEW\_STORE\_FILE**

This file is hardware dependent and cannot be read by any text editor.

Create a binary file to be used as a *work file* (also called *store file*) before any optimization can be done. The workspace used by the GIBBS, POLY and PARROT modules is stored automatically on the work file but not any experimental information. The binary work file has a default extension of PAR under MS-Windows or par under Linux/UNIX.

A work file that is created at a previous run can be used in the PARROT module with the other command SET-STORE-FILE.

Syntax	CREATE_NEW_STORE_FILE
	<*.par file name>
Prompt	A <b>Save</b> window displays. Enter a <b>File name</b> and specify the working directory in the <b>Save in</b> field. Normally the default file-type in the <b>Files of type</b> field is the correct one for the PARROT workspace format (i.e. PAR file). Click <b>Save</b> or <b>Cancel</b> as applicable.

# EDIT\_EXPERIMENTS

Initialize and make available the sub-module for editing experimental equilibria, i.e. the ED\_EXP module. All experimental equilibria compiled from the \*.POP file can be accessed in the ED\_EXP module.

The ED\_EXP module is similar to the normal POLY module, but some commands are special and some POLY commands are not available. In the ED\_EXP module, you can calculate each equilibrium separately or together, provide start values for equilibria which failed to converge, set weights, and modify the values of experiments or conditions.

"Running the EDIT\_EXPERIMENTS Command" on page 26

### Syntax EDIT\_EXPERIMENTS

You are now in this submodule and can start using the commands.

# About the ENTER\_PARAMETER Command



For the PARROT Module, this command is the same and is described here. See <u>"ENTER\_PARAMETER" on page 137</u> for the command details.

In the descriptions of the standard thermochemical properties and special physical properties for a phase, there are a number of parameters which may depend on the temperature and pressure. The expressions for these parameters can be given in a free form as a sum of terms with powers of T and P and may also include the natural logarithm and exponential function. This type of expression is called *TP-functions*. Identical parameters (in terms of parameter-names) are stored only once in the GIBBS workspaces.

The composition-dependence of the Gibbs energy is described in the GIBBS module by the internal data structure, which is created when the phase is entered (see <u>"Gibbs Energy</u> <u>System (GES) Commands" on page 76</u>). The Gibbs energy of a phase is always referred to one formula unit of the phase, i.e. the amount derived from the number of sites (i.e. the stoichiometric coefficient) for each sublattice. If vacancy is a constituent of a sublattice, the amount of matter per formula unit of the phase may vary with composition.

### **Defining the Parameter**

A valid parameter should have the general form of:

```
<identifier>(<phase>, <constituent array>; <digit>) <xxx> <expression> <yyy> <keyword Y or N> <zzz> !
```

The identifier must be followed by an opening parenthesis, a phase name, a comma and a constituent array. Optionally, the constituent array can be followed by a semicolon and a digit. The parameter name is terminated by a closing parenthesis. The parameter form is defined as:

- <identifier> is the parameter type;
- <phase> is the phase name (maximum 24 characters);
- <constituent array> is the specific constituent array in the phase;
- <digit> is the degree of composition-dependent interaction contribution (an integer number from 0 through 9), that is only for excess energy (L), Curie temperature (TC) and Bohr magneton number (BMAGN), as well as for volume-related parameters (V0 or VA or VB or VC or VK); if it is valued as zero, or if it is for the standard Gibbs energy (G) for which the degree is always zero, it can be omitted;
- <expression> is the mathematical relation to describe the parameter;
- <xxx> and <yyy> are the low and high temperature limits respectively for the applicable temperature range of the parameter expression;
- <keyword Y or N> is the indicator on if there is continuation for the parameter expression or not;
- <zzz> is the reference index/number for the assessment of this parameter;
- The exclamation point ! is used to indicate that the current parameter definition is ended.

### PARAMETER NAME

### The GES parameter name has a general form of:

<identifier>(<phase>,<constituent array>;<digit>)

### Examples of parameter names:

- G (GAS, C102): The Gibbs energy of formation of a CO2 molecule in gas.
- G (FCC, FE:VA): The Gibbs energy of formation of fcc Fe with interstitials.
- L(LIQ, Fe, Cr; 0: The regular solution parameter for Fe and Cr in liquid.
- L(LIQ, Fe, Cr; 1): The sub-regular solution parameter.
- TC (BCC, Fe:Va): The Curie temperature of bcc Fe.
- BMAGN (BCC, Fe:Va): The Bohr magneton number parameter of bcc Fe.

The parameter name consists of several parts. The first is a type-identifier and these can be used:

- G: Standard energy parameter (Gibbs energy of formation) or for interaction parameters;
- L: Excess energy parameter (Gibbs energy of interaction) always used for interaction parameters;
- TC: Curie temperature for magnetic ordering;
- BMAGN or BM: Bohr magneton number for magnetic ordering (or Born function  $\omega$ Pr,Tr for aqueous solute species).
- vo: Molar volume at 298.15 K and 1 bar (a numeric value only);

• VA: Integrated thermal expansivity; 
$$\int_{298.15}^{T} lpha(T) \mathrm{dT}$$

- VB: Bulk modulus at 1 bar;
- vc: Isothermal compressibility;
- VK: High-pressure fitting parameter.



When necessary quantities as H (enthalpy), S (entropy), V (Volume), F (Helmholtz energy), etc., can be calculated from the Gibbs energy.

### PHASE NAME

Specifying the PHASE NAME in uppercase is recommended; however, if you prefer to write it

as a mixture of uppercase and lowercase, it automatically converts all lowercase to uppercase, as the GIBBS module only recognises uppercase phase names. It is important that if a phase bears a legal phase-type (among G, A, Y, L, I, F and B) in its phase definition (already by the PHASE keyword; such as GAS:G, LIQUID:L,SLAG:L, IONIC-LIQ:Y, SPINEL:I, FCC\_ L12:F, HCP\_D021:F, BCC\_B2:B, AQUEOUS:A), such a valid phase-type code should not be attached to the phase name when ENTER\_PARAMETER is executed.

### **CONSTITUENT ARRAY**

The constituent array consists of a list of constituent names. Interaction parameters have two or more constituents from the same sublattice separated by a comma. If the phase has sublattices, at least one constituent in each sublattice must be specified. The constituents in different sublattices must be given in sublattice order and are separated by a colon.

After the component array, a sub-index digit can be specified after a semicolon. This digit must be in the range 0 to 9. The interpretation of the sub-index depends on the excess energy model used for the phase. If no semicolon and digit are given, the sub-index value is assumed to be as zero.

The excess energy parameters, e.g. the regular/subregular (binary) parameter or ternary parameters, are multiplied with two or more fractions of the constituents from the same sublattice of the solution phase. These additional constituents must be given as interacting constituents (as the following prompt).

Solution phases with sublattices can have interacting constituents in each sublattice.

### INTERACTION PARAMETER

An interaction parameter, which is used to describe the excess term of a quantity, must have two or more constituents that interact with each other on a specified sublattice site of the given phase. It is arbitrary which of these constituents is given as the first constituent and what is given as the interacting constituents. The software always sorts the constituents (in each sublattice) in alphabetical order when the parameter name is written as a prompt (for entering its parameter value) and when the parameter is listed (using the GIBBS commands "LIST\_PARAMETER" on page 141 or "LIST\_PHASE\_DATA (from GIBBS)" on page 142). This is important for all asymmetric interaction parameters where the sign of the interaction parameter must depend on the appearance order.

Use an asterisk\* to denote that the excess interaction parameter is independent of the constituents of a specific sublattice. For example,  $L(FCC_L12, AL, NI:*)$  means that the interaction parameter is for the binary interaction between constituents AL and NI on the

first sublattice in the FCC\_L12 solution phase, while it is independent of all constituents on the second sublattice. An interaction parameter in the list of constituents is always added to the Gibbs energy and the asterisk \* is calculated with the term of  $[1\sum y(specified$ constituents)], which implies that in an A-B binary system these L parameters are identical (but in higher-order systems, these are different):

- L(phase, A, B) is multiplied with X(A)\*X(B)
- L(phase, A, \*) is multiplied with X(A)\*(1-X(A))
- L(phase, B, \*) is multiplied with X(B)\*(1-X(B))

If you press <Enter> when you are asked for a parameter name or if you have improperly input the entire parameter name, you are asked for each of these items in the name.

# **ENTER\_PARAMETER**



This command is for both the GES and PARROT modules.

### "About the ENTER\_PARAMETER Command" on page 133

Use this command to enter TP-function(s) for a specific parameter for a phase interactively with this command. If there is already a parameter expression defined (for this parameter), that is deleted and replaced with newly entered one.

The entered TP-Function(s) for the parameter can be changed later with the command "AMEND\_PARAMETER" on page 79.

Syntax	ENTER_PARAMETER
	Parameter: <parameter name=""></parameter>
Prompts	As explained in <u>"About the ENTER_PARAMETER Command" on page 133</u> , specify a correct and complete Parameter Name, which should contain all the necessary parts of the general form: <identifier>(<phase>, <constituent array="">;<digit>) If a parameter name is not acceptable or <enter>, is pressed, a message displays: *** Error, please re-enter each part separately and you are prompted for input for each required part for a parameter name.</enter></digit></constituent></phase></identifier>
Options	Description or Information

Syntax	ENTER_PARAMETER
	Identifier (/X/): <g bm,="" l,="" or="" tc,="" va="" vb="" vc="" vk="" vo=""></g>
Identifier	If this command is used one or more times, the previous value on this prompt is set as default. Press <enter> for the same type identifier or specify a new type.</enter>
	Phase name (/ABCD/): <phase name=""></phase>
Phase name	Each parameter is valid for a specific phase only. The name of that phase must be supplied (maximum 24 characters). The name can be abbreviated.
	Constituent (in sublattice # /ABC/): <species name=""></species>
	A parameter is identified by the constituents on a specified sublattice site of the given phase, the stoichiometric coefficients of which are multiplied with the parameter. The name of the constituent can be abbreviated.
Constituent	It is the species name, not the stoichiometric formula required here.
	If this command is used one or more times, the previous value on this prompt is set as default. Accept it by pressing <enter> if the constituent is the same, or specify a new species name.</enter>
	For phases with several sublattices, the program asks for one constituent in each sublattice.
	<pre>Interacting constituent (in sublattice # /XYZ/): <species name=""></species></pre>
Interacting constituent	If this command is used one or more times, the previous value on this prompt is set as default. Press <enter> to accept it if the constituent is the same, or specify a new species name.</enter>
	To cancel the default value of the interacting constituent type NONE or the name of another constituent.
	This question is repeated until all the interested interacting constituent(s) on a specific sublattice in the phase are specified, and finally an <enter> is enforced.</enter>
	Degree /#/: <degree></degree>
Degree	Degree is model-dependent. Specify an integer number (a value from 0 through 9) as the degree of composition-dependent interaction

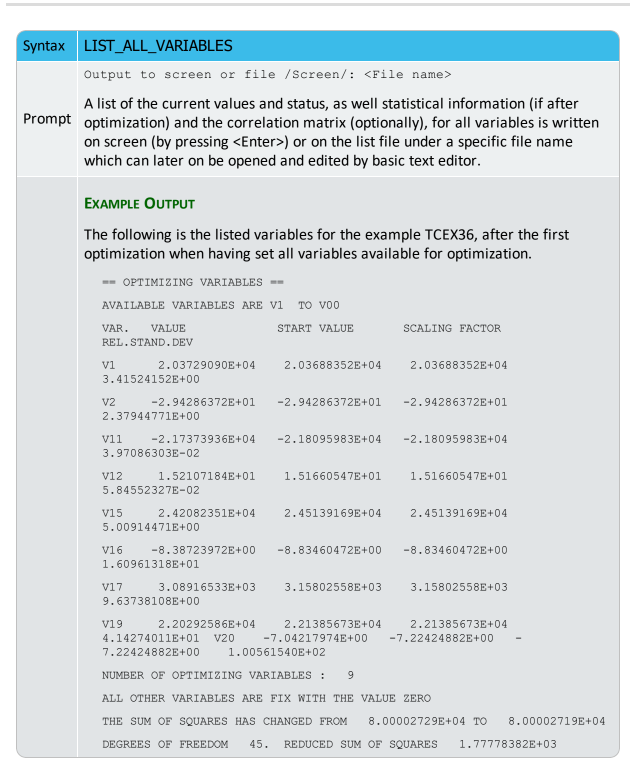
Syntax	ENTER_PARAMETER
	contribution for the phase parameter. This is valid for excess energy (L), Curie temperature (TC) and Bohr magneton number (BMAGN), as well as for volume-related parameters (V0, VA, VB, VC or VK).
	For binary interaction parameters, the degree is usually the power in the Redlich-Kister expression. For ternary interaction parameters, it is usually the Hillert ternary index.
	For a standard G parameter for a pure component (end-member) its degree should be always 0 and this prompt should not display.
	After this prompt, the program echoes on the screen the full TP-Function of the phase parameter.
	Low temperature limit /298.15/: <lowest in="" k="" limit="" temperature=""></lowest>
	Specify the lowest temperature limit (in Kelvin), or the lowest-pressure limit (in Pascal but entered as a negative number), for the current TP-Function.
	Function: <definition a="" for="" function=""></definition>
Phase	A TP-Function consists of terms in T and P.
parameter	& <continuation current="" definition="" for="" function="" of="" the=""></continuation>
	Continuation of a TP-Function definition.
	High temperature limit /6000/: <high in="" k="" limit="" temperature=""></high>
	Specify the high temperature limit (in Kelvin), or the high-pressure limit (in Pascal; only if the <i>lowest limit</i> is entered as a negative number) for the current TP-Function.
	Any more ranges? /N/:
	Enter Y for more function(s) or N to end this command.

# LIST\_ALL\_VARIABLES

This is a PARROT module command.

Create a list of the values and status of all variables in the PARROT workspace on screen or in a text file. After an optimization some statistical information is also written. The correlation matrix is written if that option is chosen.

SET\_OUTPUT\_LEVELS" on page 154.



# LIST\_CONDITIONS



This is a PARROT module command.

In the PARROT module, generate a list of the current values of optimization conditions, as well the current status of listing, either on screen or in a basic text file which can be opened and edited by a text editor.

	Syntax	LIST_CONDITIONS
	Prompt	Output to screen or file /Screen/: <file name=""></file>
		Press <enter> to output to screen or on the list file under a specific file name.</enter>

# LIST\_PARAMETER



This command is for both the GIBBS and PARROT modules.

List the TP-function(s) of a specific parameter for a phase. You must supply the name of the phase parameter:

The parameter name: <Identifier>(<Phase>,<Constituent array>;<Digit>)



For encrypted commercial databases, you may not be able to use this command to list any retrieved parameter.

Syntax	LIST_PARAMETER
	Parameter: <parameter name=""></parameter>
	Specify a correct and complete parameter name, which should contain all the necessary parts of the general form: <identifier>(<phase>,<constituent array="">;<digit>)</digit></constituent></phase></identifier>
Prompts	See <u>"About the ENTER_PARAMETER Command" on page 133</u> to learn how to define the Parameter Name.
	If a parameter name is not acceptable or <enter> is pressed, an error message displays:</enter>
	*** Error, please re-enter each part separately
	The program prompts for separate input for each required part for a parameter name.
	Identifier (/X/): <g bm,="" l,="" or="" tc,="" va="" vb="" vc="" vk="" vo=""></g>
	Specify one of these types of legal identifiers.
	Phase name (/ABCD/): <phase name=""></phase>

### Syntax LIST\_PARAMETER

Specify the phase name.

Constituent (in sublattice # /ABC/): <Species name>

Specify the constituent name on the specified sublattice site of the given phase.

Interacting constituent (in sublattice # /XYZ/): <Species name>

Specify the interacting constituent name on the specified sublattice site of the given phase; if there is no interacting constituent, press <Enter>.

Interacting constituent (in sublattice # /XYZ/): <Species name>

If there is more than one interacting constituent on the specified sublattice site of the given phase, specify them; otherwise press <Enter>.

Degree /#/: <Degree>

Specify a numerical number as the degree for the phase parameter.

## LIST\_PHASE\_DATA (from GIBBS)



This is a PARROT module command.

All data for a specific phase are written on screen. The thermochemical parameters listed for each phase are always in SI units.

#### Syntax LIST PHASE DATA

Phase name: <Phase name>
Prompt
Specify a phase name.

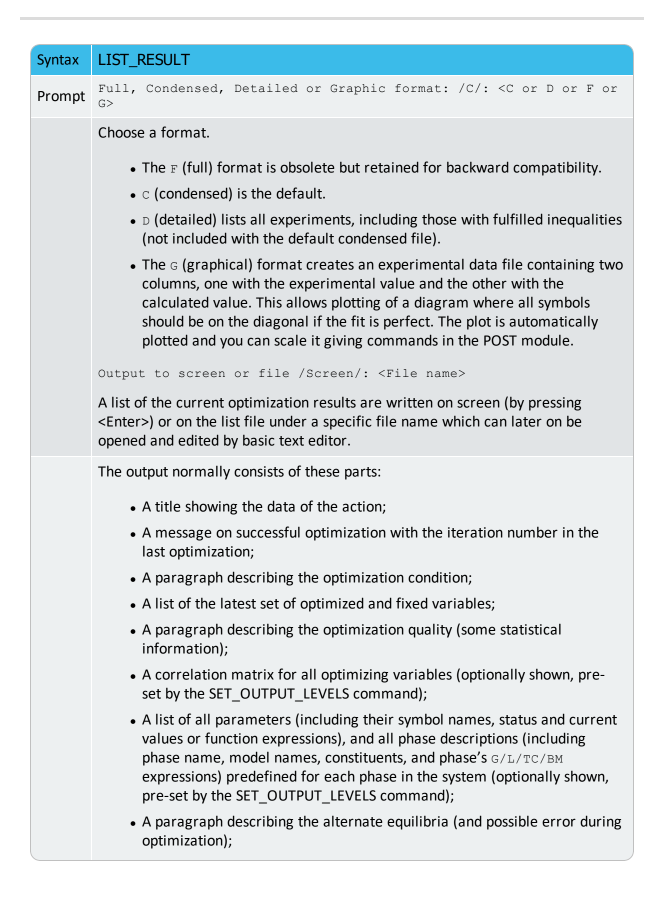
## LIST\_RESULT



This is a PARROT module command.

List the result of the current optimization run on screen or in a specified file. The level of detail listed can be prescribed by the command <u>"SET\_OUTPUT\_LEVELS" on page 154</u>.

The LIST\_RESULT command has an option G for creation of an experimental data file with two columns, one for the experimental value and the other for the calculated value. It allows plotting a diagram and visualizing the fitting results. The plot is automatically plotted and you can scale it giving commands in the POST module. The option D lists all experiments, including those with fulfilled inequalities which are suppressed with the default option C.



Syntax	LIST_RESULT
	<ul> <li>A detailed list on all the equilibrium points used in the current optimization. If the alternate-mode is used in the optimization, only an error value is shown. For normal-mode calculations, a list is shown for the experimental equilibrium numbers (first column) and corresponding original experimental data (second column, in the form quantity = value) which are used in optimization (i.e. non-zero weighted points). Such experimental data are by each point compared by the calculated value (third column) after the last optimization. Also listed are the experimental error (column 4), the difference between the calculated value and original experimental data (column 5), and the contribution to the sum of least square (column 6).</li> </ul>

#### **EXAMPLE OUTPUT**

The following is the listed result in the example <u>tcex36</u> in the *Thermo-Calc Console Mode Examples Guide*, after the first optimization based on the prescribed definitions in the TCEX36a.TCM and TCEX36b.TCM, but having set all variables available for optimization.

\_\_\_\_\_

```
OUTPUT FROM P A R R O T.
*** SUCCESSFUL OPTIMIZATION. *** NUMBER OF ITERATIONS: 10
== OPTIMIZING CONDITIONS ==
  RELATIVE STANDARD DEVIATIONS FOR EXPERIMENTS: N
  MINIMUM SAVE ON FILE: Y
  ERROR FOR INEQUALITIES = 1.0000000E+00
  RELATIVE STEP FOR CALCULATION OF DERIVATIVES = 1.0000000E-04
ARGUMENTS FOR SUBROUTINE VA05AD (HSL)
  MAXFUN = 1 DMAX = 1.00000000E+02 H = 1.00000000E-04 ACC = (INITIAL
SUM OF SQUARES) * 1.0000000E-03
== OPTIMIZING VARIABLES ==
AVAILABLE VARIABLES ARE V1 TO V00
VAR.
      VALUE
                     START VALUE
                                     SCALING FACTOR REL.STAND.DEV
      2.03749463E+04 2.03729090E+04 2.03729090E+04
V1
                                                       3.41455863E+00
V2
      -2.94286372E+01 -2.94286372E+01 -2.94286372E+01 2.37944774E+00
      -2.17395673E+04 -2.17373936E+04 -2.17373936E+04 3.98405298E-02
V11
      1.52107184E+01 1.52107184E+01 1.52107184E+01 5.82861832E-02
V12
V15 2.42106560E+04 2.42082351E+04 2.42082351E+04 5.07239609E+00
```

```
V16
      -8.38723972E+00 -8.38723972E+00 -8.38723972E+00
                                                          1.69546796E+01
V17
       3.08947424E+03
                         3.08916533E+03
                                          3.08916533E+03
                                                             9.85220694E+00
        2.20314615E+04 2.20292586E+04 2.20292586E+04
V19
                                                          4.16329629E+01 V20
-7.04217974E+00 -7.04217974E+00 -7.04217974E+00 1.03161466E+02
NUMBER OF OPTIMIZING VARIABLES :
                                 9
ALL OTHER VARIABLES ARE FIX WITH THE VALUE ZERO
THE SUM OF SQUARES HAS CHANGED FROM 8.00002719E+04 TO 8.00002709E+04
DEGREES OF FREEDOM
                   45. REDUCED SUM OF SOUARES 1.77778380E+03
Number of alternate equilibria
                                        14
SYMBOL
             STATUS VALUE/FUNCTION
         80000000 8.3145100E+00
R
RTLNP
         20000000 +R*T*LN(1E-05*P)
V1
         48000000 2.0374946E+04
         48000000 -2.9428637E+01
V2
V11
         48000000 -2.1739567E+04
V12
         48000000 1.5210718E+01
V15
         48000000 2.4210656E+04
V16
         48000000 -8.3872397E+00
V17
         48000000 3.0894742E+03
              48000000 2.2031462E+04 22 V20 48000000 -7.0421797E+00
 21 V19
LIQUID
EXCESS MODEL IS REDLICH-KISTER_MUGGIANU
                                                                   CONSTITUENTS:
A,B
     G(LIQUID,A;0)-G(BCC,A;0) = 500.00<T< 2000.00: +14000-10*T
     G(LIQUID,B;0)-G(BCC,B;0) = 500.00<T< 2000.00: +18000-12*T
     L(LIQUID, A, B; 0) =
                         500.00<T< 2000.00: +V11+V12*T
                                                            L(LIQUID, A, B; 1) =
500.00<T< 2000.00: +V13+V14*T
A2B
  2 SUBLATTICES, SITES 2: 1
                               CONSTITUENTS: A : B
      G(A2B, A:B; 0) - 2 G(BCC, A; 0) - G(BCC, B; 0) =
            500.00<T< 2000.00: +V1+V2*T+V3*T*LN(T)
BCC
EXCESS MODEL IS REDLICH-KISTER MUGGIANU
                                                                   CONSTITUENTS:
A,B
      G(BCC, A; 0) - G(BCC, A; 0) = 500.00 < T < 2000.00: 0.0
      G(BCC,B;0) - G(BCC,B;0) = 500.00 < T < 2000.00: 0.0
      L(BCC, A, B; 0) = 500.00<T< 2000.00: +V15+V16*T
```

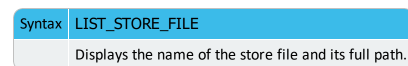
```
L(BCC,A,B;1) = 500.00<T< 2000.00: +V17+V18*T
FCC
EXCESS MODEL IS REDLICH-KISTER MUGGIANU
                                                                     CONSTITUENTS:
A,B
      G(FCC,A;0)-G(BCC,A;0) = 500.00<T< 2000.00: 408
      G(FCC,B;0) - G(BCC,B;0) = 500.00 < T < 2000.00; +3300 - 3*T
      L(FCC, A, B; 0) = 500.00<T< 2000.00: +V19+V20*T
      L(FCC, A, B; 1) = 500.00<T< 2000.00: +V21+V22*T
===== BLOCK NUMBER 1
DEFINED CONSTANTS
    DX=2E-2, P0=101325, DH=500, DT=10
DEFINED FUNCTIONS AND VARIABLES%
    HTR=HM(LIQUID)-HM(A2B)
                                                   0.4183
Alternate equilibrium calculation
Alternate equilibrium calculation
                                                   0.1932
                                                   0.1016
Alternate equilibrium calculation
Alternate equilibrium calculation
                                                   1.4354E-03
Alternate equilibrium calculation
                                                   2.5063E-02
                                                   8.3929E-03
Alternate equilibrium calculation
Alternate equilibrium calculation
                                                   141.4
                                                   141.4
Alternate equilibrium calculation
Alternate equilibrium calculation
                                                    141.4
Alternate equilibrium calculation
                                                   141.4
Alternate equilibrium calculation
                                                   1.2532E-03
Alternate equilibrium calculation
                                                   5.4781E-04
Alternate equilibrium calculation
                                                   1.5404E-03
                                                   1.2702E-03
Alternate equilibrium calculation
ACR(B) = 9.4E - 1
                             0.9397
                                         2.85E-02 -2.7745E-04 -9.7472E-03
                                         2.82E-02 -4.9038E-04 -1.7396E-02
ACR (B) =8.4E-1
                            0.8395
ACR(B)=7.4E-1
                             0.7407
                                         2.81E-02 7.3804E-04 2.6305E-02
ACR(B)=6.4E-1
                             0.6424
                                         2.81E-02 2.3935E-03 8.5272E-02
ACR(B)=5.4E-1
                             0.5434
                                         2.82E-02 3.4449E-03 0.1220
ACR(B)=4.4E-1
                             0.4428
                                         2.85E-02 2.8265E-03 9.9024E-02
ACR(B)=3.4E-1
                             0.3394
                                         2.90E-02 -5.8174E-04 -2.0040E-02
                                         2.97E-02 2.0260E-03 6.8208E-02
ACR (B) = 2.3E-1
                             0.2320
                                         3.06E-02 -6.4192E-04 -2.0981E-02
ACR (B) =1.2E-1
                             0.1194
```

HMR(LIQUID)=-1964	-1957.	5.00E+02	7.439	1.4878E-02	
HMR(LIQUID)=-3500	-3478.	5.00E+02	21.67	4.3338E-02	
HMR(LIQUID)=-4588	-4565.	5.00E+02	22.69	4.5382E-02	
HMR(LIQUID)=-5239	-5217.	5.00E+02	21.50	4.3008E-02	
HMR(LIQUID)=-5454	-5435.	5.00E+02	19.11	3.8216E-02	
HMR(LIQUID)=-5233	-5217.	5.00E+02	15.50	3.1008E-02	
HMR(LIQUID)=-4575	-4565.	5.00E+02	9.691	1.9382E-02	
HMR(LIQUID)=-3481 (LIQUID)=-1950	-3478. -1957.	5.00E+02 5.00E+02 -6.	2.669 561	5.3385E-03 -1.3122E-02	118 HMR

## LIST\_STORE\_FILE



This is a PARROT module command.



## LIST\_SYMBOL\_IN\_GES



This is a PARROT module command.

Lists TP-function(s) for the entered model parameters for phases in the system on screen. In many cases, the optimizing variables are parts of the TP-functions which in turn are entered in model parameters for various phases in the GIBBS workspace.

This command is a way to find out how the functions depend on the optimizing variables. In the PARROT module the list is only shown on screen, not written to any file (which the LIST\_SYMBOL command in the GIBBS module does).

Syntax	LIST_SYMBOL_IN_GES
Prompt	Name: <symbol name=""></symbol>
	Specify a symbol name. Only those symbols that match this name are listed. Or press
Prompt	

## **OPTIMIZE\_VARIABLES**

Perform variable optimization. All system-definition data needed for the optimization is read from the current work file (\*.PAR). The result of the optimization is automatically stored in the current work file.

Syntax	OPTIMIZE_VARIABLES	
Prompt	Number of iteration /N/: <integral iteration="" number="" of=""> Specify an integral number for the optimization iteration. The specified iteration number is shown as the default number. The PARROT module tries this number of different sets of values of the optimizing variables unless it has converged earlier, or has given up earlier.</integral>	
	It initially takes a small step in each variable to find the steepest slope. You can give zero iteration to calculate the error in all selected experiments, and then use <u>"LIST_RESULT" on page 142</u> to check how good (bad) the current fit is.	

# **READ\_PARROT\_WORKSPACES**

This command is used in the PARROT module.

Read the previous PARROT/GIBBS/POLY3 workspaces back to replace the current PARROT/GIBBS/POLY3 workspaces. It is useful if late changes made through various PARROT/GIBBS/POLY commands are not satisfactory.

### Syntax READ\_PARROT\_WORKSPACES

>\_

The previous PARROT/GIBBS/POLY3 workspaces are always associated with the latest action to either open a work file by <u>"SET\_STORE\_FILE" on page 156</u>, or to create a work file by <u>"CREATE\_NEW\_STORE\_FILE" on page 133</u> (either interactively in the PARROT module or through a <u>"MACRO\_FILE\_OPEN" on page 5</u> of an \*SETUP.TCM file), or to update the work file by <u>"SAVE\_PARROT\_</u>WORKSPACES" on the next page.

Unlike the READ commands in other modules (e.g. GIBBS or PLOY), this command in the PARROT module does not ask for the file name where to read a previously opened/created/updated PARROT/GIBBS/POLY3 workspaces, for the reason described above.

You cannot use this command if there is no work file opened or created already.

## **RECOVER\_VARIABLES**

#### Syntax RECOVER\_VARIABLES

The values of all variables are set back to the start values.

## REINITIATE

This command should not be used unless you want to destroy the current PARROT workspace. However, this PARROT command does not reinitiate the GIBBS/POLY3 workspaces.

### Syntax REINITIATE

The workspace used by the PARROT program is reinitiated. All output and optimizing conditions are given their default values. All variables are set fixed with their value equal to zero.

## **RESCALE\_VARIABLES**

### Syntax RESCALE\_VARIABLES

The current values of all the parameters are copied to their start values and the scaling factors for further optimization. Thus it should be done now and again, in particular if you think the optimization results are improved after the previous run (s), or if any variable changes more than a factor of 10.

## SAVE\_PARROT\_WORKSPACES



This command is used in the PARROT module.

### Syntax SAVE\_PARROT\_WORKSPACE

If the latest changes made through various PARROT/GIBBS/POLY commands are as required, use this command to save the current workspace (i.e. data area) used by the PARROT program (also including the current GIBBS and POLY3 workspaces), onto the present work \*.PAR file which has already been opened by "SET\_STORE\_FILE" on page 156 or been created by "CREATE\_NEW\_STORE\_FILE" on page 133 (either interactively in the PARROT module or through a MACRO\_FILE\_OPEN of an \*SETUP.TCM file).

Syntax	SAVE_F	PARROT_WORKSPACE
	5	The PARROT/GIBBS/POLY3 workspaces are updated after each PARROT/GIBBS/POLY command. This means the current PARROT/GIBBS/POLY3 workspaces are always updated onto the latest work *.PAR file that is associated with the latest SET_STORE_FILE or CREATE_NEW_STORE_FILE command.
		Unlike the SAVE commands in other modules (e.g. GIBBS or PLOY), this command in the PARROT module does not ask for the file name where to save the current PARROT/GIBBS/POLY3 workspaces, for the reason described above. You cannot use this command if there is no work *.PAR file opened or created already.

## **SET\_ALTERNATE\_MODE**

Turn the alternate mode on or off.

With the ALTERNATE mode is possible to include functions to be evaluated together with an alternate calculation and to select ALTERNATE mode for each experimental equilibrium.

The alternate mode is described in <u>"Optimizing with the PARROT Module" on</u> page 124. It should be used only to optimize start values of the model parameters in the beginning of the assessment.

Syntax	SET_ALTERNATE_MODE
Duanat	On /Y/:
Prompt	By default alternate mode is ON.

## **SET\_EXTERNAL\_PARAMETER**



This is a PARROT module command.

Optimize model parameters of the so-called external models, which may not be as fullyimplemented/integrated parts inside the GIBBS system and thus are independently defined within user-specified/written source codes. This command makes links between the PARROT optimizing variables and external model parameters. In order to be able to build the specified external model and to use this SET\_ EXTERNAL\_PARAMETER command, it requires you to first write codes for a preferred external model (as it is not within the standard GIBBS system of the Thermo-Calc software) and then provide it to the consultancy team (consult@thermocalc.se) of Thermo-Calc Software Company in Stockholm for the purpose of including the external model in a separate DLL that is interactively connected to the Thermo-Calc software. Such a special service can/should only be provided under the restrictive condition that a specific Consultancy Project Contract is bilaterally agreed and signed in advance between users and Thermo-Calc Software.

```
Syntax SET_EXTERNAL_PARAMETER
```

8

External name: <Parameter name in the external model>

Prompt Correctly specify the parameter name defined within a specified/written external model.

As optimizing variable number: <n>

Give the number of the PARROT optimizing variable, which should have a unique link to the specified parameter name of the external model.

For example, @@ Link PARROT variables to CAM parameters:

```
SET-EXTER-PAR GT (SLAG_A, O-2, SI+4, CA+2) 1
SET-EXTER-PAR ET (SLAG_A, O-2, SI+4, CA+2) 2
SET-EXTER-PAR EXT (SLAG A, O-2, SI+4, CA+2) 3
```

## SET\_FIX\_VARIABLE

Prescribe a fixed value to a variable. The variable is considered as a constant at the optimization, usually after successful optimization runs for some specific variables (for some of the phases in the system). Such fixed variables can also be set back for further optimization run by using <u>"SET\_OPTIMIZING\_VARIABLE" on page 153</u>.

Syntax	SET_FIX_VARIABLE
	Variable number: <variable number(s)=""></variable>
Prompts	Specify the number of the variable. It is possible to give a range by giving two numbers with a hyphen in between (no spaces allowed), e.g. 1. The parameters within the limits are set fixed to their current values (no question of values).
	Start value /xxxx.xxxxx/: <a be="" fixed="" to="" value=""></a>

#### Syntax SET\_FIX\_VARIABLE

Specify a numerical value to be fixed for the selected variable (only when one variable number is specified at the previous prompt). The current value for the selected variable is shown as default.

If a range of variable numbers is specified at the previous prompt, this question is not prompted, and the current values are used as fixed values.

### **SET\_OPTIMIZING\_CONDITION**

Specify the conditions for the optimization. The default values are chosen automatically by the PARROT program, and in most cases these should not be changed.

The optimization stops when the sum of errors decreases by this value, even though it might be possible to increase it further with a new OPTIMIZE command.

Syntax	SET_OPTIMIZING_CONDITION			
	Relative standard deviation for experiments? /N/: By default the standard deviation of the experimental determinations are absolute values. Enter Y if these are used as relative weighting factors.			
	The estimated standard deviations of the optimized variables might be different for the two cases.			
	Min save on file? /Y/:			
	Specify whether the program should minimize the transfer of data to and from the current work file during the optimization:			
Prompt	<ul> <li>If the default (N) is kept, the workspaces are stored on the current work file after every iteration in the optimization procedure.</li> </ul>			
	• Enter Y to speed up the optimization. After optimization use the <u>"SAVE</u> <u>PARROT_WORKSPACES" on page 149</u> command to update the progress onto the current work file.			
	Relative step for calculation of derivatives: /1e-04/: <xxx></xxx>			
	In the calculation of the correlation matrix for equilibria with inaccuracy in the independent state variables, some numerical derivatives might have to be calculated.			
	Specify the relative step ( $xxx$ ) for the calculation of these derivatives.			

#### Syntax SET\_OPTIMIZING\_CONDITION

Maxfun (va05ad): /100/: <n>

The maximum number of iterations in the optimization. The same value set by the OPTIMIZE command.

```
Dmax (va05ad): /100/: <n>
```

An estimate of the maximum distance between the start and the final values of the variables. A smaller value makes the program vary the parameters with smaller factors.

```
H (va05ad): /1e-04/: <xxx>
```

The step used in the scaled variables for calculating numerical derivatives during the optimization.

Acc/(initial sum of squares) (va05ad): /.001/: <xxx>

The break condition for the optimization. The accepted value is the difference between the true minimum and the calculated one.

## **SET\_OPTIMIZING\_VARIABLE**

Specify which variable value should be estimated at the optimization. Such a variable can have a value of zero or another start value, or is fixed by <u>"SET\_FIX\_VARIABLE" on page 151</u> prior to the previous optimization run.

For a good reference prior to this command, a list of all variables (which are used to define various parameters for phases in the optimizing system, as pre-entered in the \*SETUP.TCM file or interactively entered in the GIBBS module) are obtained by using <u>"LIST\_ALL\_VARIABLES" on page 139</u>.

Syntax	SET_OPTIMIZING_VARIABLE		
	Variable number: <variable number(s)=""></variable>		
Prompts	Specify the number(s) for the variable(s) to be set. Any non-zero parameter within the range is allowed to be optimized. A parameter with a current value equal to zero must be specified explicitly here in order to be optimized. Specify a range by giving two numbers connected by a hyphen (no space allowed!), e.g. 2–8. Under this circumstance, no question of start values are asked.		
	<pre>Start value /xxxx.xxxx/: <a guess="" value=""></a></pre>		
	Specify a start guess of the optimum value (only when one variable number is		

### Syntax SET\_OPTIMIZING\_VARIABLE

specified at the previous prompt). The current numerical value for the selected variable is shown as default.

This guess is critical as the initial guess of all parameters must make it possible to calculate the selected equilibria. If a range of variable numbers is specified at the previous prompt, this question is not prompted, and the current values (as start values) are not changed.

### **SET\_OUTPUT\_LEVELS**

8¢8

Choose the type of information the PARROT module gives during the optimization procedure and when listing results.

Syntax	SET_OUTPUT_LEVELS		
	List increment /1/: <increment in="" iteration="" number=""></increment>		
Prompts	Specify the increment in iteration number for which information is listed on screen during the optimization. By giving a larger number it shortens the list.		
	List sum of squares: /Y/:		
	Choose whether the sum of squares is listed on screen during the optimization procedure.		
	List scaled variables: /Y/:		
	Choose whether the scaled variable values are listed on the terminal during the optimization procedure.		
	List weighted residuals: /N/:		
	Choose whether the weighted residuals are listed on the terminal during the optimization procedure.		
	List all parameters: /N/:		
	Choose whether all parameters in the models are listed in <u>"LIST_RESULT" on</u> page 142.		
	List correlation matrix: /N/:		
	Choose whether the correlation matrix of the variables are listed with the commands LIST_RESULT and <u>"LIST_ALL_VARIABLES" on page 139</u> .		

# **SET\_SCALED\_VARIABLE**

This command is similar to <u>"SET\_OPTIMIZING\_VARIABLE" on page 153</u> in that it specifies start values for optimizing variables. The difference being it is functional for only one optimizing variable at one time. It also prescribes a minimum and maximum value for the variable. During the optimization runs, the variable value is limited within this min-max range.

Such a variable may have a value of zero or another start value, or may be fixed by <u>"SET\_FIX\_VARIABLE" on page 151</u> prior to the previous optimization run. For a good reference prior to using this command, a list of all variables (which are used to define various parameters for phases in the optimizing system, as entered in the \*SETUP.TCM file or interactively entered in the GIBBS module) is obtained by using <u>"LIST\_ALL\_VARIABLES" on page 139</u>.

Syntax	SET_SCALED_VARIABLE
	Variable number: <variable number(s)=""></variable>
Prompts	Specify the number for an optimizing variable to be set. Any non-zero parameter is allowed to be optimized, and a parameter with a current value equal to zero must be specified explicitly here in order to be optimized.
	<pre>Start value /xxxx.xxxxx/: <a guess="" value=""></a></pre>
	Specify a start guess of the optimum value. The current numerical value for the selected variable is shown as default.
	This guess is critical as the initial guess of all parameters must make it possible to calculate the selected equilibria.
	Min value /xxxx.xxxxx/: <a guess="" value=""></a>
	Specify a minimum guess of the optimum value. It should be smaller than, at least equal to, the current start value for the selected variable.
	Max value /xxxx.xxxxx/: <a guess="" value=""></a>
	Specify a maximum guess of the optimum value. It should be larger than, at least equal to, the current start value for the selected variable.

## **SET\_STORE\_FILE**

Specify a store file (work file) to be used for compilation and optimization. The \*.PAR work file must be created (<u>"CREATE\_NEW\_STORE\_FILE" on page 133</u>) as a store file. The GIBBS, POLY and PARROT workspaces are read from the specified store file (with a default extension of PAR).

Syntax	SET_STORE_FILE
Prompt	Enter an *.PAR file name in the <b>File name</b> field and specify the working directory in the <b>Look in</b> field. Normally the default file-type in the <b>Files of type</b> field is the correct one for the PARROT workspace format (i.e. *.PAR file). Click <b>Open</b> or <b>Cancel</b> as applicable.

# **Equilibrium Calculations with the POLY\_3 Module**

### In this section:

About POLY3 - The Equilibrium Calculation Module	
ADD_INITIAL_EQUILIBRIUM	
About the ADVANCED_OPTIONS Command	
ADVANCED_OPTIONS	
AMEND_STORED_EQUILIBRIA	
CHANGE_STATUS	
COMPUTE_EQUILIBRIUM	
COMPUTE_TRANSITION	
CREATE_NEW_EQUILIBRIUM	
DEFINE_COMPONENTS	
DEFINE_DIAGRAM	
DEFINE_MATERIAL	
DELETE_INITIAL_EQUILIBRIUM	
DELETE_SYMBOL	
ENTER_SYMBOL	
EVALUATE_FUNCTIONS	
LIST_AXIS_VARIABLE	
LIST_CONDITIONS	
LIST_EQUILIBRIUM	
LIST_INITIAL_EQUILIBRIA	
LIST_STATUS	
LIST_SYMBOLS	
LOAD_INITIAL_EQUILIBRIUM	201
MAP	201

POST	203
READ_WORKSPACES	
RECOVER_START_VALUES	204
REINITIATE_MODULE	204
SAVE_WORKSPACES	204
SELECT_EQUILIBRIUM	206
SET_ALL_START_VALUES	206
SET_AXIS_VARIABLE	208
SET_CONDITION	
SET_INPUT_AMOUNTS	
SET_NUMERICAL_LIMITS	
SET_REFERENCE_STATE	214
SET_START_CONSTITUTION	216
SET_START_VALUE	
SHOW_VALUE	
STEP_WITH_OPTIONS	
TABULATE	

# **About POLY3 - The Equilibrium Calculation Module**

Knowledge of the thermodynamic equilibrium is an important factor for understanding properties of materials and processes. With a database of thermodynamic model parameters, it is possible to predict such properties and also to obtain driving forces for diffusion-controlled phase transformations and other dynamic processes.

With the Equilibrium Calculation module, POLY3, it is possible to calculate many different kinds of equilibria and diagrams, in particular multicomponent phase diagrams.

Different types of databases can be used with the POLY module, and it can be used for alloys or ceramic systems, as well as gaseous equilibria, aqueous solution involved heterogeneous interaction systems. Up to 40 elements and 1000 species can be defined into a single system for equilibrium calculations.

All normal thermodynamic state variables can be used to set as conditions in calculating equilibria, and as axes in plotting diagrams. You can set the composition or any property of an individual phase as a condition. Any state variable can be varied along an axis in order to generate a diagram. During calculations of a diagram, complete descriptions of all calculated equilibria are stored, and in the diagram any state variable can be used as an axis.

The Global Minimization technique is used to ensure that the present minimum in an equilibrium calculation is the most stable minima for the specified conditions. This is based on the traditional GEM (Gibbs Energy Minimization) technique.

A Direct Global Minimization can be performed on conditions: N, n(comp), B, b(comp), w (comp), x(comp), T, and P, but not when combined conditions as e.g. w(a)-3\*w(b)=1 are used or when an activity or potential condition is used. For all other types of conditions where regular minimization converges, Indirect Global Minimization, i.e. global test and corrections, if necessary, are performed until the lowest minimum is found.

- Direct Global Minimization: From the mesh of Gibbs energy, find the set of grid points that gives the lowest energy solution under the specified conditions. This set of grid points provides starting combination of phases and their constitutions for regular minimization to find the exact equilibrium solution. This solution is then subject to a global test as described below.
- Indirect Global Minimization: Under certain conditions, a direct approach is impossible. In this case, regular minimization is performed first and then a check is performed in order to see if the found local minimum is a global one by checking if all grid points are above the equilibrium Gibbs energy plane. If not, then recalculate by including these grid points until no grid point is above the equilibrium Gibbs energy plane from the previous step.

The full-scale and full-scope usage of the Global Minimization Technique has been extended from for only single-point calculations for all types of calculations (of single-points, property diagram stepping and phase diagram mapping) in Thermo-Calc.

The use of Global Minimization Technique may increase the computation time. The main cost in time comes from the calculation of Gibbs energy at each grid point generated by properly meshing the composition space for each entered phase. An additional (but much smaller) cost in time comes from finding the set of grid points in the above mesh that give the lowest energy solution. This solution is where POLY starts its ordinary minimization. When POLY has found an equilibrium, the equilibrium Gibbs energy surface is compared to the mesh to assure that no grid point is below the surface, i.e. a global minimization has been reached.

Global Minimization is performed by default in single-point or stepping or mapping equilibrium calculations.

Together with the PARROT module, the POLY module is also used for critical assessment of experimental data in order to develop thermodynamic databases. The POLY module uses the Gibbs Energy System (GES) for modeling and data manipulations of the thermodynamic properties of each phase.

## ADD\_INITIAL\_EQUILIBRIUM

Add initial equilibrium points from which a phase diagram is calculated (through the <u>"MAP"</u> on page 201 command).

The ADD command with the default direction scans along the axis variables and generates start points each time the scan procedure crosses a phase boundary. In addition, it generates four start points, scanning cross the middle of each axis, if there is any solubility line that does not reach the axes. At the MAP command, a search for lines in the diagram is made along each direction of the axis variables in the diagram.

In this way, it should guarantee that all possible phase boundary lines in a phase diagram are found. It may take a little longer to execute than using the minimum number of start points, as some lines may be calculated more than once. But the POLY module remembers all node points and subsequently stops calculations along a line when it finds a known node point.

It is also possible to create a sequence of start points from one initial equilibria by appending a > after the direction at the ADD command. For example:

```
Direction /Default/: 2>
Direction /Default/: -2>
```

This generates one start point for each set of phase change in the positive direction of the axis 2 (or negative direction of the axis 2); this ensures finding all possible phase boundary lines (not just the first one) along such an axis direction.

This is particularly useful when you have a phase diagram with several lines with no intersection. It is thus possible to calculate e.g. an isopleth for a much more limited composition range. It is also useful for calculating CVD diagrams and Pourbaix diagrams.

Syntax	ADD_INITIAL_EQUILIBRIUM
	Direction /Default/: <direction code=""></direction>
	Direction code(s):
	• 1 or 2 for positive direction of axis 1 or 2, respectively.
Dromot	<ul> <li>-1 or -2 for negative direction of axis 1 or 2, respectively.</li> </ul>
Prompt	• Default for all directions.
	The direction is important when the initial equilibrium point is in a single-phase region or when the diagram is an isopleth (tie-lines not in the plane of calculation). In such cases, the program will search for a line in the diagram (i.e., a line where the amount of a phase is zero) in the given direction.

Normally, this command is not needed to calculate a property diagram (with the "STEP\_ WITH\_OPTIONS" on page 217  $\rightarrow$  NORMAL command-sequence). In many cases, the ADD\_ INITIAL\_EQUILIBRIUM command is not required for MAP commands either, as the mapping procedure starts from the current equilibrium state already calculated. In order to calculate a simple phase diagram, set the equilibrium conditions and the mapping axis variables, and then give a MAP command. But if a phase diagram has disconnected lines, the ADD\_ INITIAL\_EQUILIBRIUM command may still be needed to add two or more initial equilibria so that the MAP calculation starts from such initial equilibria at the specified directions to find all phase boundary lines.

This command becomes unnecessary in most cases, as the MAP routines that by default use the Global Minimization Technique handles all the initial equilibrium points in a robust and automatic way. Therefore, you do not need to have a good guess of the starting point and to add any initial equilibrium point prior to the calculations of various types of phase diagrams. However, if preferred, use this command to add any initial equilibrium points in certain directions, the POLY module uses the specified starting point(s) and corresponding useradded initial equilibrium point(s) for a mapping calculation; in such a case, the automatic procedure in the rewritten MAP routines are not enforced while the Global Minimization Technique can still be applied. Similar situations occur when a specific database, in which there are some definitions of initial-equilibrium adding direction(s) for accessed binary and/or ternary subsystems in its ACCESSED\_SYSTEM section, is used in the BINARY or TERNARY module for calculating a binary or ternary phase diagram.

All the initial equilibrium points generated by the ADD\_INITIAL\_EQUILIBRIUM command (previously and presently; saved in the current POLY workspace) can be easily listed out on screen with <u>"LIST\_INITIAL\_EQUILIBRIA" on page 199</u>. A certain initial equilibrium point (including its conditions and equilibrium results) can be loaded into the current equilibrium, if needed, with <u>"LOAD\_INITIAL\_EQUILIBRIUM" on page 201</u>. Any specific or all of the initial equilibrium points can be deleted from current POLY workspace, if desired, with <u>"DELETE\_INITIAL\_EQUILIBRIUM" on page 194</u>.

## **About the ADVANCED\_OPTIONS Command**

"ADVANCED\_OPTIONS" on page 167

These ADVANCED\_OPTIONS can be set:

### EQUILIBRIUM\_CALCUL

Designed to decide how to perform the Ordinary POLY Minimization (i.e. the traditional Gibbs Energy Minimization) in the POLY module. By default, the Global Minimization Technique for various single point equilibrium calculations and for stepping or mapping calculations is used. You can use the advanced option <u>"GLOBAL\_MINIMIZATION" on page 292</u> to permanently turn off (for all sequential calculations throughout the current TCC run) the Global Minimization Technique and consequently use only the Ordinary POLY Minimization, and additionally adjust the ways to proceed the normal minimization (mainly in terms of how to control the steps in reaching Gibbs energy minima in an equilibrium state). When enforcing the Global Minimization Technique in the equilibrium calculations, you can further adjust the manners for assigning grid-points and for handling new possible compositional set(s) for solution phases during stepping/mapping; since the Global Minimization Technique also uses the normal POLY optimization routine for some calculations, it is also possible to change the ways on how to control the steps in reach Gibbs energy minima in an equilibrium state, in order to improve convergence.

#### **GLOBAL\_MINIMIZATION**

Designed to decide how to perform the so-called Global Minimization in the POLY module. By default, the POLY module uses the Global Minimization technique for various single-point equilibrium calculations and for stepping or mapping calculations. You can use this to permanently (for all sequential calculations throughout the current software run) turn off the Global Minimization technique and consequently use only the Ordinary POLY Minimization, and additionally adjust the ways to proceed the normal minimization (mainly in terms of how to control the steps in reaching Gibbs energy minima in an equilibrium state). When enforcing the Global Minimization technique in the equilibrium calculations, you can adjust the manners for assigning grid-points and for handling new possible compositional set(s) for solution phases during stepping/mapping; since the Global Minimization technique also uses the normal POLY optimization routine for some calculations, it is also possible to change the ways on how to control the steps in reach Gibbs energy minima in an equilibrium state, in order to improve convergence.

### LIST\_PHASE\_ADDITION

The values set as additional contributions (given by PHASE\_ADDITION) to Gibbs energy *G*m (J/mol formula unit) to all the phases (stoichiometric or solution) are listed at the current calculated equilibrium.

### MAJOR\_CONSTITUENTS

Use this to set the major constituent(s) of a composition sets in a miscibility gap of a solution phase. Normally, the major constituents are specified when a new composition set is created by the NEW\_COMPOSITION\_SET option; but for the first composition set, this option may be needed before using NEW\_COMPOSITION\_SET.

#### NEW\_COMPOSITION\_SET

A solution phase that can exist with two (or more) different compositions simultaneously must have two (or more) composition sets in its phase descriptions. Normally the database creates as many composition sets as is necessary but use this command to add or delete more composition sets. A complex solution phase in a defined multicomponent system can have up to 9 different composition sets for the purpose of appropriately handling its possible miscibility gap(s) under various temperature-pressure-composition conditions.

This option is unnecessary if the Global Minimization Technique is in use, as it can automatically detect all possible miscibility gap(s) for complex solution phases (normal or disordered/ordered phases) and then automatically add required composition sets for such phases in the defined system.

This option can automatically create composition sets also for disordered phase when it is executed for the ordered phase.

If you have your own data file, this option must be used to indicate that a solution phase can have a miscibility gap (or may exhibit some complex phase separations as to more than two composition sets), unless that the possibly-additional composition set(s) for the solution phase must have already been added inside the TDB file (through a TYPE\_DEFINITION command to amend composition-set) or been manipulated through the GIBBS module (using

the AMEND\_PHASE\_DESCRIPTION COMPOSITION\_SET command-sequence). This option asks for default major constituents for the new sets and it is important that this is set correctly, otherwise the test for miscibility gaps may fail.

#### OUTPUT\_FILE\_FOR\_SHOW

The name of a text file is asked for and all the results output from the command SHOW\_ VALUE is written to this file.

#### PARAEQUILIBRIUM

This calculates a paraequilibrium between two specific phases in an alloy system with one or more interstitial component(s) as fast diffusion species. Under the paraequilibrium state, two partially-equilibrated phases have the same chemical potential (but different contents) for one or more interstitial components (such as C, N, O, S, etc., as individual or combined)], along varied temperature or along a composition variable (of the matrix or one substitutional component) which has already set as the stepping variable with <u>"SET\_AXIS\_VARIABLE" on page 208</u>.

To ensure a successful point calculation of paraequilibrium state between two specific phases in a defined alloy system, it is important that you first have made a single-point equilibrium calculation with an initial overall composition in the current system before performing this advanced-option calculation; however, it is unnecessary to obtain an equilibrium in which either one or both of the target phases is stable. The initial overall composition must have a reasonable setting for the desired paraequilibrium calculation for the two target phases. This is especially true for cases where there are more than one interstitial components to be considered in the paraequilibrium state, because different interstitial components (for example C and N combined) may have significant different behaviors as partitioning into different structured phases; otherwise, for one chosen interstitial component the initial overall composition is OK for the paraequilibrium calculation between the specified two phases, but for other chosen interstitial component(s) it might be impossible to calculate the paraequilibrium state.

#### Note the following:

Always check if the chosen phases A and B have the exactly same definition of elements and if the chosen interstitial components are all in the vacancy sublattice sites of the two phases; otherwise the program cannot find the paraequilibrium state (as it is impossible to correctly calculate u-fractions).

Always have a comprehensive understanding of the normal phase diagram for the currently investigated system, so that you make the appropriate choice of the phase pair and staring bulk composition for the system.

Always set the status of the chosen interstitial components as SPECIAL using the POLY command: <u>"CHANGE\_STATUS" on page 177</u> Component <interstitial component> = SPECIAL. By doing this, you get a clear picture of u-fractions of various substitutional and interstitial components, which are different from the overall composition in the system. The SPECIAL status means that specified component(s) are not included in summations for mole or mass fractions. Therefore, all the composition variables plotted from paraequilibrium calculations are u-fraction related quantities.

#### PHASE\_ADDITION

Sometimes it is interesting to add a constant contribution to the Gibbs energy of a phase (stoichiometric or solution). This can be done in the Database Module (DATA) or Gibbs Energy Module (GIBBS), for a stoichiometric phase, or for a pure end-member in a solution phase. However, if the addition is related to the equilibrium state, for example, strain energies or surface energies, interfacial energies or deformation energies, it may be more convenient to have this quantity related to the equilibrium state rather than the thermodynamic data.

Give a value of an addition to the Gibbs energy of a phase. The value should always be constant (implying that the addition is not a function of phase composition or temperature-pressure conditions in the equilibrium state) and always be given in the unit of J/mol formula unit of the phase.

#### PRESENT\_PHASE

The phase specified must be stable at all equilibria calculated during a <u>"MAP" on page 201</u> command. It is a way to limit the calculations of the monovariant lines in a ternary system to those in the liquidus surface. Normally such a calculation would have two compositional axes and a temperature axis and all monovariant lines, also those between 3 solids, would be mapped. If the liquid is set as PRESENT, only those with the liquid is mapped.

### SHOW\_FOR\_T=

Display various thermodynamic properties (state variables, derived/partial variables or entered symbols) of the currently calculated (stable/meta-stable) equilibrium state but under a different temperature condition. This is useful for knowing, for example, a volume-related property of a frozen (stable/meta-stable) equilibrium state at a certain temperature, where the equilibrated phase assemblage and all the phase compositions are not adjusted while only the temperature condition is changed.

Use this option carefully. You must have successfully calculated a real equilibrium state under one temperature condition (normally the temperature for the last heat treatment). This can then be used to obtain the value(s) of any specified state variable(s) or

derived/partial variable(s) or defined symbol(s) for thermodynamic properties of the entire system, of components, or of phases for the currently-defined system (being in a frozen state) under another temperature (normally at room temperature). No real equilibrium is re-calculated through this option, and thus the phase amounts and compositions in the system are the same as at the last real equilibrium calculation.

#### STABILITY\_CHECK

In some composition ranges of a multicomponent system, it often happens that an unstable solution phase region locates inside a miscibility gap, and the stability limit (the *spinodal curve* or *spinodal*) may be not easy to find. A system inside a spinodal is thermodynamically unstable with respect to compositional fluctuations, and the system may experience the *spinodal decomposition* (i.e. decomposing to a mixture of regions with the two stable compositions, one on each side of the miscibility gap).

This option makes it possible to automatically check internal stability of both stable and unstable phases in all subsequent single-point equilibrium and MAP/STEP calculations. It can find out if any phase is subject to spinodal decomposition during the subsequent calculations. If there is an unstable phase located inside a miscibility gap in a calculation, it gives a warning so that you suspend the unstable phase, or use FORCED automatic start values for phase constituents (i.e. "SET\_ALL\_START\_VALUES" on page 206  $\rightarrow$  FORCE command-sequence), or create other composition sets; you can also ignore the warning message if you know that the unstable phase is not formed in the current calculations.

#### STEP\_AND\_MAP

This is the default and is used to determine how to perform Global Minimization test and how to handle initial equilibrium points.

### T-ZERO\_TEMPERATURE

This calculates the temperature when two specific phases have the same Gibbs energy, i.e. the so-called T0 temperature. You must calculate an equilibrium state at an estimated temperature before performing this advanced-option calculation; however, it is unnecessary to obtain an equilibrium state in which either one or both of the target phases is stable.

### TOGGLE\_ALTERNATE\_MODE

Toggle the ALTERNATE mode for experimental equilibrium calculation between DEFAULT, ALWAYS and NEVER, during data-assessments using the PARROT optimization.

### **ADVANCED\_OPTIONS**

Sy	yntax	ADVANCED_OPTIONS	
Р	rompt	Which option? /Step_and_Map/:	<option></option>

"About the ADVANCED\_OPTIONS Command" on page 162.



LIST PHASE ADDITION has no additional prompt.

#### STEP\_AND\_MAP (THE DEFAULT)

#### Settings for MAP and STEP:

Global test interval /0/: <Integer number>

#### Settings for MAP:

Automatically add initial equilibria /Y/: Number of mesh along an axis /3/: <Integer number> Use inside meshing points /N/: Global test interval /0/: <Integer number>

The integer number determines how often the Global Minimization should be used during STEP\_WITH\_OPTIONS and MAP calculations. If it is set to 0 (zero), the recommended global test interval is used: i.e. every tenth step and at each phase change during STEP calculations, and only at node points during MAP calculations. Any other positive integer number, n, suggests it performs a Global Minimization test at every *n*-th step during STEP and MAP calculations. Of course, the Global Minimization test is always carried out at a phase change or a node point. The Thermo-Calc software is installed with 0 (zero) as the default value, but such a configuration for the default value can be changed by using SET\_TC\_OPTIONS in the SYS module.

Automatically add initial equilibria /Y/:

When <u>"ADD\_INITIAL\_EQUILIBRIUM" on page 160</u> is not used before MAP, a mesh of initial equilibrium points are added before the mapping itself takes place. This mesh is only available when two (2) axes are already defined. The default on start-up may be changed by SET\_TC\_OPTIONS in the SYS monitor.

Number of mesh along an axis /3/: <Integer number>

An integer for how many intervals of initial equilibrium points to be added along an edge in the diagram. For example, if set to 2 (two), initial equilibrium points are added at the beginning, center, and end of each axis-variable forming 2 intervals.

Use inside meshing points /N/:

If N the INITIAL\_EQUILIBRIUM mesh consists of initial equilibrium points added only along the edge/border of the diagram defined by the axis-variables. If Y, the initial equilibrium points added are also added to the inside edge.

#### NONE

Take away any set advanced option in all subsequent single-point equilibrium and MAP/STEP calculations.

If there is any convergence problem in finding a stable solution, at any stage of an advanced option calculation conducted by an ADVANCED\_OPTIONS command-sequence, these messages display:

Convergence problems, increasing smallest site-fraction from 1.00E-30 to hardware precision 2.00E-14. You can restore using SET-NUMERICAL-LIMITS

The smallest site fraction in the current POLY3 workspace is automatically increased from the default value 1.00E-30 to the hardware-dependent precision (under Linux, as 2.00E-14). For other subsequent POLY-module calculation in the current TC run, you can use the POLY command <u>"SET\_NUMERICAL\_LIMITS" on page 293</u> to restore or reset the smallest site fraction to the previous or another preferred value, as well as to reset other numerical limits.

#### EQUILIBRIUM\_CALCULATION

The following prompts affect the ways how the POLY optimization does the Ordinary POLY Minimization to reach the minimum of an equilibrium state.

Settings for the minimization of an equilibria:

```
Force positive definite phase Hessian /Y/:
Control stepsize during minimization /Y/:
```

Choose Y or N to determine how to reach the minimum of an equilibrium state in a normal POLY optimization procedure. This is related to the special quantity, the phase stability function QF (phase), for all kinds of phases in an equilibrium state [A phase stability function for a phase is negative when the phase composition is inside a spinodal, and positive everywhere else].

If the composition of a solution phase is inside its spinodal, the enforcement of positive definite eigenvalues of a Hessian matrix (i.e. by answering Y) makes the step be taken towards the minima rather than the local maxima. Furthermore, if an eigenvalue (for a phase) of the Hessian matrix is near 0.0, the step-size is large and the answering Y sets a lower limit to this eigenvalue, and reduces the step-size and improves the convergence. The POLY command-sequence SHOW\_VALUE QF(phase) shows the size of the lowest eigenvalue of a phase in an equilibrium state.

By answering Y it might help the Ordinary POLY Minimization to converge, if there is such a problem, especially if QF(phase)=0 for phases.

By answering N the Ordinary POLY Minimization routine is applied.

For each phase in a defined system, the molar Gibbs energy of the phase is a function of the temperature-pressure condition and its composition:

$$G_m(X) = G_m(T,P;y_1,y_2,\ldots,y_n)$$

Compute a Hessian matrix as which describes the curvature of the Gibbs energy curve of this phase at the defined composition X:

### $\partial^2 G_m/\partial y_i\partial y_j$

Diagonalize this matrix and call the Eigenvalues as e1,e2,...,en . The QF(phase) quantity for this phase is the lowest of these eigenvalues divided by the same values for a corresponding ideal (stoichiometric) phase:

### $QF(ph)=\min(\{e_1,e_2,\ldots,e_n\})/\min(\{se_1,se_2,\ldots,se_n\})$

Here, se1,se2,...,sen are the eigenvalues for the ideal phase. Therefore, for an ideal (stoichiometric) phase, the QF(phase) should always be 1.0.

Control stepsize during minimization /Y/:

Choose Y or N to determine whether to introduce a control of step-size while reaching the minimum of an equilibrium state in a normal POLY optimization procedure. By answering Y it might help the POLY optimization to converge, especially if when some site fractions are less than 1E-4.

### **GLOBAL\_MINIMIZATION**

The following prompts affect how the Global Minimization is done. Settings for global minimization:

Use global minimization as much as possible /Y/:

```
Use global minimization for test only? /N/:
Maximum number of gridpoints per phase /2000/: <Integer number>
Use global minimization as much as possible /Y/:
```

Choose Y or N to decide if using the Global Minimization Technique from start of calculation or not. The default value on start-up may be changed by SET\_TC\_OPTIONS in the SYS module.

If Y the calculation is done when possible (depending on the condition settings), and a Global Minimization test is always performed when an equilibrium is reached. This costs more computer time but the calculations are more accurate.

If N the calculation is not used to reach the equilibrium state. Use global minimization for test only? determines if the Global Minimization test is made against the calculated equilibrium states obtained by the Ordinary POLY Minimization calculation.

Use global minimization for test only? /N/:

If Y a calculated equilibrium state obtained by the Ordinary POLY Minimization calculation is tested against the Global Minimization Technique after it is reached; and if found it as unstable, there is an error message.

If N the Global Minimization Technique is never tested for or done for all sequential calculations throughout the current TCC run. This implies that Global Minimization Technique is permanently turned off, and that consequently only the Ordinary POLY Minimization routine is used.

Maximum number of gridpoints per phase /2000/: <Integer number>

Specify the maximum number of grid points that are computed for each of the phases in the currently defined system, during the calculations enforced by the global minimization. More grid points give a higher robustness and take more computation time. The total number of computed grid points in all phases is limited to 2E6.

#### IGNORE\_COMPOSI\_SET\_ORDER

Ignore composition set order /N/

#### NEW\_COMPOSITION\_SET

The following prompts are to specify the additional composition set(s) to handle possible miscibility gap(s) of a certain solution phase:

Phase with miscibility gap: <Name of a phase>

Specify the name of the phase with miscibility gap.

New highest composition set number /2/: <#>

The default value is usually one higher than the current value. Each phase has initially one composition set. If a lower value is given, composition sets are deleted. You cannot take away the first composition set.

A message displays to show that you need to specify the composition for the composition set # (2,3,...).

Major constituent(s) for sublattice 1: /XX/: <YY>

The major constituents (YY) in each sublattice can be given. This may simplify giving start values when calculating the equilibrium as phases with miscibility gaps should have different major constituents for each composition set.

This prompt is repeated for each sublattice in the phase, sometimes even for all sublattices in the first composition set if such major constituents have not been specified.

#### MAJOR\_CONSTITUENT

The following prompts are to specify the major constituent(s) on each of the sublattice sites for a composition set of a certain solution phase:

Phase name: <name of a phase>

Specify the name of the solution phase with a new set of major constituents.

Composition set number /1/: <#>

The default value for the composition set number (#) is usually /1/ as the other composition sets are given major constituents when creating them. Each phase has initially one composition set.

Major constituent(s) for sublattice 1: /XX/: <YY>

The major constituents (YY) on each sublattice of a solution phase can be given. This may simplify giving start values when calculating the equilibrium as phases with miscibility gaps should have different major constituents for each composition set.

This question is repeated for each sublattice in the phase.

#### PHASE\_ADDITION

The following prompts are to specify the additional energy term (always being a constant) of a given phase.

Phase name: <Name of a phase>

Specify the name of the (stoichiometric or solution) phase with the addition.

Addition to G per mole formula unit: <xxxxx>

The value (xxxxx) given is added to the Gibbs energy of the (stoichiometric or solution) phase. It can represent a nucleation barrier, surface tension, elastic energy or whatsoever.



It is not composition-, temperature- or pressure-dependent.

#### PRESENT\_PHASE

Specify the name of the present phase a the name of the phase that should be present at all calculated equilibria:

```
Present phase:
Phase name: <Name of a phase>
```



Console Mode examples tcex45, tcex46 and tcex47 use this advanced option. See <u>"About the Thermo-Calc Console Mode Examples</u>" on page 2 in the *Console Examples Guide*.

#### **T-ZERO\_TEMPERATURE**

```
Name of first phase: <Phase A>
Name of second phase: <Phase B>
```

The names of phases A and B must be given, for which the T0 temperature (where the Gibbs energies are equal) is to be calculated.

If the T0 temperature between the two specified phases is successfully calculated, a message displays, e.g.

The TO temperature is 840.82 K. Note: LIST-EQUILIBRIUM is not relevant

The first message shows the calculated T0 temperature between the two specified phases.

The second message indicates that after this option calculation with LIST\_EQUILIBRIUM is irrelevant and does not list the equilibrium for the system at the TO-temperature.

#### PARAEQUILIBRIUM

```
Name of first phase: <Phase A>
Name of second phase: <Phase B>
```

The names of the two target phases A and B, between which the paraequilibrium state is to be calculated, must be entered subsequently or on the same (first) line at once then separated by an empty space, e.g. FCC#1 BCC or FCC#2 M23C6.

You need to understand what you are dealing with in terms of calculating a paraequilibrium state between the two specified phases.

Specifically, there are four distinguished cases to understand: (1) both chosen phases must have similar interstitial/vacancy sublattices where the fast-diffusion interstitial component (s) occupy; (2) the choice on the target phase pair must be reasonable for the defined system and specified initial overall composition; (3) both target phases should have phase constitution definitions that cover all the defined substitutional and interstitial components of the current alloy system; or (4) it is impossible to calculate the paraequilibrium state between the target phase pairs with given interstitial component(s) in the currently defined system.

```
Fast diffusing component: /c/: <interstitial component(s)>
Fast diffusing component: /none/: <interstitial component(s)>
```

The name(s) of the fast-diffusing component(s) (C as the default single component) must be given at the above prompts subsequently or at the same (first) prompt. It is possible to specify more than one interstitial component as fast diffusion species.

Note the following:

Such specified interstitial component(s) must be appropriately defined according to the phase constitution definitions of the two selected phases: these must be located on the interstitial/vacancy sublattices in both chosen phases;

If there is only one fast-diffusing component which is carbon, press the <Enter> key to accept the default input at the first prompt; if the single fast-diffusing component is another element (e.g. N), type its name at the first prompt;

If there are two or more fast-diffusing components (e.g. c and N), type their names at the above prompts subsequently or at the same (first) prompt (separated by an empty space, such as c N);

To finish the input of fast-diffusing elements, accept NONE at a repeated prompt, i.e. by pressing <Enter> key to start the paraequilibrium point calculation;

If NONE or a non-existing component name is typed at the first prompt, it means no back diffusion is to be considered, and the para-equilibrium calculation is thus canceled entirely.

If the paraequilibrium state between the two specified phases is successfully calculated, the messages displays e.g.

```
NP(FCC) = 0.3586 with U-fractions C = 2.71821E-02 N = 4.1548129E-03
```

```
NP(BCC) = 0.6414 with U-fractions C = 7.10061E-04 N = 2.3781027E-04
All other compositions the same in both phases
Note: LIST-EQUILIBRIUM is not relevant
```

The first and second lines list the phase amounts expressed in mole-percent [NP(phase)] and the contents of the interstitial components C and N in a specific phase expressed in the socalled u-fractions [u-f(phase,C) and u-f(phase,N)], for the phase A (in this case as FCC) and phase B (in this case as BCC), respectively. The third line states that the compositions of the matrix component and all the remaining compositions (regarding substitutional components) in both the target phase A and target phase B are the same at the current paraequilibrium state, while these are not shown on screen. The last line indicates that after this advanced-option calculation the LIST\_EQUILIBRIUM command is irrelevant and does not list the paraequilibrium state for the system at the current condition.

However, if the single-point calculation of the paraequilibrium state between the two specified phases has failed, these messages display:

```
*** Error 4 in ns01ad
*** Numerical error
```

This implies that the chosen target phase pair may be unreasonable for the defined alloy system or for the defined initial overall composition, or one or both phases may have inappropriate phase constitution definitions regarding the specified interstitial component (s). Then, you must either modify the settings of initial overall composition or specify the reasonable target phase pair with an appropriate choice of the fast diffusion interstitials in the defined alloy system.

#### STABILITY\_CHECK

```
Stability check on? /Y/:
Check also for unstable phases? /Y/:
```

If the automatic stability-check option is switched on choose to also check the stability for unstable phases. The default is Y and if an unstable phase is found to be located in a miscibility gas during a subsequent single-point equilibrium or MAP/STEP calculation, a warning message informs you to selectively make adjustments in the calculation settings [e.g. suspending the unstable phase, or using FORCED automatic start values for phase constituents by the S\_A\_S\_V F command-sequence, or creating other composition sets, etc.]. By answering N the stability check is enforced only to stable phases in the system.

#### TOGGLE\_ALTERNATE\_MODE

```
Set alternate toggle to DEFAULT, ALWAYS, or NEVER?
Set global toggle to DEFAULT, ALWAYS, or NEVER?
```

- DEFAULT: the experiment points are calculated according to the Alternate Technique depending on the SET\_ALTERNATE command.
- ALWAYS: the experiment points are always calculated according to the Alternate Technique even if the Alternate Mode is switched off in the PARROT module.
- NEVER: the experiment points are calculated as normal equilibria even if you SET\_ ALTERNATE in the PARROT module.

### SHOW\_FOR\_T=

Temperature (K) /298.15/: <temperature in K>

Specify the new temperature condition (in K) under which the values of some specific state variable(s) or derived/partial variable(s) or defined symbol(s) for various thermodynamic properties of the entire system, of components, or of phases) in the currently-defined system (being in a *frozen* state) is shown on screen.

State Variable or symbol /VM/: <State variable or symbol name(s)>

Specify the name(s) of the desired state variable(s) or derived/partial variable(s) or defined symbol(s) for various thermodynamic properties of the entire system, of components, or of phases) in the currently-defined system. More than one state variable or symbol of interest can be simultaneously specified on the same line. For example, you can choose to show the values of VM or GM (i.e. molar volume or molar Gibbs energy of the entire system), or of VM(\*) or HM(\*).T (i.e. molar volumes or isobaric heat capacity of all phases), or of ACR(\*), DGM(\*) and LNACR(\*,FCC) [i.e. activities of all system components, driving forces for all phases, and activities (in logarithm) of all system components in the FCC solution phase], under the new temperature condition (being in a frozen state).

#### OUTPUT\_FILE\_FOR\_SHOW

A window opens with a default name for the file tc\_show.dat. Choose a location to save the file and click **Open**.

### AMEND\_STORED\_EQUILIBRIA

This command gives information about the calculated blocks (and phase regions included in blocks) after the STEP or MAP calculation(s). It allows you to list all or part of the calculation results, to suspend all or parts of the calculation results that are redundant or where metastable equilibria are calculated, and to restore all or parts of the calculation results (if having been suspended by another AMEND\_STORED\_EQUILIBRIA command).

The workspace for storing equilibria may overflow during stepping or mapping, and is then written to a file as blocks. Each block usually contains one or more ranges of equilibrium regions.

Use one these options:

- L to list the calculated equilibria (all or a specified block)
- s to suspend everything (all blocks and their regions)
- Q to suspend each set of equilibria individually (specified blocks and/or regions)
- R to restore everything (all blocks and their regions)

Syntax	AMEND_STORED_EQUILIBRIA
	Name: <name a="" defined="" of="" table=""></name>
Prompts	List (L) the calculated equilibria, suspend everything (S) or suspend each set of equilibria individually (Q), or restore everything (R).
	Block /*/: <block number=""></block>
	Specify a block number in the option $L$ , $S$ , $Q$ or $R$ . Or include all the blocks in the amending option, by accepting the wildcard $*$ (press <enter>).</enter>
	For $L$ , a block (if a block number is specified) or all the blocks (if the wildcard $*$ is accepted/used), with the regions and the equilibrium details, that are calculated during stepping or mapping is listed on screen or in a textual file.
	For $Q$ , each ranger in a block (if a block number is specified) or in all blocks (if the wildcard * is accepted/used) then choose:
	S(uspend) K(eep) /K/: <s k="" or=""></s>
	For s or R, if a block is specified, you are asked what region in the block should be suspended or restored. However, if the wildcard $*$ is accepted/used, you are asked the following:
	Really suspend all /N/:
	Really restore all /N/:
	In both cases the default is N and you are prompted for which block(s) and which region(s) in a certain block to execute the $s$ or $r$ action.
	Really suspend all /N/:
	Y suspends everything.
	Really restore all /N/:
	Y restores everything.
	Range: <range(s) of="" region=""></range(s)>
	Specify one or more ranges to be suspended (the ${\rm s}$ option) or restored (the ${\rm R}$

#### Syntax AMEND\_STORED\_EQUILIBRIA

option). The wildcard \* suspends or restores all ranges in the specified block.

In order to know the ranges the LIST command must first be used.

S(uspend) K(eep) /K/: <S or K>

The Query suspend option (Q) asks for each region in a certain block if it should be suspended or kept. Suspended regions are not included on sequentially generated plots.

```
Output file: /Screen/: <File name>
```

This prompt is only for listing (the L option), and after it the command is terminated. The file name must be given here; or accept the default Screen (terminal) by pressing <Enter>. A list of stored equilibria as various blocks (with all their ranges) are shown out on screen or the file.

## CHANGE\_STATUS

In the POLY module, set the status for components, species and phases in the defined system for all the sequential calculations (single-point, stepping and mapping) in equilibrium or local/partial equilibrium state. Each component, species and phase has a status. The default status is ENTERED.

The most important use is to calculate metastable equilibria and metastable phase diagrams by setting some phases (that would otherwise be stable) to the SUSPENDED or DORMANT phase-status. Another important applications is to calculate paraequilibria by setting some components to the SPECIAL component-status.

For a component and for a species, the status can be one of the following:

- ENTERED: the component(s) or species are included in the calculation. This is the default status.
- SUSPENDED: the component(s) or species are not considered in the calculation.
- SPECIAL: the specified component(s) are not included in summations for mole or mass fractions. It only works for component(s).



Only component(s) can have the status SPECIAL, which implies that these are not included in summations for mole or mass fractions.

For example, for the *u*-fractions or other normalized fractions, when one or more of the components are excluded from the summation, you must specify which component(s) should be excluded from the calculation of mole or mass fraction. This component status is

particularly useful when calculating paraequilibrium states. Such component(s) are normally interstitial component, and must have the status SPECIAL. This is assigned by the CHANGE\_STATUS command.

Syntax	CHANGE_STATUS
Dromoto	For phases, species or components? /Phases/: <keyword></keyword>
Prompts	Keyword = phase or species or components
	<pre>Phase name(s): <name(s) of="" phase(s)="" the=""></name(s)></pre>
	For phase as the keyword, the names of the phases that have their status changed must be given (all on one line). A comma or space must be used as separator. The status to be assigned to the phases can also be given on the same line if preceded with an equal sign (=).
	An asterisk, *, can be used to denote all phases. The special notations *s, i.e. a wildcard * directly followed by an s, sign, means all suspended phases. In the same way, *D means all dormant phases, and *E means all entered phases.
	<pre>Name(s): <name(s) component(s)="" of="" or="" species="" the=""></name(s)></pre>
	For species or components as the keyword, the names of the species or components that have their status changed must be given (all on one line). A comma or space must be used as separator. Similarly to the case of phase as the keyword, the status to be assigned to the species or components can also be given on the same line if preceded with an equal sign =.
	An asterisk, *, can be used to denote all species or components. The special notations *s, i.e. a wildcard * directly followed by an s, sign, means all suspended species or components. In the same way, *E means all entered species or components.
	Status /Entered/: <new status=""></new>
	The new status to be assigned must be given.
	For species, the values ENTERED or SUSPENDED can be used.
	For components, the status ENTERED, SUSPENDED or SPECIAL can be given. SPECIAL means that this component is excluded from sums for mole fractions and mass fractions, which is useful when calculating the <i>ufractions</i> or other normalized fractions of system components.
	For phases, the status entered, suspended, dormant or fixed can be given.

Syntax	CHANGE_STATUS
	DORMANT means the same as suspended but the driving force is calculated. FIXED means that it is a condition that the phase is stable at a certain amount.
	For example, for the ufractions, when one or more of the components are excluded from the summation, you must specify which component should be excluded from the calculation of the mole fraction. This component must have the status SPECIAL. This is assigned by the CHANGE_STATUS command: Change_Status comp C=special.
	Start value, number of mole formula units /0/: <initial amount=""></initial>
	For ENTERED phases, an initial amount of the phase can be given. Normally, 0 is given if the phase is not likely to be stable, and 0.5 or 1 or any positive number if the phase could be stable, but such an initial amount is only used as the rough starting estimation in the equilibrium calculations.
	Number of mole formula units /0/: <equilibrium amount=""></equilibrium>
	For FIXED phases, the equilibrium amount of the phase [always using an initial estimation being the NPF (phase) value which it is the normalized mole number of components (per mole formula unit) of the specific status-fixed phase] must be given. If the equilibrium amount is zero, then the phase is at its stability limit.
	Note when specifying a FIXED phase status
	Special attention should be paid when specifying a FIXED phase status in equilibrium calculations (for single points, stepping or mapping calculations), as described below.
	The phase amount variables, NP(phase), BP(phase) and VP(phase), as well as all their M/W/V-suffixed quantities, should not be used as conditions. Instead, use the CHANGE_STATUS command to set a relevant condition, e.g. CHANGE_STATUS phase <pre>sphase&gt;=fix <amount> where the fixed <amount> is roughly the same as the F-suffixed quantity NPF(phase).</amount></amount></pre>
	The NPF (phase) quantity is the normalized mole number of components (per mole formula unit) of the specific phase in the defined system, which unlike other F-suffixed state variables [e.g. GF(phase), HF(phase) and DGF(phase)] cannot be directly applied in any POLY command, implying that it cannot be directly evaluated or listed/shown. If intended to shown such a normalized phase amount value in an equilibrium state, you should use a properly-entered symbol (function or variable), for example: NPFabc = NP(abc) /NA or NPFabc = NPM(abc) /NA*N. N is the total system size (in mole). The NA value is a quantity that is phase-dependent (and sometimes also equilibrium-dependent for ionic

# SyntaxCHANGE\_STATUSsolution phases), and is the total atomic number in a mole-formula-unit of the<br/>specific phase abc (excluding interstitial component and, of course, vacancy).<br/>For example, the SIGMA, FCC, BCC and LIQUID phases (among others) in a<br/>defined Fe-Cr-Ni-C-N-O system (retrieved from a specific database) may be<br/>modeled by certain models, and their NA values must be evaluated in different<br/>ways, as described below:<br/>LIQUID (C,Cr,Cr03/2,Fe,Fe0,FE03/2,N,Ni,NiO)1 NA = 1<br/>FCC\_A1 (Cr,Fe,Ni)1(Va,C,N,O)1 NA = 1<br/>BCC\_A2 (Cr,Fe,Ni)1(Va,C,N,O)3 NA = 1<br/>SIGMA (Fe,Ni)8(Cr)4(Cr,Fe,Ni)18 NA = 30If in the same Fe-Cr-Ni-C-N-O system the liquid solution phase is modeled by the<br/>Two-Sublattice lonic Liquid Model, i.e.:<br/>IONIC\_LIQ (Cr+3,Fe+2,Ni+2)p(VA,C,N,O-2,FE03/2)q,7

then the evaluation of its NA value becomes even more complicated:

NA = p + q\*yC2 + q\*yN2 + q\*yO262 + q\*yFeO2 3/ 2

where the stoichiometric coefficients *p* and *q* are also dependent upon the real equilibrium state (rather than having fixed values in the system). Similar situations occur for other (solid) phases which are described by a multiple sublattice model with ionic constituents, such as SPINEL and HALITE phases in some databases.

There is no strange thing when using a zero value [i.e. 0] in a FIXED phasestatus, since it means the specified phase is stable in equilibrium state but has a zero-amount of mass in the equilibrium calculations; in other words, on a phase diagram, the specific phase is on a zero-fraction line (ZFL), i.e. it starts becoming stable on one side of a corresponding phase-boundary line or unstable on the other side of the same boundary. It is often and efficient to do so when calculating e.g. solidus equilibrium states.

However, when a non-zero value [it must always be positive; e.g. 1 or 0.5 or 0.3 or 1.5] is to be specified in a FIXED phase-status, it is unnecessarily the exactly same stable amount of the specific FIXED-status phase in a calculated equilibrium state any longer; instead, the <equilibrium amount> value is the NPF(phase) value that is only roughly used as the estimated starting-value of the FIXED-status phase in the equilibrium calculations.

Therefore, a FIXED-status for a liquid phase being unity does not necessarily imply that it is a liquidus equilibrium state (where the liquid phase is in

SyntaxCHANGE\_STATUSequilibrium with some solid phases but the liquid phase takes all the mass in the  
defined system). A unity value for setting the liquid phase status in calculating  
liquidus equilibrium state can only be used when the liquid mixture phase is  
predefined as a single-sublattice solution phase (such as metallic liquid phase in  
multicomponent alloy systems) and the total system size as one mole (i.e. N=1).When a phase is described by a solution model in which two or more sublattices  
are considered and these sublattice sites may also have different stoichiometric  
coefficients [meaning that the mixture phase could have more than one atom in  
formula [NA>1; see some examples above], the unity value should not be used  
when setting the FIXED status for the phase; instead, you should use an  
appropriate value that ranges from 0 to a NPF(phase) value that equals to or is  
smaller than 1/NA (if the total system size N=1) or 1/NA\*N (if N differs from  
unity). For this reason, if a multicomponent system bears an IONIC\_LIQUID  
phase that is described by the Two-Sublattice lonic Liquid Model (or any other  
multiple-sublattice ionic solution phases), it is difficult to use a proper NPF (ION\_  
LIQ) value in setting its FIXED phase-status, because that should be less than (or  
equal to) the complex value of  
N/[
$$p + q^{*y}C2 + q^{*y}_{D}22 + q^*_{Y_{FeO}}2$$
 3/ 2].

#### EXAMPLES

For example, to obtain the metallic fraction in a system with carbon as an interstitial component, you can set the component status for carbon as SPECIAL:

Change\_Status comp C=special



The **SUSPENDED** status for components and species does not always work as expected.

For a phase, it may have one of these statues:

• ENTERED: the phase(s) are included in the equilibrium calculations and these are stable if that minimizes the total Gibbs energy in the defined system. This is the default status for all phases already retrieved from the chosen database(s). An ENTERED phase-status is always associated with an initially-estimated amount [in mole number; normally, as 0 if the phase is not likely to be stable, and as 0.5 or 1 or any positive number if the phase could be stable] but it is only used as the rough starting value in the equilibrium calculations.

- SUSPENDED: the phase(s) are not considered in the equilibrium calculations.
- DORMANT: the phase(s) are not considered in the equilibrium calculations but their driving forces for precipitation are calculated.
- FIXED: it is an equilibrium condition that the status-fixed phase must be stable, and be in equilibrium at a specified amount [always using an initial estimation being the NPF (phase) value which it is the normalized mole number of components (per mole formula unit) of the specific status-fixed phase]. See more descriptions at the end of this command.

#### **COMPUTE\_EQUILIBRIUM**

The full equilibrium state is calculated for the given set of conditions. The Global Minimization Technique is by default enforced in this command (C\_E), while it can be disabled temporarily (for the current single-point equilibrium calculation) if using C\_E – or C\_E \* command-combination, or permanently (for all the sub sequential single-point calculations or stepping/mapping calculations within the current TCC run) if having decisively switched it off by a user (or possibly in some special modules) through changing the minimization option using the "ADVANCED\_OPTIONS" on page 167  $\rightarrow$  MINIMIZATION\_OPTION command-sequence.

Syntax	COMPUTE_EQUILIBRIUM
Prompt	The C_E command can be used, in order to enforce the ordinary POLY minimization routines in an equilibrium calculation; this is because of that the ordinary C_E command is now associated with the Global Minimization Technique, and only after the Global Minimization
	If technique is permanently switched off the C_E command makes no difference from the C_E – command-combination.
	Only certain types of equilibrium conditions [e.g. T, P, N, N( <component>), X (<component>), B, B(<component>), and W(<component>)] are fully supported in the Global Minimization mode (called <i>Direct Global Minimization</i>); and when other types of equilibrium conditions are used, after the initial POLY, a Global Minimization test and corrections are performed until the lowest minimum is found (called <i>Indirect Global Minimization</i>).</component></component></component></component>
	If there is any problem with convergence, you may try the C_E * command- combination. The character * enforces the command to use an advanced technique to obtain a complex equilibrium. However, after a successful C_E *calculation, you may repeat the C_E command and can check the status of phases, species or components (with <u>"LIST_STATUS" on page 200</u> $\rightarrow$ CPS command-sequence) and equilibrium conditions (with <u>"LIST_CONDITIONS" on</u>

Syntax	COMPUTE_EQUILIBRIUM
	page 198) and list out the calculation results (with "LIST_EQUILIBRIUM" on page 198), because such actions may tell you how to further modify various settings for your current calculation. This command-combination is not that useful anymore, because the Global Minimization Technique that is always associated with the C_E command is even more powerful and more precise in finding the most-stable equilibrium state in a complex heterogeneous interaction system; therefore, the C_E * command-combination is functional and can be used only after the Global Minimization mode has already been disabled temporarily or permanently.
	Some phases that are not stable in the current equilibrium state may not have their most favorable composition after this command, and thus their driving forces may not be correct. You can force the program to correctly calculate the driving forces of metastable phases, by giving repeated C_E commands until the number of iterations (that is shown on screen after this command) is reduced to 2.
	Also see the POLY command <u>"SET_NUMERICAL_LIMITS" on page 293</u> which can set the approximate driving force for metastable phases option on or off in all the subsequent POLY calculations within the current TCC run.
	If an equilibrium state for the defined system is not found, an error message is given.
	You can try repeating this command a few times, or change some of settings for the numerical limits, for starting variables and starting values, for starting constitutions of certain phases and for reference states of certain components, or to verify some of the defined conditions.

#### **COMPUTE\_TRANSITION**

This command is a combination of the CHANGE\_STATUS, SET\_CONDITION and COMPUTE\_ EQUILIBRIUM commands. It allows a direct calculation when a new phase may form by varying one of the already-set conditions. It can be used only after at least one equilibrium is calculated successfully; otherwise, you are informed on the necessity of first making an equilibrium calculation to find out the stable phases under the current conditions.

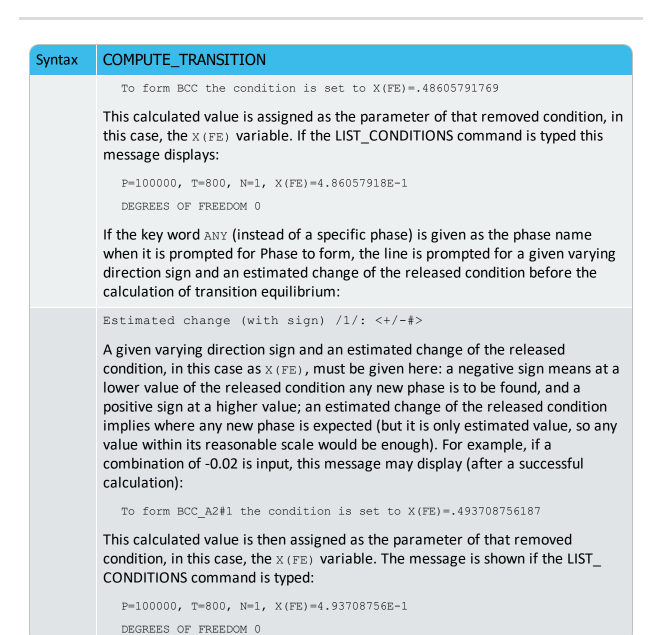
When this command is used, the program calls the command <u>"CHANGE\_STATUS" on page 96</u> to temporarily change the phase status of a specified phase as FIXED at the *zero* amount, and at the same time to temporarily release one of the existing equilibrium conditions

(which is chosen by you). The program calculates a new equilibrium in which that specific phase is stable but its equilibrium amount in the system is zero. The released condition is then assigned with a calculated value that ensures the calculated equilibrium. After wards, the program automatically changes the phase status of that specific phase back to ENTERED, and resets the temporarily released condition as one of the conditions and assigns it with the value that is calculated to ensure the zero-amount formation of that specific phase.

This command is useful to find melting temperature, boiling temperature, or solubility limits, and generally when you want to set the most optimal conditions for calculating an equilibrium where a specific phase becomes stable. It can also be used when you want to know exactly how far away the defined conditions are from the value that can ensure a zeroamount of a specific phase in the system when other conditions remain the same.

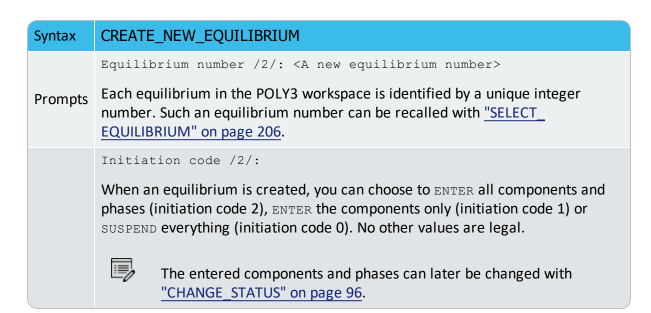
After a successful COMPUTE\_TRANSITION calculation, you can issue a <u>"COMPUTE\_EQUILIBRIUM" on page 182</u> calculation to assure the calculated transitional equilibrium is a really stable one, and can also use <u>"LIST\_EQUILIBRIUM" on page 198</u> to see the details of transitional equilibrium state.

Syntax	COMPUTE_TRANSITION
	Phase to form: <phase name=""></phase>
	A new phase name, e.g. BCC that is expected to form, is specified here. This changes the status of this new phase to be FIXED as 0 amount, and the program shows the information such as:
	You must remove one of the these conditions
	P=100000, T=800, N=1, X(FE)=.5 DEGREE OF FREEDOM 0
Prompts	If the key word ANY is used when prompted for Phase to form (instead of a specific phase name), it is possible to find out any new phase to be formed, in a given varying direction sign and at an estimated change of the released condition: a negative sign means at a lower value of the released condition any new phase is to be found, and a positive sign at a higher value; an estimated change of the released condition implies where any new phase is expected (but it is only estimated value, so any value within its reasonable scale would be enough). Such calculations can be repeated if required. This feature is useful to find out all possible phase transformations along a certain released condition.
	Give the state variable to be removed /T/: <one condition=""></one>
	One condition must be removed, in order to calculate the transition equilibrium where the specified (or any) new phase to be formed at a calculated value of this released variable.
	Therefore, the message may display (after a successful calculation) if, for example, $x (Fe)$ is entered:



#### **CREATE\_NEW\_EQUILIBRIUM**

During data-assessments using the PARROT/ED\_EXP modules, you can, in the POLY module, create several equilibria with different sets of conditions and phases (but normally with the same set of components). By default, there is one equilibrium. To keep the set of conditions and phase for this equilibrium, create another one using this command, and use another set of conditions for that. Two equilibria may be useful to calculate easily the enthalpy difference between two states. In the PARROT module, the experimental information is stored as a sequence of equilibria.



# **DEFINE\_COMPONENTS**

Change the set of components. By default, the elements are used as components. The set of components can be important because some conditions are set using components, for example, the amounts, activities or chemical potentials.

For example, in the system Fe-Si-O, you can define FEO, FE2O3 and SIO2 as components, thus replacing the default FE, SI and O.

This implies a command <u>"REINITIATE\_MODULE" on page 204</u> and it should be given as the first command in the POLY module.

Syntax	DEFINE_COMPONENTS
	Give all new components / existing components/: <new components=""></new>
Prompt	The new components must be specified all on one line. These replace the existing component definitions.
	The number of components cannot be changed with this command. Use <u>"CHANGE_STATUS" on page 96</u> instead.
	To keep some existing components, it is recommended that you also enter these on the line. Otherwise, the added new components may not be defined correctly. This is especially important if some components are built out of several elements.
	The new components do not have to be present as species.

# **DEFINE\_DIAGRAM**

This allows automatic calculation and plotting of a diagram with a single command. It is the same as the DEFINE\_MATERIAL command up to when the first equilibrium is calculated. The alloy OPTION feature is also available in this command to specify alloying compositions for a special alloy predefined by the OPTION keyword in a selected database (e.g. the TCNI Nibased Superalloys Database).

Use this command to calculate all types of phase diagrams after specifying all composition value and an initial temperature (if temperature is used as an axis). However, for binary and ternary diagrams, the special BIN and TERN modules may be preferred.

It then lists all the independent variables for the defined system (i.e. temperature and the components) and asks for a variable as the X-axis. You must also specify a maximum and minimum for the X-axis. The second axis (Y-axis) can be another composition (or the temperature if that is not on the X-axis) from the independent variable list. The program then calculates and plots a *phase diagram*, as there are two independent quantities on the axes.

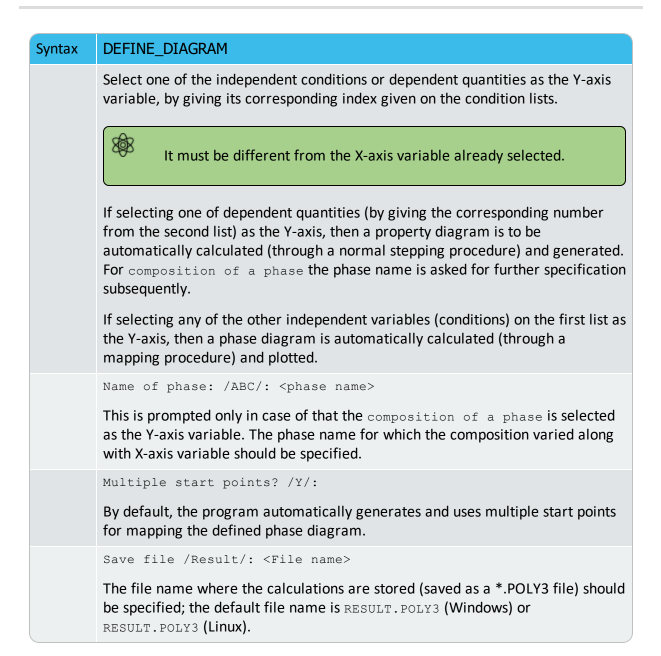
Alternatively, select a dependent quantity as the Y-axis variable from the second list on screen (e.g. the amount of all phases, composition of a specific phase, or fractions of a component in all phases), and the program calculates and plots how this quantity depends on the condition on the X-axis. This is a *property diagram*.

This command ends up within the POST module monitor. You can refine the calculated phase diagram or property diagram.

Moreover, many more property diagrams with axes other than compositions can also be plotted (after the calculation), using <u>"SET\_AXIS\_VARIABLE" on page 208</u> in the sequent POST monitor.

Syntax	DEFINE_DIAGRAM
	These prompts are given:
	Same elements as before /Y/?
	Mole percent of <element> /##/: <value></value></element>
	or
Prompts	Mass percent of <element> /##/: <value></value></element>
	Database /ABCDE/: <database name=""></database>
	Major element or alloy: <element name=""></element>
	Composition in mass (weight) percent? /Y/:
	1st alloying element: <element name=""></element>

Syntax	DEFINE_DIAGRAM
	Mass (weight) percent: <amount above="" element="" of="" specified="" the=""></amount>
	2nd alloying element: <element name=""></element>
	Next alloying element: <element name=""></element>
	Mass (weight) percent: <amount above="" element="" of="" specified="" the=""></amount>
	Temperature (c) /1000/: <temperature in="" interest="" oc="" of=""></temperature>
	Reject phase(s) /none/: <list be="" of="" phase(s)="" rejected="" to=""></list>
	Restore phase(s) /none/: <list be="" of="" phase(s)="" restored="" to=""></list>
	OK? /Y/: <y n="" or=""></y>
	Should any phase have a miscibility gap check? /N/:
	Phase with miscibility gap: <phase name=""></phase>
	<pre>Major constituent(s) for sublattice #: /aa/: <constituent(s)></constituent(s)></pre>
	Phase with miscibility gap: <phase name=""></phase>
	The first equilibrium is calculated, as with <u>"DEFINE_MATERIAL" on the next</u> page. Then a list of all independent conditions suitable to be chosen as X/Y-axis variables is given by the program.
	Quit? /Y/: <y n="" or=""></y>
	This question is asked only when the axis variables are already defined, or if the command DEFINE_DIAGRAM is used. It then offers an opportunity to quit (Y) the calculation or to continue (N) the calculation but by defining other axes.
	Give the number of the condition to vary /1/: <a condition="" index=""> <math display="inline">% \left( \left( A_{1}^{2}\right) \right) =\left( \left( A_{1}^{2}\right) \right) \left( \left( A_{1}^{2}\right) \right) \left( A_{1}^{2}\right) \right) \left( \left( A_{1}^{2}\right) \right) \left( A_{1}^{2}\right) \right) \left( \left( A_{1}^{2}\right) \right) \left( A_{1}^{2}\right) \right) \left( A_{1}^{2}\right) \left</math></a>
	Select one of the independent conditions by giving its index on the condition list as the X-axis variable.
	Minimum value /XXX/: <minimum for="" value="" x-axis=""></minimum>
	Specify the minimum value of the chosen X-axis variable. A default value is shown automatically by the program; press <enter> to accept it or input another value.</enter>
	Maximum value /YYY/: <maximum for="" value="" x-axis=""></maximum>
	Specify the maximum value of the chosen X-axis variable. A default value is shown automatically by the program; press <enter> to accept it or input another value. Then another list with some dependent quantities is given by the program, which can be selected as the Y-axis variable.</enter>
	Give the number of the quantity on the second axis /#/: <##>



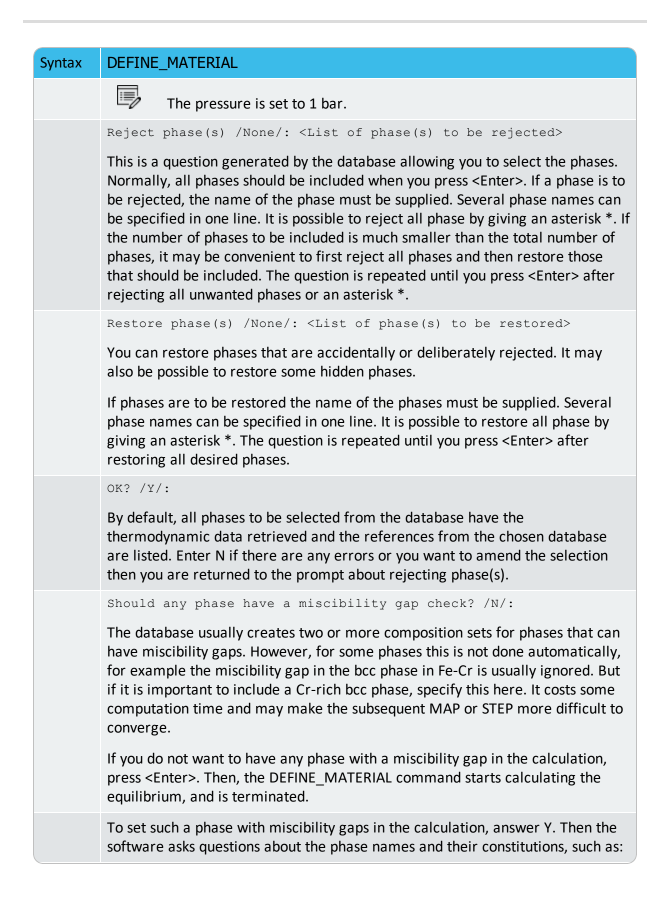
#### **DEFINE\_MATERIAL**

Read data for a system from a database in the POLY module. It is convenient to use for alloys when there is a major component and the amount of the other elements is known in mass (weight) fraction. The command reads the system from the specified database, sets the composition and temperature (and pressure equal to 1 bar) and calculates the equilibrium state before the prompting for a new command. You can list the results with "LIST\_EQUILIBRIUM" on page 198 or set a new composition or set axis for a STEP or MAP command.

You cannot append data from different databases in this way. Use this command with data from a USER database.

Syntax	DEFINE_MATERIAL
	Same elements as before /Y/?
Prompts	This question is asked only if some data is already read from the database, or if the command DEFINE_MATERIAL or DEFINE_DIAGRAM is used. It then offers a convenient way to change the composition and temperature with one command.
	This command only works properly in cases where the composition of the material system is already defined as in the mole-percent or mass-percent unit.
	Mole percent of <element> /##/: <value></value></element>
	or
	Mass percent of <element> /##/: <value></value></element>
	If you have decided to use the same materials system (available in the current POLY3 workspace) by accepting the default (Y) to the previous prompt Same elements as before /Y/?, one of the alternative prompts display for each of the components in the defined system, depending on how the composition is defined (either in mole-percent, or in mass-percent).
	Prompts are repeated until all the defined components are completed. Then, the program prompts to specify the temperature condition.
	Database /ABCDE/: <database name=""></database>
	The database with the description for the material must be given, or press <enter> if using the current database. It is possible to give a USER database.</enter>
	Major element or alloy: <element name=""></element>
	The material must have a major element, usually the element which is present in the largest amount. The fraction of this element is not set but is the rest.
	In some databases there are the alloys predefined. An alloy has a default major element and have limits of the amounts of the alloying elements. If you stay within these limits the calculation gives reasonable results.
	Composition in mass (weight) percent? /Y/:
	By default the input is taken as mass percent. Choose N to change to mole percent.

Syntax	DEFINE_MATERIAL
	Composition should be given in PERCENT not FRACTION, as it is required for the w and x state variables in the <u>"SET_CONDITION" on page 209</u> command.
	1st alloying element: <element name=""></element>
	The first alloying element must be given.
	All alloying elements are asked for in a sequence. These can be given in any order. You must know if you are present as assessed systems in the database. There is no error or warning messages if data are missing. Check the documentation of the database selected.
	If an alloy is selected, a list of legal alloying elements and their maximum percent is listed on-line.
	Mass (weight) percent: <amount above="" element="" of="" specified="" the=""></amount>
	The amount of the alloying element in mass (weight) percent. Using the DEFINE_MATERIAL command you cannot use the normal flexibility of Thermo-Calc for conditions, but all must be given in mass percent. However, you can after wards change the conditions using the SET_CONDITION command.
	2nd alloying element: <element name=""></element>
	The second alloying element must be given. If only one, press <enter>. If an element name is given then the program asks for its mass fraction.</enter>
	Mass (weight) percent: <amount above="" element="" of="" specified="" the=""></amount>
	The amount of the above specified alloying element in mass (weight) percent.
	Next alloying element: <element name=""></element>
	Continue giving elements and mass (weight) fractions until all elements specified. When all alloying elements and their compositions (as in the above prompt) are specified, press <enter> as answer to this question to finish the materials definition.</enter>
	Temperature (C) /1000/: <temperature in="" interest="" oc="" of=""></temperature>
	POLY makes the first calculation after retrieving the data for this temperature. By pressing <enter> to accept the default temperature. The value should be given in Celsius (°C).</enter>



Syntax	DEFINE_MATERIAL
	Phase with miscibility gap: <phase name=""></phase>
	You must supply the phase name, which has a miscibility gap under the specified system and conditions.
	<pre>Major constituent(s) for sublattice #: /AA/: <constituent(s)></constituent(s)></pre>
	The software shows a default constituent in the sublattice $\#(1, 2, 3,)$ , according to the existing phase definition in the chosen database. You can specify one or more major constituents for the sublattice $\#$ in the phase.
	This question is repeated until all sublattices are specified.
	Phase with miscibility gap: <phase name=""></phase>
	You can supply another phase name with a miscibility gap under the specified system and conditions, and answer the questions concerning the major constituent(s) in associated sublattice(s).
	By pressing <enter>, the command starts calculating the equilibrium, and then terminates.</enter>
	It is also possible to use this command to select an alloy from a specific database (e.g. the TCNI Ni-based Superalloys Database). Such alloys are predefined by the OPTION keyword inside the database, and have their default major elements and composition limits of their alloy elements.
	Such alloys available in the selected database (at the prompt Database /ABCDE/ can be listed on the screen if typing a ? mark on the prompt Major element or alloy. When a specific predefined alloy (instead of a major element) is selected, the major element is staked from the alloy definition and shown on the screen (with a message like Alloy found with major element NI).
	You can only specify alloying elements and the compositions (weight percent or mole percent). Typing a ? at any of the prompts for the alloying element names, e.g. 1st alloying element, 2nd alloying element, lists all the alloying elements and the composition limits in the alloy. If the composition of an alloying element is outside of its limit, there is a message (such as Amount above limit: 30.0000) and a prompt override limit ? /N/. If you decide to enforce the override by answering Y on this prompt (i.e. accepting the overlimit alloying composition), another warning message (such as Amount of major element below limit: 70.0000) and prompt override limit ?

#### Syntax **DEFINE\_MATERIAL**

/N/. Then further decide if enforcing the overriding: if Y then accepting the major element's composition below the limit; if N then using the predefined major element composition limit.

#### DELETE\_INITIAL\_EQUILIBRIUM

Delete ONE specific initial equilibrium point or ALL of the initial equilibria. The initial equilibria are used as starting points for all the sub-sequential MAP and STEP calculations.

"ADD\_INITIAL\_EQUILIBRIUM" on page 160.

#### Syntax DELETE\_INITIAL\_EQUILIBRIUM

Prompt Number /All/: <Number of an initial equilibrium>

Specify the number of an initial equilibrium (as a specific digit e.g. 3, or ALL) to be deleted from the POLY3 workspaces. It is recommended to use "LIST\_INITIAL\_ EQUILIBRIA" on page 199 to figure out the numbers for all the existing initial equilibrium points that have already been added (*manually* through "ADD\_ INITIAL\_EQUILIBRIUM" on page 160, or *automatically* by some other POLY commands) and stored in the current POLY3 workspace. By default, ALL the initial equilibrium points are deleted.

#### DELETE\_SYMBOL



This is a POLY3 module command.

Delete symbols (i.e. constants, variables, functions or tables. These symbols are entered with <u>"ENTER\_SYMBOL" on page 109</u>.

# Syntax DELETE\_SYMBOL Name: <Name of a symbol> Prompt Specify the name of the symbol to be deleted. Only one symbol can be deleted each time.

#### **ENTER\_SYMBOL**



This command is for both the POST and POLY modules in Console Mode.

Symbols are a useful feature of the POLY and POST modules to define quantities that are convenient. Symbols can be constants, variables, functions or tables.

Functions or tables (with defined functions as values) can be entered in the POST module after a stepping or mapping calculation, for purposes of plotting such entered functions or tables as axis variables.

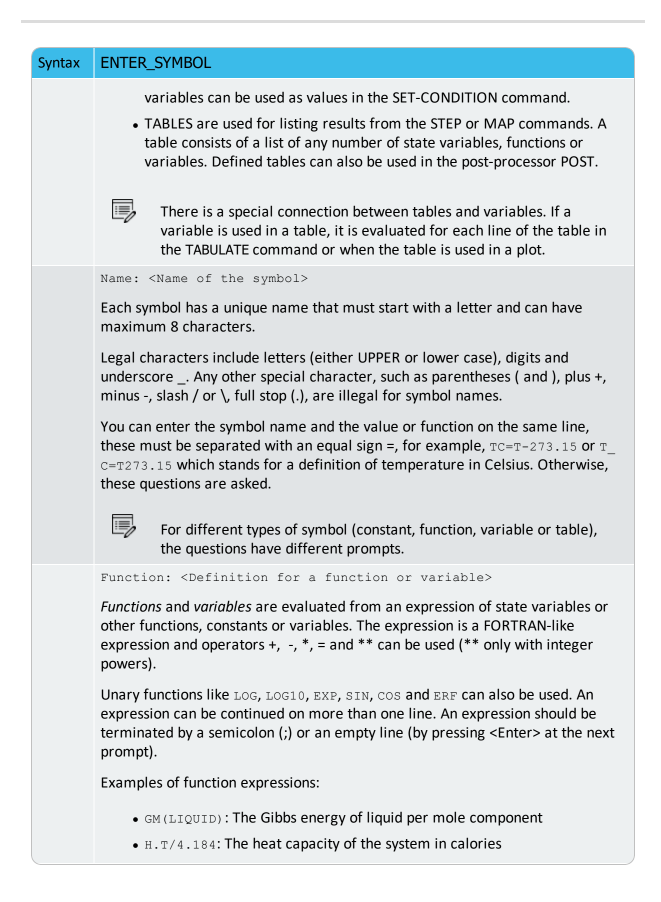
Within the POLY module, symbols are normally defined prior to an equilibrium calculation (enforced by a C\_E command), stepping calculation (enforced by the command <u>"STEP</u><u>WITH\_OPTIONS" on page 217</u>) or mapping calculation (enforced by <u>"MAP" on page 201</u>). These can be entered after an equilibrium calculation; however, for defined functions, variable or tables, it requires using <u>"EVALUATE\_FUNCTIONS" on page 197</u> before showing the corresponding values in the calculated equilibrium state.

The symbols entered in the POST module are not saved in the currently-loaded POLY3 workspaces. Therefore, if you want to apply such symbols in other similar calculations for the same defined system, you must use the ENTER\_SYMBOL command prior to the STEPPING or MAPPING calculation in the POLY module.

- <u>"ENTER\_SYMBOL" on page 109</u> for the GIBBS Module.
- .:::

See <u>example 44</u> in the *Thermo-Calc Console Mode Examples Guide* for an example of using variables and functions.

Syntax	ENTER_SYMBOL
	Constant, Variable, Function or Table? /Function/: <keyword></keyword>
Prompt	The Keyword can be specified as CONSTANT, VARIABLE, FUNCTION or TABLE.
	<ul> <li>CONSTANTS can only be entered once and is a means of using a name for a numeric value. For example, the value of 1 atmosphere in Pascal can be denoted by P0 after the command ENTER CONSTANT P0=101325. Defined constants can be used as values in condition assignments, for example, SET-COND P=P0.</li> </ul>
	<ul> <li>FUNCTIONS are expressions of state variables or other functions. These expressions are saved, and whenever a function value is requested all functions are evaluated. The reason for this is that they may depend on each other.</li> </ul>
	<ul> <li>VARIABLES are similar to functions because they can also be expressions of state variables. However, contrary to functions, they are only evaluated when they are entered or if they are explicitly named in an <u>"EVALUATE_FUNCTIONS" on page 197</u> command. It is possible to enter a variable with a new expression any time. This expression is evaluated directly and its value stored as the value of the variable. Defined</li> </ul>



```
ENTER_SYMBOL
Syntax
              • ACR(CR)/X(FCC,CR): The activity coefficient for Cr in FCC

    T-273.15: The temperature in Celsius

         &: <Continuation of the definition for the symbol>
         The ampersand & means that you can continue to write the function on the new
         line if one line is not enough for the function. If you finish the function press
         <Enter> again.
         Value: <Value for a constant>
         A constant can only be assigned a numeric value once.
         Value or expression: <Value of expression for a variable>
         A variable can be assigned a numeric value or an expression. An expression is
         evaluated immediately and discarded. Only the numeric value is kept. This gives
         a possibility to save a value between calculations with different conditions
         because all state variables and functions are evaluated for the new conditions.
         Variable(s): <Variable(s) in a table>
         A table consists of a list of state variables or functions. One way to obtain results
         from a STEP command is through a table.
         Example:
            ENTER TABLE K=T, X(LIQ, C), X(LIQ, CR), ACR(C)
         Which means that the table called K contains four columns, i.e. the temperature,
         the mole fractions of C and Cr in the LIQUID phase, and the activity of C.
         To show the temperature in Celsius in a table, give the command ENTER
         FUNCTION TC=T-273; and then use the symbol TC in the table.
         & <Continuation of the definition for the table>
         The ampersand a means that you can continue to write the table on the new line
         if one line is not enough for the table. If you finish the table press <Enter> again.
```

#### **EVALUATE\_FUNCTIONS**

The value of one or more or all entered functions or variables are evaluated and listed.

Syntax	EVALUATE_FUNCTIONS
Prompt	<pre>Name(s): <name(s) defined="" function(s)="" of=""></name(s)></pre>

#### Syntax EVALUATE\_FUNCTIONS

The names of one or more entered functions or variables must be specified. By typing a wildcard \*, all functions and variables are evaluated.

Variables are evaluated only if these are explicitly named.

# LIST\_AXIS\_VARIABLE



This is a POLY3 module command.

#### Syntax LIST\_AXIS\_VARIABLE

Lists all the axis variables for a stepping or mapping calculation that have already been set by "SET\_AXIS\_VARIABLE" on page 208.

# LIST\_CONDITIONS

1	
	>
	-

This is a POLY3 module command.

All the conditions that are set by the command SET\_CONDITION and the command-sequence CHANGE\_STATUS PHASE =FIXED <0 or 1 or alike>, are listed. The current conditions are also listed by the command LIST\_EQUILIBRIUM. The degree of freedom in the defined system is also shown up. If this is zero, you can perform a COMPUTE\_EQUILIBRIUM command. If it is larger than zero, some more conditions are required, and you must further set additional ones, using <u>"SET\_CONDITION" on page 209</u> or <u>"CHANGE\_STATUS" on page 177</u>. If it is negative, a user has defined too many conditions and needs to take away the unnecessary ones, using the SET\_CONDITION command (with a value of NONE for the to-be-deleted condition) or CHANGE\_STATUS command (i.e. changing a FIXED status of a phase to another type of phase status, ENTERED or DORMANT or SUSPENDED).

Syntax	LIST_CONDITIONS
	P=100000, T=800, N(NI)=1E-1, N=1
Example	FIXED PHASES
	FCC_A1=1 LIQUID=0 DEGREE OF FREEDOM 0

# LIST\_EQUILIBRIUM



This is a POLY3 module command.

The result (always in SI units) from the last calculated equilibrium is listed on screen or in a textual file.

You can also execute this command if no calculation is made or if the calculation fails. It is your responsibility to interpret the results accordingly.

Syntax	LIST_EQUILIBRIUM
Duovout	Output to screen or file /Screen/: <file name=""></file>
Prompt	The name of the text file where the list of the calculation results shall be written.
	Options /VWCS/: <option(s)></option(s)>
	Select the output units and formats by optionally specifying a combination of these letters.
	• Fraction order: w means VALUE ORDER; A means ALPHABETICAL ORDER.
	• Fraction type: w means MASS FRACTION; x means MOLE FRACTION.
	<ul> <li>Composition: c means only COMPOSITION; N means CONSTITUTION and COMPOSITION.</li> </ul>
	<ul> <li>Phase: s means including only STABLE PHASES; P means including ALL NON-SUSPENDED PHASES.</li> </ul>
	Default options are vwcs. If the output fraction type should be in mole fraction (rather than mass fraction), then give vxcs or type x (implying that in this case the options V, C and S are accepted as the listing manners for fraction order, composition and phase).
	If accepting all the default options, or if accepting all the altered options that had already changed when using this command previously, you can type $L_E$ , , , or $LIST_EQUILIBRIUM$ , , ,
	LIST_EQUILIBRIUM displays ORD (ordered) or DISORD (disordered) (when available). See <u>"CHANGE_LEGEND" on page 237</u> for details.

# LIST\_INITIAL\_EQUILIBRIA

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l	>_	

This is a POLY3 module command.

List all the equilibria added with <u>"ADD\_INITIAL\_EQUILIBRIUM" on page 160</u>.

#### Syntax LIST\_INITIAL EQUILIBRIA

All the initial equilibria are used for the MAP (and STEP) calculations.

#### LIST\_STATUS



This command is for the POLY and ED-EXP modules.

List the status of components, species or phases.

Syntax	LIST_STATUS
	Specify what to list:
	Option /CPS/: <keyword(s)></keyword(s)>
	keyword = C or P or S, or any combination
Prompt	• c: list component status
	• P: list phase status
	• s: list species status
	<ul> <li>cs: list both components and species</li> </ul>
	Default is CPS.
	The results depend on the key word specified in the options for <u>"CHANGE</u> <u>STATUS" on page 96</u> , a table with the current status of phases or species or components, or the combinations, is shown:
	<ul> <li>For components, the statuses and reference states are listed.</li> </ul>
	• For species, only the status is listed.
	<ul> <li>For ENTERED and FIXED phases, the status, driving forces and equilibrium amounts are listed.</li> </ul>
	The metastable phases are listed in descending order of stability. To avoid long outputs only 10 metastable phases (in ENTERED status) are listed by lines, while all other less stable phases are merged onto one line. For DORMANT phases, their phase names and driving forces are listed. For SUSPENDED phases, only the phase names are listed.

# LIST\_SYMBOLS



For both the POLY and POST modules, list the definitions for all constants, functions, variables and tables.

In order to find the value of a function or variable, use <u>"SHOW\_VALUE" on page 217</u> or <u>"EVALUATE\_FUNCTIONS" on page 197</u>. A table is tabulated with the TABULATE command.

#### Syntax LIST\_SYMBOLS

The defined variables are listed up together with the defined functions, but variable names are followed by a percentage sign %.

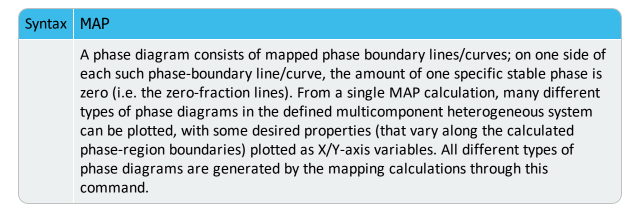
# LOAD\_INITIAL\_EQUILIBRIUM

Copies all conditions and calculated results from a specific added initial equilibrium to the current equilibrium. The current conditions and calculation results are lost, and the newly loaded initial equilibrium point gets into the POLY workspace.

Syntax	LOAD_INITIAL_EQUILIBRIUM
	Number: <number an="" equilibrium="" initial="" of=""></number>
Prompt	Specify the number of an initial equilibrium to be loaded as current. The number can be found with <u>"LIST_INITIAL_EQUILIBRIA" on page 199</u> .

#### MAP

This command starts the mapping procedure for making a calculation of phase diagrams in a defined multicomponent heterogeneous system, from one or more initial equilibria. A phase diagram is usually mapped within a specific space that is constructed by two (or more) defined independent mapping axis-variables.



Normally, you need to have calculated at least one initial equilibrium point and have also defined at least two independent varying variables (i.e. the controlling conditions in the system) that are set with <u>"SET\_AXIS\_VARIABLE" on page 208</u>. You can also have three, or four or maximum five independent varying variables that are defined by the SET\_AXIS\_VARIABLE command).

This lists the current values of each of the independent axis variables for each of the calculated equilibrium points along each of the mapped phase boundaries, and also lists the corresponding information when the set of stable phases changes.

The Global Minimization Technique is by default enforced in this command while it can be disabled if having decisively switched it off (or possibly in some special modules) through changing the minimization option using the <u>"ADVANCED\_OPTIONS" on page 167</u>  $\rightarrow$  MINIMIZATION\_OPTION command-sequence. You can choose how often to do a Global Minimization test (via the Global Test Interval option) in the ADVANCED\_OPTIONS STEP\_AND\_MAP command-sequence.

During a MAP calculation, the values of mapped axis-variables for presenting each phase boundary (lines/points) of the calculated phase-regions are listed, and the corresponding stable-phase sets are shown up.

You can terminate the mapping of a line by pressing a single <Ctrl-A> (Windows) or <Ctrl-C> (Linux). This can be useful in order to stop a longish calculation without losing what is already calculated.

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If there is any convergence problem in finding a stable solution at a certain stage during a calculation procedure enforced by this command, these messages display on screen:

```
Convergence problems, increasing smallest site-fraction from 1.00E-30 to hardware precision 2.00E-14. You can restore using SET-NUMERICAL-LIMITS
```

This implies that smallest site fraction in the current POLY3 workspace is automatically increased from the default value 1.00E-30 to the hardware-dependent precision (under Linux, as 2.00E-14). For other subsequent POLY-module calculation in the current TC run, you can use the POLY command <u>"SET\_NUMERICAL\_LIMITS" on page 293</u> to restore or reset the smallest site fraction to the previous or another preferred value, as well as to reset other numerical limits.

In particular, for phase diagrams with tie-lines in the plane (i.e. most binary systems and ternary isotherms), there is a MAP procedure which checks for the best phase to use as axis variables in order to ensure reasonable increments between the tie-lines. This produces smoother curves and also gives a better stability in finding adjacent phase-regions.

Complex miscibility gaps of various solution phases are automatically detected during a mapping calculation, and two or more composition sets for each of such solution phases are automatically generated as well, through an automatic Global Minimization Test procedure (by specifying a Global Test Interval value and confirming the Automatically add initial equilibria in the ADVANCED\_OPTION STEP\_AND\_MAP command-sequence). As a particular case, such an automatic Global Minimization Test ensures that you can start calculating from high temperatures in an austenitic steel (metallic FCC\_A1 solution phase) where the MC carbides/nitrides/carbonitrides (i.e. the C-/N-rich sides of the FCC\_A1 miscibility-gap, often (while not always) being referred to as FCC\_A1#2, FCC\_A1#3, etc.) are not stable, and during the MAP command the MC carbides/nitrides/carbonitrides may first become metastable with a composition different from the metallic FCC\_A1 phase and later also stable. The advanced mapping procedure (enforced by the Global Minimization Technique) inside the Thermo-Calc software (both the Console Mode and Graphical Mode) can efficiently and effectively handle complex miscibility gaps in multicomponent heterogeneous systems.

To get a complete phase diagram, sometimes it may be necessary to have multiple starting equilibrium-points, and/or to have added multiple initial-equilibrium points (through <u>"ADD\_INITIAL\_EQUILIBRIUM" on page 160</u>) inside various phase-region boundaries under certain direction(s).

# POST

Syntax	POST
	Switches to the post-processor (the POST module), which has its own command set.
	Plotting and Visualization with the POST Module " on page 231

# **READ\_WORKSPACES**



This command is used in both the POLY and GES (GIBBS) modules.

The POLY3 and GIBBS workspaces and the calculated results from the MAP and STEP commands can be READ from a file where they are saved with <u>"SAVE\_WORKSPACES" on page 52</u>. Such an \*.POLY3 file is not printable.

Syntax	READ_WORKSPACES
Options	File name is the name of a saved POLY3-file where the POLY3 and GIBBS workspaces shall be read from must be specified. You do not need to type the extension if it is the default *.POLY3 (Windows) or *.poly3 (Linux), otherwise type the whole POLY-file name.

#### Syntax READ\_WORKSPACES

A window opens so that the path (in the **Look in** field) and **File name** can be specified. The **Files of type** (i.e. POLY3) cannot be changed. Click **Open** to open the POLY3 and GIBBS workspaces from the saved \*.POLY3 file.

When reading back an original POLY3 workspace that has already been saved as an \*.POLY3 file in the current Thermo-Calc (Console Mode) run or had been read from an existing POLY3 file under the current work area, while some additional changes in the settings may be made but do not need to be kept in further steps in the current Thermo-Calc (Console Mode) run or any diagram is plotted in the POST module, you can type READ, , or READ\_WORKSPACE , ,

#### **RECOVER\_START\_VALUES**

#### Syntax RECOVER\_START\_VALUES

Recovers the start values in equilibrium calculations.

#### **REINITIATE\_MODULE**

Ensures that the whole POLY module (and thus the entire POLY workspace) is reinitiated to the state as it was when first entered.

#### Syntax REINITIATE\_MODULE

All the defined components, defined conditions, changed status, entered symbols, defined independent axis-variables, calculated starting equilibrium-points, added initial equilibrium points, stepped/mapped equilibrium data, and so forth, are removed completely. The saved file name is restored to the default.

#### SAVE\_WORKSPACES

>\_

This command is for both the POLY3 and GES (GIBBS) modules.

Thermo-Calc allows saving the current status and workspaces of the program, including thermodynamic data, conditions, options and results from a single, stepping or mapping calculation on an \*.POLY3 file. Do this for later use or when you need to terminate the current Console Mode session.

The POLY3 and GIBBS workspaces are saved on a file with this command. In the GIBBS workspace, all thermochemical data are stored. In the POLY3 workspace, all thermochemical data, all the last set of conditions and equilibrium state, changed status,

entered symbols, advanced options, defined stepping/mapping variables, added initial equilibria, stepped/mapped results, etc., are stored, so it also contain the GIBBS workspace. After a SAVE\_WORKSPACE command, you can always come back to exactly the state you had when you issued the SAVE\_WORKSPACE command by giving a READ\_WORKSPACE command.

After saving the POLY3 and GIBBS workspaces on a file, you can leave the program and at a later time READ the file and continue from the saved state.



A STEP or MAP command automatically saves the work file with the most lately specified name. Do not SAVE after a MAP or STEP.

The results from the STEP or MAP commands are destroyed by the SAVE\_WORKSPACE command. You may append several results obtained by sequential STEP or MAP calculations without destroying the previous results, whilst SAVE\_WORKSPACE erases them all. Keeping this in mind is important and useful particularly for calculating various isothermal (or isoplethal) sections and plotting them on the same diagram in a single Thermo-Calc run.

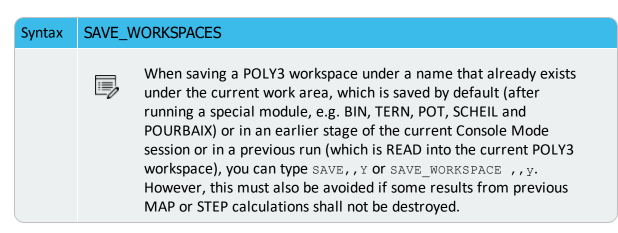
You may append several results by STEP or MAP without destroying the previous results but the SAVE command erases them all. To suspend some of the STEP or MAP results, use the AMEND\_STORED\_EQUILIBRIA command.

When you go to a response-driven module such as POTENTIAL or SCHEIL for example, a workspace file is automatically opened. In the workspace file, system definitions, conditions for the calculation, calculation results, and plot settings is saved. The file is saved in the current working directory, and is named after the name of the module that created it. For example, the POTENTIAL module saves a workspace file called POT.POLY3 (or POT.poly3 in Linux); the POURBAIX module saves a file called POURBAIX.POLY3; etc.

#### Syntax SAVE\_WORKSPACES

A file name must be specified. The default extension of the POLY workspace file is \*.POLY3 (Windows) or \*.poly3 (Linux), while you can have any other options extension.

A **Save** window displays, so that the path (in the **Save in** field) and **File** name can be specified. The file type (i.e. POLY3) in the **Files of type** cannot be changed.



#### **SELECT\_EQUILIBRIUM**

If you create more than one initial equilibrium (during data-assessments using the PARROT/ED\_EXP modules), you can switch between them using this command.

Syntax	SELECT_EQUILIBRIUM
	Number /Next/: <choice equilibrium="" on=""></choice>
Prompt	Options: FIRST, LAST, NEXT, PREVIOUS OF PRESENT. Most commands affect only the PRESENT equilibrium. However, the commands "REINITIATE_MODULE" on page 204 and "DEFINE_COMPONENTS" on page 186 remove all the stored equilibria.

# SET\_ALL\_START\_VALUES

Set start values, e.g. if the calculation fails or if you have a miscibility gap or ordering. If temperature and pressure are not conditions, you are asked for values of them. Then for each phase prompt on if it should be stable and on its constitution.

Syntax	SET_ALL_START_VALUES
	T /XXXX/: <temperature in="" k=""></temperature>
	If the temperature is not a condition, supply a guess of its final value (in K).
	P /100000/: <pressure in="" pa=""></pressure>
	If the pressure is not a condition supply a guess of its final value (in Pa).
	Automatic start values for phase constitutions? /N/: <y, f="" n="" or=""> <math display="inline"></math></y,>
Prompts	<ul> <li>For N, enter an initial amount of each entered phase and the major constituents or site fraction of each constituent.</li> </ul>

Syntax	SET_ALL_START_VALUES
	<ul> <li>For Y, the program automatically sets the start values for phase constitutions in all possible phases.</li> </ul>
	<ul> <li>F is for force. In some cases the calculation fails because impossible conditions are set, e.g. W(C) =1.5 [W(C) is mass fraction and must thus be less than unity. The program tries to fulfill this condition by putting maximum amount of carbon in all phases, but can fail anyway. When you detect the error and set W(C) to 0.015, the calculation may still fail because it could start from the previous values. To get back to 'fresh' start values, choose F.</li> </ul>
	Should <phase> be stable /N/: <y 1="" 2="" n="" or=""></y></phase>
	A guess if this phase should be stable or not is requested. All entered phases are prompted for this question and next two.
	You cannot have more phases stable than you have components, but at least one (which dissolves all constituents) must be set as stable. For backward compatibility, this question can be answered by 1 (for Yes) or 0 (for No).
	The phase name may have a hash sign # followed by a digit, e.g. BCC_A2#2. For phases with miscibility gaps, there should be two phases with the same name but with different numbers after the hash sign.
	<pre>Major constituent(s): <name constituent(s)="" in="" major="" of="" phase="" the=""></name></pre>
	The constituent with the largest fraction in the phase should be specified. If there are more than one constituent with a large fraction, give them all on the same line. If the default major constituents should be used answer with an asterisk (*). By giving \$ the constitution is not changed. If there should be no major constituent give NONE and or if the major constituent(s) are improperly specified, you are asked for individual fractions in the phase.
	Y( <phase>,<constituent>) /.XXXXXXXXX/: &lt;.YYYYY&gt;</constituent></phase>
	The current value (.xxxxxxxxx) is default. You can accept the default by pressing <enter> or give a new value (.YYYYY).</enter>
	The phase name or constituent name may have a hash sign # followed by a digit, e.g. $Y(BCC_A2#2, FE)$ , $Y(BCC_A2#2, C#2)$ . For phases with miscibility gaps, there should be two phases with the same name but with different

#### Syntax SET\_ALL\_START\_VALUES

numbers after the hash sign. For phases with sublattices, the constituents in sublattice 2 or higher is also be suffixed with a hash sign # followed by a digit.

# SET\_AXIS\_VARIABLE

In order to calculate a diagram, set at least one axis variable in a stepping calculation, or at least two axis variables in a mapping calculation. For property diagrams, one axis is enough; for phase diagrams two or more are necessary. Any condition that can be set to calculate an equilibrium can be used as an axis variable (with its lower and upper limits and step length) by using the SET\_AXIS\_VARIABLE command, and the POLY program does, after a STEP or MAP command, vary the value of the condition between the limits set on its related axis variable. As an extraordinarily unique and powerful feature of the Thermo-Calc software, up to five independent axis variables can possibly be used in a mapping calculation of a multicomponent heterogeneous system; however, the axis numbers 3, 4 and 5 must have chemical potentials of components (or temperature or pressure) as conditions.

Syntax	SET_AXIS_VARIABLE
	Axis number /#/: <an axis="" number=""></an>
	Specify a number between 1 and 5. The axis numbers 3, 4 and 5 must have chemical potentials of components (or temperature or pressure) as conditions.
	Condition /None/: <one condition=""></one>
Prompts	Here the condition that should be varied along the axis must be given. The condition is specified as in <u>"SET_CONDITION" on the next page</u> , for example $W$ (C) for mass fraction of carbon. By accepting NONE, the axis is removed.
	Min value /0/: <min value=""></min>
	Specify the minimum value of the axis condition.
	Max value /1/: <max value=""></max>
	Specify the maximum value of the axis condition.
	<pre>Increment /.025/: <step length=""></step></pre>
	Specify the maximum step length. By default, this is 1/40 of the total axis length.

You can give the SET\_AXIS\_VARIABLE command without having set a condition on the axis variable. Under such circumstances, the relevant condition is automatically created and the

value set between the minimum and maximum axis limits; however, as a side effect, the POLY module creates two conditions, P=1e5 and N=1 (these have not been defined as a condition yet), in case you set an axis variable which is not already a condition.

You can use a logarithmic axis during calculations. This is useful for low fractions like in a gas phase where 1e-7 to 1e-2 might be an interesting range. The pressure is also suitable for logarithmic step. You specify the logarithmic axis by giving an asterisk \* after the increment value.



The increment in this case is treated as a factor.

For example,

S-A-V 1 P 1E5 1E25 5\*

This makes axis 1 a logarithmic axis where the difference between two calculated values makes a factor 5.



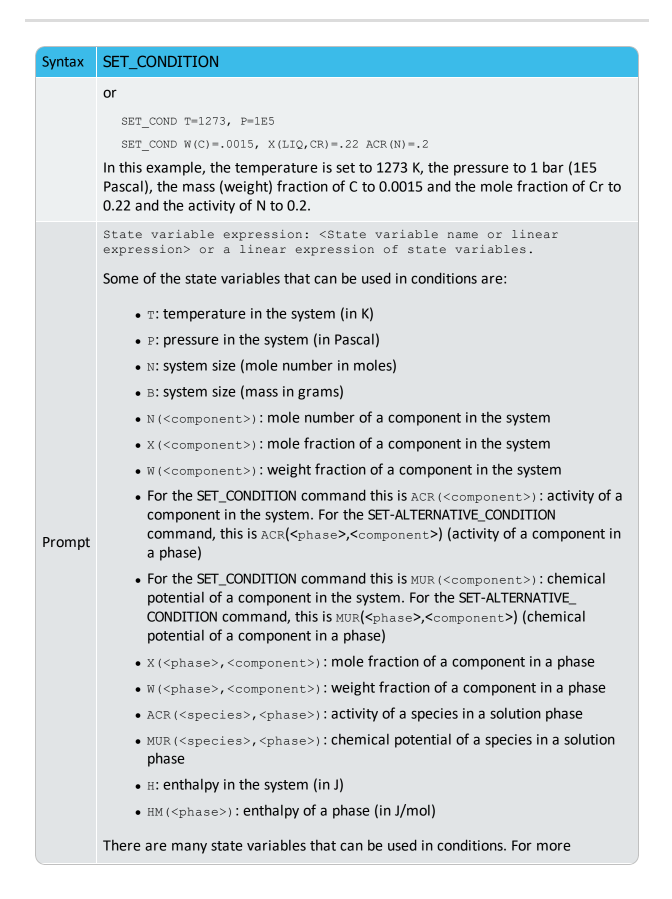
The factor must be larger than 1.0.

In some cases, such as when the DEFINE\_DIAGRAM command is used or a special advanced module (e.g. BIN, TERN, POT, SCHEIL or POURBAIX) is called, some axis variables are automatically set by the program, not necessarily by this command.

# **SET\_CONDITION**

Specify the equilibrium conditions for the calculation. All kinds of state variables, as well as most of the M/W/V/F-suffixed state variables (for normalization) and R-suffixed state variables (with respect to chosen reference states), can be used as conditions. Beside temperature and pressure conditions, a multicomponent system can have mixed types of conditions; this brings the extreme flexibility and powerfulness to the complex calculations with the Thermo-Calc software system. You should repeat this command for a multicomponent system, till the degree of freedom in the defined system becomes zero. When a FIXED phase status is used on a specific phase (use <u>"CHANGE\_STATUS" on page 177</u>), it is equivalent to one condition (implying that particular phase is stable in the defined system).

Syntax	SET_CONDITION
	Each condition must be given explicitly, but can be given on the same line (separated by a comma (,) or an empty space) or on separate lines with each one started with the command.
	EXAMPLE
	SET_COND T=1273, P=1E5, $W(C) = .0015$ , $X(LIQ, CR) = .22$ , ACR(N) = .2



Syntax	SET_CONDITION
	information, enter an INFO STATE_VARIABLES command. A condition is normally a value of a single state variable with its value.
	Example
	T=1273.15 P=1E5
	X(C)=.002
	W(CR)=0.5
	ACR (CR) =0.85
	X(FCC,C)=.001
	H=-250000
	HM(BCC)=-225000
	A condition can also be a value of a linear expression involving more than one state variable. For example,
	X(LIQ,S)-X(PYRR,S)=0
	This means that it is a condition that the mole fraction of $ s$ component should be the same in the LIQUID and PYRRHOTITE phases. In practice, that should be the congruent melting point.
	After the equal sign only a numeric value is allowed in the expression.
	Factor: <a a="" continuation="" factor="" for="" or="" state="" the="" variable,=""></a>
	This question means you did not answer the previous question. The program is then expecting a single state variable or a complete state variable expression, or the numeric factor in an expression with only one state variable. In a state variable expression a state variable may be preceded by a constant factor. An example of this is:
	2*MUR(FE)+3*MUR(O)=-35000
	This means that it should be a condition that two times the chemical potential of ${\tt FE}$ plus three times the chemical potential of $\circ$ should be -35000 J/mol.
	State variable: <a a="" continuation="" or="" specified="" state="" variable,=""></a>
	This question is prompted if a single state variable name has not given in either the prompt State variable expression or Factor, or a state variable expression is given but the expression is incomplete, for example, T- or 2*MUR

Syntax	SET_CONDITION
	(FE) +, for which the program is then expecting a continuation of the unfinished expression. You need to specify a state variable or a complete state variable expression, or complete the unfinished state variable expression. If a numeric factor is given before this prompt, only one state variable can be specified; otherwise, the program only takes the first state variable to complete the expression (i.e. the factor times the state variable).
	Value /X/: <a a="" constant="" numeric="" or="" value,="" variable=""></a>
	The value of the condition. This can be a numeric value, a constant or a variable. A suggestion is given as the default value. The special value NONE means that the condition is removed; for example $T=NONE$ takes away the temperature condition.

#### SET\_INPUT\_AMOUNTS

Specify how a system is made up from mixing of various substances. It is useful with a substance database. In the C-H-O-N system, you can, for example, give:

S-I-A N(H2)=10, N(H2O)=25, N(C1O2)=5, N(N2)=100

The POLY module automatically converts this into conditions for the current set of components. In the case when the elements are defined as the components, the command above is equivalent to

SET-CONDITION N(H)=70, N(O)=35, N(C)=5, N(N)=200

Syntax	SET_INPUT_AMOUNTS
	Quantity: <n(<specie>) or B(<specie>)&gt;</specie></n(<specie>
Prompts	You can give the amount also preceded with an equal sign = $[e.g. N(H2)=10$ or $B(H20)=1000$ ], or press <enter> for the next prompt on the amount of the quantity.</enter>
	Amount: <value of="" quantity="" the=""></value>
	Specify the numerical value of the quantity.
	You can give negative amounts in the SET-INPUT-AMOUNTS command.

#### **SET\_NUMERICAL\_LIMITS**

Change the criteria for convergence. This is to speed up a calculation in a complex system.

If there is any convergence problem in finding a stable solution at any stage during a calculation procedure enforced by <u>"COMPUTE\_EQUILIBRIUM" on page 182</u>, <u>"STEP\_WITH\_</u> <u>OPTIONS" on page 217</u>, <u>"MAP" on page 201</u> or <u>"ADVANCED\_OPTIONS" on page 167</u>), this message displays:

Convergence problems, increasing smallest site-fraction from 1.00E-30 to hardware precision 2.00E-14. You can restore using SET-NUMERICAL-LIMITS

This implies that smallest site fraction in the current POLY3 workspace is automatically increased from the default value 1.00E-30 to the hardware-dependent precision (under Linux, as 2.00E-14). For other subsequent POLY-module calculation in Console Mode, you can use this command to restore or reset the smallest site fraction to the previous or another preferred value, as well as to reset other numerical limits.

Syntax	SET_NUMERICAL_LIMITS
	Maximum number of iterations /200/: <xxx></xxx>
Prompt	By default, the program tries 500 iterations before it gives up. As some models give computation times of more than 1 CPU second/iteration, this number is also used to check the CPU time and the calculation stops if 500 CPU seconds/iterations are used.
	Required accuracy /1E-6/: <yyy></yyy>
	This is a relative accuracy, and the program requires that the relative difference in each variable must be lower than this value before it has converged. A larger value normally means fewer iterations but less accurate solutions. The value should be at least one order of magnitude larger than the machine precision.
	Smallest fraction /1E-12/: <zzz></zzz>
	This is the value assigned to constituents that are unstable. It is normally only in the gas phase you can find such low fractions.
	The default value for the smallest site-fractions is $1E-12$ for all phases, except for IDEAL phase with one sublattice site (such as the GAS mixture phase in many databases) for which the default value is always as $1E-30$ (unless you have used the SET_NUMERICAL_LIMITS command to reset an even-lower value (e.g. $1E-45$ , that is naturally enforced to all the phases in the system).
	Approximate driving force calculation for metastable phases /Y/:
	Normally the POLY module only requires that the set of stable phases is in equilibrium in order to terminate the iterations. By default, the metastable phases are included in all iterations but these may not have reached their most favourable composition and thus their driving forces may be only approximate.

### Syntax SET\_NUMERICAL\_LIMITS

8

If it is important that these driving forces are correct, choose N to force the calculation to converge for the metastable phases. This may require more iterations, and the STEP and MAP command may also terminate due to bad convergence in a metastable phase.

## **SET\_REFERENCE\_STATE**

The reference state for a component is important when calculating activities, chemical potentials and enthalpies and is determined by the database being used. For each component the data must be referred to a selected phase, temperature and pressure, i.e. the reference state.

All data in all phases where this component dissolves must use the same reference state. However, different databases can use different reference states for the same element/component. It is important to be careful when combining data obtained from different databases.

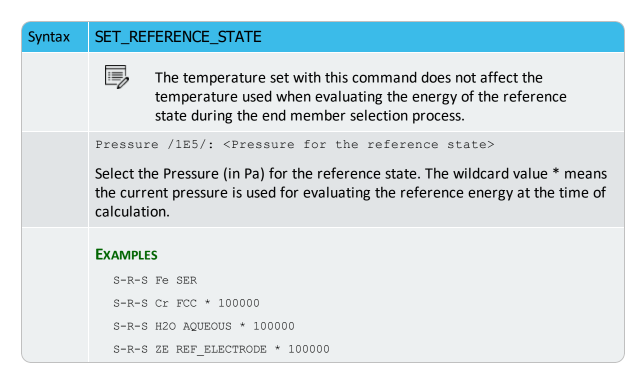
By default, activities, chemical potentials and so forth are computed relative to the reference state used by the database. If the reference state in the database is not suitable for your purposes, use this command to set the reference state for a component using SER, i.e. the Stable Element Reference (which is usually set as default for a major component in alloys dominated by the component). In such cases, the temperature and pressure for the reference state is not needed.

In order to specify conditions in the specific reference state, you can append an R to the state variables. Thus, AC is the activity (of a system component or of a species in a phase) with respect to the default reference state, and ACR is the activity with respect to the selected reference state; MU is the chemical potential of a system component with respect to the default reference state, and MUR stands for the chemical potential with respect to the selected reference state. The AC and ACR variables, for both components in a system and species in a phase, can be specified in natural logarithm, e.g. LNAC(Fe), LNACR(C), LNAC (O2,GAS), LNACR(O2,GAS).

For the POST module, and after calculating with the <u>"STEP\_WITH\_OPTIONS" on page 217</u> or <u>"MAP" on page 201</u> commands, the reference state for a component can also be changed to plot various properties of the component in the entire system or in a specific phase. Then you can set the diagram axes as the chemical potential or activity quantities with the R suffix, i.e. MUR(comp), MUR(comp,ph), ACR(comp), ACR(comp,ph) or the common logarithms [e.g. LNACR(comp,ph)].

Syntax	SET_REFERENCE_STATE
Dromoto	Component: <name component="" of="" the=""></name>
Prompts	The name of the component must be given.
	Reference phase: <name a="" as="" new="" of="" phase="" reference="" state="" the="" used=""></name>
	Enter the Name of a phase for the system.
	For a phase to be usable as a reference for a component, the component needs to have the same composition as an end member of the phase. The reference state is an end member of a phase. The selection of the end member associated with the reference state is only performed once this command is executed.
	If a component has the same composition as several end members of the chosen reference phase, then the end member that is selected at the specified temperature and pressure will have the lowest Gibbs energy.
	The following are example using temperature. Pressure is used in the same way:
	<ul> <li>In the Fe-C system, BCC can be a reference state for Fe but not for C since BCC can exist as pure Fe but cannot exist as pure C.</li> </ul>
	<ul> <li>If GAS is chosen as the reference phase of component O at such a high temperature that O1 has the lowest energy, then O1 remains the reference state even at calculations performed at lower temperatures where O2 has a lower energy than O1.</li> </ul>
	<ul> <li>Setting the reference state for component O as gas (one sublattice) with constituents O1, O2 and O3 results in O2 being the reference state if, at the present temperature, pure O2 has the lowest energy of O1, O2 and O3. If the reference state is set above a critical higher temperature, then O1 has the lowest energy and consequently becomes the reference state.</li> </ul>
	Temperature /*/: <temperature for="" reference="" state="" the=""> Select the Temperature (in K) for the reference state. The wildcard value * means the current temperature is used at the time of evaluation of the</temperature>

reference energy for the calculation.



# **SET\_START\_CONSTITUTION**

This command is similar to <u>"SET\_ALL\_START\_VALUES" on page 206</u>, but is used for an individual phase that may have a wrong constitution.

Syntax	SET_START_CONSTITUTION	
	Phase name: <name a="" and="" constituent(s)="" major="" of="" phase,="" possible=""></name>	
	Specify the name of a phase for which the constitution shall be set.	
Prompts	If there is a major constituent of the phase, you must specify this on the same line as the phase name. By giving an*, select the default major constituents. A \$ means keeping the same constitution, and NONE means to give individual site-fractions.	
	Y( <phase>#<composition_set>,<constituent>#<sublattice>) /xxx/: <sf></sf></sublattice></constituent></composition_set></phase>	
	The site fraction (SF) of the constituent shall be given. The default value $xxx$ is the last calculated one.	

## **SET\_START\_VALUE**

Set a start value of an individual variable.

This command is not often required as the automatic start values can be handled for most problems.

Syntax	SET_START_VALUE	
Prompts	State variable: <name a="" of="" state="" variable=""></name>	
	The state variable must be given.	
	Value	
	Specify the value to be assigned the state variable as start value.	

### SHOW\_VALUE

Show the current value of any state variable, function or variable.

Syntax	SHOW_VALUE
Prompt	<pre>State variable or symbol: <name(s) (s)="" of="" or="" state="" symbol="" variable(s)=""></name(s)></pre>
	A single or several state variables or symbols (entered functions or variables) can be specified.
	The wildcard * can be used to denote all phases or all components.
You can use the dollar-sign \$ to denote all stable phases. Thus, the com SHOW W(*,*)lists the mass fraction of all phases, whereas SHOW W(\$, *) mass fraction of all stable phases.	
	If you SHOW a function, all functions are evaluated using the current values of the state variables. But if you SHOW a variable, it retains its value from the time it was ENTERED or last EVALUATED.
	Therefore to save a value from one calculation to another, ENTER it to a variable. This is frequently done in the PARROT module to transfer values between equilibria.

# **STEP\_WITH\_OPTIONS**

The STEP\_WITH\_OPTIONS (short name, STEP) command initiates the stepping procedure to make a calculation of property diagrams in a multicomponent heterogeneous system. A

property diagram is usually stepped under one defined independent stepping axis-variable, and is often constructed by the defined independent mapping axis-variable against a chosen property (for the system, or for a specific phase, or for a certain species).

In the POLY module, it is possible to compute a series of equilibria with one condition as the stepping axis variable. This is achieved by first calculating a single equilibrium, and then selecting one of the conditions as the axis variable. The minimum and maximum value and the incremental step along the axis must also be given. By default, the global minimization technique is used .

- A property diagram consists of property lines that change (normally) along with the stepping axis-variable. From a single STEP calculation, many different types of property diagrams in the defined multicomponent heterogeneous systems can be plotted. Various types of property diagrams are generated by the stepping calculations.
- "About Property Diagrams" on page 187 in the Thermo-Calc User Guide

Normally, before the command can be invoked, you need to have already calculated an initial equilibrium point and set one independent variable (i.e. the controlling condition in the system). This is done using <u>"SET\_AXIS\_VARIABLE" on page 208</u>. You then list the current values of the axis variable for each calculated equilibrium, as well as the corresponding information when the set of stable phases changes.

By default, the global minimization technique is enabled, You can turn this off by using the command-sequence "ADVANCED\_OPTIONS" on page  $167 \rightarrow$  GLOBAL\_MINIMIZATION.

You can choose how often to do a global minimization test using the commandsequence ADVANCED\_OPTIONS  $\rightarrow$  STEP\_AND\_MAP  $\rightarrow$  GLOBAL TEST INTERVAL.

Syntax	STEP_WITH_OPTIONS
	Option? /Normal/: <a chosen="" e,="" i,="" m,="" n,="" o,="" option,="" or="" p="" s,="" t,=""></a>
Prompt	N = Normal, I = Initial equilibria, S = Separate phases, O = One phase at a time, E = Evaluate, M = Mixed Scheil, T= T0, and P = Paraequilibrium

#### NORMAL (N)

When you accept the default, Normal (N), it allows a stepping calculation with the chosen independently-varying equilibrium condition (axis-variable). Only the step axis value is changed between each step.

When you repeat the STEP  $\rightarrow$  NORMAL command-sequence, you can make several stepping calculations while still keep all the stepping-calculation results in the current POLY3 workspaces.

The calculation results are saved in the default RESULT. POLY3 file or you can specify to save it in the <MyName>. POLY3 file. This is as long as there is no SAVE\_ WORKSPACES command before the second set of command sequences (STEP  $\rightarrow$  NORMAL). Repeated stepped results can be restored into the same current POLY3 workspaces/file.

The addition of several steps enables you to generate special types of property diagrams and to calculate and plot special phase boundary lines on phase diagrams, such as:

- The missing part(s) of a specific property [that are calculated under another controlling condition inside the range of the stepping-variable] on the same property diagram.
- Two or more sets of a specific property [that are calculated under different controlling conditions and stepped over the same stepping-variable range] on the same property diagram for the same system.
- Calculate some special phase-boundary lines [especially for some defined secondary phase transformations or alike, e.g. the BCC\_A1 <--> BCC\_B2 phase-boundary, or equal-Gm for two specific phases, or equal-fraction or equal-activity for a certain species in two specific phases, that shall be calculated under different controlling-conditions and also be stepped over different stepping variable(s)], and then plot (or impose) them onto a corresponding phase diagram (which can be resulted from several such STEP → NORMAL calculations or even from a separate MAP calculation) for the same system.

### INITIAL\_EQUILIBRIA (I)

An initial equilibria is added at each calculated point. This can be useful, for instance, to generate a set of isothermal calculations.

When you choose I, an initial equilibrium is stored at each calculated equilibria. It is intended to generate a matrix of calculated initial equilibria, first stepped under a STEP INITIAL\_EQUILIBRIA command-sequence and then repeated by another STEP NORMAL command-sequence (or even continued by a MAP command).

For example, you can first use the temperature condition as the stepping axis-variable with this option, which calculates a number of initial equilibrium points and automatically adds each one as an initial equilibria for another STEP  $\rightarrow$  NORMAL command-sequence (or even for a MAP command). Before the second STEP command (i.e. the STEP  $\rightarrow$  NORMAL

command-sequence), you can chose a compositional condition as the new stepping axisvariable; then the STEP  $\rightarrow$  NORMAL command-sequence uses those created starting equilibrium points (at different temperatures) and steps in such a new compositional variable. As a result, these two STEP commands together give a matrix of values.

### SEPARATE\_PHASES (S)

In this way, you can calculate how the Gibbs energy for a number of phases varies for different compositions. This is particularly useful to calculate Gibbs energies for complex phases with miscibility gaps and for an ordered phase that is never disordered (e.g. SIMGMA-phase, G-phase, MU-phase, etc.).

### ONE\_PHASE\_AT\_TIME (O)

This option is recommended for STEPPING heterogeneous systems with both ordered phases and the disordered pairs.

When you choose  $\circ$ , the stepping calculation is conducted individually for one phase at a time. That is, it is stepped over the whole stepping variable range, but the stepping is repeated for all the entered phases in the defined system. This is particularly useful to properly step in composition for an equilibrium heterogeneous system with both ordered phases and the disordered pairs. Thus, instead of calculating for the same composition for each of the phases at each composition step, it starts the stepping in each of the phases in the default most-stable compositions (the major constitutions).

### EVALUATE (E)

\$

For Scheil-Gulliver simulations it is recommended you use the SCHEIL module.

Evaluate E is an advanced option and is useful when additional conditions (instead of the stepping variable) require changing during a stepping calculation. It allows a stepping calculation in a single axis with simultaneous evaluation of one or more variables after each step. As variables can be used as conditions, it means that you can change the conditions during the stepping. After a successful EVALUATE calculation, you can go to the POST module to manually define and plot property diagrams.

The variables given are evaluated after each step, which may change other conditions. This can be useful to perform the Ordinary ScheilGulliver Approach (non-equilibrium transformations) for simulating alloy solidification without back diffusion of fast-diffusing interstitial components in solid phases. However, it is recommended you use the "SCHEIL Module Commands" on page 290.

Specify the prompt:

Variable name(s): <Variable name>

The names of the variables that shall be evaluated after each step must be typed here.



The miscibility gap test is automatically used during stepping (see <u>"ADVANCED</u> <u>OPTIONS" on page 167</u>) if a phase has two or more composition sets.

The miscibility gap test means that you can start calculating from high temperatures in a steel where the MC carbide is not stable. During the calculation, the MC carbide first becomes metastable with a composition different from the metallic FCC phase and later also becomes stable.



88

See Console Mode example tcex30A.

### MIXED\_SCHEIL (M)

For Scheil-Gulliver simulations it is recommended you use the <u>SCHEIL module</u>.

This is an advanced stepping calculation designed for the Extended Scheil-Gulliver Approach (partial-equilibrium transformations) to simulate alloy solidifications with back-diffusion of one or more fast-diffusing interstitial components (such as C, N, O, S, etc.) in solid phases is performed. It also will consider BCC-->FCC phase transformation (practically for steels) along the alloy solidification process.

To ensure a successful stepping calculation of mixed Scheil-Gulliver simulation of solidification process of a defined alloy system with a certain initial overall composition, it is important to note the following:

- For solution phases with possibility of miscibility gap existence, you have appropriately added the necessary composition set(s);
- You have already set the temperature condition as the stepping SET\_AXIS\_VARIABLE

command, with minimum and maximum temperature points as well as an appropriate s temperature step for the cooling process).

- There must be a composition condition for each of the back-diffusion components (i.e. fast-diffusing interstitial components, such as C, N and/or other elements) that are already defined in terms of mole-fraction or mass-fraction, e.g. X (C) and W (N). Otherwise, a stepping calculation with this option can fail.
- An initial equilibrium in which the LIQUID mixture phase is the only stable must be calculated. This is why it is good to start with a relatively high temperature condition for the initial equilibrium calculation. It is always necessary to reject or suspend the GAS mixture phase (if it exists) before the C-E and STEP commands, to avoid its formation along with LIQUID.
- In cases where fast-diffusing interstitial components are specified as back-diffusion components, you can choose to allow BCC>FCC phase transformation (practically for steels) during the alloy solidification process.

These are the prompts for this option:

```
Fast diffusing components: <Fast diffusion interstitial(s)>
```

Specify one or more interstitial component(s) as the back-diffusion component(s).

- If there is only one fast-diffusing interstitial component (e.g. C or N), type its name;
- If there are two or more fast-diffusing interstitial components (e.g. C, N and S), always type the names on the same line at once (separated by an empty space in between, e.g. C N S);
- If there is no fast-diffusing interstitial component to be considered, type NONE to ignore back diffusion entirely, meaning a normal Scheil-Gulliver simulation is performed.



Such specified back-diffusion interstitial component(s) must be appropriately defined according to the phase constitution definitions of some major alloy solution phases (e.g. FCC, BCC, HCP, etc.). These are located on the interstitial/vacancy sublattices in such alloy solution phases; otherwise, such a Scheil-Gulliver simulation may not make sense at all.

```
Allow BCC -> FCC ? /Y/:
```

By default the BCC-->FCC phase transformation is allowed. This is usually a typical phenomenon along the cooling processes of steels/Fe-alloys.

During the calculation the simulated solidification path, including the locally-equilibrated phase assemblage (region), captured temperature condition (T in K), remaining liquid phase (NL, mole fraction), overall formed solid phases (NS, mole fraction) and latent heat formation (NH, J/mol) along the cooling process are shown after the information on starting temperature point of the alloy solidification process, e.g.

Solidification starts at 1743.15 K				
Phase Region from	1.744150E+03 for:	LIQUID		
Phase Region from	1.742525E+03 for:			
LIQUID				
FCC_A1#1				
1742.5250	0.9960	0.0040	-1.1824	
1742.4000	0.9795	0.0205	-202.1585	
Phase Region from	1.733150E+03 for:			
LIQUID				
BCC_A2				
FCC_A1#1				
1733.1500	0.3294	0.6707	-8032.6240	
1733.0250	0.3237	0.6763	-8095.1490	

After a successful calculation, you can go to the POST module to manually define and plot property diagrams for the alloy solidification process according to the chosen Scheil-Gulliver model, or to impose existing relevant experimental information or other types of calculated solidification results [e.g. Scheil-Gulliver simulation with or without considering fast-diffusing interstitial component(s), full-equilibrium approach, or Diffusion Module (DICTRA)-type simulation with moving-boundary conditions] onto the same plotted property diagrams. Normally, the solidus temperature (T in Celsius) is set as one of the axis variables (usually as Y-axis in most cases), while the other plotted quantity on the other axis can be the amount of overall formed solid alloy phases (NS in mole-fraction or BS in mass-fraction), the amount of remaining liquid phase (NL in mole-fraction or BL in mass fraction), the heat of latent along the solidification process (NH in J/mol or BH in J/gram), among many other properties in the solidified alloy phases or in the whole alloy system.

### T-ZERO (T)



See Console Mode example 41.

This advanced option allows a stepping calculation of the so-called T0 (T-zero) line in a diffusionless transformation (where two specific partially-equilibrated phases have the same Gibbs energy) along a composition variable which has already set as the stepping variable with <u>"SET\_AXIS\_VARIABLE" on page 208</u>.

The temperature condition cannot be set as the stepping variable if you want to make a STEP T-ZERO calculation.



8

Prior to performing the step T0 command sequence, and to ensure a successful calculation of T0 line between two specific phases after an initial equilibrium calculation in the current system, it is sometimes recommended to make a single T0 point calculation using <u>ADVANCED\_OPTIONS</u>  $\rightarrow$  T-ZERO.

Specify these prompts:

```
Name of first phase: <Phase A>
Name of second phase: <Phase B>
```

The names of the target phases, for which the Gibbs energies equal to each other at each point on the T0 line, must be entered.

During the calculation, the TO values are shown after the corresponding scanned conditions (of the stepping composition variable), e.g.

```
Phase Region from
                   1.000000E-01 for:
    BCC A2
    FCC A1
  1.000000E-01
                        940.24
  9.250000E-02
                        941.20
   . . . . . .
  2.500000E-03
                        977.61
  7.500000E-09
                        979.34
Phase Region from 1.000000E-01 for:
    BCC A2
    FCC A1
  1.000000E-01
                        940.24
  1.075000E-01
                        939.62
   . . . . . .
   2.950000E-01
                       1084.87
```

3.000000E-01 1080.99

After a successful calculation, you can go to the POST module to plot the T0 line against the stepping composition variable or another varying axis, or to impose the calculated T0 line onto a normal phase diagram [normally plotted as a T-x isopleth].

### PARAEQUILIBRIUM (P)

See Console Mode examples 42 and 43.

This is an advanced option. Select P to calculate the paraequilibrium state between two specific phases (the *paraequilibrium lines*). It allows a stepping calculation of paraequilibrium lines where the chemical potential for one or more interstitial components (that are considered as back-diffusion elements, such as C and N) but not for the other (s) in two partially equilibrated phases are equal.

More than one interstitial component can be treated as fast-diffusion species (such as C, N, O, S, etc., as individual or combined) in a paraequilibrium stepping calculation.



The composition condition for the interstitial components cannot be set as the stepping variable if you want to make this calculation.

To ensure a successful stepping calculation of paraequilibrium states between two specific phases in a defined alloy system, it is important to first make a starting-point equilibrium calculation with an initial overall composition in the current system. However, it is unnecessary to obtain an equilibrium in which either one or both of the target phases is stable.

The initial overall composition must have a reasonable setting for the desired paraequilibrium calculation for the two target phases. This is especially true where there is more than one interstitial component to be considered in the paraequilibrium state, because different interstitial components (for example C and N combined) may have significant different behaviours as partitioning into different structured phases; otherwise, for one chosen interstitial component the initial overall composition is OK for the paraequilibrium calculation between the specified two phases, but for other chosen interstitial component(s) it might be impossible to calculate the paraequilibrium state.

Always check if the chosen phaseA and phaseB have the exact same definition of elements and if the chosen interstitial components are all in the vacancy sublattice sites of the two phases; otherwise the program cannot find the paraequilibrium state (as it is impossible to correctly calculate u-fractions).

Always have a comprehensive understanding of the normal phase diagram for the currently investigated system, so that you make the appropriate choice of the phase pair and staring bulk composition for the system.

Always set the status of the chosen interstitial components as SPECIAL using the command:

CHANGE\_STATUS COMPONENT <interstitial component> = SPECIAL

By doing this, you get a clear picture on u-fractions of various substitutional and interstitial components, which are different from the overall composition in the system. The SPECIAL status means that specified component(s) are not included in summations for mole or mass fractions. Therefore, all the composition variables plotted from the paraequilibrium calculations are u-fraction related quantities.

Always make a single-point paraequilibrium calculation by the command-sequence of ADVANCED\_OPTIONS  $\rightarrow$  PARAEQUILIBRIUM, prior to performing this STEP  $\rightarrow$  PARAEQUILIBRIUM command-sequence, although this might not be always necessary for some systems.

With this stepping calculation specify the following:

```
Name of first phase: <Phase A>
Name of second phase: <Phase B>
```

8\$B

The names of the two target phases A and B, between which the paraequilibrium states establish, must be typed at the above two prompts subsequently or on the same (first) line at once (separated by an empty space, e.g. FCC BCC, FCC#2 M23C6).

It is important to understand that this is the calculation of a paraequilibrium state between the two specified phases.

Particularly, there are these issues to consider:

- 1. Both chosen phases must have similar interstitial/vacancy sublattices where the fastdiffusion interstitial component(s) occupy.
- 2. The choice on the target phase pair must be reasonable for the defined system and specified initial overall composition.
- 3. Both target phases should have phase constitution definitions that cover all the defined substitutional and interstitial components of the current alloy system; or
- 4. It is impossible to calculate the paraequilibrium state between the target phase pairs with given interstitial component(s) in the currently defined system.

Fast diffusing component: /C/: <Interstitial component(s)>

```
Fast diffusing component: /None/: <Interstitial component(s)>
```

The name(s) of the fast-diffusing interstitial component(s) (c as the default single component) must be given either one by one at each of the above prompts subsequently or in a combination (separated by an empty space in between, e.g.  $c \ge s$ ) at the same (first) prompt; otherwise, press <Enter> key to start the paraequilibrium line stepping-calculation.

Note the following:

Such specified fast-diffusing interstitial component(s) must be appropriately defined according to the phase constitution definitions of the two selected phases: These must be located on the interstitial/vacancy sublattices in both chosen phases.

If there is only one fast-diffusing component which is carbon, press <Enter> to accept the default value (c) at the first prompt; if the single fast-diffusing interstitial component is another element (e.g. N), appropriately type interstitial component the name at the first prompt;

If there are two or more fast-diffusing components (e.g. c and N), type their names at the above prompts subsequently or at the same (first) prompt (separated by an empty space, such as c N);

To finish the input of fast-diffusing elements, accept NONE at a repeated prompt;

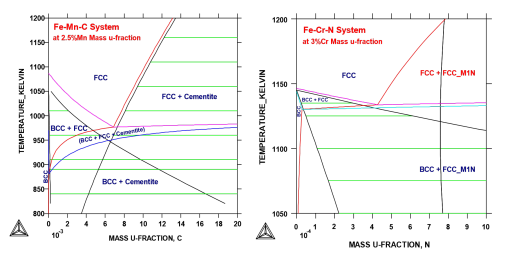
If NONE or a non-existing component name is typed at the first prompt, it means no back diffusion is to be considered, and the para-equilibrium calculation is thus cancelled entirely.

During the STEP  $\rightarrow$  PARAEQUILIBRIUM calculation procedure, the calculated paraequilibrium states include the:

- corresponding stepping conditions (e.g. temperature in K, or the stepping composition variable of the matrix component or of one of substitutional components) which has already set as the stepping variable by the SET\_AXIS\_VAR command);
- amounts of the phase A in mole number [i.e. NP (phaseA)];
- amounts of the phase B in mole number [i.e. NP(phaseB)];
- contents of the interstitial component(s) in the phase A expressed in u-fractions [i.e. u-f(phaseA,component(s))];
- contents of the interstitial component(s) in the phase B expressed in u-fractions [i.e. u-f(phaseB, component(s))];
- LNACR(component) value(s) for the interstitial(s).

After a successful STEP  $\rightarrow$  PARAEQUILIBRIUM calculation, you can go to the POST module to plot a paraequilibrium phase diagram, or to impose the calculated paraequilibrium states onto a normal phase diagram [normally plotted as a T-X isopleth with one of the considered interstitial component(s) as the x-axis variable].

However, the stepping calculation procedure for the currently specified paraequilibrium states between the two specified phases may find difficulties at some steps due to some possible numerical problems (normally regarding to the u-fractions of the interstitial components under some composition ranges), or it may completely fails. In the latter case, it implies that the chosen target phase pair may be unreasonable for the defined alloy system or for the defined initial overall composition, or one or both phases may have inappropriate phase constitution definitions regarding the specified interstitial component(s) and substitutional components. Then, you must either modify the settings of initial overall composition or specify the reasonable target phase pair with an appropriate choice of the fast diffusion interstitials in the defined alloy system.



The figure above shows the plot of a paraequilibrium calculation of isopleths: Formation of para-pearlite in two Fe-based alloys, with equilibrium results appended.

The calculated paraequilibrium phase boundaries (black lines) are shown with tie-lines (in green-colour). In the Fe-2.5Mn-C alloy system, C is considered as fast diffusing interstitial component in the BCC+FCC and FCC+Cementite paraequilibrium stepping calculations; while in the Fe-3Cr-N alloy system, N is treated as fast diffusing interstitial components in the BCC+FCC and FCC+FCC\_M1N (i.e. FCC#1+FCC#2) paraequilibrium stepping calculations.

During a STEP calculation, the value of the stepping axis variable for each calculated equilibrium is listed and also the set of stable phases.

You can terminate the mapping of a line by pressing a single <Ctrl-A> (Windows) or <Ctrl-C> (Linux). This can be useful in order to stop a longish calculation without losing what is already calculated.

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If there is any convergence problem in finding a stable solution at any stage during a calculation procedure enforced by a STEP\_WITH\_OPTIONS command-sequence, this message displays:

Convergence problems, increasing smallest site-fraction from 1.00E-30 to hardware precision 2.00E-14. You can restore using SET-NUMERICAL-LIMITS

This implies that smallest site fraction in the current POLY3 workspace is automatically increased from the default value 1.00E-30 to the hardware-dependent precision (under Linux, as 2.00E-14). For other subsequent POLY-module calculation in the current TC run, you can use the POLY command <u>"SET\_NUMERICAL\_LIMITS" on page 293</u> to restore or reset the smallest site fraction to the previous or another preferred value, as well as to reset other numerical limits.

Complex miscibility gaps of various solution phases are automatically detected during a stepping calculation, and two or more composition sets for each of such solution phases are automatically generated as well, through an automatic Global Minimization Test procedure (by specifying a Global Test Interval value and confirming the Automatically add initial equilibria in the ADVANCED\_OPTION STEP\_AND\_MAP command-sequence). As a particular case, such an automatic Global Minimization Test ensures that you can now start calculating from high temperatures in an austenitic steel (metallic FCC\_A1 solution phase) where the MC carbides/nitrides/carbonitrides (i.e. the C-/N-rich sides of the FCC\_A1 miscibility-gap, often (while not always) being referred to as FCC\_A1#2, FCC\_A1#3, etc.) are not stable, and during the MAP command the MC carbides/nitrides/carbonitrides may first become metastable with a composition different from the metallic FCC\_A1 phase and later also stable.

Occasionally, it may be necessary to have a specified increment other than the default value for the defined stepping axis-variable, in order to calculate/plot a smoother property diagram.

To get a complete property diagram, sometimes (especially in some earlier Thermo-Calc versions) it may also be required to have more than one starting equilibrium-points (under another starting equilibrium condition), and repeat the stepping calculation over the same stepping axis-variable range (while without over-writing the same POLY3 workspaces via a SAVE\_WORKSPACE command). However, using the current STEP procedure such a requirement become less necessary.

## **TABULATE**

For any entered table, this command gives a table of values from equilibria calculated with a STEP command. It works only for a table that is entered in either the POLY or POST module.



Unlike the TAB module's command (also called <u>"TABULATE" on page 311</u>) it is not possible to plot columns from the tabulated tables. For this use normal POST commands.

Syntax	TABULATE
Prompts	Name: <name a="" either="" entered="" in="" of="" or="" poly="" post="" table=""></name>
	Specify a table name that is entered in either the POLY or POST module.
	Output on screen or file /Screen/:
	Specify a file name if you want to save on a file the table values along the defined STEP calculation, or press <enter> if want to see the table values displayed.</enter>

### Plotting and Visualization with the POST Module

The POST module (post-processor) is available from the POLY module. See <u>"Equilibrium</u> <u>Calculations with the POLY\_3 Module" on page 157</u> for other information about the POLY module. Read <u>"Using the POST Module" on page 233</u> before beginning.

#### In this section:

Using the POST Module	. 233
ADD_LABEL_TEXT	234
APPEND_EXPERIMENTAL_DATA	.236
CHANGE_LEGEND	237
DUMP_DIAGRAM	.237
ENTER_SYMBOL	238
FIND_LINE	. 241
LIST_DATA_TABLE	.241
LIST_PLOT_SETTINGS	. 242
LIST_SYMBOLS	242
MAKE_EXPERIMENTAL_DATAFILE	. 243
MODIFY_LABEL_TEXT	243
PLOT_DIAGRAM	244
PRINT_DIAGRAM	244
QUICK_EXPERIMENTAL_PLOT	. 245
REINITIATE_PLOT_SETTINGS	. 245
RESTORE_PHASE_IN_PLOT	. 245
SELECT_PLOT	. 246
SET_AXIS_LENGTH	.246
SET_AXIS_PLOT_STATUS	. 246
SET_AXIS_TEXT_STATUS	.247
SET_AXIS_TYPE	. 247

SET_COLOR	
SET_CORNER_TEXT	249
SET_DIAGRAM_AXIS	249
SET_DIAGRAM_TYPE	252
SET_FONT	252
SET_INTERACTIVE_MODE	
SET_LABEL_CURVE_OPTION	
SET_PLOT_FORMAT	254
SET_PLOT_OPTIONS	255
SET_PLOT_SIZE	
SET_PREFIX_SCALING	
SET_RASTER_STATUS	
SET_REFERENCE_STATE	258
SET_SCALING_STATUS	
SET_TIC_TYPE	
SET_TIELINE_STATUS	
SET_TITLE	
SET_TRUE_MANUAL_SCALING	
SUSPEND_PHASE_IN_PLOT	
TABULATE	

# **Using the POST Module**

Use the POST module after a <u>"STEP\_WITH\_OPTIONS" on page 217</u> or <u>"MAP" on page 201</u> calculation to generate *graphical* or *tabulated* presentations on screen or into files.

The graphical files are available in different formats (e.g. PS, EMF, PNG, BMP, PDF, JPG, TIF) or data forms (e.g. EXP, TAB and WRL). The graphical output can be phase diagrams, property diagrams, diffusion profiles and other types of plots. The tabulated presentation is into a basic text file (TXT) or a spreadsheet (e.g. an MS Excel file with an extension of .XLS).

The POST module is available through these modules:

- POLY
- TAB
- BIN
- TERN
- POT
- SCHEIL
- POURBAIX

Within the POST module, you can choose any state variable, any derived variable, or any entered symbol (functions or variable) as the X/Y-axis.

#### "Thermodynamic Variables and Units" on page 327

When a diagram is plotted, appearance parameters for defining a high-standard graph can be further specified, e.g. curve labelling options, diagram titles and subtitles, plot size, axis length, axis types, axis-tic type, tie-line status, automatic or manual scaling and zooming, semi-automatic or manual labelling on phase boundaries and phase regions, graphic formats, text fonts, colours, raster plot, etc.

You can append experimental data onto a plotted diagram. You can also save the coordinates onto a textual file, which can be edited and used as an experimental data file to be merged onto another diagram or as a part of a setup file for PARROT-module assessment.

Phases can be suspended or restored in a plotted diagram. The reference states for components can also be modified for the resulted diagrams. Moreover, you can translate a plotted property diagram (after a stepping calculation) into a tabulated form, such as a simple list on screen or a textual file, and a spreadsheet .

# ADD\_LABEL\_TEXT

Add a text to an area in a phase diagram or a property diagram. The label is written starting from the specified coordinates. Optionally, you can also let the program automatically add the text by first calculating an equilibrium state at the specified coordinates and then making a label out of the stable phase names.

The label texts and coordinates, either added by this command or modified with <u>"MODIFY</u> <u>LABEL\_TEXT" on page 243</u> (or manually edited in the **Console Results** window), are stored in the workspace, and these may later be saved onto an \*.EXP file (with <u>"MAKE\_</u> <u>EXPERIMENTAL\_DATAFILE" on page 243</u>).

In some cases such an optional calculation for automatic phase region labeling may not work. It is not possible to automatically add labels for certain sets of axes, like entropy, enthalpy, pH, Eh, etc.

<u>"Plot Labels " on page 137</u> in the Thermo-Calc User Guide.

Syntax	ADD_LABEL_TEXT	
Prompts	Give X coordinate in axis units: <value coordinate="" of="" the="" x=""></value>	
	Specify the value of the X coordinate where the label is to start.	
	Give Y coordinate in axis units: <value coordinate="" of="" the="" x=""></value>	
	The labels disappear if any axis is changed after wards.	
	Specify the value of the Y coordinate where the label is to start.	
	Automatic phase labels? /Y/:	
	By default the program automatically calculates the equilibrium at the given coordinates and generates a label with the names of the stable phases. However, the automatic calculation works only for phase diagrams that are mapped with two axes in the POLY module, and sometimes it may not work correctly especially for complex heterogeneous interaction systems; in such cases, a message displays.	
	Text: <text for="" label="" the=""></text>	
	If you answer N enter text at the prompt. You can enter simple text, use LaTeX, rotate the text, as well as adjust the font size.	
	LaTeX: You can use LaTeX commands to enter text and symbols. First enter the	

Syntax	ADD_LABEL_TEXT	
	command followed by the expression or string, for example to di subscripts:	splay Al2O3 as
	\latex Mole-fraction Al_20_3	
	For a list of supported LaTeX commands, see <u>http://www2.ph.ed.ac.uk/snuggletex/documentation/m</u> <u>mode.html</u> .	ath-
	By default when using the LaTeX command, you are in a and Thermo-Calc automatically interprets the entered to math. The above link uses the dollar sign (\$) in all the e this is NOT required. For example, if you enter \$E=mc^2 displayed including the \$ instead of the equation.	text as xamples but
	To add a mixture of plain text and equation text, you can mode using the string \textrm followed by the text to conclosed in curly brackets. For example \latex \textr fraction }A1_20_3 displays this label Mole-fraction $Al_20_3$ .	lisplay
	<b>Rotate the text</b> : If you want to rotate the label, start a text string rotation command (which is case sensitive and must be in lower then enter the number of degrees to rotate the label. For examp label 45 degrees enter:	case letters)
	\rotation 45	
	To rotate the text and use LaTeX, the rotation command must be example:	used first. For
	\rotation 45 \latex Mole-fraction Al_20_3	
	Rotates the text 45° and includes subscripts in the title Mole-frac	tion Al2O3.
	Text size: /.35/: <size for="" label="" the=""></size>	
	A smaller size of the text may be necessary to fit the label text in A default size is given using the latest size in adding label texts, o command is called for the first time. A recommended size is e.g.	or .35 if the

### Syntax ADD\_LABEL\_TEXT

These changes to plot labels can also be manually edited in the **Console Results** window.

# APPEND\_EXPERIMENTAL\_DATA

Add experimental data and text onto a calculated diagram. This is done by placing the experimental data and text on a formatted textual file (always having an extension of EXP) that is prepared outside of Thermo-Calc software (Console Mode and Graphical Mode), according to the syntax of the DATAPLOT Graphical Language.

The picture generated from the data is superimposed on the ordinary graphical output from the postprocessor. Such an experimental data file (\*.EXP) may be created by an ordinary textual editor.

Another important usage of this command is to superimpose plots from several independent calculations. For this purpose, there is a command <u>"MAKE\_EXPERIMENTAL\_DATAFILE" on page 243</u>, which dumps a calculated diagram on an EXP file according to the DATAPLOT syntax. With the aid of a normal text editor, many such files may be merged together. Remember that you can only have one prologue section on each such EXP file.

**DATAPLOT** User Guide included with this documentation set.

Syntax	APPEND_EXPERIMENTAL_DATA	
	Use experimental data (Y or N) /N/:	
Prompts	By default, no experimental data are plotted. Enter Y to use the data from an experimental data file in the next plot. An <b>Open</b> window displays to specify the name of the file with the experimental data.	
	Prologue number: /0/: <prologue number=""></prologue>	
	Select which prologue to use. In a prologue, you can, for example, give the scaling of an axis, the axis texts, and so on.	
	By answering -1, a list of all prologues on the file is given; and by 0, no experimental data are appended.	
	<pre>Dataset number(s) /-1/: <dataset number(s)=""></dataset></pre>	
	Select from which dataset(s) data should be read. Several datasets may be given separated with commas or spaces. By answering -1, a list of all datasets on the file is given; and by 0, no experimental data are appended.	

# CHANGE\_LEGEND

When there is a legend in the diagram you can append constitution and/or ordering descriptions to the phase name.



8

"LIST\_EQUILIBRIUM" on page 198 also displays ORD (ordered) or DISORD (disordered) (when available).

In some cases the legend only shows the phase name for a line when the <u>"SET</u> <u>LABEL\_CURVE\_OPTION" on page 253</u> option is set to F.

SyntaxCHANGE\_LEGENDPromptSet suffix for phase names in legend? (All, None, Constitution,<br/>Ordering)/None/:Prompt• None: the legend includes the standard information about the phases.<br/>• All: the Constitution description and the Ordering description (when<br/>there is ordering) are also included.<br/>• Constitution: it is added to the standard description.<br/>• Ordering (when applicable): it is added to the standard description.Image: Constitution of the standard description of the standard description of the standard description.<br/>• Ordering (when applicable): it is added to the standard description.

## **DUMP\_DIAGRAM**

After a Thermo-Calc graph is created (and displays with the PLOT\_DIAGRAM command) you can save (dump) a diagram to a file so you can quickly print it. It is similar to the PRINT\_DIAGRAM command.

	-	_	- 1	
13	-	_		
	-	-	/	2

Similar to <u>"PRINT\_DIAGRAM" on page 244</u>, it does not matter what graphic device is set up for the graph, this command is always able to convert the graph shown on screen to the selected graphic format.

Syntax	DUMP_DIAGRAM
Deserved	Output format/: <format or="" return=""></format>
Prompt	Specify an appropriate graphic format for dumping the graph or pressing

### Syntax DUMP\_DIAGRAM

<Enter> to accept the default format.

A **Save** window opens if you have not specified a file name. Select a directory path and enter a **File name**. If a file with the specified name already exists, then you are prompted whether to overwrite the old file or not. The third argument of the command (Y) allows you to specify whether to overwrite the old file or not (N).

After the printing process is finished, the temporarily set graphic device for conversion and printing is switched back to the one set by the command <u>"SET\_PLOT\_FORMAT" on page 254</u>, so that you can save the graph in file(s) using "PLOT\_DIAGRAM" on page 244.

## **ENTER\_SYMBOL**

8

>\_

This command is for both the POST and POLY modules in Console Mode.

Symbols are a useful feature of the POLY and POST modules to define quantities that are convenient. Symbols can be constants, variables, functions or tables.

Functions or tables (with defined functions as values) can be entered in the POST module after a stepping or mapping calculation, for purposes of plotting such entered functions or tables as axis variables.

Within the POLY module, symbols are normally defined prior to an equilibrium calculation (enforced by a C\_E command), stepping calculation (enforced by the command <u>"STEP\_WITH\_OPTIONS" on page 217</u>) or mapping calculation (enforced by <u>"MAP" on page 201</u>). These can be entered after an equilibrium calculation; however, for defined functions, variable or tables, it requires using <u>"EVALUATE\_FUNCTIONS" on page 197</u> before showing the corresponding values in the calculated equilibrium state.

The symbols entered in the POST module are not saved in the currently-loaded POLY3 workspaces. Therefore, if you want to apply such symbols in other similar calculations for the same defined system, you must use the ENTER\_SYMBOL command prior to the STEPPING or MAPPING calculation in the POLY module.

**<u>"ENTER\_SYMBOL" on page 109</u>** for the GIBBS Module.



See <u>example 44</u> in the *Thermo-Calc Console Mode Examples Guide* for an example of using variables and functions.

Syntax	ENTER_SYMBOL
	Constant, Variable, Function or Table? /Function/: <keyword></keyword>
	The Keyword can be specified as CONSTANT, VARIABLE, FUNCTION or TABLE.
	• CONSTANTS can only be entered once and is a means of using a name for a numeric value. For example, the value of 1 atmosphere in Pascal can be denoted by P0 after the command ENTER CONSTANT P0=101325. Defined constants can be used as values in condition assignments, for example, SET-COND P=P0.
	<ul> <li>FUNCTIONS are expressions of state variables or other functions. These expressions are saved, and whenever a function value is requested all functions are evaluated. The reason for this is that they may depend on each other.</li> </ul>
Prompt	<ul> <li>VARIABLES are similar to functions because they can also be expressions of state variables. However, contrary to functions, they are only evaluated when they are entered or if they are explicitly named in an <u>"EVALUATE_FUNCTIONS" on page 197</u> command. It is possible to enter a variable with a new expression any time. This expression is evaluated directly and its value stored as the value of the variable. Defined variables can be used as values in the SET-CONDITION command.</li> </ul>
	• TABLES are used for listing results from the STEP or MAP commands. A table consists of a list of any number of state variables, functions or variables. Defined tables can also be used in the post-processor POST.
	There is a special connection between tables and variables. If a variable is used in a table, it is evaluated for each line of the table in the TABULATE command or when the table is used in a plot.
	Name: <name of="" symbol="" the=""></name>
	Each symbol has a unique name that must start with a letter and can have maximum 8 characters.
	Legal characters include letters (either UPPER or lower case), digits and underscore Any other special character, such as parentheses ( and ), plus +, minus -, slash / or  full stop (.), are illegal for symbol names.
	You can enter the symbol name and the value or function on the same line, these must be separated with an equal sign =, for example, $TC=T-273.15$ or $T_c=T273.15$ which stands for a definition of temperature in Celsius. Otherwise, these questions are asked.

Syntax	ENTER_SYMBOL
	For different types of symbol (constant, function, variable or table), the questions have different prompts.
	Function: <definition a="" for="" function="" or="" variable=""></definition>
	<i>Functions</i> and <i>variables</i> are evaluated from an expression of state variables or other functions, constants or variables. The expression is a FORTRAN-like expression and operators +, -, *, = and ** can be used (** only with integer powers).
	Unary functions like LOG, LOG10, EXP, SIN, COS and ERF can also be used. An expression can be continued on more than one line. An expression should be terminated by a semicolon (;) or an empty line (by pressing <enter> at the next prompt).</enter>
	Examples of function expressions:
	• GM(LIQUID): The Gibbs energy of liquid per mole component
	• H.T/4.184: The heat capacity of the system in calories
	• ACR (CR) /X (FCC, CR): The activity coefficient for Cr in FCC
	• T-273.15: The temperature in Celsius
	&: <continuation definition="" for="" of="" symbol="" the=""></continuation>
	The ampersand & means that you can continue to write the function on the new line if one line is not enough for the function. If you finish the function press <enter> again.</enter>
	Value: <value a="" constant="" for=""></value>
	A constant can only be assigned a numeric value once.
	Value or expression: <value a="" expression="" for="" of="" variable=""></value>
	A <i>variable</i> can be assigned a numeric value or an expression. An expression is evaluated immediately and discarded. Only the numeric value is kept. This gives a possibility to save a value between calculations with different conditions because all state variables and functions are evaluated for the new conditions.
	<pre>Variable(s): <variable(s) a="" in="" table=""></variable(s)></pre>
	A <i>table</i> consists of a list of state variables or functions. One way to obtain results from a STEP command is through a table.
	Example:

#### Syntax ENTER\_SYMBOL

ENTER TABLE K=T,X(LIQ,C),X(LIQ,CR),ACR(C)

Which means that the table called K contains four columns, i.e. the temperature, the mole fractions of C and Cr in the LIQUID phase, and the activity of C.

To show the temperature in Celsius in a table, give the command ENTER FUNCTION TC=T-273; and then use the symbol TC in the table.

& <Continuation of the definition for the table>

The ampersand & means that you can continue to write the table on the new line if one line is not enough for the table. If you finish the table press <Enter> again.

## FIND\_LINE

Identifies various property curves on a plotted property diagram (after a stepping calculation); it also works well in find stable phase regions on a phase diagram (after a mapping calculation).

Syntax	FIND_LINE
	X coordinate /.484623611/: <value coordinate="" of="" the="" x=""></value>
Prompts	Specify the value of the X coordinate where you want to find the details of property curve (or phase region).
	Y coordinate /.5153737962/: <value coordinate="" of="" the="" y=""></value>
	Specify the value of the Y coordinate where you want to find the details of property curve (or phase region).
	The POST module displays the identified property (for a property curve on a property diagram), or the identified stable phase-assemblage (for a phase region on a phase diagram).

### LIST\_DATA\_TABLE

>\_

This is a POST module command.

This command is used for listing various properties (i.e. the Y-axis variable on a property diagram) that change with an independent variable (i.e. the X-axis variable on a property diagram), which have already been defined in the POST module (and often are plotted on the **Console Results** window), but only after a STEP calculation.

The listing output can be either on screen or in a spreadsheet file in the MS Excel format (under specified file name with the default file extension of XLS), which can be further edited or used for graphical handling by opening the file (using the MS-Excel program) outside of the Thermo-Calc software system.

Syntax	LIST_DATA_TABLE
	Output to screen or file /Screen/: <file name="" or="" return=""></file>
Prompt	Specify a file name if you want to save on a spreadsheet file (in the MS Excel format, with the default extension of XLS) on which the already-defined properties on a property diagram (after a STEP calculation) is saved, or press <enter> if want to see a simple textual table on screen (for the already-defined properties that change alone with an independent variable).</enter>
	This command only works for a property diagram after a stepping calculation but not for any phase diagram after a mapping calculation.

## LIST\_PLOT\_SETTINGS



This is a POST module command.

### Syntax LIST\_PLOT\_SETTINGS

Lists the present values of most parameters specifying the type of diagram to be plotted.

# LIST\_SYMBOLS



For both the POLY and POST modules, list the definitions for all constants, functions, variables and tables.

In order to find the value of a function or variable, use <u>"SHOW\_VALUE" on page 217</u> or <u>"EVALUATE\_FUNCTIONS" on page 197</u>. A table is tabulated with the TABULATE command.

### Syntax LIST\_SYMBOLS

The defined variables are listed up together with the defined functions, but variable names are followed by a percentage sign %.

# MAKE\_EXPERIMENTAL\_DATAFILE

Write the graphical information onto screen or a formatted text file (always having an extension of .EXP) using the DATAPLOT graphical format. In order to merge two or more diagrams from separate calculations, you can write them out on files with this command, and then add/edit them together with a normal text editor.

**DATPLOT User Guide** included with this documentation set.

### Syntax MAKE\_EXPERIMENTAL\_DATAFILE

Output to screen or file /Screen/: <Name of a file or return>

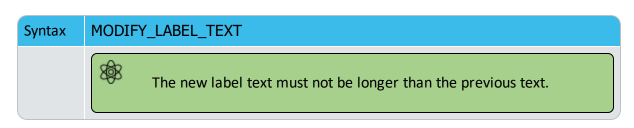
Prompt Specify the name of the desired file. Press <Enter> for a list. Otherwise enter a file name where the graphical information is written. The default file extension is EXP (for Windows) or exp (for Linux).

# MODIFY\_LABEL\_TEXT

Move a label created with the command <u>"ADD\_LABEL\_TEXT" on page 234</u> to another position, or replace its text with another one.

The label texts and the coordinates, either added by ADD\_LABEL\_TEXT command or modified by this command or manually edited on the Console Results window, are stored in the workspace, and can be saved onto an \*.EXP file (through <u>"MAKE\_EXPERIMENTAL\_</u>DATAFILE" above).

Syntax	MODIFY_LABEL_TEXT
	Which label to modify? /#/: <number label="" of="" the=""></number>
Prompts	Before this prompt, all the labels created by the ADD_LABEL_TEXT command is listed with an identifying number. Specify the Number of the label you want to modify. The default one (#) is the last added label.
	New X coordinate: /xxx/: <new position="" x=""></new>
	Specify the ${\tt New \ X \ position}.$ The previous X coordinate is displayed. Press <enter> to accept.</enter>
	New Y coordinate: /yyy/: <new position="" y=""></new>
	Specify the New Y position. The previous Y coordinate is displayed. Press <enter> to accept.</enter>
	New text /ABCDEFGH/: <new labeling="" text=""></new>
	Specify the New labeling text. The previous text is displayed. Press <enter> to accept.</enter>



# PLOT\_DIAGRAM

With this command, a calculated/plotted diagram is displayed if the default graphic device has already been defined by the <u>"SET\_PLOT\_FORMAT" on page 254</u> command. It can also be saved onto a file.



The diagram axes must be set first.

The functionality of this command is shared in two ways:

- in PLOT\_DIAGRAM for normal screen display and file saving and
- in <u>"PRINT\_DIAGRAM" below</u> (under MS-Windows environment) for direct hard-copy printing.

Syntax	PLOT_DIAGRAM	
Prompt	Output to screen or file /Screen/: <name a="" file="" for="" of="" or="" return="" screen=""></name>	
	Specify the name of the desired file (Note that under Linux, the prompt is PLOTFILE /SCREEN/). Consequently, a file with a proper extension for the selected graphic format (set by the SET_PLOT_FORMAT command) is saved under the current working directory. If no extension is given in the file name, the default extension e.g. ps for Postscript portrait/landscape modes, P7 or p7 for HPGL landscape, P8 or p8 for HPGL portrait, etc.) is automatically given to the file name. If the default device is set (by the command SET_PLOT_FORMAT), press <enter></enter>	

### PRINT\_DIAGRAM

Syntax PRINT\_DIAGRAM

Print with high quality to a file after a Thermo-Calc graph is created (and is displayed with <u>"PLOT\_DIAGRAM" above</u>). It has the same function as the Print Plot

Syntax PRINT\_DIAGRAM

option on the Console Results windows.

## QUICK\_EXPERIMENTAL\_PLOT

This is similar to <u>"APPEND\_EXPERIMENTAL\_DATA" on page 236</u> but can be used when there is no graphical information to be plotted in the POLY3 workspace. It defines a pair of axis, sets the axis labels to X and Y, and scales both X- and Y-axes between 0.0 and 1.0 unless a prologue is read from the DATAPLOT (\*.EXP) data file.

"Plotting Experimental Data Files" on page 4 in the DATPLOT User Guide included with this documentation set.

Syntax	QUICK_EXPERIMENTAL_PLOT	
	Prologue number: /0/: <prologue number=""></prologue>	
Prompts	An <b>Open</b> window displays. Select the prologue to use. In a prologue, you can, for example, give the scaling of an axis, the axis texts, and so on. By answering -1, a list of all prologues on the file is given; and with 0, no experimental data are appended.	
	<pre>Dataset number(s) /-1/: <dataset number(s)=""></dataset></pre>	
	Select from which dataset(s) data should be read. Several datasets may be given separated with commas or spaces. By answering -1, a list of all datasets on the file is given; and with 0, no experimental data are appended.	

### **REINITIATE\_PLOT\_SETTINGS**

All parameters describing the diagram are given default values.

### Syntax REINITIATE\_PLOT\_SETTINGS

All plot settings made in the POST module are deleted, and it returns to the initial settings when the POST module is entered.

### **RESTORE\_PHASE\_IN\_PLOT**

Restores a phase previously suspended from the plot with <u>"SUSPEND\_PHASE\_IN\_PLOT" on</u> page 262. It works only after at least one phase is suspended.

Syntax	RESTORE_PHASE_IN_PLOT	
Prompt	Phase name: <name a="" of="" phase=""></name>	

Syntax	RESTORE_PHASE_IN_PLOT
--------	-----------------------

Give the name of the phase to be restored.

# SELECT\_PLOT

Select a plot within the current Results tab in the Console Results window.

Syntax	SELECT_PLOT
	Plot <n new=""></n>
Prompt	Where $N$ is a positive integer. Enter the number of the plot that is selected in the <b>Console Results</b> window (in the currently selected <b>Results</b> tab). If a plot with that number does not exist, it is created.
	The default parameter value is NEW. If this is entered, then a new plot is created. Create a new plot can also be done by pressing <ctrl+shift+t>.</ctrl+shift+t>

## SET\_AXIS\_LENGTH

Change the real length of an axis in inches. The default number of tic-marks on an axis is 10 when the relative length is 1. The number and units per tic-mark must be a multiple of 1, 2 or 5 to obtain a reasonable scaling of an axis.

Syntax	SET_AXIS_LENGTH
<u> </u>	Axis (X, Y or Z): <name an="" axis="" of=""></name>
Prompts	Specify which axis to set the length.
	Axis length /11.5/: <new axis="" length="" relative=""></new>
	Specify the new real axis length in inches. The relative length 1 corresponds to 10 tic-marks on the axis.

# SET\_AXIS\_PLOT\_STATUS

Specify whether the axis title texts and axis label texts are to be plotted or not on a diagram. It does not work for the axis lines and tic-marks. If you want to plot a diagram without tic-marks, use "SET\_TIC\_TYPE" on page 260 first.

This can be used to merge different diagrams on a pen-plotter or to obtain the diagram faster. The default status is that all the set axes are to be plotted.

SyntaxSET\_AXIS\_PLOT\_STATUSPromptAxis plot (Y or N) /Y/: (Y or N)

# SET\_AXIS\_TEXT\_STATUS

Change the axis text from the automatic text given by the axis specification to a text.

"ADD\_LABEL\_TEXT" on page 234

Syntax	SET_AXIS_TEXT_STATUS
Prompts	Axis (X, Y or Z): <name an="" axis="" of=""></name>
	Specify which axis text status to change.
	Automatic axis text (Y or N) /N/:
	Specify if automatic axis text is to be used.
	Axis text: <enter axis="" text=""></enter>
	You can enter simple text, use LaTeX, rotate the text, as well as adjust the font size.

## SET\_AXIS\_TYPE

Change the axis type among linear, logarithmic and inverse axis.

Syntax	SET_AXIS_TYPE
Deserves	Axis (X, Y or Z): <name an="" axis="" of=""></name>
Prompts	Specify which axis to change the axis type.
	Axis type /Linear/: <new axis="" type=""></new>
	Specify which new axis type to set. Select LINear (default), LOGarithmic or INVerse. Only the three first characters are relevant.

## **SET\_COLOR**

Some devices support colours or line thickness. With this command, select different colors or line types on some types of information on a diagram.



This command can fail if a chosen device does not support colors or line thickness.

This command gives four sequential prompts (i.e. Text and axis, Invariant equilibria, Tielines, and All other lines), with the Keyword and default option (both listed below). Depending on if the switched-on graphic device (by the SET\_PLOT\_FORMAT command) supports colors or line types, the POST automatically switches on the appropriate Keyword.

Syntax	SET_COLOR		
Prompt	Text and axis keyword /Default option/		
	<return new="" option="" or=""></return>		
	Invariant equilibria keyword /Default option/: <return new="" option="" or=""></return>		
	Tie-lines color keyword /Default option/: <return new="" option="" or=""></return>		
	Keyword of all other lines /Default option/: <return new="" option="" or=""></return>		
	This is usually for the solubility lines.		
	The Keyword is either Color or LineType, depending on if color or line type is supported by the switched-on graphic device (by the <u>"SET_PLOT_FORMAT" on page 254</u> command); the POST module <i>automatically</i> switches on the appropriate Keyword.		

#### **COLOR OPTIONS**

The options for Color are:

BACKGROUND	BLUE	PURPLE	GRAY
FOREGROUND (default)	YELLOW	GOLD4	ORANGERED3
RED (default for invariant equilibria)	MAGENTA	TURQUOISE4	MAROON
GREEN (default for tie lines)	CYAN	PINK	PLUM
	SIENNA	OLIVEDRAB	SEAGREEN
	ORANGE1	CORAL1	USERDEF

#### LINETYPE OPTIONS

The options for LineType are:

INVISIBLE	VERY_THICK (default for invariant equilibria)
NORMAL (default)	THIN (default for tie lines)
ТНІСК	VERY_THIN
DASHED	DOTTED

## **SET\_CORNER\_TEXT**

Adds text to the corners on a plotted diagram. Normally, you can write such texts as a subtitle.

The primary database (not the appended ones) used in calculations and the calculated conditions are automatically plotted at the upper-left corner as a part of the header on all plotted diagram. Unless the plotting option WRITE CONDITIONS is switched off by the command <u>"SET\_PLOT\_OPTIONS" on page 255</u>, the calculated conditions are always written on plotted diagrams; and only when the plotting option PLOT HEADER is switched off by the command SET\_PLOT\_OPTION, the used database also always appear. Under such circumstance, avoid writing texts at the UPPER\_LEFT corners.

Syntax	SET_CORNER_TEXT
	Corner/Lower_Left/: <return new="" option="" or=""></return>
	Choose an option:
	• LOWER_LEFT
Prompt	UPPER_LEFT
	UPPER_RIGHT
	LOWER_RIGHT
	TOP_OF_TRIANGLE
	Text: <texts a="" as="" note="" or="" subtitle=""></texts>
	Write the texts that are to be written on the specified diagram corner.

# SET\_DIAGRAM\_AXIS

Specify an axis variable for a plot or a listing. To be able to plot a diagram, at least two axis variables (X and Y) must be specified. In Console Mode, three axes (X, Y, and Z) can be specified for a diagram.



The axis variables in the plot can be different from the number of axis used for the mapping.

If you want to plot a phase diagram with tie-lines in the plane after a MAP calculation, then the composition axis must be mole-fraction or weight-fraction, or mole-percent or weight-percent of one specific component. You must not use the stable variable x(comp) or w(comp) even if that is what you used when calculating the diagram, because that only gives one side of the two-phase region. It may be sufficient to indicate that mole-fraction is the same as the stable variable x(\*, comp), and weight-fraction is the same as the stable variable x(\*, comp).

If you plot a phase diagram where the tie-lines are not in the plane after a MAP calculation, then mole-fraction and x(comp) are identical.

When you plot a property diagram after a STEP calculation and want a composition axis variable, you should normally use x(comp) and not mole-fraction.

The axis must have exactly the same number of columns, or one axis must have one column. In the first case the columns are plotted matching them one by one; in the latter all columns on one axis is plotted against the single column. For example, you can have the temperature on one axis and the amount of each phase on the other. The amount of each phase is the state variable NP(\*) in mole fraction or BPW(\*) for mass fractions of phases.

Automatic diagram axis: The POST module can set automatic diagram axis identical to those used in the <u>"MAP" on page 201</u> command.

If the state variable x(comp) is used in a MAP calculation, then mole-fraction of the component is used as diagram axis; if a potential or some other state variables are used, then that is used as a diagram axis.

After a STEP calculation, automatic diagram axis is not set as there is only one axis used in the calculation.

Syntax	SET_DIAGRAM_AXIS
	Axis (X, Y or Z): <axis name=""></axis>
	Specify which axis (X or Y or Z) to set with a variable.
	Variable type: <variable name=""></variable>
	Specify a variable for the chosen axis here.
Prompts	The valid variables are:
	<ul> <li>TEMPERATURE-CELSIUS as temperature in °C</li> </ul>
	TEMPERATURE as temperature in K

Syntax	SET_DIAGRAM_AXIS
	PRESSURE as pressure in Pa
	ACTIVITY for a component
	<ul> <li>LNACTIVITY for a component [In(ACR(component)]</li> </ul>
	MOLE-FRACTION for a component
	<ul> <li>MOLE-PERCENT for a component</li> </ul>
	<ul> <li>WEIGHT-FRACTION for a component</li> </ul>
	WEIGHT-PERCENT for a component
	<ul> <li>ENTHALPY as enthalpy in J/mol</li> </ul>
	ENTROPY as entropy in J/mol/K
	NONE to clear the axis
	<ul> <li>Any valid state variable including those with wildcards, e.g. NP(*) or x (*,component). Due to the use of wildcard * in such a state variable, the program asks for a column number.</li> </ul>
	Any entered function or variable.
	Any entered table.
	If you specify an entered table as the variable, then you are given these prompts:
	For component: <component name=""></component>
	When an activity, mole or weight fraction or percent of a component shall be plotted, the component name must be supplied here.
	Column number: <column number(s)=""></column>
	Specify the column number(s) in the chosen table that are to be plotted onto the earlier specified axis. For example:
	• 1 for column 1
	• 2, 3 for columns 2 and 3
	• 2, 3 >5 for column 2,3 and all columns above column 5
	• * for all columns

# SET\_DIAGRAM\_TYPE

Choose the diagram type as perpendicular plot or triangular plot (Gibbs triangle, especially for ternary systems). The default is with a perpendicular axes.

For phase diagrams of ternary or pseudo-ternary systems, it is usually need to plot isothermal sections as triangular grams. If desired, all lines outside the region limited by a line joining the end points of the X- and Y-axis is removed.



In order to create a 3D-plot file (in the VRML format, \*.WRL) for a tetrahedron diagram, this command in the sequence SET\_DIAGRAM\_TYPE N Y should be used first.

Syntax	SET_DIAGRAM_TYPE
	Triangular diagram (Y or N) /N/:
	The default is to set a perpendicular axis. Perpendicular diagrams have (almost) the same scaling on both axes.
Prompts	Create tetrahedron WRML file (Y or N) $/N/:$
Trompts	For a perpendicular axis, you can enter Y to use a tetrahedron diagram to generate a VRML (* . $WRL$ ) file by sequentially using the CREATE_3D_PLOTFILE command.
	Enter Y to specify a triangular plot instead.
	Plot 3rd axis (Y or N) /N/:
	If a triangular plot is selected, enter Y to specify a 3rd axis and to connect (and plot) the end points of the X- and Y-axes.
	Clip along 3rd axis (Y or N) $/N/:$
	If a triangular plot is selected, enter Y so that all lines outside the region limited by a line joining the end points of the X- and Y-axis are removed.

# **SET\_FONT**

Select the font to use for labels and numbers when plotting the diagram under the currently selected graphic device (<u>"SET\_PLOT\_FORMAT" on page 254</u>). For some devices there may be other fonts available and these are selected by the SET\_PLOT\_FORMAT command.

Syntax	SET_FONT
Prompts	Select font number /1/: <#>

Syntax	SET_FONT
	Specify an available font number # as the default font for the current graphic device, or accept the font number 1 by pressing <enter>. By typing a question mark ? here, the program lists the available fonts under the currently selected graphic device.</enter>
	Font size /.34/: <##>
	The size of the chosen font is specified. A value around 0.34 is recommended.

# **SET\_INTERACTIVE\_MODE**

### Syntax SET\_INTERACTIVE\_MODE

Resets the input and output units to their initial values, i.e. keyboard and screen.



Remember to add this as the last command to your MACRO files, in order to stop execution of the command file in the POST module. It has no meaning in interactive mode.

# SET\_LABEL\_CURVE\_OPTION

Identify the curves drawn in the post-processor by marking each curve with a digit and then list the meaning of these digits beside the plot.



The number of digits can be quite large in some cases. If so, use the SET\_FONT command and reduce the font size; usually, 0.2 is sufficient.

Syntax	SET_LABEL_CURVE_OPTION
Prompt	Curve label option (A, B, C, D, E, F OR N) /A/: <option></option>
	The options:
	• A: List stable phases along line
	• B: As A but curves with same fixed phase have same number
	• c: List axis quantities
	• ${ iny D}$ : As ${ iny C}$ but curves with same quantities have same number
	• E: As B with changing colors
	• F: As D with changing colors

Syntax	SET_LABEL_CURVE_OPTION
	• N: No labels
	This question is rather cryptic but usually the option B or E is good for phase diagrams (after a MAP command) and option D or F for property diagrams (after a STEP command). Those interested can try out the option A and C by themselves.
	Option B or E lists the fixed phases along each curve, whereas option D or F gives the axis variables used along each curve. E provides varied colors for different stable-phase curves on phase diagrams (after a MAP command); F provides varied colors for different property curves on property diagrams (after a STEP command). The option N (NONE) disables all curve labels and lists.
	If, for example, you use $T$ (temperature in K) as the variable axis in a STEP command and then plot the amount of stable phases (with <u>"SET_DIAGRAM_AXIS"</u> on page 249 Y NP(*)) on the Y-axis versus $T-C$ on the X-axis, then the list may have lines like:
	1: T-273.15, NP(LIQUID)
	2: T-273.15, NP(FCC_A1)
	This means that for curve 1 the X-axis is $T-273.15$ (the same for all curves, of course) and NP(LIQUID) on the Y-axis. Curve 2 has the same X-axis but NP(FCC_

#### A1) on the Y-axis.

# SET\_PLOT\_FORMAT

Set the format of the graphical output to different graphical devices. The default graphic device is normally for screen output (number 22 - TCUNITE Driver). Legacy devices can still be used (1 for Windows and 9 for XWindow for Linux). This default graphic device value may also be altered with <u>"SET\_PLOT\_ENVIRONMENT" on page 300</u> in the system monitor or by your TC.INI file.

Syntax	SET_PLOT_FORMAT
	Graphic device number /#/: <number device="" of="" the=""></number>
Prompt	Specify a graphic device number. Depending on the hardware, different plot formats (graphic devices) may be available. These are listed online by giving a question mark ?.
	For some formats there can be additional sub-prompts (Y or N) about the

### Syntax SET\_PLOT\_FORMAT

available fonts being used for the chosen format. Also enter Y for further specifications, e.g. font type and size.

## **SET\_PLOT\_OPTIONS**

You are prompted for and can toggle on (Y) or off (N) the plotting of some options on all subsequently generated diagrams.

For the Always initiate POST on re-entering prompt, the default is Y which implies that the POST module automatically reinitiates whenever the POLY3 workspace is changed upon re-entering. If answered N the same diagram axis variables are kept even if the workspaces in the POLY module are read from or written to an \*.POLY3 file. The N answer is useful when there are several sets of \*.POLY3 files with the same calculation but for different sets of model parameters.

For the Always solid line prompt, the default answer is N which means that the POST module plots solid lines for stable phases but dashed lines for metastable phases. This is normally for plotting property diagram after a STEPPING calculation; however, when there is a metastable extrapolation in a phase region on a phase diagram after a MAPPING calculation, this is also enforced. When the alternative answer Y is chosen, all the lines are always plotted in solid lines.

Syntax	SET_PLOT_OPTIONS
Dromoto	Plot header /Y/:
Prompts	Toggle the plotting of the Thermo-Calc header text above a diagram.
	Plot logo /Y/:
	Toggle the plotting of the Thermo-Calc logotype at the lower-left corner of a diagram.
	Plot footer /Y/:
	Toggle the plotting of the footer identifier text at the right margin of a diagram.
	White-Contoured-PS-CHARS /Y/:
	By default, the PostScript characters are written as a white contoured status.
	<pre>Plot remote exponent(s) /Y/:</pre>
	By default the remote exponent on the axis is plotted.

```
SET_PLOT_OPTIONS
Syntax
          Plot symbols at node points /0/: <#>
          This makes it possible to plot symbols at the node points on plotted
          lines on a diagram.
          Symbol size /.1/: <.#>
          Set the size of symbols which are plotted at the node points.
          Write condition? /Y/:
          By default, the initial equilibrium conditions for the current
          calculations are automatically plotted at the upper-left corner as a
          part of the header. If you enter N, the calculated conditions are not
          included on a plotted diagram.
          Write database name? /Y/:
          By default, the primary database (not appended ones) used in the
          current calculations are plotted at the upper-left corner as a part of
          the header. If you enter N, the database name is not included on a
          plotted diagram.
          Always initiate POST on re-entering: /Y/:
          By default the POST module automatically reinitiates whenever the
          POLY3 workspaces are changed upon re-entering.
          If you enter N, the same diagram axis variables are kept even if the
          workspaces in the POLY module is read from or written to an
          *.POLY3 file. This can be useful when there are several sets of
          *.POLY3 files with the same calculation but for different set of model
          parameters.
          Always solid line: /N/:
          By default the POST module plots solid lines for stable phases and
          dashed lines for metastable phases. This is normally for plotting
          property diagrams after a STEPPING calculation. However, when
          there is a metastable extrapolation in a phase region on a phase
          diagram after a MAPPING calculation, this is also enforced. Enter Y
          to always plot solid lines in this case, for example.
```

# SET\_PLOT\_SIZE

Change the size of the diagram by specifying a global plot size (as a relative scaling factor). The default value of the relative scaling factor is 1, while the real size of the plotted diagram depends on what output device is chosen with <u>"SET\_PLOT\_FORMAT" on page 254</u>. The default plot size (corresponding to the default global plot size 1) is adjusted to the chosen device.

Syntax	SET_PLOT_SIZE
	Global plot size /1/: <relative factor="" scaling=""></relative>
Prompt	Enter the relative scaling factor as a numerical number (e.g. 0.5, 0.8, 1.0, 1.5, etc.). By pressing <enter>, the default relative scaling factor (1) is accepted.</enter>

# SET\_PREFIX\_SCALING

Specify on which axis is to have a prefix scaling then set the scaling of remote exponents with certain powers.

Syntax	SET_PREFIX_SCALING
Dramat	Axis (X, Y or Z): <name an="" axis="" of=""></name>
Prompt	Specify which axis is to have a prefix scaling.
	Use prefix scaling /Y/: (Y or N or #)
	• NO switches off its action.
	• YES arranges the prefix scaling so that the axis scaling is done with the remote exponents being powers of three, i.e, -6, -3, 0, 3, 6, The default is 3.
	• # is an integer number that sets the remote exponents as being powers of # (as the powers for the remote exponents).

# SET\_RASTER\_STATUS

Plot a raster (i.e. with gridlines on both axis directions) in the diagram.

Syntax	SET_RASTER_STATUS
Prompt	Raster plot (Y or N) /N/:
	By default no raster is plotted. Enter Y to plot using a raster plot.

# **SET\_REFERENCE\_STATE**

8

The reference state for a component is important when calculating activities, chemical potentials and enthalpies and is determined by the database being used. For each component the data must be referred to a selected phase, temperature and pressure, i.e. the reference state.

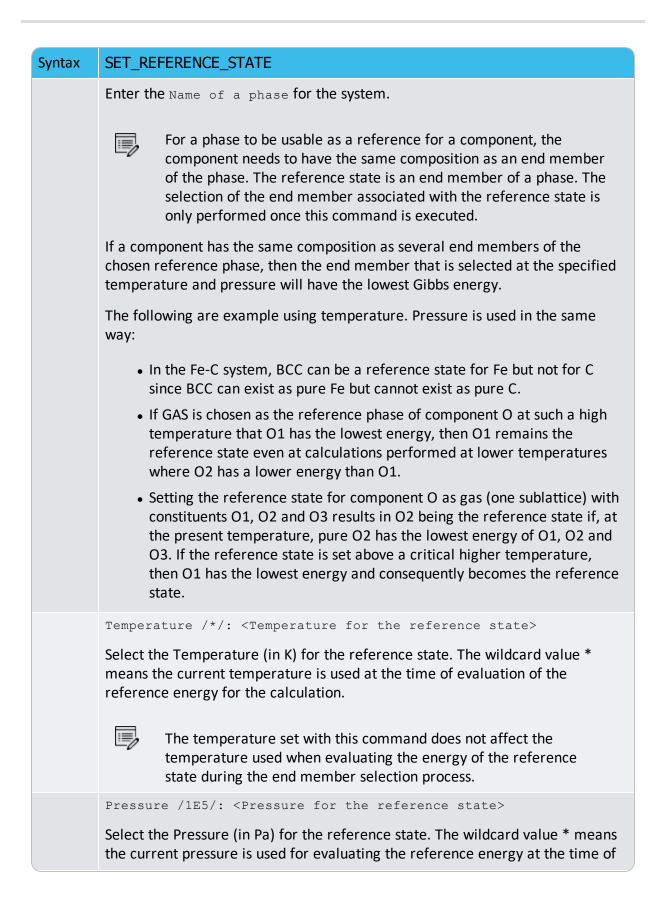
All data in all phases where this component dissolves must use the same reference state. However, different databases can use different reference states for the same element/component. It is important to be careful when combining data obtained from different databases.

By default, activities, chemical potentials and so forth are computed relative to the reference state used by the database. If the reference state in the database is not suitable for your purposes, use this command to set the reference state for a component using SER, i.e. the Stable Element Reference (which is usually set as default for a major component in alloys dominated by the component). In such cases, the temperature and pressure for the reference state is not needed.

In order to specify conditions in the specific reference state, you can append an R to the state variables. Thus, AC is the activity (of a system component or of a species in a phase) with respect to the default reference state, and ACR is the activity with respect to the selected reference state; MU is the chemical potential of a system component with respect to the default reference state, and MUR stands for the chemical potential with respect to the selected reference state. The AC and ACR variables, for both components in a system and species in a phase, can be specified in natural logarithm, e.g. LNAC(Fe), LNACR(C), LNAC (O2,GAS), LNACR(O2,GAS).

For the POST module, and after calculating with the <u>"STEP\_WITH\_OPTIONS" on page 217</u> or <u>"MAP" on page 201</u> commands, the reference state for a component can also be changed to plot various properties of the component in the entire system or in a specific phase. Then you can set the diagram axes as the chemical potential or activity quantities with the R suffix, i.e. MUR(comp), MUR(comp,ph), ACR(comp), ACR(comp,ph) or the common logarithms [e.g. LNACR(comp,ph)].

Syntax	SET_REFERENCE_STATE
Dramanta	Component: <name component="" of="" the=""></name>
Prompts	The name of the component must be given.
	Reference phase: <name a="" as="" new="" of="" phase="" reference="" state="" the="" used=""></name>



Syntax	SET_REFERENCE_STATE
	calculation.
	EXAMPLES
	S-R-S Fe SER
	S-R-S Cr FCC * 100000
	S-R-S H2O AQUEOUS * 100000
	S-R-S ZE REF_ELECTRODE * 100000

# SET\_SCALING\_STATUS

When an axis variable is selected by <u>"SET\_DIAGRAM\_AXIS" on page 249</u>, the scaling status for the axis is always set to automatic scaling. By this command, choose between manual or automatic scaling on a specified axis. If manual scaling is chosen, specify a minimum and a maximum value. Manual scaling can be used to magnify interesting parts of a diagram.

Syntax	SET_SCALING_STATUS
Drevente	Axis (X, Y or Z): <name an="" axis="" of=""></name>
Prompts	Specify which axis that you want to set the scaling status.
	Automatic scaling (Y or N) /N/:
	The default is manual scaling. Enter Y for automatic scaling (N).
	For manual scaling specify the Min and Max values.
	Min value: <minimum value=""></minimum>
	The minimum value at the start point of the specified axis.
	Max value: <maximum value=""></maximum>
	The maximum value at the end point of the specified axis.

# SET\_TIC\_TYPE

Change the drawing of the axis tics. You may alter the placement of tic marks on the axis, e.g. inside or outside or no tic marks. The size of tic marks cannot be altered by this command, but these are adjusted based on the selected graphic device and the defined relative scaling factor (global plot size).

SyntaxSET\_TIC\_TYPEPromptTIC Type /1/: <1 or -1 or 0>PromptThe tic type 1 is default, i.e. the tics are drawn outside the diagram. -1 means<br/>that the tics are inside the diagram and 0 means no tics.

# **SET\_TIELINE\_STATUS**

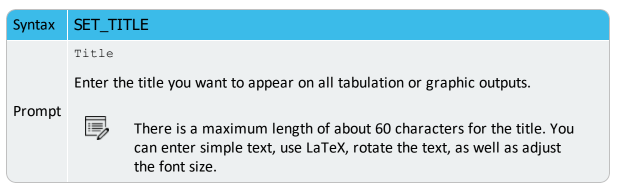
If the tie-lines are in the plane of the calculation, you can select to plot the tie-lines in twophase fields with this command.

S	yntax	SET_TIELINE_STATUS
		Plotting every tie-line no. /0/: <number of="" tie-lines=""></number>
Ρ	rompt	The tie-lines plotted are not equally spaced by the graphics software. Instead, you can select to plot a subset of the calculated tie-lines, i.e. every one (1), every second (2), every three (3), etc. By accepting the default value 0, no tie-line is plotted.

# **SET\_TITLE**

Specify a title that displays on all listed tables and plotted diagrams from the POST module in the same run of Console Mode.

"ADD\_LABEL\_TEXT" on page 234



# SET\_TRUE\_MANUAL\_SCALING

The tic-marks on the axes are normally placed in even intervals using the whole axis length. The scaling routine adjusts the given minimum and maximum values slightly to accomplish this. If such an automatic adjustment behaviour is unwanted, it can be avoided by using this command (followed by a message showing the TRUE MANUAL SCALING set for a specific axis).

The command works like a toggle. To reset the automatic scaling behaviour, repeat the command a second time for that specific axis (followed by a message saying the SEMI\_MANUAL SCALING set for the specific axis.

Syntax	SET_TRUE_MANUAL_SCALING
Drement	Axis (X, Y or Z): <name an="" axis="" of=""> Specify which axis (X or Y or Z) you want to toggle between automatic</name>
Prompt	adjustments or avoid the adjustment of the given maximum and minimum values.

# SUSPEND\_PHASE\_IN\_PLOT

Specify that lines originating from the presence of a certain phase shall not be plotted on a diagram. If you want to bring the suspended phase back to the diagram, use <u>"RESTORE</u> PHASE\_IN\_PLOT" on page 245.

Syntax	SUSPEND_PHASE_IN_PLOT
Description	Phase name: <name a="" of="" phase=""></name>
Prompt	Specify the name of the phase to be suspended from the diagram.

# **TABULATE**

For any entered table, this command gives a table of values from equilibria calculated with a STEP command. It works only for a table that is entered in either the POLY or POST module.



Unlike the TAB module's command (also called <u>"TABULATE" on page 311</u>) it is not possible to plot columns from the tabulated tables. For this use normal POST commands.

Syntax	TABULATE
Dremente	Name: <name a="" either="" entered="" in="" of="" or="" poly="" post="" table=""></name>
Prompts	Specify a table name that is entered in either the POLY or POST module.
	Output on screen or file /Screen/:
	Specify a file name if you want to save on a file the table values along the defined STEP calculation, or press <enter> if want to see the table values displayed.</enter>

### **POTENTIAL\_DIAGRAM Module**

### In this section:

POTENTIAL	DIAGRAM		

# **POTENTIAL\_DIAGRAM**

8

The POTENTIAL\_DIAGRAM module (short name, the POTENTIAL module) is a potential phase diagram calculation module.

If you have a metal-oxide/sulphide-gas interaction system, you might want to calculate a socalled potential phase diagram given certain temperature and pressure. A potential diagram uses the activities of two major species in the gaseous mixture phase as mapping variables. The activities (that is, the fugacities at 1 bar) of these two species are typically set as the Xand Y-axis when the diagram is plotted. The phase relations between the gaseous mixture and various metal forms, metal-oxides, metal-sulphides, or other metal-bearing solids, are represented by different fields controlled by these two potential quantities.

To enter the module, at the SYS prompt type GOTO\_MODULE POTENTIAL.

There are no commands for this module. Follow the prompts and see <u>"About Potential Diagrams" on page 211</u> in the *Thermo-Calc User Guide* for details about how to calculate using potential diagrams.

See example <u>tcex35</u> in the *Thermo-Calc Console Mode Examples Guide*.

Syntax	POTENTIAL_DIAGRAM
	Database: /POT/
Prompts	Press <enter> and the program lists the available databases. Type one of the database names and press <enter> to select. The following default elements are then based on the chosen database.</enter></enter>
	Matrix element: /Fe/
	First potential species: /S102/
	Second potential species: /02/
	Temperature: /1000/

### **POURBAIX\_DIAGRAM Commands**



Also search the online help or see the Pourbaix diagrams educational material and the *Thermo-Calc User Guide*.

### In this section:

POURBAIX_DIAGRAM Commands	266
Getting started with the Pourbaix module	266
Pourbaix Main Options	269

# **POURBAIX\_DIAGRAM Commands**



Also search the online help or see the *Pourbaix Diagrams Educational Material*, the *Thermo-Calc User Guide* and the *Thermo-Calc Console Mode Examples Guide* all included in this documentation set.

### Getting started with the Pourbaix module

After you enter the POURBAIX\_DIAGRAM module, you are asked if you want to have step-bystep instuctions to help walk you through the module.

Need any instruction on the Pourbaix module? /N/: Y

If you answer  ${\tt Y}$  a series of notes are displayed. The following is a more comprehensive summary of these same notes.

### What is a Pourbaix diagram?

The Pourbaix diagram is a calculated equilibrium phase diagram mapped and plotted with the independently-varied electropotential (Eh, as defined with regard to the standard hydrogen electrode as its reference) and acidity (pH), that represents all the equilibrated phase boundaries among aqueous solution, gaseous mixture, and various primary and secondary solids (modeled as either complex solution or simple stoichiometric phases) in a certain multicomponent heterogeneous interaction system, under a defined T-P condition and a specific initial bulk composition (which is, by default, always set as 1 kg of water dissolving a specified amount of metals/alloys and other acids/alkalines/salts).

### What database should I use?

At least one database is required that contains an aqueous solution phase (with thermodynamic data for water and various aqueous solutes), that shall be selected from TCAQ (PAQ) or AQS (PAQS) [which use the SIT Model or the Complete Revised HKF Model, respectively] within the Thermo-Calc databases.

### Choosing a custom aqueous solution database

You can also use a custom database but it must be in the Thermo-Calc TDB format. Due to the restrictions of aqueous solution models used within Thermo-Calc, such a database must be designed in the same format as in the default TCAQ (PAQ2) for aqueous solution phases. Among other things, keep the following requirements in mind:

The ELECTRON is defined as an special element (ZE) and as the only constituent in its reference phase REFERENCE\_ELECTRODE (for determining the electro-potential that is defined as Eh with the standard hydrogen electrode as the reference), but it is not defined as an aqueous species;

The vacancy (VA) is unnecessary for AQUEOUS solution phase and it should be avoided in the definition of phase-constituents in the AQUEOUS phase;

The AQUEOUS solution phase should always be defined as a constitutional solution phase, implying that all the aqueous solution species must be included in a single site, rather than in two or multiple sublattices.

Beside the AQUEOUS solution phase, there shall exist a GAS mixture phase containing at least H2O1, O2 and H2; and for multicomponent systems, normally there shall also contain some solid (stoichiometric or solution) phases. Of course, if desired, you could also choose to calculate and generate a Pourbaix diagram without considering the GAS mixture phase entirely; however, such a plot is not really a complete Pourbaix diagram, due to the fact that thermodynamically-stable phase boundaries between the AQUEOUS solution phase and GAS mixture phase will then not be calculated at all!

All the required thermodynamic data for calculations of Pourbaix diagrams or other diagrams must be retrieved either from one (Single) database which consists of an AQUEOUS solution phase, a GASeous mixture phase, a REF\_ELECTRODE phase, and some SOLID phases (being solutions and/or stoichiometric compounds; for primary metals/alloys and for secondary products formed from heterogeneous chemical/electrochemical interactions, or from several (Multiple) databases that respectively contain various solutions/compounds (as listed above).

Such databases suitable for calculations of aqueous-bearing heterogeneous interaction systems can be those default-prompted ones [i.e., in the Single-Database case, the PAQ or PAQS; and in the Multiple-Database case, the PAQ2 or as primarily-switched database, plus the SSUBx as firstly-appended one and the SSOLx as secondly-appended one if it is necessary; even more databases can be appended). Of course, you could also choose to append required data from other appropriate databases (such as TCFE, TCSLD, TCNI/TTNi, TCAL/TTAI, TCMG/TTMg, TTZr, TCMP, SLAG, etc.) for GASeous mixture phase and for varous solid solution and stoichiometric compound phases. Furthermore, an experienced user can also utilize his/her own USER-specified databases in various cases.

The advanced POURBAIX module is designed so that it requires you to just answer some necessary questions, rather than to go through the other Console Mode modules (i.e., TDB, GES5, POLY3 and POST) although you still enter commands at the prompts. Beside the default plotted Pourbaix diagram, it also allows the user to easily and quickly plot many different properties of the system, stable phases and aqueous species, varied along the calculated phase boundaries for the same defined heterogeneous interaction system. Moreover, it permits the user to directly change some plotting settings and manipulate all kinds of plotted diagrams.

The current advanced POURBAIX-Module has been extended so that it is additionally able to directly perform a normal STEPPING calculation (varied with a specified independent variable) and to easily generate various types of property diagrams, for the same heterogeneous interaction system that has been defined in a previous POURBAIX or TDB-GES5-PLOY3-POST calculation.

### Copy and Rename the POLY3 file

At the end of a Pourbaix module calculation, a new POLY3 file is automatically saved. For Windows users this is saved to the Documents folder.

If you are using a POLY-3 file more than once (as is done in the extended examples), it is recommended that you copy and rename the previouslygenerated POLY3 file. This is so that the required POLY3 file structure and Pourbaix results are not lost. Do this outside of Thermo-Calc i.e. do not save from within Thermo-Calc.

>\_

8

The reason you do not save from within Thermo-Calc is because of how the file is saved. When the command sequence (GO POLY3>SAVE *<FILE NAME>*.POLY3>Y) is used, the POLY3 workspace of the POURBAIX calculation results in the POLY3 module monitor and all the initial equilibrium points and the mapping/stepping calculation results are lost as well as parts of the POLY3 file structure (e.g., some definitions of the previously defined POURBAIX-type calculation system).

#### Enforce a Pause

After the initial set of information is displayed, you are asked:

Enforce a PAUSE after plotting when running a MACRO? /N/

- If you choose x a pause is always enforced after each diagram is plotted;
- If you choose N, all the command-lines (answers) for the session continue without stopping or until the SET\_INTERACTIVE\_MODE line in the macro file is reached.

Whenever running a Thermo-Calc macro (TCM) file, you may want to pause after a specific diagram is plotted on screen. This gives you the flexibility to:

- Print it (of EMF/PS format) to a printer
- Convert it (of EMF format) to (PDF) graphical files;
- Save it as an EMF graphical file;
- Dump it as a PNG or BMP graphical file;

- Set background colour for the current diagram
- You can also set the default font/size and change plot layers for sequential plots.

### **Pourbaix Main Options**

If you answer y to the Pause prompt above, these options are displayed:

- 1. Start a completely new Pourbaix diagram calculation.
- 2. Open an old file and plot other property diagrams.
- 3. Open an old file and make another Pourbaix calculation.
- 4. Open an old file and make another stepping calculation.
- "POURBAIX\_DIAGRAM Commands for Options 1 to 4" below

### **POURBAIX\_DIAGRAM Commands for Options 1 to 4**



8

Also search the online help or see the *Pourbaix Diagrams Educational Material*, the *Thermo-Calc User Guide* and the *Thermo-Calc Console Mode Examples Guide* all included in this documentation set.

### **Option 1: Start a completely new Pourbaix diagram calculation**

Option 1 is used to make a completely new Pourbaix diagram calculation and automatically plot a pH-Eh diagram. i.e., define a new chemical system; specify the T-P-X conditions; calculate the initial equilibria; perform the pH-Eh mapping calculation; plot pH-Eh and various property diagrams.

Many of the commands in Option 1 are also repeated for Options 2 to 4. The details for each command are included with this section unless it is specific to one of the other option choices. The order that the commands display on screen is also different and based on what is entered at the prompts. These prompts are also available in other commands and when applicable a link to that command is included.

Consider the GAS phase in calculating a Pourbaix diagram? /Y/

You can choose to ignore the GAS mixture phase on a calculated/plotted Pourbaix diagram. However, such a plot is actually not a complete Pourbaix diagram because the thermodynamically-stable phase boundaries between the AQUEOUS solution phase and GAS mixture phase will not be calculated.

- If you enter Y (the default), GAS mixture phase is always considered.
- If you enter N the GAS mixture phase is completely ignored.

Use single database? /Y/

If you keep the default you are prompted for a database.

Combined database: /PAQ2/

The default is PAQ2, a database included with all installations.

The following information is provided on screen if you chose Y at the first prompt, Need any instruction on the POURBAIX module? /N/.

#### DEFINE A CHEMICAL SYSTEM AND ITS INITIAL BULK COMPOSITION

Normally a Pourbaix diagram and its related equilibrium property in a heterogeneous interaction system are calculated under a certain bulk composition, which is usually 1 kg of water with defined amounts of dissolving solute substances. The solutes may either be completely dissolved into the aqueous solution, or be partially dissolved and simultaneously form some solid phases.

#### Chemical system (elements)

- Default defined elements (solvent H2O): H and O and specially assigned ZE (electron) and VA(vacancy).
- Prompt specified elements (solutes ELEM): Fe Ni Co Cr C N S Na Cl

#### Initial bulk composition

- Default defined composition (solvent): 1.0 kg of H2O
- Prompt specified composition (solutes): x mole of ELEM
- Mass of Water (weight) = 1 kg

When entering solutes in chemical formulas, the first element's letter must be in UPPER case, and second in lower case, for example NaCl, CO2, CoCl3, Fe0.93S, NaSO4-1, and H2SO4.

First solute

8\$B

Enter a solute, e.g. Fe.

Molality of <first solute> [mol/kg] /.001/

Enter a molality for the first solute, e.g. Fe.

Second solute

Enter a second solute and its molality on the same line, e.g. NaCl .1.

Next solute

You can continue to enter solutes. Press <Enter> to skip to the next prompt.

The following information is provided on screen if you chose Y at the first prompt, Need any instruction on the POURBAIX module? /N/.

#### DEFINED CHEMICAL SYSTEM AND INITIAL BULK COMPOSITION

Solutes have been split up into chemical elements and their mole numbers. If confirmed, the Pourbaix module will, in further steps, count the initial bulk composition in terms of chemical elements.

Confirm defined system and initial bulk composition? /Y/

If you keep the default (Y) thermodynamic data is retrieved from the database. Then it lists the database's phases for the defined system.

You can then reject or restore phases as required with the next prompts.

Reject phase(s) /NONE/
Restore phase(s): /NONE/

Then the program lists the phases retained in the system and you prompted whether this is OK or not.

OK? /Y/

If you answer Y, then a list of references for the assessed data is displayed on the screen. The next prompt is related to miscibility gap checks, global minimization, saving the information to a POLY3 file, and setting numerical limits.

Should any phase have a miscibility gap check? /N/

The database usually creates two or more "composition sets" for phases that can have miscibility gaps. However, for some phases this is not done automatically, for example the miscibility gap in the bcc phase in Fe-Cr is usually ignored. But if it is important to include a Cr-rich bcc phase specify this here. It costs some computation time and may make subsequent MAP or STEP processes more difficult to converge. To keep the default (not to have any such phase) press <Enter>. Otherwise answer Y and follow the additional prompts.

Enforce Global Minimization Technique in calculations? /N/

If you choose Y, the Global Minimization is done when possible (depending on the conditions setting), and a Global Minimization test is performed when an equilibrium is reached. This costs more computer time but the calculations are more robust.

Save all functions, variables and tables in POLY3 file? /Y/

Choose x if you want to save all the defined symbols (functions, variables and tables) into the POURBAIX.POLY3 file, for further normal POLY3 calculations or normal POST post-processing. Choosing yes may result in a large POLY3 file.

Set numerical limits? /N/

"SET\_NUMERICAL\_LIMITS" on page 293

The Pourbaix module has a default setting for numerical limits.

- Maximum number of iterations: 20000
- Required accuracy: 1E-6
- Smallest fraction: 1E-15
- Approximate driving force for metastable phases: Y

If you kept the defaults for the above prompts, the program lists the default and pre-defined calculation conditions and you are prompted to confirm.

Confirm defined conditions? /Y/

Now Thermo-Calc calculates the start points. This can take a few moments. Near the end of the calculation there is a list of the default axis variables. The maximum pH limit is calculated precisely and determined automatically by the Pourbaix module as a function of the temperature-pressure conditions and initial bulk compositions of the current defined interaction system. You are prompted to accept these settings.

Accept the default settings for two mapping variables? /Y/

The program now lists the initial equilibria for mapping and continues to calculate the mapping. Then it sets the automatic diagram axes and starts to plot the diagram. But first there are several prompts to guide you.

Any missing phase boundary you could possibly think of? /N/

Sometimes the Pourbaix module may miss some phase boundaries during the mapping process. If you answer Y you can add additional starting point(s) into the calculation. You are now prompted to specify a pH-Eh coordinate pair, for instance (4.0, -0.15), which is close to the expected phase boundary. However, such additional pH-Eh point should not outside the AQUEOUS/GAS(O2) or AQUEOUS/GAS(H2) boundaries; otherwise, the module will fail in finding any new starting point.

The next set of prompts relate to plotting.

```
Change the pH/Eh steps for smoother curves? /N/ \,
```

You can modify the mapping steps of X/Y axes for smoother curves; sometimes it may help finding missing phase boundaries by decreasing the pH and Eh increments.

Zoom in? /N/

Change the scaling status for both X and Y axes. The program then asks for minimum and maximum values for the axes. In some cases values are suggested.

Change Curve-Label Option for the diagram? /N/

Change the Curve-Label option to one of the following:

- A: List stable phases along phase boundaries
- B: As A, but curves with same fix phase have same number
- C: List axis quantities
- D: As C, but curves with same quantities have same number
- E: As B, but curves with changing colours
- F: As D, but curves with changing colours
- N: No labels

Usually, Option E or B is good for phase diagrams such as pH-Eh plots, and F or D is good for property diagrams like pH-RC(Fe+2).

"SET\_LABEL\_CURVE\_OPTION" on page 253

Add Label-Texts onto the Pourbaix diagram? /N/

Add label text to an area in the diagram. The label is written starting from the specified coordinates. Optionally, let the program automatically add the text by calculating an equilibrium at some specific coordinates and also make a label as the stable phase names in the area. However, such an automatic calculation may fail, especially for a Pourbaix diagram where either pH or Eh is not an intensive variables.

In most cases, the program is able to write the label if the coordinates and texts are specified.

Change the Subtitle of the diagram? /N/

This is the same as the POST module command <u>"SET\_CORNER\_TEXT" on page 249</u>  $\rightarrow$  UPPER\_RIGHT <SUBTITLE>. The default subtitle is automatically assigned by the POURBAIX module and is based on the defined conditions in the system (temperature, pressure and initial bulk compositions).

If you choose Y then the existing subtitle is displayed and you are prompted to enter a new subtitle or to have None.

New Subtitle /NONE/

You can change the subtitle for all the diagrams in the same POURBAIX run (i.e., within the same Option). The new subtitle that can be altered later at the same option, when further modifying the present graph or when refining any sequentially defined plot. By entering text (empty space at any position is allowed), the new subtitle is defined. By typing OLD, the existing subtitle is kept.

By pressing <Enter> (i.e., accepting None), the subtitle is deleted from all sequential plots, which can be set back later at the same option, too.

```
Change Axis-Texts? /N/
```

Change the axis text for both X and Y axes.

Further Refine the diagram in POST Module? /N/

With this option, you can further change the appearance the plot and use any command in the normal POST Module.

Plotting and Visualization with the POST Module " on page 231 Hard copy of the diagram? /N/

### **RECOMMENDED RESPONSES**

- Hard copy of the diagram? /N/: Y
- CURRENT DEVICE: MS-Windows
- GRAPHIC DEVICE NUMBER /1/: 5
- NEW DEVICE: Postscript portrait mode
- USE POSTSCRIPT FONTS /YES/: Y
- SELECT FONTNUMBER /9/: 27
- NEW FONT: Times-Bold
- FONT SIZE /.35833/: .32
- PLOTFILE : /defaultname/: yourfilename

Save X-Y coordinates of curve on text file?  $/\ensuremath{\,\text{N}}/$ 

Write all the current graphic information (generated and modified in the above procedure) on a text file using the DATAPLOT format (see the DATAPLOT User Guide or search the online help). This also makes it possible to merge two or more diagrams from separate but similar calculations, or put some experimental data points onto a calculated diagram. You do this by first writing them onto text files with this option and then edit them together (using a general text editor). The program asks for a file name. Accept the default (usually POURBAIX) or use another name. If an extension is not given, the written file is given an extension of \*.EXP.

```
Modify the diagram? /N/
```

Further modify the appearance of the created plot. This brings you back to the above prompts for changing various features.

Any more diagram? /N/

Make many other plots using the same POURBAIX module calculation file.

#### Option 2: Open an existing file and plot other property diagrams

#### COPY AND RENAME THE POLY3 FILE

At the end of a Pourbaix module calculation, a new POLY3 file is automatically saved. For Windows users this is saved to the Documents folder.

8 <b>9</b> 8	If you are using a POLY-3 file more than once (as is done in the extended
-00-	examples), it is recommended that you copy and rename the previously-
	generated POLY3 file. This is so that the required POLY3 file structure and
	Pourbaix results are not lost. Do this outside of Thermo-Calc i.e. do not save
	from within Thermo-Calc.

The reason you do not save from within Thermo-Calc is because of how the file is saved. When the command sequence (GO POLY3>SAVE *<FILE NAME*>.POLY3>Y) is used, the POLY3 workspace of the POURBAIX calculation results in the POLY3 module monitor and all the initial equilibrium points and the mapping/stepping calculation results are lost as well as parts of the POLY3 file structure (e.g., some definitions of the previously defined POURBAIX-type calculation system).

Open an existing POLY3 file created by the Pourbaix module (from a previous Pourbaix calculation Option 1 or 3 or a previous Stepping calculation Option 4), and selectively plotting other property diagrams. i.e., open the old GES and POLY3 workspaces; plot pH-Eh or various property diagrams.

File name /ABCDEF/

The name of a previously saved POLY3 file where the POLY3 and GES5 workspaces is to be read must be specified. You do not need to enter the file extension if it is the default .POLY3 name. The default file name (e.g. ABCDEF) is the most recent specified POLY3-file name or RESULT if there is no workspace already saved in the run. You can press <Enter> or type a specific file name to read the POLY3 and GES5 workspaces from the file.

For Windows, and at the prompt, an **Open file** window displays if a file name is not given or the file path is incorrect. The file type (i.e. POLY3) cannot be changed. Click **Open** or **Cancel** to continue with the selected file.

When the file opens, it confirms on screen what kind of calculation is contained in the POLY3 file. For example: *The loaded POLY3 file is of POURBAIX-MAPPING Calculation*.

On screen it also lists the system components and the default and predefined calculation conditions to enable you to review these.

Next you are given a set of properties that are available to be selected as axis variables for making other diagrams from the same calculation.

Abbreviation	Definition
рН	Acidity
Eh	Electronic potential (V)
Ah	Electronic affinity (kJ)
ре	Electronic activity [log10ACRe)
IS	Ionic strength
ТМ	Total concentration
Aw	Activity of water
Oc	Osmotic coefficient
MF (AQsp)	Mole fractions of aqueous species
ML (AQsp)	Molalities of aqueous species
AI (AQsp)	Activities of aqueous species
RC (AQsp)	Activity coefficients of Aq species

Where AQsp is the name of a specific aqueous species, and "\*" can be used as a wild sign for all AQSp. It is not necessary to give a complete aqueous species name. However, the entered AQsp name must be unique. If neither AQsp nor \* is given, all species are searched.

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The rest of the prompts are the same as with <u>Option 1</u> and relate to plotting. Enter ? at any prompt for details.

### **Option 3: Open an existing file and make another Pourbaix calculation**

#### COPY AND RENAME THE POLY3 FILE

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At the end of a Pourbaix module calculation, a new POLY3 file is automatically saved. For Windows users this is saved to the Documents folder.

If you are using a POLY-3 file more than once (as is done in the extended examples), it is recommended that you copy and rename the previouslygenerated POLY3 file. This is so that the required POLY3 file structure and Pourbaix results are not lost. Do this outside of Thermo-Calc i.e. do not save from within Thermo-Calc.

The reason you do not save from within Thermo-Calc is because of how the file is saved. When the command sequence (GO POLY3>SAVE *<FILE NAME*>.POLY3>Y) is used, the POLY3 workspace of the POURBAIX calculation results in the POLY3 module monitor and all the initial equilibrium points and the mapping/stepping calculation results are lost as well as parts of the POLY3 file structure (e.g., some definitions of the previously defined POURBAIX-type calculation system).

Open an existing POLY3 file created by the Pourbaix module and make another Pourbaix diagram calculation i.e. open the old GES and POLY3 workspaces, adopt the defined chemical system, modify the T-P-X conditions, calculate the initial equilibria, perform the pH-Eh mapping calculation and plot pH-Eh or other property diagrams.

The rest of the prompts are the same as with <u>Option 1</u> or <u>Option 2</u>. Enter ? at any prompt for details.

Modify Label-Texts on the Pourbaix diagram? /N/

Once some labels are written by the Add-Label-Text command onto the diagram, you can also modify the labels (changing their positions and text contents). The text-size cannot be changed. The program asks which label and you enter its new X/Y coordinates with the new text. The new text must not be longer than the old one.

Modify Label-Curve-Option on the Pourbaix diagram? /N/

Because some phase-region labels have been written onto the diagram, the Label-Curve-Option might need to be modified, normally to the Option N.

#### **IMPORTANT INFORMATION ABOUT OPTION 3**

The default definitions of the Eh and pH quantities in the advanced Pourbaix module (and in the ordinary TDB-GES-POLY calculation routines) should ALWAYS be as below:

```
Eh = MUR(ZE)/RNF
pH = -log10[AI(H+1,AQUEOUS)]
= -log10[ACR(H+1,AQUEOUS)*AH20]
```

where RNF is the Faraday constant (96485.309 C/mol), and AH2O is the molecular weight of H2O (55.508435 g). MUR(ZE) is the electrochemical potential (ECP; in the unit of J/mol; w.r.t. the standard hydrogen electrode). ACR(H+1,AQUEOUS) is the site-fraction-based activity of the H+1 aqueous species in AQUEOUS solution phase, but AI(H+1,AQUEOUS) [that equals ACR(H+1,AQUEOUS)\*AH2O] is the molality-based activity of the H+1 aqueous species that should be used for defining the pH quantity.

Within an aqueous-bearing heterogeneous interaction system, the fundamental systemcomponents must be H2O, H+1 and ZE, which are corresponding to the basic elements O & H and the hypothetical electron (ZE) in the aqueous solution phase. For the additional chemical elements in the system, their corresponding system-components shall be defined as in their element forms (such as Fe, Cr, Mn, Ni, Na, Cl, S) or (for some) in their molecular forms (e.g., NaCl, H2S). The reference state for the H2O component must always be defined as the solvent species H2O in the AQUEOUS solution phase under the current temperature (\*) and 100000 Pascal (i.e., 1 bar). The reference states for the H+1 and ZE components are by default set as their SER.

Various conventional properties of aqueous solute species I are converted in the following manners:

```
ML = Y(AQUEOUS, I) *AH2O/YH2O
```

- RC = ACR(I,AQUEOUS)\*YH2O/Y(AQUEOUS,I)
- AI = RC\*ML
- = ACR(I, AQUEOUS)\*AH2O

# where YH2O [i.e., Y(AQUEOUS,H2O)] and Y(AQUEOUS,I) are the site-fractions of solvent H2O and solute species I.

#### LIST THE DEFINED AXIS-VARIABLES:

```
Axis No 1: LNACR(H+1)Min: -34.532525Max: 2.3025851 Inc: 0.8
Axis No 2: MUR(ZE) Min: -150000 Max: 200000 Inc: 7718.85
```



The default settings (listed above) for two mapping variables [in terms of InACR (H+1) and MUR(ZE), and their minimum/maximum values and increment steps] are covering the following pH-Eh ranges/steps:

pH: 0.00 -> 14.00, at a step of 0.35 Eh: -1.55 -> 2.07, at a step of 0.08 (V) The maximum pH limit has been calculated precisely and determined automatically by the Pourbaix module, as a function of the temperature-pressure conditions and initial bulk compositions of the current defined interaction system.

### **Option 4: Open an Existing file and make another Stepping calculation**

Open an existing POLY3 file created by POURBAIX Module (from a previous POURBAIX calculation Option 1 or 3 or a previous STEPPING calculation Option 4), and make a normal STEPPING calculation. i.e., open the old GES and POLY3 workspaces; adopt the defined chemical system; specify one of the T-P-X conditions as the stepping variable; calculate the initial equilibria; perform the stepping calculation; plot various property diagrams.

Many prompts are the same as with <u>Option 1</u> or <u>Option 2</u>. Enter ? at any prompt for details.

#### COPY AND RENAME THE POLY3 FILE

At the end of a Pourbaix module calculation, a new POLY3 file is automatically saved. For Windows users this is saved to the Documents folder.

If you are using a POLY-3 file more than once (as is done in the extended examples), it is recommended that you copy and rename the previously-generated POLY3 file. This is so that the required POLY3 file structure and Pourbaix results are not lost. Do this outside of Thermo-Calc i.e. do not save from within Thermo-Calc.

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The reason you do not save from within Thermo-Calc is because of how the file is saved. When the command sequence (GO POLY3>SAVE *<FILE NAME*>.POLY3>Y) is used, the POLY3 workspace of the POURBAIX calculation results in the POLY3 module monitor and all the initial equilibrium points and the mapping/stepping calculation results are lost as well as parts of the POLY3 file structure (e.g., some definitions of the previously defined POURBAIX-type calculation system).



The stepping calculation modifies the POLY3 file but this can be used for other Stepping calculations inside the Pourbaix module (e.g. repeating Option 4) or in normal POLY3 module.

#### **STEPPING PARAMETERS**

The following parameters are suitable for a STEPPING calculation for the defined aqueousbearing system. Choose one of them as the stepping variable:

```
T Temperature (C)
P Pressure (bar)
pH Acidity
Eh Electronic Potential (V)
N(Comp) Mole of One Component
```

where "Comp" is the name of a specific component as listed below. One should not use any key component (H2O or H+1 or ZE) as such a stepping variable, because of that their compositional definitions and variations are determined by the AQUEOUS solution phase in the POURBAIX Module.

An interacting component defined as an element (such as Fe or Na or Cl) can be selected as a stepping variable.

However, if you want to make a stepping calculation with a non-element chemical formula (such as NaCl or CaS2) as the stepping variable, one must have made some necessary modifications in the POLY3 Module, before reaching this point, by conducting some POLY3 commands as given below (as an example for the system Fe-Na-Cl-H-O):

```
POLY_3: def-component H2O H+1 ZE Fe NaCl Cl
POLY_3: set-ref-state H2O AQUEOUS * 1E5
POLY_3: set-ref-state ZE REF_ELEC * 1E5
POLY_3: change-status phase REF-ELEC=SUSPEND
POLY_3: set-cond b(H2O)=1000 n(H+1)=0 n(ZE)=0
POLY_3: set-cond P=1E5 T=298.15**
POLY_3: set-cond n(Fe)=1E-3 n(Cl)=0 n(NaCl)=3 **
```

\*\* Here the P-T-n(Fe) conditions can be set as what you wanted, while the initial n(NaCl) value must be set as between its minimum and maximum values that you are interested in. Later on, it will be prompted for specifying the n(NaCl) Min-Max-Step values of the stepping calculation.

```
Stepping variable: /T/
```

The chosen stepping variable must be one of the suitable parameters listed above.

```
Minimum temperature (C) /15/
Maximum temperature (C) /35/
Increment in temperature (C) /1/
```

The AQUEOUS solution phase under the defined bulk composition and pressure condition is stable only between 0 and 99° C. However, you can make a stepping calculation outside this temperature range, where the AQUEOUS solution phase become metastable. In this case, the resulting POLY3 file is possibly able to be opened by any other option within the Pourbaix module.

# **REACTOR\_SIMULATOR\_3 Commands**

### In this section:

CREATE_DIVIDERS	283
CREATE_PIPES	283
CREATE_STAGE_BOXES	284
EXECUTE_POLY3_COMMAND	285
LIST_RECORDS	286
READ_WORKSPACE	286
SAVE_WORKSPACE	286
START_SIMULATION	. 287

# **CREATE\_DIVIDERS**

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This is a command used with the REACTOR module.

Define the segment boundaries and the ways on how to distribute phases between various segments and to split the feeds of heat and mass.

Syntax	CREATE_DIVIDERS
	Number of dividers /4/: <the dividers="" number="" of=""></the>
Prompts	Specify the number of dividers (boundaries between various segments) in the reactor. It is normally the segment number plus one: for example, if the stage box number is defined as 4, the divider number should be defined as 5.
	Number of outputs for divider # /3/: <the number="" of="" outputs=""></the>
	Specify the number of outputs for a specific divider #.
	Percent of input to output 1 /100/: <percent input="" of=""></percent>
	Define the percentage of input to the output 1 in the current divider #.
	<pre>Percent of input to output ## /100/: <percent(s) input(s)="" of=""></percent(s)></pre>
	Define the percentages of inputs to the outputs ## in the current divider #. It is repeated till the last output is specified.

# **CREATE\_PIPES**



This is a command used with the REACTOR module.

Define the pipes between various segments and the ways on how the mass and heat transport between the surroundings (with the record number as 0) and the reactor, among various segments in the reactor, and along various dividers (segment boundaries in the reactor).

Syntax	CREATE_PIPES
Prompts	From record: <a number="" record=""></a>
	Specify a record number. If there are four segments in the reactor, these records are available:
	<ul> <li>0 surroundings</li> <li>1-4 stage boxes (segments)</li> </ul>

Syntax CREATE\_PIPES

• 5-9 dividers (segment boundaries)

```
Pipe to record: <Another record number>
```

Specify another record number to which the mass or heat flows from the current record (specified above). Before this prompt, a message shown on the screen indicates what kind of mass or heat flow (according to the definitions of stage boxes and dividers, which have already been created prior to this command) should be specified in the pipe. This is repeated for all inputs and/or outputs for mass and/or heat in current record.

### **CREATE\_STAGE\_BOXES**

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This is a command used with the REACTOR module.

Create several reaction stages as boxes (segments), where the you define the feed in either mass input and initial temperature or heat input from surroundings, as well as the mass out and initial temperature in each of the segments.

Syntax	CREATE_STAGE_BOXES
	Number of stage boxes /4/: <the boxes="" number="" of="" stage=""></the>
Prompts	Specify the number of stage boxes in the reactor. Then define the feed of mass and/or energy (from surroundings to) the rector.
	Give feed to system: <feed></feed>
	One feed (of mass or heat) to the reactor must be specified here. Two types of feeds can be used:
	• As input amount of heat (enthalpy): H=Value.
	• As input amount of specific species or element and its initial temperature: N(SPECIES)=Value, Input temperature (in one line, or the input temperature in the next prompt).
	<pre>Input temperature /298.15/: <input temperature=""/></pre>
	Give feed to system: <feed></feed>
	Specify other feeds (mass or energy) to the reactor. Press <enter> to finish the feed inputs, and then:</enter>
	Give for stage box #

Syntax	CREATE_STAGE_BOXES
	Now start to define the type of stage box (segment is specifically numbered as #), and to specify a guess of initial temperature and an output for each phase in the segment.
	Type of box/Equilibrium/: <type box="" of=""></type>
	Only the EQUILIBRIUM type is allowed. Press <enter> to accept it.</enter>
	Is the stage heat controlled? /Y/: <yes no="" or=""></yes>
	Y means the current segment is heat controlled.
	Initial guess of temperature? /1000/: <initial temperature=""></initial>
	Give a guess of initial temperature (in K) for the current segment; press <enter> to accept the default.</enter>
	Give initial amount: <initial amount="" heat="" mass="" of="" or=""></initial>
	The initial amount of mass and/or heat in the current segment can be specified here. Two types of initial amount can be used:
	• As initial amount of heat (enthalpy): H=Value.
	• As initial amount of specific species or element and its initial temperature: N(SPECIES)=Value, Input temperature (in one line, or the input temperature in the next prompt).
	<pre>Input temperature /298.15/: <input temperature=""/></pre>
	Give initial amount: <initial amount="" heat="" mass="" of="" or=""></initial>
	Specify initial amount (mass or energy) in the current segment. Press <enter> to finish the initial amount inputs, and then this message displays:</enter>
	Each phase may have a separate output, give these
	Now start to define the output for each phase.
	Phase name /Rest/: <name of="" phase=""></name>
	Specify a phase that has a separate output, e.g. GAS; by pressing <enter> to accept that all REST phases do not have separate outputs. Then the program asks for further definitions of all the other segments (one by one), repeating</enter>

### some of the above questions, till the last segment in the reactor is defined.

# EXECUTE\_POLY3\_COMMAND



This is a command used with the REACTOR module.

### Syntax EXECUTE\_POLY3\_COMMAND

To directly use all kinds of POLY commands inside the REACTOR module. Any legal <u>"Equilibrium Calculations with the POLY\_3 Module" on page 157</u> can be entered.

# LIST\_RECORDS

This is a REACTOR\_SIMULATOR\_3 module command.

### Syntax LIST\_RECORDS

Lists all the already-created records for the reactor (including stage boxes, dividers, pipes, etc.).

### **READ\_WORKSPACE**

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This is a command used with the REACTOR module.

The REACTOR workspace (along with the related POLY3 and GIBBS workspaces) and the simulated results from the SIMULATE command can be read by this command from a binary file where they must be saved with <u>"SAVE\_WORKSPACE" below</u>. The \*.RCT file is not printable.

Syntax	READ_WORKSPACE
Options	File name: The name of a saved *.RCT file where the REACTOR and POLY3/GIBBS workspaces shall be read from must be specified. You do not need to type the extension if it is the default .RCT, otherwise type the whole *.RCT file name.
	An <b>Open</b> window displays if a file name is not given after the command or its path is incorrect, so that the path (in the <b>Look in</b> field) and <b>File name</b> can be specified. The <b>Files of type</b> (i.e. RCT) cannot be changed.

### SAVE\_WORKSPACE



This command is for the POLY3, GES (GIBBS), and REACTOR\_SIMULATOR\_3 modules.

The REACTOR (and POLY3/GIBBS) workspaces are saved on a file with this command. In the GIBBS workspace, all thermochemical data are stored. In the POLY3 workspace, all thermochemical data, all conditions, changed status, entered symbols, etc., are stored. In the REACTOR workspace, all definitions of multi-stage steady-state rector and its distribution coefficients are saved, together with the GIBBS and POLY3 workspaces. After a SAVE command, you can always return to the state when you issued the SAVE command by giving a READ command.

After saving the REACTOR (and POLY3/GIBBS) workspaces on a file, you can leave the program and at a later time READ the file and continue from the saved state.

A START\_SIMULATE command automatically saves on the work file with the most lately specified name. To avoid destroying the simulated results, do not SAVE after a START\_SIMULATE command is enforced, similar to that you should not SAVE after a STEP or MAP command! You may append several results by START\_SIMULATE without destroying the previous results but SAVE erases them all.

### Syntax SAVE\_WORKSPACE

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File name

A file name must be specified. The default extension of the RCT file is .RCT, while or any other extension as required.

A **Save** window displays if a file name is not given after the command, so that the path (in the **Save in** field) and **File name** can be specified. The **Files of type** (i.e. RCT) cannot be changed. If there is already a file with the same file name under the directory a warming message displays. Click **OK** to overwrite the current REACTOR (and POLY3/GIBBS) workspaces onto the existing \*.RCT file. Click **Cancel** to return to the REACTOR module. You can use the SAVE command with an unspecified name (i.e. through SAVE , , , command-sequence) to save the REACTOR (and POLY3/GIBBS) workspaces.

## **START\_SIMULATION**

This is a command used with the REACTOR module.

Start the reactor simulation, and list the results for all iterations. The output for each iteration consists of the conditions set in each segment, and you can also select some state variables to list. After each loop, the temperatures in all segments are listed.

```
      Syntax
      START_SIMULATION

      Prompts
      Max number of loops /10/: <The max number of loops>

      Give the max number of loops. The REACTOR module repeats the reactor simulation until this loop number is reached.

      Output to screen or file /Screen/: <File name or return>

      Decide on either listing simulation results on the screen (by pressing <Enter>) or saving them on an experimental file (with the default extension .EXP).

      Image: The conditions in each segment are not saved in the file, only on the screen.

      Output variables /T BP($)/: <Variable name(s)>

      Specify the variable name(s) that is also listed on the screen together with the conditions in each segment, or in the output experimental file.
```

#### **EXAMPLE OUTPUT**

For a four-stage reactor, the output at a certain iteration may look like this:

```
H=-991745.1, P=100000, N(C)=2.8468936, N(N)=6.78399966E-4, N(O)=3.8171615,
N(SI)=1.77031
DEGREE OF FREEDOM 0
BP(GAS)=43.464341, BP(C1SI1 BETA)=25.175981, BP(C S)=8.1588099,
BP(QUARTZ S3)=68.193896
1.78803E+03
BP(GAS)=61.971213, BP(C1SI1 BETA)=38.144346, BP(QUARTZ L)=66.230717 2.06569E+03
BP(GAS)=67.66262, BP(C1SI1 BETA)=35.480983, BP(QUARTZ L)=58.219292
2.08691E+03
BP(GAS)=71.23129, BP(SI L)=22.471787
2.20831E+03
H=-924751,87, P=100000, N(C)=2.0594558, N(N)=5.17999978E-4, N(O)=4.0142358,
N(SI)=2.7551438
DEGREE OF FREEDOM 0
BP(GAS)=43.464341, BP(C1SI1 BETA)=25.175981, BP(C S)=8.1588098,
BP(QUARTZ S3)=68.193896
1.78803E+03
```

```
BP(GAS)=61.971213, BP(C1SI1 BETA)=38.144346, BP(QUARTZ L)=66.230717 2.06569E+03
BP(GAS)=67.66262, BP(C1SI1_BETA)=35.480983, BP(QUARTZ_L)=58.219292
2.08691E+03
BP(GAS)=71.23129, BP(SI L)=22.471787
2.20831E+03
H=-813239., P=100000, N(C)=1.6592668, N(N)=3.59999988E-4, N(O)=3.7549293,
N(SI)=2.896635
DEGREE OF FREEDOM 0
BP(GAS)=43.464341, BP(C1SI1_BETA)=25.175981, BP(C_S)=8.1588098,
BP(QUARTZ S3)=68.193896
1.78803E+03
BP(GAS)=61.971213, BP(C1SI1 BETA)=38.144346, BP(QUARTZ L)=66.230717 2.06569E+03
BP(GAS)=67.66262, BP(C1SI1 BETA)=35.480983, BP(QUARTZ L)=58.219292
2.08691E+03
BP(GAS)=71.23129, BP(SI L)=22.471787
2.20831E+03
H=7374.6403, P=100000, N(C)=8.84911857E-1, N(N)=1.99999996E-4, N(O)=1.9379203, N
(SI)=1.853872
DEGREE OF FREEDOM 0
BP(GAS)=43.464341, BP(C1SI1 BETA)=25.175982, BP(C S)=8.1588097,
BP(QUARTZ S3)=68.193896
1.78803E+03
BP(GAS)=61.971213, BP(C1SI1 BETA)=38.144346, BP(QUARTZ L)=66.230717 2.06569E+03
BP(GAS)=67.66262, BP(C1SI1 BETA)=35.480983, BP(QUARTZ L)=58.219292
2.08691E+03
BP(GAS)=71.23129, BP(SI L)=22.471787
2.20831E+03
10 1788.03 2065.69 2086.91 2208.31 0.00 0.00 0.00
```

### **SCHEIL Module Commands**

Search the online help or see the *Thermo-Calc User Guide* included with this PDF documentation set for details about how to calculate using Scheil either in Graphical Mode (the Scheil Calculator) and Console Mode (the wizard Scheil module).



If you also have a license for the Diffusion Module (DICTRA), you can import a previously calculated Scheil segregation profile into the software using the command INPUT\_SCHEIL\_PROFILE. Search the online help or see the *Diffusion Module (DICTRA) Command Reference Guide*.

#### In this section:

BCC_TO_FCC_TRANSFORMATION	291
EVALUATE_SEGREGATION_PROFILE	291
GLOBAL_MINIMIZATION	292
LIQUID_PHASE_NAME	292
SAVE_FILE_NAME	292
SET_INTERACTIVE	292
SET_NUMERICAL_LIMITS	293
START_WIZARD	294
TEMPERATURE_STEP	295
TERMINATION_CRITERIA	295

# **BCC\_TO\_FCC\_TRANSFORMATION**

Use this command for the Scheil module to allow transformations in the solidified part of the alloy caused by each of the components specified to be Fast Diffuser. It is recommended that you only select this for steels.

Syntax	BCC_TO_FCC_TRANSFORMATION
Descent	Allow bcc $\rightarrow$ fcc ? /N/
Prompt	Enter Y to allow transformations.

# **EVALUATE\_SEGREGATION\_PROFILE**

Use this command in order to evaluate a segregation profile while performing the Scheil calculation. The Scheil calculation itself does not contain a spatial dimension. However, if it is assumed that the molar volume is constant and equal for all elements, and if a length is provided over which solidification occurs (for example the secondary dendrite arm spacing) then the generated profile corresponds to the segregation that would be obtained under the assumptions of a Scheil calculation. The grid is then equidistant and can be used as input for a DICTRAsimulation.

The generated text file contains, on each line, the mole fractions of all elements in alphabetical order at a certain grid point. If the composition is the same on several consecutive lines it means that the composition of the then remaining liquid has reached a state in which there are no degrees of freedom.

Syntax	EVALUATE_SEGRATION_PROFILE			
	Evaluate segregation profile? /N/			
Prompt If the answer is Y, a segregation profile will be evaluated and to disc. Note that this feature is not supported with back diffusion is enabled no segregation profile will be generated and the segreg				
	Number of grid points /100/			
	<ul> <li>Enter the number of grid points for the segregation profile. A file dialogue window opens and allows you to specify the name of the file in which the segregation profile is written.</li> <li>In a macro file it is possible to enter all arguments on a single line, e.g.</li> <li>EVALUATE_SEGREGATION_PROFILE Y 100 XF.TXT, meaning that the segregation profile is evaluated over 100 grid points and saved to a file (XF.TXT) in the current working directory.</li> </ul>			

## **GLOBAL\_MINIMIZATION**

Use this command for the Scheil module to use global minimization.



Also see the POLY commands topic <u>"About the ADVANCED\_OPTIONS Command"</u> on page 162 for information about that GLOBAL\_MINIMIZATION command.

Syntax	GLOBAL_MINIMIZATION
	Use global minimization? /N/ The default (N) does not use global minimization. If you choose Y, the Global
Prompt	

# LIQUID\_PHASE\_NAME

Use this command for the Scheil module to use the liquid phase name from the database definitions or enter another phase name.

	Syntax	Syntax LIQUID_PHASE_NAME			
		Liquid phase name /Default/			
	Prompt	Default selects the liquid phase from the database definitions, otherwise enter a phase name.			

# SAVE\_FILE\_NAME

Use this command for the Scheil module to save the file. The files get the console number added to this name, e.g. 005.->Scheil\_006.poly3 and Scheil\_segregation\_profile.txt).

Syntax	SAVE_FILE_NAME
Prompt Name of files to be saved, poly3 and segregation profiles.	

# **SET\_INTERACTIVE**

Syntax	SET_INTERACTIVE
	Resets the input and output units to the initial values, i.e. keyboard and screen. Add this as the last command to the macro files.

# **SET\_NUMERICAL\_LIMITS**

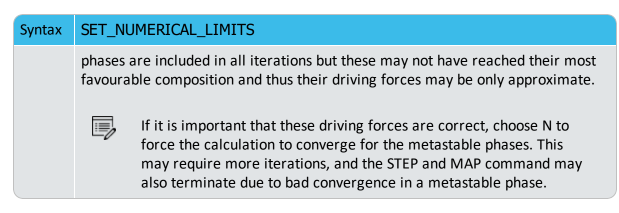
Change the criteria for convergence. This is to speed up a calculation in a complex system.

If there is any convergence problem in finding a stable solution at any stage during a calculation procedure enforced by <u>"COMPUTE\_EQUILIBRIUM" on page 182</u>, <u>"STEP\_WITH\_</u> <u>OPTIONS" on page 217</u>, <u>"MAP" on page 201</u> or <u>"ADVANCED\_OPTIONS" on page 167</u>), this message displays:

Convergence problems, increasing smallest site-fraction from 1.00E-30 to hardware precision 2.00E-14. You can restore using SET-NUMERICAL-LIMITS

This implies that smallest site fraction in the current POLY3 workspace is automatically increased from the default value 1.00E-30 to the hardware-dependent precision (under Linux, as 2.00E-14). For other subsequent POLY-module calculation in Console Mode, you can use this command to restore or reset the smallest site fraction to the previous or another preferred value, as well as to reset other numerical limits.

Syntax	SET_NUMERICAL_LIMITS			
	Maximum number of iterations /200/: <xxx></xxx>			
Prompt	By default, the program tries 500 iterations before it gives up. As some models give computation times of more than 1 CPU second/iteration, this number is also used to check the CPU time and the calculation stops if 500 CPU seconds/iterations are used.			
	Required accuracy /1E-6/: <yyy></yyy>			
	This is a relative accuracy, and the program requires that the relative difference in each variable must be lower than this value before it has converged. A larger value normally means fewer iterations but less accurate solutions. The value should be at least one order of magnitude larger than the machine precision.			
	<pre>Smallest fraction /1E-12/: <zzz></zzz></pre>			
	This is the value assigned to constituents that are unstable. It is normally only in the gas phase you can find such low fractions.			
	The default value for the smallest site-fractions is $1E-12$ for all phases, except for IDEAL phase with one sublattice site (such as the GAS mixture phase in many databases) for which the default value is always as $1E-30$ (unless you have used the SET_NUMERICAL_LIMITS command to reset an even-lower value (e.g. $1E-45$ , that is naturally enforced to all the phases in the system).			
	Approximate driving force calculation for metastable phases /Y/:			
	Normally the POLY module only requires that the set of stable phases is in equilibrium in order to terminate the iterations. By default, the metastable			



### START\_WIZARD

Use this command for the Scheil module to use the wizard with step-by-step prompts to create a Scheil simulation.

This wizard walks you through a basic Scheil simulation. You are prompted for settings including the major element or alloy, composition input in mass (weight) percent, the alloying elements, the temperature, fast diffusing components, and whether any phase should have a miscibility gap check.

Use the other commands to make changes to settings that are not included in this wizard.

- BCC\_TO\_FCC\_TRANSFORMATION
- EVALUATE\_SEGREGATION\_PROFILE
- GLOBAL\_MINIMIZATION
- LIQUID\_PHASE\_NAME
- SAVE\_FILE\_NAME
- STORED\_COMPOSITION\_TYPE
- TEMPERATURE\_STEP
- TERMINATION\_CRITERIA

### Syntax START\_WIZARD

	Database / <default>/:</default>
	Major element or alloy:
	Composition input in mass (weight) percent? /Y/:
Prompt	1st alloying element: Cr 10 C 1
	Next alloying element:
	Temperature (C) /2000/:
	Fast diffusing components: /None/: C

Syntax	START_WIZARD				
	Should any phase have a miscibility gap check? /N/:				

## **TEMPERATURE\_STEP**

Use this command for the Scheil module to define a temperature step.

Syntax	TEMPERATURE_STEP		
Prompt	Set temperature step	(C)	/1/

### **TERMINATION\_CRITERIA**

Use this command for the Scheil module to define the termination criteria for the fraction of liquid or temperature.

```
      Syntax
      TERMINATION_CRITERIA

      Prompts
      Fraction of liquid or Temperature (F or T)? /F/

      Fraction of liquid /.01/
```

# System Utilities Commands

### In this section:

ABOUT	
CLOSE_FILE	
DISPLAY_LICENSE_INFO	
HP_CALCULATOR	
OPEN_FILE	
SET_COMMAND_UNITS	
SET_ECHO	
SET_GES_VERSION	
SET_INTERACTIVE_MODE	
SET_LOG_FILE	
SET_PLOT_ENVIRONMENT	
SET_TC_OPTIONS	
STOP_ON_ERROR	

# ABOUT

Syntax	ABOUT
	Lists basic information about the development history and ownership of the Thermo-Calc software as well as version information.

# **CLOSE\_FILE**

Close an opened text file.

Syntax	CLOSE_FILE
Duanat	Unit number: <unit number=""></unit>
Prompt	A unit number given in an OPEN_FILE command must be specified.

# DISPLAY\_LICENSE\_INFO

Use this command to launch a license diagnostics program implemented in the SYS module. It detects the license information for the Thermo-Calc installations on the current client computer (and on the connected license server), and lists the details, including the current status of available license file and all its included license features for the current installation (s), as well as the Safenet Environment Variables and detailed Local (Client) Information, on screen or into a textual file (\*.TXT).

This is useful for a list of license details, and to obtain basic technical support by sending the \*.TXT file to <u>support@thermocalc.se</u>. If you are running a network client installation of Thermo-Calc, you can see how many of the client example licenses that are currently checked out and how many of them are left. You can also see which clients computers it is that have checked these client example licenses out.

Syntax	DISPLAY_LICENSE_INFO
	Output to screen or file /Screen/: <mylicenseinfo.txt></mylicenseinfo.txt>
Prompt	Press <enter> to display a list of license information or into a *.TXT file under a specific file name which can later on be opened and edited by any basic text editor.</enter>

# **HP\_CALCULATOR**

This is the QBA simple interactive calculator using reverse Polish notations.

### Syntax HP\_CALCULATOR

Available OPCODEs (HPC codes) are listed by entering the command HELP.

Use the BACK command to quit the HP\_CALCULATOR and go back to the SYS module.

Input are numbers, + - \* / and  $^ and OPCODEs$ . Several numbers an operations can be given on one line. The content of the X register is displayed after each operation.

For example, to compute 2\*EXP(1.5\*\*3-30000/(8\*1273)), you should input the following: 30000 8/1273/chs 1.5 3 ^ + exp 2 \*

### **OPEN\_FILE**

A text file is opened for use in other commands where a unit number is necessary. The program automatically assigns a unit number.

Syntax	OPEN_FILE
Durant	File name: <file name=""></file>
Prompt	A legal file name must be specified.

# **SET\_COMMAND\_UNITS**

This command is useful for reading inputs already prepared by a textual editor on a file. Such inputs can be a table of values or a large number of parameters. The file must be opened with "OPEN\_FILE" above.



The first two lines of an input file are skipped before any input is read.

Syntax	SET_COMMAND_UNIT
Prompt	Input unit number /5/: <input number="" unit=""/> Specify the input unit number returned from an OPEN_FILE command. The next command is taken from the file connected to this unit number. The last command on such a file must be EXIT or SET_INTERACTIVE when you are in POLY, POST or SYS module, in order to go back to read input from the keyboard. The default value is the current input unit.
	Output unit number /6/: <output name="" unit=""> Specify the output unit number returned from an OPEN_FILE command.</output>

## **SET\_ECHO**

Syntax	SET_ECHO
	Add this command at the beginning of a Console Mode MACRO *.TCM file (or to the primary MACRO file on the top level if any sub-level(s) of MACRO files are used).
	Automatically displays on screen the complete details of various commands in all the sequential operations in the software, that are enforced according to the MACRO file(s).
	This command is not applicable if MACRO files are not used.

# **SET\_GES\_VERSION**

8

This command enables switching between the default standard version (5) Gibbs Energy System (GES) calculation engine and an experimental version (6).

If you are interested in participating and helping to test the experimental version of the calculation engine, please contact Thermo-Calc Software AB at <u>info@thermocalc.se</u> for more information about the changes. Make sure you compare/check your results as GES6 continues to be developed and improved.

### Syntax SET\_GES\_VERSION

	USE	GES	VERSION	5	OR	6	/5/:
Prompt	The	defa	ult is versi	on	5.		

# **SET\_INTERACTIVE\_MODE**

### Syntax SET\_INTERACTIVE\_MODE

Resets the input and output units to their initial values, i.e. keyboard and screen.

Remember to add this as the last command to your MACRO files, in order to stop execution of the command file in the POST module. It has no meaning in interactive mode.

# SET\_LOG\_FILE

Use it to save, in a basic text file, everything typed into the Thermo-Calc software in Console Mode. When having problems executing a command sequence, this command can be used to save the command typed onto a log file with the file extension \*.TCM.

This command also makes the system echo the full command for all commands typed. This feature is useful when demonstrating the system since the abbreviated commands are often difficult.

The saved log file can then be edited as a macro file by using a basic text editor. This is useful to run the macro file(s) for similar calculations (the same system but slightly varied temperature-pressure-composition conditions), or run the example macro files.

▶ <u>"M</u>	acro Files" on page 174 in the Thermo-Calc User Guide.
Syntax	SET_LOG_FILE
Promot	A <b>Save</b> window opens to specify a file name for the *.TCM file, so that the path (in the <b>Save in</b> field) and <b>File name</b> can be specified. If a log file with the same

(i.e. TCM) cannot be changed. Click Save or Cancel as applicable.

## SET\_PLOT\_ENVIRONMENT

8\$B

Allows you, preferably in the initiation file tc.ini, to set the plot devices you normally have access to as this can vary for different Thermo-Calc installations.

name exists in the current working directory, it is overwritten. The Files of type

This command must be terminated with an empty line or two commas.

Syntax	SET_PLOT_ENVIRONMENT
	Default plotdevice number /Default number/: <device number=""></device>
Prompts	The number given is unique for each type of graphical device. The default graphical device number should normally be selected. A question mark ? gives you a list of all available graphical devices used in Console Mode.
	Pseudo file name: <pseudo-file name=""></pseudo-file>
	The name given here is a symbol which can be used to refer to a physical graphics device when asked for graphical output file.
	Plotdevice number /1/: <device number=""></device>

### Syntax SET\_PLOT\_ENVIRONMENT

The number that specifies the type of graphical device.

Plot file name: <File name or printer name>

The name of the graphical device on system level (name of file or printer).

#### **EXAMPLE**

SET\_PLOT\_ENVIRONMENT 1 lasp 5 a0tr,,

This example sets the default plot device number to 1, and defines an alias named lasp with plot device 5 and connected to a printer named a0tr.

### **SET\_TC\_OPTIONS**

Predefine the default values of three general Thermo-Calc options for performing all the sequential single-points, stepping and mapping calculations.

Syntax	SET_TC_OPTIONS			
	Use_global_minimization /Y/:			
Prompts	By default it global minimization is used where possible (depending on the conditions setting). A global minimization test is performed when an equilibrium is reached. This costs more computer time but the calculations are more robust. Enter N to prevent global minimization being used to reach the equilibrium.			
	Global_test_interval /0/: <an integer="" value=""></an>			
	The integer number determines how often the Global Minimization should be used during calculations with <u>"STEP_WITH_OPTIONS" on page 217</u> and <u>"MAP"</u> on page 201. If it is set to 0 (zero), the recommended global test interval is used: i.e. every tenth step and at each phase change during STEP calculations, and only at node points during MAP calculations. Any other positive integer number, n, suggests it performs Global Minimization tests at every n <sup>th</sup> step during STEP and MAP calculations. Of course, the Global Minimization test is always carried out at a phase change or a node point. The Thermo-Calc Console Mode is initially configured with 0 (zero) as the default value, but such a configuration for the default value can be changed by this option.			
	Automatically_add_initial_equilibria /Y/: When <u>"ADD_INITIAL_EQUILIBRIUM" on page 160</u> is not used before a MAP calculation, a mesh of INITIAL_EQUILIBRIUM is automatically added before the mapping procedure itself takes place. This mesh is only available when the two independent axis-variables have already been defined by the POLY command.			

### Syntax SET\_TC\_OPTIONS

Set\_axis\_variable

If Y is entered, the POLY module always enforces the automatic procedures for adding initial equilibrium points during mapping. For N, no initial equilibrium points are then added during mapping.

## **STOP\_ON\_ERROR**

Useful for batch jobs in order to prevent that an erroneous sequence of commands to the program causes waste of computer activity.

Syntax	STOP_ON_ERROR
	Give the preferred argument ( $ON$ or $OFF$ ) after the STOP command. If the default argument ON is given, the program terminates after an illegal or ambiguous command. It is possible to reset the effect of this command by giving the argument OFF.

### **TABULATION\_REACTION Commands**



"Tabulation of Chemical Substances, Phases or Reactions" on page 224 in the *Thermo-Calc User Guide*.

#### In this section:

ENTER_FUNCTION	304
ENTER_REACTION	305
LIST_SUBSTANCES	307
SET_ENERGY_UNIT	309
SET_PLOT_FORMAT	309
SWITCH_DATABASE	310
TABULATE	311
TABULATE_DERIVATIVES	311
TABULATE_REACTION	316
TABULATE_SUBSTANCE	319

## **ENTER\_FUNCTION**

Used to define a new thermodynamic function to be included in the sixth column of tabular output. The last entered function is listed in the 6th column of all tables, and can be plotted (if choosing the Plot Column value 6) in all graphs, for all subsequently tabulated substances or reactions.

Syntax	ENTER_FUNCTION
Prompt	Name: <name> Function: <definition></definition></name>
	A limited number of state variables, G, H, S, T, P, V and H298 can be used in the function definition. For example, -(G-H)/T can be defined as a function which is identical to S for a substance or $\Delta$ S for a reaction, G+T*S-P*V can be defined as a function which is U (internal energy) for a substance or $\Delta$ U (internal energy change) for a reaction.
Options	Name of the function (maximum 8 characters) and Definition of the function. A long function can be typed over several lines. A function should be terminated by a semicolon (;) or an empty line.

#### EXAMPLE OUTPUT

For the pure substance Fe (using the SSUB database), by entering a function as G+T\*S-P\*V, the following table is obtained by typing <Enter> in the prompt Output file /Screen/:, and the figure by typing 6 in the prompt Plot column /2/:

```
TAB: ENTER-FUNCTION
Name: InEnergy
Function: G+T*S-P*V;
TAB: TABULATE_SUBSTANCE
Substance (phase): Fe
Pressure /100000/: <Enter>
Low temperature limit /298.15/: <Enter>
High temperature limit /2000/: <Enter>
Step in temperature /100/: <Enter>
Output file /try1/: try2
Graphical output? /Y/: <Enter>
Plot column? /2/: 6
```

#### This gives the following output:

Ουτρυ	IT FROM T	HERMO-C	CALC		
Column 6:	InEnergy G+T*	S-P*V			
Phase : F	`E_S	Pr	essure :	100000.00	
Specie: F	Έ				
******	*****	*****	****	* * * * * * * * * * * * *	
Т	Ср	Н	S	G	InEnergy
(K)	(Joule/K)	(Joule)	(Joule/K)	(Joule)	
******	*****	* * * * * * * * * * * * * * *	*****	*****	
298.15	2.48446E+01	2.17972E-06	2.72800E+01	-8.13353E+03	2.17972E-06
300.00	2.48905E+01	4.60049E+01	2.74338E+01	-8.18414E+03	4.60049E+01
400.00	2.71299E+01	2.64957E+03	3.49085E+01	-1.13138E+04	2.64957E+03
500.00	2.93561E+01	5.47211E+03	4.11976E+01	-1.51267E+04	5.47211E+03
600.00	3.19293E+01	8.53245E+03	4.67701E+01	-1.95296E+04	8.53245E+03
700.00	3.50985E+01	1.18777E+04	5.19207E+01	-2.44667E+04	1.18777E+04
800.00	3.92042E+01	1.55830E+04	5.68623E+01	-2.99068E+04	1.55830E+04
900.00	4.49645E+01	1.97726E+04	6.17903E+01	-3.58387E+04	1.97726E+04
1000.00	5.42147E+01	2.46891E+04	6.69619E+01	-4.22728E+04	2.46891E+04
1100.00 Stable ph	4.55851E+01 ase is FE_S2	2.99025E+04	7.19412E+01	-4.92328E+04	2.99025E+04 \$
1200.00	3.40840E+01	3.51037E+04	7.64466E+01	-5.66322E+04	3.51037E+04
1300.00 1400.00	3.49398E+01 3.57994E+01	3.85549E+04 4.20918E+04	7.92086E+01 8.18293E+01	-6.44162E+04 -7.24692E+04	3.85549E+04 4.20918E+04
1500.00	3.66636E+01	4.57149E+04	8.43287E+01	-8.07780E+04	4.57149E+04
1600.00 Stable ph	3.75330E+01 ase is FE_S3	4.94247E+04	8.67226E+01	-8.93314E+04	4.94247E+04 \$
1700.00	4.05217E+01	5.41173E+04	8.95609E+01	-9.81363E+04	5.41173E+04
1800.00 Stable ph	4.12595E+01 ase is FE_L	5.82055E+04	9.18975E+01	-1.07210E+05	5.82055E+04 \$
1900.00 2000.00	4.60000E+01 4.60000E+01	7.74165E+04 8.20165E+04	1.02377E+02 1.04736E+02	-1.17099E+05 -1.27456E+05	7.74165E+04 8.20165E+04

## **ENTER\_REACTION**

This is the same as <u>"TABULATE\_REACTION" on page 316</u> in that both the chemical reaction can be specified and the thermodynamic property changes for the reaction can be generated as a table, and plotted as a graph or saved as an experimental file (with an extension of .EXP). The thermodynamic data are automatically retrieved from the SSUB5 database (by default), or the current database set by <u>"SWITCH\_DATABASE" on page 22</u> (thus

there is no need to use the DATA module before this command). You can select any other database by using the SWITCH\_DATABASE command in the TAB module. The rules to specify a reaction are the same as in the command TABULATE\_REACTION.

Syntax	ENTER_REACTION
	Same reaction? /Y/:
Prompts	This displays if there is at least one reaction already defined either by the command ENTER_REACTION or TABULATE_REACTION. For Y the next prompt for defining reaction is not asked.
	Reaction: <chemical equation="" reaction=""> &amp; <ra+rb=pc+pd></ra+rb=pc+pd></chemical>
	A long reaction can be typed over several lines. A reaction should be terminated by a semicolon (;) or an empty line. A list showing the used database, defined elements, getting-data sequence, references, etc.:
	Pressure /100000/: <pressure in="" interest,="" of="" pa=""></pressure>
	Low temperature limit /298.15/: <t-low, in="" k=""></t-low,>
	High temperature limit /2000/: <t-high, in="" k=""> Step in temperature /100/: <t-step></t-step></t-high,>
	Output file /Screen: <return a="" file="" for="" name="" or="" screen,="" type=""></return>
	If you type Return for screen a list of thermodynamic property changes for the defined/chosen reaction is shown up as a table (which is demonstrated in the example outputs given below), and the command is terminated.
	If typing a file name, the table is displayed and saved as an *.EXP or an *.TAB file. Then prompts:
	Graphical output /Y/:
	For N a table is output to the screen and this table is saved as a basic text file with the default extension .TAB under the current working directory. In this case, the graph cannot be plotted.
	By default a table is created with all thermodynamic functions as normal (which is displayed on screen) and generates a graph with temperature on the X-axis and a chosen property on a certain column in the table on the Y-axis (which is plotted on screen and saved as an *.EXP file). You are also prompted for the plot column.
	Plot column? /2/: <1 or 2 or 3 or 4 or 5 or 6>
	Specify which property (as column number) to be plotted as the Y-axis (versus temperature as X-axis) as a graph on the screen. Simultaneously, all tabulated properties and the Y-axis setting (i.e. the plotted column) for the graph is

### Syntax ENTER\_REACTION

written into an \*.EXP file using the DATAPLOT format. The default column 2 is heat capacity, 3 enthalpy, 4 entropy and 5 Gibbs energy, and the additional column 6 is a user-entered function. The table displays on the screen. The plot then displays on the screen, followed by a POST prompt. The POST module (postprocessor) automatically opens, and all types of the POST-module commands to refine the plotted diagram may be used. Possibilities include scaling the X/Y-axis, changing the X/Y-axis texts, etc. The command BACK or EXIT at the POST prompt always takes you back to the TAB module.

## LIST\_SUBSTANCES

>\_

This is a TAB module command.

List all species in the current database with a certain set of elements. This is useful to specify species as pure substances in the command <u>"TABULATE\_SUBSTANCE" on page 319</u>.

Syntax	LIST_SUBSTANCES
	With elements /*/ <* or a set of elements>
Prompts	The elements building up various species must be specified. The wildcard * means all elements in the current database.
	If some elements are specified, it also prompts for how to list species with such elements.
	Exclusively with those elements /Y/:
	All species (not phases) in the current database are searched. Keep the default (Y) if you do not want other elements included in the search except for those specified. Enter N to list all species containing at least one of the specified elements.

#### EXAMPLE OUTPUT

For the SSOL database, there are different lists, as shown below:

TAB: 1	sub	
With e	elements /*/: <enter></enter>	
VA	AG	AL
AM	AS	AU

В	ВА	BE BI BR C
СА	CD	CE
CL	CO	CR
CS	CU	DY
ER	EU	F
FE	GA	GD
GE	Н	HF
HG	НО	I
IN	IR	K
LA	LI	LU
MG	MN	MO
NA	NB	
ND	NI	NP
OS	P	
PA	PB	PD
PR	PT	PU
RB	RE	RH
RU	S	SB
SC	SE	SI
SM	SN	SR
ТА	TB	TC
TE	TH	TI
TL	ТМ	U
V	W	Y
YB	ZN	ZR
C1	C2	C3
C4	C5	C6
C7	N2	02
P1	P2	P4
NA/+1	K/+1	CL/-1
H2O	H1/+	LI1/+
CS1/+	RB1/+	01H1/-
F1/-	BR1/-	I1/-
H1/+1	H101/-1	SI102 CA101 TAB:
TAB: l-sub		
With elements /*/: fe	cr ni c n o	

Exclusively wit	th those elements?	/Y/:	<enter></enter>
С	CR		FE
Ν	NI		0
С	C2		C3
C C5		C6	
С	N2		02
TAB:l-sub fe c	rnicno		
Exclusively wit	th those elements?	/Y/:	n
С	CR		FE N NI O
C1	C2		C3
C4	C5		C6
C7	N2		02
Н2О	01H1/-		H101/-1 SI102 CA101 TAB:

# **SET\_ENERGY\_UNIT**

Set the energy unit as calories or Joule, in all the subsequent outputs (tables, graphs, and files) in the current calculation operation.

Syntax	SET_ENERGY	_UNIT			
Prompt	Energy unit	(C=Cal,	J=Joule)	/J/:	<unit></unit>

# SET\_PLOT\_FORMAT

Set the plotting format while the result is plotted on screen or the file is saved as EXP file [using DATAPLOT graphical format] and TAB file (as a basic text file and as displayed on screen).

This command is different from <u>"SET\_PLOT\_FORMAT" on page 254</u> in the POST module, but is identical to <u>"SET\_PLOT\_ENVIRONMENT" on page 300</u> in the SYS (system utility) monitor.

Syntax	SET_PLOT_FORMAT
	The default set Unitis 1 (Windows) or 9 (Linux) for graphical presentation on screen, or all other units for outputs as *.EXP and *.TAB files.
Options	Also set the format of the graphical output to different graphical devices. The default unit 22 (1 for legacy Windows, and 9 for legacy Linux) may be altered with the SET_PLOT_ENVIRONMENT command in the SYS monitor or by the TC.INI file.

# SWITCH\_DATABASE

By default, the TAB module always automatically retrieves thermodynamic data for substances or reactant/product substances in a defined reaction, or for pure phases or ideal solution phases, from the SSUB Substances Database. However, you can select thermodynamic data for complex non-ideal solution phases from another database, which is possible with this command.

Syntax	SWITCH_DATABASE
	Database /XYZ/: <new database="" name=""></new>
	Specify a New database name. XYZ stands for the default SSUB database or the current database which is switched on.
Prompt	If an appropriate database name is not given and the USER option is not used, a list displays of available databases.
riompe	Use one of these databases:
	• PURE = SGTE Pure Elements Database
	<ul> <li>SSUB = SGTE Substances Database</li> </ul>
	<ul> <li>USER = user defined database</li> </ul>

This command switches (or changes) from the current database to a new one, and reinitializes the entire TAB module for defining a substance or reaction and the GIBBS workspace for storing retrieved data. All the directly connected databases as predefined by the original database initiation file (TC\_INITD.TDB file in the /DATA/ area for Windows environments, or the initd.tdb file in the \data\ area for Linux) are listed by pressing the <Enter> key without giving any argument. You can also supply your own database by giving the argument USER and then the database name and, if it is not located in the current working directory, its correct path.

This command (when required to use another database for subsequent tabulations) must be used before executing the TAB commands <u>"TABULATE\_SUBSTANCE" on page 319</u> for substances, and <u>"ENTER\_REACTION" on page 305</u> or <u>"TABULATE\_REACTION" on page 316</u> for reactions.

When a predefined database or an appropriate USER database is switched on, the DATA command <u>"GET\_DATA" on page 16</u> is automatically executed.

DATABASE NAME: The new database is specified by typing the abbreviation given in the front of one of the available predefined databases. For convenience when switching/appending databases you can add them to the predefined database list in the database initiation file TC\_INITD.TDB or initd.tdb of your installed software.

When this command is used in a MACRO (\*.TCM) file, if the USER option is selected, the database setup file name (\*setup.TDB) containing the setup definitions of the USER database, and its correct path, must be provided.

If a USER database name or its path is not given on the same line of the SWITCH\_DATABASE command, or if it is incomplete or incorrect, an Open window displays to specify the filename of the database setup file name (\*setup.TDB) of the to-be-switched USER database.



Unlike in the DATA module, if a USER database is chosen, the Gibbs energy system is also reinitialized and only data from the USER databases are retrieved for tabulations.

After this command, those commands to define systems (in terms of either elements or species), to reject/restore phases or species, retrieve data, as well as to append data from additional database(s), can be done.

## **TABULATE**

Gives the values of an already-defined table (with various properties varied with the stepping axis variable) from equilibria calculated with a STEP command. You can tabulate a table even if the table is entered after the STEP calculation.

Syntax	TABULATE
Dromoto	Name: <name a="" defined="" of="" table=""></name>
Prompts The name of the table must be given. The table must be enter	
	Output to Screen or file /Screen/: <file name=""></file>
	The table can be listed on the screen (by pressing <enter>), on a file (the file name must be specified here).</enter>

# TABULATE\_DERIVATIVES

This command is mainly for system debugging purposes and for experienced users. It automatically calculates all partial derivatives for a pure phase or solution phase at a given composition.



This is not the same as the chemical potential.

Before using this command, the system must be defined and thermodynamic data retrieved from an appropriate solution database in the DATA module. The phase name can be given in either upper, lower or mixed cases, and can be abbreviated until it is unique, e.g. fcc, BCC, cem, Liq, etc. as the SSOL database is used for the tabulations, or GAS, FE-S, wustite, Fe2O3-

hem, etc. as the SSUB database is used. The module then optionally goes through a specific sublattice or all the possible sublattices for the specified phase, as well as through the whole framework of a currently defined system (including all the defined elements/species, as well as the default defined vacancy and electron if necessary for the current database), when it prompts for inputs of site fractions for various species possibly occupying such sublattices. Therefore, thermodynamic properties can be tabulated for a pure phase, or for a solution phase as a pure end-member (with a composition definition for the corresponding non-interacting species on all necessary sublattices in the end-member), or for a solution phase as a real solution (with a composition definition for related interacting species on all necessary sublattices in the solution).

TABULATE_DERIVATIVES
Phase name /XXXX/: <name of="" phase="" the=""></name>
Specify the phase name of a pure phase or a solution phase.
xxxx is the last phase the TAB module accounted for. It automatically prompts for other options and necessary inputs for the definition of the composition of the specified phase.
For a pure phase (such as Fe-S, Wustite, and Fe2O3-Hematite), no further composition definition is needed. For a solution phase with one sublattice (such as AQUEOUS solution, GAS mixture and SLAG solution), it requests <i>n</i> -1 site-fraction inputs for the phase constituents (if there are <i>n</i> species in the defined phase within the whole framework of currently defined system, including all the defined elements/species, as well as the default defined vacancy and electron if necessary for the current database); the <i>n</i> th species is automatically assigned as the rest.
The sum of input site fractions must not exceed unity.
Example
For example, these prompts and inputs can be seen for a LIQUID solution phase (from the SSOL database) within the Fe-Cr-Ni-C-N-O framework (Note that by default, the unprompted Ni species are assigned as the rest):
FRACTION OF CONSTITUENT (RETURN FOR PROMPT): <return> C /1/: .05</return>
CR /1/: .1
FE /1/: < RETURN>
SUM OF FRACTIONS EXCEED UNITY, PLEASE REENTER

### Syntax TABULATE\_DERIVATIVES N /1/: .005 For a solution phase with two or more sublattices (such as FCC alloy solution, and ION LIQ ionic liquid solution), it first asks for which sublattice the sitefractions of constituents shall be specified: the default value 0 for all possible sublattices, and a given positive number for a specific sublattice (of course, this number must be reasonable for the currently specified phase, i.e. it must be smaller than the total sublattice number of the phase). Then, for the given sublattice or for all sublattices, it prompts for all the necessary inputs of sitefractions for the possible constituents on each sublattice n-1 times if there are n species on such a sublattice in the defined phase; the nth species on each of the sublattices is automatically assigned as the rest. For example, the following prompts and inputs can be seen for an FCC solution phase (from the SSOL5 database) within the Fe-Cr-Ni-C-N-O framework (Note that by default, the unprompted O species on its sublattice 1 and VA on its sublattice 2 are assigned as the rest): SPECIFY SUBLATTICE (0 FOR ALL) /0/: <RETURN> FRACTIONS IN SUBLATTICE 1 CR /1/: .1 FE /1/: .8 NI /1/: .0995 FRACTIONS IN SUBLATTICE 2 C /1/: .05 N /1/: .05 Temperature /2000/: <Temperature of interest, in K> Specify the temperature of interest in K. Pressure /100000/: <Pressure of interest, in pa>

Specify the pressure of interest in Pa.

### EXAMPLE OUTPUT

For the FCC phase in the Fe-Cr-Ni-C-N-O system (using the SSOL database), the following table is obtained at a specified composition:

TAB: TAB\_DER Phase name /BCC/: FCC SPECIFY SUBLATTICE (0 FOR ALL) /0/: <Enter>

```
FRACTIONS IN SUBLATTICE 1
CR /1/: .1
FE /1/: .8
NI /1/: .3
SUM OF FRACTIONS EXCEED UNITY, PLEASE REENTER
NI /0/: .0995
FRACTIONS IN SUBLATTICE 2
C /1/: .05
N /1/: .05
Temperature /1800/: <Enter>
Pressure /100000/: <Enter>
Gibbs energy: ..... -1.27432533E+05
Helmholz energy: ..... -1.27433205E+05
Enthalpy: ..... 5.95773994E+04
Internal energy: ..... 5.95767279E+04
Entropy: ..... 1.03894407E+02
Molar volume: ..... 6.71473258E-06
Thermal expansivity: ..... 7.63886416E-05
Isothermal compressibility: ..... 6.02925387E-12
Heat capacity at constant pressure: 4.33555074E+01
First partial derivative with respect to CR in sublattice 1 of Gibbs energy:
..... -1.26034739E+05 of enthalpy: .....
                                                         of volume:
4.63000206E+04 of entropy: ..... 9.57415334E+01
..... 6.87203263E-06
Second partial derivative of Gibbs energy with respect to also
 CR
                     in 1: 1.54392858E+05
 FΕ
                     in 1: -1.53353158E+04
                     in 1: -1.71750366E+04
 NI
 0
                     in 1: 0.0000000E+00
 С
                     in 2: -1.82016870E+05
in 2: -3.73062665E+05
 VA
                     in 2: -9.36260878E+04
irst partial derivative with respect to FE in sublattice 1 of Gibbs energy:
.....-1.02869265E+05
   of enthalpy: ..... 6.11738912E+04
                                             of entropy:
7.53441165E-06
Second partial derivative of Gibbs energy with respect to also
```

FE in 1: 1.82508696E+04 ΝT in 1: -3.07043434E+03 in 1: 0.0000000E+00 С in 2: -1.36027071E+05 in 2: -2.13007485E+05 in 2: -1.11741180E+05 VA First partial derivative with respect to NI in sublattice 1 of Gibbs energy: ..... -1.32427029E+05 of enthalpy: ..... 5.21563580E+04 of entropy: ..... 1.02546326E+02 of volume: Second partial derivative of Gibbs energy with respect to also ΝT in 1: 1.48390257E+05 in 1: 0.0000000E+00 С in 2: -7.70620431E+04 in 2: -1.61551726E+05 in 2: -1.12772206E+05 VA First partial derivative with respect to 0 in sublattice 1 of Gibbs energy: ..... -2.62929308E+05 of enthalpy: ..... of entropy: ..... 1.73991581E+02 of volume: 5.02555370E+04 .....0.0000000E+00 Second partial derivative of Gibbs energy with respect to also in 1: 2.99322360E+07 С in 2: 0.0000000E+00 in 2: 0.0000000E+00 Ν in 2: -1.82377137E+05 VΑ First partial derivative with respect to C in sublattice 2 of Gibbs energy: of enthalpy: ..... 1.21269096E+05 of entropy: ..... 1.55987507E+02 of volume: ..... 1.06885187E-05 Second partial derivative of Gibbs energy with respect to also С in 2: 2.99322360E+05 Ν in 2: -1.90144000E+04 in 2: -2.78465070E+04 VA First partial derivative with respect to N in sublattice 2 of Gibbs energy: ..... -2.39671400E+05 of enthalpy: ..... of entropy: ..... 1.55620608E+02 of volume: 4.04456947E+04 .....0.0000000E+00 Second partial derivative of Gibbs energy with respect to also Ν in 2: 2.99322360E+05 VA in 2: -3.30383665E+04

## **TABULATE\_REACTION**

Tabulate thermodynamic property changes for a chemical reaction as a table, or variation of one chosen property can be plotted against temperature as a graph, or saved as an experimental file (with an extension of .EXP). The thermodynamic data are automatically retrieved from the SSUB5 database (by default), or the current database set by the SWITCH\_DATABASE command (thus there is no need to use the DATA module before this command). You can select any other database by using <u>"SWITCH\_DATABASE" on page 310</u> in the TAB module.

It is important to remember the rules to specify a reaction:

- The reaction is specified, by giving the reactants and products, such as Ga+S=GaS, 5A1+2Fe=A15Fe2.
- Under Linux environments, the notation for a chemical formula treats upper and lower cases as the same (such as ga+s=GaS, 5Al+2Fe=al5fe2); but under Windows, the TAB module only accepts upper cases, e.g. GA+S=GAS, 5AL+2FE=AL5FE2.
- Elements with a single letter symbol must be followed by a stoichiometry factor, even if it is 1; thus, the symbol co is taken as cobalt, carbon monoxide must be given as c101 (or 01c1).
- All the reactants and products must be defined as species-stoichiometries in the currently-used database; in other words, the reaction definition only accepts alreadydefined species-stoichiometries as reactants and products. For example, if a database does not have a species defined with a stoichiometry of Al5Fe2 {e.g. the Al and Fe (but not Al5Fe2) may be defined as species, and the Al5Fe2 phase may be defined as a two-sublattice stoichiometric phase [Al]5[Fe]2, rather than a single sublattice stoichiometric phase [Al5Fe2]}, then the reaction 5Al+2Fe=Al5Fe2 cannot be tabulated.
- Make sure that the Gibbs free energy expression for a stoichiometric phase is really corresponding to the species-stoichiometry. For example, if a database has a definition of Function GAL5FE2 for 1/7 of the AL5FE2 formula, then the relation G (Al5Fe2)=7\*GAL5FE2 must be defined; otherwise, the tabulated thermodynamic properties for the reaction 5Al+2Fe=Al5Fe2 is wrong.

Syntax	TABULATE_REACTION
	Same reaction? /Y/:
Prompts	This prompt displays if there is at least one reaction already defined either with the commands <u>"ENTER_REACTION" on page 305</u> or TABULATE_REACTION. The following prompts only display if N is entered.
	Reaction: <chemical equation="" reaction=""> &amp; <ra+rb=pc+pd></ra+rb=pc+pd></chemical>
	A long reaction can be typed over several lines. A reaction should be terminated by a semicolon (;) or an empty line. A list showing the used database, defined elements, getting-data sequence, references, etc.
	Pressure /100000/: <pressure in="" interest,="" of="" pa=""></pressure>
	Low temperature limit /298.15/: <t-low, in="" k=""> Specify the starting temperature K.</t-low,>
	High temperature limit /298.15/: <t-high, in="" k=""> Specify the ending temperature K. Step in temperature /100/: <t-step> Specify the temperature step for the tabulation.</t-step></t-high,>
	Output file /Screen/: <return a="" file="" for="" name="" or="" screen,="" type=""></return>
	If you type Return for screen, a list of thermodynamic property changes for the defined/chosen reaction is shown as a table (which is demonstrated in the example outputs given below), and the command is terminated.
	If typing a file name, the table is both displayed on screen and saved as an *.EXP or a *.TAB file, and the program prompts:
	Graphical output /Y/:
	<ul> <li>For N a table is output to the screen, and the same table is saved as a basic text file with the default extension .TAB under the current working directory. In this case, the graph cannot be plotted.</li> </ul>
	• For Y a table is created with all thermodynamic functions as normal (which is displayed on screen) and generates a graph with temperature on the X-axis and a chosen property on a certain column in the table on the Y-axis (which is plotted on screen and saved as an *.EXP file), and it also asks which column is to be plotted on the resulting graph:
	Plot column? /2/: <1 or 2 or 3 or 4 or 5 or 6>
	Specify which property (as column number) to be plotted as the Y-axis (versus temperature as X-axis) as a graph on the screen. Simultaneously, all tabulated properties and the Y-axis setting (i.e. the plotted column) for the graph is written into an *.EXP file using the DATAPLOT format. The default column 2 is

Syntax	TABULATE_REACTION
	heat capacity, 3 enthalpy, 4 entropy and 5 Gibbs energy, and the additional column 6 is a user-entered function. The table displays on the screen. The plot then displays on the screen, followed by a POST: prompt. The POST module (postprocessor) automatically opens, and all types of the POST-module commands to refine the plotted diagram may be used. Possibilities include scaling the X/Y-axis, changing the X/Y-axis texts, etc. The command BACK or EXIT at the POST: prompt always takes you back to the TAB module.
	<b>DATPLOT User Guide</b> included with this documentation set.

#### **EXAMPLE OUTPUT**

For the reaction Ga+S=GaS (using the SSUB database), the following table is obtained by typing <Enter> in the prompt Output file /SCREEN/, and the figure by typing 2 in the prompt Plot column /2/:

```
OUTPUT FROM THERMO-CALC
  Reaction: S+GA=GA1S1
stable as S S
GA stable as GA S
GA1S1 stable as GA1S1 S
Delta-Cp
          Delta-H Delta-S Delta-G
    (Joule/K) (Joule) (Joule/K) (Joule)
(K)
298.15 -2.79489E+00 -2.09200E+05 -1.50580E+01 -2.04710E+05
300.00 -2.87516E+00 -2.09205E+05 -1.50755E+01 -2.04683E+05
302. ---- GA becomes GA L ,delta-H = 5589.80
367. ---- S becomes S_S2 ,delta-H =
                                  401.00
389.
     ---- S becomes S L ,delta-H = 1721.00
400.00 -1.22278E+01 -2.17521E+05 -4.07488E+01 -2.01222E+05
500.00 -1.57686E+01 -2.19323E+05 -4.47280E+01 -1.96959E+05
600.00 -1.02518E+01 -2.20583E+05 -4.70410E+01 -1.92359E+05
700.00 -6.58954E+00 -2.21430E+05 -4.83534E+01 -1.87583E+05
800.00 -4.46246E+00 -2.21963E+05 -4.90686E+01 -1.82709E+05
900.00 -2.82671E+00 -2.22328E+05 -4.94995E+01 -1.77778E+05
1000.00 -1.21787E+00 -2.22530E+05 -4.97138E+01 -1.72816E+05
```

1100.00 3.71702E-01 -2.22572E+05 -4.97552E+01 -1.67841E+05

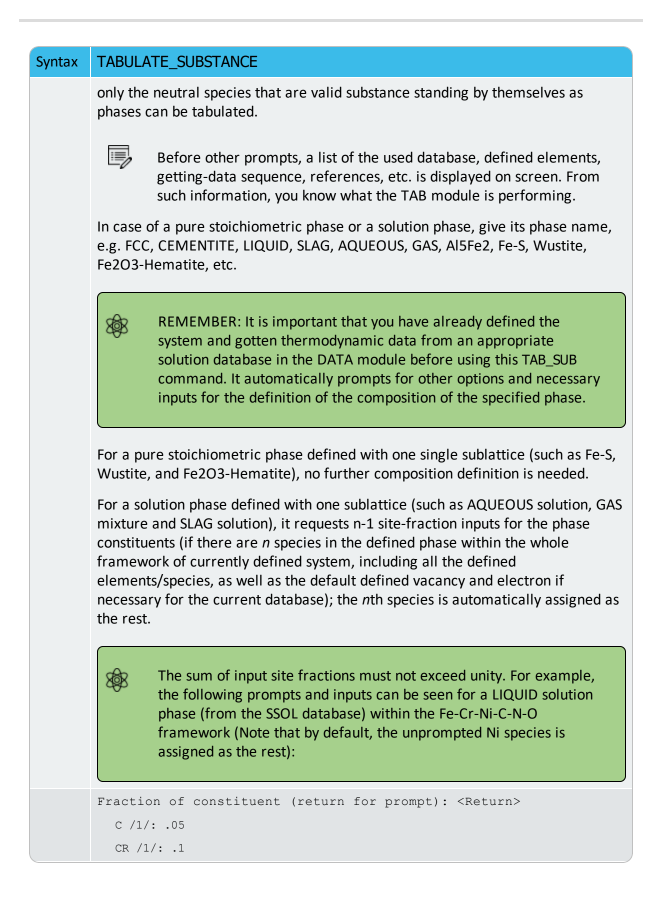
### TABULATE\_SUBSTANCE

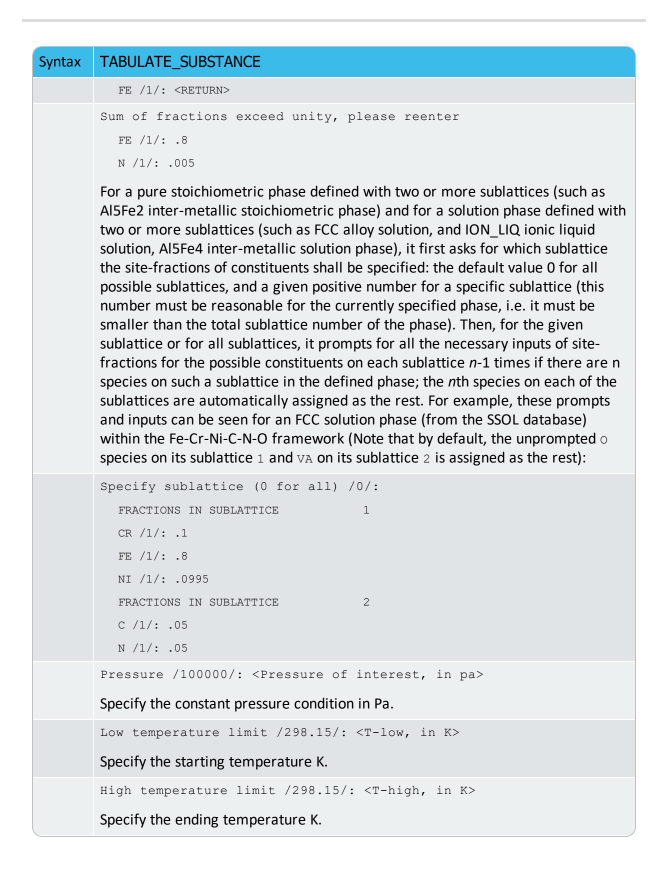
Tabulate thermodynamic properties of a substance (with a given chemical formula but it is not certain in which phase/state it may form) or a pure stoichiometric phase (getting data from a specific substance database such as SSUB5, or from a specific solution database such as SSOL5 and TCFE8), or of a solution phase with a fixed composition (getting data from a specific solution database such as SSOL5 and TCFE8), under a constant pressure and various temperatures.

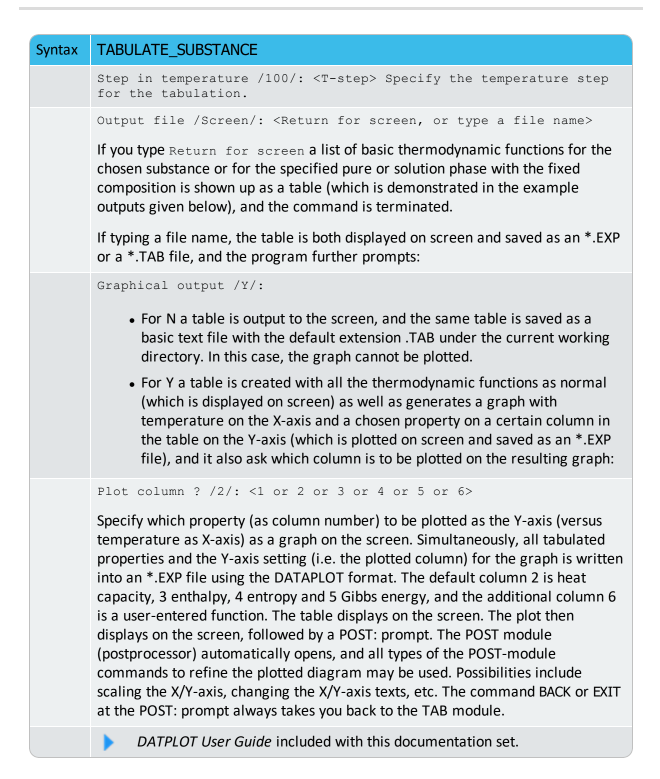
In case of a pure stoichiometric phase or a solution phase with a fixed composition, you must already define the system and get thermodynamic data from an appropriate solution database in the DATA module before using this TAB command. The phase name can be given either upper or lower or mixed cases, and can be abbreviated until it is unique, e.g. fcc, BCC, Liq, cem, A15Fe2, etc. as the SSOL database is used for the tabulations, or GAS, FE-S, wustite, Fe203-hem, etc. as the SSUB database is used. The module then optionally goes through a specific sublattice or all the possible sublattices for the specified phase, as well as through the whole framework of a currently defined system (including all the defined elements/species, as well as the default defined vacancy and electron if necessary for the current database), when it prompts for inputs of site fractions for various species possibly occupying such sublattices. Therefore, you can tabulate thermodynamic properties for a pure phase, or for a solution phase as a pure end-member (with a composition definition for the corresponding non-interacting species on all necessary sublattices in the end-member), or for a solution phase as a real solution (with a composition definition for related interacting species on all necessary sublattices in the solution).

Moreover, the variation for one of the tabulated properties for a substance, a pure stoichiometric phase, or a solution phase with a fixed composition can be plotted against temperature as a graph or saved as an experimental file (with an extension of .EXP).

Syntax	TABULATE_SUBSTANCE
Prompt	Substance (phase): <name of="" or="" phase="" species="" the=""> In case of a substance, give its chemical formula, e.g. Fe, H2, C1H6, FeC1, CaC1O3, MgSiO3, etc. When the TABULATE_SUBSTANCE <substance> command sequence is used for the first time in the TAB module, it always uses the SSUB Substances Database as the default database. If the SSUB database is not available for the current Thermo-Calc installation and a specific database has not been switched on yet, it may prompt to specify an appropriate substance or solution database (e.g. typing SSUB2, SSUB3, SSUB4, SSUB5; SSOL2, SSOL4, SSOL5; TCFE3, TCFE4, TCFE5, TCFE6, TCFE7, TCFE8 etc.). Prior to this command, you can also use the <u>"SWITCH_DATABASE" on page 310</u> command in the TAB module to set the current database. If a solution database is set as the current database,</substance></name>







#### EXAMPLE OUTPUT 1

For the FCC alloy phase as a non-ideal solution with a fixed composition

[Fe0.80,Cr0.10,Ni0.0995,O0.005][C0.05,N0.05,VA0.90]

The following table is obtained by typing Return for screen at the prompt Output file /SCREEN/:

ΟυΤΡ	UT FROM	THERMO-	CALC	
Phase :	FCC	P	ressure :	100000.00
Specie:	CR1/2			
******	* * * * * * * * * * * * * * *	****	* * * * * * * * * * * * * *	* * * * * * * * * * * * * * *
Т	Ср	Н	S	G
(K)	(Joule/K)	(Joule)	(Joule/K)	(Joule)
******	*****	*****	* * * * * * * * * * * * * *	* * * * * * * * * * * * * * *
298.15	2.70517E+01	6.23824E+03	4.40241E+01	-6.88755E+03
300.00	2.70889E+01	6.28832E+03	4.41916E+01	-6.96915E+03
400.00	2.87304E+01	9.08420E+03	5.22235E+01	-1.18052E+04
500.00	2.99904E+01	1.20222E+04	5.87742E+01	-1.73649E+04
600.00	3.10889E+01	1.50770E+04	6.43408E+01	-2.35275E+04
700.00	3.21116E+01	1.82375E+04	6.92106E+01	-3.02100E+04
800.00	3.30994E+01	2.14982E+04	7.35633E+01	-3.73524E+04
900.00	3.40742E+01	2.48569E+04	7.75182E+01	-4.49094E+04
1000.00	3.50483E+01	2.83130E+04	8.11586E+01	-5.28456E+04
1100.00	3.60268E+01	3.18667E+04	8.45449E+01	-6.11327E+04
1200.00	3.70143E+01	3.55187E+04	8.77219E+01	-6.97476E+04
1300.00	3.80149E+01	3.92700E+04	9.07241E+01	-7.86713E+04
1400.00	3.90311E+01	4.31222E+04	9.35784E+01	-8.78875E+04
1500.00	4.00649E+01	4.70768E+04	9.63064E+01	-9.73827E+04
1600.00	4.11174E+01	5.11358E+04	9.89256E+01	-1.07145E+05
1700.00	4.21896E+01	5.53010E+04	1.01450E+02	-1.17165E+05
1800.00	4.33555E+01	5.95774E+04	1.03894E+02	-1.27433E+05
1900.00	4.58528E+01	6.40379E+04	1.06306E+02	-1.37943E+05
2000.00	4.75402E+01	6.87138E+04	1.08704E+02	-1.48694E+05

#### EXAMPLE OUTPUT 2



For a phase, as either an end-member or real solution, the listed species name is irrelevant.

For the H2 species as a pure substance (using the SSUB5 database), the following table is obtained by typing Return for screen at the prompt Output file /SCREEN/, whilst the figure is generated by typing 5 at the prompt Plot column /2/:

	OUTPUTE	"ROMTHER	MO-CALO	C		
Phase	: GAS		Pressure :	100000.00		
Specie:	Н2					
		*****	*****	* * * * * * * * * * * * * * *	Т	Ср
Н	S	G				
(K)	(Joule/K)	(Joule)	(Joule/K)	(Joule)		
******* 2.88369E		L-06 1.30680E	+02 -3.896221		298.15	
300.00	2.88473E+01	5.33580E+01	1.30858E+02	-3.92042E+04		
400.00	2.91591E+01	2.95686E+03	1.39209E+02	-5.27268E+04		
500.00	2.92650E+01	5.87874E+03	1.45729E+02	-6.69856E+04		
600.00	2.93441E+01	8.80908E+03	1.51071E+02	-8.18336E+04		
700.00	2.94579E+01	1.17488E+04	1.55602E+02	-9.71730E+04		
800.00	2.96320E+01	1.47027E+04	1.59547E+02	-1.12935E+05		
900.00	2.98786E+01	1.76776E+04	1.63050E+02	-1.29068E+05		
1000.00	3.02043E+01	2.06810E+04	1.66214E+02	-1.45533E+05		
1100.00	3.05319E+01	2.37171E+04	1.69108E+02	-1.62302E+05		
1200.00	3.09281E+01	2.67897E+04	1.71781E+02	-1.79348E+05		
1300.00	3.13615E+01	2.99040E+04	1.74274E+02	-1.96652E+05		
1400.00	3.18115E+01	3.30625E+04	1.76614E+02	-2.14197E+05		
1500.00	3.22641E+01	3.62664E+04	1.78824E+02	-2.31970E+05		
1600.00	3.27094E+01	3.95151E+04	1.80921E+02	-2.49958E+05		
1700.00	3.31406E+01	4.28078E+04	1.82917E+02	-2.68151E+05		
1800.00	3.35526E+01	4.61426E+04	1.84823E+02	-2.86539E+05		
1900.00	3.39415E+01	4.95175E+04	1.86648E+02	-3.05113E+05		
2000.00	3.43045E+01	5.29300E+04	1.88398E+02	-3.23866E+05		

# **TERNARY\_DIAGRAM Commands**

In	thic	section:	
	LIIIS	section.	

TERNARY	DIAGRAM	Module			
		1110000	 	 	

# **TERNARY\_DIAGRAM Module**

The TERNARY\_DIAGRAM module (short name, the TERN module) enables you to quickly calculate a simple ternary phase diagram.

To enter the module, at the SYS prompt type GOTO\_MODULE TERNARY. There are no other commands for this module. Follow the prompts to plot a diagram.



Also see the Graphical Mode equivalent to this command, <u>"Ternary Calculator"</u> on page 152 as described in the *Thermo-Calc User Guide*.

Syntax	TERNARY_DIAGRAM
Prompts	Database: /FEDEMO/
	First element:
	Second element:
	Third element:
	Phase diagram, Monovariants, or Liquidus Surface: /Phase_Diagram/
	Temperture (C) /1000/
	Global minimization on: /Y/

## **Thermodynamic Variables and Units**

The following topics are about the different thermodynamic variables recognized by Thermo-Calc. The topics have information about how to define new derived variables and functions and how to express thermodynamic variables in non-default units. The POLY module only recognizes default units, which are the SI units.



See <u>example 44</u> in the *Thermo-Calc Console Mode Examples* to learn about using variables and functions.

#### In this section:

Thermodynamic Variables	328
Common Thermodynamic Variables	. 328
How to Read the Tables	. 329
Derived Variables and Partial Derivatives	. 342
Defining New Derived Variables and Functions	344
Derived Variables for Aqueous Solutions	.345
Defining New Derived Variables for Aqueous Solutions	349
Thermodynamic Variables and Wildcard Characters	349
Units of State Variables and Derived Variables	.350

# **Thermodynamic Variables**

There are different thermodynamic variables in Thermo-Calc. Some are *state variables*, for example temperature, pressure and mole fraction. These characterize the equilibrium state of a system. You set them when you define your system in the POLY module before performing a calculation. They are also used in other modules such as DATA, GIBBS and PARROT. State variables that involve components can be used for the defined components, but not for any species. (To define new components in a defined system, use <u>"DEFINE</u> <u>COMPONENTS" on page 186</u>).

Other useful thermodynamic variables are derive from state variables with a mathematical function. Some derived variables are pre-defined by Thermo-Calc, such as normalized energetic and compositional extensive state variables for example. You use these by appending various normalizing suffixes to abbreviations that are associated with the state variables.

Thermodynamic variables can also be divided into extensive variables and intensive variables. An extensive variable is a variable whose value depends on the size of the system, whereas an intensive variable is a variable whose value is independent of system size.

More information about thermodynamic variables is available in the online help in Thermo-Calc Console Mode. Use the command <u>"INFORMATION" on page 5</u>, with a subject keyword such as one of the following as argument: STATE VARIABLES, INTENSIVE VARIABLES, EXTENSIVE VARIABLES, DERIVED VARIABLES, CONDITIONS (for condition settings), AXIS-VARIABLES (for stepping/mapping variable settings).

#### **Common Thermodynamic Variables**

The tables below list common thermodynamic variables. Most of these can be used to define equilibrium conditions in the POLY module unless otherwise indicated.

The variables are divided into the following tables:

- Intensive variables
- Energy-related extensive variables for whole system or for a phase
- Compositional extensive variables (overall amount of components in the whole system, or amount of a component in system or in a phase)
- Constitutional composition-related extensive variables (amount of a constituent/species on a sublattice site in a phase)
- Special quantities

#### How to Read the Tables

- The *Abbrev*. column shows the abbreviation that you use in the POLY module for referring to the variable. If the variable must be given any arguments, then these are given in parenthesis directly after the abbreviation. For most variables, this is also the abbreviation that you use to refer to the variable in other modules. However, some variables are referred to with a different abbreviation in the POST module. If this is the case, then this is noted at the bottom of the table.
- Variables that are specific to a species, in which case the species is specified as an argument (*sp* in the table below). If there are two or more sublattices in a given phase, then this argument is given as constituent#sublattice. This indicates a constituent on a specific sublattice or site in the phase (referred to by the # sign and a digit). For example, y (BCC, C#2) stands for the site fraction of the C species on the second sublattice site of the BCC\_A2 solution phase.
- The *Unit* column show which unit that you can use to express the variable in the POLY module. This unit is always the SI unit. In other modules, state variables may be expressed in other units.
- The *Suffix* column shows the suffixes you can append to the variables. The R-suffix can be used with all compositional extensive state variables, but the suffix does not always change the value of the variable. This is indicated by putting the R in parenthesis.

These types of variables are listed in the following topics:

- "Intensive Variables" below
- "Extensive Variables" on the next page
- "Special Quantities" on page 335
- "The u-Fraction Variable" on page 335
- "Suffixes" on page 336

#### **Intensive Variables**

v	Abbrev.	Unit	Descript.	Domain	Suffix
Т	T <sup>1</sup>	К	Temperature	System	
Р	Ρ	Ра	Pressure	System	

v	Abbrev.	Unit	Descript.	Domain	Suffix
	MU(comp)			Component	R
μ	MU(sp,ph) <sup>2</sup>	J/mol	Chemical potential	Species relative to a solution phase	R
a	AC(comp)		Activity	Component	R
	AC(sp,ph) <sup>2</sup>			Species relative to a solution phase	R
	LNAC (comp) <sup>3</sup>	N/A	In(Activity)	Component	R
	LNAC (sp,ph) <sup>2</sup>			Species relative to a solution phase	R

 $^1$  When plotting in POST, besides T (Kelvin), you can also use T\_C or T\_F to plot temperature in °C or °F.

<sup>2</sup> Only for single-substitutional-lattice solution phases such as AQUEOUS solution and GASeous mixture phases and for interacting species on the substitutional sublattice of two-sublattice solution phases (such as the phases BCC\_A2 and M6C phases).

<sup>3</sup> In natural logarithm (InAC=MU/RT)

#### **Extensive Variables**

#### ENERGY-RELATED EXTENSIVE VARIABLES FOR WHOLE SYSTEM OR FOR A PHASE



See <u>example tcex44</u> in the *Thermo-Calc Console Mode Examples Guide* for an example of using variables and functions.

V	Abbrev.	Unit	Descript.	Domain	Suffix
	V			System	M, W, V, R
v	V(ph) <sup>1</sup>	m <sup>3</sup>	Volume	Phase	M, W, F, R
	VP(ph) <sup>2</sup>	-		Phase	M, W, V, R
	G			System	M, W, V, R
G	G(ph) <sup>1</sup>	J	Gibbs energy	Phase	M, W, V, F, R
	А	J	Helmholtz	System	M, W, V, R
A	A(ph) <sup>1</sup>		energy	Phase	M, W, V, F, R
	U		Internal energy	System	M, W, V, R
U	U(ph) <sup>1</sup>	J		Phase	M, W, V, F, R
Н	Н	J	Enthalpy <sup>6</sup>	System	M, W, V, R
	H(ph)	J		Phase	M, W, V, F
	S			System	M, W, V, R
S	S(ph) <sup>1</sup>	J/K	Entropy <sup>6</sup>	Phase	M, W, V, F, R
6.7	HM.T <sup>3</sup>	1/201/1/	Heat capacity at	System	R
Ср	HM(ph).T <sup>3</sup>	J/mol/K	constant pressure	Phase	R
Cv	HM.T <sup>4</sup>	J/mol/K	Heat capacity at constant volume	System	R

V	Abbrev.	Unit	Descript.	Domain	Suffix
	HM(ph).T <sup>4</sup>			Phase	R
D	DG(ph) <sup>5</sup>	N/A	Driving force (thermodynamic factor)	Phase	Always use with M, W, V or F. R can also be used.

<sup>1</sup> For this variable, if the phase is unstable, then the variable gets a value of zero.

<sup>2</sup> Do not use VP(ph) as a condition in POLY. Instead, use CHANGE\_STATUS PHASE <PHASE>=FIX <AMOUNT> where the fixed <AMOUNT> is roughly equal to VPF(ph), which cannot be directly evaluated or listed. If the phase is unstable, then VP(ph) and its normalized quantities get zero value.

- <sup>3</sup> Use only if pressure has been set as a condition.
- <sup>4</sup> Use only if volume has been set as a condition.
- <sup>5</sup> Already divided by RT.

<sup>6</sup> When working in the POST module you can also enter the abbreviation for this variable into the Console, for example, enth or enthalpy, which corresponds to HM, and entr and entropy, which corresponds to SM.

#### COMPOSITIONAL EXTENSIVE VARIABLES

These are for the overall amount of components in whole system, or amount of a component in system or in a phase.

Do not use NP(ph) or BP(ph) as conditions in the POLY module. Instead use CHANGE\_STATUS PHASE <PHASE>=FIX <AMOUNT> where the fixed <AMOUNT> is roughly equal to NPF(ph) or BPF(ph), which cannot be directly evaluated or listed. If the phase is unstable, then NP(ph) or BP(ph) (and its normalized quantities) get zero value.

When plotting the following quantities in POST, refer to each as indicated:

- X(comp) -> Mole-Fraction <comp> (Or M F)
- W(comp) -> Weight-Fraction <comp> (or W\_F)

- X%(comp) -> Mole-Percent <comp> (or M\_P)
- W%(comp) -> Weight-Percent <comp> (or W\_P).
- u-f(ph,comp) -> U-F

v	Abbrev.	Unit	Descript.	Domain	Suffix	Comment
	N			All components	M, W, V, (R)	
2	N(comp)	mole	Moloc	Component	M, W, V, (R)	
n	N (ph,comp)	mole	Moles	Component in a phase	M, W, V, (R)	
	NP(ph) <sup>1</sup>			Phase	M, W, V, (R)	See above
	В			All components	M, W, V, (R)	
b	B(comp)	aram		Component	M, W, V, (R)	
b	B (ph,comp)	gram	Mass	Component in a phase	M, W, V, (R)	
	BP(ph) <sup>1</sup>			Phase	M, W, V, (R)	See above
	X(comp)			Component	(R)	See above
x	X (ph,comp)	N/A	Mole fraction	Component in a phase	(R)	

V	Abbrev.	Unit	Descript.	Domain	Suffix	Comment
	W(comp)		Mass	Component	(R)	See above
w	W (ph,comp)	N/A	(weight) fraction	Component in a phase	(R)	
x%	X% (comp) <sup>2</sup>	N/A	Mole percent	Component	(R)	See above
w%	W% (comp) <sup>2</sup>	N/A	Mass (weight) percent	Component	(R)	See above
u	u-f (ph,comp) 1	N/A	u-fraction	Component in a stable phase	(R)	See above
in	IN(sp) <sup>2</sup>	mole	Input mole number	Phase species in the system	(R)	See above
im	IM(sp) <sup>2</sup>	gram	Input mass unit	Phase species in the system	(R)	See above

#### CONSTITUTIONAL COMPOSITION-RELATED EXTENSIVE VARIABLE

This is the amount of a constituent/species on a sublattice site in a phase.

v	Abbrev.	Unit	Descript.	Domain
у	Y (ph,cons#sub)	N/A	Site fraction	Constituent on a sublattice site (denoted by # and a digit) in a phase

v	Abbrev.	Unit	Descript.	Domain	Suffix
Q	QF(ph) <sup>1</sup>	N/A	Phase stability function	Phase	R Under a condition and system definition, QFR(ph) = QF(ph)
Тс	TC(ph) <sup>2</sup>	К	Curie temperature	Phase	R Under a condition and system definition, TCR(ph) = TC(ph)
Mb	BMAG (ph) <sup>3</sup>	N/A	Bohr magneton number	Phase	R Under a condition and system definition, BMAGR (ph) = BMAG(ph)

## Special Quantities

<sup>1</sup> Negative when phase composition is inside a spinodal, otherwise positive. Can be used to find out if an equilibrium is within the miscibility gap for a solution phase. Cannot be used as a condition.

<sup>2</sup> Calculated for phases with magnetic contributions in an equilibrium state. Cannot be used as a condition, but can be plotted in POST or evaluated in POLY using SHOW\_VALUE.

<sup>3</sup> Calculated for phases with magnetic contributions in an equilibrium state.

#### The u-Fraction Variable

The u-fraction is an important quantity in some single-point or stepping paraequilibrium calculations in the POLY module and in the DICTRA module. You can apply the variable in a paraequilibrium calculation to the substitutional matrix component, the substitutional alloying components or the interstitial component. The u-fraction is denoted as ui for the i-th system component *n* of a whole system or of a specific phase in various equilibrium states (such as full-equilibria, partial-equilibria, para-equilibria, local-equilibria).

Generally, the u-fraction is defined as:



The sum of  $x_j$  in the denominator is calculated for all substitutional alloying elements in the whole system or in a specific phase. The variable  $x_i$  in the numerator stands for the mole-fraction of the *i*-th component in the whole system or in the specific phase.

#### **Suffixes**

Suffixes may be appended to all extensive variables and to some intensive variables. Some suffixes can be used to enter the value of variables in normalized form. These are referred to as normalizing suffixes. If the variable that you create with a normalizing suffix is based on a state variable that can be set as a condition in POLY, then the normalized variable can also be set as a condition in POLY.

The normalizing suffixes are M (per mole), W (per mass in gram), V (per volume in  $m^3$ ) and F (per mole formula unit). There is also a reference state suffix R, which you can use if you want the value of a thermodynamic variable to be calculated with respect to a reference state that have previously set.

#### Normalizing Suffixes

When variables that express system and phase quantities are normalized, the following general rules are used:

- System quantities are normalized by the total system size (in terms of N, B or V).
- Phase quantities are normalized by the phase amount [in terms of NP(ph), BP(ph) or VP(ph)].

The normalized quantities of G(ph), A(ph), U(ph), H(ph), S(ph) and V(ph) are calculated according to the thermodynamic model used for the phase (e.g. GM(ph), AM(ph), UM(ph), HM(ph), SM(ph) and VM(ph)). These quantities are calculated using the first derivatives of the Gibbs energy expressed for the phase with respect to the current composition of the system.

The tables describe the normalizing suffixes for the different state variables.

#### G, A, U, H, S AND V OF A WHOLE SYSTEM

Suffix	Description	Example
M (per mole)	Total system size in terms of N.	GM is the Gibbs energy per mole of the system (J/mol).

Suffix	Description	Example
		GM = G/N
W (per mass in gram)	The total system size in terms of B.	GW is the Gibbs energy per mass of the system (J/g). $\mathbf{GW} = \mathbf{G}/\mathbf{B}$
V (per volume in m <sup>3</sup> )	The total system size in terms of V. VV does not have to be evaluated.	GV is the Gibbs energy per volume of the system (J/m3). ${f GV}={f G}/{f V}$

## G(PH), A(PH), U(PH), H(PH), S(PH) AND V(PH) OF A PHASE

Suffix	Example
M (per mole)	For example, GM(ph) is Gibbs energy of the phase per mole of the phase (J/mol). GM(ph) = G(ph)/NP(ph)
W (per mass in gram)	For example, GW(ph) is Gibbs energy of the phase per mass of the phase (J/mol). GW(ph) = G(ph)/BP(ph)
V (per volume in m <sup>3</sup> )	For example, GV(ph) is Gibbs energy of the phase per volume of the phase (J/mol). $GV(ph) = G(ph)/V(ph)$
F (per mole formula unit)	For example, GF(ph) is the Gibbs energy of the phase per formula unit of the phase (J/mol). $GF(ph) = G(ph)/NP(ph)^*NA$

## DG(PH) OF A PHASE

Suffix	Description	Example(s)
M (per mole)	Theoretically, the first derivative of the variable with regard to the phase amount in terms of NP (ph). Since DG(ph) is not directly calculated, the second derivative of the Gibbs energy expressed for the phase in question with respect to the current compositions in the equilibrium state of the system is calculated instead. DGM(ph) cannot not be set as a condition since it is only calculated under a certain type of equilibrium state.	DGM(ph) is driving force for precipitation of the phase per mole of components. DGM(ph) = $\partial$ DG(ph)/ $\partial$ NP(ph)
W (per mass in gram)	Theoretically, the first derivative of the variable with regard to the phase amount in terms of BP (ph). Since DG(ph) is not directly calculated, the second derivative of the Gibbs energy expressed for the phase in question with respect to the current compositions in the equilibrium state of the system is calculated instead. DGW(ph) cannot not be set as a condition since it is only calculated under a certain type of equilibrium state.	DGW(ph) is driving force for precipitation of the phase per mass of components. DGW(ph) = $\partial$ DG(ph)/ $\partial$ BP(ph)
V (per volume in m <sup>3</sup> )	Theoretically, the first derivative of the variable with regard to the phase amount in terms of VP (ph). Since DG(ph) is not directly calculated, the second derivative of the Gibbs energy expressed for the phase in question with respect to the current compositions in the equilibrium state of the system is calculated instead. DGV(ph) cannot not be set as a condition since it is only calculated under a certain type of equilibrium state.	DGV(ph) is driving force for precipitation of the phase per volume of components. DGV(ph) = $\partial DG(ph) / \partial VP(ph)$

Suffix	Description	Example(s)
F (per mole formula unit)	Theoretically, the first derivative of the variable with regard to the phase amount in terms of NP(ph) and NA (NA is the total atomic number in the phase formula). Since DG(ph) is not directly calculated, the second derivative of the Gibbs energy expressed for the phase in question with respect to the current compositions in the equilibrium state of the system is calculated instead. DGF(ph) cannot not be set as a condition since it is only calculated under a certain type of equilibrium state.	DGF(ph) is driving force for precipitation of the phase per formula unit of components. DGF(ph) = $\partial DG(ph)/\partial NP(ph)^* \partial NA$

#### N AND B OF A SYSTEM

Suffix	Description	Example(s)
M (per mole)	The total system size in terms of N. NM does not have to be evaluated. BM cannot be set as a condition.	BM is mass (gram) of components per mole of the system (g/mol). BM = $B/N$
W (per mass in gram)	The total system size in terms of B. BW does not have to be evaluated. NW cannot be set as a condition.	NW is mole number of components per mass of the system (mol/g). $\mathbf{NW} = \mathbf{N}/\mathbf{B}$
V (per volume in m <sup>3</sup> )	The total system size in terms of V. BV is the density of the entire system.	NV is mole number of components per volume of the system (mol/m <sup>3</sup> ). NV = N/VBV is the density of the entire system (g/m <sup>3</sup> ). BV = B/V

## N (COMP) AND B(COMP) OF A COMPONENT IN THE SYSTEM

Suffix	Description	Example(s)
M (per mole)	The total system size in terms of N. BM(comp) cannot be set as a condition.	NM(comp) is the mole of a component per mole of the system (i.e. mole fraction, which also is expressed as X (comp)). NM(comp) = N(comp)/N
W (per mass in gram)	The total system size in terms of B. NW(comp) cannot be set as a condition.	BW(comp) is mass (gram) of a component per mass of the system (i.e. mass fraction, which also is expressed as W (comp)). BW(comp) = B(comp)/B
V (per volume in m <sup>3</sup> )	The total system size in terms of V.	NV(comp) is mole number of a component per volume of the system (mol/m3). NV(comp) = N(comp)/V

## NP(PH), BP(PH) AND VP(PH) OF A PHASE IN THE SYSTEM

Suffix	Description	Example(s)
M (per mole)	BPM(ph) and VPM(ph): The phase amount in terms of NP (ph). NPM(ph): The total system size in terms of N.	$\begin{array}{l} \mbox{BPM(ph) is mass (gram) of a phase per moles of the system (g/mol).} \\ \mbox{BPM(ph)} = \mbox{BP(ph)}/\mbox{N} \\ \mbox{NPM(ph) is number of moles of a phase per moles of the system (mole fraction).} \\ \mbox{NPM(ph)} = \mbox{NPM(ph)} \\ \mbox{NPM(ph)} = \mbox{NP(ph)}/\mbox{N} \end{array}$
W (per mass in gram)	NPW(ph) and VPW(ph): The phase amount in terms of BP (ph). BPW(ph):The total system size in terms of B.	VPW(ph) is volume (m3) of a phase per mass of the system (m3/g) VPW(ph) = VP(ph)/B BPW(ph) is mass (gram) of a phase per mass of the system (mass fraction) BPW(ph) = BP(ph)/B

Suffix	Description	Example(s)
V (per volume in m <sup>3</sup> )	NPV(ph) and BPV(ph): The phase amount in terms of VP(ph). VPV(ph): The total system size in terms of V.	NPV(ph) is number of moles of a phase per volume of the system (mol/m3). NPV(ph) = NP(ph)/V VPV(ph) is volume (m3) of a phase per volume of the system (volume fraction). VPV(ph) = VP(ph)/V

#### N(PH,COMP) AND B(PH,COMP) OF A COMPONENT IN A PHASE

Suffix	Description	Example(s)
M (per mole)	The phase amount in terms of NP(ph) of the phase.	NM(ph,comp)is number of moles of a component per mole of a phase (i.e. mole fraction of a component in a phase, which is equivalent to X(ph, comp)). NM(ph,comp) = N(ph,comp)/NP(ph)
W (per mass in gram)	The phase amount in terms of BP(ph) of the phase.	BW(ph,comp) is mass (gram) of a component per mass of a phase (i.e. mass fraction of a component in a phase, which is equivalent to W(ph,comp)). BW(ph,comp) = B(ph,comp)/BP(ph)
V (per volume in m <sup>3</sup> )	The phase amount in terms of VP(ph) of the phase.	NV(ph,comp) is mole number of a component per volume of a phase (mol/m3). NV(ph,comp) = N(ph,comp)/VP(ph)

#### THE REFERENCE STATE SUFFIX R

You can use the reference state suffix R for some thermodynamic variables to calculate their value with respect to a reference state that you have previously set for a system component with the <u>"SET\_REFERENCE\_STATE" on page 258</u> command in POLY (or in a response-driven modules such as the POURBAIX module). The value of energy-related variables that are used with the R suffix depends on the reference states of all the components in the defined system.

It is possible to use an R suffix on all compositional extensive state variables as well, but the value of the state variable is always the same, with or without the suffix.

If the reference state for a system component is the default reference state (the stable reference state (SER) which is defined in a Thermo-Calc database), then MUR(comp)= MU (comp), ACR(comp)= AC(comp) and LNACR(comp)= LNAC(comp).

In the case of some thermodynamic variables, you can also use the R suffix to express chemical potentials and activities of species relative to some single-substitutional-lattice solution phases (such as aqueous solution, gaseous mixture, metallic liquid solution, slag mixture or MO solid solution). These state variables are MU(sp,ph), MUR(sp,ph), AC(sp,ph), ACR(sp,ph), LNAC(sp,ph) and LNACR(sp,ph).

The reference states and standard states of various solution species are pre-defined for some solution phases in some databases. For all solution species in any solution model in any database, it is always the case that MUR(sp,ph)= MU(sp,ph), ACR(sp,ph)= AC(sp,ph) and LNACR(sp,ph)= LNAC(sp,ph).

# **Derived Variables and Partial Derivatives**

Many derived variables can be obtained easily by using partial derivatives of state variables, such as heat capacity, thermal expansivity, isothermal compressibility, among others.

Derivatives of state variables can be defined or evaluated by putting a dot (.) between two state variables. The variable after the dot must be a condition that you have set when you defined your system. For example, the heat capacity (at constant pressure or volume) of a system is the partial derivative of the state variable enthalpy with respect to temperature. To evaluate this derivative, use the designation HM.T, where temperature is a state variable that you have set already.

Any partial derivative of a state variable with respect to another can be defined or evaluated as a derived variable by using the dot (.) notation.

#### **DERIVED VARIABLES DOT NOTATION EXAMPLES**

The following table shows some examples:

Derived variable	Description
$\mathrm{HM.T}=\partial\mathrm{HM}/\partial\mathrm{T}$	Heat capacity for the system at either constant P if pressure is a condition or constant V if volume is a condition.

Derived variable	Description
${ m HM}({ m ph}).{ m T}=\partial { m HM}({ m ph})/\partial { m T}$	Heat capacity for a phase at either constant P if pressure is a condition or constant V if volume is a condition.
$\mathrm{H.T}=\partial\mathrm{H}/\partial\mathrm{T}$	Heat capacity for the system multiplied by total mole number of components, i.e., $\partial H/\partial T = C_p * N_{if}$ pressure is a condition in a closed system or $\partial H/\partial T = C_v * N_{if}$ volume is a condition in a closed system.
$ m H(ph).T=\partial H(ph)/\partial T$	Heat capacity for a phase multiplied by the sum of $NP(ph)_{and} HM(ph)^* \partial NP(ph) / \partial T_{, i.e.,}$ $C_p(ph)^*NP(ph) + HM(ph)^* \partial NP(ph) / \partial T_{if \ pressure}$ is a condition or $C_{v(ph)}^*NP(ph) + HM(ph)^* \partial NP(ph) / \partial T_{if \ volume}$ is a condition.
$VM.T = \partial VM / \partial T$	Thermal expansivity of the system (already multiplied by the total molar volume), i.e., $\partial VM / \partial T = \alpha^* VM$ .
${ m VM(ph).T}=\partial { m VM(ph)}/\partial { m T}$	Thermal expansivity of a phase (already multiplied by its molar volume), i.e., $\partial VM(ph) / \partial T = \alpha (ph)^* VM(ph)$ .
${ m V.T}=\partial { m V}/\partial { m T}$	Thermal expansivity of the system (already multiplied by the total volume), i.e., $\partial V / \partial T = \alpha^* V$ .
$ m V(ph).T=\partial V(ph)/\partial T$	Thermal expansivity of a phase (already multiplied by the phase volume) plus the $VM(ph)^* \partial NP(ph) / \partial T$ term, i.e., $\partial V(ph) / \partial T = \alpha(ph)^* V(ph) = \alpha(ph)^* VM(ph)^* NP(ph) + VM(ph)^* \partial NP(ph) / \partial T$ .
$\mathrm{VM.P}=-\partial\mathrm{VM}/\partial\mathrm{P}$	Isothermal compressibility of the system (already multiplied by the total molar volume), i.e., $\partial VM / \partial P - \kappa^* VM$ .
${ m VM}({ m ph}).{ m P}=-\partial { m VM}({ m ph})/\partial { m P}$	Isothermal compressibility of a phase (already multiplied by its molar volume), i.e., $\partial VM(ph)/\partial P = -\kappa (ph)^* VM(ph)$ .

Derived variable	Description
$V.P = -\partial V / \partial P$	Isothermal compressibility of the system (already multiplied by the total volume), i.e., $\partial { m V}/\partial { m P}=-\kappa^*{ m V}$
$ m V(ph).P = -\partial V(ph)/\partial P$	Isothermal compressibility of a phase (already multiplied by the phase volume) plus the $VM(ph)^* \partial NP(ph) / \partial P_{term, i.e.,}$ $\partial V(ph) / \partial P = -\kappa (ph)^* V(ph) = -\kappa (ph)^* VM(ph)^* NP(ph) + VM(ph)^* \partial NP(ph) / \partial P.$
${ m T.X(comp)}=\partial { m T}/\partial { m X(comp)}$	Slope of a phase boundary on a T-X(comp) phase diagram with respect to mole fraction of the component in the system.
${ m T.W(comp)}=\partial { m T}/\partial { m W(comp)}$	Slope of a phase boundary on a T-W(comp) phase diagram with respect to mass of the component in the system.
$T.X(ph,comp) = \partial T/\partial X(ph,comp)$	Slope of a phase boundary on a T-X(ph,comp) phase diagram with respect to mole fraction of the component in the phase.
${ m T.W(ph,comp)}=\partial { m T}/\partial { m W(ph,comp)}$	Slope of a phase boundary on a T-W(ph,comp) phase diagram with respect to mole fraction of the component in the phase.
$\mathrm{P.T}=\partial\mathrm{P}/\partial\mathrm{T}$	Slope of a phase boundary on a P-T phase diagram (Note that the equilibrium with phase assemblage must be calculated first).

#### **Defining New Derived Variables and Functions**

You can define additional derived variables or functions based on a state variable by using <u>"ENTER\_SYMBOL" on page 238</u>. Any derived variable or function must have a unique name that starts with a letter and has a maximum number of eight characters. These characters can include both uppercase and lowercase letters, numerical digits and the underscore (\_). No other characters are allowed.

The difference between new derived variables introduced with the ENTER\_SYMBOL command and new functions introduced with the same command is as follows. Whenever Thermo-Calc calculates the value of a function (as part of an equilibrium calculation for example), all functions related to the defined system is evaluated. An entered derived variable, on the other hand, is only evaluated when it is defined and when it is re-evaluated with <u>"EVALUATE\_FUNCTIONS" on page 197</u>.

### **Derived Variables for Aqueous Solutions**

Standard state variables used for other phases can be directly applied to the aqueous solution phase. In addition, there are pre-defined derived variables for the aqueous solution phase in particular.

Examples of predefined derived variables for the aqueous solution phase are listed and briefly described in the following tables. Some derived variables for a gaseous mixture phase are also shown. This is because EOS (Equation of State) expressions, standard thermodynamic properties and transport properties of the pure solvent H2O are also expressions and properties of the pure gaseous species H2O.

Name	Abbrev.	Units	Description	Comments
рН	РН	N/A	Acidity	of the aqueous solution phase pH = -log10(AC(H+)) = -log10(ACR(H+,AQ))
Eh	EH	V, mV	Hypothetical electric potential	of the aqueous solution phase Eh = u(ZE)/96485.309
Pe	PE	N/A	Logarithm of the hypothetical electron activity	of the aqueous solution phase pe = u(ZE)/ (2.3025851*RT)
Ah	АН	kJ, kcal	Thermodynamic affinity per electron	of a redox couple wrt the standard hydrogen electrode in the aqueous solution phase Ah = u(ZE)
yw	YH2O	N/A	Mole fraction	of the solvent H2O in the aqueous solution phase YH2O = Y(AQ,H2O)

#### FOR AQUEOUS SOLUTION PHASE

Name	Abbrev.	Units	Description	Comments
Nw	AH2O	mole	Mole number	of 1.0 kg of solvent H2O AH2O = 55.508435
Non	NSH2O	mala	Mole number, NS(AQ,H2O)	of the solvent (H2O) in the aqueous solution phase NS(AQ,H2O) = YH2O*NP(AQ)
Nsp	NS#	mole	Mole number, NS(AQ,sp)	of a solute species in the aqueous solution phase NS(AQ,sp) = Y (AQ,sp)*NP(AQ)
Μ	ML#	mol/kg_ H2O	Molality, ML(AQ,sp)	of a solute species in the aqueous phase ML(AQ,sp) = Y (AQ,sp)*AH2O/YH2O
m*	TIM	equivalent molality	Total molality	of all solute species in the aqueous solution phase TIM = sum[ML (AQ,sp)]ions + sum [ML (AQ,sp)]complexes
mt	TIC#	equivalent molality	Total ionic concentration, TIC (AQ,sp)	of a cation I in the aqueous phase TIC(AQ,spI) = sum[ML (AQ,spJ)*V(spI-in-spJ)]
I	ISTR	N/A	Ionic strength	of the aqueous solution phase ISTR=1/2*sum[ML (AQ,sp)*Z(AQ,sp)**2]
γ	RCH2O	N/A	Activity coefficient,	of the solvent (H2O)

Name	Abbrev.	Units	Description	Comments
			RC(H2O,AQ)	RC(H2O,AQ) = ACR (H2O,AQ)/YH2O
	RC#		Activity coefficient, RC(sp,AQ)	of a solute species RC(sp,AQ) = ACR (sp,AQ)/Y (AQ,sp)*YH2O
	AIH2O		Activity, AI(H2O,AQ)	of the solvent (H2O), AI(H2O,AQ) = ACR (H2O,AQ)
αί	AI#	N/A	Activity, Al(sp,AQ)	of a solute species related to the aqueous solution phase, AI(sp,AQ) = ACR (sp,AQ)*AH2O
	LogAI#		Log10 Activity, LogAI(H2O,AQ), LogAI(sp,AQ)	of the solvent or a solute species related to the aqueous solution phase in common logarithm, LogAl(H2O,AQ) = log10[Al(H2O,AQ)] LogAl(sp,AQ) = log10[Al(sp,AQ)]
αw	AW	N/A	Activity	of H2O in the aqueous solution phase AW = ACR(H2O,AQ)
ф	OS	N/A	Osmotic coefficient	of aqueous solution phase OS = - 55.508435*InAW/TIM

Name	Abbrev.	Units	Description	Comments
				of the aqueous solution phase
At1	AT1	equil_ mol/kg_ H2O	Titration alkalinity (definition 1)	Generally defined as the equivalent molality of carbonate and bicarbonate at the methyl orange endpoint (pH=4.5).
				of the aqueous solution phase
At2	AT2	equil_ mol/kg_ H2O	Titration alkalinity (definition 2)	Generally defined as the equivalent molality of carbonate and bicarbonate, plus sulfide, at the methyl orange endpoint (pH=4.5).

### FOR GASEOUS MIXTURE PHASE

Name	Abbrev.	Units	Description	Comments
γ	RA#	N/A	Activity coefficient,	of a gaseous species in the gaseous mixture
			RA(sp,GAS)	RA(sp,GAS) = function(Y,T,P)
γ*	RF#	N/A	Fugacity coefficient, RF(sp,GAS)	of a pure gaseous species under TP RF(sp,GAS) = function(T,P/V)
F	FUG#	pa, bar, psi	Fugacity, FUG(sp,GAS)	of a gaseous species in the gaseous mixture FUG(sp,GAS) = RA(sp,GAS)*RF(sp,GAS)*Y (GAS,sp)*P
ft	TFUG	pa, bar, psi	Total gas fugacity	of the gaseous mixture phase TFUG = sum[FUG(sp,GAS)]

#### **Defining New Derived Variables for Aqueous Solutions**

You can define many different additional derived variables and functions. For instance, the partition coefficient of a component between two phases is defined as follows:

```
Pc<name> = X(phase1, component) /X(phase2, component)
```

Another example is the activity coefficient of a component in the system. It is defined as follows:

```
Rc<name> = ACR(component)/X(component)
```

The activity coefficient of a species in a solution phase depends on which model definitions on the reference states that are used and it depends on the standard states for the species. If the standard state of a species is defined to be the same as the pure species in a substitutional phase, then the activity coefficient of the species can be calculated as:

```
RC<name> = ACR(species, phase) /Y(phase, species)
```

In some cases, there are more than one sublattice in the phase and the reference states for the phase species cannot be presented by the pure species, as in the case of C in FCC [(Fe)1 (C,N,VA)1], where the model may end at fifty percent C and fifty percent Fe. In those cases, the chemical potential and activity of all phase constituents is not well defined and you cannot properly define the activity coefficients of the species in the phase.

For an aqueous solution phase, no matter what model is used, the reference state and standard state for both the solvent and solutes are defined in a special way in the Thermo-Calc software. The reference state for the solvent is set as pure water, the same as its standard state (according to Raoults's Law). The standard state for a solute species is set the hypothetical solute species, whilst its reference state is defined as the hypothetical state at unit molality (one mole dissolved into 1.0 kg of solvent) but in which the environment of each molecular is the same as at infinite dilution (according to Henry's Law). Under these definitions, the activity coefficients for solvent species can be calculated as follows:

RC<H2O> = ACR(H2O, aqueous) / Y(aqueous, H2O)

The coefficients for the solute species can be calculated as follows:

```
RC<spec> = AI(species,aqueous)/ML(aqueous,species)
= ACR(species,aqueous)/ Y(aqueous,species)*Y(aqueous,H20)
```

Al is here the activity of solute species calculated from the model, and ML is the molality of the species.

#### **Thermodynamic Variables and Wildcard Characters**

The ACR (species, phase) and the MUR (species, phase) quantities can only be appropriately and meaningfully applied to substitutional solution phases (such as GAS, AQUEOUS, LIQUID)

and to solution phases without an interstitial sublattice set. This means that the wild card characters \* and \$ normally don't work properly for the activity and/or chemical potential properties.

For example, do not use ACR(\*,\*) or MUR(\*,\*) for all the species relative to all phases, nor ACR(\*,phase) or MUR(\*,phase) for all the species relative to a specific phase, nor ACR (species,\*) or MUR(species,\*) for a specific species relative to all phases.

Instead, use specific names for species and (applicable) phase, as in the following examples:

```
SHOW_VALUE acr(fe+2, aqueous) Lists activity of the Fe+2 species in AQUEOUS solution phase
```

SHOW\_VALUE mur(c2,gas)Lists chemical potential of the C2 species in GAS mixture phase

<code>SET\_AXIS\_VAR</code> y ac(o2,gas)Sets activity of the O2 species in GAS mixture phase as Y-axis

# **Units of State Variables and Derived Variables**

The units for the state variables in the Units column in the table in <u>"User-Specified Units"</u> <u>below</u> are the SI units that are used in the POLY module. However, in the POST module, you can both use pre-defined symbols and define your own symbols which are based on state variables and derived variables. By doing this, you can express the value of the thermodynamic variables in non-default units. The following topic briefly describes how you can create your own symbols and suggests expressions that you can use to convert default units into other standard units which are not default units in Thermo-Calc.

The topic <u>"Examples of User-Defined Units" on the next page</u> describes some variables of specific species in a phase and the units in which they are expressed. These variables are not used in POLY or POST, but in the GIBBS, DATA, PARROT and ED\_EXP modules.

#### **User-Specified Units**

You do not always need to use the default units for variables. Using <u>"ENTER\_SYMBOL" on</u> <u>page 238</u>, you can convert the value of variable from the default unit to your preferred unit. (Some response-driven modules also allow you to set conditions in non-default units.) Userdefined symbols in the form of functions can be used for plotting in the POST module, but they cannot be used as conditions in POLY. User defined symbols in the form of variables can be used as conditions.

For example, if you want pressure to be plotted in bar rather than Pa, then you define the following function PB:

Enter symbol function PB = P/1E5;

If you wanted heat capacity of a system to be shown as in terms of J/mol/K, then you can define the following function Cp:

```
Enter_symbol function Cp = Hm.T ;
```

If the variable you want to convert is a function of one or more derivatives, then you must enter the symbol as a variable rather as a function. For instance, a symbol expressing heat capacity of a system in a unit of cal/mol/K can be defined as the following variable  $c_{p2}$ :

```
Enter_symbol variable Cp2 = Hm.T/4.1858 ;
```



Also see <u>"Examples of User-Defined Units" below</u> for information about entering conventional functions and the associated units.

#### **Examples of User-Defined Units**

The following tables give examples of expressions you can use to enter symbols. This enables you to show the values of thermodynamic variables in a variety of units.



<u>"Units of State Variables and Derived Variables" on the previous page</u> and <u>"User-Specified Units" on the previous page</u>



You can also use thermodynamic variables with normalizing suffixes in the expression when you enter a symbol. However, the tables do not have examples of these expressions.

In the expressions these are defined as follows:

- ${\tt i}$  is for an auxiliary index or letter that corresponds to the component  ${\tt c}$
- $\tt j$  is for an auxiliary index or letter that corresponds to the phase  $\tt ph$  , and
- ij stands for an auxiliary index or letter that corresponds to the component  ${\tt c}$  in the phase  ${\tt ph}.$

#### INTENSIVE AND EXTENSIVE VARIABLES OF A DEFINED SYSTEM

Quantity	Suggested name and expression	Unit	
For intensive variables of a	iables of a defined system		
Temperature	TempC = T-273.15	Celsius (C)	
	TempF = 1.8*T-459.67	Fabraphait (F)	
	TempF = 1.8*TempC+32	Fahrenheit (F)	

Quantity	Suggested name and expression	Unit
	PB = P/1E5	bars (bar)
	PKb = P/1E8	kilobars (kbar)
Processo	PAtm = P/101325	atmosphere (atm)
Pressure	PSI = P/6894.76	pounds/sq. inch (psi)
	PIHg = P/3342.11	inches of Hg
	PTor = P/133.322	Tors (millimeters of Hg)
For extensive variables of a defined system		
	BKg = 1E-3*B	kilograms (kg)
Amount	BTon = 1E-6*B	tons
	VDM = 1E-3*V	cubic decimeters (dm <sup>3</sup> , I)
Volume	VCM = 1E-6*V	cubic centimeters (cm <sup>3</sup> )
	VMM = 1E-9*V	cubic millimeters (mm <sup>3</sup> )
	VCI = 1.6387064E-5*V	cubic inches (in <sup>3</sup> )
	VLi = 1E-3*V	liters (I)

Quantity	Suggested name and expression	Unit
	GCal = G/4.1858	cal
	ACal = A/4.1858	cal
Energy	UCal = U/4.1858	cal
	HCal = H/4.1858	cal
	SCal = S/4.1858	Cal/K
	Cp2 = HM.T/4.1858	cal/mol/K
Heat Capacity	Cp3 = HW.T/4.1858	cal/g/K
	Cp4 = HV.T/4.1858	cal/m <sup>3</sup> /K
The survey of Francessien	A2 = VM.T*1E-6*VM	cm <sup>3</sup> /mol/K
Thermal Expansion	A3 = VW.T*1E-6*VM	cm <sup>3</sup> /g/K
Isothermal Compressibility	B2 = -VM.P*1E-9*VM	mm <sup>3</sup> /mol/Pa
	B3 = -VW.P*1E-9*VM	mm <sup>3</sup> /g/Pa

#### INTENSIVE AND EXTENSIVE VARIABLES OF A SYSTEM COMPONENT

Quantity	Suggested name and expression	Unit
For intensive variables of a system component		
	MUi = MU(c)/4.1858	cal/mol
Chemical Potential	MURi = MUR(c)/4.1858	cal/mol

Quantity	Suggested name and expression	Unit	
Special quantity for aqueous solution	pH = -LOG10(ACR(H+1))	dimensionless	
	Eh = MUR(ZE)/RNF where RNF=96485.309	volts (V)	
For extensive variables of a system component			
Amount	MFi = N(c)/N = X(c)	dimensionless	
	MPi = N(c)/N*100	dimensionless	
	WFi = B(c)/B = W(c)	dimensionless	
	WPi = B(c)/B*100	dimensionless	
	BKgi = 1E-3*B(c)	kilograms (kg)	
	BToni = 1E-6*B(c)	tons	

**EXTENSIVE VARIABLES OF A PHASE** 

Quantity	Suggested name and expression	Unit
Amount	MFj = NP(ph)/N	dimensionless
	MPj = NP(ph)/N*100	dimensionless
	WFj = BP(ph)/B	dimensionless
	WPj = BP(ph)/B*100	dimensionless
	BKgj = 1E-3*BP(ph)	kilograms (kg)
	BTonj = 1E-6*BP(ph)	tons
Volume	VCMj = 1E-3*V(ph)	cubic decimeters (dm <sup>3</sup> , l)
	VCMj = 1E-6*V(ph)	cubic centimeters (cm <sup>3</sup> )
	VMMj = 1E-9*V(ph)	cubic millimeters (mm <sup>3</sup> )
Energy	GCalj = G(ph)/4.1858	cal
	ACalj = A(ph)/4.1858	cal
	UCalj = U(ph)/4.1858	cal
	HCalj = H(ph)/4.1858	cal
	SCalj = S(ph)/4.1858	Cal/K

Quantity	Suggested name and expression	Unit
Driving Force	DG2j = DGM(ph)/4.1858	cal/mol
	DG3j = DGW(ph)/4.1858	cal/g
	DG4j = DGV(ph)/4.1858	Cal/m <sup>3</sup>
	DG5j = DGF(ph)/4.1858	cal/mole_formula_unit
Heat Capacity	Cp2j = HM(ph).T/4.1858	cal/mol/K
	Cp3j = HW(ph).T/4.1858	cal/g/K
	Cp4j = HV(ph)/.T/4.1858	cal/m <sup>3</sup> /K
	Cp5j = HF(ph)/.T/4.1858	cal/mole_formula_unit/K
Thermal Expansion	A2j = VM(ph).T*1E-6*VM	cm <sup>3</sup> /mol/K
	A3j = VW(ph).T*1E-6*VM	cm <sup>3</sup> /g/K
	A4j = VF(ph).T*1E-6*VM	cm <sup>3</sup> /mole_formula_ unit/K
Isothermal Compressibility	B2j = -VM(ph).P*1E-9*VM	mm <sup>3</sup> /mol/Pa
	B3j = -VW(ph).P*1E-9*VM	mm <sup>3</sup> /g/Pa
	B4j = -VF(ph).P*1E-9*VM	mm <sup>3</sup> /mole_formula_ unit/Pa

EXTENSIVE VARIABLES OF A COMPONENT IN A PHASE

Quantity	Suggested name and expression	Unit
Amount	MFij = N(ph,c)/NP(ph) = X(ph,c)	dimensionless
	MPij = N(ph,c)/NP(ph)*100	dimensionless
	WFi = B(ph,c)/BP(ph) = W(ph,c)	dimensionless
	WPij = B(ph,c)/BP(ph)*100	dimensionless
	BKgij = 1E-3*B(ph,c)	kilograms (kg)
	BTonij = 1E-6*B(ph,c)	tons

### INTENSIVE AND EXTENSIVE VARIABLES OF A SPECIES IN A PHASE

Quantity	Suggested name and expression	Unit	
For intensive variables of a species in a phase			
Chemical Potential	MUCalij = MU(sp,ph)/4.1858	cal/mol	
	MUErgij = MU(sp,ph)*1E7	cal/mol	
Fugacity	FAij = AC(sp,ph)*PATM	atmosphere (atm)	
	FBij = AC(sp,ph)*PB	bars (bar)	
	FCij = AC(sp,ph)*PKB	kilobars (kbar)	
	FTij = AC(sp,ph)*PTOR	Tors (millimeters of Hg)	

Quantity	Suggested name and expression	Unit
	LnFAij = LNAC(sp,ph)+LN(PATM)	atmosphere (atm)
	LnFBij = LNAC(sp,ph)+LN(PB)	bars (bar)
In(Fugacity)	LnFCij = LNAC(sp,ph)+LN(PKB)	kilobars (kbar)
	LnFTij = LNAC(sp,ph)+LN(PTOR)	Tors (millimeters of Hg)
	Alij=ACR(FE+3,AQ)*AH2O	dimensionless
Special quantity for	RCij=ACR(FE+3,AQ)*YH2O/Y(AQ,FE+3) where AH2O=55.508435 and YH2O=Y(AQ,H2O)	dimensionless
aqueous solution	WRCalij = WR(AQ,sp)/4.1858 (WR(AQ,sp) is only valid for aqueous species in a system in which the AQUEOUS solution phase is considered)	cal/mol
For extensive va	riables of a species in a phase	
Special	MLij=Y(AQ,FE+3)*AH2O/YH2O	molality (m)
quantity for aqueous solution	ISTR=1*IS1+1*IS2+1*IS3 where AH2O=55.508435, YH2O=Y(AQ,H2O) and ISn=.5*MLi*Zi**2+.5*MLj*Zj**2+.5*MLk*Zk**2	molality (m)

### UNITS FOR VARIABLES OF SPECIFIC SPECIES IN A PHASE

In the GIBBS, DATA, PARROT and ED\_EXP modules, there are quantities that describe various thermodynamic, physical, chemical and transport properties for a compound phase, and for a certain species or a given species combination in a specific solution phase. The following

table describes the units for those variables.

Variables	Description and unit
G(ph,sp) G(ph,sp;0) G(ph,sp1;sp2;;0)	Gibbs energy (J/mol) of a pure substance or end-member.
G(ph,sp1,sp2,) L(ph,sp1,sp2,) G(ph,sp1,sp2,;0) L(ph,sp1,sp2,;0) G (ph,sp1,sp2,;;0) L (ph,sp1,sp2,;;0)	Zero-order interaction parameter (J/mol) of a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
G(ph,sp1,sp2,;i) L(ph,sp1,sp2,;i) G (ph,sp1,sp2,;;i) L (ph,sp1,sp2,;;i)	The ith -order interaction parameter (J/mol) of a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
TC(ph,sp) TC(ph,sp;0) TC (ph,sp1;sp2;;0)	Curie temperature (K) of a pure substance or end-member.
TC(ph,sp1,sp2,) TC (ph,sp1,sp2,;0) TC (ph,sp1,sp2,;;0)	Zero-order Curie-temperature contribution (K) to a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.

These variables cannot be directly used in the POLY or POST modules.

Variables	Description and unit
TC(ph,sp1,sp2,;i) TC (ph,sp1,sp2,;;i)	The ith -order Curie-temperature contribution (K) to a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
BMAGN(ph,sp) BMAGN(ph,sp;0) BMAGN (ph,sp1;sp2;;0)	Bohr magneton number (dimensionless) of a pure substance or end-member.
BMAGN (ph,sp1,sp2,) BMAGN (ph,sp1,sp2,;0) BMAGN (ph,;sp1,sp2,;0)	Zero-order Bohr magneton number (dimensionless) to a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
BMAGN (ph,sp1,sp2,;i) BMAGN (ph,sp1,sp2,;;i)	The ith -order Bohr magneton number (dimensionless) to a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
V0(ph,sp) V0(ph,sp;0) V0 (ph,sp1;sp2;;0)	Molar volume (m <sup>3</sup> ) at 1 bar and reference T0 of a pure substance or end-member.
V0(ph,sp1,sp2,) V0 (ph,sp1,sp2,;0) V0 (ph,sp1,sp2,;;0)	Zero-order composition-dependent molar volume (m <sup>3</sup> ) of a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
V0(ph,sp1,sp2,;i) V0 (ph,sp1,sp2,;;i)	The ith-order composition-dependent molar volume (m <sup>3</sup> ) of a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.

Variables	Description and unit
VA(ph,sp) VA(ph,sp;0) VA (ph,sp1;sp2;;0)	Integrated thermal expansivity (m <sup>3</sup> /mol/K) of a pure substance or end-member.
VA(ph,sp1,sp2,) VA (ph,sp1,sp2,;0) VA (ph,sp1,sp2,;;0)	Zero-order composition-dependent thermal expansivity (m <sup>3</sup> /mol/K) of a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
VA(ph,sp1,sp2,;i) VA (ph,sp1,sp2,;;i)	The ith -order composition-dependent thermal expansivity (m <sup>3</sup> /mol/K) of a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
VC(ph,sp) VB(ph,sp;0) VB (ph,sp1;sp2;;0)	Isothermal compressibility (m <sup>3</sup> /mol/Pa) of a pure substance or end-member.
VC(ph,sp1,sp2,) VB (ph,sp1,sp2,;0) VB (ph,sp1,sp2,;;0)	Zero-order composition-dependent isothermal compressibility (m <sup>3</sup> /mol/Pa) of a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
VC(ph,sp1,sp2,;i) VB (ph,sp1,sp2,;;i)	The ith -order composition-dependent isothermal compressibility (m <sup>3</sup> /mol/Pa) of a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
VK(ph,sp) VB(ph,sp;0) VB (ph,sp1;sp2;;0)	High-pressure fitting parameter (m <sup>3</sup> /mol) of a pure substance or end-member.

Variables	Description and unit
VK(ph,sp1,sp2,) VB (ph,sp1,sp2,;0) VB (ph,sp1,sp2,;;0)	Zero-order composition-dependent high-pressure fitting parameter (m <sup>3</sup> /mol) of a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
VK(ph,sp1,sp2,;i) VB (ph,sp1,sp2,;;i)	The ith -order composition-dependent high-pressure fitting parameter (m <sup>3</sup> /mol) of a specified binary, ternary or higher-order interactions on a certain sublattice site in a solution phase.
WR(ph,sp) WR(ph,sp;0)	Standard Born function (J/mol) of a specific aqueous species in the AQUEOUS solution phase (always with a single sublattice) under the reference-state temperature and pressure.

# **Graphical Mode Examples Guide**

**Thermo-Calc Version 2018b** 





# **About the Graphical Mode Examples**

Graphical Mode examples are available for Thermo-Calc and the add-on Diffusion Module (DICTRA) and Precipitation Module (TC-PRISMA). These are in the format of project files (\*.tcu).

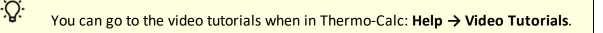


Also search the online help for "Opening Graphical Mode Project Files", which is in the *Thermo-Calc User Guide*.

This guide includes the example descriptions and information about how to open a Graphical Mode *project* file.

# Watch the Video Tutorials

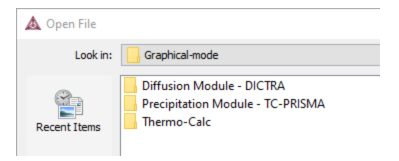
Many of the Graphical Mode examples have video tutorials available both on <u>our website</u> and our <u>YouTube channel</u>.



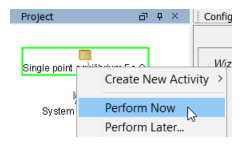
# **Opening a Project File**

To open an example project file for Thermo-Calc and the add-on Diffusion Module (DICTRA) and/or Precipitation Module (TC-PRISMA):

- 1. Open Thermo-Calc.
- 2. Confirm you are in Graphical Mode. If not, click the **Switch to Graphical Mode** button.
- 3. From the main menu, select File  $\rightarrow$  or Help  $\rightarrow$  Examples Files.
- 4. The examples are divided into folders by product (the add-on Diffusion Module (DICTRA) and/or Precipitation Module (TC-PRISMA) and Thermo-Calc). Navigate to the folder and project file you want and double-click to open.



5. The Graphical Mode examples are stored without calculated results so you will want to run the example calculation and generate a plot or table. To do this, click the first node in the tree at the top of the **Project** window. Then right-click and select **Perform now**.



8¢8

Some add-on module examples require additional licenses to run for more than three elements.

# **Thermo-Calc Graphical Mode Examples**

This guide includes descriptions of the following examples.

Category	Example number
Thermo-Calc including binary and ternary system examples, Scheil, and the Property Model and Equilibrium Calculators plus more	T_01 to T_12
"Diffusion Calculator Examples" on page 32	D_01, D_02, D_03 (the Quick Start Guide Examples) D_04 to D_07
"Precipitation Calculator Examples" on page 52	P_01 to P_11

# T\_01: Calculating a Single-Point Equilibrium

This example shows the result from a single-point equilibrium calculation in the Fe-C system. It demonstrates the use of the Equilibrium Calculator and generates a Table Renderer in the **Results** window. The number of equilibrium conditions is C+2 where C is the number of components, meaning that four conditions are needed in this example:

- Temperature is 1000 K
- Pressure is 101325 Pa
- System size is 1 mole
- Mass percent carbon is 0.1%

### **PROJECT FILE AND VIDEO TUTORIAL INFORMATION**

- Folder: Thermo-Calc
- File name: T\_01\_Single-point\_equilibrium.tcu
- This example is included in the tutorial on our website and YouTube channel.

### TABLE RESULT

The displayed results of the calculation show that the BCC\_A2 (ferrite) and GRAPHITE phases are stable for this set of equilibrium conditions.

Results					L X
Table Renderer 1					
System					
Moles	1.00000				
Mass	55.64392	[9]			
Temperature	1000.00000	[K]			
Total Gibbs Energy					
Enthalpy	24706.54116	[1]			
Volume	7.30619E-6	[m3]			
Component	Mole Fraction	Maga Eraction	Activity	Potential	
-					
c	0.00463	0.00100		-12658.89428	
Fe	0.99537	0.99900	0.00619	-42278.50542	
Stable Phases					
	Moles	Mass	Volume Fraction		
BCC_A2#1	0.99609	55.59692	0.99711	Composition 🗸	
Composition					
Component	Mole Fraction	Mass Fraction			
Fe	0.99928	0.99984			
с	0.00072	0.00016			
	Moles	Mass	Volume Fraction		
GRAPHITE#1	0.00391	0.04700	0.00289	Composition 🗸	
Composition					
Component	Mole Fraction	Mass Fraction			
с	1.00000	1.00000			
Fe	0.00000	0.00000			
		-			

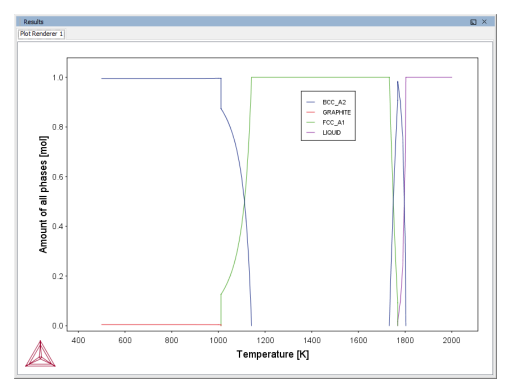
# T\_02: Stepping in Temperature in the Fe-C System

This example shows how the fractions of stable phases vary for an Fe-0.1 mass-% C alloy when the temperature is varied between 500 and 2000 K. It demonstrates the use of the Equilibrium Calculator. To allow temperature to vary, the corresponding **Axis Definition** check box is selected.

### **PROJECT FILE AND VIDEO TUTORIAL INFORMATION**

- Folder: Thermo-Calc
- File name: T\_02\_Step\_in\_temperature\_in\_Fe-C.tcu
- This example is included in the tutorial on our website and YouTube channel.

### **PLOT RESULT**



In this example, results are displayed graphically using a Plot Renderer activity.

If you want, you can use a Table Renderer to generate text results instead, as shown in this example after adding a Table Renderer to the Equilibrium Calculator and clicking **Perform**.

Results				E	. ×
not render er 1	ble Renderer 1		1	1	
Temperature [K]	Amount of BCC_A2 [mol]	Amount of FCC_A1 [mol]	Amount of GRAPHITE [mol]	Amount of LIQUID [mol]	
1000.00000	0.99609		0.00391		
1010.00000	0.99618		0.00382		
1011.17630	0.99620		0.00380		
1011.17630	0.99620	0.00000	0.00380		
1011.17630	0.87367	0.12633	0.00000		
1011.17630	0.87367	0.12633			
1020.00000	0.86050	0.13950			
1030.00000	0.84335	0.15665			
1040.00000	0.82359	0.17641			
1050.00000	0.80085	0.19915			
1060.00000	0.77399	0.22601			
1070.00000	0.74184	0.25816			
1080.00000	0.70282	0.29718			
1090.00000	0.65470	0.34530			
1100.00000	0.59418	0.40582			
1110.00000	0.51618	0.48382			
1120.00000	0.41250	0.58750			
1130.00000	0.26894	0.73106			
1140.00000	0.05863	0.94137			
1142.15538	0.00000	1.00000			
1142.15538		1.00000			
1150.00000		1.00000			
1160.00000		1.00000			
1170.00000		1.00000			
1180.00000		1.00000			
1190.00000		1.00000			
1200.00000		1.00000			
210.00000		1.00000			
220.00000		1.00000			

# T\_03: Fe-C Phase Diagrams

This example shows the stable Fe-C phase diagram (stable meaning that the graphite phase is entered in the calculation). The same diagram is calculated using the Binary Calculator activity and then using a System Definer and Equilibrium Calculator.

In both cases, a Plot Renderer is used to display results. The purpose of the Binary Calculator is to simplify common calculations for binary systems. In the Equilibrium Calculator two axes are defined: *Mass percent C* and *Temperature*. An axis variable must also be an equilibrium condition.

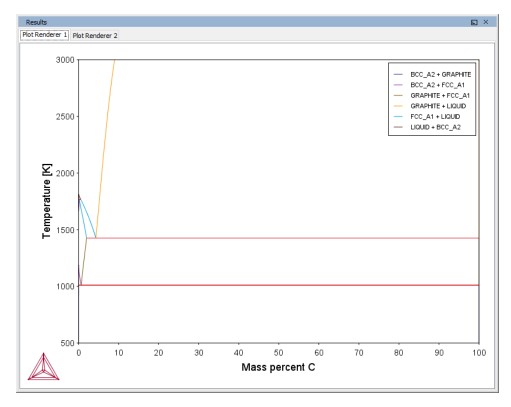
### PROJECT FILE AND VIDEO TUTORIAL INFORMATION

- Folder: Thermo-Calc
- File name: T\_03\_Fe-C\_phase\_diagram.tcu
- This example is included in the tutorial on our website and our YouTube channel.

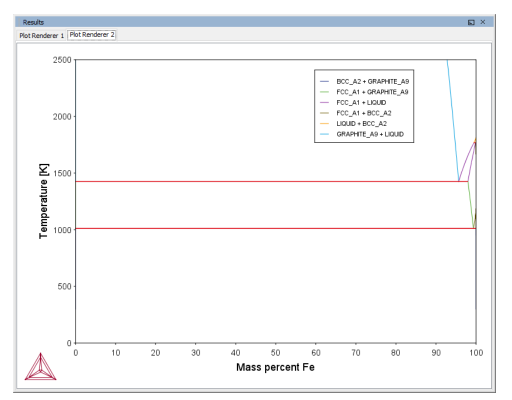
### **PLOT RESULTS**

### Equilibrium Calculator Results

This shows the plot result of the System Definer and Equilibrium Calculator system.



#### Binary Calculator Results



# T\_04: Fe-Cr-C Ternary Phase Diagram at 1000 K

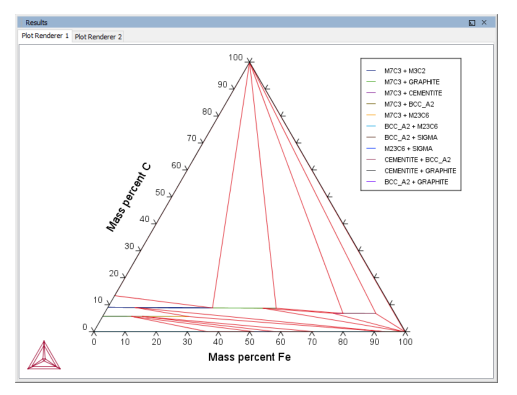
This example shows a ternary phase diagram in the Fe-Cr-C system at 1000 K. Similar to example <u>"T\_03: Fe-C Phase Diagrams" on page 8</u>, the same diagram is calculated using a Ternary Calculator and then using a System Definer and Equilibrium Calculator. Two Plot Renderer activities are added to create two plots.

#### **PROJECT FILE AND VIDEO TUTORIAL INFORMATION**

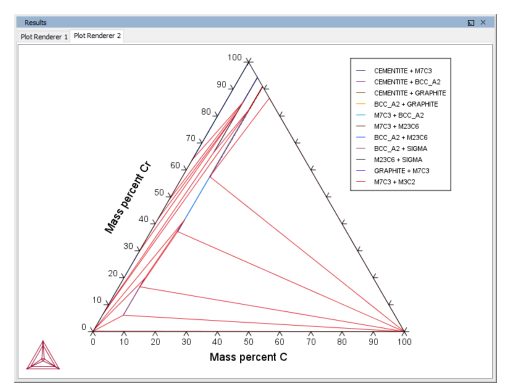
- Folder: Thermo-Calc
- File name: T\_04\_Fe-Cr-C\_ternary\_phase\_diagram.tcu
- This example is included in the tutorial on our website and YouTube channel.

### **PLOT RESULTS**

#### Ternary Calculator



#### Equilibrium Calculator



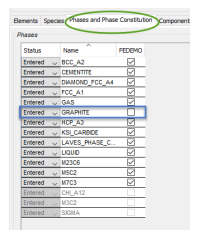


Try practicing with two features on the Plot Renderer Configuration window. To toggle between a triangular and a rectangular diagram, click the **Show Triangular** button then click **Perform** to see what happens. To toggle the X and Y axis variables, click the **Switch Axes** button and then click **Perform**.

# T\_05: Stable and the Metastable Fe-C Phase Diagrams

This example shows how to overlay results from two calculations in the same plot using the Equilibrium Calculator.

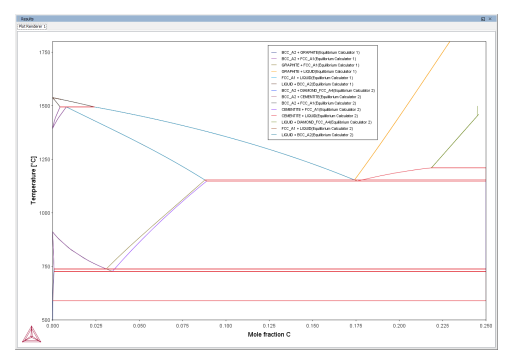
- 1. The Plot Renderer activity is first created as a successor to Equilibrium Calculator 1.
- 2. Then right-click the Plot Renderer node and select **Add predecessor** and the results from both equilibrium calculators are in the same plot. In *Equilibrium Calculator 1* the stable phase diagram is calculated and in *Equilibrium Calculator 2* the metastable phase.
- 3. The metastable diagram is obtained by deselecting the graphite phase on System Definer  $2 \rightarrow$  Phases and phase constitution tab.



### **PROJECT FILE NAME**

- Folder: Thermo-Calc
- File name: T\_05\_Fe-C\_stable\_and\_metastable\_phase\_diagram.tcu

#### **PLOT RESULT**



# **T\_06: Serially Coupled Equilibrium Calculators**

Sometimes there are multiple solutions for a given set of equilibrium conditions. In other cases the equilibrium calculation does not converge. You can then aid the final calculation by, in effect, telling the software where it should start the search for the equilibrium.

This example shows how to serially couple two Equilibrium Calculator nodes together for more complex equilibrium conditions. For each calculation, the output is to a Table Renderer.

#### **PROJECT FILE NAME**

- Folder: Thermo-Calc
- File name: *T\_06\_Serial\_equilibrium\_calculators.tcu*

### **Table Renderer Results**

#### **Equilibrium Calculator 1**

In the first example, the objective is to calculate the equilibrium at 850° C where the mole

fractions of the FCC A1 and BCC A2 phases are 0.5 each for an Fe-Cr-C steel with 10% chromium. In Equilibrium Calculator 1 a simple set of equilibrium conditions (temperature, pressure, system size and composition) are used to find a carbon content where only the FCC A1 and BCC A2 phases are stable. The results from this preliminary calculation are displayed in Table Renderer 1.

Results					
Table Renderer 1	Table Rendere	er 2 Table Ren	derer 3 Table R	enderer 4	
System					
Moles	1.00000				
Mass	55.41634	[a]			
Temperature	1123, 15000	K			
Total Gibbs Energy	-52361.90905	[1]			
	32589.15580				
Volume	7.34406E-6	[m3]			
Component	Mole Fraction	Mass Fraction	Activity	Potential	
с	0.00046	0.00010	0.00137	-61592.91681	
Cr	0.10658	0.10000	0.00235	-56531.30746	
Fe	0.89296	0.89990	0.00387	-51859.50665	
Stable Phases					
	Moles	Mass	Volume Fraction		
BCC_A2#1	0.76777	42.55315	0.76992	Composition ~	
Composition					
		Mass Fraction			
Fe	0.89183	0.89864			
Cr	0.10801	0.10133			
с	0.00016	0.00003			
	Moles	Mass	Volume Fraction		
FCC_A1#1	0.23223	12.86320	0.23008	Composition ~	
Composition					
		Mass Fraction			
	0.89669	0.90408			
	0.10185	0.09560			
с	0.00147	0.00032			

#### **Equilibrium Calculator 2**

In Equilibrium Calculator 2 the carbon content equilibrium condition is replaced by the condition that the numbers of moles of the BCC A2 phase should be 0.5. The final result is displayed in Table Renderer 2. In this case, the final objective is to calculate the solidus temperature of an Fe-Cr-C steel with 10% Cr and 0.01% C.

Results						E		E ×	
Table Renderer 1	Table Rendere	r 2 Table Ren	derer 3 Table R	enderer 4					
System									
Moles	1.00000								
Mass	55.39810	[9]							
Temperature	1123.15000	[K]							
Total Gibbs Energy									
Enthalpy	32827.91821								
Volume	7.32017E-6	[m3]							
Component	Mole Fraction	Mass Fraction	Activity	Potential					
C	0.00088			-60967.89084					
Cr	0.10654			-56409.41447					
Fe	0.89258			-51874.52376					
-									
Stable Phases									
	Moles	Mass	Volume Fraction						
BCC_A2#1	0.50000	27.70824	0.50303	Composition	/				
Composition									
Component	Mole Fraction	Mass Fraction							
Fe	0.89000	0.89692							
Cr	0.10982	0.10304							
с	0.00017	0.00004							
	Moles	Mass	Volume Fraction						
FCC_A1#1	0.50000	27.68987	0.49697	Composition 🗸	/				
Composition									
Component	Mole Fraction	Mass Fraction							
Fe	0.89515	0.90270							
Cr	0.10326	0.09695							
с	0.00159	0.00034							

### Equilibrium Calculator 3

In Equilibrium Calculator 3 the state at 2000 K is calculated and the result is displayed in Table Renderer 3.

Results						
Table Renderer 1	Table Rendere	er 2 Table Ren	derer 3 Table R	enderer 4		
System	4 00000					
Moles	1.00000					
Mass	55.41634	[g]				
		[K]				
Total Gibbs Energy Enthalpy	-52361.90905 32589.15580					
Volume	7.34406E-6	[m3]				
Component	Mole Fraction	Mass Fraction	Activity	Potential		
c	0.00046			-61592.91681		
Cr	0.10658			-56531.30746		
Fe	0.89296	0.89990		-51859.50665		
Stable Phases						
	Moles	Mass	Volume Fraction			
BCC_A2#1	0.76777	42.55315	0.76992	Composition	$\sim$	
Composition						
Component	Mole Fraction	Mass Fraction				
Fe	0.89183	0.89864				
Cr	0.10801	0.10133				
с	0.00016	0.00003				
	Moles	Mass	Volume Fraction			
FCC_A1#1	0.23223	12.86320	0.23008	Composition	$\sim$	
Composition						
Component	Mole Fraction	Mass Fraction				
Fe	0.89669	0.90408				
Cr	0.10185	0.09560				
с	0.00147	0.00032				

### **Equilibrium Calculator 4**

In Equilibrium Calculator 4 the temperature equilibrium condition is replaced by the condition *Fix phase / liquid / 0.0*, meaning that liquid should be stable in an amount of zero moles, i.e. the solidus temperature. The final result is displayed in Table Renderer 4.

Results					
Table Renderer 1	Table Render	er 2 Table Re	nderer 3	able Renderer 4	
System					
Moles	1.00000				
Mass	55.41634	[g]			
Temperature	1791.51760	[K]			
Total Gibbs Energy	-1.09463E5	[0]			
	59070.77313				
Volume	7.63568E-6	[m3]			
Component	Mole Fraction	Mass Fraction	Activity	Potential	
C	0.00046	0.00010	0.00012	-1.34945E5	
Cr	0.10658	0.10000	0.00030	-1.20950E5	
Fe	0.89296	0.89990	0.00071	-1.08079E5	
Stable Phases					
	Moles	Mass	Volume Fra	ction	
BCC_A2#1	1.00000	55.41634	1.00000	Composition	~
Composition					
Component		Mass Fraction			
Fe	0.89296	0.89990			
Cr	0.10658	0.10000			
C	0.00046	0.00010			
Fixed Phases					
	Moles	Mass	Volume Fra	oction	
LIQUID#1	0.00000	0.00000	0.00000	Composition	~
Composition					
		Mass Fraction			
	0.88607	0.89447			
	0.11178	0.10506			
C	0.00216	0.00047			

# **T\_07: User-Defined Functions**

This example shows how to implement user-defined functions.

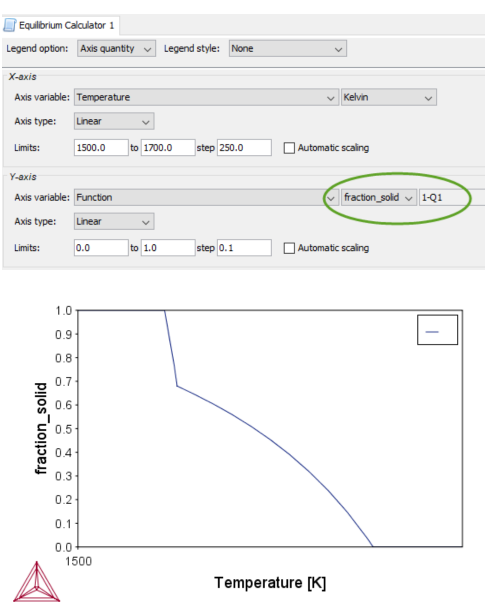
A series of equilibria for an Fe-Cr-C alloy are calculated by varying temperature between 500 and 3000 K. In the configuration window of the Equilibrium Calculator under the **Functions** tab, two identically meaning functions are defined, *fraction solid* and *f solid*, the values of which are plotted against temperature in two Plot Renderer activities. Functions can be entered in terms of *QuantitiesQ1*, *Q2*, *Q3* and so forth, or by using the Thermo-Calc syntax.

#### **PROJECT FILE INFORMATION**

- Folder: Thermo-Calc
- File name: T\_07\_User\_defined\_functions.tcu

### **PLOT RESULT**

*This plot example shows the result of the fraction solid function.* 



# T\_08: Scheil and Equilibrium Solidification

This example shows a comparison for an Al-Si alloy solidified under full local equilibrium and under the Scheil assumptions, i.e. zero diffusion in the solidified material and infinitely fast diffusion in the liquid. The example uses the Scheil Calculator and adds a Plot and Table Renderer.

### PROJECT FILE AND VIDEO TUTORIAL INFORMATION

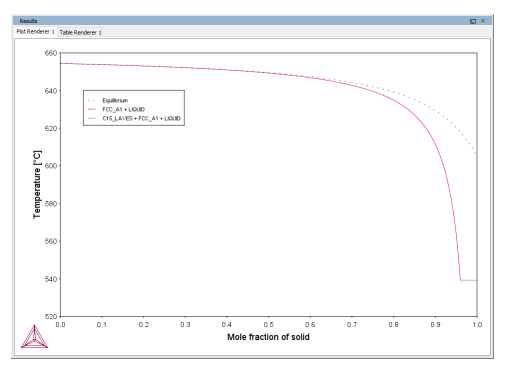
- Folder: Thermo-Calc
- File name: *T\_08\_Scheil\_and\_equilibrium\_solidification.tcu*.

For other examples of using Scheil, see our website and YouTube channel:

- Modified Austenitic Stainless Steel 18/8 Scheil Solidification Simulation
- Applications to Welding and Joining with Bonus Scheil Example

#### TABLE AND PLOT RESULTS

The plot compares the mole fraction of a solid vs temperature. The table provides the data, which you can also export.



Results	[	
Plot Renderer 1 Table Renderer 1		
Mole fraction of solid	Temperature [°C]	
-2.22045E-16	654.52000	^
-2.22045E-16	654.52000	
-2.22045E-16	654.52000	
0.00105	654.42400	
0.16114	653.42400	
0.27839	652.42400	
0.36780	651.42400	
0.43812	650.42400	
0.49481	649.42400	
0.54143	648.42400	

#### TO SAVE THE TABLE

There are two ways to save the table.

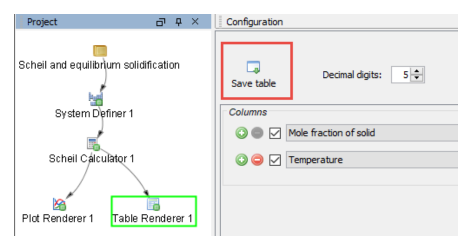
#### Method 1

In the Results window, right click the table and select Save As...

enderer 1		
Temperature [°	c]	
654.52000		
654.52000		
654.52000	Сору	
654.42400		
653.42400	Copy all	
652.42400	Save As	
651.42400	Print	
650.42400		
649.42400	Properties	
648.42400		
	654.52000           654.52000           654.52000           654.52000           654.42400           653.42400           652.42400           651.42400           650.42400           649.42400	

#### Method 2

- 1. In the **Project** window, click the **Table Renderer**.
- 2. In the **Configuration** window, click **Save table**.

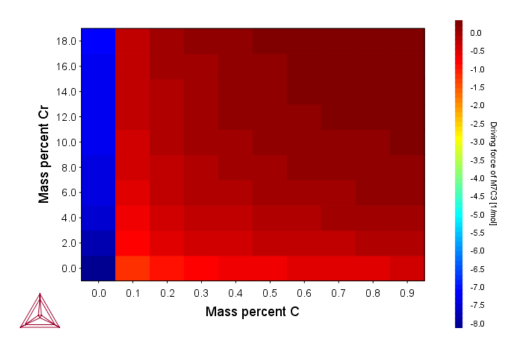


# T\_09: Carbide Driving Force Heat Map

This is an example of using *property grid* calculations to plot the driving force for a carbide as a function of two composition variables. With the property grid calculation type, a 2D grid is generated from the two calculation axes. After the calculation is done, an equilibrium is calculated in each grid point. A Plot Renderer connected to a property grid calculator plots the *z*-axis property for each equilibrium as a function of the two calculation axes. The final plot can be either a heat map or a contour plot.

### **PROJECT FILE NAME**

- Folder: Thermo-Calc
- File name: T\_09\_Heat\_map\_of\_carbide\_driving\_force.tcu



## **PLOT RESULT**

# T\_10: Phase Transition

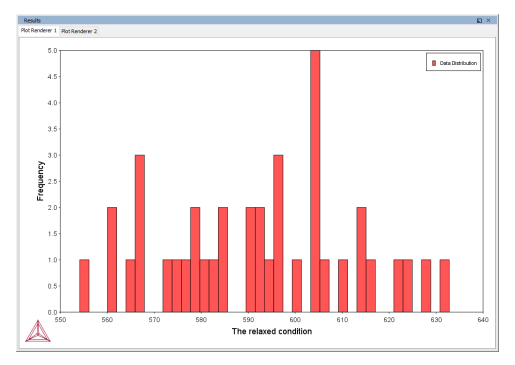
The example uses the Property Model Calculator to predict the transition temperature to the unwanted brittle Sigma phase. The example shows how the temperature is influenced by changes to a steel alloy's composition using the *uncertainty* calculation type and how to create either a *histogram* or *probability* plot.

### **PROJECT FILE AND VIDEO TUTORIAL INFORMATION**

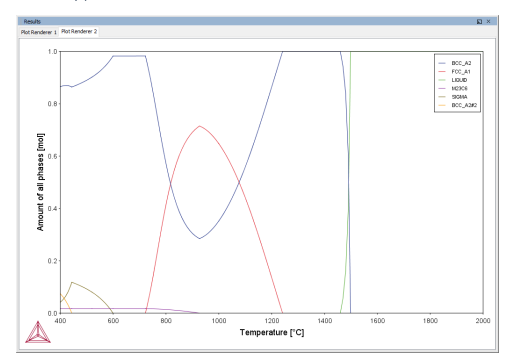
- Folder: Thermo-Calc
- File name: T\_10\_Phase\_Transition.tcu
- This example is included in the tutorial on our <u>website</u> and <u>YouTube</u> channel. Also see the <u>video introduction</u> to this feature.

### **PLOT RESULTS**

### Histogram plot



### Probability plot



Experienced users can also create custom property models for use with the Property Model Calculator.

Search for Property Model Calculator and Development Framework in the help.

# T\_11: Coarsening and Interfacial Energy

The example uses the Property Model Calculator and both thermodynamic and kinetic demonstration steel databases. Using a *grid* calculation type it produces these plots: a *heat map*, a *contour plot* and a *cross plot with contour* (where both the interfacial energy and coarsening rate is shown). It also creates a property diagram to show the phase fractions vs time and a 3D plot comparing the coarsening rate coefficient.

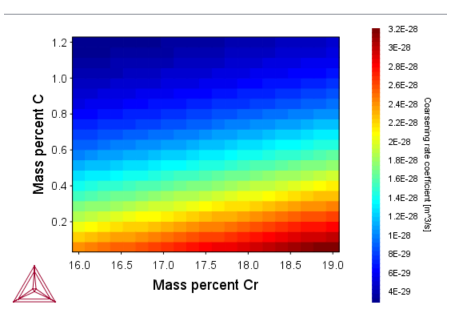
### **PROJECT FILE AND VIDEO TUTORIAL INFORMATION**

- Folder: Thermo-Calc
- File name: T\_11\_Coarsening\_and\_Interfacial\_energy.tcu
- This example is included in the tutorial on our <u>website</u> and our <u>YouTube</u> channel. Also see the <u>video introduction</u> to this feature.

## **Plot Results**

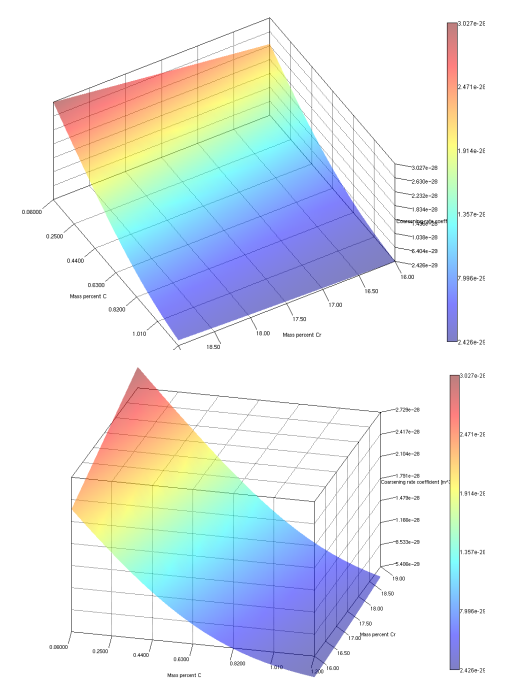
÷Ω

#### Coarsening rate coefficient (heat map)



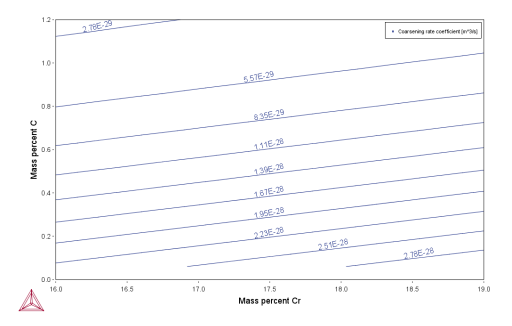
### Coarsening rate coefficient (3D plot)

Two different rotation angles. When viewing in Thermo-Calc you can rotate the plot.

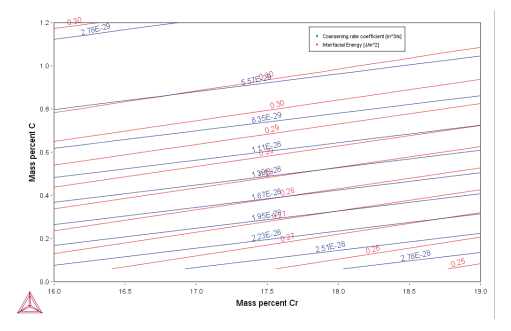




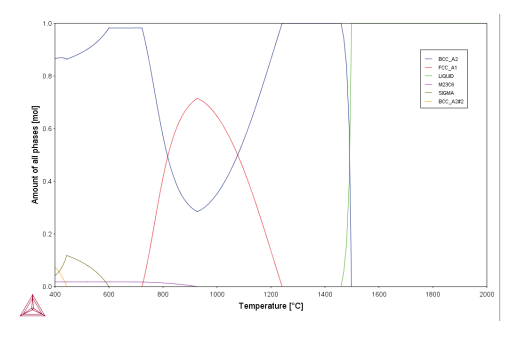
### Coarsening rate coefficient



### Coarsening rate coefficient and Interfacial energy (overlayed plots)



Phase fractions vs T (Property diagram)



Experienced users can also create custom property models for use with the Property Model Calculator.

Search for Property Model Calculator and Development Framework in the help.

# T\_12: Driving Force and Interfacial Energy

The example uses the Property Model Calculator and a thermodynamic demonstration steel database. Using a *grid* calculation type it produces a *contour* plot comparing the driving force and interfacial energy.

#### **PROJECT FILE AND VIDEO TUTORIAL INFORMATION**

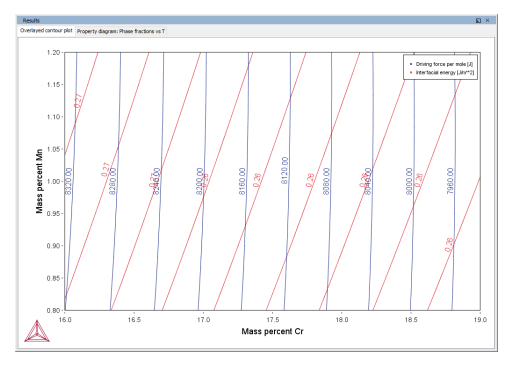
• Folder: Thermo-Calc

÷Ω

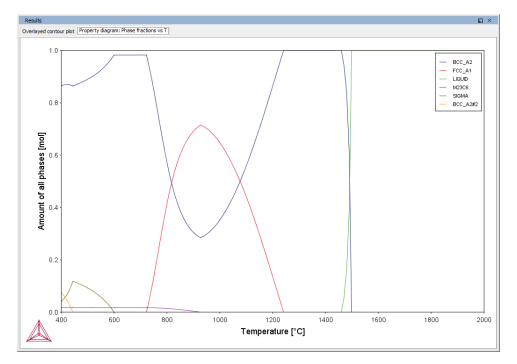
- File name: T\_12\_Driving\_force\_and\_Interfacial\_energy.tcu
- See the video introduction to this feature.

### **PLOT RESULT**

A contour plot



Phase Fraction



Graphical Mode Examples Guide | 30 of 87

# Experienced users can also create custom property models for use with the Property Model Calculator.

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Search for Property Model Calculator and Development Framework in the help.

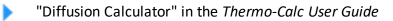
# **Diffusion Calculator Examples**

Examples using up to three elements are available to all users. The other examples require a Diffusion Module (DICTRA) license to calculate and plot results.



8

All examples use demonstration database packages included with your installation no matter what licenses you have.



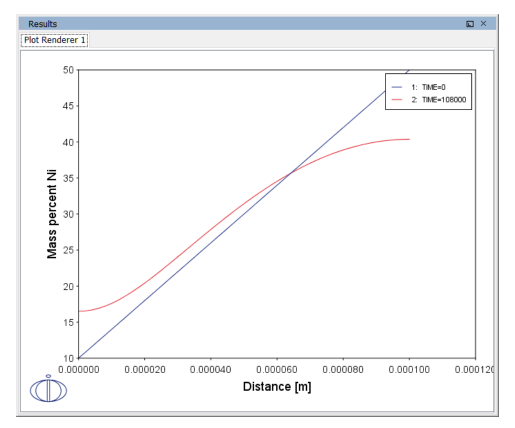
# **D\_01** Homogenization of a Binary Fe-Ni Alloy

The single phase example simulates the diffusion of Fe and Ni at a temperature of 1400 K in a planar domain. At this temperature the material is fully austenitic, i.e. the only phase present is the so-called fcc (face centered cubic) phase. Initially, there is a linear variation in Ni going from 10 mass-% on the left-hand side to 50 mass-% on the right-hand side.



This example is available as a video tutorial on <u>our website</u> and <u>YouTube</u> <u>channel</u>. In previous versions of the software (prior to 2017b) it is example 19.

### Composition of Ni vs Distance



## **PROJECT FILE NAME**

- Folder: Diffusion Module DICTRA
- File name: *D\_01\_Diffusion\_Single\_Phase.tcu*
- This example is part of the *Diffusion Module (DICTRA) Quick Start Guide* available to all users. A version of the example is also available for Console Mode.

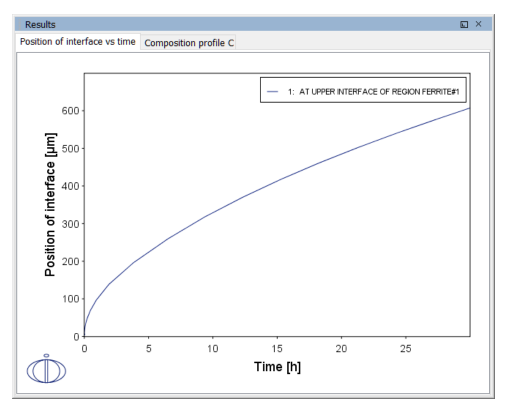
# D\_02 Ferrite(bcc)/Austenite(fcc) Transformation in a Binary Fe-C Alloy

The moving phase boundary example simulates the growth of ferrite (bcc) into austenite (fcc). The austenite is assumed to be initially homogeneous with the composition Fe-0.15 mass-% C. The transformation temperature is 1050 K. The initial thickness of the austenite is 2 mm and an initially very thin ferrite (1 nm) is also present at the start of the simulation.

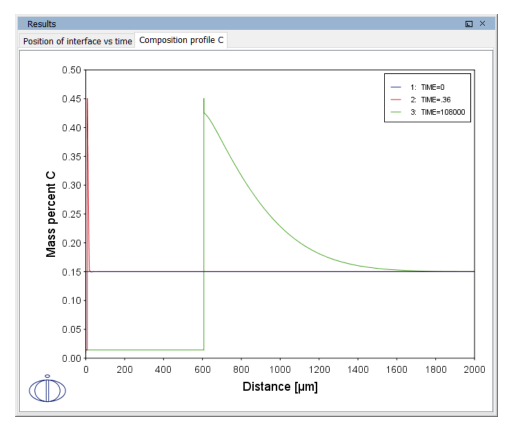


This example is available as a video tutorial on <u>our website</u> and <u>YouTube</u> <u>channel</u>. In previous versions of the software (prior to 2017b) it is example 20.

# Position of interface vs time



## Composition profile C



## **PROJECT FILE NAME**

- Folder: Diffusion Module DICTRA
- File name: *D\_02\_Diffusion\_Moving\_Boundary.tcu*
- This example is part of the *Diffusion Module (DICTRA) Quick Start Guide* available to all users. A version of the example is also available for Console Mode.

# **D\_03** Evolution of an Fe-Cr-Ni Diffusion Couple

The multiphase example simulates the evolution of an Fe-Cr-Ni diffusion couple during a 100 hour heat treatment at 1100° C (1373.15 K). Both end members of the diffusion couple are duplex ferrite plus austenite, but the majority phase is ferrite on the left-hand side and austenite on the right.

With this type of simulation it is assumed that the material is fully equilibrated at each grid point, i.e. the local phase fractions, phase compositions and so forth are obtained from an equilibrium calculation with the local overall composition as a condition.

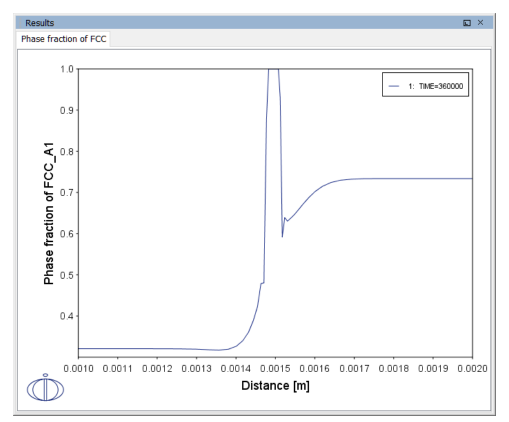
More details about the homogenization model for multiphase simulations can be found in Larsson and Engström (2006)<sup>1.</sup> and Larsson and Höglund (2009)<sup>2.</sup>. Experimental data is from Engström (1995)<sup>3.</sup>.

<sup>1. &</sup>quot;A homogenization approach to diffusion simulations applied to  $\alpha + \gamma$  Fe–Cr–Ni diffusion couples", *Acta Mater.*, 54 (9), pp. 2431–2439.

<sup>&</sup>lt;sup>2</sup>. "Multiphase diffusion simulations in 1D using the DICTRA homogenization model", *Calphad*, 33 (3), pp. 495–501.

<sup>3. &</sup>quot;Interdiffusion in multiphase, Fe-Cr-Ni diffusion couples", Scand. J. Metall., 24(1), 12–20.

### Phase fraction of FCC



### **PROJECT FILE NAME**

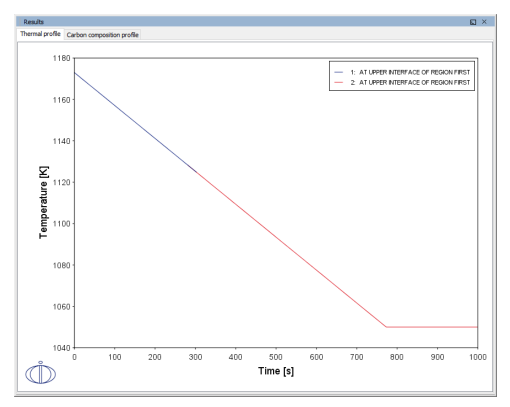
- Folder: Diffusion Module DICTRA
- File name: D\_03\_Diffusion\_Multiphase.tcu

This example is part of the Diffusion Module (DICTRA) Quick Start Guide available to all users. A version of the example is also available for Console Mode.

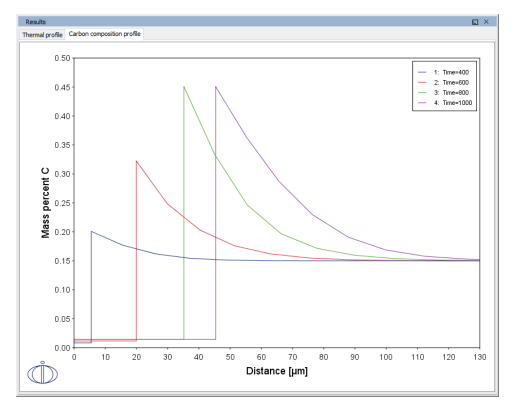
# **D\_04 Fe-C Moving Boundary: Austenite to Ferrite**

This example simulates the austenite to ferrite transformation in a Fe-0.01 mass% C steel during continuous cooling. The simulation starts at a temperature where only austenite is stable, ferrite nucleates and grows into the austenite during cooling in the two phase region. Plots of thermal and carbon composition profiles are generated.

## Thermal profile



## Carbon composition profile



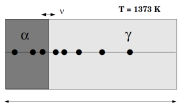
## **PROJECT FILE NAME**

- Folder: Diffusion Module DICTRA
- File name: D\_04\_Diffusion\_Fe-C\_Moving\_Boundary\_Austenite\_to\_Ferrite.tcu
- This example is available to all users.

# D\_05 $\gamma/\alpha/\gamma$ Diffusion couple of Fe-Ni-Cr alloys

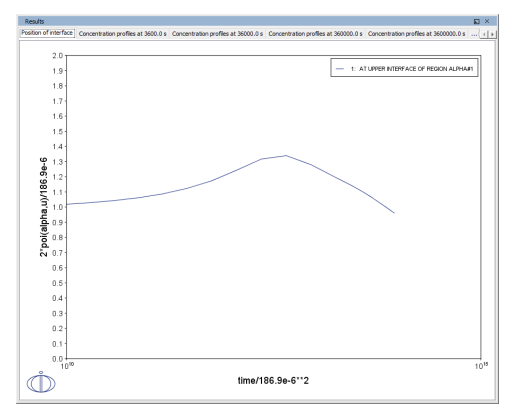
This example demonstrates the evolution of a ternary Fe-Cr-Ni diffusion couple. A thin slice of ferrite ( $\alpha$  phase) (38%Cr,0%Ni) is clamped between two thicker slices of austenite ( $\gamma$  phase) (27%Cr, 20%Ni). The assembly is subsequently heat treated at 1373 K.

This set up corresponds to diffusion couple in Kajihara, Lim, and Kikuchi (1993)<sup>1</sup>. Also see Kajihara and Kikuchi (1993)<sup>2</sup>.





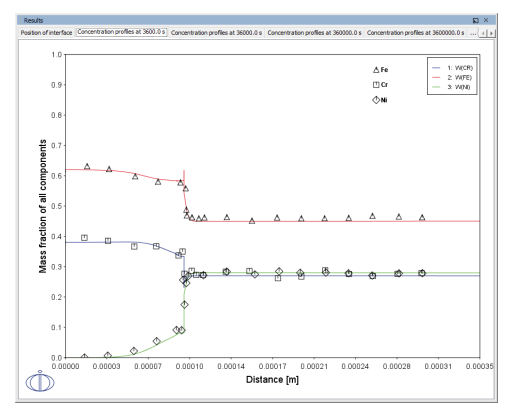
#### Interface Position

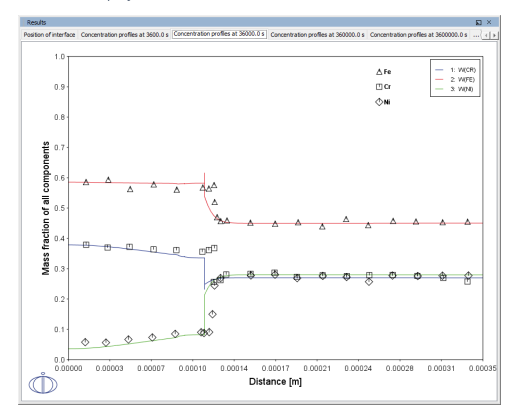


<sup>&</sup>lt;sup>1.</sup> "Experimental Study on Dissolution of ALPHA Phase in GAMMA/ALPHA/GAMMA Diffusion Couples of the Fe-Cr-Ni System", *ISIJ Int.*, 33(4), 498–507.

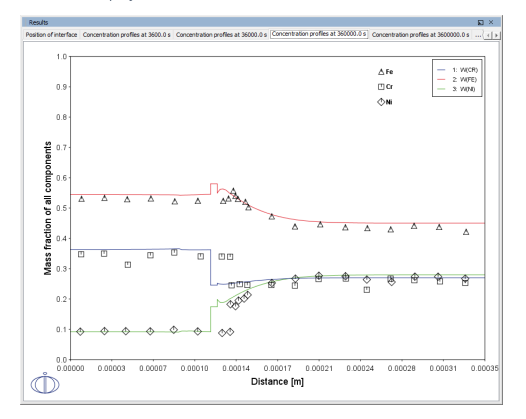
<sup>&</sup>lt;sup>2.</sup> "Numerical analysis of dissolution of  $\alpha$  phase in  $\gamma/\alpha/\gamma$  diffusion couples of the Fe-Cr-Ni system", *Acta Metall. Mat.*, 41 (7), 2045–2059.



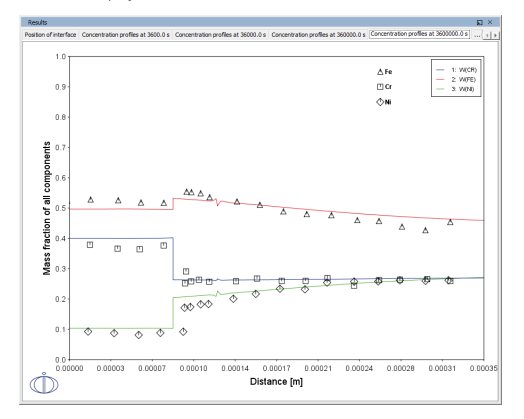




## Concentration profiles at 36000 s

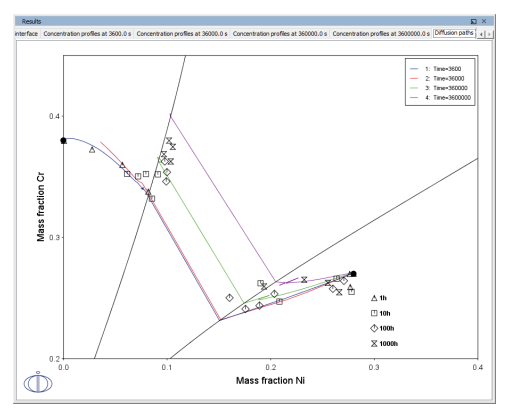


## Concentration profiles at 36 0000 s



## Concentration profiles at 3 600 000 s

### Diffusion paths



### **PROJECT FILE NAME**

- Folder: Diffusion Module DICTRA
- File name: D\_05\_Diffusion\_Fe\_Ni\_Cr\_Moving\_Boundary\_Diffusion\_Couple.tcu

# **D\_06 Diffusion Through a Tube Wall**

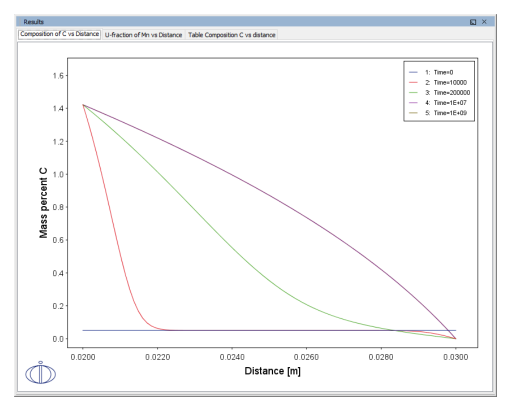
This is a simple example of diffusion through a tube wall . The tube material is an Fe-0.06Mn-0.05C alloy. Two plots comparing distance to the U-fraction of manganese and composition of carbon are generated to visualize the austenite region. A *cylindrical* geometry is used with *mixed zero flux and activity* boundary conditions.

On the inside wall a carbon activity of 0.9 is maintained whereas on the outside the carbon activity is very low. This example demonstrates the use of boundary conditions, advanced plotting and tables.

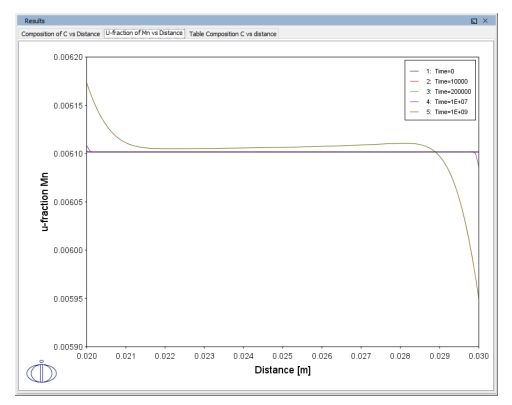


This example is based on Console Mode exab6 (see in the *Diffusion Module* (*DICTRA*) Console Mode Examples Guide). When in Console Mode, you can open the example from Thermo-Calc (File  $\rightarrow$  Examples Files  $\rightarrow$  Diffusion Module).

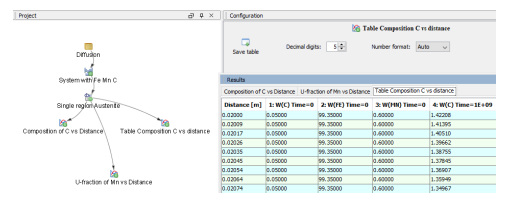
# Composition of C vs Distance



## U-fraction of Mn vs Distance



#### Table Composition C vs Distance



### **PROJECT FILE NAME**

- Folder: Diffusion Module DICTRA
- File name: D\_06\_Diffusion\_Carburization\_Tube.tcu

# D\_07 Multiphase Carburization of a Ni-25Cr-0.0001C alloy

This example is about carburization of a Ni-25Cr-0.0001C alloy. In this case the M7C3 and M3C2 carbides are entered as spheroid phases in an FCC\_A1 matrix. It is similar to Graphical Mode example D\_06 except the default **Simulation condition** is automatically set to use the **Homogenization** model.

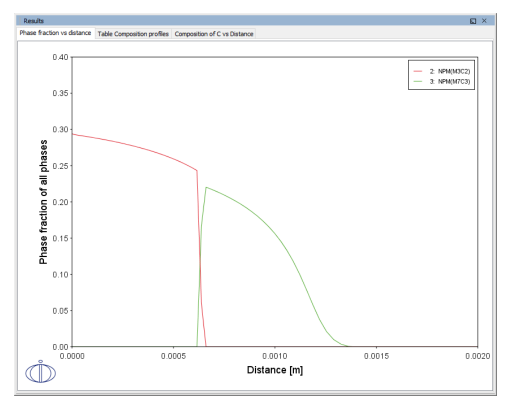
The Isothermal calculation is run for 1000 hours at a temperature of 1123 K using the **Mixed zero flux and activity** left boundary condition. Results are plotted using two Plot Renderers and a Table Renderer.

This case is from Engström, Höglund, and Ågren (1994), "Computer simulation of diffusion in multiphase systems", *Met. Mat. Trans. A*, 25(6), 1127–1134.



This example is based on Console Mode exd1b (see in the *Diffusion Module* (*DICTRA*) Console Mode Examples Guide). When in Console Mode, you can open the example from Thermo-Calc (File  $\rightarrow$  Examples Files  $\rightarrow$  Diffusion Module).

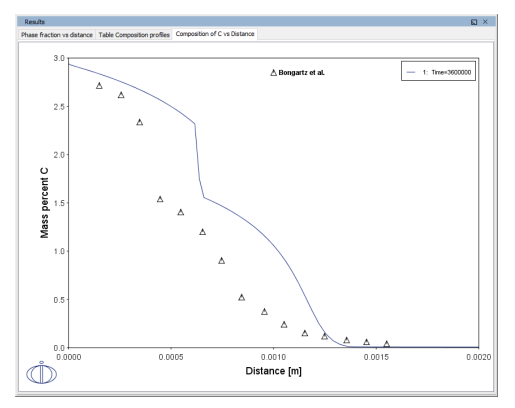
Phase fraction vs distance



# Table of Composition profiles

Results					×
Phase fraction vs distance Table Compos		sition profiles	Composition of C vs Distance		
Distance [m]	1: W(C)	2: W(CR)	3: W(NI)		
0.00000	2.93966	24.35835	72.70199		~
9.88104E-6	2.92919	24.27458	72.79623		
0.00002	2.92153	24.27324	72.80522		-
0.00003	2.91543	24.27447	72.81011		
0.00004	2.90899	24.27584	72.81517		-
0.00005	2.90235	24.27733	72.82032		
0.00006	2.89549	24.27886	72.82565		
0.00007	2.88841	24.28053	72.83106		-
0.00008	2.88107	24.28222	72.83670		
0.00010	2.87349	24.28403	72.84248		
0.00011	2.86564	24.28588	72.84849		
0.00012	2.85750	24.28783	72.85467		
0.00013	2.84907	24.28984	72.86110		
0.00014	2.84033	24.29194	72.86773		
0.00016	2.83124	24.29412	72.87464		
0.00017	2.82181	24.29640	72.88179		
0.00018	2.81201	24.29877	72.88922		
0.00020	2.80183	24.30124	72.89693		
0.00021	2.79122	24.30381	72.90497		
0.00022	2.78018	24.30650	72.91332		
0.00024	2.76867	24.30930	72.92203		
0.00025	2.75666	24.31222	72.93112		
0.00027	2.74412	24.31528	72.94060		
0.00028	2.73101	24.31848	72.95051		
0.00030	2.71731	24.32182	72.96087		
0.00031	2.70295	24.32533	72.97171		
0.00033	2.68790	24.32901	72.98308		
0.00035	2.67210	24.33288	72.99502		
0.00036	2.65550	24.33694	73.00756		
0.00038	2.63803	24.34122	73.02075		
0.00040	2.61961	24.34573	73.03466		
0.00042	2.60016	24.35050	73.04934		
0.00043	2.57960	24.35554	73.06486		~

## *Composition of C vs Distance*



### **PROJECT FILE NAME**

- Folder: Diffusion Module DICTRA
- File name: *D\_07\_Diffusion\_Carburization\_Multiphase.tcu*

# **Precipitation Calculator Examples**

Examples that use three elements are available to all users. The other examples require a Precipitation Module (TC-PRISMA) license to calculate and plot results.



8

All examples use demonstration database packages included with your installation. You can open the examples from the main menu: File  $\rightarrow$  or Help  $\rightarrow$  Examples Files.



Unless specified in tables for each example, all the numerical parameters are assumed default values.

# P\_01: Isothermal Precipitation of Al<sub>3</sub>Sc

This example simulates the kinetics of precipitation of  $Al_3Sc$  from an FCC\_A1 solution phase. The simulation results can be compared with experimental data collected from Marquis and Seidman<sup>1.</sup> and Novotny and Ardell (2001)<sup>2.</sup>.



This example is available as a video tutorial on <u>our website</u> and <u>YouTube</u> channel.

System (System Definer)		
Database package	Demo: Aluminum-based alloys (ALDEMO, MALDEMO)	
Elements	Al, Sc	
Conditions (Precipitation Calculator)		
Composition	Al-0.18Sc Mole percent	
Matrix phase	FCC_A1	

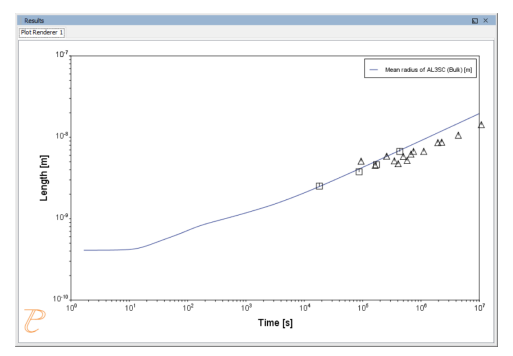
<sup>&</sup>lt;sup>1.</sup> Marquis, E.A, and D.N Seidman. 2001. "Nanoscale Structural Evolution of Al3Sc Precipitates in Al(Sc) Alloys." Acta Materialia 49 (11): 1909–19.

<sup>&</sup>lt;sup>2.</sup> Novotny, Gabriel M., and Alan J. Ardell. 2001. "Precipitation of Al3Sc in Binary Al–Sc Alloys." Materials Science & Engineering, A: Structural Materials: Properties, Microstructure and Processing 318 (1–2): 144–54.

Precipitate phase	AL3SC	
Precipitate Phase Data Parameters (Precipitation Calculator)		
Nucleation sites	Bulk	
Interfacial energy	Calculated	
Calculation Type (Precipitation Calculator)		
Calculation type	Isothermal	
Temperature	350° C	
Simulation time	1.0E7 seconds	
Other		
File name and location	Folder: Precipitation Module - TC-PRISMA File name: P_01_Precipitation_Al-Sc_AL3SC.tcu	

# **Plot Results**

### Mean radius



# P\_02: Stable and Metastable Carbides - Isothermal

This example simulates the kinetics of precipitation of both stable and metastable carbides from ferrite phase. It demonstrates that metastable carbides (cementite, M7C3) may first emerge and then disappear and the stable phase (M23C6) prevails.

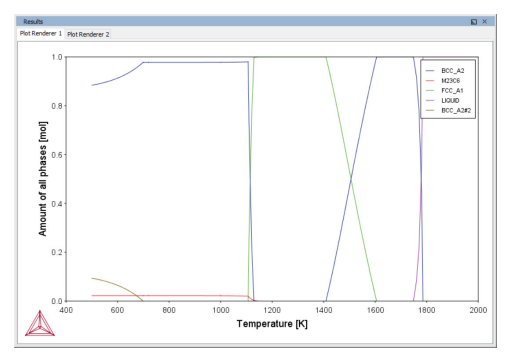
This example uses the Equilibrium Calculator and a property diagram to determine how the phases change with temperature. We are interested in the carbide precipitation at 1053K where only the carbide M23C6 is stable according to the equilibrium calculation. The Precipitation Calculator is used to do an isothermal calculation of the three phases (cementite, M23C6 and M7C3) where cementite and M7C3 are metastable phases.

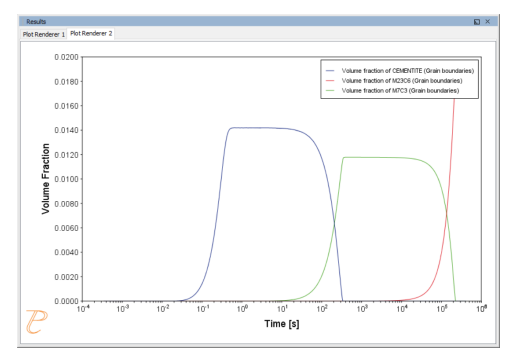
System (System Definer)			
Database package	Demo: Steels and Fe-alloys (FEDEMO,MFEDEMO)		
Elements	Fe, C, Cr		
Conditions (Precipitation Calculator)			
Composition	Fe-0.1C-12Cr Mass percent		
Matrix phase	BCC_A2		
Precipitate phases	Cementite, M23C6 and M7C3		
Matrix Phase Data Parameters (Precipitation Calculator)			
Grain size (click <b>Show details</b> to display this setting)	1.0E-4 m		
Precipitate Phase Data Parameters (Precipitation Calculator)			
Nucleation sites	Grain boundaries		
Interfacial energy	Cementite 0.167 J/m <sup>2</sup> , M23C6 0.252 J/m <sup>2</sup> , M7C3 0.282 J/m <sup>2</sup>		
Calculation Type (Precipitation Calculator)			
Calculation type	lsothermal		
Temperature	1053 К		

Simulation time	400 000 seconds
Other	
File name and location	Folder: Precipitation Module - TC-PRISMA File name: P_02_Precipitation_Fe-C-Cr_Cementite-M7C3- M23C6.tcu

# **Plot Results**

Equilibrium Calculator: How the phases change with temperature





Precipitation Calculator: Volume fractions

# P\_03: Stable and Metastable Carbides - TTT Diagram

In this example, the kinetics of precipitation of both stable and metastable carbides is calculated from the ferrite phase. It demonstrates that metastable carbides may first emerge and then disappear and the stable phase prevails.

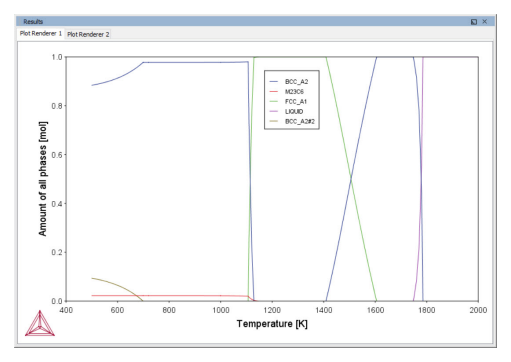
This example uses the Equilibrium Calculator and a property diagram to determine how the phases change with temperature. Using this result, the Precipitation Calculator is used to do a TTT (Time-Temperature-Transformation) diagram calculation of the three phases (cementite, M23C6 and M7C3) at the grain boundaries.

For a TTT diagram calculation, select **TTT diagram** in **Calculation Type**, then enter **Min**, **Max**, and **Step** of **Tempeature**, as well as **Max annealing time**. In **Stop criterion**, choose **Volume fraction of phase** and enter the value.

System (System Definer)		
Database package	Demo: Steels and Fe-alloys (FEDEMO and MFEDEMO)	

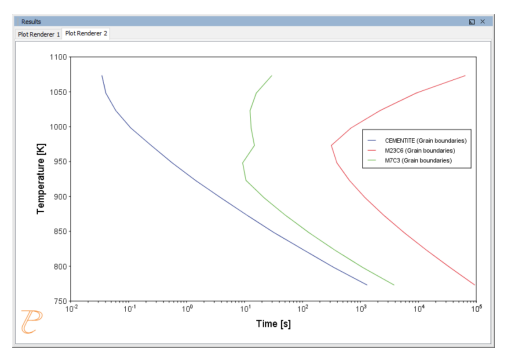
Elements	Fe, C, Cr	
Conditions (Precipitation Calculator)		
Composition	Fe-0.1C-12Cr Mass percent	
Matrix phase	BCC_A2	
Precipitate phases	Cementite, M23C6 and M7C3	
Matrix Phase Data Parameters (Precipi	tation Calculator)	
Grain size (click <b>Show details</b> to display this setting)	1.0E-4 m	
Precipitate Phase Data Parameters		
Nucleation sites	Grain boundaries	
Interfacial energy	Cementite 0.167 J/m <sup>2</sup> , M23C6 0.252 J/m <sup>2</sup> , M7C3 0.282 J/m <sup>2</sup>	
Calculation Type (Precipitation Calculator)		
Calculation type	TTT diagram	
Temperature	500° to 800° C with 25° C steps	
Max. annealing time	1.0E8 seconds	
Stop criteria	Volume fraction of phase is set to 0.0001	
Other		
File name and location	Folder: Precipitation Module - TC-PRISMA File name: P_03_Precipitation_Fe-C-Cr_TTT_Cementite- M7C3-M23C6.tcu	

# **Plot Results**



Equilibrium Calculator: How the phases change with temperature

Precipitation Calculator: TTT diagram



# **P\_04:** Precipitation of Iron Carbon Cementite

Simulate the kinetics of precipitation of carbides from a BCC Fe solution phase. This example is based on Wert (1949)<sup>1.</sup>.

This example uses two Experimental File Reader activities with the Precipitation Calculator and an isothermal calculation, to plot the volume fraction of the cementite phase.

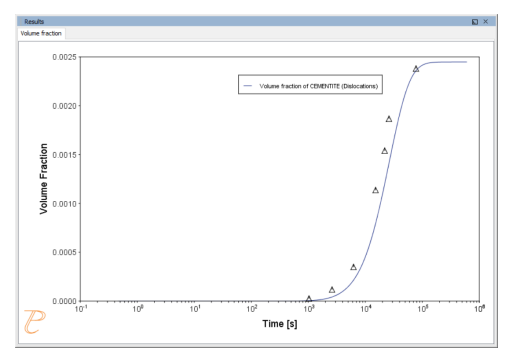
Demo: Steels and Fe-alloys (FEDEMO and MFEDEMO)		
Fe, C		
Fe-0.016C mass percent		
BCC_A2		
Cementite		
Matrix Phase Data Parameters (Precipitation Calculator)		
1.0		
1.5e11m <sup>-3</sup>		
Precipitate Phase Parameters (Precipitation Calculator)		
Dislocations		
0.24 J/m <sup>2</sup>		
Calculation Type (Precipitation Calculator)		
lsothermal		
102° C		

<sup>1.</sup> Wert, Charles A. 1949. "Precipitation from Solid Solutions of C and N in  $\alpha$ -Iron." Journal of Applied Physics 20 (10). AIP Publishing: 943.

Simulation time	600 000 seconds		
Options (Precipitation Calculator)			
Growth rate model	Advanced		
Other			
File name and location	Folder: Precipitation Module - TC-PRISMA File name: P_04_Precipitation_Fe-C_ Cemetite.tcu		

# **Plot Results**

# Volume Fraction



# $P\_05$ : Precipitation of $\gamma^\prime$ in Ni Superalloys - Isothermal

P

This example simulates the kinetics of precipitation of  $\gamma'$  phase from  $\gamma$  phase. The simulation results can be compared with experimental data collected from Sudbrack et al. (2008)<sup>1</sup>.

This example uses three Experimental File Reader activities with the Precipitation Calculator. It does an isothermal calculation to plot the volume fraction, mean radius and number density of the cementite phase.



DIS\_FCC\_A1 needs to be selected on the System Definer. Search the online help for **Selecting the Disordered Phase as a Matrix Phase** in the *Precipitation Module (TC-PRISMA) User Guide* for details.

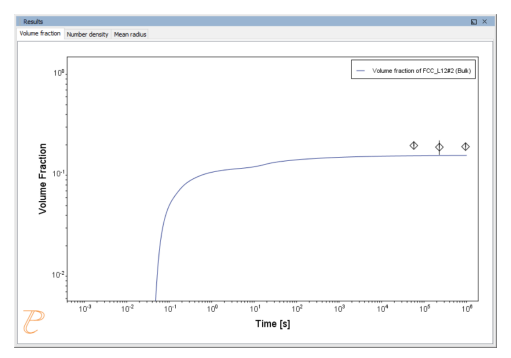
System (System Definer)		
Database package	Demo: Nickel-based Super Alloys (NIDEMO and MNIDEMO)	
Elements	Ni, Al Cr	
Conditions (Precipitation Calculator)		
Composition	Ni-9.8Al-8.3Cr Mole percent	
Matrix phase	DIS-FCC_A1 (see note above about how to select this phase)	
Precipitate phase	FCC_L12#2	
Precipitate Phase Data Parameters (Precipitation Calculator)		
Nucleation sites	Bulk	
Interfacial energy	0.012 J/m <sup>2</sup>	
Calculation Type (Precipitation Calculator)		
Calculation type	Isothermal	
Temperature	800° C	

<sup>1.</sup> Sudbrack, Chantal K., Tiffany D. Ziebell, Ronald D. Noebe, and David N. Seidman. 2008. "Effects of a Tungsten Addition on the Morphological Evolution, Spatial Correlations and Temporal Evolution of a Model Ni–Al–Cr Superalloy." Acta Materialia 56 (3): 448–63.

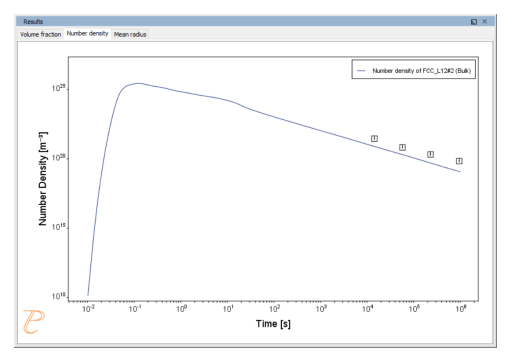
Simulation time	1 000 000 seconds
Other	
File name and location	Folder: Precipitation Module - TC-PRISMA File name: P_05_Precipitation_Ni-Al-Cr_Isothermal_Gamma-Gamma_ prime.tcu

# **Plot Results**

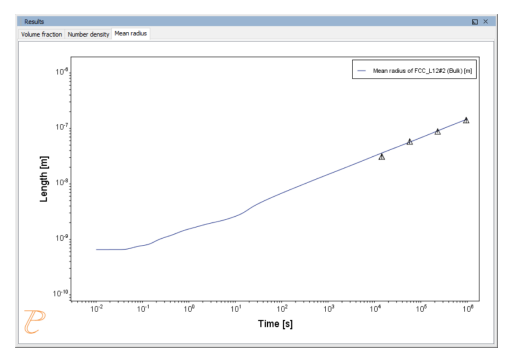
Volume Fraction



## Number Density



### Mean Radius



# P\_06: Precipitation of $\gamma'$ in Ni Superalloys - Non-isothermal

This example simulates the kinetics of precipitation of  $\gamma'$  phase from  $\gamma$  phase in Ni-8Al-8Cr and Ni-10Al-10Cr at.% alloys during continuous cooling. The simulation results can be compared with experimental results from Rojhirunsakool et al. (2013)<sup>1</sup>.



When you run (Perform) this example, it takes about 10 minutes for the calculations to complete.

In these examples a linear, continuously cooling condition is set up. This is done on the Precipitation Calculator settings window in the **Calculation Type** section:

- 1. Select a Non-Isothermal.
- 2. Select a Temperature Unit and Time Unit from the lists.
- Click Thermal Profile, where Time-Temperature profile points are entered to describe the stepwise heat treatment schedule with linear segments. Up to 999 rows of data points can be entered.

In these examples two rows of data points are needed for starting time 0s and temperature 1150° C, as well as end time 3300s and temperature 380° C.

4. Click the **Plot Renderer** node to select the **Separate multimodal PSD** check box. This plots the mean radius and size distributions of the two compositions.

Plotting the size distribution from the final simulation time of 3300 seconds, you can see there are several peaks, although these are not completely separated. As above, select the **Separate multimodal PSD** check box to separate the peaks.

- Adjust the Valley depth ratio setting to 0.05 to separate into two peaks as shown in the <u>Ni-10Al-10Cr plot example</u>. You can experiment with this setting to see how the size distribution evolves with time, for example, try entering several values as plot times 400 600 3300.

DIS\_FCC\_A1 needs to be selected on the System Definer. Search the online help for **Selecting the Disordered Phase as a Matrix Phase** in the *Precipitation Module (TC-PRISMA) User Guide* for details.

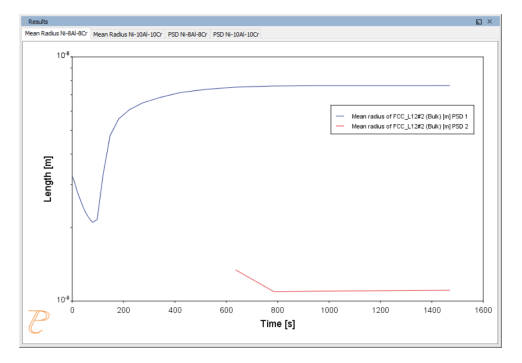
<sup>1.</sup> Rojhirunsakool, Tanaporn, S. Meher, J. Y. Hwang, S. Nag, J. Tiley, and Rajarshi Banerjee. 2013. "Influence of Composition on Monomodal versus Multimodal Γ' Precipitation in Ni–Al–Cr Alloys." Journal of Materials Science 48 (2): 825–31.

System (System Definer)			
Database package	Demo: Nickel-based Super Alloys (NIDEMO and MNIDEMO)		
Elements	Ni, Al, Cr		
Conditions (Precipitation Calculator)			
Composition (Ni-8Al-8Cr)	Ni-8Al-8Cr Mole percent		
Composition (Ni-10Al-10Cr)	Ni-10Al-10Cr Mole percent		
Matrix phase	DIS_FCC_A1 (see note above about how to select this phase)		
Precipitate phase	FCC_L12#2		
Matrix Phase Data Parameters (Prec	ipitation Calculator)		
Mobility enhancement prefactor (click <b>Show details</b> to display this setting)	5.0		
Precipitate Phase Data Parameters (Precipitation Calculator)			
Nucleation sites	Bulk		
Interfacial energy	0.023 J/m <sup>2</sup>		
Calculation Type (Precipitation Calculator)			
Calculation type	Non-isothermal		
Temperature unit	Celsius		
Time unit	Seconds		
Simulation time (Ni-8Al-8Cr)	1470 s		
Simulation time (Ni-10Al-10Cr)	3300 s		
Multimodal PSD (Plot Renderer)			
Separate multimodal PSD for 8AI-8Cr	The <b>Valley depth ratio</b> is set to 0.05 for both plots. The number of <b>Points</b> is increased to 200 for an average radius plot.		

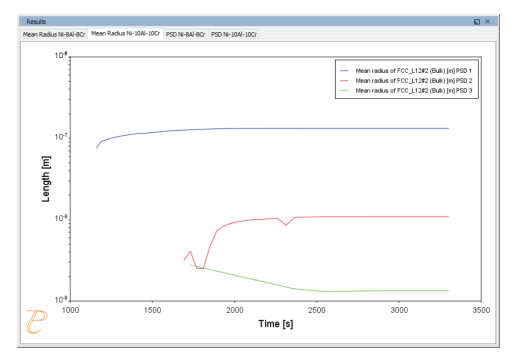
Separate multimodal PSD for 10Al- 10Cr	The <b>Valley depth ratio</b> is set to 0.18 for both plots.
Other	
File name and location	Folder: Precipitation Module - TC-PRISMA File name: P_06_Precipitation_Ni-Al-Cr_Non-isothermal_ Gamma-Gamma_prime.tcu

# **Plot Results**

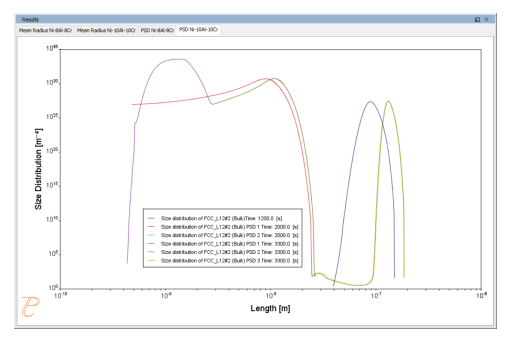
Mean Radius Ni-8Al-8Cr

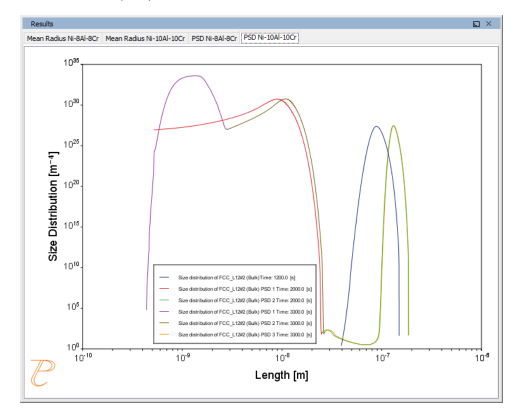


### Mean Radius Ni-10Al-10CR



## Size Distribution (PSD) Ni-8Al-8Cr





#### Size Distribution (PSD) Ni-10Al-10Cr

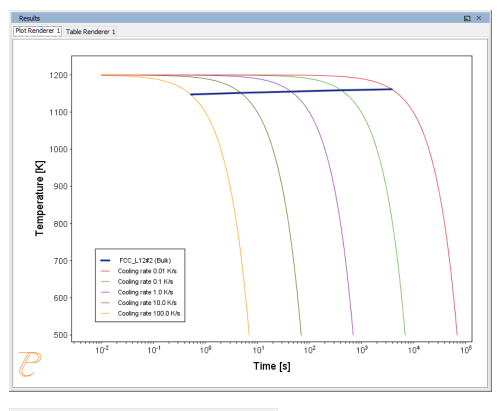
# P\_07: Cooling Rate (CCT) Diagram of Ni-Al-Cr γ-γ'

In this example, a CCT diagram for a Ni-10Al-10Cr  $\gamma$ - $\gamma'$  alloy is calculated and plotted with superimposition of the cooling rate values.

System (System Definer)		
Database package	Demo: Nickel-based Super Alloys (NIDEMO and MNIDEMO)	
Elements	Ni, Al, Cr	
Conditions (Precipitation Calculator)		
Composition	Ni-10Al-10Cr Mole percent	
Matrix phase	DIS_FCC_A1	

Precipitate phase	FCC_L12#2	
Precipitate Phase Data Parameters (Precipitation Calculator)		
Nucleation sites	Bulk	
Interfacial energy	0.023 J/m <sup>2</sup>	
Calculation Type (Precipitation Calculator)		
Calculation type	CCT Diagram	
Temperature Min to Max	500 to 1200 Kelvin	
Cooling rate(s)	.01 .1 1 10 100 K/s	
Stop criteria	Volume fraction of phase 1.0E-4	
Other		
File name and location	Folder: Precipitation Module - TC-PRISMA File name: P_07_Precipitation_Ni-Al-Cr_CCT_Gamma-Gamma_ prime.tcu	

### Plot and Table Results



Results			
Plot Renderer 1 Table Renderer 1			
Temperature [K]	FCC_L12#2 (Bulk)		
1147.52034	0.53291		
1151.74192	4.83553		
1155.24636	44.76534		
1158.85368	411.49173		
1161.48750	3851.47779		

# P\_08: Precipitation of Cu-Ti CU4TI1 with Assumptions of Sphere and Needle Morphologies

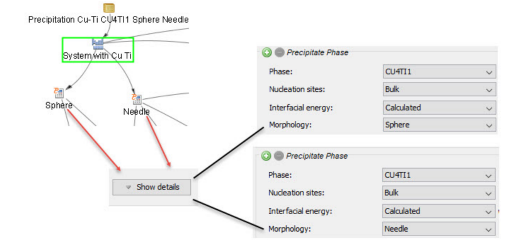
In this isothermal calculation example, the precipitation of Cu4Ti phase in a Cu-Ti binary alloy is calculated. To make a comparison, two separate simulations are performed, one assuming spherical morphology without elastic strain energy, and the other assuming needle

morphology whose shape, determined by competition between interfacial energy and elastic strain energy, is changed during the simulation. The transformation strain is obtained from Borchers<sup>1</sup>. The results are compared with experiment results from Kampmann et al<sup>2</sup>.

This example takes a few minutes to run.

To ensure that the settings are done on the correct Precipitation Calculators, the **Sphere** and **Needle** nodes are renamed from **Precipitation Calculator** to match their morphology. The morphology is set in the **Precipitate Phase** section when you click **Show details**.

Search the help for "precipitation morphology".



System (System Definer)		
Database package	Demo: Copper-based alloys (CUDEMO and MCUDEMO)	
Elements	Cu, Ti	
Sphere and Needle Conditions (Precipitation Calculator)		
Composition	Cu-1.9Ti Mole percent	
Matrix phase	FCC_L12	

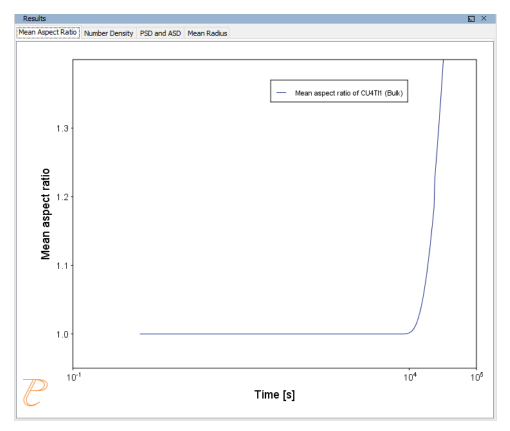
<sup>&</sup>lt;sup>1.</sup> Borchers C. 1999. "Catastrophic Nucleation during Decomposition of Cu-0.9at.%Ti." Phil. Mag. A 79(3):537-547

<sup>&</sup>lt;sup>2.</sup> Kampmann R., Eckerlebe H., and Wagner R. 1987. "Precipitation Kinetics in Metastab le Solid Solutions - Theoretical Considerations and Application to Cu-Ti Alloys." Mat. Res. Soc. Symp. Proc. 57: 525-542.

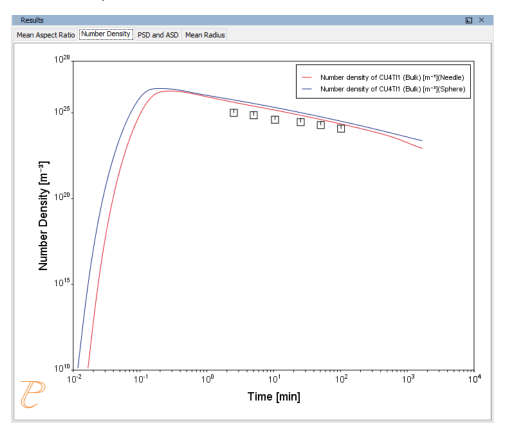
Precipitate phase	CU4TI1	
Matrix Phase Data Paramete	rs (Precipitation Calculator)	
Mobility enhancement prefactor (click <b>Show details</b> to display this setting)	100	
Precipitate Phase Data Parameters (Precipitation Calculator)		
Nucleation sites	Bulk	
Interfacial energy	The default	
Morphology (click <b>Show</b>	For the <b>Sphere</b> node (renamed from Precipitation Calculator), keep the default.	
<b>details</b> to display this setting)	For the <b>Needle</b> node (renamed from Precipitation Calculator), <b>Needle</b> is selected.	
	For the <b>Sphere</b> node (renamed from Precipitation Calculator), keep the default.	
Transformation strain (click <b>Show details</b> to display this	For the <b>Needle</b> node (renamed from Precipitation Calculator), <b>User</b> <b>defined</b> is selected. In this example, the following settings are defined:	
setting)	<ul> <li>ε11 and ε22 are set to 0.022</li> <li>ε33 is set to 0.003</li> </ul>	
Calculation Type (Precipitati	on Calculator)	
Calculation type	Isothermal	
Temperature	350° C	
Simulation time	10,000 seconds	
Datasets (Experimental File Reader)		
Borchers Mean radius vs Time and Borchers Number density vs Time	Data sets included with this example and imported to two Experimental File Readers. These data sets are used for the Mean Radius and Number Density plots, respectively.	
Other		
File name and location	Folder: Precipitation Module - TC-PRISMA File name: P_08_Precipitation_Cu-Ti_CU4TI1_Sphere_Needle.tcu	

### **Plot Results**

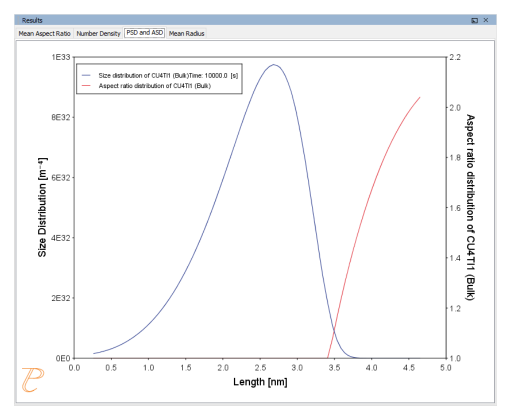
Mean Aspect Ratio



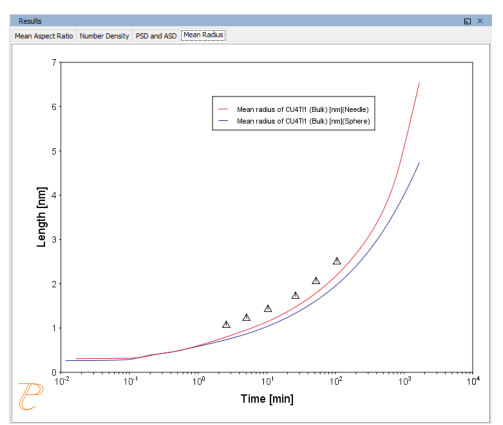
### Number Density



### PSD and ASD



#### Mean Radius



# P\_09: Precipitation of Al-Sc AL3SC with Assumption of Sphere and Cuboid Morphologies

In this isothermal calculation example, the precipitation of Al<sub>3</sub>Sc phase from FCC\_A1 matrix phase in an Al-Sc binary alloy is simulated. To make a comparison, two separate calculations are performed, one assuming spherical morphology without elastic strain energy, and the other assuming cuboid morphology whose shape is determined by competition between interfacial energy and elastic strain energy. The simulation results are compared with experimental data collected from Marquis and Seidman<sup>1.</sup> and Novotny and Ardell (2001)<sup>2.</sup>. In addition, mean cubic factor and cubic factor distribution are also plotted for cuboid shape to illustrate the spherical-cuboidal transition during precipitation.

<sup>&</sup>lt;sup>1.</sup> Marquis, E.A, and D.N Seidman. 2001. "Nanoscale Structural Evolution of Al3Sc Precipitates in Al(Sc) Alloys." Acta Materialia 49 (11): 1909–19.

<sup>&</sup>lt;sup>2.</sup> Novotny, Gabriel M., and Alan J. Ardell. 2001. "Precipitation of Al3Sc in Binary Al–Sc Alloys." Materials Science & Engineering, A: Structural Materials: Properties, Microstructure and Processing 318 (1–2): 144–54.

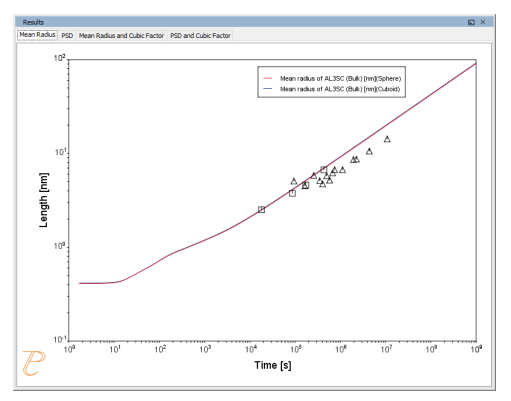
To ensure that the settings are done on the correct Precipitation Calculators, the **Sphere** and **Cuboid** nodes are renamed from **Precipitation Calculator** to match their morphology. The morphology is set in the **Precipitate Phase** section when you click **Show details**. See P\_08 for an example of this.

System (System Defi	ner)	
Database package	Demo: Aluminum-based alloys (ALDEMO, MALDEMO)	
Elements	Al, Sc	
Sphere and Cuboid C	Conditions (Precipitation Calculator)	
Composition	Al-0.18Sc Mole percent	
Matrix phase	FCC_A1	
Precipitate phase	AL3SC	
Matrix Phase Data P	arameters (Precipitation Calculator)	
Elastic properties (click <b>Show details</b> to display this setting)	<ul> <li>For the Sphere node (renamed from Precipitation Calculator), the default, Disregard is kept.</li> <li>For the Cuboid node (renamed from Precipitation Calculator), choose Cubic. Then enter the elastic constants accordingly. Default elastic constants are given based on the major element of the alloy system. In this example that is <ul> <li>c11 is 108.2 GPa</li> <li>c12 is 61.3 GPa</li> <li>c44 is 28.5 GPa</li> </ul> </li> </ul>	
Precipitate Phase Da	ta Parameters (Precipitation Calculator)	
Nucleation sites	Bulk	
Interfacial energy	The default	
Morphology (click <b>Show details</b> to display this setting)	For the <b>Sphere</b> node (renamed from Precipitation Calculator), keep the default. For the <b>Cuboid</b> node (renamed from Precipitation Calculator), <b>Cuboid</b> is selected.	

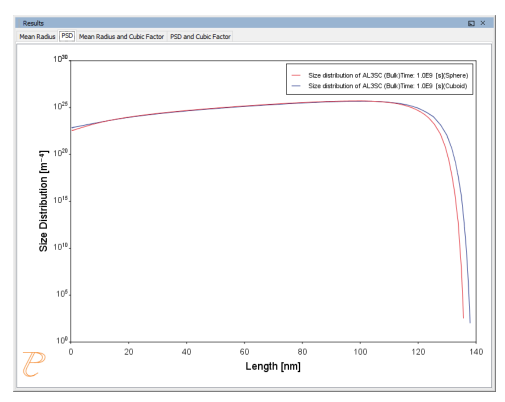
Transformation strain (click <b>Show</b> <b>details</b> to display this setting)	For the <b>Sphere</b> node (renamed from Precipitation Calculator), keep the default. For the <b>Cuboid</b> node (renamed from Precipitation Calculator), <b>Calculate from</b> <b>molar volume</b> is selected to obtain a purely dilatational strain.		
Calculation Type (Pro	ecipitation Calculator)		
Calculation type	Isothermal		
Temperature	350° C		
Simulation time	1.0E9 seconds		
Datasets (Experimen	Datasets (Experimental File Reader)		
Dataset 1 and Dataset 2	Data sets included with this example and imported to one Experimental File Reader. It is used for the Mean Radius plot.		
Other			
File name and location	Folder: Precipitation Module - TC-PRISMA File name: P_09_Precipitation_Al-Sc_AL3SC_Sphere_Cuboid.tcu		

### **Plot Results**

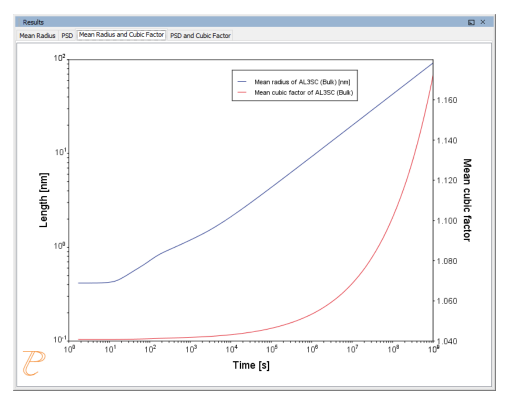
Mean Radius



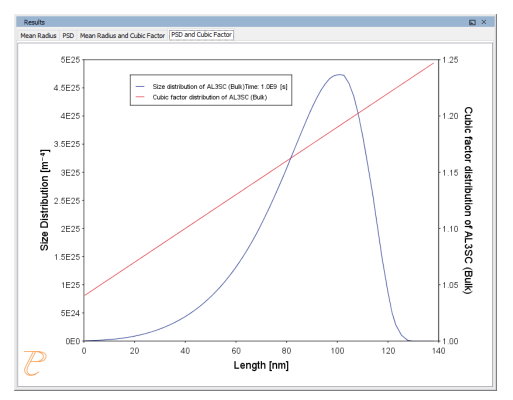
#### PSD



#### Mean Radius and Cubic Factor



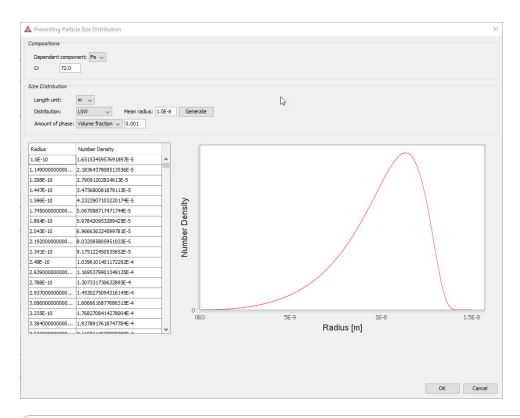
#### **PSD and Cubic Factor**



# P\_10: Initial Particle Size Distribution of Fe-Cr-C

This example demonstrates the effect of initial particle size distribution of the precipitate phases on the overall precipitation kinetics. It uses two Precipitation Calculators to simulate and compare carbide precipitations from a ferritic BCC\_A2 matrix in a Fe-0.1C-12Cr alloy. Three carbides, CEMENTITE, M23C6 and M7C3, are included in the calculations for competitive precipitations, and the precipitation kinetics are compared with or without initial particle size distribution.

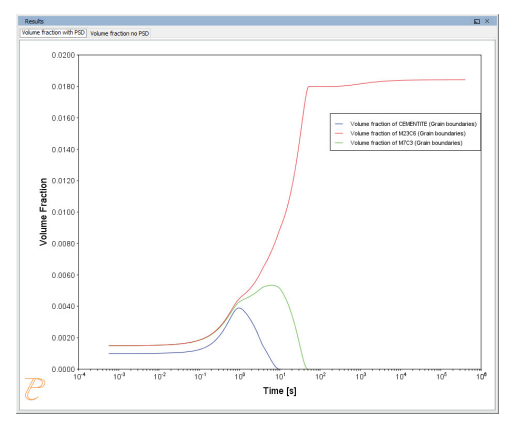
The example illustrates the use of the particle size distribution setting. You can import data from a spreadsheet or text file (.xls, .xlsx, .csv or .txt formats are acceptable). The Preexisting Particle Size Distribution window shown below, provides a graphical representation of the radius versus corresponding frequencies.



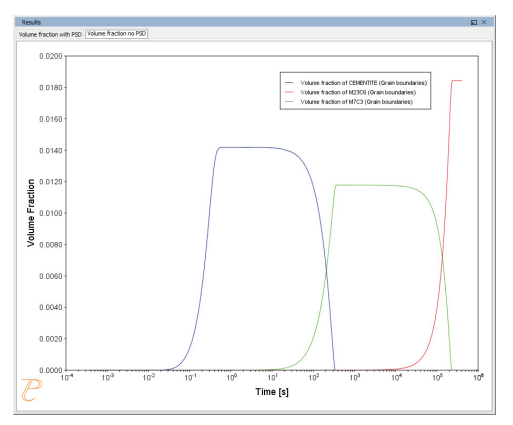
System (System Definer)		
Database package	Demo: Steels and Fe-alloys (FEDEMO and MFEDEMO)	
Elements	Fe, C, Cr	
Conditions (Precipitation Calculator)		
Composition	Fe-0.1C-12Cr Mass percent	
Matrix phase	BCC_A2	
	All other defaults are kept.	
Precipitate phases	CEMENTITE, M23C6 and M7C3	
Precipitate Phase Data Parameters (Precipitation Calculator)		
Nucleation sites	Grain boundaries (all calculations): Calculated from the matrix settings with a wetting angle of 90°	
	User-defined function f(r,T) (all calculations):	
Interfacial energy	• CEMENTITE: 0.167 J/m <sup>2</sup>	
	• M23C6 0.252 J/m <sup>2</sup>	

	• M7C3 0.282 J/m <sup>2</sup>	
Preexisting size distribution (click <b>Show</b> <b>details</b> to display this setting)	For the Precipitation Calculator including particle size distribution, and for all precipitate phases, this check box is selected. For each precipitate phase (CEMENTITE, M23C6 and M7C3), click <b>Edit particle size distribution</b> to make changes to the parameters. A window opens with a graphical representation of the radius vs number density.	
Calculation Type (Precipitation Calculator)		
Calculation type	Isothermal	
Temperature	1053 К	
Simulation time	400 000 seconds	
Other		
File name and location	Folder: Precipitation Module - TC-PRISMA File name: P_10_Precipitation_Initial_PSD_FeCrC.tcu	

### **Plot Results**



Volume Fraction with Initial Particle Size Distribution



Volume Fraction with No Initial Particle Size Distribution

### **P\_11: Interfacial Energy Function**

In some cases, interfacial energy may be a function of temperature or particle radius. This example uses four Precipitation Calculators at four temperature points in 30 K increments: 673 K, 703 K, 733 K, and 763 K. It is an isothermal calculation to examine the mean radius of an Al-0.12Sc system. It uses an FCC\_A1 matrix phase and AL3SC precipitate phase with bulk nucleation sites and user-defined interfacial energy function. The user defined interfacial energy function uses an error function to set a smooth transition of the interfacial energy from 0.065 J/m<sup>2</sup> to 0.085 J/m<sup>2</sup> for particle radii below and above 1e<sup>-8</sup>m and 5e<sup>-8</sup>m, respectively.

A dataset based on Iwamura and Miura (<sup>1.</sup>) data is compared with the calculated results.

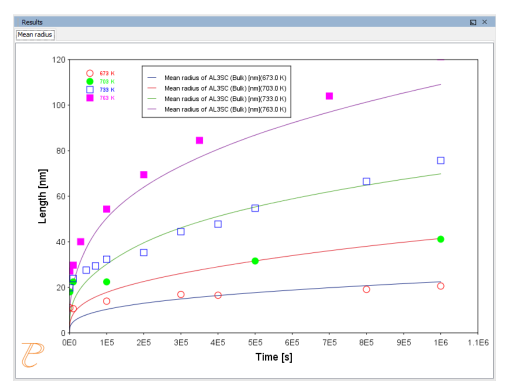
System (System Definer)	
Database package	Demo: Aluminum-based Alloys (ALDEMO, MALDEMO)

<sup>1.</sup> Iwamura, S, and Y Miura. 2004. "Loss in Coherency and Coarsening Behavior of Al3Sc Precipitates." Acta Materialia 52 (3): 591–600.

Al, Sc		
Conditions (Precipitation Calculator)		
Al-0.12Sc Mole percent		
FCC_A1		
All other defaults are kept.		
AL3SC		
Nucleation sites (all calculations): Bulk (6.025E28 m <sup>-3</sup> )		
Interfacial energy (all calculations): User-defined function f(r,T): 0.075+0.011*erf ((r-3e-8)/1e-8 J/m <sup>2</sup> )		
ecipitation Calculator)		
Isothermal (all calculations)		
Four temperature points in 30 K increments: 673 K, 703 K, 733 K, and 763 K.		
1 000 000 seconds (all calculations)		
ital File Reader)		
Data set included with this example and imported to one Experimental File Reader.		
Folder: Precipitation Module - TC-PRISMA		
File name: P_11_Interfacial_energy_function.tcu		

### **Plot Results**

### Mean Radius



# Thermo-Calc Console Mode Examples Guide

**Thermo-Calc Version 2018b** 





# **About the Thermo-Calc Console Mode Examples**

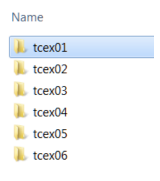
"About the Graphical Mode Examples" on page 2

There are many examples to examine and to help you learn Thermo-Calc in Console Mode.

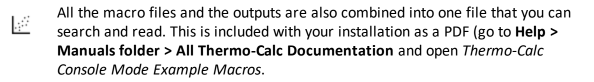
### **OPENING THE MACRO FILE**

To open an example macro file for Thermo-Calc:

- 1. Open Thermo-Calc.
- 2. Confirm you are in Console Mode. If not, click the **Switch to Console Mode** button.
- 3. From the main menu, select File → or Help → Examples Files. The Console Mode folder opens.
- 4. Open the **Thermo-Calc** folder to see the available examples contained in separate folders.



- For the Diffusion Module (DICTRA) examples (\*.DCM macro files), open the Diffusion Module - DICTRA folder.
  - 5. In the folder, double-click the \*.TCM file or click **Open**.
  - 6. Follow the prompts in the Console window i.e. keep pressing <Enter> to run the macro and produce a plot in the **Console Results** window.



Each topic below has a brief description of the example and includes some or all of the commands used in the macros. As you progress through the examples, sometimes only new commands are highlighted.

Торіс	TCEX Examples included
"Examples of Binary Systems" on the next page	1, 4, 10, 11, 13, 16, 18, 26, 34, 51
"Examples Using Functions and Variables" on page 18	2, 44
"Examples of the Gibbs Energy System (GES) Module" on page 20	31, 38, 39, 49, 52
"Examples of Multicomponent Systems" on page 16	6, 7, 47
"Example using the PARROT and EDIT_ EXPERIMENT Modules" on page 26	36a
"Pourbaix Module Examples" on page 33	40, 53
"Pourbaix Module Examples TCEX40A to 40E" on page 35	40A to 40E (additional examples related to TCEX40)
"Examples of Property and Potential Diagrams" on page 15	8, 35
"Examples of Scheil-Gulliver Calculations" on page 24	15, 30a, 30b, 48
"Examples with Specialized Databases" on page 30	20, 22, 25, 27, 29, 50,
"General Examples Using the Steel Database" on page 13	9, 23, 28, 32, 37, 41
"Example of the Tabulate Reaction (TAB) Module" on page 28	12
"Example of the Reactor Module" on page 29	24
"Examples of Ternary Systems" on page 9	3, 5, 14, 17, 19a, 19b, 21, 33, 42, 43, 45, 46,

# **Examples of Binary Systems**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

### TCEX01 - CALCULATING THE BINARY FE-C PHASE DIAGRAM

This example calculates the binary Fe-C phase diagram using the Binary module. The TCBIN database is used and included with your installation.

Some of the main commands used:

- "BINARY\_DIAGRAM Module" on page 10
- "LIST\_EQUILIBRIUM" on page 198
- <u>"LIST\_STATUS" on page 200</u>
- "LIST\_CONDITIONS" on page 198
- <u>"CHANGE\_STATUS" on page 177</u>
- <u>"COMPUTE\_EQUILIBRIUM"</u> on page 182
- "SET\_DIAGRAM\_AXIS" on page 249
- <u>"SET\_TIELINE\_STATUS" on page 261</u>
- "PLOT\_DIAGRAM" on page 244
- <u>"SET\_SCALING\_STATUS" on page 260</u>
- "SET\_LABEL\_CURVE\_OPTION" on page 253
- "ADD\_LABEL\_TEXT" on page 234
- <u>"MODIFY\_LABEL\_TEXT" on page 243</u>

### TCEX04 - CALCULATING THE FE-CR MISCIBILITY GAP

This example calculates the miscibility gap in the Fe-Cr system. The FEDEMO database is used and included with your installation.

These commands are used:

"SWITCH_DATABASE" on page 22	"DEFINE_SYSTEM" on page 15	"REJECT" on page 18
"GET_DATA" on page 16	"AMEND_PHASE_DESCRIPTION" on page 89	"LIST_STATUS" on page 200
"LIST_EQUILIBRIUM" on page 198	"SET_CONDITION" on page 209	"COMPUTE_EQUILIBRIUM" on page 182
"SET_AXIS_VARIABLE" on page 208	"SAVE_WORKSPACES " on page 52	"SET_TIELINE_STATUS" on page 261

### TCEX10 - PREVENTING CR2O3 CLOGGING IN A CONTINUOUS CASTING PROCESS

The example calculates an equilibrium with suspended or dormant phases and shows how to avoid  $Cr_2O_3$  clogging in a continuous casting process.

This uses the SLAG database. A license is required to run the example.

The origin of this example is that a manufacturer wanted to increase the Cr content of a material from 18 to 25 weight percent but the continuous casting of this material was problematic because solid  $Cr_2O_3$  was formed. Using Thermo-Calc to calculate the equilibria in the steel/slag system, a simple correction was found: modify the Mn or Si content to decrease the oxygen potential.

In Thermo-Calc, you can FIX a phase with zero amount to simulate how to avoid forming this phase. You then release one of the conditions, usually one of the compositions, and this composition is determined by the equilibrium calculation.

These commands are used:

"DEFINE_SYSTEM" on	"AMEND_PHASE_DESCRIPTION"	"GET DATA" on page 16
page 15	on page 89	GET_DATA OF page 10
"SET_CONDITION" on page 209	"LIST_CONDITIONS" on page 198	"LIST_STATUS" on page 200
"CHANGE_STATUS" on	"COMPUTE_EQUILIBRIUM" on	"SET_NUMERICAL_LIMITS" on
page 177	page 182	page 293
"LIST_EQUILIBRIUM" on	"SET_REFERENCE_STATE" on	"SHOW_VALUE" on page 217
page 198	page 258	SHOW_VALUE ON page 217

### TCEX11 - OXIDATION OF CU2S WITH AN H2O/O2 GAS MIXTURE

This example demonstrates the oxidation of  $Cu_2S$  with an  $H_2O/O_2$  gas mixture. Thermo-Calc is used to find the optimum O/H ratio (i.e. oxygen potential) as certain oxygen potential values can desulphurize  $Cu_2S$  without forming copper oxides.

In Thermo-Calc, the problem reduces to perform equilibria calculations in a Cu-S-H-O system. The amounts of the components should be kept to correct ratio corresponding to  $Cu_2S$  and H2O using a command SET\_INPUT\_AMOUNTS in POLY3. Initially, O/H = 0.5 is given. Optimum O/H ratio is calculated with the desired calculation conditions. For example, to simulate one phase disappearing, you can FIX the phase with zero amount.

The *tcex11.TDB* file is used to access data and this is included with your installation.

These commands are of also of note and used in this example:

- "SET\_NUMERICAL\_LIMITS" on page 293
- "SET\_REFERENCE\_STATE" on page 258
- <u>"SHOW\_VALUE" on page 217</u>

### TCEX13 - THE BINARY AL-TI PHASE DIAGRAM AND ITS G CURVE

This example calculates an Al-Ti binary phase diagram and G curve using the BINARY module. The TCBIN database is used and included with your installation.

These commands are of note and used in this example:

- "BINARY\_DIAGRAM Module" on page 10
- "SET\_ALL\_START\_VALUES" on page 206

#### TCEX16 - SECOND ORDER TRANSITION OF THE AL-FE SYSTEM

This example calculates the second order transition line in the Bcc field of the Al-Fe system.



This uses the SSOL database. A license is required to run the example.

These commands are of note and used in this example:

- "REJECT" on page 18
- "RESTORE" on page 20
- "ADVANCED\_OPTIONS" on page 167
- "SET\_SCALING\_STATUS" on page 260
- "MAKE\_EXPERIMENTAL\_DATAFILE" on page 243
- "LIST\_EQUILIBRIUM" on page 198
- "APPEND\_EXPERIMENTAL\_DATA" on page 236

### • <u>"STEP\_WITH\_OPTIONS" on page 217</u>

### TCEX18 - A3 TEMPERATURE OF A STEEL

This example calculates the A3 temperature of a steel and examines the influence of each alloying element on this temperature.

A3 temperature is the temperature where ferrite starts to form from austenite. You can easily read A3 from an Fe-C phase diagram. However, for complex multicomponent steels no simple diagram can be used. Using POLY, it is easy to find out the influence of each alloying element on the A3 temperature. This information is useful if you want to modify the compositions of a steel but keep A3 unchanged.



This uses the TCFE database. A license is required to run the example.

These commands are of note and used in this example:

- <u>"DEFINE\_ELEMENTS" on page 14</u>
- "AMEND\_PHASE\_DESCRIPTION" on page 89
- "COMPUTE\_TRANSITION" on page 183
- "DEFINE\_MATERIAL" on page 189
- "ENTER\_SYMBOL" on page 238
- "SHOW\_VALUE" on page 217

#### TCEX26 - THE AS-GA PHASE DIAGRAM: PLOTTING THE PARTIAL PRESSURES OF A GAS SPECIES

This is an example of plotting the partial pressures of a gas species along the solubility lines in the As-Ga phase diagram. The calculation makes it possible to monitor the input gases to a process of depositing solid As-Ga.

These commands are of note and used in this example:

- "REJECT" on page 18
- "RESTORE" on page 20
- "SET\_DIAGRAM\_AXIS" on page 249
- <u>"SET\_SCALING\_STATUS" on page 260</u>
- "SET\_AXIS\_TEXT\_STATUS" on page 247
- "SET\_LABEL\_CURVE\_OPTION" on page 253

### TCEX34 - THE AL-ZN PHASE DIAGRAM AND ITS G CURVE

This example uses the BINARY module to calculate the phase diagram and G curves in the Al-Zn system. It uses the TCBIN database included with your installation.

### TCEX51 - CALCULATION OF MOLAR VOLUME, THERMAL EXPANSIVITY AND DENSITY

This example uses the POLY3 module to calculate the molar volume, thermal expansivity and density of the FCC\_A1, BCC\_A2 LIQUID and liquid phases of C-Fe.



This uses the TCFE database. A license is required to run the example.

# **Examples of Ternary Systems**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



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### TCEX03 - CALCULATING AN ISOTHERMAL SECTION WITH THE TERNARY MODULE

This example calculates an isothermal section using the Ternary module. The FEDEMO database is used and included with your installation.

These commands are used:

- "TERNARY\_DIAGRAM Module" on page 326
- "AMEND\_PHASE\_DESCRIPTION" on page 89
- "ADD\_LABEL\_TEXT" on page 234
- "PLOT\_DIAGRAM" on page 244
- <u>"SET\_DIAGRAM\_AXIS" on page 249;</u> <u>"SET\_DIAGRAM\_TYPE" on page 252;</u> and <u>"SET\_</u> <u>AXIS\_TYPE" on page 247</u>
- <u>"SET\_REFERENCE\_STATE"</u> on page 258
- "CHANGE\_STATUS" on page 177
- <u>"SET\_SCALING\_STATUS" on page 260</u>

### TCEX05 - CALCULATING A VERTICAL SECTION IN THE AL-CU-SI SYSTEM

This example calculates a vertical section in the Al-Cu-Si system and of a vertical section from Al to 10% Cu<sub>2</sub>Si. The ALDEMO database is used and included with your installation.

In addition to various commands used in the previous examples, the LIST\_AXIS\_VARIABLE command is used.

#### TCEX14 - VARIATIONS IN SOLIDIFICATION OF AN AL-MG-SI ALLOY

This example calculates the heat and heat capacity variations during solidification of an Al-

Mg-Si alloy. The ALDEMO database is used and included with your installation.

These commands are of note and used in this example:

- "LIST\_SYSTEM" on page 17
- <u>"LIST\_CONDITIONS" on page 198</u>
- "SET\_AXIS\_TEXT\_STATUS" on page 247

### TCEX17 - THE PSEUDO-BINARY SYSTEM - CAO-SIO2

This example calculates the pseudo-binary system CaO-SiO<sub>2</sub> using the OXDEMO database, which is included with your installation.

The oxide demo (Oxdemo) database can be used both for pseudo-binary systems like the one in this case, CaO-SiO<sub>2</sub>, or for full ternary systems like Ca-Fe-O.

These commands are of note and used in this example:

- "LIST\_SYSTEM" on page 17
- "REJECT" on page 18 and "RESTORE" on page 20
- "AMEND\_PHASE\_DESCRIPTION" on page 89
- "LIST\_STATUS" on page 200
- "DEFINE\_COMPONENTS" on page 186
- "SET\_CONDITION" on page 209
- "COMPUTE\_EQUILIBRIUM" on page 182 and "LIST\_EQUILIBRIUM" on page 198

TCEX19A AND TCEX19B - MAPPING OF UNIVARIANT EQUILIBRIA WITH THE LIQUID IN AL-CU-SI

- Part A. Step-by-step calculation using the POLY-3 module.
- Part B. Using the TERNARY module, you can get the information on invariant reactions, such as temperature and compositions.

These commands are of note and used in these examples:

- <u>"DEFINE\_SYSTEM" on page 15</u>
- <u>"DEFINE\_SYSTEM" on page 15</u>
- <u>"GET\_DATA" on page 16</u>
- "AMEND\_PHASE\_DESCRIPTION" on page 89

- "SET\_CONDITION" on page 209 and "LIST\_CONDITIONS" on page 198
- "ADVANCED\_OPTIONS" on page 167
- "SAVE\_WORKSPACES " on page 52 and "READ\_WORKSPACES" on page 203
- "APPEND\_EXPERIMENTAL\_DATA" on page 236
- "MAKE\_EXPERIMENTAL\_DATAFILE" on page 243

### TCEX21 - CALCULATING A TERNARY ISOTHERM IN FE-CR-NI

This example calculates a ternary isotherm in Fe-Cr-Ni with a user-defined database included with the example in the file named *tcex21.TDB*.

These commands are of note and used in this example:

- <u>"DEFINE\_SYSTEM" on page 15</u>
- <u>"GET\_DATA" on page 16</u>
- "SET\_CONDITION" on page 209
- "SET\_DIAGRAM\_TYPE" on page 252

### TCEX33 - AN FE-CR-C ISOPLETH

This example is a benchmark calculation of an isopleth in the Fe-Cr-C system. The FEDEMO database is used and included with your installation.

#### TCEX42 - PARAEQUILIBRIUM CALCULATION - FORMATION OF PARA-PEARLITE - ISOPLETH

(This is the Drop-down text)

This example uses an Fe-Mn-C system at 2.5%Mn Mass u-fraction to show a paraequilibrium calculation where there is formation of para-pearlite. It is an isopleth calculation and shows the <u>STEP\_WITH\_OPTIONS</u> command using the Paraequilibrium option.





This uses the TCFE database. A license is required to run the example.

### TCEX43 - PARAEQUILIBRIUM CALCULATION - FORMATION OF PARA-PEARLITE - ISOTHERMAL

This example uses an Fe-Mn-C system at 700 C to show a paraequilibrium calculation where there is formation of para-pearlite. It is an isothermal calculation and shows the STEP\_

WITH\_OPTIONS command using the Paraequilibrium option.



This uses the TCFE database. A license is required to run the example.

"About Paraequilibrium" on page 207 in the Thermo-Calc User Guide.

### TCEX45 - 3D DIAGRAM WITH THE GAMMA VOLUME IN THE FE-CR-C SYSTEM

This example calculates the gamma volume in the Fe-Cr-C system and plots it in 3D. The FEDEMO database is used and included with your installation.



To view the generated file, *tcex45.wrl*, install a WRML (Wirtual Reality Modelling Language) viewer to the web browser. WRML viewers can be downloaded from various sites, e.g. www.parallelgraphics.com or www.sim.no.

The <u>"ADVANCED\_OPTIONS" on page 167</u> (present phase option) command is of note and used in this example.

### TCEX46 - 3D DIAGRAM WITH THE LIQUIDUS SURFACE OF THE FE-CR-C SYSTEM

This example calculates the liquidus surface of the Fe-Cr-C system and plots it in 3D.

The <u>"ADVANCED\_OPTIONS" on page 167</u> (present phase option) command is of note and used in this example.

To view the generated files, *tcex46\_tri.wrl* and *tcex46\_sqrt*, install a WRML (Wirtual Reality Modelling Language) viewer to the web browser. WRML viewers can be downloaded from various sites, e.g. <u>www.parallelgraphics.com</u> or <u>www.sim.no</u>.



This uses the TCFE database. A license is required to run the example.

# **General Examples Using the Steel Database**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

### TCEX09 - CALCULATING A DEW POINT WITH THE POLY3 MODULE

This is an example calculating the dew point in the POLY3 module. The SUBDEMO database is used and included with your installation.

The <u>"STEP\_WITH\_OPTIONS" on page 217</u> and <u>"MAP" on page 201</u> commands are of note and used in this example.

### TCEX23 - CALCULATING A PARAEQUILIBRIUM AND THE TO TEMPERATURE IN A LOW ALLOYED STEEL

This example calculates a paraequilibrium and the T0 temperature in a low alloyed steel.



This uses the TCFE database. A license is required to run the example.

### TCEX28 - PITTING RESISTANCE EQUIVALENCE (PRE) FOR A DUPLEX STAINLESS STEEL

This example calculates the Pitting Resistance Equivalence (PRE) for a duplex stainless steel.



This uses the TCFE database. A license is required to run the example.

These commands are of note and used in this example:

- "REJECT" on page 18
- "RESTORE" on page 20
- <u>"DEFINE\_ELEMENTS" on page 14</u>
- "AMEND\_PHASE\_DESCRIPTION" on page 89
- "EVALUATE\_FUNCTIONS" on page 197
- "SET\_START\_CONSTITUTION" on page 216

### • <u>"COMPUTE\_TRANSITION" on page 183</u>

### TCEX32 - CALCULATING OXIDE LAYERS ON STEEL

This example calculates oxide layers on a steel and shows how to append databases.



This uses the TCFE and SSUB databases. These licenses are required to run the example.

These commands are of note and used in this example:

- "SET\_START\_CONSTITUTION" on page 216
- "SET\_REFERENCE\_STATE" on page 258
- <u>"APPEND\_DATABASE" on page 12</u>

### TCEX37 - CALCULATING AN ISOTHERMAL SECTION IN CONSOLE MODE

This example shows how to use command lines to enter and calculate an isothermal section using the POLY3 module. The FEDEMO database is used and included with your installation.

#### TCEX41 - CALCULATING A SOLUBILITY PRODUCT

Calculation of a solubility product. Shows the <u>STEP\_WITH\_OPTIONS</u> command using the T-zero option.



This uses the TCFE database. A license is required to run the example.

These commands are of note and used in this example:

- "DEFINE\_ELEMENTS" on page 14
- "REINITIATE\_PLOT\_SETTINGS" on page 245
- "SET\_DIAGRAM\_AXIS" on page 249

### **Examples of Property and Potential Diagrams**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

### **Property Diagrams**

### TCEX08 - PROPERTY DIAGRAMS FOR A HIGH SPEED STEEL

This example shows how to calculate property diagrams for a high speed steel i.e. phase fraction plots, activity vs temperature, and so forth.



This uses the TCFE database. A license is required to run the example.

These commands are of note and used in this example:

- "DEFINE\_DIAGRAM" on page 187
- <u>"DEFINE\_ELEMENTS" on page 14</u>
- "AMEND\_PHASE\_DESCRIPTION" on page 89
- "REINITIATE\_PLOT\_SETTINGS" on page 245

### **Potential Diagram**

### TCEX35 - CALCULATING A POTENTIAL DIAGRAM

"About Potential Diagrams" on page 211 in the Thermo-Calc User Guide.

This example uses the POTENTIAL module to calculate a potential diagram. Note that there are no commands for this module, you just follow the prompts. The SUBDEMO database is used and included with your installation.

### **Examples of Multicomponent Systems**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

### TCEX06 - AN ISOPLETH IN LOW ALLOYED FE-MN-SI-CR-NI-C STEEL

This example calculates a multicomponent phase diagram using the <u>DEFINE\_MATERIAL</u> command in POLY and the TCFE steel database. The material contains 1.5 %Cr + 0.4 %Mn + 3.5 %Ni + 0.3 %Si and 1 %C (by weight). These conditions and the temperature are set by the DEFINE\_MATERIAL command.

After the first equilibrium results, a phase diagram is calculated with one axis variable as temperature and the other as the carbon content.



This uses the TCFE database. A license is required to run the example.

### TCEX07 - SINGLE EQUILIBRIA IN LOW ALLOYED FE-MN-SI-CR-NI-C STEEL

This example shows how to calculate a single equilibria in a low alloyed Fe-Mn-Si-Cr-Ni-C steel. There are two common ways to perform a single equilibrium calculation:

- Get data from a database, then in POLY use and COMPUTE\_EQUILIBRIUM.
- Go directly to POLY and use DEFINE\_MATERIAL.

The <u>COMPUTE\_TRANSITION</u> command is also used to determine the temperature or composition where one phase forms or disappears. It is the same as the CHANGE\_STATUS  $\rightarrow$  SET\_CONDITION $\rightarrow$  COMPUTE\_EQUILIBRIUM sequence of commands.

A <u>"STEP\_WITH\_OPTIONS" on page 217</u> command using the option *normal* is also used in addition to the other commands that should be becoming more familiar as you work through the examples.



This uses the TCFE database. A license is required to run the example.

### TCEX47 - 3D-QUATERNARY DIAGRAM WITH THE GAMMA VOLUME IN THE FE-CR-NI-C SYSTEM AT

### 1373K

The example calculates the gamma volume in the Fe-Cr-Ni-C system at 1373K and plots a 3D-quaternary diagram. The FEDEMO database is used and included with your installation.



To view the generated file, *tcex47.wrl*, install a WRML (Wirtual Reality Modelling Language) viewer to the web browser. WRML viewers can be downloaded from various sites, e.g. <u>www.parallelgraphics.com</u> or <u>www.sim.no</u>.

# **Examples Using Functions and Variables**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

### TCEX02 - PLOTTING THERMODYNAMIC FUNCTIONS

This example shows how to plot thermodynamic functions in unary, binary and ternary systems. It also involves working with partial derivatives and partial quantities.



This uses the SSOL database. A license is required to run the example.

These commands are used:

"GOTO_MODULE" on page 4	"GET_DATA" on page 16	<u>"SET_CONDITION" on</u> page 209
"SWITCH_DATABASE" on page 22	"AMEND_PHASE_ DESCRIPTION" on page 89	"LIST_EQUILIBRIUM" on page 198
"DEFINE_SYSTEM" on page 15	"COMPUTE_EQUILIBRIUM" on page 182	"SET_AXIS_VARIABLE" on page 208
"SAVE_WORKSPACES " on page 52	"STEP_WITH_OPTIONS" on page 217	"SET_DIAGRAM_AXIS" on page 249
"ENTER_SYMBOL" on page 238	"PLOT_DIAGRAM" on page 244	"SET_LABEL_CURVE_ OPTION" on page 253
"REJECT" on page 18	"RESTORE" on page 20	"LIST_SYSTEM" on page 17
"LIST_STATUS" on page 200	"CHANGE_STATUS" on page 177	"SHOW_VALUE" on page 217

"SET_INTERACTIVE" on page 292	
page 202	

#### TCEX44 - EXPLORING VARIABLES AND FUNCTIONS

This example uses variables and functions to predict properties e.g. proof strength for an austenitic stainless steel at elevated temperatures (20-550° C). The example was created using an expression from Eliasson and Sandström (2000)<sup>1</sup>.



This uses the TCFE database. A license is required to run the example.

<sup>1. &</sup>quot;Proof strength values for austenitic stainless steels at elevated temperatures", *Steel Res.*, 71(6–7), 249–254.

# **Examples of the Gibbs Energy System (GES) Module**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

### TCEX31 - USING THE GES MODULE TO CALCULATE CVM

Using the Gibbs energy system (GES) module, this example calculates the CVM and compares it with the sublattices of a fictitious A B system. You also learn how to overlay diagrams from two calculations.

These <u>GES commands</u> are of note and used in this example:

- <u>"ENTER\_ELEMENT" on page 96</u>
- <u>"ENTER\_SPECIES" on page 108</u>
- <u>"ENTER\_SYMBOL" on page 109</u>
- <u>"ENTER\_PHASE" on page 106</u>
- "ENTER\_PARAMETER" on page 137
- "LIST\_PHASE\_DATA (from GIBBS)" on page 142

### TCEX38 - CALCULATING THE MORRAL "ROSE"

This example uses the Gibbs energy system (GES) module to calculate the Morral rose, which are miscibility gaps.

These commands are of note and used in this example:

- "REINITIATE" on page 121
- <u>"ENTER\_ELEMENT" on page 96</u>
- <u>"AMEND\_ELEMENT\_DATA" on page 78</u>
- "ENTER\_PARAMETER" on page 137
- <u>"ENTER\_PHASE" on page 106</u>

#### TCEX39 - CALCULATING REVERSIBLE CARNOT CYCLES OF A HEAT ENGINE

This example shows how to calculate the reversible Carnot cycle of a heat engine using one mole of an ideal gas with two fictitious species A and A2. The GES, POLY3 and POST modules are used, as well as the SUBDEMO database, which is included with your installation.

One application of the Second Law is to the efficiencies of heat engines, pumps and refrigerators. Whenever there is a difference of temperature, work can be produced - the principle of heat engines. The Gibbs energy also enables the prediction of the maximum work that a process may achieve. The goal of this example is to help relate the results to different thermodynamic quantities.

These commands are of note and used in this example:

- "REINITIATE" on page 121
- <u>"ENTER\_PARAMETER" on page 137</u>
- <u>"ENTER\_SPECIES" on page 108</u>
- "SET\_PLOT\_FORMAT" on page 254
- <u>"ENTER\_PHASE" on page 106</u>

#### TCEX49 - QUASICHEMICAL MODEL USING THE GIBBS ENERGY SYSTEM (GES) MODULE

This example shows how to enter parameters for a FACT quasichemical liquid model and how to calculate the sulfur activity using the GIBBS\_ENERGY\_SYSTEM (GES) module commands.

#### Solution Models in the GIBBS Module" on page 82

These commands are of note and used in this example:

- "ENTER\_ELEMENT" on page 96
- <u>"ENTER\_SYMBOL" on page 109</u>
- "AMEND\_ELEMENT\_DATA" on page 78
- <u>"ENTER\_PARAMETER" on page 137</u>
- <u>"ENTER\_SPECIES" on page 108</u>
- "SET\_REFERENCE\_STATE" on page 258
- **<u>GIBBS\_ENERGY\_SYSTEM</u>**
- "SHOW\_VALUE" on page 217

#### TCEX52 - CHANGING THE EXCESS MODELS FOR INTERACTION PARAMETERS IN A SOLUTION PHASE

Solution Models in the GIBBS Module" on page 82

This example shows how to change the excess models for binary and ternary interactions in a solution phase, either

- through direct interactive amendments of phase descriptions within the GIBBS\_ ENERGY\_SYSTEM (GES) module, or
- enforced by specific type-definitions given in a database file retrieved by the DATABASE\_RETRIEVAL module (also called the TDB module).

For binary excess models, the example changes it from the default *R*-*K* model to a Mixed-Excess-Model. The phase has to be a substitutional phase in this case.

For ternary extrapolation models, the example changes it from the default *R*-*K*-*M* model to a *Toop\_Kohler model*.

### Notes related to the example

The following is included in the macro file as part of the instructions but may be of use to new users:

#### Amending the binary excess model

The default binary excess model is the Redlich-Kister Model for all the three associated binary interaction pairs (A-B, A-C and B-C) in the substitutional LIQUID solution phase (without sublattice) that consists of three elements (A, B and C).

Before changing this default binary excess model for the ternary LIQUID solution phase, one must have already entered the G parameters (for standard Gibbs energies of all pure end-members) and L parameters (for binary R-K excess interaction energies), as shown in the example.

In this particular example, we want to change from the default R-K binary excess model to the Mixed-Excess-Model (with three different binary excess models, namely Legendre, Polynom and Redlich-Kister models, applied to the A-B, A-C and B-C binaries, respectively), as demonstrated below:

For the A-B interaction, the Legendre binary excess model should be used (rather than the default Redlich-Kister Model), with the first species (i.e. A) as the independent constituent and the second species (i.e. B) as the dependent constituent, while the L parameters for the A-B interaction shall remain the same as those handled by the R-K model.

For the A-C interaction, the Polynom binary excess model should be used (rather than the default Redlich-Kister Model), with the second species (i.e. C) as the independent constituent and the first species (i.e. A) as the dependent constituent, while the L parameters for the A-C interaction shall remain the same as those handled by the R-K model.

#### Amending the ternary extrapolation model

The default ternary excess model is the Redlich-Kister\_Muggianu Model (i.e., the MUGGIANU\_RESTOR method for ternary extrapolation based on binary parameters) for the associated ternary interaction terms; when no ternary L parameter is entered for that, such a default Redlich-Kister\_Muggianu Model is thus to be used for extrapolation from binary excess energies to ternary interactions in the substitutional LIQUID solution phase (without sublattice) that consists of three elements (A, B and C).

However, in this particular example as illustrated in the following, we shall change from this default R-K-M ternary excess model to the TOOP-KOHLER method for the ternary extrapolation method, with the species C as the Toop constituent, while the species A and B as the Kohler constituents (entering A and B, or B and A, as the basis constituent and first interacting constituent). This will implicitly enforce that, during the ternary extrapolation, only the A-B binary interaction parameters are utilized in accordance with the Kohler ternary extrapolation formula for A-B-C ternary interaction, while any other binary interaction parameters used in line with the Toop-Kohler ternary extrapolation formula (for the A-C-B and B-C-A ternary interactions). This makes the extrapolated ternary excess interaction terms different from those handled either by the default MUGGIANU\_RESTOR method or by the alternative KOHLER-ALL method.

Note that only when all the relevant binary excess energies in a ternary system are treated by the default Redlich-Kister Model (i.e., the Mixed-Excess-Model should have not been used), the MUGGIANU\_RESTOR method for ternary extrapolations is equivalent to the Redlich-Kister\_Muggianu Model, or the KOHLER-ALL method to the Redlich-Kister\_Kohler Model.

## **Examples of Scheil-Gulliver Calculations**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

#### TCEX15 - SIMULATING THE SOLIDIFICATION OF A CR-NI ALLOY WITH THE SCHEIL MODULE

This is an example of a solidification simulation of a Cr-Ni alloy. There is no back diffusion in the solid, i.e. Scheil-Gulliver model is used.



This uses the TCFE database. A license is required to run the example.

These commands are of note and used in this example:

- <u>"START\_WIZARD" on page 294</u>
- <u>"COMPUTE\_TRANSITION" on page 183</u>
- "ADD\_INITIAL\_EQUILIBRIUM" on page 160
- <u>"ENTER\_SYMBOL" on page 238</u>
- "ADVANCED\_OPTIONS" on page 167
- "STEP\_WITH\_OPTIONS" on page 217
- "MAKE\_EXPERIMENTAL\_DATAFILE" on page 243
- "CHANGE\_STATUS" on page 177

#### TCEX30A AND 30B - SCHEIL CALCULATION FOR AN AL-4MG-2SI-2CU ALLOY

<u>"About Equilibrium Calculations" on page 182</u> and <u>"About Scheil Simulations" on page 197</u> in the *Thermo-Calc User Guide*.

These examples show two ways to do a Scheil calculation for an Al-4Mg-2Si-2Cu alloy. In part A, you use the POLY3 module and the <u>STEP\_WITH\_OPTIONS</u> command with an EVALUATE option. Then in part B you use the SCHEIL module commands to do the same thing. Data from the COST2 database is used and included with the example in the file named *tcex30\_cost2.TDB*.

#### TCEX48 - SCHEIL SOLIDIFICATION WITH C BACK DIFFUSION

This is an example of Scheil solidification with C back diffusion in solid phases. The results are compared between a simple Scheil and equilibrium calculation. The FEDEMO database is used and included with your installation.



See <u>"SCHEIL Module Commands" on page 290</u> for a list of the commands for this module.

# Example using the PARROT and EDIT\_EXPERIMENT Modules

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



8

All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

### TCEX36A - USING THE PARROT AND EDIT EXPERIMENTS MODULES

The example is divided into parts corresponding to the files in the **tcex36a** folder. It is recommended you read *tcex.36.readme* before continuing with this advanced exercise.

Also see the Data Optimization User Guide.

Each problem has "selected" experimental data. The example uses a fictitious binary A-B system where element A is BCC until it melts. Element B is BCC up to 1100 K and FCC above. There is a compound A2B stable in a limited temperature range.

- The creation of the "setup" file is described (*tcex36a.TCM*) where you enter the elements and phases into the Gibbs Energy System (GES) module
- The creation of the experimental data file is described (*tcex 36b.TCM*). When you run the TCEX36a macro, the contents of this macro file is automatically launched and the experimental data file created.
- The actual run in the PARROT module in order to obtain the result (*tcex 36cpd.TCM*)

There are many commands used in this advanced example (listed in alphabetical order):

"ADVANCED_OPTIONS" on	"AMEND_ELEMENT_DATA"	"AMEND_PHASE_
page 167	on page 78	DESCRIPTION" on page 89
"APPEND_EXPERIMENTAL_	"COMPILE_EXPERIMENTS"	COMPUTE_ALL_EQUILIBRIA
DATA" on page 236	on page 131	on page 36
"CONTINUE_OPTIMIZATION"	"CREATE_NEW_	"CREATE_NEW_STORE_FILE"
on page 132	EQUILIBRIUM" on page 185	on page 133

"DEFINE_COMPONENTS" on	"EDIT_EXPERIMENTS" on	"ENTER_PARAMETER" on
page 186	<u>page 133</u>	page 137
"ENTER_PHASE" on page 106	"LABEL_DATA" on page 46	"LIST_ALL_VARIABLES" on page 139
"LIST_PHASE_DATA" on	"LIST RESULT" on page 142	"MACRO_FILE_OPEN" on
page 117		page 5
"OPTIMIZE_VARIABLES" on	<u>"QUICK_EXPERIMENTAL_</u>	"READ_WORKSPACES" on
page 148	PLOT" on page 245	page 203
"RESCALE_VARIABLES" on	"SAVE_PARROT_	"SAVE_WORKSPACES " on
page 149	WORKSPACES" on page 149	page 52
"SELECT_EQUILIBRIUM" on	"SET_ALL_START_VALUES"	"SET_ALTERNATE_CONDITION"
page 206	on page 206	on page 55
"SET_ALTERNATE_MODE" on	"SET_FIX_VARIABLE" on	"SET_OPTIMIZING_VARIABLE"
page 150	<u>page 151</u>	on page 153
"SET_STORE_FILE" on page 156	"SET_WEIGHT" on page 65	

## **Example of the Tabulate Reaction (TAB) Module**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

#### TCEX12 - TABULATION OF THERMODYNAMIC DATA FOR REACTIONS

This example shows a number of independent cases using the TABULATE\_REACTION (TAB) module to tabulate thermodynamic data for reactions.



This uses the SSUB database. A license is required to run the example.

These commands are of note and used in this example:

- "TABULATE\_REACTION" on page 316
- <u>"DEFINE\_SPECIES" on page 14</u>
- <u>"ENTER\_FUNCTION" on page 304</u>
- "QUICK\_EXPERIMENTAL\_PLOT" on page 245
- "SET\_SCALING\_STATUS" on page 260
- "LIST\_SUBSTANCES" on page 307
- <u>"TABULATE\_SUBSTANCE" on page 319</u>

# **Example of the Reactor Module**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

#### TCEX24 - SIMULATION OF THE SILICON ARC FURNACE USING THE REACTOR MODULE

This is a simple reactor model with output of gases at the top and output of condensed phases at the bottom. The gas phase from one segment flows to higher segments, 80% reacts in the first above, 15% in the second above and 5% in the third above.

The condensed phases flow downwards and all of it goes to the next lowest segment. Heat can be added at any module. The only way to specify the initial state of the reactants added to the reactor is to specify the heat content.



This uses the SSUB database. A license is required to run the example.

These commands are of note and used in this example:

- <u>"DEFINE\_SPECIES" on page 14</u>
- <u>"CREATE\_STAGE\_BOXES" on page 284</u>
- <u>"CREATE\_DIVIDERS" on page 283</u>
- <u>"CREATE\_PIPES" on page 283</u>
- "START\_SIMULATION" on page 287

## **Examples with Specialized Databases**

The commands are described in the *Thermo-Calc Console Mode Command Reference*. All the links are to that guide (unless indicated otherwise), which is part of this documentation set. Or use the online help to navigate more easily.



All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

#### TCEX20 - ADIABATIC DECOMPRESSION IN A GEOLOGICAL SYSTEM

This example calculates the adiabatic decompression in a geological system using the geochemical database. The database is included with the file named PGEO.TDB, which is in the same folder as this example.

These commands are of note and used in this example:

- <u>"DEFINE\_SYSTEM" on page 15</u>
- <u>"DEFINE\_COMPONENTS" on page 186</u>
- "SET\_INPUT\_AMOUNTS" on page 212
- "LIST\_CONDITIONS" on page 198
- "SET\_CONDITION" on page 209
- <u>"SHOW\_VALUE" on page 217</u>

#### TCEX22 - CALCULATING AN ADIABATIC FLAME TEMPERATURE

This example examines a heat balance when  $C_3H_8$  is burned in oxygen by calculating the adiabatic flame temperature.



This uses the SSUB database. A license is required to run the example.

#### TCEX25 - SIMULATING THE REFINEMENT OF STEEL

This example is based on a typical steel refining process. Raw iron used to produce steel usually has high carbon and silicon content, which is why oxygen is blown into the furnace to burn off carbon. Lime (CaO) is added to form a slag with silica, and the slag can be removed. Alloying elements, such as Mn, Ni, Cr and V are added to produce the desired

steel. Since the reaction between O and C increases the temperature, scrap iron is added to keep the temperature constant (it is assumed the furnace is isolated and no heat is lost to the environment).

This example simulates blowing oxygen into a liquid steel of one metric ton (1e6 grams) with 4 w/o C, 2 w/o Si and 1 w/o Mn. 100 moles of CaO (equivalent to 5.6 kg) is added. Keeping the enthalpy constant is the way to simulate the isolation of the furnace. The oxygen reacts with carbon and increases the temperature. After blowing a certain amount of oxygen, scrap iron is added to keep the temperature constant.



This uses the SLAG database. A license is required to run the example.

#### TCEX27 - CALCULATING CHEMICAL VAPOR DEPOSITIONS (CVD)

This example calculates the chemical vapor deposition (CVD).



This uses the SSUB database. A license is required to run the example.

These commands are of note and used in this example:

- "DEFINE\_COMPONENTS" on page 186
- "SET\_CONDITION" on page 209
- "ADD\_INITIAL\_EQUILIBRIUM" on page 160
- "LIST\_AXIS\_VARIABLE" on page 198
- "LIST\_INITIAL\_EQUILIBRIA" on page 199

#### TCEX29 - CALCULATING THE SPECIATION OF A GAS

This example shows how to calculate the speciation of a gas.



This uses the SSUB database. A license is required to run the example.

These commands are of note and used in this example:

- "SET\_INPUT\_AMOUNTS" on page 212
- "LIST\_CONDITIONS" on page 198
- <u>"SET\_CONDITION" on page 209</u>

TCEX50 - QUASICHEMICAL MODEL USING THE DATABASE RETRIEVAL (TDB) MODULE

This example shows the quasichemical model using the <u>DATABASE\_RETRIEVAL</u> module (also called the TDB module).

The example replicates figures 3 and 4 from Kongoli, Pelton and Dessureault (1998)<sup>1</sup>.

<sup>1. &</sup>quot;Thermodynamic modeling of liquid Fe-Ni-Cu-Co-S mattes", Metall. Mat. Trans. B, 29(3), 591-601.

# **Pourbaix Module Examples**

All the macro files and the outputs are also combined into one file that you can search and read. This is included with your installation as a PDF (go to Help > Manuals folder > All Thermo-Calc Documentation and open Thermo-Calc Console Mode Example Macros.

#### TCEX40 - USING THE POURBAIX MODULE

This is the first in a series of examples to demonstrate the advanced POURBAIX module calculations and plotting. This example automatically calculates and plots a Pourbaix diagram for 0.001 m Fe in a 0.1 m NaCl aqueous solution at 25° C and 1 bar. Other diagrams, along various phase boundaries for the same interactions resulted from the same Pourbaix module calculation, are also plotted.

It is recommended you read the text file *TCEX40-README.txt*, which is located in the **Extended** subfolder.

Also see the Pourbaix Diagrams Educational Material Guide.

A so-called Pourbaix diagram is actually a phase diagram with independently-varied acidity (pH) and electropotential (Eh), for a heterogeneous interaction system at a certain bulk composition (that is by default always set as 1 kg of water dissolving a specified amount of metals and other solutes), under a defined temperature and pressure condition.

# The PAQ2 Database

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The example uses the PAQ2 (TCS Public Aqueous Solution Database using the SIT aqueous solution model) included with your installation. The database contains an AQUEOUS solution phase and REF\_ELECTRODE phase (as a reference for electron in aqueous electrolyte systems), as well as some data for various solid phases (solution or stoichiometric) and gaseous mixture phase. The PAQ2 is specially designed for these example calculations of the so-called Pourbaix diagrams (i.e., pH-Eh plots) and other types of phase diagrams or property diagrams in some aqueous-bearing multicomponent heterogeneous interaction systems within the limited chemical framework of Fe-Co-Cr-Ni-C-N-S-H2O-NaCl, via the Single-Database Option in the advanced POURBAIX module or through the normal TDB-GES-PLOY-POST routine.

# **Additional Examples in this Series**

Additional examples (TCEX40A, TCEX40B, TCEX40C, TCEX40D and TCEX40E) are provided to demonstrate options and features in the Pourbaix module and to show more advanced applications of the TCAQ and AQS aqueous solution databases [in the Multiple-Database Option, i.e., with appended data from the SSOL and SSUB databases]. See <u>"Pourbaix Module Examples TCEX40A to 40E" on the next page</u>.

### TCEX53 - POURBAIX DIAGRAM CALCULATIONS FOR A HETEROGENEOUS INTERACTION SYSTEM

Also see the Pourbaix Diagrams Educational Material Guide.

This example uses the PAQ2 database to calculate some Pourbaix diagrams. The DATABASE\_ RETRIEVAL (TDB), GIBBS\_ENERGY\_SYSTEM (GES), POLY3, and POST modules are used for the Fe-X-H2O-NaCl heterogeneous interaction system, where X = Cr-Ni-Co.



The initial bulk composition of Fe-based alloy in this calculation is just preliminarily assigned, in which the BCC\_A2 and/or FCC\_A1 solution phase(s) are considered as of primarily interest. For practical calculations, one shall have more precise inputs for the initial bulk compositions of alloys.

### Step 1: Single-Point Calculations for H2O-NaCl system

To demonstrate how to define the molality of NaCl in an aqueous-bearing heterogeneous interaction system.

### Step 2: Single-Point Calculations for Fe-X (X = Cr-Ni-Co) system

To demonstrate how to define the initial amount of alloy in an aqueous-bearing heterogeneous interaction system. In this example, we are interested in only the BCC\_A2 and FCC\_A1 phases in the Fe-based alloy.

Step 3: Single-Point Calculations for Fe-Cr-Ni-Co + H2O-NaCl system

Step 4: Pourbaix Diagram Mapping for Fe-Cr-Ni-Co + H2O-NaCl system - define the variables

From the same mapping calculations you arrive at by the end of the example, you can plot more diagrams using a variety of X-Y axis variables.



For a list of pre-defined symbols, go to the end of the tcex53.TCM macro file (open it in a text editor).

# **Pourbaix Module Examples TCEX40A to 40E**



Also search the online help or see the Pourbaix diagrams educational material and the *Thermo-Calc User Guide*.

These are the extended examples that are described in <u>"Pourbaix Module Examples" on</u> page 33.

### TCEX 40A

This example shows you how to use Option 1 where you **start a completely new calculation**. It uses a single database. The Pourbaix diagram for 0.001 m Fe in a 0.1 m NaCl solution at 25C and 1 bar is automatically calculated and plotted.



A more advanced version of 40A can be found in TCEX40E, which also uses Option 1 but with multiple databases. However, licenses are required for three types of commercial Thermo-Calc databases.



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See <u>"POURBAIX\_DIAGRAM Commands for Options 1 to 4" on page 269</u> in the *Console Mode Command Reference*.

A so-called Pourbaix diagram is actually a phase diagram with independently-varied electropotential (Eh) and acidity (pH), for an heterogeneous interaction system at a certain bulk composition (that is by default always set as 1 kg of water solving a specified amount of metals and other solutes), under defined temperature and pressure conditions. Some property diagrams, along various phase boundaries for the same interactions resulting from the same Pourbaix module calculation, are also plotted, without leaving Option 1.

At the end of this example, a new POLY3 file called *POURBAIX.POLY3* is automatically saved. For Windows users this is saved to the Documents folder. The following examples (TCEX40B, TCEX40C and TCEX40D) use this same POLY3 file for the calculations. Therefore it is important to make four copies of this file and rename these to TCEX40A.POLY3, TCEX40B.POLY3, TCEX40C.POLY3 and TCEX40D.POLY3. This must be done outside of the Thermo-Calc software and after the TCEX40A calculation and plotting is complete. This is so that the required POLY3 file structure is not lost. The reason you do not save from within Thermo-Calc is because of how the file is saved. When the command sequence (GO POLY3>SAVE TCEX40A.POLY3>Y) is used, the POLY3 workspace of the POURBAIX calculation results in the POLY3 module monitor and all the initial equilibrium points and the mapping/stepping calculation results are lost as well as parts of the POLY3 file structure (e.g., some definitions of the previously defined POURBAIX-type calculation system).

### TCEX 40B

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This example shows you how to use Option 2 where you **Open an old (existing) file and plot other property diagrams**.

Example 40A calculated and plotted the Pourbaix diagram for 0.001 m Fe in a 0.1 m NaCl solution at 25° C and 1 bar. Example 40B reads the original POURBAIX.POLY3 file , and uses the calculated results to plot property diagrams for the same heterogeneous interaction system.



For example 40B, the POLY3 file is not modified. However, it is still recommended that a copy of the original POLY3 file is used so you do not lose the file structure and calculated results.



If you have a POLY3 file generated after choosing Option 4, that file can also be used for Option 2, and vice versa.

### **TCEX 40C**

This example shows you how to use Option 3 where you **Open an old file and make another Pourbaix calculation**. The previously calculated (and copied) POURBAIX.POLY3 file from TCEX40A is edited to make another Pourbaix calculation and plot.

The same chemical system is used (Fe-H2O-NaCl) but with different P-T-X (pressure-temperature-composition) conditions and without the GASeous mixture phase. The new conditions are 0.01 m Fe in a 1 m NaCl solution at 200° C and 1 kbar.



In this example, the GASeous mixture phase for the Pourbaix diagram is not considered.



The POLY3 file is modified and rewritten with the new conditions. As previously recommended, make a copy of the original POLY3 file and rename it (e.g.TCEX40C.POLY3) so you do not lose the file structure and new calculated results.

After the default Pourbaix diagram is generated and refined, you can also use Option 3 to plot property diagrams along different phase boundaries for the same system. These settings are possible from within Option 3 and you do not need to go back to Option 2.

### **TCEX 40D**

This example shows you how to use Option 4 where you **Open an old file and make another stepping calculation**. The previously calculated (and copied) POURBAIX.POLY3 file from TCEX40A is edited to make a stepping calculation and plot.

The same chemical system is used (Fe-H2O-NaCl) but with different P-T-X (pressuretemperature-composition) conditions and using a POLY3 stepping calculation. The new conditions are 0.01 m Fe in a 0.1 m NaCl solution at 1 kbar and under varied temperature (from 0 to 400° C). The equilibrium conditions are modified and two stepping calculations are done. The same stepping calculation is used to plot property diagrams for the heterogeneous interaction system.

The example also explores settings such as editing the definitions for some system components and their initial bulk composition conditions. This is done through the direct and internal POURBAIX-POLY3 intersection, for a Stepping (or Mapping) calculation inside the Pourbaix module.



The POLY3 file is modified and rewritten with the new conditions. As previously recommended, make a copy of the original POLY3 file (from example TCEX40A) and rename it (e.g. to TCEX40D.POLY3) so you do not lose the file structure and new calculated results.



If you have a POLY3 file generated after choosing Option 2, that file can also be used for Option 4 and vice versa.

### **TCEX 40E**

This example shows you how to use Option 1 where you **start a completely new calculation**. It uses multiple databases (TCAQ, SSUBand SSOL). The Pourbaix diagram for 0.001 m  $Fe_{90}Co_5Ni_5$  alloy (9E-3 m Fe + 5E-4 m Co + 5E-4 m Ni) in an aqueous solution with 2E-2 m H<sub>2</sub>S and 3 m NaCl at 25° C and 1 bar is automatically calculated and plotted.

You need licenses for three Thermo-Calc databases, i.e.

- A: AQS (TGG Aqueous Solution Database; HKF model) or TCAQ (TCS Aqueous Solution Database; SIT model);
- B: SSUB (SGTE Pure Substances Database)

• C: SSOL (SGTE Alloy Solutions Database) or TCFE (TCS Steels/Fe-Alloys Database) or similar.

The TCAQ database contains only an AQUEOUS solution phase (using the SIT model) and REF\_ELECTRODE phase (as a reference state for electron in aqueous electrolyte systems). The SSUB database is appended for the GASeous mixture phase and some pure solid phases (so-called secondary phases formed during aqueous-involving interaction processes), and the SSOL database is appended for various solid solution phases.

The Pourbaix module inside the software also accepts the AQS2 aqueous solution database (using the complete Revised HKF model) in the Multiple Database Option, in addition to the TCAQ2 aqueous solution database (using the SIT model).

# **Data Optimization User Guide**

Thermo-Calc Version 2018b





### Introduction

#### In this section:

Introduction to Data Optimization	3
Data Optimization	3
The Least-Squares Method	4
The CALPHAD Approach	5

# **Introduction to Data Optimization**

This user guide describes the basic idea behind data optimization, contains brief descriptions of the modules and file types that you work with while doing your optimization, as well as rough outline of overarching workflow that you typically follow when working with optimization.

This guide assumes that you are familiar with using the Thermo-Calc Console Mode. It also assumes that you understand the principles of thermodynamics and are familiar with the theoretical foundations of computational thermodynamics.



Also see the *Thermo-Calc Console Mode Command Reference* included with this documentation set, which has information about all the commands available in the Thermo-Calc Console Mode, including PARROT and ED\_EXP. Note that the commands listed under PARROT and ED\_EXP are only commands that are specific to PARROT and/or ED\_EXP. These modules also contain some commands that are also available in POLY.

Thermo-Calc Console Mode <u>example tcex36</u> in the *Console Mode Examples Guide* is a detailed example of assessment and optimization in Thermo-Calc.

# **Data Optimization**

In a thermodynamic database, each phase in a system is characterised using a mathematical model of its Gibbs energy. The Gibbs energy of a phase depends on various state variables. It can be defined individually even in a heterogeneous system with many stable phases since the properties of one phase are completely independent of the properties of the other phases in the system. In most alloy systems, the thermodynamic properties of a phase can be modelled by expressing how the Gibbs energy depends on temperature and composition. State variables such as pressure, volume or entropy can also serve as parameters in the Gibbs energy expression. The Gibbs energy of the whole system is the sum of the products of the Gibbs energy of each phase multiplied by the amount of that phase.

How the Gibbs energy of the phase varies with various state variables is determined by a mathematical model and how various adjustable parameters of the model is set. By optimizing these parameters, you can calculate the thermodynamic properties of a system under various conditions (by calculating phase diagrams for example). The thermodynamic properties themselves are functions of temperature, pressure or composition. They include, among other properties, standard enthalpy of formation, entropy, heat capacity, molar volume, thermal expansivity, compressibility and Curie temperature for magnetic transformations.

Data optimization is about adjusting the model parameters so that calculated equilibria fit well with experimental data. An important part of a data optimization is therefore collecting and assessing available experimental and theoretical information about phase equilibria and thermochemical properties of a system.

When you optimize the parameters of the model of a phase, you put what is called optimizing variables into the Gibbs energy expressions of those parameters. During the optimization, the values of these variables are varied in order to find a fit between the calculated equilibria and the experimental data that you base the optimization on. Once you have settled on a Gibbs energy expression for a certain parameter of a phase of the element, then you must stick to this expression in all future optimizations involving the element. If you do not do this, then you must re-optimize all previous systems that involve this element when the expression is changed.

The Thermo-Calc software allows you to optimize parameters that characterize not only binary systems, but also ternary system and systems of even higher orders. You can even optimize parameters that characterise systems of different orders at the same time.

### **The Least-Squares Method**

Data optimization in Thermo-Calc is based on the least-squares method for fitting values calculated on the basis of a model with observed quantities. The software is accordingly trying to find the optimizing variable values that lead the minimized sum of the squares of the differences between the calculated values and the observed quantities (that is, of the errors or residuals).

The least-squares method works best under the following conditions:

- The observed quantities have a Gaussian probability distribution.
- The observed quantities are only subject to random errors.
- The different observations (experiments) are uncorrelated
- The standard deviation of each observation can be estimated.
- The number of observations is large.
- The models used give precise predictions.

Of course, these conditions are usually not all met in a normal thermodynamic assessment. But even in non-ideal conditions, there is no known method that works better than the leastsquares method.

# **The CALPHAD Approach**

The data optimization functionality in Thermo-Calc works according to the CALPHAD approach to computational thermodynamics. This approach builds on the development of models that accurately represent thermodynamic properties for various phases. These models enable you to predict the thermodynamic properties of multicomponent systems-based data concerning binary and ternary subsystems. The predictions can take many factors into accounts, such as, for example, crystallography, type of bonding, order-disorder transitions and magnetic properties.

CALPHAD is originally an abbreviation of *CALculation of PHAse Diagrams*, but as the approach has expanded in scope, it now stands for *computer coupling of phase diagrams and thermochemistry*.

#### REFERENCES

For in-depth information about CALPHAD as well as data optimization, see *Computational Thermodynamics* – *The Calphad Method* by Hans Leo Lukas, Suzana G. Fries and Bo Sundman (Cambridge University Press, 2007). CALPHAD was given its first general description in *Computer Calculations of Phase Diagrams* by Larry Kaufman and Harold Bernstein (Academic Press, 1970). The book contains some case studies of optimizations using Thermo-Calc.

Information about the concept of 'lattice stability' (which is important for the development of multicomponent thermodynamic databases) can also be found in the paper *Hume-Rothery and Calphad Thermodynamics* by Larry Kaufman, published in *Calphad and Alloy Thermodynamics*, edited by Patrice E.A. Turchi, Antonios Gonis, Robert Shull (The Minerals, Metals & Materials Society, 2002), pp. 3-19.

# Working with Data Optimization in Thermo-Calc

### In this section:

Data Optimization in Thermo-Calc	. 7
Data Optimization Workflow	. 7
The PARROT Module	. 8
The EDIT_EXPERIMENTS Module	8
Data Optimization in Other Modules	9
Optimization File Types	9
Optimization Workspaces	. 10
Method for Optimization and Simulation	.11

# **Data Optimization in Thermo-Calc**

The actual performance of the optimization of a system is primarily done in the PARROT module, and in its sub-module ED\_EXP. The POLY and POST modules are used to calculate and plot property or phase diagrams based on the optimized variables. This allows you to visually assess how good a fit you have achieved between your calculated results and your experimental data.

The end result of a successful optimization is typically an updated Thermo-Calc database file (with filename extension \*.TDB) or a new database file.

# **Data Optimization Workflow**

When you carry out an optimization using Thermo-Calc, you typically follow a workflow as outlined below.

- 1. Collect experimental data about your system from various sources such as journal articles and reports.
- 2. Create a POP-file in which you enter the experimental data you have collected.
- 3. Create a setup macro file in which you define your system and the variables you want to optimize. A useful general procedure is to first find a minimum set of variables that allows you to calculate most of the experiments.
- 4. Run the setup macro file.
- 5. Compile your POP-file. This file often contains errors, which need to be corrected, and the file recompiled several times before the compilation is completed without errors.
- 6. In the ED\_EXP module, check that the experiments in your POP-file reach equilibrium with reasonable results. If any experiments do not reach equilibrium with reasonable results, then try changing the starting values on equilibrium conditions (such as composition) to see whether the equilibrium can be computed after all. If this does not work, then you can temporarily exclude the experiment from the optimization. At a later stage in the optimization, when the optimizing variables have different values, you can check whether the experiments can be computed with reasonable results.
- 7. Back in the PARROT module, run the first optimization cycle and evaluate the feedback in the console. You can also plot diagrams in the POLY module that allow you to visually inspect the fit between calculated optimization results and the experimental data. If you are not satisfied with the fit, then enter the ED\_EXP module again and adjust the weights of the experiments in a way that is likely to improve the fit and run another optimization cycle. Reiterate until you are satisfied with the fit.

- 8. Once you get the optimization stable and smooth with the minimum set of variables chosen in step 3 above, try using different sets of variables to see whether improvements are possible.
- 9. When you are satisfied with the fit between calculated results and experimental data with your final set of optimizing variables, update your setup file and POP-file. In the setup file, enter the calculated optimizing variable values as the variables' start values. In the POP-file, enter the final weights of the experiments. With these files updated, you can easily regenerate the parameter values of your optimized system. Having the files updated also makes it easier to optimize the system again in light of new data or new theoretical models.
- 10. Finally, update the database with your calculated results or create a new database with information about the system that you have optimized.

## **The PARROT Module**

The PARROT module consists of a comprehensive subroutine package for data evaluation of thermodynamic model parameters from experimental data. The module has 99 variables that you can use for optimization and the module can handle a thousand experimental measurements in an optimization. However, there are limits on the simultaneous numbers of variables and experiments. These limits are listed at each optimization.

The PARROT module is part of the Thermo-Calc software. The module is not included in the software development kits TQ-Interface, TC-API or TC-Toolbox for MATLAB<sup>®</sup>.

The experimental data is entered and edited in a sub-module of PARROT: <u>"The EDIT</u> EXPERIMENTS Module" below.

# The EDIT\_EXPERIMENTS Module

The EDIT\_EXPERIMENTS module, or ED\_EXP for short, is used for entering, evaluating and editing experimental data. To enter the module, go to <u>"The PARROT Module" above</u> and then issue the command <u>"EDIT\_EXPERIMENTS" on page 133</u>.

Since ED\_EXP uses the POLY module for doing equilibrium calculations of experimental data points, many POLY commands are also available in the ED\_EXP module. However, the commands may work somewhat differently in the two modules. There are also unique ED\_EXP commands which are not found in POLY.

When you execute ED\_EXP commands, the PARROT workspace is modified. However, note that you must always first use <u>"READ\_WORKSPACES" on page 203</u> in ED\_EXP before doing

anything in the module. The command loads the experimental data that you are going to work with during the optimization. Furthermore, before leaving ED\_EXP you must use <u>"SAVE\_WORKSPACES " on page 52</u> or the results of your work in the ED\_EXP module is lost.

# **Data Optimization in Other Modules**

All kinds of thermodynamic data, calculated equilibrium states or dynamic parameters are transferred back and forth between the PARROT and the GIBBS module as well between these and the POLY module. Whenever an optimization run is performed, PARROT calls the GIBBS module for stored system definition data and model parameters.

In Thermo-Calc, the GIBBS module (the Gibbs Energy System) handles the models of the various phases that can form in a multicomponent system. It stores thermodynamic data and performs various Gibbs energy equilibrium calculations. It contains subroutines to analytically calculate the first and second partial derivatives of integral Gibbs energy with respect to any set of variables. Many thermodynamic models for various types of substances are implemented in the module. Some commands that are available in the PARROT module are also available in the GIBBS module.

In the GIBBS module, parameters of the Gibbs energy models are referred to as *TP*-*functions*.

PARROT calls the POLY module for equilibrium calculations. The POLY module is used for calculating and storing complex heterogeneous equilibria.

# **Optimization File Types**

File type	Description
POP-file	The POP-file (Parrot OPtimization file) is the file that contains all the experimental data that you use for optimizing Gibbs energy values for your system. The POP-file is a plain text file that contains ED_EXP commands. By default, the file has the filename extension *.POP in Thermo-Calc.
Setup file	The setup file is a Thermo-Calc macro file (*.TCM). This file typically contains commands that define your system, opens the PARROT workspace (and associated GIBBS and POLY workspaces) and sets optimizing variables.

These are the important files you work with when optimizing.

File type	Description
Other macro files	It is often useful to have other macro files than the setup file at hand while you are doing your optimization. For example, it is convenient to have a macro file that automatically plots phase diagrams or other diagrams based on the latest values of your optimizing variables. An optimization process typically involves many optimization cycles, so this operation needs to be done many times during a typical optimization.
Workspace files	The PARROT module has its own workspace with dynamic memories, similar to the workspaces of POLY and GIBBS. It is stored and updated in a PARROT workspace file with the filename extension *.PAR. The workspace file is a binary file whose format is hardware dependent. This means that the format is unique for each type of CPU. Hence, a workspace file saved on a computer with one type of CPU cannot be used on a computer with another type of CPU.
EXP-file	The EXP-file is a plain text file with information in the form of DATAPLOT Graphical Language commands. These commands can specify some or all of the data points in your POP-file. This allows you to plot these data points on top of your plotted optimization results (with the POST commands <u>"APPEND_ EXPERIMENTAL_DATA" on page 236</u> or <u>"QUICK_EXPERIMENTAL_PLOT" on</u> page 245). You can then visually assess the fit between these results and the experimental data. The PARROT workspace is not influenced in any way by the creation or use of EXP-files. DATPLOT User Guide included with this documentation set.

# **Optimization Workspaces**

When writing the setup file, you enter a <u>"CREATE\_NEW\_STORE\_FILE" on page 133</u> command which creates a PARROT workspace file. Alternatively, select an existing PARROT workspace file with <u>"SET\_STORE\_FILE" on page 156</u>. The workspace file is automatically updated and saved with the latest optimization results (after each use of the <u>"OPTIMIZE\_VARIABLES" on</u> <u>page 148</u> command). You can also explicitly instruct Thermo-Calc to save the workspace using <u>"SAVE\_PARROT\_WORKSPACES" on page 149</u>. If you want to get rid of your latest changes and return to the state of your workspace when it was last saved, use <u>"READ\_PARROT\_WORKSPACES" on page 148</u>.

When the PARROT workspace is saved, parts of the POLY and GIBBS workspaces are also saved to the PARROT workspace file, along with the PARROT workspace itself. Using the POLY command <u>"SAVE\_WORKSPACES" on page 204</u> creates a new POLY3 file, nothing is saved to the PARROT workspace file.

If you do any POLY calculations based on data in the current workspace file, then a \*.POLY3 file is created, which contains a copy of the current set of parameters. If you continue on your optimization and read the old \*.POLY3 file, then the new set of parameters is overwritten with the old set. It is therefore recommended that you never read a \*.POLY3 or \*.GES5 file while you are doing an optimization.

# **Method for Optimization and Simulation**

The PARROT module typically uses ordinary POLY minimization for equilibrium calculations. The global minimization technique that is used in POLY cannot be used because it automatically creates new composition sets which corrupt the PARROT workspace. Furthermore, the equilibrium definitions that you use as input for your optimization must each be attributed to a specific phase. Some of these definitions may specify some local or partial equilibrium state rather than a global minimum. However, it is possible to use the global minimization technique for some equilibrium calculations by using the ED\_EXP command "ADVANCED\_OPTIONS" on page 167 with the Toggle Alternate option.

Optimization in PARROT is performed on the basis of a 'maximum likelihood' principle. According to this principle, the best fit between various calculated results and all the input of experimental data is found where the sum of the square of the weighted residuals is at its minimum.

Typically, the results of an optimization is considered to be better the fewer optimizing variables that are needed to get the same level of fit between computed results and experimental data. If you can get almost the same fit using eight instead of twelve parameters, then this should be considered to be a significant improvement. When more parameters are used, the values of the individual parameters tend to become unrealistic. This is particularly true when it comes to temperature-dependent parameters. However, relatively small differences in the number of parameters are often insignificant. For example, it does not typically matter whether you used 24 or 25 parameters to reach a certain degree of fit.

It is often difficult to compare the quality of different optimizations based on the number of optimizing variables that have been optimized. Since you may rely more or less heavily on different pieces of available experimental data (and do so with good reason), it is possible that two different users could end up with very different numbers of parameters for the same system. For example, suppose you optimize the Fe-Al system to incorporate it into an Al database, while someone else optimizes the same system to incorporate it into an Fe database. In this situation, the two of you would probably make different judgements about which experimental information your calculated results must have a good fit with.

### **Creating the POP-File**

Before starting an optimization in Thermo-Calc, you must collect relevant experimental data about your system that is available in the research literature. This data is then entered in the POP-file. The experimental data points and measurements are entered in this plain text file in the form of ED\_EXP commands.

The syntax of the POP-file is in principle independent of the models used for describing the phases in the system.

#### In this section:

Experimental Information	13
POP-File Syntax	. 15
Entering Experimental Information	. 16
Using Stability Conditions	22
Grouping Equilibria with Labels	23
Dividing the POP-File into Several Blocks	23

# **Experimental Information**

An optimization of thermodynamic model parameters should be based on a range of reliable experimental results that is as wide as possible, as well as on empirical or semi-empirical correlations and theoretical considerations. During an optimization in Thermo-Calc, many kinds of thermochemical and thermophysical data can be mixed and used together.

The information needed for an optimization represent measurements of thermodynamic parameters in the system at equilibrium. These can be measurements of any of the thermodynamic quantities that can be set as conditions in Thermo-Calc. The information may represent measurements of activities or enthalpies in single-phase regions for example, or solubilities or transformation temperatures in a multi-phase region.

For each equilibrium that is used in the optimization, at least one quantity must have been measured and you also need to know the conditions that must be set to compute the system's equilibrium state. For example, consider a binary system in a single-phase region. Suppose that you have measured the temperature, pressure, composition and the chemical potential. Three of these quantities are necessary to specify the equilibrium state and the forth can be used as experimental information to model the phase.

If you have a two-phase region and know the temperature and pressure conditions, as well as which the stable phases are, then you could use the compositions of one or both phases as experimental data. The former information about the conditions is sufficient to determine the equilibrium. The experimental information can then be used to optimize the parameters that are used to characterise and model the system.



For examples how to enter various kinds of experimental data in the POP-file, see "Experiments and Experimental Data" on page 53.

### CONFLICTING OR MISSING INFORMATION

You are likely to sometimes have inconsistent or conflicting experimental information. In principle, all available experimental data should still be entered into the POP-file, unless you have good reasons to exclude some information. For example, if you have reason to believe that the samples used in some experiments were not pure, then these experiments could be excluded from the POP-file.

For some systems, you have very little experimental data to go on. When this is the case, do not try to fit the available data uncritically with high accuracy without considering possible errors. You can look for data in systems with similar elements to get an idea about what the reasonable estimates of data that is lacking could be. Or you can use calculations from first principles or semi-empirical methods to get reasonable estimates that can be used as experimental data.

#### **INVARIANT EQUILIBRIA INFORMATION**

The most valuable experimental information for an optimization is information about stable invariant equilibria. An invariant equilibrium is such that neither pressure, temperature nor composition can be changed without there being a change in the set of phases present in the system. It is recommended that all information about invariant equilibria for a system is included in the POP-file. These equilibria need not all be based on explicit experimental measurements of thermodynamic properties.

Reasonable estimates from available experimental data are often useful to include in the POP-file. Such estimates, which should be based on the available experimental data, helps Thermo-Calc find a good set of start values on the optimization variables. However, at the end of the assessment, these estimated equilibria should be excluded and you should complete the optimization based only on the experimental information that is based on measurements.

With some experience from phase diagram evaluation, it is possible to make reasonable estimates of metastable invariant equilibria. Such estimates helps reduce the number of phases that are assessed simultaneously during the optimization. For example, you may assume that a certain intermediate phase does not form in a system. Then extrapolate the liquidus curves below the stable three-phase equilibria and finally, estimate temperatures and compositions of metastable three-phase equilibria between two other phases and the liquid.

Another useful technique is to extrapolate a liquidus line from a peritectic equilibrium to estimate the congruent melting temperature of a compound. This estimate may be more useful than the information about the peritectic equilibrium itself as the equilibrium involves only two phases.

### EQUILIBRIA IN HIGHER-ORDER SYSTEMS INFORMATION

When you optimize a binary system, the available experimental information can often be described equally well by very different sets of parameters. Information about ternary and higher order systems that include the components of the binary system are then often useful for optimizing the binary system (the same is true of information about quaternary system with respect to optimizing ternary systems). Often, it is only when you extrapolate the optimized parameters to a higher-order system that you can judge which of the sets of model parameters that is best. Sometimes information from several ternary systems is required to make a reliable judgement about which set of parameters best describes a particular binary system.

#### HEAT CAPACITIES INFORMATION

If you want to optimize your system based on data about heat capacities, then the corresponding parameters should first be optimized separately before you perform the

general optimization of your system. The parameters should be kept fixed during the general optimization. This is because it is normally not possible to optimize heat capacity data together with other data in a meaningful way. If you do this, then it is very likely that the parameters related to the heat capacity are badly determined.

### **POP-File Syntax**

It is recommended that you specify an experimental equilibrium as close as possible to actual experimental conditions. Ideally, there is information about the set of stable phases, temperature, pressure and some or all compositions for the equilibrium.

In addition, it is recommended that you keep the POP-file well-organised and extensively commented. This is especially important if someone else reassesses the system when new experimental information is available.

If you have too many equilibria defined in your file, or if you define equilibria with different sets of components in the same file, then you must divide the POP-file in *blocks* (using the <u>"FLUSH\_BUFFER" on page 75</u> command). You know whether you have defined too many equilibria for one block when you compile the file. A too large number of equilibria leads to an error message when the file is compiled.

If you use the <u>"DEFINE\_COMPONENTS" on page 186</u> command, then it must be the first command in the POP-file (or if you have several blocks, DEFINE\_COMPONENTS can also be put as the first command after FLUSH\_BUFFER). This is because the command automatically reinitiates the current workspace, so the effect of any commands placed before it are not saved.



The last command in the POP-file should always be <u>"SAVE\_WORKSPACES</u>" on page 52.

### LEGAL COMMANDS IN A POP-FILE

The following commands can be used in a POP-file.



Also see the *Thermo-Calc Console Mode Command Reference Guide* for information about all these commands.

- "ADVANCED\_OPTIONS" on page 167
- <u>"CHANGE\_STATUS" on page 177</u>
- <u>"COMMENT" on page 75</u>
- "CREATE\_NEW\_EQUILIBRIUM" on page 185
- "DEFINE\_COMPONENTS" on page 186

- <u>"ENTER\_SYMBOL" on page 238</u>
- "EVALUATE\_FUNCTIONS" on page 197
- <u>"EXPERIMENT" on page 43</u>
- "EXPORT" on page 44
- <u>"FLUSH\_BUFFER" on page 75</u>
- "IMPORT" on page 46
- "LABEL\_DATA" on page 46
- "SAVE\_WORKSPACE" on page 286
- "SET\_ALL\_START\_VALUES" on page 206
- "SET\_ALTERNATE\_CONDITION" on page 55
- <u>"SET\_CONDITION" on page 209</u>
- "SET\_NUMERICAL\_LIMITS" on page 293
- "SET\_REFERENCE\_STATE" on page 258
- <u>"SET\_START\_VALUE" on page 216</u>
- <u>"SET\_WEIGHT" on page 65</u>
- "TABLE\_HEAD, TABLE\_VALUES and TABLE\_END" on page 73

### **Entering Experimental Information**

A POP-file in Thermo-Calc (\*.POP) mainly consists of descriptions of equilibria, each equilibrium describing a data point that is based on an experiment from the existing literature on your system. An equilibrium is specified using ED\_EXP commands.

The following is an example of a specification of an equilibrium with a single stable phase in the Au-Cu system. Equilibria with a single stable phase are often from experiments where enthalpies of mixing or chemical potentials have been measured.

```
CREATE_NEW_EQUILIBRIUM 1 1
CHANGE_STATUS PHASE LIQUID=FIX 1
SET_CONDITION T=1379 P=1E5 X(LIQUID,AU)=0.0563
SET_REFERENCE_STATE AU LIQ * 1E5
SET_REFERENCE_STATE CU LIQ * 1E5
EXPERIMENT HMR=-1520:200
COMMENT Measurement by Topor and Kleppa, Met trans 1984
```

### Creating a New Equilibrium

The first command in an equilibrium specification is always <u>"CREATE\_NEW\_EQUILIBRIUM"</u> on page 185. This command takes two integers as arguments. The first integer should specify a unique identifier that can be used later to refer to the equilibrium in question. The second integer is an initialisation code 0, 1 or 2:

- Code 1: This means that all components are entered but all phases are suspended. This code is appropriate in most cases.
- Code 0: This means that all components and phases in the equilibrium are suspended initially. This code is needed if you are using experimental data about systems of different orders simultaneously, such as data about both binary and ternary systems for example.
- Code 2: This means that all components and phases are initially entered.

#### EXAMPLE

For example, the first command in the example above creates an equilibrium data point with the unique identifier 1. All components of the system are entered but all phases are suspended:

CREATE\_NEW\_EQUILIBRIUM 1 1

### Setting Equilibrium Conditions and its Uncertainty

All the equilibrium-related commands that are placed in between two <u>"CREATE\_NEW\_</u> <u>EQUILIBRIUM" on page 185</u> commands are interpreted as specifying the conditions of the equilibrium created by the first command. The values of these conditions should be based on the conditions specified in the experiment that you base the equilibrium definition on.

#### EXAMPLE

In the example, the <u>"CHANGE\_STATUS" on page 96</u> command specifies that the equilibrium is a single-phase equilibrium with the liquid phase.

```
CHANGE STATUS PHASE LIQUID=FIX 1
```

Furthermore, the temperature is 1379 K, the pressure is 1 bar (1E5 Pa) and the mole fraction of Au is 0.0563.

SET\_CONDITION T=1379 P=1E5 X(LIQUID,AU)=0.0563

Sometimes you have reason to think that some information about an equilibrium condition may not be accurate. If this is the case, then you specify the degree of uncertainty for a condition with a colon after the condition quantity, directly followed by either an absolute value or a percentage. For example, if you have reason to doubt the reliability of the Au mole fraction measurement, then you could write:

SET\_CONDITION T=1379 P=1E5 X(LIQUID,AU)=0.0563:10%

This indicates that you think the Au mole fraction is between 0.0507 and 0.0619. PARROT calculates the equilibrium twice, once for 0.0507 and once for 0.0619, and then computes the change of each experimental value between these two equilibria. This change is used to modify the uncertainty specified for the experimental value.

### "Entering the Experimental Value and its Uncertainty" on the next page

The possibility of specifying uncertainties on conditions may be useful when you have experimental data about a ternary system. In a binary system, a tie-line is determined by the two phases, the temperature and the pressure. Measurements of the compositions of both phases can then be used as experimental data. To specify the tie-line in a ternary two-phase equilibrium one of the four compositions must also be set as a condition. If the measurement of this composition has the same uncertainty as the other compositions, then you can assign this same degree of uncertainty both to the composition selected as a condition and the compositions selected as experimental data. Alternatively, the sample (overall) composition could be used as a condition and all four phase compositions could be used as experiment values. These phases must have the status *entered* (not fixed) since the relative amount of each is unknown.

#### Setting Reference States

If you want to specify a non-default reference state for one or several of your system components, then you do this with the <u>"SET\_REFERENCE\_STATE" on page 258</u> command. In the following example, the reference state for the liquid phase is set to current temperature (\*) and a pressure of 1E5 Pa.

#### EXAMPLE

SET\_REFERENCE\_STATE AU LIQ \* 1E5 SET\_REFERENCE\_STATE CU LIQ \* 1E5

Note that in order for these reference states to be taken into account when the measurement of the enthalpy per mole of the system is entered with experiment, the R-suffix must be used.

### Entering the Experimental Value and its Uncertainty

The <u>"EXPERIMENT" on page 43</u> command is used to specify the quantity that the calculated results should be fitted to. When specifying this quantity, you both specify the measured quantity itself and your estimation of the uncertainty of this value.

### EXAMPLE

In the example, the EXPERIMENT command specifies the enthalpy per mole of the system, where this value is specified with respect to non-default reference states that you have set in the equilibrium specification:

EXPERIMENT HMR=-1520:200

The value before the colon is the quantity of the variable. The value after the colon is an assessment of the uncertainty of this quantity. A higher value means a greater uncertainty. In the example, the uncertainty has been specified with an absolute value but it could also be specified in percent of the quantity:

EXPERIMENT HMR=-1520:13%

Several experiments can be specified after the experiment command. Besides assigning values to conditions, it is also possible to specify that a certain quantity is greater than or less than a certain value. For instance, the following command would specify that the activity of the C-component is less than 0.01 and the mass fraction of the BCC phase of Ag is greater than 0.05:

EXPERIMENT ACR(C)<0.01:0.001 W(BCC,AG)>0.05:10%

Note that an experimental quantity can typically also be treated as an equilibrium condition, and vice versa. Which measured quantities that you treat as conditions and which quantities you treat as experimental data should be based on the accuracy of the different measurements. In most cases, the quantity that is based on the experimental technique with the lowest accuracy should be used as the experimental value.

### Entering Many Equilibria as a Table

When one thermodynamic variable (e.g. heat capacity) has been measured as a function of another (e.g. temperature), then experimental data is often presented as a table. By using the TABLE\_HEAD, TABLE\_VALUES and TABLE\_END commands, you can enter such experimental information into the POP-file in the form of a table. These three commands should always be used together and must be entered in the right order. See <u>"TABLE\_HEAD, TABLE\_VALUES</u> and TABLE\_END" on page 73.

The TABLE\_HEAD command is followed by a number. This number identifies the table as a whole, but also uniquely identifies the first equilibrium that is included in the table. This numeric identifier is incremented by one for each additional equilibrium in the table. Each

row in the table represents one equilibrium. Thus, the equilibrium defined by the first row in the table above (on the line immediately below TABLE\_VALUE) is uniquely identified with the number 1, the equilibrium defined by the second row by number 2, and so on until the last equilibrium on the sixth row, which is identified by the number 6.

Since the unique identifiers of the equilibria in the table are automatically generated from the unique numerical identifier that you give to the table, the unique numerical identifier that is normally given to the <u>"CREATE\_NEW\_EQUILIBRIUM" on page 185</u> command is replaced by two at signs (@@).

At least one of the equilibria conditions specified with the <u>"SET\_CONDITION" on page 209</u> command must be a condition whose value is set by the numbers in one of the table columns. To assign the numbers in a column to a condition, use the syntax SET\_CONDITION <condition>=@<column number>. For instance, in the example above, the mole fraction of X(ALO3/2) is for each equilibrium given by the number in the first column of the table.

At least one of the measured variables that are entered with the experiment command must be a variable whose value is set by the numbers in a table column. To assign the numbers in a column to a measured variable, use the syntax EXPERIMENT <variable>=@<column number>. In the example above, the measured temperature for each equilibrium is given by the number in the second table column.

The actual table should be entered between TABLE\_VALUES and TABLE\_END. Separate the columns with tabs or any number of spaces. The columns do not have to contain numbers, but can also contain other types of values, such as phase names for example. If a column contains text with spaces or special characters, then the text must be surrounded by double quotes ("ABC DEF&ghi", for example). The table may have columns that are not used, but each row must have exactly the same number of columns.

#### EXAMPLE

The following block of commands exemplifies how a table can be entered into a POP-file:

```
TABLE_HEAD 1

CREATE_NEW_EQUILIBRIUM 00 1

CHANGE_STATUS PHASE TETR=FIX 0

CHANGE_STATUS PHASE ION=ENT 1

SET_CONDITION P=101325 N=1 ACR(02,GAS)=.21 X(AL03/2)=01

EXPERIMENT T=02:5

TABLE_VALUE

0.46082476

0.57142380
```

0.66822276 0.7496 2176 0.50382426 0.63132249 TABLE END

### Setting Site Fraction Start Values

In some systems, you have to set non-default composition start values on some experiments in order to get them to converge properly during optimization. This may be important if your system has a miscibility gap for example, in which case several equilibria with different compositions exist at certain temperatures. In such cases, you have to make sure it is specified in the POP-file which side of the miscibility gap the equilibrium is on. The calculated equilibrium could otherwise end up on the wrong side, resulting in a large error.

In the POP-file, you can set the site fraction of a constituent on an equilibrium with <u>"SET\_START\_VALUE</u>" on page 216.

### EXAMPLE

For example, the following command sets the composition start value of VA constituent of the FCC\_A1 phase to 0.99:

SET START VALUE Y(FCC A1,VA#2)=0.99

Setting the mole fraction for a component in a phase is not as reliable as setting the site fraction.

Additional equilibria entered after the last SET\_START\_VALUE command are automatically given a similar start value if the following command appears previously in the POP-file:

```
SET_ALL_START_VALUES Y
```

If start values are not set automatically for phase constituents, then you may have to set the composition start value for each equilibrium (or table of equilibria) separately.

### **Commenting About an Equilibrium**

The COMMENT command inserts a comment about the equilibrium that is saved to the PARROT workspace file when the POP-file is compiled. The comment must be entered on one line.

Comments can also be entered immediately after a dollar sign (\$), but these are not saved to the PARROT workspace. Instead, these comments are ignored by Thermo-Calc when the POP-file is compiled.

### Simultaneous Use of Binary and Ternary Experiments

You can use experimental information about binary and ternary systems (as well as systems of even higher order) in the same POP-file and optimization. To do this, you must inform the PARROT module that not all components of the ternary system (or system of even higher order) should be considered. This allows you, for example, to use experimental information about a binary equilibrium that exists within a ternary system.

Use <u>"CHANGE\_STATUS" on page 96</u> with the keyword component and which components that should be entered. You should also give the initialization code 0 (rather than 1) to the <u>"CREATE\_NEW\_EQUILIBRIUM" on page 185</u> command. This code indicates that all components must be entered.

### EXAMPLE

For example, the following set of ED\_EXP commands characterises a binary (A-B) threephase equilibrium (FCC-BCC-LIQ) in a ternary system (A-B-C). It can be used in a POP-file that otherwise only contain experimental information about the ternary system (A-B-C):

```
CREATE_NEW_EQUILIBRIUM 1 0
CHANGE_STATUS COMPONENT A B = ENTERED
CHANGE_STATUS PHASE FCC BCC LIQ=FIX 1
SET_CONDITION P=1E5
EXPERIMENT T=1177:10
COMMENT from A-B
```

## **Using Stability Conditions**

Early in the optimization when parameter values are not so good it is often useful to set the driving force for precipitation of a phase (per mole of components) to make sure that it appears where it should. The DGM condition is very useful for making sure that phases appear where they should. It can be removed when the optimization becomes stable.

The DGM condition is also useful for suppressing phases that appear where they should not appear. Do this by setting the value and the uncertainty to something reasonable. For example, the driving force for precipitation of the BCC phase could be set as follows:

```
DGM(BCC)<-0.1:0.1
```

This produces an error even before BCC becomes stable and the weight of the experiment in question can be adjusted as needed. If you specify that the driving force for the BCC phase should be below 0 (DGM(BCC)<0:1E-4 for example), then the optimization converges extremely slowly (if at all). With such a sharp error condition, calculating the equilibrium is like finding the minimum point on a lawn that gently slopes towards a rock wall. The optimization keeps bouncing into the wall.

Another useful stability condition is the phase stability function, abbreviated QF. This can be used to specify that a phase is outside the miscibility gap for a solution phase. If QF(phase) is negative, then the phase is inside the miscibility gap; if QF(phase) is positive, then it is outside the miscibility gap.

# **Grouping Equilibria with Labels**

If you have several equilibria describing experiments that you want to be able to treat collectively, you can give all of the equilibria one and the same label.

Use LABEL to give an equilibrium a label. The label must start with the letter A and can only be up to four characters long. For example, you can insert the following command in several equilibria specifications:

LABEL ALH

During the optimization, you could use this label to, for example, set all the equilibria that have it to the same weight in the ED\_EXP module.

### **Dividing the POP-File into Several Blocks**

For two different reasons, you may have to divide your POP-file into different blocks:

- PARROT uses a buffer for storing compiled experimental data. If there is not enough memory in this buffer for storing all the experiments from the POP-file, then the POP-file can be divided into several blocks. When PARROT encounters the end of a POP-file block, it saves the experimental data that has been compiled to PARROT workspace, clears out the memory buffer and then proceeds to read the next block in the POP-file. This goes on until the experimental data from all the blocks have been saved to the PARROT workspace.
- If you want to put equilibria with different sets of components in one and the same POP-file, then you must divide the file up into blocks. Each block should only contain equilibria that all have the same set of components.

To divide the equilibria in the POP-file into different blocks, enter the <u>"FLUSH\_BUFFER" on</u> page 75 command between two equilibria. This command marks the beginning of new block. When the command is encountered during the compilation of the POP-file, the compilation of the current block is terminated, the equilibria saved to the PARROT workspace file and a new block is initiated. That the buffer is reinitiated means that all functions and constants that were entered in the previous block must be entered again in new block in order to be used. When editing your experimental data in the ED\_EXP module, you can select which block to load and edit using READ\_WORKSPACES <block number>. The block before the first instance of FLUSH\_BUFFER is block number 1, the block after the first instance and before the second instance is block number 2, and so on.

# The Setup File

### In this section:

Creating the Setup File	26
Defining the System	26
Creating Additional Composition Sets of a Phase	27
Entering the Optimizing Variables	27
Initializing Optimizing Variables in PARROT	. 29
Creating the Workspace File	30
Compiling the POP-file	30
Returning Control to the Console	31

## **Creating the Setup File**

The second step is to create the setup file. This is an ordinary Thermo-Calc macro file (\*.TCM). Typically, your setup file should contain the following:

- A system definition.
- A section where you enter model parameters and optimizing variables.
- A GO PARROT command.
- A section where initial values are assigned to the optimizing variables for the first optimization cycle.
- A CREATE\_NEW\_STORE\_FILE <filename.par> which creates the PARROT workspace file onto which the results of your optimization are continuously saved and updated. By default this workspace file has the same filename as the setup file, but with the \*.PAR filename extension.
- A COMPILE\_EXPERIMENTS <filename.pop> command. Since you often encounter syntax errors when you first try to compile a POP-file, it may be useful to execute this command from the console rather than as part of the setup macro file.
- Normally, you also have a <u>"SET\_INTERACTIVE" on page 292</u> at the end of the setup file. This command returns control over Thermo-Calc to the console.

It is possible to interactively execute all these commands one at a time in the console. It may be instructive to enter the commands that would normally go into the setup file interactively while using the on-line help while you are learning to use the PARROT module.

## **Defining the System**

If you have an existing database with information about your system, then you can write the system definition part of your setup file in two different ways.

- If you have an unencrypted user database, then you can define your system by entering DATA module commands directly in the setup file. Use the DATA commands that you normally use to define a system when doing equilibrium calculations in Thermo-Calc Console Mode.
- Define your system by entering DATA module commands directly in the console, then generate a macro file with the GIBBS command LIST\_DATA <name of setup file>.tcm p . This command saves all the data that has been loaded in the GIBBS workspace as a macro file. This file can then be modified as required into an optimization setup file.

### **Creating Additional Composition Sets of a Phase**

If you need several composition sets of a phase, then these should be created with the GIBBS command AMEND\_PHASE\_DESCRIPTION <phase name> composition\_sets <composition set number> in the setup file.



If you add composition sets of a phase after the POP-file has been compiled, the data structure in the workspace is corrupted and you have to recompile the POP-file.

## **Entering the Optimizing Variables**

When you have retrieved the data about your system for the setup file, you must manually enter the model parameters that you want to optimize. This is done using the GIBBS command <u>"ENTER\_PARAMETER" on page 137</u>, but when you specify the Gibbs energy expression for the parameter, you enter variables that can be given various values into the expression. It is the values of these variables (and thus, the Gibbs energy of the parameter) that you are optimizing.

How you in detail specify the optimizing variables in your set up file depends on which phase you are trying to optimize and what thermodynamic model that is used in the optimization.

"Thermodynamic Models" on page 62

### **Entering Optimizing Variables in Parameter Specifications**

There are ninety-nine predefined variables in the PARROT module which you can use in your optimization. These variables are referred to as v1, v2, v3, ... v98 and v 99. When the GIBBS workspace is initialised, all optimizing variables are set to a fixed numerical value of zero. When you do an optimization, there is always some variable for which you want to evaluate the best value, but you can assign fixed values to optimizing variables too. It is often useful to enter all the variables that you can conceivably want to optimize, and then you can interactively select and change which variable to focus at various stages of the optimization.

The setup file excerpts that follow show how optimizing variables are entered. However, before the optimizing variables are introduced in this example, the following command enters an ionic liquid solution phase (a phase for which you have some experimental information):

```
ENTER_PHASE IONIC_LIQ Y,
AL+3,ZR+4 ; ALO2-1,O-2 ; N N
```

This <u>"ENTER\_PHASE" on page 106</u> command enters a phase called IONIC\_LIQ into the GIBBS workspace. This phase is of the type ionic liquid solution, which is specified by type code Y. Such a phase is modelled by the *Ionic Two Sublattice Liquid Model*. The next two arguments specify the two sublattices, in this case the cations AL+3,ZR+4 and the anions ALO2-1,O-2.

Components that interact in a sublattice are separated by a comma. The first N at the end means that no additional constituents are added to the phase. The second N means that a list of possible parameters for the phase should not be shown in the console.

There are model parameters that describe the phase which you do not want to optimize, in which case the Gibbs energy expression of the parameter does not contain any optimizing variables.

#### EXAMPLE 1

For example, the following parameters of the IONIC\_LIQ phase do not contain any optimizing variables:

```
ENTER_PARAMETER G(IONIC_LIQ,AL+3:ALO2-1;0) 298.15 +2*GAL2O3_L;
6000 N
ENTER_PARAMETER G(IONIC_LIQ,AL+3:O-2;0) 298.15 +GAL2O3_L+400000;
6000 N
ENTER PARAMETER G(IONIC LIQ,ZR+4:O-2;0) 298.15 +2*GZRO2L; 6000 N
```

These parameters define the Gibbs energy expressions for three constituents in the constituent array, namely AL+3:ALO2-1, AL+3:O-2 and ZR+4:O-2. The names that occur in these expressions—that is, GAL2O3\_L and GZRO2L—are the names of complex functions of temperature that are entered in the GIBBS workspace with the <u>"ENTER\_SYMBOL" on</u> page 109 command.

The numbers that frame the Gibbs energy expressions specify the temperature range within which the expression should be applied. In the above, the expressions are applied in a temperature range from 298.15 K to 6000 K.

#### EXAMPLE 2

In this example, suppose that you want to optimize the model parameter for the ZR+4:ALO2-1 combination, then you could enter the variables V1 and V2 in the Gibbs energy expression for that parameter in the following way:

```
ENTER_PARAMETER G(IONIC_LIQ,ZR+4:ALO2-1;0) 298.15 +GZRO2L +2*GAL2O3 L +V1+V2*T; 6000 N
```

You normally have different optimizing variables in different parameters, but in some cases you may want several parameters to vary together, in which case you can use the same variable in the Gibbs energy expressions of several parameters.

#### **Entering Interaction Parameters**

For some systems, you have to consider interaction parameters in order to take excess Gibbs energy into account and correctly model the phase. By default, the software uses a Redlich-Kister expression for the excess Gibbs energy. The coefficient is entered after the constituent array when you use the <u>"ENTER\_PARAMETER" on page 137</u> command. In the last section, all the parameters were entered with a coefficient of zero (0).

#### EXAMPLE

To take account of the excess Gibbs energy, you might enter the following:

```
ENTER_PARAMETER L(IONIC_LIQ,ZR+4:ALO2-1,O-2;0) 298.15 +V11+V12*T;
6000 N
ENTER_PARAMETER L(IONIC_LIQ,ZR+4:ALO2-1,O-2;1) 298.15 +V13+V14*T;
6000 N
ENTER_PARAMETER L(IONIC_LIQ,ZR+4:ALO2-1,O-2;2) 298.15 +V15+V16*T;
6000 N
```

Parameters with interaction coefficients are referred to as *interaction parameters*. The regular parameter has a coefficient of 0. The subregular parameter has a coefficient of 1. It is possible to interaction coefficients up to a value of 9, but you are advised not to use more than three coefficients (that is, with coefficient values of 0, 1 and 2).

### **Initializing Optimizing Variables in PARROT**

Variables that have been entered into Gibbs energy expressions for model parameters in the setup file has to be initialized and given start values before they are optimized.

You have to estimate what start values to give to the optimizing variables based on your knowledge of the system or of relevantly similar systems. There are no general rules for what start values your optimizing variables should have. If you are not able to make any informed estimation about what start values to use, then you may not be able to find a good fit between your experimental data and your calculated results. In this case, it is often best to start the optimization in the so-called alternate mode. In alternate mode, the PARROT module helps you find some reasonable start values for your optimizing variables.

#### "About Alternate Mode" on page 46

Even if you are starting your optimization in alternate mode, you still have to initialize the optimizing variables though (you can all give them a start value of 0).

To initialize an optimizing variable and to give it a start value, use either <u>"SET\_OPTIMIZING\_</u> <u>VARIABLE</u>" on page 153 or <u>"SET\_SCALED\_VARIABLE</u>" on page 155. Using SET-OPTIMZING\_ VARIABLE results in a scaling factor equal to the current value of the parameter unless the current value is zero, in which case the scaling factor is 1000. This is typically appropriate for enthalpic parameters, but not necessarily for other parameters. The scaling factor is the factor by which the optimizing variable value is varied during the optimization.

#### **EXAMPLE**

For example, if you want to optimize the variables V1, V2, V11, V13 and V17, then you could enter the following in the setup file:

```
SET_OPTIMIZING_VARIABLE 1 +43000
SET_OPTIMIZING_VARIABLE 2 -35.2
SET_OPTIMIZING_VARIABLE 11 +72000
SET_OPTIMIZING_VARIABLE 13 +27000
SET OPTIMIZING VARIABLE 17 -100000
```

The value given to each optimizing variable (+43000 for variable V1 for example) is also that variable's initial scaling factor. If you want to set the initial scaling factor of a variable to a value that is lower or higher than its start value, then use SET-SCALED\_VARIABLE <start value><scaling factor><min value><max value>.

### **Creating the Workspace File**

In the setup file, after the system definition and the entered optimizing variables, use the PARROT command <u>"CREATE\_NEW\_STORE\_FILE" on page 133</u> to create and save a PARROT workspace file for the system defined in the setup file. For example, the following command saves all current data on system definitions (elements, species, components, phases), symbol definitions (constants, variables, functions, tables) and parameters that are stored in the workspace used by GIBBS, POLY and PARROT into a file called AL2O3-ZRO2.PAR:

CREATE NEW STORE FILE AL203-ZR02



The experimental information in the POP-file is not saved to this PARROT workspace file.

### **Compiling the POP-file**

At the end of the setup file, use the command <u>"COMPILE\_EXPERIMENTS" on page 131</u> followed by the name of your POP-file to compile the data in that file and save it to the current PARROT workspace file. Note that you must first have opened or created a new PARROT workspace file before you compile your POP-file. The other argument to the command can typically be given default values with comma signs (,):

```
COMPILE_EXPERIMENTS AL203-ZR02.POP,,,
```

If Thermo-Calc encounters errors in the syntax of the POP-file, then error messages are displayed in the console. Normally, the compilation also stops when syntax errors are encountered. If this happens, then you must correct the errors and recompile the file. It is useful to have two console windows open when you do this. That way, you can use one console for editing the experimental data and another console for compiling and reading any further error messages.

# **Returning Control to the Console**

Put the command <u>"SET\_INTERACTIVE" on page 292</u> at the end of the setup file if you want to return control to the console when the commands in the setup file have all been executed.

# **Optimizing in PARROT**

### In this section:

Optimizing the System in PARROT	33
Changes that Require POP-file Recompilation	33
Optimization Workflow	33
Ensuring Computation of all Equilibria	34
Setting Weights	35
Optimizing and Evaluating Results	36
The Critical Set of Experiments	37
Continuing the Optimization and Resetting Variables	37
Updating the Set of Optimizing Variables	39
Reducing the Number of Optimizing Variables	40
Plotting Intermediate Results	40

# **Optimizing the System in PARROT**

When you have created a POP-file, a setup macro file and (optionally) EXP-files, you should start doing optimization runs interactively in the console. This process can be divided into a number of steps (such as running optimization, evaluating optimization results, modifying weights and models and adding new information), but usually, you have to go through many optimization cycles before achieving a satisfactory fit between calculation results and experimental data.

Typically, it is not obvious when an optimization is finished. You must exercise your judgement about when the fit you have achieved is good enough.



It is possible to interactively change almost everything in the initial setup macro file and in the POP-file. For example, you can add more parameters to be optimized and modify or add more experimental information.

# **Changes that Require POP-file Recompilation**

Some changes that you do interactively in the PARROT module or the ED\_EXP sub-module destroys the data structure in the workspace. For example, if you add more composition sets to a phase, then the number of phases actually change and the links break between the experimental data and the thermodynamic models. Note that if the global minimization technique is used in POLY, then new composition sets that corrupts the PARROT workspace may be created automatically. If this happens, then you have to compile the POP-file again.

If you recompile the POP-file, then the experimental equilibria are calculated with the default start values.

## **Optimization Workflow**

Normally, the workflow during an optimization is the following:

- 1. In the PARROT module, run the setup macro file using <u>"MACRO\_FILE\_OPEN" on</u> page 5.
- 2. Go to ED\_EXP with <u>"EDIT\_EXPERIMENTS" on page 133</u>, use <u>"READ\_WORKSPACES" on page 203</u> to load your experimental data and then execute <u>"COMPUTE\_ALL\_EQUILIBRIA" on page 36</u>.
- 3. If an equilibrium cannot be computed, then this either means that the start values (site fractions, for example) were inappropriate or that the equilibrium cannot be calculated with the current set of parameters. If the latter, then the equilibrium may have to be excluded from the optimization. However, you should make an effort to exclude the possibility that the reason is just that you do not have the appropriate start values for the equilibrium.

- 4. When each equilibrium is either successfully computed or excluded from the calculation, go back to the PARROT module and run the <u>"OPTIMIZE\_VARIABLES" on page 148</u> command. Give the number of optimization iterations you want the Thermo-Calc program to carry out as an argument to the command. This number specifies how many different sets of values for the optimizing variables that PARROT tries to fit with the experimental data. It is recommended that you run 0 iterations first to see the discrepancy between the experimental data and the initial calculated results. The current workspace file is automatically updated with the last set of optimized variables and calculated results.
- 5. List and evaluate the result of the optimization using <u>"LIST\_RESULT" on page 142</u>. Typically, what is of interest in the results are the relative standard deviation of the optimizing variables and the error values in the list of calculated experimental data results. If the relative standard deviation for an optimizing variable is very large, this means that the variable doesn't have any effect on the computed results and that the variable should not be used (you can make PARROT ignore the variable by setting its value to 0 with <u>"SET\_FIX\_VARIABLE" on page 151</u>). If the relative standard deviation is close to zero for a variable, then this might indicate that you need to use more variables. When it comes to errors in the calculated equilibrium results, these can be decreased by lowering the weight that PARROT gives the experiment in the calculation (the more weight, the more the experiment affects the calculated results).
- 6. Use <u>"OPTIMIZE\_VARIABLES" on page 148</u> or <u>"CONTINUE\_OPTIMIZATION" on</u> <u>page 132</u> to carry out a new optimization cycle. Go back to step 5 above and continue to do optimization cycles until you judge that there is a good enough fit between your calculated results and your experimental data. It is recommended that you do not only rely on feedback from <u>"LIST\_RESULT" on page 142</u> but also frequently plot various diagrams and compare them visually to some of your experimental data.

# **Ensuring Computation of all Equilibria**

When you have run the setup macro file, you should start by trying to compute all the equilibria that were created when your POP-file was compiled. To do this, you must first go to the ED\_EXP submodule, and load the experimental data from the current PARROT workspace file with the READ\_WORKSPACES <block #> command (normally, the block number is 1).

When you have loaded all equilibria, you should try to compute them all using the ED\_EXP command <u>"COMPUTE\_ALL\_EQUILIBRIA" on page 36</u>. A list of calculated equilibrium results are shown in the console, but the software is not able to reach an equilibrium for an experiment, then the program cancels the calculations and displays an error message.

The following list shows an example output where the first two experiments in the POP-file have been successfully computed, but an equilibrium is not reached for the third experiment. No other equilibria are computed, since the calculations are aborted when the error occurs.

Eq Lab Iter Weight Temp Exp Fix phases or comments 1 A1 22 1. 2575.3 IONIC\_ FLUORI TETR 2 A1 18 1. 2156.9 IONIC\_ CORUND TETR \*\*\* ERROR 1614 IN QTHISS \*\*\* CONDITIONS CAN NOT BE FULLFILLED

When this happens, try to find out why the equilibrium calculation fail. Are the condition start values that you have given the equilibrium inappropriate for example? If you have a similar equilibrium that has already been successfully computed, then you can select the equilibrium that does not work, use the ED\_EXP command <u>"SET\_ALL\_START\_VALUES" on page 206</u> and accept all the suggestions for the site fractions. Alternatively, use TRANSFER\_START\_VALUE <equilibrium number> to make the currently selected equilibrium inherit the start values of the indicated equilibrium. Run <u>"COMPUTE\_EQUILIBRIUM" on page 182</u> again to see if the equilibrium can be computed with the inherited start value.

If you cannot find an explanation of why the equilibrium cannot be computed, then it may be that the experiment is faulty. If so, then this experiment should be excluded from the optimization by being given a weight of 0. You should then run COMPUTE\_EQUILIBRIUM again to make sure that all equilibria can be computed. However, experiments that fail to reach equilibria could succeed in doing so if the optimizing variables have other values. It is therefore sometimes worth trying to include experiments later in the optimization process (when the optimizing variables are relatively close to their final values), even if they had to be excluded at its start. In some cases, it might also be appropriate to change the error tolerance for in an experiment (using the ED\_EXP command <u>"EXPERIMENT" on page 43</u>).

You must save the PARROT workspace with <u>"SAVE\_WORKSPACES</u>" on page 52 before you exit the ED\_EXP module.

# **Setting Weights**

8\$B

To change the weight of an experiment, first enter the ED\_EXP module from PARROT using "EDIT\_EXPERIMENTS" on page 133. Then use the "SET\_WEIGHT" on page 65 command to set the weight for a specific equilibrium (equilibrium number 3 for example), for a range of equilibria (the equilibria 10-19) or all the equilibria that have a certain label (such as all the equilibria with the label A1). To save the new weight(s), use "SAVE\_WORKSPACES" on page 52.

By default, all the experimental information in your POP-file is treated equally: each experiment has a weight of 1. However, experiments that you excluded because they did not reach equilibrium now have a weight of 0. The higher the weight of an experiment is, the more Thermo-Calc tries to ensure that the computed result fits the condition measured in that experiment.

The contribution of an experiment's error to the sum of errors is multiplied by the square of the weight set for that experiment. In other words, if you want to cut the error of an experiment with weight 1.0 by half, then you should use a weight of 0.7. This makes the experiment's error 0.49. Setting the weight to 1.4 instead makes the error twice as large (1.96).

Note that if you use <u>"READ\_WORKSPACES" on page 203</u> after you have changed weights on the experiments, your changes are lost unless you have saved those changes. In the ED\_EXP module, you can get a list of all the experiments in the console if you first read the experimental data from the workspace file (with READ\_WORKSPACES) and then use <u>"LIST\_ALL\_EQUILIBRIA" on page 47</u>.

## **Optimizing and Evaluating Results**

When you have ensured that all the experimental data points can reach equilibrium, go back to the PARROT module and use <u>"OPTIMIZE\_VARIABLES" on page 148</u> <number of iterations> with 0 iterations. Present the result of this initial optimization by using <u>"LIST\_RESULT" on page 142</u>.

When evaluating the optimization results, the most important information is found in the list of experiments at the end of the listed results. The rightmost column shows the difference between the value of the experimental data and the value computed by Thermo-Calc in the last optimization cycle.

The following shows part of list of experiments in the output of a LIST\_RESULT command:

106	ACR(B)=0.34	0.3127	2.89E-02	-2.7282E-02	2 -0 9444		
107	ACR(B)=0.23	0.2085	2.89E-02	-2.1522E-02	2 0.7450	$\mathbf{N}$	
108	ACR(B)=0.12	0.1042	2.89E-02	-1.5761E-02	-0.5455	- \	
110	HMR(LIQUID)=-1964	3.6380E-12	5.00E+02	1964.	3.928	۱ ا	1
111	HMR(LIQUID)=-3500	0.000	5.00E+02	3500.	7.000	*	۱
112	HMR(LIQUID)=-4588	0.000	5.00E+02	4588.	9.176	*	۱
113	HMR(LIQUID)=-5239	7.2760E-12	5.00E+02	5239.	10.48	*	
114	HMR(LIQUID)=-5454	-3.6380E-12	5.00E+02	5454.	10.91	*	
115	HMR(LIQUID)=-5233	1.8190E-12	5.00E+02	5233.	10.47	*	I
116	HMR(LIQUID)=-4575	-1.8190E-12	5.00E+02	4575.	9.150	*	I
117	HMR(LIQUID)=-3481	-1.8190E-12	5.00E+02	3481.	6.962	- * <b>/</b>	
118	HMR(LIQUID)=-1950	0.000	5.00E+02	1950.	3.900		
						/	

#### PARROT:

The experiments for which the optimization has not found a good fit are marked with an asterisk (\*) or a hash sign (#) in the rightmost column. An asterisk in this column indicates that the experiment has an error that is larger than the error tolerance and a hash sign indicates that the error is much larger. The error tolerance depends on the degree of accuracy set for that particular piece of experimental information. The error values are presented in the next to rightmost column.

Note that if you are optimizing in *alternate mode*, then you usually do not need to change the weights for the experiments. Instead, you can simply run OPTIMIZE\_VARIABLES again, specifying, say, thirty iterations, evaluate the results again, and continue in that way until you are satisfied with the results.

### "About Alternate Mode" on page 46

### **The Critical Set of Experiments**

The set of weighted experiments that you end up with as a result of your optimization is referred to as the *critical set* of experiments. When you work to determine this set and the weights of the experiments, the following factors should be reflected in both the selection of experiments and the weights of those experiments:

- The reliability of the experimental technique.
- The extent of agreement/disagreement between independent measurements of the same quantity.
- The extent of agreement/disagreement between data obtained with different experimental methods.

## **Continuing the Optimization and Resetting Variables**

To continue the optimization after you have used <u>"OPTIMIZE\_VARIABLES" on page 148</u>, you can either use this command and enter <Number of iterations> again or <u>"CONTINUE\_</u> <u>OPTIMIZATION" on page 132</u> <Number of iterations>.

If you use CONTINUE\_OPTIMIZATION, then PARROT continues the optimization using the same Hessian matrix. If the optimization is going well but you run out of iterations, then continuing with the same Hessian matrix may be useful.

As before, use <u>"LIST\_RESULT" on page 142</u> to inspect the fit between the calculated and the experimental values for your experiments. If necessary, adjust the weights of the experiments again in the ED\_EXP module. You may even discover that you have to exclude additional experiments from the optimization.

Besides looking at the rightmost column in the list of experiments (that is produced when you use LIST\_RESULT), it is also useful to look at the *sum of squares* of the errors of all the variables. This is shown in the output of OPTIMIZE\_VARIABLES and CONTINUE\_ OPTIMIZATION. This sum should be as low as possible.

The following shows part of the output that you get when using LIST\_RESULT:

```
== OPTIMIZING VARIABLES ==
AVAILABLE VARIABLES ARE V1 TO V00
VAR. VALUE
                    START VALUE
                                    SCALING FACTOR REL.STAND.DEV
V1 6.94641846E+05 4.3000000E+04 4.3000000E+04 1.42299558E+00
v2
    -3.32619842E+02 -3.5200000E+01 -3.52000000E+01 7.91369573E-01
V11
     8.37422071E+04 7.2000000E+04 7.2000000E+04 2.02212692E-01
V13 2.65253341E+04 2.7000000E+04 2.7000000E+04 1.45201828E-01
V17 -3.66637428E+05 -1.0000000E+05 -1.0000000E+05 8.04279315E-01
NUMBER OF OPTIMIZING VARIABLES : 5
ALL OTHER VARIABLES ARE FIX WITH THE VALUE ZERO
THE SUM OF SQUARES HAS CHANGED FROM 3.50196879E+05 TO 3.49096720E+05
DEGREES OF FREEDOM 14. REDUCED SUM OF SQUARES 2.49354800E+04
```

In the example you can see that the sum of squares have decreased slightly. (The final sum of squares that you get after an optimization run is also presented directly in the output from OPTIMIZE\_VARIABLES and CONTINUE\_OPTIMIZATION.) As long as the sum of squares figure is decreasing, the optimization is working as it should.

When your calculated results have been improving for a while, that is, when the sum of squares has been decreasing, and you are confident that you are making definitive progress, then you should use the PARROT command <u>"RESCALE\_VARIABLES" on page 149</u> to reset the starting values of the optimizing variables to their current values. When you do this, the start values and scaling factors of the optimizing variables (shown in the table's third column in the preceding screen shot), are all set to the current values (shown in the table's second column).

If your optimization is not going well, then you may want to reset the values of your optimization variables to their current start values. To do this, use <u>"RECOVER\_VARIABLES"</u> on page 149.

After rescaling the variables, continue the optimization using OPTIMIZE\_VARIABLES again. Cycle through calls of the commands OPTIMIZE\_VARIABLES/CONTINUE\_OPTIMIZATION, CONTINUE\_OPTIMIZATION, RESCALE\_VARIABLES and OPTIMIZE\_VARIABLES again until your set of optimizing variables does not seem to improve anymore. Even when it seems to you that you have reached this point, it is recommended that you use OPTIMIZE\_ VARIABLES/CONTINUE\_OPTIMIZATION a few extra times to make sure that the variable values cannot be further improved, that is, that the sum of squares of the errors of all the variables cannot be reduced further. If the final solution is repeatedly calculated after the same number of iterations and reaching the same result, then it is generally reasonable to accept the current variable values.

## **Updating the Set of Optimizing Variables**

Besides evaluating the fit between the calculated results and the experimental data, you should also ensure that you have the right number of optimizing variables. If you have too many or too few optimizing variables, then you may not be able to use your model to accurately extrapolate thermodynamic properties in a wide range of temperature, pressure and composition conditions.

With more optimization parameters, the sum of errors usually decrease, but the parameters also tend to become less precisely determined. A measure of this precision is the Relative Standard Deviation (RSD) of each optimizing variable. However, the RSD is only significant if you have used <u>"RESCALE\_VARIABLES" on page 149</u> followed by an optimization which converges and the values of the variables do not change much. The RSD tells you how much the parameter can be changed in either a positive or in a negative direction without changing the reduced sum of errors with more than one unit.

A large RSD value thus means that the parameter has not been determined well.

```
== OPTIMIZING VARIABLES ==
AVAILABLE VARIABLES ARE V1 TO V00
VAR. VALUE
                    START VALUE
                                     SCALING FACTOR
                                                     REL.STAND.DEV
     2.01820948E+04 2.01874462E+04 2.01874462E+04
                                                      2.60519742E-02
V1
     -2.90936164E+01 -2.90969472E+01 -2.90969472E+01
V2
                                                       1.41576516E-02
V11
     -2.18127453E+04
V12
      1.55559524E+01
v15
      2.36701148E+04 2.37258719E+04 2.37258719E+04 1.00528931E-01
V16
     -7.56541897E+00 -7.64417807E+00 -7.64417807E+00 3.17218507E-01
     3.00342429E+03 3.02274918E+03 3.02274918E+03 2.60675550E-01
v17
     2.20133190E+04 2.14534347E+04 2.14534347E+04
V19
                                                      6.92923849E-01
v20
     -6.72498276E+00 -6.31334932E+00 -6.31334932E+00 1.79242248E+00
NUMBER OF OPTIMIZING VARIABLES : 7
ALL OTHER VARIABLES ARE FIX WITH THE VALUE ZERO
THE SUM OF SQUARES HAS CHANGED FROM 1.51360340E-01 TO 1.44420287E-01
DEGREES OF FREEDOM 18. REDUCED SUM OF SQUARES 8.02334928E-03
```

If such a significant RSD has a value larger than 1, then this typically means that you are using too many optimizing variables. However, the RSD values also depend on the weighting of the experiments, so it is sometimes possible to reduce the RSD by changing weights.

The RSD should not be too low either. If one or more of them is very close to 0, then this suggests that you are trying to optimize the system with too few optimizing variables. You can add optimizing variables interactively using <u>"ENTER\_PARAMETER" on page 137</u> in GIBBS. However, it is recommended that you add a sufficient number of variables in the setup file from the start so that do not need to do this. You can fix variables to a value of 0 until you discover that they are needed.

Besides the RSD, you should also keep an eye on the variable values themselves (shown in the VALUE column, next to the leftmost column in the table shown in the preceding screenshot). If the value of a variable is in the order of 1E5 or higher and this variable has been put in the temperature-independent part of an interaction parameter's Gibbs energy expression, then you may not have the right weights on your experiments or you are using too many optimizing variables. This is also typically the case if a variable with a value in the order of 10 or more (per mole atoms) is in the temperature-dependent part of the Gibbs energy expression of an interaction parameter. Such a high parameter value may lead to inverted miscibility gaps or re-stabilisation of a phase at high temperature. It is important to keep an eye on this since the temperature-independent and the temperature-dependent interaction parameter often varies together. If the temperature-dependent interaction parameter cannot be reliably determined, you may have to fix its value.

Note that even if you have found a good enough fit between calculated and experimental values after an optimization run, a parameter value may suddenly start to change by several orders of magnitude when you use <u>"OPTIMIZE\_VARIABLES" on page 148</u> is used again. If this happens, then careful reconsideration of the weighting of the experiments of the set of optimizing variables is required. However, when the optimization has converged and the parameter values remain stable after repeated use of OPTIMZE\_VARIALBES, this should usually not happen.

## **Reducing the Number of Optimizing Variables**

If you have one or more optimizing variables with RSD larger than 1, then you should remove one or more of the variables by setting them to a value of 0, or alternately, set them to a reliably estimated value (estimated based on, for example, semi-empirical methods).

To fix an optimizing variable to certain value, use the PARROT command SET\_FIX\_VARIABLE <variable number> <value>.

## **Plotting Intermediate Results**

It is typically not sufficient to look at the plain text output of <u>"LIST\_RESULT" on page 142</u> to reliably judge how the optimization is progressing. Plotting a phase or property diagram of the system you are optimizing using the POLY and POST modules is often very useful.

It is recommended that you create a macro file for plotting the diagram since you probably want to repeatedly plot the diagram as you progress with the optimization. This macro file can then be called directly from the PARROT module with the <u>"MACRO\_FILE\_OPEN" on</u> page 5 command.

### Superimposing Experimental Data from EXP-files

If you create one or several EXP-files based on the experiments in your POP-file, you can visually compare the fit your plot and the experimental data you are optimizing the system against. An EXP-file contains data points specified in the DATAPLOT format. You could make an EXP-file for each type of data that you are using in the optimization.

Use the POST command <u>"APPEND\_EXPERIMENTAL\_DATA" on page 236</u> to superimpose the data points in an EXP-file on a phase or property diagram that you have plotted. For example, to put the experimental data points from a file entitled DP.EXP onto a plotted diagram, you could enter the following:

```
APPEND EXPERIMENTAL DATA Y DP.EXP 0 1
```

Such a command call tells Thermo-Calc to superimpose the data points in dataset 1 of DP.EXP onto the plot in the Console Results window, but not to impose any information about axis scaling, labels and other information which is normally found in the prologue of the DP.EXP file.

The POST command <u>"QUICK\_EXPERIMENTAL\_PLOT" on page 245</u> can also be used to superimpose data points on a plotted phase or property diagram.

DATPLOT User Guide included with this documentation set.

# Finalizing the Optimization

### In this section:

Finishing the Optimization	43
Rounding off Optimizing Variable Values	43
Updating the Database File	. 43
Creating a Database File	44
Updating the Setup File and the POP-file	. 44

## **Finishing the Optimization**

When to stop optimizing is a matter of judgement. It is rarely the case that you finish optimizing a system with the feeling that it cannot be improved.

The final step is to update your files with the final results. Besides updating your database file (or alternatively, creating a new database file), it is advised that you also update your setup and POP-files with the final weights and optimizing variable values.

## **Rounding off Optimizing Variable Values**

When you save the optimizing variable values, it is important that you round off the values correctly. When you have a metallic system, it is usually best to keep whatever number of decimal digits that you need to make less than one J/mol difference at 1000 K. When you have an aqueous system or a system that you are optimizing based on data from very different temperature ranges, then this is not always the only appropriate guideline. However, rounding off the values in such a way that it gives a difference that is larger than one J/mol may lead to differences in the phase diagram.

Another approach you can take when rounding off variable values is by progressively fixing the value of more and more of the variables. This allows you to round-off all but one of the variables values so that each only has a few significant digits. Variables whose values have been rounded off in this way are easier to handle compared with those that that are rounded off so that they continue to have many significant (non-zero) digits.

To round off the variable values according to this second approach, first use <u>"SET\_FIX\_VARIABLE" on page 151</u> to set the variable(s) with the highest RSD to a rounded off value (such as, say, 0.4). When you then re-optimize using <u>"OPTIMIZE\_VARIABLES" on page 148</u>, the sum or errors changes. However, you should get almost exactly the same sum of errors as before after you rescale the variable values using <u>"RESCALE\_VARIABLES" on page 149</u>. (If the sum of errors is different, then this means that the variable with the highest RSD was not rounded off in a good way.) If you successfully round off the first variable value, then continue to round of the optimizing variable which now has the highest RSD. Continue doing this until you have fixed all but one of the optimizing variables to their rounded-off values. The final sum of errors that you get after having fixed the variable values should not deviate significantly from the initial sum of errors you had before starting to round-off the variable values.

# **Updating the Database File**

To update the database file (with filename extension TDB) open it in a text editor and enter or update each of the parameters that you have optimized.

For example, suppose you have the following command in your setup file:

ENTER\_PARAMETER L(IONIC\_LIQ,AL+3,ZR+4:O-2;0) 298.15 +V17; 6000 N

If you then end up with a value for optimizing variable V17 of -100000, then you should enter the following under the IONIC\_LIQ phase in the database file:

```
PARAM L(IONIC_LIQ,AL+3,ZR+4:0-2;0) 298.15 -100000; 6000 N REF !
```



If you often need to round-off the optimizing variable value in some way, see "Rounding off Optimizing Variable Values" on the previous page.

When you have updated or entered the parameters that you optimized, save the database file.

# **Creating a Database File**

In some circumstances you may want to save all the information about your system into a new database file. To do this, go to the GIBBS module and use LIST\_DATA <Filename> N. This creates a new user database file that contains all the information about the system that is in the workspace.

The N-argument tells the <u>"LIST\_DATA" on page 115</u> command to save the output into database file rather than some other type of file, such as a macro file for example. Note that the output database file that you get from LIST\_DATA have to undergo some manual editing before it can be used as proper database file.

# Updating the Setup File and the POP-file

It may be useful for you to be able to recreate your optimization from scratch. It is therefore recommended that you do the following when you are done with the optimization:

- Update your POP-file so that it contains the weights of your critical set of experiments. Use <u>"SET\_WEIGHT" on page 65</u> to set the weights directly in the POP-file. You may also have to use <u>"SET\_START\_VALUE" on page 216</u> for some conditions in the POP-file. Otherwise, you may not be able to compute the equilibria with <u>"COMPUTE\_ALL\_EQUILIBRIA" on page 36</u>.
- Copy the final values of your optimizing variables and set them as start values of the variables in the setup macro file (with <u>"SET\_FIX\_VARIABLE" on page 151</u>).
- It is important that you round off the values of the optimizing variables correctly. See <u>"Rounding off Optimizing Variable Values" on the previous page</u>.
   When you have done this, you can run the setup macro file to recreate the final result of the optimization.

## Alternate Mode

### In this section:

About Alternate Mode	46
Optimizing in Alternate Mode	47
Preparing the POP-file for Alternate Mode	. 47
Examples of SET_ALTERNATE_CONDITION	47
Entering Other Experiments in the POP with Alternate Mode	48

## **About Alternate Mode**

To turn Alternate Mode ON or OFF, use the PARROT command <u>"SET\_ALTERNATE\_MODE" on</u> page 150.

When you have one or more multi-phase equilibria (that is, with the fixed phase-status) in your POP-file and the optimizing values lack appropriate starting values (such as the default value of 0 for example), then Thermo-Calc may not be able to compute the equilibria for the experiments. In these cases, a common tangent cannot be found, that is, a state where the chemical potentials for all components are the same in all phases cannot be found. A common tangent may not exist or it be found at the wrong composition or temperature given the initial set of model parameters.

If you are not able to estimate reasonable starting values for the optimizing variables, then this problem can be overcome by initially running your optimization in alternate mode. The alternate mode in PARROT module should help you find reasonable starting values for the optimizing variables. Note that you are only supposed to use alternate mode in the beginning of the optimization. Once you have achieved a reasonable fit between calculated results and experimental data, turn off alternate mode and calculate all equilibria again in the normal mode.

In alternate mode, PARROT modifies multi-phase equilibria so that the chemical potentials for each stable phases is calculated separately. For example, consider the following experimental equilibrium:

```
CREATE_NEW_EQUILIBRIUM 1 1
CHANGE_STATUS PHASE LIQ FCC=FIX 1
SET_CONDITION T=1000 P=1E5
EXPERIMENT X(LIQ,B)=.2:.01 X(FCC,B)=.1
```

When PARROT optimizes in alternate mode, the computation of this equilibrium is done by way of the following two calculations:

```
CHANGE_STATUS PHASE LIQ=FIX 1
SET_CONDITION T=1000 P=1E5 X(LIQ,B)=.2
CHANGE_STATUS PHASE FCC=FIX 1
SET CONDITION T=1000 P=1E5 X(FCC,B)=.1
```

The difference between the chemical potentials in every pair of phases is calculated and PARROT tries to adjust the parameters of the phases so that these differences are minimized. In other words, PARROT tries to make the chemical potentials equal in all phases. For PARROT to be able to do this, some additional information about the experimental multi-phase equilibria is required, at least for all invariant equilibria.

## **Optimizing in Alternate Mode**

When you are optimizing in the alternate mode, you usually do not need to enter the ED\_EXP module and manually change the weights of experiments that the computed results have not been fitted well with. Instead, after using "OPTIMIZE\_VARIABLES" on page 148 initially with no iterations, and listing the result with "LIST\_RESULT" on page 142, simply run OPTIMIZE\_VARIABLES again specifying, say, thirty iterations. Continue to optimize until your optimization seems to be going in the right direction, then set the optimizing variables' starting values to their current values use "RESCALE\_VARIABLES" on page 149 before you turn alternate mode off by answering N at the "SET\_ALTERNATE\_MODE" on page 150 prompt.

# **Preparing the POP-file for Alternate Mode**

Information in your POP-file that concerns equilibria with ENTERED or DORMANT phases is ignored when you optimize in alternate mode.

To use the alternate mode, ensure that there is enough information about each phase in a given equilibrium. In addition to the temperature and pressure of the system, you should specify the composition of all phases in a given equilibrium. This is done with, <u>"SET\_</u><u>CONDITION" on page 209</u>, <u>"EXPERIMENT" on page 43</u>or <u>"SET\_ALTERNATE\_CONDITION" on page 55</u>.

If both compositions of a binary tie-line have been measured, then you could specify this in the POP-file in the following way:

```
CREATE_NEW_EQUILIBRIUM 1 1
CHANGE_STATUS PHASE FCC BCC=FIX 1
SET_CONDITION P=1E5 T=1000
EXPERIMENT X(BCC,B)=.2:.01 X(FCC,B)=.3:.01
```

This equilibrium could be calculated with alternate mode without any modification. The compositions specified with the experiment command is used as condition when the thermodynamic properties of each phase are calculated.

## **Examples of SET\_ALTERNATE\_CONDITION**

The following topics demonstrate how <u>"SET\_ALTERNATE\_CONDITION" on page 55</u> is used to prepare different kinds of equilibria for optimization in alternate mode.

### SPECIFYING THE COMPOSITION OF ONE SIDE OF A BINARY TIE-LINE

If only one side of the tie-line has been measured, then you must provide an estimate of the

composition of the other phase. You can make such an estimate with SET\_ALTERNATE\_ CONDITION. This command has no effect unless alternate mode is turned on. The equilibrium described could thus be specified in the following way:

```
CREATE_NEW_EQUILIBRIUM 1 1
CHANGE_STATUS PHASE FCC BCC=FIX 1
SET_CONDITION P=1E5 T=1000
EXPERIMENT X(BCC,B)=.2:.01
SET ALTERNATE CONDITION X(FCC,B)=.3
```

When the alternate mode is turned on, the composition given by the SET\_ALTERNATE\_ CONDITION command is used for the FCC phase. The properties of the BCC phase is calculated using the composition specified by experiment as a condition.

#### SPECIFYING THE COMPOSITIONS OF A THREE-PHASE EQUILIBRIUM

If you have a three-phase equilibrium in your POP-file, then one of its compositions can be specified as experimentally determined, while two can be provided as alternate conditions:

```
CREATE_NEW_EQUILIBRIUM 1 1
CHANGE_STATUS PHASE FCC BCC LIQ=FIX 1
SET_CONDITION P=1E5
EXPERIMENT T=912:5 X(LIQ,B)=0.2:.02
SET ALTERNATE CONDITION X(FCC,B)=0.1 X(BCC,B)=.4
```

#### SPECIFYING THE COMPOSITION OF A STOICHIOMETRIC PHASE

When you specify the composition of stoichiometric phase with SET\_ALTERNATE\_ CONDITION, the value must be given with at least seven decimal digits, as in the following example:

```
CREATE_NEW_EQUILIBRIUM 1 1
CHANGE_STATUS PHASE LIQ A2B=FIX 1
SET_CONDITION P=1E5 X(LIQ,B)=0.2
EXPERIMENT T=992:5
SET_ALTERNATE_CONDITION X(A2B,B)=.66666667
```

### **Entering Other Experiments in the POP with Alternate Mode**

The following table shows some additional examples of how to enter different kinds of experiments in the POP-file when you are using the alternate mode.

Example	ED_EXP commands
Two-phase equilibrium: The melting temperature of an Au- Cu alloy	CHANGE_STATUS PHASE LIQUID FCC=FIX 1 SET_CONDITION X (FCC,CU)=0.14 P=1E5 EXPERIMENT T=970:2 SET_ALTERNATE_ CONDITION X(LIQUID,CU)=0.16
Invariant equilibrium experiment: A three- phase equilibrium in a binary system	CREATE_NEW_EQUILIBRIUM 1 1 CHANGE_STATUS PHASE FCC BCC LIQUID=FIX 1 SET_CONDITION P=1E5 EXPERIMENT T=912:5 SET_ALTERNATE_CONDITION X (FCC,B)=0.1SET_ALTERNATE_CONDITION X(BCC,B)=0.4SET_ ALTERNATE_CONDITION X(LIQ,B)=0.2
Ternary system, with two compositions measured (both with uncertainty 0.02)	CREATE_NEW_EQUILIBRUM 1 1 CHANGE_STATUS PHASE FCC BCC=FIX 1 SET_CONDITION T=1273 P=1E5 SET_CONDITION X (FCC,B)=0.1:0.02 EXPERIMENT X(FCC,C)=0.12:.02 SET_ ALTERNATE_CONDITION X(BCC,B)=0.17 SET_ALTERNATE_ CONDITION X(BCC,C)=0.07

## Troubleshooting

During an optimization, you are likely to run into many different problems and challenges. This topic gives some general guidelines that can help avoid common mistakes and problems.

### In this section:

Useful Guidelines	51
Excluding and Including the Correct Equilibria	52
Conflicting Data	52

# **Useful Guidelines**

The following rules of thumb are generally good to adhere to when you are planning and executing an optimization:

- Do not use data from thermochemical tables of unknown origin.
- Use experimentally determined properties; avoid converted quantities.
- Carefully estimate the accuracy of the experiments.
- Correct systematic errors (by adjusting a temperature scale for example).
- Use negative information such as, for example, the information that a phase should not be stable in a composition or temperature region.
- With a hundred activity measurements in a system but only ten composition points from the phase diagram, it is often appropriate to decrease the weights on the activity experiments.
- If you initially exclude any intermediate phases to only optimize the liquid and the terminal or end-member phases for the pure components, then it is often useful to compute a metastable phase diagram with just these phases. In the metastable phase diagram, the metastable solubility lines should not have any strange kinks or turns. Such a diagram is often also useful to compute at later stages in the optimization.
- When you have fitted the liquid phase and some solution phases reasonably well, then you can fix the optimizing variables that characterise these phases. You can then go on and optimize the variables for the intermediate phases.
- Phases with miscibility gaps are always difficult to optimize. Try to keep control of the miscibility gaps by using real or estimated experimental information. (Note that it is not possible to calculate the top of a miscibility gap as a single equilibrium.)
- Phases with order/disorder transformation are often difficult to optimize. It may be difficult to find the right start values on the optimizing variables. The TABULATION module (see <u>"TABULATION\_REACTION Commands" on page 303</u>) can be useful sometimes. Properties for specified site fractions can be calculated in the TABULATION module. This is not possible in the POLY module since this module only calculates for equilibrium site fractions. The ordered state may also disappear during the optimization of these phases. If this happens, you may be able to add an experiment that controls the state of order.
- Only use the alternate mode to find an initial set of model parameters that can then be make it possible to calculate the experimental equilibria in the normal mode.
- When you have achieved a satisfactory fit between calculated results and experimental data with an appropriate number of optimization variables, perform a couple of final optimization runs to make sure that your results are stable.

• Make sure that the values of the optimizing variables keep within a reasonable range. If a variable starts to change several orders of magnitudes, then you must consider whether you have the right weights on the experiments and whether you are using too many optimizing variables.

### **Excluding and Including the Correct Equilibria**

Use as few experiments as possible initially to get a reasonable overall fit. It is generally best to first focus on the invariant equilibria, as well as metastable states that can be estimated by excluding some phases. Crucially, any experiments for which the computed results are clearly faulty should be excluded from the final critical set, while all important invariant equilibria are calculated and included in this critical set.

If you cannot compute some invariant equilibria with intermediate phases, then it may be best to first exclude those phases from the optimization and only optimize the liquid and the most important solution phases. After you have obtained reasonable results for these phases, you can enter the intermediate phases back in again. With the variables for the optimized liquid and solid phases set fixed, you can finally optimize the intermediate phases.

### **Conflicting Data**

It is important that you do not include conflicting sets of data at the same time during an optimization. After all, if they are genuinely conflicting, then at least one of the sets must be incorrect. If you have conflicting sets of data, then only use one of these sets at a time together with the rest of the data. Often, the optimization makes it clear which dataset coheres best with the other information that is available about the system.

Sometimes, conflicting data cannot be detected directly. For example, you may have activity data which is inconsistent with solubility data from the phase diagram. This would be indicated by large errors in the fit when both the activity data and the solubility data are included. To find these inconsistencies, let alone correct the errors, you may thus have to optimize with some datasets excluded.

### **Experiments and Experimental Data**

These topics contain examples of different kinds of equilibria in different kinds of systems. They are intended to help you figure out how to use various kinds of experimental data when creating POP-files. The topics that follow contain examples how to enter various kinds of experimental data into a POP-file.

#### In this section:

Phase Diagram Data	54
Data for Individual Compounds	55
Single Phase Mixing Enthalpies or Partial Enthalpies	57
Enthalpies of Formation and Enthalpies of Reactions	57
Chemical Potentials via EMF Measurements	58
Driving Force for Metastable Phases	58
Pressure-Temperature-Volume EOS Data	59
Crystal Structure, Lattice Parameters and Site-Occupancy	59
Magnetism and Other Atomistic Properties	60
Data About Systems of Different Orders	61

### **Phase Diagram Data**

P-T-X phase diagram data for binary and ternary systems, or systems of even higher order can be used in the POP-file. This includes data of the following types:

- Data about the temperature- and composition-dependence of various properties (obtained from differential thermal analysis (DTA) measurements for example), including enthalpy, lattice parameter, dilatometric length, electronic conductivity and magnetic susceptibility;
- Data about reaction types (invariant/monovariant) and phase relations obtained from qualitative or quantitative metallographic studies;
- Data about phase amounts or compositions obtained from microprobe measurements (using X-ray spectroscopy or transmission electron microscopy (TEM));
- Data about the positions or directions of two-phase tie-lines and three-phase equilibria for ternary systems.

### EXAMPLE

The following table shows a POP-file excerpt that demonstrates how some kinds of phase diagram data can be entered:

Example	ED_EXP commands
Solidus (melting temperature) of an Au- Cu alloy	CREATE_NEW_EQUILIBRIUM 1 1 CHANGE_STATUS PHASE LIQUID FCC=FIX 1 SET_CONDITION X(FCC,CU)=0.14 P=1E5
	EXPERIMENT T=970:2
Three-phase equilibrium in a binary A-B system	CREATE_NEW_EQUILIBRIUM 1 1 CHANGE_STATUS PHASE FCC BCC LIQUID=FIX 1 SET_CONDITION P=1E5 EXPERIMENT T=912:5
Congruent transformation in a binary A-B system	CREATE_NEW_EQUILIBRIUM 1 1 CHANGE_STATUS PHASE BCC LIQUID=FIX 1 SET_CONDITION P=1E5 X(BCC,B)-X(LIQ,B)=0 EXPERIMENT T=1213:10

Example	ED_EXP commands	
	An uncertainty factor is set not only for the experimental measurement, but also for the X(FCC,B) condition. The factor is 0.02.	
Tie-line in a ternary A-	CREATE_NEW_EQUILIBRUM 1 1	
B-C system	CHANGE_STATUS PHASE FCC BCC=FIX 1	
	SET_CONDITION T=1273 P=1E5 X(FCC,B)=0.1:0.02	
	EXPERIMENT X(FCC,C)=0.12:.02	

# **Data for Individual Compounds**

You can use data about individual components in the POP-file. Such data includes activity, heat capacity, heat content, entropy, Gibbs energy of phase transformations, heat of transition and heat of melting.

Example	ED_EXP commands
	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE FCC_A1=FIX 1
Coulou outivity data in the	CHANGE_STATUS PHASE GRAPHITE=D
Carbon activity data in the fcc phase	SET_REFERENCE_STATE C GRAPHITE,,,,
	SET_CONDITION P=101325 T=1273 X(MN)=0.03
	SET_CONDITION X(C)=0.03
	EXPERIMENT ACR(C)=0.29:5%
	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE IONIC_LIQ=FIX 1
	CHANGE_STATUS PHASE GAS=D
Ln(activity coefficient for O)	SET_CONDITION X(0)=0.02 T=1523 P=101325
	SET_REFERENCE_STATE O GAS,,,,
	ENTER_SYMBOL FUNCTION LNFO=LOG(ACR(O)/X(O));
	EXPERIMENT LNFO=-1.5:0.01
	CREATE_NEW_EQUILIBRIUM 1 1
Entropy at 298.15 K of Cu2O	CHANGE_STATUS PHASE CU2O=FIX 1
	CHANGE_STATUS PHASE CUO=FIX 0

Example	ED_EXP commands
	SET_CONDITION P=101325 T=298.15
	EXPERIMENT S=92.36:1
	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE SPINEL=ENT 1
	CHANGE_STATUS PHASE FCC O2GAS=DORM
	SET_REFERENCE_STATE NI FCC,,,,
Gibbs energy of formation of NiAl2O4	SET_REFERENCE_STATE AL FCC,,,,
	SET_REFERENCE_STATE O O2GAS,,,,
	SET_CONDITION P=101325 T=1000 N(NI)=1 N(AL)=2
	SET_CONDITION N(O)=4
	EXPERIMENT GM=-298911:5%
	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE SPINEL=ENT 1
Light compains of Mara 204	SET_CONDITION P=101325 N(FE)=2 N(MG)=1 N(O)=4
Heat capacity of MgFe2O4	SET_CONDITION T=800
	ENTER_SYMBOL FUNCTION CP=H.T;
	EXPERIMENT CP=207:5%
	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE SPINEL=ENT 1
	SET_CONDITION P=101325 T=298.15 N(MN)=1 N(FE)=2
	SET_CONDITION N(O)=4
	ENTER_SYMBOL VAR H298=H;
Heat content (H-H298) for	ENTER_SYMBOL FUN HMNFE204=H-H298;
MnFe2O4	CREATE_NEW_EQUILIBRIUM 2 1
	CHANGE_STATUS PHASE SPINEL=ENT 1
	SET_CONDITION $P=101325 \text{ N}(MN)=1 \text{ N}(FE)=2 \text{ N}(O)=4$
	SET_CONDITION T=400
	EXPERIMENT HMNFE204=16610:5%

# **Single Phase Mixing Enthalpies or Partial Enthalpies**

You can use calorimetric data for mixing in the POP-file. This data could be the result of, for example, drop calorimetry or scanning calorimetry. This includes data about, for example, enthalpy of mixing of liquids, series of mixing-enthalpy and about partial enthalpy. Information about enthalpies of mixing can also be derived from theoretical principles using special quasi-random structures (SQSs).

Example	ED_EXP commands
Enthalpy of mixing in the liquid state of an Au-Cu alloy	CREATE_NEW_EQUILIBRIUM 1 1 CHANGE_STATUS PHASE LIQUID=FIX 1 SET_CONDITION T=1379 P=1E5 SET_CONDITION X(LIQUID,AU)=0.0563 SET_REFERENCE_STATE AU LIQ * 1E5 SET_REFERENCE_STATE CU LIQ * 1E5
	EXPERIMENT HMR=-1520:200

# **Enthalpies of Formation and Enthalpies of Reactions**

You can use calorimetric data for phase transformations in the POP-file. This data could be the result of, for example, direct-reaction calorimetry, solution calorimetry or combustion calorimetry. Information about enthalpies of formation can also be derived from theoretical principles.

Example	ED_EXP commands
	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE CORUNDUM ZRO2_TETR=FIX 0
Enthalpy of melting of the	CHANGE_STATUS PHASE IONIC_LIQ=ENT 1
eutectic in the Al2O3-ZrO2 binary (J/g)	SET_CONDITION P=101325 N=1 LNAC(O)=-80
	SET_REFERENCE_STATE ZRO2 TETR,,,,,,
	SET_REFERENCE_STATE AL203 CORUND,,,,,,
	EXPERIMENT HWR=1080:90
Enthalpy of formation of o-	CREATE_NEW_EQUILIBRIUM 1 1
CaZrO3 from the component oxides	CHANGE_STATUS PHASE ZRO2_MONO HALITE CAZRO3_O=ENT 1

Example	ED_EXP commands
	SET_CONDITION T=298.15 P=101325 N(CAO)=.5
	SET_CONDITION N(ZRO2)=.5 LNAC(O)=-20
	SET_REFERENCE_STATE CAO HALITE,,,,
	<pre>SET_REFERENCE_STATE ZRO2 ZRO2_MONO,,,,</pre>
	EXPERIMENT HMR(CAZRO3_0) =-15960:5%

### **Chemical Potentials via EMF Measurements**

You can use chemical potentials and activities/activity coefficients from electromagnetic field (EMF) measurements in the POP-file.

#### EXAMPLE

Example	ED_EXP commands
	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE FCC CU2O=FIX 1
Emf data on Cu	CHANGE_STATUS PHASE O2GAS=DORM
(s)/Cu2O(s)	SET_CONDITION P=101325 T=1000
	SET_REFERENCE_STATE O O2GAS,, 100000
	EXPERIMENT MUR(0) = -95387:1000

# **Driving Force for Metastable Phases**

You can use chemical driving forces for metastable phases in the POP-file.

Example	ED_EXP commands
To prevent BCC_A2 to be stable	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE FCC_A1 M7C3 M23C6=FIX 1
	CHANGE_STATUS PHASE BCC_A2=D
	SET_CONDITION P=101325 T=1473
	EXPERIMENT DGM(BCC) <-0.1:0.1

Example	ED_EXP commands
	CHANGE_STATUS PHASE BCC M7C3 GRAPHITE=FIX 1
To promote CEMENTITE to be stable	CHANGE_STATUS PHASE CEMENTITE=DORMANT
	SET_CONDITION P=P0 T=973
	EXPERIMENT DGM(CEM)>0.1:0.01

### **Pressure-Temperature-Volume EOS Data**

You can use Pressure-Temperature-Volume Equation of State (EOS) data in the POP-file. Such data includes, for example, molar volume, density, thermal expansion and bulkmodulus/compressibility.

Example	ED_EXP commands
	CREATE_NEW_EQUILIBRIUM 1 1
Volume of liquid Pt	CHANGE_STATUS PHASE LIQUID=ENT 1
	SET_CONDITION P=101325 N=1 T=1873
	EXPERIMENT VM=1.00758E-5:5%
	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE FCC_A1=ENT 1
Thermal expansion	SET_CONDITION P=101325 N=1 T=400
	ENTER_SYMBOL FUNCTION ALPHA=VM.T/VM/3;
	EXPERIMENT ALPHA=2.35E-5:5%

# **Crystal Structure, Lattice Parameters and Site-Occupancy**

You use information about defects, ordering or site occupancy, as well as information about crystal structures in the form of lattice parameter values. This information may come from measurements made by X-ray, neutron diffraction, Mössbauer spectroscopy or perturbed angular-correlation.

Example	ED_EXP commands
Lattice parameter for fcc	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE FCC_A1=ENT 1
	SET_CONDITION P=101325 N=1 T=298.15
	SET_CONDITION X(CR)=0.05
	<pre>ENTER_SYMBOL FUNCTION LPFCC=((4*VM/6.02214179E23)** (1/3))*1E10;</pre>
	EXPERIMENT LPFCC=4.02:5%
Lattice parameter for bcc	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE BCC_A2=ENT 1
	SET_CONDITION P=101325 N=1 T=298.15
	SET_CONDITION X(AL)=0.08
	ENTER_SYMBOL FUNCTION LPBCC=((2*VM/6.02214179E23)**(1/3))*1E10;
	EXPERIMENT LPBCC=2.90:5%
Site-occupancy, degree of inversion for spinel	CREATE_NEW_EQUILIBRIUM 1 1
	CHANGE_STATUS PHASE SPINEL=ENT 1
	SET_CONDITION $P=101325 \text{ N}(MG)=1 \text{ N}(AL)=2 \text{ N}(O)=4$
	SET_CONDITION T=1073
	EXPERIMENT Y(SPINEL,AL+3#1)=0.31:5%

# **Magnetism and Other Atomistic Properties**

You can use data about atomistic properties such as magnetism and Curie temperatures in the POP-file.

Example	ED_EXP commands
	CREATE_NEW_EQUILIBRIUM 1 1
Curio tomporaturo of	CHANGE_STATUS PHASE SPINEL=ENT 1
Curie temperature of MnxFe3-xO4, x=1.26	SET_CONDITION T=520 P=101325 N(O)=4
	SET_CONDITION N(FE)+N(MN)=3 N(MN)=1.26
	EXPERIMENT TC(SPINEL)=520:5

# **Data About Systems of Different Orders**

You might want to use data about systems of different orders in the same optimization, such as data about a binary equilibrium with components A and B in a ternary system with components A, B and C, together with data about equilibria in the binary system with components A and B. To do this, you must first suspend all components in the ternary system (A, B and C), and then set the two components you want binary information on (A and B) to status ENTERED.

Suspending all components is done by giving the initialization code 0 to the <u>"CREATE\_NEW\_</u> <u>EQUILIBRIUM" on page 185</u> command. The components about which you want to use binary information must then be entered again using the <u>"CHANGE\_STATUS" on page 96</u> command.

Example	ED_EXP commands
A binary (A-B) three-phase equilibrium (FCC-BCC-LIQ) in a ternary system (A-	CREATE_NEW_EQUILIBRIUM 1 0
	CHANGE_STATUS COMPONENT A B = ENTERED CHANGE_STATUS PHASE FCC BCC LIQ=FIX 1
B-C)	SET_CONDITION P=1E5
	EXPERIMENT T=1177:10

### **Thermodynamic Models**

The most common models used in assessments are described briefly in these topics. Examples of how to write the phase definitions in the setup-file are also given for each of these models. In addition to the models described here, a large number of thermodynamic models for various phases in different states have been implemented.

#### In this section:

Gas	63
Compound Energy Formalism	63
Substitutional Liquid	64
Ionic Two-Sublattice Liquid Model	64
Models for Ordered Phases	65
4SL Model for FCC, HCP and BCC	66
2SL Model for FCC, HCP and BCC	67
Magnetic Ordering	67
Molar Volumes and High Pressure	68
Excess Models	69

# Gas

The gaseous mixture phase is usually treated as a substitutional phase without sublattice. In the gaseous mixture phase, there are usually molecules formed, and the number of constituents of the gas phase is often much larger than the number of elements. These complex constituents have to be defined as species. The ideal gas model may handle a gas phase under low pressures (and low temperatures). This implies that the P-V-T relations and thermodynamic properties of pure gaseous species are calculated as for ideal gas, and that there is interaction between gaseous species in the mixture. In a defined system, there may only exist one gaseous mixture phase. Type code G is used for the gas phase:

ENTER PHASE GAS G, 1 H, H101, H102, H2, H201, H202, O, O2, O3; N N

### **Compound Energy Formalism**

If the atoms are sufficiently different in size, electronegativity or charge, they may prefer different types of sites in the lattice of crystalline solids. In some cases, a solute atom may even occupy interstitial sites between the normal lattice sites. All such phenomena are treated in Thermo-Calc by the sublattice concept. The Compound Energy Formalism (CEF) has proved to be the most general formalism for many different types of solution phases. It is therefore by default applied to various solid solution phases in the Thermo-Calc software package. CEF can also take ionic constraints (charged cation/anion species) into account. For a comprehensive description on the CEF, please see Hillert (2001), Frisk and Selleby (2001), Sundman and Ågren (1981).

Since the CEF is used by default, no special notation is needed in the phase definition, except for ionic phases, where type code I is given.

### EXAMPLES

Examples of phase definitions in a setup-file using the CEF:

```
ENTER_PHASE SPINEL I, 4 1 2 2 4
FE+2,FE+3; FE+2,FE+3,VA; FE+2,VA; O-2; N N
ENTER_PHASE SIGMA, 3 10 4 16
AL,CO,CR,FE,NB; AL,CO,CR,FE,NB; AL,CO,CR,FE,NB; N N
ENTER_PHASE FCC_A1, 2 1 1
CR,FE,MN; C,N,VA; N N
ENTER_PHASE BCC_A2, 2 1 3
CR,FE,MN,VA; C,N,VA; N N
```

# Substitutional Liquid

The liquid phase is normally (but not always) modelled as a substitutional phase with no sublattice, which is occupied by two or more constituents. The constituents are normally the elements, but molecules or associates can also be used. Type code L is used for the liquid phase.

### EXAMPLE

```
ENTER_PHASE LIQUID L, 1
AL,C,CR,FE,MO,NI,SI ; N N
ENTER_PHASE LIQUID L, 1
C,FE,FEO,FEO3/2,NI,NIO ; N N
```

# **Ionic Two-Sublattice Liquid Model**

Within the framework of the CEF, the ionic two-sublattice liquid (I2SL) model was developed to be used when there is a tendency for ionization in the liquid, which happens in liquid oxides and sulphides for example. The same model can be used both for metallic and oxide melts. At low levels of oxygen, the model becomes equivalent to a substitutional solution model between metallic atoms. Two sublattices are assumed, one containing charged cations and one containing charged anions, neutrals and vacancies. The complication for an ionic liquid is that the numbers of sites on the cation sublattice and anion sublattice vary with the composition to preserve electroneutrality. For a comprehensive description on the I2SL model, see Hillert et al. (1985), Sundman (1991). In these references, the possible excess parameters are also discussed. Type code  $\underline{x}$  is used for the I2SL model.

### EXAMPLE

Examples of phase definitions in a setup-file:

```
ENTER_PHASE LIQUID Y,

AL+3,CR+2,FE+2,NI+2; VA; N N

ENTER_PHASE IONIC_LIQ Y,

CA+2,MG+2; O-2; N N

ENTER_PHASE IONIC_LIQ Y,

AL+3,CA+2,FE+2,MG+2,SI+4;

ALO2-1,O-2,SIO4-4,VA,FEO3/2,SIO2; N N
```

### **Models for Ordered Phases**

Some solution phases have chemical order-disorder transformations: for example, the disordered FCC\_A1 phase can transform to the ordered  $L1_2$  or  $L1_0$  structure where the atoms of different kinds occupy different sublattices. In some cases, it may be advantageous and easy to describe such ordered structures as completely different phases; but in some other cases, like the B2-ordered BCC structure in Al-Fe, the disordered and ordered structures must be described as one singe phase because the transformation is of second order along a line in temperature and composition.

The ordering can easily be described by the CEF. For instance, an ordered FCC phase with only  $L1_2$  or  $L1_0$  ordering can be efficiently handled by the so-called Two Substitutional-Sublattice Ordering Model (2SL model), while an ordered FCC with both  $L1_2$  and  $L1_0$  ordering must be described with four sublattices using the so-called Four Substitutional-Sublattice Ordering Model (4SL model). Please note that in both these models, all the substitutional constituents must enter into these sublattices. In addition, you may also have one sublattice for interstitials.

Thermo-Calc supports a feature of splitting the parameters of a chemically ordered phase onto two different phase descriptions. This is very convenient. In this way, you can have one phase for all parameters describing the disordered state (the configuration independent state) and in the other phase only those parameters needed to describe the remaining part (the configuration dependent part), that is, the Gibbs energy contribution due to the ordering transformation. This has been implemented in such a way that the contribution from the configuration dependent part is zero when the solution phase is disordered. The optimization of the disordered phase is independent of the order-disorder transformation; the ordering energy is an add-on to the disordered part. This is particularly useful in higher order systems, as an ordered phase can dissolve several elements that have no particular contribution to the actual ordering. The phases Gibbs energy add together by giving this command in the setup-file: AMEND\_PHASE\_DESCRIPTION <ordered phase> disordered\_ part <disordered phase>

Since PARROT does not automatically create new composition sets, it is necessary to create composition sets for both the ordered and disordered phases in the setup-file: Amend\_phase\_description <phase> composition\_sets <new highest set number>,,,,

### EXAMPLE

Example of phase definitions for the corresponding disordered and ordered parts in a setupfile:

> ENTER\_PHASE FCC\_A1, 2 1 1 AL,CR,FE,NI; VA; N N

```
ENTER_PHASE FCC_L12, 3 .75 .25 1
AL,CR,FE,NI ; AL,CR,FE,NI; VA; N N
AMEND_PHASE_DESCRIPTION FCC_L12 DIS_PART FCC_A1
```

### **4SL Model for FCC, HCP and BCC**

Ordered FCC, HCP and BCC solution phases handled by the 4SL model requires four sublattices for substitutional ordering and can additionally have an interstitial sublattice. A 4SL model is especially useful for modelling different kinds of ordered phases that are based on the same disordered phase, such as  $L1_0$  and  $L1_2$  based on FCC\_A1, and B2,  $D0_3$ ,  $L2_1$  and B32 based on BCC\_A2 etc.

For ordered FCC or HCP phases, these four substitutional sublattices represent four corners of the regular tetrahedron on these lattices, all of which are the nearest neighbours. These corners are equivalent lattice points, thus all G parameters for each end-member with the same elements, but distributed on different sites, must be identical. It should be emphasized that the end-member energy here represents the ordering energy rather than the formation energy of the compound.

The constraints on the parameters in the 4SL model can be derived based on the symmetry of the lattice:

 $G_{A:B:B:B} = G_{B:A:B:B} = G_{B:B:A:B} = G_{B:B:A:B} = G_{B:B:B:A}$   $G_{A:A:B:B} = G_{A:B:A:B} = G_{A:B:A:B} = G_{B:A:A:B} = G_{B:A:A:B} = G_{B:A:A:A} = G_{A:B:A:A} = G_{A:B:A:A} = G_{A:A:A:B} = G_{A:A:A:B} = G_{A:A:A:B} = G_{A:A:A:B} = L_{A,B:*:*} = L_{*:A,B:*} = L_{*:*:A,B}$ 

The asterisk \* means that the interaction parameter is independent of the occupation of that sublattice. For the disordered phase to be completely disordered, i.e. that all site fractions are equal on all four sublattices, all constraints must be correct.

For ordered BCC phases, the situation is a bit more complicated, as the four sublattice ordering phase represents an irregular tetrahedron with two pairs of sites that are next nearest neighbours. Thus, for an A-B binary solution phase, with A located on two sublattice sites and B on the other two, the end-member described by  $G_{A:A:B:B}$  has four nearest neighbour bonds between A and B atoms, whereas the end-member described by  $G_{A:B:A:B}$  has two nearest neighbour bonds between A and B atoms and two next nearest neighbour bonds. Many parameters thus have a relation:

 $\mathsf{G}_{\mathsf{A}:\mathsf{B}:\mathsf{B}:\mathsf{B}}{=}\mathsf{G}_{\mathsf{B}:\mathsf{A}:\mathsf{B}:\mathsf{B}}{=}\mathsf{G}_{\mathsf{B}:\mathsf{B}:\mathsf{A}:\mathsf{B}}{=}\mathsf{G}_{\mathsf{B}:\mathsf{B}:\mathsf{B}:\mathsf{A}}$ 

 $\mathsf{G}_{\mathsf{B}:\mathsf{A}:\mathsf{A}:\mathsf{A}}{=}\mathsf{G}_{\mathsf{A}:\mathsf{B}:\mathsf{A}}{=}\mathsf{G}_{\mathsf{A}:\mathsf{A}:\mathsf{B}:\mathsf{A}}{=}\mathsf{G}_{\mathsf{A}:\mathsf{A}:\mathsf{A}:\mathsf{B}}$ 

 $G_{A:A:B:B}=G_{B:B:A:A}$ 

 $G_{A:B:A:B} = G_{A:B:B:A} = G_{B:A:A:B} = G_{B:A:B:A}$ 

 $L_{A,B:*:*:*} = L_{*:A,B:*:*} = L_{*:*:A,B:*} = L_{*:*:A,B}$ 

The  $G_{A:B:B:B}$  term represents D0<sub>3</sub>,  $G_{A:A:B:B}$  B2 and the  $G_{A:B:A:B}$  B32 ordering. There are also two kinds of reciprocal interaction parameters:

L<sub>A,B:A,B</sub>:\*:\*;0...9

L<sub>A,B:\*:A,B:\*;0...9</sub>

Kusoffsky et al. (2001) have shown the influence of different parameters available in the 4SL model applied to fcc ordering. The possibility to use the 4SL model to BCC ordering has been studied by Sundman et al. (2009) in the Al-Fe system.

### **2SL Model for FCC, HCP and BCC**

The reason for using a 2SL model instead of a 4SL model is that the calculations are faster, but on the other hand it can only model one kind of ordered phase(s). For instance, it is not possible to model FCC with both  $L1_0$  and  $L2_1$ , or BCC with D0<sub>3</sub> and B2, etc.

In order to get the fully disordered phase to form, constraints are needed as for the 4SL model. For the symmetric phases, e.g. B2 and  $L1_0$ , the relation between parameters are:

G<sub>A:B</sub>=G<sub>B:A</sub>

 $L_{A,B}$ :\*=L\*:A,B

For a 2SL asymmetric model including, but not limited to,  $L1_2$ , more constraints are needed. This model has many complicated relations between the parameters. The relation between the parameters for the 2SL  $L1_2$  model can be derived from a 4SL model. The conversion from 4SL model parameters to 2SL parameters for  $L1_2$  can be found in Dupin et al. (2001) and for  $L1_0$  in Yuan et al. (2012). Higher-order systems with an  $L1_2$  phase modelled with two sublattices require a lot of ternary and some quaternary interaction parameters in order to make the disordered state stable. These parameters have been derived by e.g. Dupin (1995), also found in Kusoffsky et al. (2001).

Many publications exist on how to model order/disorder transformations using a twosublattice model, see e.g. Dupin, Ansara (1999), De Keyzer et al. (2009).

### **Magnetic Ordering**

The magnetic contribution to the Gibbs energy is given by a model proposed by Inden (1975) and adapted by Hillert and Jarl (1978), and is described by:

G=RTln(b+1)f(t)

where  $t=T/T_C$ .

The values of the Curie temperature, T<sub>C</sub>, and Bohr magneton number, b, at a certain composition are then used in an optimization procedure to calculate the contribution to Gibbs energy due to magnetic ordering. Curie temperature and Bohr magneton number are given in the command <u>"ENTER\_PARAMETER" on page 137</u> for phases that undergo a magnetic transition:

```
AMEND_PHASE_DESCRIPTION BCC_A2 MAGNETIC -1.0 0.4
ENTER_PARAMETER TC(BCC_A2,FE:VA;0) 298.15 +V1; 6000 N
ENTER_PARAMETER BMAGN(BCC_A2,FE:VA;0) 298.15 +V2; 6000 N
```

The arguments -1 and 0.4 specify the values of parameters in the magnetic model: -1 is the antiferromagnetic factor and 0.4 is the structure factor. Usually the antiferromagnetic factor =-1 for bcc and =-3 for fcc (and other non-bcc phases). The structure factor corresponds to the short range order fraction of the enthalpy. This factor is 0.4 for bcc and 0.28 for non-bcc phases.

### **Molar Volumes and High Pressure**

Including volumes in a thermodynamic database enable calculations of volume changes, volume fraction, lattice mismatch in cubic structures and phase diagrams at increased pressure. Pressure independent volumes can be used up to about 1 GPa. At pressures above 1 GPa the pressure dependence of the volumes must be included by using a proper equation of state (EOS). An EOS has been developed by Lu et al. (2005), based on the EOS from Jacobs and Oonk (2000). Four parameters are used in the implementation of the model in Thermo-Calc:

- vo: Volume at 1 bar and at reference temperature T<sub>0</sub>
- VA: Integrated thermal expansivity
- VK: Isothermal compressibility at 1 bar
- vc: Parameter to fit high pressure data

All parameters can be composition dependent in the same way as all other model parameters in Thermo-Calc. Experimental information about the temperature, pressure and composition dependence of the volume can be assessed using these model parameters. To evaluate V0, VA and VK experimental data at 1 bar only are needed.

#### EXAMPLE

Examples on modelling molar volume in a setup-file:

ENTER\_PARAMETER V0(BCC\_A2,FE:VA;0) 298.15 +V1; 6000 N

```
ENTER_PARAMETER VA(BCC_A2,FE:VA;0) 298.15 +V2*T+V3*T**2; 6000 N
ENTER_PARAMETER VC(BCC_A2,FE:VA;0) 298.15 +V4+V5*T; 6000 N
ENTER_PARAMETER VK(BCC_A2,FE:VA;0) 298.15 +V6+V7*T; 6000 N
ENTER_PARAMETER V0(FCC_A1,CR,FE:VA;0) 298.15 +V8; 6000 N
ENTER_PARAMETER VA(FCC_A1,CR,FE:VA;0) 298.15 +V9*(T-298.15); 6000 N
```

### **Excess Models**

Unless specifically amended, the excess Gibbs energy terms are modelled by the Redlich-Kister-Muggianu Model. Other excess models for the Gibbs energy are also implemented in Thermo-Calc, such as e.g. Kohler and Toop-Kohler extrapolation models.

More information about different excess models can be found in <u>example</u> <u>tcex51</u> in the *Thermo-Calc Console Mode Examples Guide*.

Thermo-Calc is not restricted to binary interaction parameters. Ternary, quaternary and higher-order interaction parameters can also be used if there is enough information to evaluate these from experimental data. The ternary interaction parameters are identified by the degrees of the so-called ternary L parameter (i.e. 0, 1 and 2) in the command "ENTER\_PARAMETER" on page 137. If the parameter is composition-independent, you only need a single interaction parameter with a degree of zero, i.e. only the <sup>0</sup>L term. If there is a ternary composition-dependent interaction parameter, you must enter all three parameters (0,1 and 2). If such a ternary interaction parameters should be zero, then assign it a function which is zero. If the three interaction parameters have the same value, then that is identical to having a composition-independent ternary parameter.

#### **EXAMPLE**

```
ENTER_PARAMETER L(LIQUID,A,B,C;0) 298.15 +1000; 6000 N
ENTER_PARAMETER L(LIQUID,A,B,C;1) 298.15 +1000; 6000 N
ENTER PARAMETER L(LIQUID,A,B,C;2) 298.15 +1000; 6000 N
```

#### is identical to

ENTER PARAMETER L(LIQUID, A, B, C; 0) 298.15 +1000; 6000 N

#### This also means that

ENTER\_PARAMETER L(LIQUID,A,B,C;0) 298.15 +1000; 6000 N ENTER\_PARAMETER L(LIQUID,A,B,C;1) 298.15 +ZERO#; 6000 N ENTER\_PARAMETER L(LIQUID,A,B,C;2) 298.15 +ZERO#; 6000 N

#### is NOT identical to

```
ENTER_PARAMETER L(LIQUID, A, B, C; 0) 298.15 +1000; 6000 N
```

# **Pourbaix Diagrams**

**Educational Material** 





# **Introduction to Pourbaix Diagrams**

#### In this section:

About Pourbaix Diagrams in Thermo-Calc	3
Pourbaix Diagrams	3
Working with Pourbaix Diagrams	5
Key Concepts	6
Effective Interaction Ratio	6
Solubility	7
Basic Settings and Definitions	7

# **About Pourbaix Diagrams in Thermo-Calc**

This guide is for Thermo-Calc software users who know the basics of thermodynamics and want to know more about <u>"Pourbaix Diagrams" below</u>.

You are introduced to the concept of a Pourbaix diagram and shown how to interpret such diagrams through a series of examples.

Materials corrosion occurs almost everywhere. It may lead to serious material damages, unexpected application failures, tremendous economic costs and environmental degradations. Consequently, scientists and engineers must often conduct expensive and time-consuming corrosion experiments as part of failure analyses, risk evaluations, quality improvements and application enhancements.

Under certain conditions, when a metal or alloy is exposed to an aqueous solution with a concentration of inorganic/organic mixture, corrosion phenomena occur at a corresponding degree. During corrosion, some metallic phases dissolve, the metal or alloy surface gets damaged and some secondary solid phases form at the solid-liquid interfaces (such as oxides, hydroxides, silicates, sulphides, sulphates, carbonates, nitrates, phosphates, borates, or halides). Such corrosive chemical or electrochemical reactions can be studied by means of the so-called Pourbaix diagrams if the reactions reach their equilibrium states (Pourbaix, 1973<sup>1</sup>. 1974<sup>2</sup>, 2003<sup>3</sup>, 2011<sup>4</sup>; Thompson et al. 2011<sup>5</sup>.).

# **Pourbaix Diagrams**

Marcel Pourbaix applied thermodynamics to predict materials corrosion resistance. He determined the phase stability relations in terms of varied pH and Eh values for an interaction system of metal and pure water or dilute aqueous solution. He presented the stability regions of metal and secondary phases (such as metal-oxides/hydroxides) on a pH-Eh diagram, which is now known as a Pourbaix diagram.

A Pourbaix diagram is a kind of phase diagram that shows the stability boundaries for a metal-aqueous interaction system. The phase boundaries are shown as a function of pH (acidity) and Eh (standard hydrogen electronic potential). An aqueous solution phase is always present in such a system. At a given pH and Eh, a metal may lose its stability to a soluble or corrosive aqueous solution, or be in equilibrium with either the aqueous solution

<sup>1.</sup> Lectures on Electrochemical Corrosion. Plenum Press, New York.

<sup>&</sup>lt;sup>2.</sup> Atlas of Electrochemical Equilibria in Aqueous Solutions, 2nd Ed, Houston, Tex., National Association of Corrosion Engineers; Cramer and Covino.

<sup>&</sup>lt;sup>3.</sup> Corrosion: Fundamentals, Testing, and Protection. ASM Handbook, Volume 13A; Verink.

<sup>&</sup>lt;sup>4.</sup> "Simplified procedure for constructing Pourbaix diagrams", iln: *Uhlig's Corrosion Handbook, 3rd Edition (Revie RW, Ed.),* The Electrochemical Society and Jon Wiley & Sons, New Jersey

<sup>&</sup>lt;sup>5.</sup> "Pourbaix diagrams for multielement systems", in: *Uhlig's Corrosion Handbook, 3rd Edition (Revie RW, Ed.)*, The Electrochemical Society and Jon Wiley & Sons, New Jersey

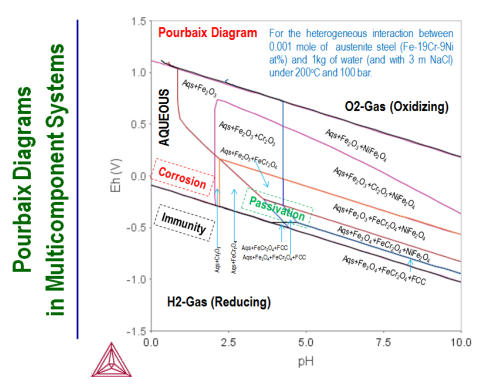
(insoluble/immune) or with a secondary-phase file that has formed (consisting of oxides, hydroxides, sulphides or other solids). In the latter case, further dissolution of the passive or protective metal is prevented.

The speciation and partition in the aqueous solution and the interacting phases depend not only on pH and Eh, but also on other factors such as the bulk composition, temperature and pressure in the system. The interacting phases may be gas mixtures, stoichiometric solids or solid solutions.

A Pourbaix diagram is divided in regions of *immunity, corrosion* and *passivity*. These regions provides information about the stability of a particular metal or alloy in a specific aqueous electrochemical environment under certain pH, Eh, pressure and temperature conditions.

- The immunity region is the region in which there is no metal dissolution.
- The corrosion region is the region in which there is active metal dissolution.
- The passivation region is the region in which a protective metal-oxide film that prevents metal dissolution is formed.

The following shows these three regions in a Pourbaix diagram for the heterogeneous interaction between 0.001 mole of austenite steel (Fe-19Cr-9Ni [at%]) and 1 kg of water (with 3 m NaCl), at 200° C and 100 bar:



Depending on the system's bulk chemical composition, secondary phases may form during the transformation process. These can not only be oxides, but also be, for example, hydroxides, sulphides, sulphates, carbonates, nitrates, phosphates, borates, silicates, hydrous silicates and halides. Information about these secondary phases may help one understand the passivation behaviours in corrosion processes. For example, it may be important to understand whether any secondary phases are formed in different pH-Eh regions during hydrolysis, oxidation, reduction or other reaction processes.

In a Pourbaix diagram, the predominant ion boundaries are often represented by dashed and straight lines imposed on a traditional Pourbaix diagram for a simple interaction system (Pourbaix, 1973, 1974; Cramer and Covino, 2003; Verink, 2011; Thompson et al., 2011). Such a predominance boundary is normally an equal molality (or equal activity) line for a pair of two most-concentrated aqueous solute species (charged or neutral) involving a common metal. A line is then simply determined by a reaction constant of its corresponding electrochemical reaction in the homogeneous aqueous solution phase. However, this is not the case in Thermo-Calc.



You cannot get any information about predominant ion boundaries from a Pourbaix diagram in Thermo-Calc.

# Working with Pourbaix Diagrams

With the Thermo-Calc software package, Pourbaix diagrams can be calculated over a wide range of pressure, temperature and composition conditions for complex heterogeneous interactions between multicomponent primary alloy solution or compound phases, concentrated aqueous solutions (with dissolved inorganic and organic substances), complicated secondary solids and oxidizing/reducing gaseous mixtures. In the Console Mode, diagrams can be calculated and plotted either using the response-driven POURBAIX module or directly using the DATA, GIBBS, POLY and POST modules.

Note that it is possible in Thermo-Calc to set aqueous properties other than pH and Eh as axis variables. For example, the following properties can also be set as axis variables: Ah (electronic affinity), pe (electronic activity log10ACRe), IS (ionic strength), TM (total aqueous concentration, in molality), oc (Osmotic coefficient), activity or activity coefficient of solvent H2O, and activities or activity coefficients or concentrations in m (molality) of charged or neutral solute species.

Beside a concentrated aqueous solution phase handled by a proper thermodynamic model named SIT, HKF or PITZ, it is also possible to consider complex alloy solution phases or compound phases, as well as secondary solid phases and gaseous mixtures, which are treated by specific solution models.

Most frequently, a Poubaix diagram is calculated for a specific alloyed phase with the initial alloying composition, such as the FCC solution phase in an austenite steel. However, it is possible to calculate a Pourbaix diagram for two co-existing phases, such as the FCC+BCC solution phases in a duplex steel for example, or even for more than two co-existing phases in a steel/alloy.

Note that each point on the lines in a pH-Eh diagram in Thermo-Calc represents a certain minimum state of Gibbs energy. When a line is crossed, a phase transformation should occur. This means that a Pourbaix diagram in Thermo-Calc does not provide any information about any predominant ion boundary.

# **Key Concepts**

The following are two key concepts important for understanding how aqueous solutions behave and how an aqueous-bearing heterogeneous interaction system is properly defined and calculated in Thermo-Calc.

### **Effective Interaction Ratio**

A pH-Eh plot is always related to a certain amount of initial alloys or other condensed materials that has effectively reacted with an aqueous solution in the system. This is the amount of condensed material that is fully in equilibrium with the defined aqueous solution phase. The amount is specified relative to an aqueous solution that is normally comprised of 1 kg of water with certain specified solute concentrations at certain temperature and pressure conditions. (This is why a calculated Pourbaix diagram is typically presented for an initial amount of the interacting metal or alloy at a certain level, such as 10-6, 1E-3, 0.1 or 1 mole of metal or alloy.) It is called the *Effective Interaction Ratio* (between the initial alloy or alloyed phases and the initial aqueous solution) and it is expressed in terms of molality (mol/kg).

The Effective Interaction Ratio is important for two reasons. First, the ratio has implications for kinetic or dynamic effects such as chemical reaction mechanism and kinetics, fluid flow dynamics, surface area and interaction time. Secondly, the ratio, being expressed as 10-6 mole of metal (or alloy), is the solubility limit that can be detected for cathodic corrosion protection by immunity.

One should always be careful when setting initial amounts and compositions in the original condensed materials, as well as when setting the initial concentrations of dissolved solutes in the original aqueous solution phase. It is often useful to make a series of calculations for different levels of initial amount of the interacting metal/alloy while the conditions are fixed for other settings (such as initial aqueous concentration, pressure, temperature, pH, and Eh).

### Solubility

When a heterogeneous equilibrium has a dissolving solution or mixture phase and a stoichiometric or solution phase, then the concept of solubility becomes important. A solubility of a phase (the solute) is its property of dissolving in the solvent phase. This concept concerns the constitution of a phase and is applied where one or several of the constituents are dominant (which is highly concentrated and dissolving) while there are only small amounts of the other remaining species (which are less concentrated and dissolved). The dissolving solution or mixture phase can be liquid, gas, aqueous or solid, as long as it has dissolving capacity. The stoichiometric or solution phase has some constituents which tend to be dissolved into the dissolving solution or mixture phase.

#### **KEY POINTS ABOUT SOLUBILITY**

- Under certain temperature, pressure and composition conditions, a Fe- or Crdominant BCC phase can dissolve certain amounts of Ni and C from a carbide phase such as M23C6, M7C3 or M3C. The Ni and C elements in the carbides have the solubility defined relative to the BCC phase.
- An aqueous solution phase is always dominant in the solvent water, that is, H2O. Under specific temperature, pressure and aqueous composition conditions, any other element (such as Fe and C) or substance (such as a pure SO2 gas, stoichiometric phase Cu2S and solution phase (Fe,Ni)1(O,Va)1) have a certain solubility limit in the defined aqueous solution.
- Under certain temperature and pressure conditions, and under given certain concentrations of other dissolved species in the mixture, an O2-dominant gaseous mixture phase can dissolve certain amounts of Fe+2 or Fe+3 species from magnetite (Fe3O4). The magnetite solid has solubility defined relative to the gaseous mixture under the given conditions.
- Under certain temperature and pressure condition, under specific concentrations of other dissolved species in the liquid phase, a Fe-dominant liquid mixture phase can dissolve certain amounts of, for example, Cr and O. The Cr and O components have the solubility defined in the liquid mixture.

### **Basic Settings and Definitions**

In a homogeneous aqueous solution or an aqueous-bearing heterogeneous interaction system, the most essential definition is for system-components there must be H2O, H+1 and ZE (electron) plus those for elements dissolved in aqueous solution (such as Na, C1, S) and associated in interacting metals/alloys (such as Fe, Cr, Mn, Mg, Ni, Al, Si, Zn). Three fundamental phases in a system are the AQUEOUS solution, the GAS mixture and the REF\_ELECTRODE. The REF\_ELECTRODE phase is used for setting the reference state for electrostatic potential in the system and for calculating the Eh condition (defined as MUR

(ZE) /RNF). Other phases should be appropriately selected and retrieved from criticallyassessed databases that cover not only the target phases (solution or stoichiometric) in interacting metals/alloys but also the secondary phases (solution or stoichiometric). The two public databases PAQ and PAQS are specially designed as single-database choices that cover all kinds of phases necessary for calculations. However, these databases are each limited to a framework of a very small number of elements. When it comes to simulations of complex multicomponent systems with a wide variety of elements and phases, the thermodynamic data must be selected and retrieved from several databases: AQUEOUS solution and REF ELCTRODE phases must be retrieved from TCAQ or AQS; alloy phases (such as FCC A1, BCC A2, HCP A3, CEMENTITE) from alloy solution databases (such as SSOL for general alloy phases, TCFE for steel/Fe-alloy phases, TCAL for Al-based alloy phases; TCMG for Mg-based alloy phases, and TCNI for Ni-based superalloy phases); gaseous mixture phase and secondary phases from specific substance or solution databases (such as SSUB for GAS phase and various solid compound phases or TCOX for oxide solution phases). Note that the REF ELECTRODE phase should always be suspended in equilibrium calculations, while GAS phase could be set as ENTERED, SUSPENDED or DORMANT, depending on the purpose of the calculation.

When defining an interaction system, the initial condition for the H2O component is always set as 1 kg of water. The initial composition conditions for dissolving and interacting elements are normally defined in moles (such as n (Fe) = 0.009, n (Cr) = 5E-4, n (Ni) = 3E-4, n (Mn) = 5E-5, n (S) = 5E-5, n (Na) = 3, n (C1) = 3). This makes it straightforward and convenient to count various related aqueous solution properties based on molality. The initial conditions for the H+1 and ZE components can be given as molar compositions (such as n (H+1) = 0, n (ZE) = 0) or their activities or potentials (such as lnACR (H+1) = -9.21, MUR (ZE) = 8400).

The pH and Eh properties of the aqueous solution in the interacting system are always defined in the following way:

```
pH = -log10[ACR(H+1,AQUEOUS)*AH2O]]Eh = MUR(ZE)/RNF
```

The symbol AH20 is the molecular weight of solvent H2O (equals 55.508435) and RNF is the Faraday constant (equals 96485.309).

The activity of the solvent water (ACRH2O, Aw), the osmatic coefficient of aqueous solution (OSMC, Os), electronic affinity (Ah), electronic activity log10ACRe (pe), ionic strength (Is), total aqueous concentration, in molality (TM) and total alkaline concentrations under two definitions (At1/AT2) are calculated and listed for each equilibrium state.

POLY3 calculations for mass balances in Thermo-Calc are always based on site-fractions. Consequently, when functions for describing various properties of aqueous solutes are defined, such as molality (MLi), activity coefficient (RCi) and activity (AIi), they should be converted to molality-based quantities:

MLi = Y(AQUEOUS,i)\*AH2O/YH2O RCi = ACR(i,AQUEOUS)\*YH2O/Y(AQUEOUS,i) AIi = ACR(i, AQUEOUS) \*AH2O

Here, YH20 is the site-fraction of solvent H2O and AH20 equals 55.508435.

Many more variables, functions and tables can be entered for various purposes. For instance, an equilibrium constant for a homogeneous reaction or a solubility product for a solid dissolution reaction can be entered.

# **Thermodynamic Data Requirements**

### In this section:

Required Thermodynamic Data	11
Including or Excluding the Gas Phase	11

# **Required Thermodynamic Data**

To calculate a Pourbaix diagram, thermodynamic data for at least the following four types of phases must be available:

- The aqueous solution phase which applies a certain aqueous solution model. In Thermo-Calc, information about this phase could be retrieved from, for instance, the TCAQ or AQS database (these also include data for the REF\_ELECTRODE phase that is used for setting the reference state of the standard hydrogen electrode).
- The alloy solution and/or compound phases for the primary matrix phases (and possibly also for the precipitated phases in the alloy. Examples of such phases include FCC\_A1, BCC\_A2, HCP\_A3, SIGMA and CEMENTITE. In Thermo-Calc, information about such phases could be retrieved from, for example, the TCFE database for steels/Fe-alloys, the TCNI database for Ni-based superalloys, the TCAL database for Al-based alloys, the TCMG database for Mg-based alloys, the TCSLD for Sn-/Au-/Bi-/Zn-based solder alloys or the SSOL for general alloys.
- The secondary solid phases that would form as a result of the heterogeneous chemical or electrochemical reactions. These phases could be, for example, oxides, hydroxides, silicates, hydrous silicates, sulphides, sulphates, carbonates, nitrates, phosphates, borates or halides. In Thermo-Calc, information about such phases could be retrieved from, for example, the SSUB database for pure solid compounds or the TCOX database for complex oxide solids.
- The gaseous mixture phase. Information about this phase could be retrieved from, for example, the SSUB database. However, note that is also possible to perform Pourbaix diagram calculations that ignore the gaseous mixture phase.

All this thermodynamic data about the various phases must be critically assessed and internally consistent. Furthermore, when the data is retrieved from several databases, the data taken from each database must be consistent with the data taken from the other databases. When Pourbaix diagrams and other diagrams of steel corrosion processes are calculated, it is typically recommended that the TCAQ, TCFE, SSUB and TCOX databases or the AQS, TCFE, SSUB and TCOX are used in combination.

The public PAQ database contains thermodynamic data for all four phase types and can be used for calculating Pourbaix diagrams. They have been designed specifically for the purpose of demonstrating the POURBAIX module functionality in Thermo-Calc. However, these databases only allow you to perform test calculations for simple cases with major phases in which only a few elements can be considered.

# **Including or Excluding the Gas Phase**

Under defined pressure and temperature conditions, the solvent water's thermodynamic stability limits are determined by the following two electrochemical reactions:

```
H2O(water) + 2e - = H2(gas) + 2OH - 1
2H2O(water) = O2(gas) + 4H + 1 + 4e -
```

The first reaction describes the formation of H2-dominated gaseous mixture, under reducing conditions. The second reaction describes the formation of O2-dominated gaseous mixture, under oxidising conditions.

If the system reaches global equilibrium, then the water component is electrolyzed to H+ and O-2 at all pH conditions. The degree of electrolysis depends on the pH value in the aqueous solution phase. If Eh gets high enough, then the O-2 anion is oxidized to O2 (aqs). On the other hand, if Eh gets low enough, then the H+ cation is reduced to H2 (aqs). The major electrolysis and redox reactions are the following:

```
H2O (water) = H+ + OH- \rightarrow Electrolysis of water at all pH.
H2O (water) = 2H+ + O-2 \rightarrow Electrolysis of water at all pH.
O-2 - 2e- = 0.502 (aqs)\rightarrow Oxidation or de-electronization of O-2 at high Eh.
2H+ + 2e- = H2 (aqs) \rightarrow Reduction or electronization of H+ at low Eh.
```

At a critically high Eh value under a given pH condition, an aqueous solution phase with a high enough O2 activity becomes less stable than an O2-dominated gaseous mixture phase. At this point, the gas phase replaces the aqueous solution phase in the system. The replacement process can be characterised by the following phase-transformation on the aqueous-gas phase boundary and oxidation of remaining water:

O2 (aqs) = O2 (gas) H2O (water) - 2e- = 2H+(gas) + 0.5O2 (gas)

Similarly, at a critically low Eh value under a given pH condition, an aqueous solution phase with a high enough H2 activity becomes less stable than a H2-dominated gaseous mixture phase. At this point, the gas phase replaces the aqueous solution phase in the system, through the following phase-transformation on the aqueous-gas phase boundary and reduction of remaining water:

```
H2 (aqs) = H2 (gas)
H2O (water) + 2e^- = 0^-2 (gas) + H2 (gas)
```

The phase transformation from an aqueous solution phase to an O2- or H2-dominated gaseous mixture phase also depends on the total molar Gibbs energies of the phases (which are complex functions of the phase constituents, the temperature and the pressure). The Gibbs energy minimization technique used in Thermo-Calc ensures that the phase transformation is accurately simulated.

The following Pourbaix diagrams show the result of a calculation where the gaseous mixture phase was included. In the upper region (high Eh) and lower region (low Eh), the aqueous solution phase has been oxidized and reduced, respectively, to an O2- or H2-dominated gaseous mixture phase.

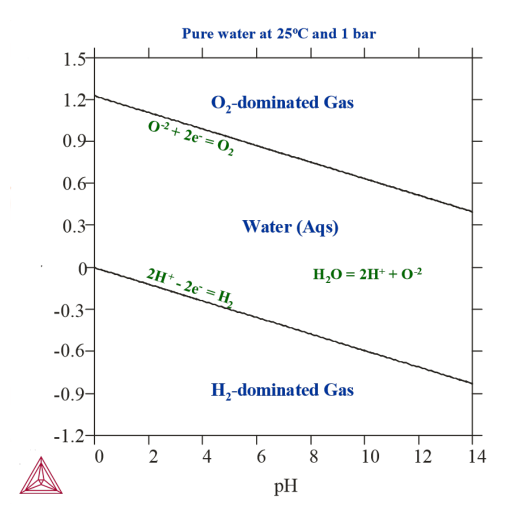
The two example diagrams show that the upper and lower boundaries between the aqueous phase (water) and the gas phase (dominated by either O2 or H2) can shift when solutes dissolve or when the temperature and pressure change.

Accordingly, if one does not take a gaseous mixture phase into account when performing a calculation of Pourbaix diagram (as above), then the aqueous solution phase may end up with an extremely high O2 (aqs) concentration at high Eh condition, or an extremely high H2 (aqs) concentration at low Eh condition. Under either of these two extreme circumstances, the concept of "aqueous solution phase" is no longer valid and consequently no proper aqueous solution model can actually be applied. Therefore, from a restrictive thermodynamic equilibrium point of view, one shall normally include a gaseous mixture phase in an aqueous-bearing heterogeneous interaction system. This is true for all types of equilibrium calculations for an aqueous-bearing heterogeneous interaction system, not only of Pourbaix diagram calculations.

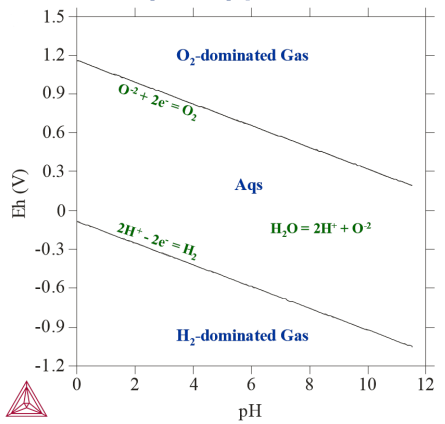
Hence, to perform a completely accurate Pourbaix diagram calculation, the gaseous mixture phase must be taken into account. However, since these electrochemical reactions normally have much higher kinetic barriers and are slower than other electrochemical and chemical reactions in the interaction system, one can sometimes ignore the gaseous mixture phase in the calculation. Note that if the gaseous mixture phase is ignored in this way, then the Pourbaix diagram does not show the H2O-stability limiting lines.

#### **EXAMPLE DIAGRAMS**

The first diagram is a pH-Eh diagram for pure water at 25° C and 1 bar:



The second diagram is a pH-Eh diagram for 3mNaCl-0.001mCO2-0.001mSO2 aqueous solution at  $150^{\circ}$  C and 100 bar.



3mNaCl-0.001mCO<sub>2</sub>-0.001m SO<sub>2</sub> aqueous solution at 150°C and 100 bar

### **Examples of Pourbaix Diagrams for Fe**

The following two topics are examples of Pourbaix diagrams for Fe. For the first example, the calculations on which the diagrams are based have not taken the gas phase into account. The diagrams in the second example are based on calculations that have taken the gas phase into account.

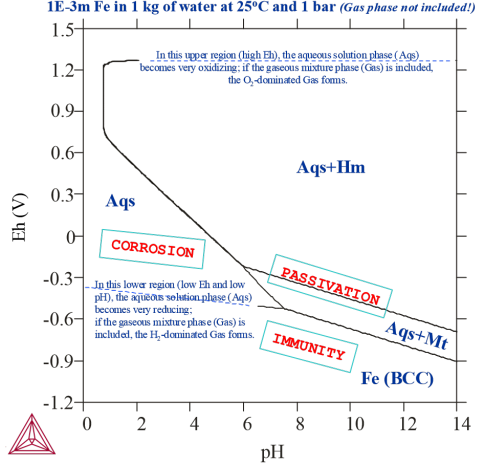
#### In this section:

Pourbaix Diagrams with Gas Phase Excluded	
Pourbaix Diagrams with Gas Phase Included	

# **Pourbaix Diagrams with Gas Phase Excluded**

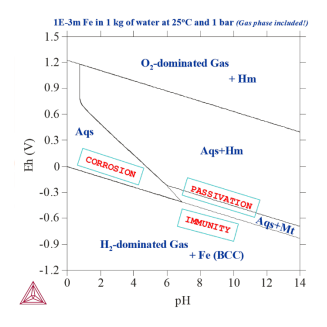
This is an example of a Pourbaix diagram of Fe, in which the gas phase has not been considered in the calculation. The system has 0.001 mole of Fe in 1 kg of pure water at a temperature of 25oC and a pressure of 1 bar. Magnetite (Mt, Fe3O4) and hematite (Hm, Fe2O3) co-exist with the Fe-containing dilute aqueous solution. The formation of magnetite and hematite represent the passivation of iron in the upper right pH-Eh field.

If pH is low and Eh is relatively high, then Fe eventually completely dissolves into the aqueous solution phase. Under conditions of low Eh, Fe remains in its stable solid state, the BCC phase, and neither dissolves into water nor into alternates to Fe (the immunity of Fe).



# **Pourbaix Diagrams with Gas Phase Included**

With all the possible redox reactions involving the aqueous solution phase and gaseous mixture phase being considered in the equilibrium system, that is, the gaseous mixture phase is included in the calculation, the Pourbaix diagram of Fe in a system with 0.001 mole of Fe in 1 kg of pure water at 25oC and 1 bar.



Comparing this Pourbaix diagram with that from <u>"Basic Settings and Definitions" on page 7</u>, you can see that both diagrams are for the same interaction system under the same temperature-pressure condition, but the present diagram shows the results of calculations where the gaseous mixture phase has been included. It presents the complete Pourbaix diagram for the defined interaction system, in a full thermodynamic equilibrium. In the upper region (high Eh) and lower region (low Eh), the aqueous solution phase is oxidised and reduced to the O2- or H2-dominated gas phase. Hematite coexists with the O2-dominated gas, while Fe-BCC is stable with the H2-dominated gas. The stability field of magnetite may extend slightly into the H2-dominated gas region, where it coexists with the gaseous mixture instead of the aqueous solution. Its boundary with Fe (BCC) cannot be drawn due to the absence of aqueous solution phase.

It is not possible to calculate the pH value in both the O2- and H2-dominated gaseous mixture phase stability regions where the aqueous solution phase is absent. Hence, in a normal Pourbaix diagram, no line is drawn that indicates a phase boundary between the gas phase and the metallic phases or secondary phases (that is, metal-oxides).

# **Other Examples of Pourbaix Diagrams**

#### In this section:

Variations of Pourbaix Diagrams	. 20
Pourbaix Diagrams for Complex Alloys	. 22
Pourbaix Diagram References	24

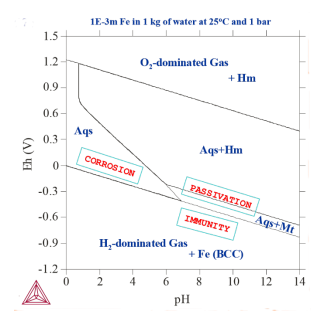
# Variations of Pourbaix Diagrams

The shape of a Pourbaix diagram of an alloy or condensed material and the stability relations of various secondary phases depend upon the following system factors:

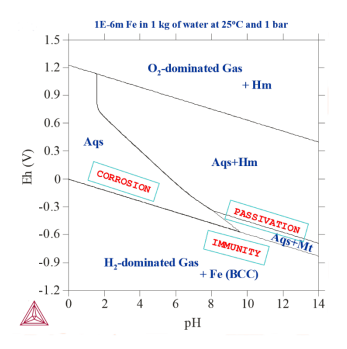
- Initial amount of the alloy or other condensed materials
- Initial composition of the alloy or other condensed materials
- Initial amount of the interacting aqueous solution phase
- Initial composition of the interacting aqueous solution phase
- Temperature and pressure conditions

In the interaction system that the diagrams in <u>"Pourbaix Diagrams with Gas Phase Excluded"</u> on page 17 and <u>"Pourbaix Diagrams with Gas Phase Included" on page 17</u> are based on, the initial amount of pure Fe that was taken to have effectively reacted with 1 kg of pure water at 25° C and 1 bar was 0.001 m. The following diagrams are Pourbaix diagrams of Fe where other initial amounts of pure Fe have been used, or where the interacting aqueous solution compositions have been alternated, or the temperature and/or pressure have been changed. Gaseous mixture phases have been included in all the calculations that these diagrams are based on.

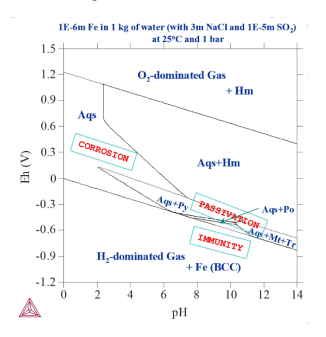
In the first diagram, 1E-3 m Fe actively reacted with 1 kg of pure water at 25° C and 1 bar (as in the calculation in Pourbaix Diagrams with Gas Phase Included):



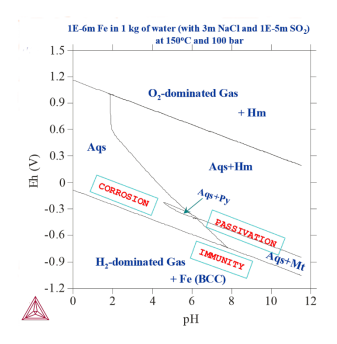
In the second diagram, 1E-6 m Fe actively reacted with 1 kg of pure water at 25° C and 1 bar. Note that the active metal corrosion region gets enlarged as the initial Fe amount decreases from 1E-3m to 1E-6m:



In the third diagram, 1E-6 m Fe actively reacted with 1 kg of water, 3m NaCl and 1E-5 m SO2 at 25° C and 1 bar. Introducing SO2 into the system leads to the formation of various metal-sulphides (Py-pyrite, Popyrrhotite, Tr-troilite). In addition, the passivation region becomes larger.



Finally, in the fourth diagram, 1E-6 m Fe actively reacted with 1 kg of water, 3m NaCl and 1E-5 m SO2 at 150° C and 100 bar. Here, changing the temperature and pressure affects the stability fields of various Fe-oxides/sulphides.

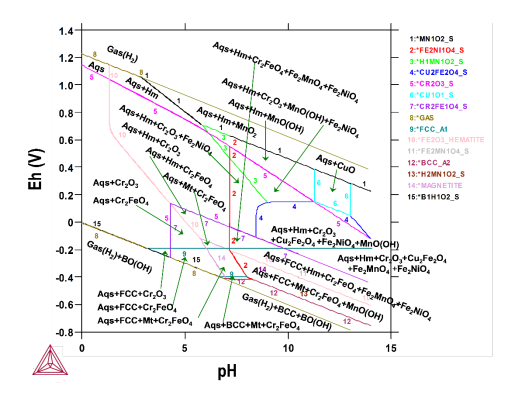


As you can see from the four diagrams, the aqueous-gas phase boundaries shift as the initial bulk compositions, pressure and temperature conditions in the interaction system change.

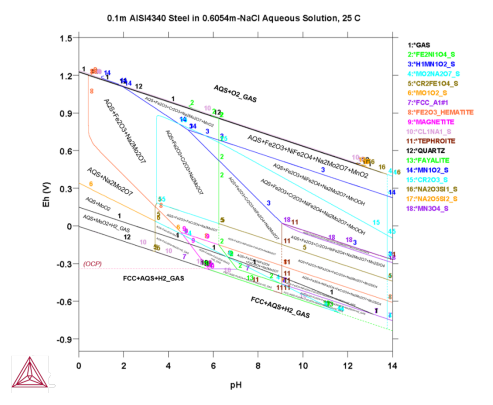
# **Pourbaix Diagrams for Complex Alloys**

Thermo-Calc can not only be used to simulate how pure metals interact with pure water or simple aqueous solutions under normal pressure and temperature conditions. It can also be used to calculate how complex alloys and concentrated aqueous solutions interact over a very wide pressure, temperature and composition ranges. This is illustrated with the two examples.

The first example is a Pourbaix diagram calculated for the heterogeneous interaction between 0.001 mole of steel (Fe-7.676Cr-5.0Ni-2.1887Mn-1.0Cu [at%]) and 1 kg of water (and with 1.2 m H3BO3, 0.022 m Li and 0.001 m NH3), at 25° C and 1 bar. This application is particularly useful for safety assessments of nuclear reactors and nuclear waste repositories.



The next example is a Pourbaix diagram calculated for the heterogeneous interaction systems between 0.1 mole of AISI4340 stainless steel (Fe-0.80Cr-1.85Ni-0.70Mn-0.25Mo-0.25Si-0.40C [wt%]) and 1 kg of seawater (with the equivalent of 0.6054 m NaCl), at 25° C and 1 bar. This application is particularly useful for failure analysis of petroleum exploitation industry.



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- 13. Shi, P., Engström, A., Sundman, B., and Ågren, J. (2011b). Thermodynamic calculations and kinetic simulations of some advanced materials. *Materials Science Forum*, 675-677: 961-974.
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- 15. Verink, E.D. (2011). Simplified procedure for constructing Pourbaix diagrams. In: *Uhlig's Corrosion Handbook, 3rd Edition (Revie RW, Ed.),* The Electrochemical Society and Jon Wiley & Sons, New Jersey.

# **DATAPLOT User Guide**

**Thermo-Calc Version 2018b** 



# Introduction to the DATAPLOT User Guide

#### In this section:

Using this Guide	. 3
About the DATAPLOT Graphical Language	. 3
DATAPLOT Examples	. 3
Plotting Experimental Data Files	4
Graphical Mode	. 4
Console Mode	. 4

# **Using this Guide**

The *DATAPLOT User Guide* gives all the details on the graphical language DATAPLOT that a phase diagram or property diagram calculated by Thermo-Calc or a kinetic profile simulated by the Diffusion Module (DICTRA) can be defined.

This guide

- Discusses the important features of the DATAPLOT language.
- Describes the commands to define PROLOGUE and DATASET.
- Several examples and the standard codes for formatting are provided.

One topic introduces the method of formatting DIGLIB symbols in LaTeX documents, for the purpose of necessarily/appropriately referring to the corresponding LaTeX symbols (closest to those DIGLIB symbols which have been plotted on a diagram using the DATAPLOT Graphical Language) in the texts of LaTeX documents for publications/reports.

# About the DATAPLOT Graphical Language

In order to obtain graphical output of any numerical data and informative strings, a graphical language called DATAPLOT was developed in connection with the graphical software DIGLIB. Using this graphical language, you can store information in a normal text file (\*.EXP), which can be plotted as graphical symbols, lines, texts or Greek letters on any plot device support by DIGLIB.

You can generate and plot DATAPLOT (\*.EXP) files together with various calculation and/or experimental results from, for example, phase and property diagrams calculated with the Thermo-Calc software composition profiles or other diagrams simulated by the Diffusion Module (DICTRA).

An EXP file automatically generated by the POST-processor (using the <u>"MAKE\_EXPERIMENTAL\_DATAFILE" on page 243</u> command) in the Thermo-Calc software is a DATAPLOT (\*.EXP) file and may contain all types of legal DATAPLOT commands and parameters. With a text editor, you can modify or add some DATAPLOT commands and related parameters in an existing EXP file. This is useful when appending experimental information to calculated/simulated plots, and when specifying user-defined texts, symbols, colors, fonts, filled patterns, diagram types, diagram sizes, symbol/character sizes, titles, special characters, etc.

#### **DATAPLOT Examples**

To help you prepare your EXP (experimental) files to use for your plotted diagrams generated by Thermo-Calc simulations, there are examples of EXP files using the DATAPLOT graphical language standard, which are normally independent of software versions.

# **Plotting Experimental Data Files**

You can read and plot experimental data files in both Console Mode and Graphical Mode.

#### **Graphical Mode**

In Graphical Mode, the Experimental File Reader activity can only be created from the top My Project node. This activity allows you to read an experimental data file (\*.EXP). This type of file has information specifying a plotted diagram, written in the DATAPLOT graphical language. From this activity node you can plot the EXP file or save a copy of the file to another location.

#### PLOTTING AN EXP FILE WITH THE EXPERIMENTAL FILE READER

- 1. Right-click **My Project** and from the **Create New Activity** submenu choose **Experimental File Reader**.
- 2. In the **Configuration** window in the **EXP file** field, enter the name of the EXP file to plot. Or click the **Select Exp file** button to navigate to a location on the computer.
- 3. Right-click the **Experimental File Reader** node and from the **Create New Successor** submenu choose **Plot Renderer**.

If the source EXP file is changed, you can also click the **Reload the selected EXP file from disk** button to refresh the data. If you want to save or export the loaded file, click the **Save EXP file** button, then navigate to where you want to save it and give it a file name.

4. In the **Configuration** window click **Perform** or right-click the **Plot Renderer** node and select **Perform**.

#### **Console Mode**

In Console Mode you use the QUICK EXPERIMENTAL PLOT command found in the POLY-3 POST module (as well as in the Diffusion Module (DICTRA) POST module) to plot the .EXP files. This command can be used when there is no graphical information to be plotted in the POLY3 workspace. It defines a pair of axes, sets the axis labels to X and Y, and scales both X-and Y-axes between 0.0 and 1.0 unless a prologue is read from the DATAPLOT (\*.EXP) data file.

#### PLOTTING AN EXP FILE IN THE POST MODULE

1. Go to the POLY-3 Module: At the SYS prompt type  ${\tt Go}~{\tt Poly}.$ 

- 2. Go to the POST module: At the POLY\_3 prompt type Post.
- 3. At the POST prompt, type Quick.
- 4. Enter Prologue and Dataset numbers (if applicable). See <u>"QUICK\_EXPERIMENTAL\_</u> <u>PLOT" on page 245</u>.
- 5. AT the POST prompt, type Plot.

# Using DATAPLOT Graphical Language

#### In this section:

DATAPLOT File Structure	7
DATAPLOT Language Syntax	7
Coordinate Systems	7
Graphical Operation Codes	8
Tables or Blocks	9
Drawing a Polygon	10
Drawing an Analytical Function	10
Painting of an Enclosed Area	10
Writing a Text	10
Plotting a Symbol	11
Other Commands	11
Interactive Plotting	12
Formatting DIGLAB Symbols in LaTeX Documents	12
LaTeX Formatting Codes	14
LTEXT vs LaTeX Commands for Labels	14

# **DATAPLOT File Structure**

A DATAPLOT file is a normal text file with the extension EXP that can be created with a text editor or by a program. The file must contain one or more DATASETS and possibly also one or more PROLOGUES. Each PROLOGUE/DATASET is an entity that can be individually selected for plotting.

A PROLOGUE/DATASET is identified by a unique positive number in the file. A PROLOGUE normally contains various DATASET commands for defining information about axis scaling, axis text, axis length, title and so on. A PROLOGUE is terminated by another PROLOGUE or by the first DATASET. This means that all PROLOGUES must be placed at the beginning of the file, before the first DATASET.

A DATASET normally contains DATASET commands that are associated to separate data points, as well as with one or more **BLOCKS** of data (calculated or experimental). A DATASET is terminated by another DATASET command or the end of file.

# **DATAPLOT Language Syntax**

The DATAPLOT language consists of commands with or without parameters:

```
COMMAND {parameter(s)}
```

The basic graphical command consists of an X/Y coordinate pair and a Graphical Operation Code (GOC). With other commands, the interpretation of this basic command can be modified. There are separate commands to draw a polygon or a function and ways to get texts in different fonts.

To edit a DATAPLOT file, the graphical commands can be abbreviated.



A command (with parameters) must not exceed 80 characters. If it is too long (normally as writing necessary codes in a command's parameters for a complex expression), two or more lines can be edited.

# **Coordinate Systems**

The DATAPLOT language accepts coordinates in three different coordinate systems, which are called *word*, *virtual* and *normalized*.

- The *word* coordinates are selected to represent any kind of data and be of "any" magnitude.
- The virtual coordinate system uses centimeters as units. However, the actual size of one unit is dependent on the implementation of the device driver in DIGLIB. It is not recommended to use this coordinate system if different output devices are used for preliminary and final plots.

• The normalized coordinate system goes from zero to one. When plotting, you interactively scale each axis by selecting the minimum and maximum word coordinates on the axis. In the normalized coordinate system, the minimum axis value is represented by zero and the maximum by one. DIGLIB draws a square between the four points (of the X and Y axes) that are determined by the coordinates zero and one in the normalized coordinates.



It is also possible to draw triangular plots as described below. However, in most places, it only references square diagrams. All data points within the minimum and maximum word coordinates are plotted inside this square. DIGLIB also writes tic marks and corresponding word values at such tic marks.

It may be convenient to use normalized coordinates to draw boxes and texts. You can give normalized coordinates outside zero and one if you want to write texts outside the area enclosed by the square. To ensure proper operation outside the normalized box, the clipping must be turned off.

# **Graphical Operation Codes**

The Graphical Operation Code (GOC) determines how the coordinates are interpreted and what is done at the point determined by the coordinates. For an individual data point, its GOC codes must be given. For each data <u>"BLOCK" on page 20</u>, you define a default GOC that is used for the whole BLOCK, unless a GOC is explicitly defined for a specific point.

- If s is omitted, no symbol is plotted. A number selecting the symbol can optionally follow the character s. The DIGLIB software determines which symbol the number represents. See <u>"Plotting a Symbol" on page 11</u>.
- If any of the other characters are omitted, the default defined for the BLOCK is used. Absolute values imply the current coordinates; relative values mean that these are added to the current coordinates. The GOC must be written without any space between various characters, but the order for the characters is irrelevant.
- If no GOC is defined for a BLOCK, the system default is MWA, (Move Word Absolute), i.e., move to the given point that is interpreted as absolute word coordinates. GOC=DNA means draw a line from the current point to the new one interpreting the coordinates as normalized. After each draw or move operation, the so-called current point is the new point.
- The <u>"TEXT" on page 29</u> following the ' command is interpreted and expanded by the DIGLIB software, if it includes any ^ operator or <u>"STRING" on page 27</u> names using the ~ descriptor. To get PostScript outputs, the TEXT must be edited by the LaTeX Text Formatting Program instead (see <u>"LaTeX Formatting Codes" on page 14</u>).

#### **GOC CHARACTER DEFINITIONS**

The GOC is a combination of the following characters:

Character	Meaning	
W, V Or N	To use word (w), virtual (v) or normalized (N) coordinates. w is the default.	
A <b>OF</b> R	XY are absolute (A) or relative (R) values. A is the default.	
M <b>Or</b> D	o perform a move (M) or draw (D) operation to XY. ${\mbox{\tiny M}}$ is the default.	
В	To apply "soft" splines on a line drawn between the coordinate pairs (used only on BLOCK data).	
S	To plot default symbol at XY.	
S#	To change default symbol to No # symbol and plot it at XY.	
'TEXT	To plot TEXT at XY. It must be the last operation in a GOC code. The TEXT following the ' command is interpreted and expanded by the DIGLIB software, if it includes any ^ operator or STRING names using the ~ descriptor.	

## **Tables or Blocks**

In many cases, you have (calculated or experimental) data in the form of tables, and want to plot one or several columns as X-axis and one or more columns as Y-axis. A single table can contain many sets of data records of the same kind of information for a specific data <u>"BLOCK" on page 20</u>. There can be many data BLOCKS in a <u>"DATASET" on page 23</u>.

By enclosing these tables in a data BLOCK, you can select which column(s) is the X-axis and which is the Y-axis. For example,

```
BLOCK X=C1; Y1=C3; Y2=C2; GOC=C4, DAW
```

Where the X-axis values are in column 1, the Y-axis values are in columns 2 and 3, and any GOC codes are in column 4. The default GOC for this BLOCK is Draw Absolute Word. The GOC code inside the table is only necessary if the default GOC is not applicable.

A more elaborate use of the table is shown in this example:

```
BLOCK X=C3; Y=1E3/(C3+273); GOC=C8,MAWS1
```

Where the X-axis values are in column 3, the Y-axis values are in column 1 with some additional calculations applied. Any GOC is in column 8, and the default GOC is Move Absolute Word and plot the symbol no. 1.

Columns in a table must be separated by one or more space characters and do not have to be justified.

It is possible to have tables with mixed text and numbers, but each word followed by a space is counted as one column. The columns used for plotting also must be numerical. An example of a legal line in a table is

298.15 This\_is\_the\_second\_column 11.5 This\_is\_the\_fourth\_column

A line in the table must not exceed 80 characters. A BLOCK must be terminated by a line with the "BLOCKEND" on page 21 command.

## **Drawing a Polygon**

8

Normally, each point is written on a separate line. But in order to draw a line in a more compact way, use the command <u>"DRAWLINE" on page 24</u>. DRAWLINE is followed by a couple of X/Y number pairs. The X/Y pairs must be separated by a space, and there must be a comma sign between the X and Y values. DRAWLINE makes a move operation to the first pair of X/Y coordinates, and then draws a line among all pairs up to the last one. All pairs must fit on one line of 80 characters, but there can be several consequent DRAWLINE commands.

"Example 2 – Draw Polygons and Symbols" on page 32

## **Drawing an Analytical Function**

Use a <u>"BLOCK" on page 20</u> command to set an axis to a function. It is not necessary to use a value from any column in order to compute the function value to be plotted. Use the <u>"FUNCTION" on page 25</u> command to plot a function with an even increment of the independent variable.

"Example 4 – Draw Curves Defined by Functions" on page 36

## Painting of an Enclosed Area

Use the <u>"PAINT" on page 26</u> command to paint or fill an area in a specified pattern in the plot. Available patterns are determined by the DIGLIB software. A related command to PAINT is <u>"PCFUNCTION" on page 27</u>.

## Writing a Text

Use the <u>"TEXT" on page 29</u> command to write a text at the current point. You can write a text at any X/Y pair by appending a single quote followed by the text on the same line. For example:

1.1 1.0 NAM'This is a text

```
writes This is a text at the normalized coordinates (1.1, 1.0).
```

Use the command <u>"FONT" on page 24</u> to select the font used for the text, and <u>"CHARSIZE" on page 21</u> to select the size of the characters. Set this size of the symbols with <u>"SYMBOLSIZE"</u> on page 29.

If a text or a single character should be of a different font than all the other text, or to use subscripts or superscripts in a text, use the  $\land$  operators or STRING command to create the text. The command <u>"STRING" on page 27</u> stores the text in a specified variable that includes all text-formatting information defined by the DIGLIB software.

<u>"Example 3 – Using Strings and Various Line Types" on page 34 and "Example 5</u> – Use Included Files for Predefined Symbols" on page 38.

However, if a graphical output is done on a PostScript device using the PostScript hardcopy fonts, special text formatting codes as <u>"LaTeX Formatting Codes" on page 14</u> should be used and the STRING formatting syntax is then not valid.

# **Plotting a Symbol**

As described in <u>"Graphical Operation Codes" on page 8</u>, a GOC code in a data BLOCK may contain an s option to plot a symbol for an X/Y pair or the same symbols for the data BLOCK. A number selecting the symbol can optionally follow the character s. You can plot a symbol at any current X/Y position by appending a quote specified by the symbol number in the GOC code (e.g., 1.1 1.0 MANS5' This is a text).

You can also insert DIGLIB symbols into LaTeX-edited documents.

<u>"Example 1 – Draw Lines and Symbols" on page 31, "Example 2 – Draw Polygons and Symbols" on page 32 and "Example 8 - DIGLIB Symbols" on page 51</u>

## **Other Commands**

#### General

- When plotting symbols representing various experimental data, it is important that the symbols are centered around the coordinate values.
- The dollar sign \$ as the first character of a line stands for a comment character, which means the whole line is ignored when plotting.

#### **ATTRIBUTE**

Use the <u>"ATTRIBUTE" on page 20</u> command to change the coordinates where text is displayed. The default is the lower left corner of the first character in the text.

#### CLIP

Use the <u>"CLIP" on page 21</u> command to change the default where all data outside the normalized coordinates zero and one are not plotted.

#### **COLOR**

If you are using a color device, use the <u>"COLOR" on page 22</u> command to change color of the lines. On some black and white devices, colors are simulated with different width and dashing of the lines.

#### **INCLUDE**

Use the <u>"INCLUDE" on page 25</u> command to create libraries with texts and include these in similar plots.

#### LINETYPE

Use the <u>"LINETYPE" on page 26</u> command to define line types as solid, long dashed, short dashed or dotted.

## **Interactive Plotting**

The DATAPLOT file is read into the POST module in the workspace with <u>"APPEND</u> <u>EXPERIMENTAL\_DATA" on page 236</u> or <u>"QUICK\_EXPERIMENTAL\_PLOT" on page 245</u>. These commands ask for the name of the DATAPLOT file and also which PROLOGUE(S) and DATASET(S) are to be plotted.

See the Thermo-Calc Command Mode Reference Guide for details about these commands.

By giving the PROLOGUE/DATASET number as -1, you get a list of the available PROLOGUES/DATASETS in the file. Note that if DATASET 0 is in a DATAPLOT file, its data is always used even if other DATASETS are chosen.

## Formatting DIGLAB Symbols in LaTeX Documents

When using the LaTeX editor, you can use the DIGLIB symbols in texts and in figures. This then refers to the corresponding LaTeX symbols (closest to those DIGLIB symbols which are plotted on a plot using the DATAPLOT Graphical Language) in the texts of LaTeX documents for publications/reports.

This topic shows how to generate some DIGLIB symbols in texts, through the attached LaTeX source file and its converted jpg file.

"Example 8 - DIGLIB Symbols" on page 51

DIGLIB\_SYM.TEX

```
\documentclass[dvips,12pt]{article}
\textwidth 165mm
\textheight 225mm
\oddsidemargin 1mm
\evensidemargin 1mm
\topmargin 1mm
%%\usepackage{amssymb}
%% next replace amssymb and to get udtimes
\usepackage[utopia] {mathdesign}
\usepackage{rotating}
\usepackage[latin1] {inputenc}
\usepackage{graphics}
\usepackage{graphicx,subfigure}% with figures
%\usepackage[draft]{graphicx}% without figures
\usepackage{subfigure}% with figures
\topmargin 1mm
\oddsidemargin 1mm
\evensidemargin 1mm
\begin{document}
{\Large \bf Diglib symbols and their corresponding LaTeX symbols}
\vspace{5mm}
```

The information below gives the closest corresponding LaTeX symbol. All symbols (except +) must be generated in math mode. Most of these require the package <code>amssymb</code>, i.e., you need a directive <code>usepackage{amssymb}</code> in the preamble. Two of the symbols require the more extensive <code>mathdesign</code> that can be included with <code>usepackage[utopia]{mathdesign}</code>.

```
\vspace{5mm}
{\Large
\begin{tabular}{1111}
Diglib & Latex & Latex name & Note\\
1& $\vartriangle$& $\backslash$vartriangle & amssymb \\
2& $\square$& $\backslash$square\\
3& {\Huge $\diamond$} & $\backslash$diamond & size $\backslash$Large \\
4& $\udtimes$& $\backslash$udtimes & mathdesign \\
5& $\triangledown$& $\backslash$triangledown & amssymb\\ 6 & +& normal + \\
7& $\ast$& $\backslash$ast & amssymb\\
8& $\times$& $\backslash$times \\
```

```
9& {\Huge $\circ$}& $\backslash$circ & size $\backslash$Large \\
10& {\Huge $\star$}& $\backslash$star & size $\backslash$Large, amssymb\\
11& $\curlyvee$& $\backslash$curlyvee & amssymb\\
12& $\Join$& $\backslash$Join\\
13&& - & nothing similar, overlapping $> <$\\
14&& - & nothing similar, 10-edged star \\
15& $\maltese$& $\backslash$maltese& mathdesign \\
16&& - & nothing similar, a pentagon \\
17& $\curlywedge$& $\backslash$curlywedge & amssymb\\
\\end{tabular}}</pre>
```

## **LaTeX Formatting Codes**



8

For a list of supported LaTeX commands, see: http://www2.ph.ed.ac.uk/snuggletex/documentation/math-mode.html.

By default when using the LaTeX command, you are in math mode and Thermo-Calc automatically interprets the entered text as math. The above link uses the dollar sign (\$) in all the examples but this is NOT required. For example, if you enter \$E=mc^2\$ this is displayed including the \$ instead of the equation.

As of Thermo-Calc 2015a the LTEXT text formatting is replaced by LaTeX. For existing users who may have plots and files that use the LTEXT format, and especially when using Console Mode, the following examples show the command changes from LTEXT and its LaTeX equivalent:

#### LTEXT vs LaTeX Commands for Labels

Symbol or text format	LTEXT	LaTeX
Subscripts in an axis text	S-A-TEXT Y N Mole-fraction Al^DO2\$O^DO3\$	S-A-TEXT Y N \latex Mole- fraction Al_2O_3
Label with subscript	add .13 .15 n c-ZrO^DO2\$	add .13 .15 n \latex c-ZrO_2
Greek symbol	ADD .05 .50 N ^GRd\$	ADD .05 .50 N \latex \delta

## **PROLOGUE Commands**

#### In this section:

PROLOG	
DIAGRAM_TYPE	16
TIC_TYPE	16
TITLE	16
XLENGTH	
XSCALE	17
XTEXT	17
ХТҮРЕ	17
YLENGTH	
YSCALE	
YTEXT	
YTYPE	

# PROLOG

PROLOG is the beginning of a number of consequent lines of user-defined diagram layout commands. The lines are displayed with the <code>optional text</code> when using the <code>-1</code> option in the POST-processor as prompted for the PROLOG number with <u>"APPEND\_EXPERIMENTAL\_DATA"</u> on page 236 or <u>"QUICK\_EXPERIMENTAL\_PLOT"</u> on page 245.



# Syntax PROLOGUE # optional text # is an unsigned integer identifying the PROLOG.

## DIAGRAM\_TYPE

DIAGRAM\_TYPE sets the diagram type to square (the default) or triangular.



This is available for Console Mode.

Example 6 – Triangular Diagrams for Ternary Systems" on page 40

Syntax	DIAGRAM_TYPE type plot_3rd_axis clip_along_third-axis
type is a character string reading SQUARE (default) or TRIANGUL If type reads TRIANGULAR, then two additional parameters are given: plot_3rd_axis and clip_along_third-axis that are characters strings reading YES or NO.	

# TIC\_TYPE

TIC\_TYPE sets the relative length of the tic marks.

Syntax	TIC_TYPE #
	# is a real number. Default value is 1. Negative number gives tics on the inside of the diagram frame. O gives no tics.

## TITLE

TITLE sets the title text string to be printed above the diagram.

Syntax	TITLE text
	${\tt text}$ is an arbitrary text string that may contain text-formatting codes

# **XLENGTH**

XLENGTH sets the X-axis length to approximately # centimeters.

Syntax	XLENGTH #
	# is a positive real number (the approximate X-axis length in centimeters).

## **XSCALE**

XSCALE sets the scaling in word coordinates of the X-axis.

Syntax	XSCALE min max
	min and max are real numbers.

## XTEXT

XTEXT sets the X-axis text.

Syntax	XTEXT text
	${\tt text}$ is an arbitrary text string that may contain text-formatting codes.

## **XTYPE**

XTYPE sets the X-axis type as linear (default), logarithmic or inverse.

Syntax	XTYPE type
	${\tt type}$ is a character string reading LIN, LOG or INV.

## **YLENGTH**

YLENGTH sets the Y-axis length to approximately # centimeters.

Syntax	Y LENGTH #
	# is a positive real number (the approximate Y-axis length in

Syntax	Y LENGTH #
	centimeters).

# **YSCALE**

YSCALE sets the scaling in word coordinates of the Y-axis.

Syntax	YSCALE min max
	min and max are real numbers.

## **YTEXT**

YTEXT sets the Y-axis text.

Syntax	Y TEXT text
	${\tt text}$ is an arbitrary text string that may contain text-formatting codes.

# **YTYPE**

YTYPE sets the Y-axis type as linear (default), logarithmic or inverse.

Syntax	Y TYPE type
	${\tt type}$ is a character string reading LIN, LOG or INV.

## **DATASET Commands**

#### In this section:

ATTRIBUTE	20
BLOCK	20
BLOCKEND	21
CHARSIZE	21
CLIP	21
COLOR	22
DATAPOINT	23
DATASET	23
DRAWLINE	24
FONT	24
Font Numbers	24
FUNCTION	25
GLOBALSIZE	25
INCLUDE	25
LINETYPE	26
Line Types	26
PAINT	26
PCFUNCTION	27
STRING	27
SYMBOLSIZE	29
TEXT	29

## ATTRIBUTE

ATTRIBUTE specifies where the current XY position is in the character or symbol plotbox.

- "Example 2 Draw Polygons and Symbols" on page 32, "Example 3 Using
  - <u>Strings and Various Line Types" on page 34</u>, and <u>"Example 5 Use Included Files</u> for Predefined Symbols" on page 38

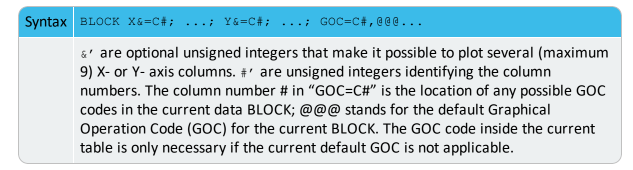
Syntax	ATTRIBUTE
Prompts	attribute
	attribute may be TOP, CENTER or BOTTOM. ATTRIBUTE CENTER is default for symbols; ATTRIBUTE BOTTOM is default for characters.
*	

# BLOCK

TOP

BLOCK defines how the following numeric data block shall be interpreted. The definitions of X and Y coordinates may also be expressed as a function of the column values, making it possible to perform transformations.

BOTTOM



#### **GOC CHARACTERS**

The GOC is a combination of the following characters:

CENTER

Character	Meaning
W, V <b>Or</b> N	To use word (w), virtual (v) or normalized (N) coordinates. w is the default.
A <b>Or</b> R	XY are absolute (A) or relative (R) values. A is the default.
м <b>ог</b> D	To perform a move (M) or draw (D) operation to XY. ${\mbox{\tiny M}}$ is the default.
В	To apply "soft" splines on a line drawn between the coordinate pairs (used only on BLOCK data).
S	To plot default symbol at XY.
S#	To change default symbol to No # symbol and plot it at XY.
`TEXT	To plot $TEXT$ at XY. It must be the last operation in a GOC code. The TEXT following the ' command is interpreted and expanded by the DIGLIB software, if it includes any ^ operator or STRING names using the ~ descriptor.

## **BLOCKEND**

Syntax	BLOCKEND
	BLOCKEND terminates the local definition of the graphical operation code defined by the earlier BLOCK command.

## **CHARSIZE**

CHARSIZE redefines the default size of the characters in the plot. The character size has an initial default value, which may vary with the current font setting and the output device (the plot format).



"Examples of DATAPLOT Files and Outputs" on page 30

Syntax	CHARSIZE \$\$
	<pre>\$\$ must be an unsigned real.</pre>

## **CLIP**

CLIP turns clipping on or off. If it is OFF, it allows output outside the ordinary plot area defined by normalized coordinates zero and one.



"Example 2 – Draw Polygons and Symbols" on page 32 and "Example 9 - DIGLIB Fonts and Lines" on page 59

 Syntax
 CLIP CLP

 clp is a character string reading ON or OFF.

# **COLOR**

COLOR redefines the current color setting. There are 22 legal color codes (from 0 to 21).

.::: .:: "Example 7 – Color Codes" on page 46

Syntax	COLOR code
	code is an unsigned integer number (from 0 to 21) identifying the color, or a character string specifying the color.

#### **DIGLIB COLOR CODES**

Code	Color	Equivalent Character
0	Background	Invisible
1	Black	Normal
2	Red	Very_Thick
3	Green	Thin
4	Blue	Thick
5	Yellow	Very_Thin
6	Magenta	Dashed
7	Cyan	Dotted
8	Purple	
9	Gold	
10	Turquoise	
11	Pink	
12	Gray	
13	Orange-red	

Code	Color	Equivalent Character
14	Maroon	
15	Plum	
16	Sea green	
17	Olive	
18	Sienna	
19	Orange	
20	Coral	
21	UserDef	
22 and higher	Any digit larger tha 1 (black).	n 21 has the color effect as

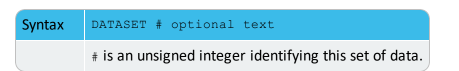
# DATAPOINT

DATAPOINT is not a DATASET command, but the basic DATAPLOT command (see synopsis below) performs an action at the current point determined by the specified X/Y- coordinates. A DATASET may contain various data points, in addition to one or more data <u>"BLOCK" on page 20</u>s. Such data points are separated and independent on each other.

Syntax	X Y GOC
	X and Y are unsigned real numbers identifying the X/Y-coordinates for the current data point. GOC stands for Graphical Operation Code (GOC) for the current point. Legal GOC characters.
	Example
	0.7 0.95 N'Example 6
	0.5 0.08 MNA'E^FS18^SQ(^SK^FS10A+5#8*C#^FS10 -!a^FS18)^FS11+B^DIa#b#\$

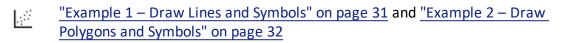
## DATASET

DATASET indicates the beginning of a number of consequent lines comprising a set of userdefined data. The DATASET lines are displayed on the terminal along with the <code>optional</code> <code>text</code> when using the <code>-1</code> option in the POST-processor as prompted for the DATASET number in the "APPEND\_EXPERIMENTAL\_DATA" on page 236 or "QUICK\_EXPERIMENTAL\_PLOT" on page 245 command.



# DRAWLINE

DRAWLINE draws a line starting at (x1,y1) to (xn,yn) through (x2,y2)  $\rightarrow$  (x(n-1), y(n-1)).



Syntax	DRAWLINE x1,y1 x2,y2 xn,yn
	${\bf x}$ and ${\bf y}$ may be reals or integers of any value. This is the same as connecting all points in a table:
	x1 y1 M x2 y2 D D xn yn D

# FONT

FONT redefines the default font setting in the POST-processor.

"Example 9 - DIGLIB Fonts and Lines" on page 59

Syntax	FONT #
	# is an unsigned integer.

#### **Font Numbers**

Number	Font
1	Bold
2	Normal
3	Italic
5	Bold and Italic
10	Monotype Normal

Number	Font
11	Monotype Italic
12	Monotype Bold
13	Monotype Bold and Italic
14	Narrow Normal
15	Narrow Italic

## **FUNCTION**

FUNCTION defines and plots a user-defined function.

"Example 4 – Draw Curves Defined by Functions" on page 36

Syntax	<pre>FUNCTION Y=f(X); start end number_of_XY_pairs; GOC;</pre>
	<pre>FUNCTION X=f(Y); start end number_of_XY_pairs; GOC;</pre>
	<pre>f(X) or f(Y) are legal mathematical functions of X or Y, understandable by a FORTRAN program. start and end are unsigned real numbers, and number_of_XY_pairs an unsigned integer. GOC is a legal graphical operation code as defined with <u>"BLOCK" on</u> page 20.</pre>

# **GLOBALSIZE**

GLOBALSIZE redefines the default global size of the plot. The global size has an initial default value.

Syntax	GLOBALSIZE \$\$
	\$\$ must be an unsigned real.

## **INCLUDE**

INCLUDE adds a file into the current input stream.

"Example 5 – Use Included Files for Predefined Symbols" on page 38.

#### Syntax INCLUDE filename

filename is a legal filename (with its correct path) for the operation system.

# LINETYPE

LINETYPE redefines the current linetype in the plot.



"Example 3 – Using Strings and Various Line Types" on page 34, "Example 8 -DIGLIB Symbols" on page 51 and "Example 9 - DIGLIB Fonts and Lines" on page 59

Syntax	LINETYPE #
	# must be an unsigned integer. Legal linetypes are (for both normal graphical outputs and for PostScript formats):
	<ul> <li>Solid (default)</li> </ul>
	Long dashed
	Short dashed
	• Dotted

#### **Line Types**

Number	Туре
1	Straight line
2	Dashed line
3	Dotted line
4	Dash-Dot line

# PAINT

PAINT paints the area enclosed by the current path in the current pattern. The current path starts at the last MOVETO given and includes all subsequent DRAWs. This command only works for the PostScript format (as graphical files or on printed hardcopy).

"PCFUNCTION" on the next page

Syntax	PAINT <code> <video> <mode></mode></video></code>
	<> denotes optional parameters. To set a new current pattern, supply any or all of the optional parameters.
	<ul> <li><code> is a single letter 0-9, A-Z or a-t (if <code>=t, also supply a space and a number in the range 0.00-1.00). The default is 0.</code></code></li> </ul>
	<ul> <li><video> is a string reading NORMAL (the default) or INVERSE.</video></li> </ul>
	<ul> <li><mode> is a string reading TRANSPARENT (the default) or OPAQUE.</mode></li> </ul>

## **PCFUNCTION**

PCFUNCTION appends a user-defined function to the current path. It is used with <u>"PAINT" on</u> the previous page.

Syntax	<pre>PCFUNCTION Y=f(X); start end number_of_XY_pairs; GOC;</pre>
	<pre>PCFUNCTION X=f(Y); start end number_of_XY_pairs; GOC;</pre>
	f(X) or $f(Y)$ are legal mathematical functions of X or Y, understandable by a FORTRAN program. start and end are unsigned real numbers, and number_of_XY_pairs an unsigned integer. GOC is a legal graphical operation code as defined with <u>"BLOCK" on</u> page 20.

# **STRING**

STRING defines a string containing a text and operation codes (e.g., to change the default font settings).

"Example 3 – Using Strings and Various Line Types" on page 34 and "Example 5 – Use Included Files for Predefined Symbols" on page 38

Syntax	STRING name text_with_each_character_in_ ^S#^G^F#^U#^D#^R#^L#^N
	name is a valid alphanumeric name (variable) to represent the text including all text formatting codes. It is highly recommended that all the characters in the name are in CAPITAL CASE (e.g. ACA2SO4); otherwise, the defined STRING can be incorrectly plotted.

#### Syntax STRING name text\_with\_each\_character\_in\_ ^S#^G^F#^U#^D#^R#^L#^N

text\_with\_each\_character\_in\_^S#^G^F#^U#^D#^R#^L#^N

means the text is coded with each of its characters (and/or numeric numbers) that are formatted with various operators ( $^S\#$ ,  $^G$ ,  $^F\#$ ,  $^U\#$ ,  $^D\#$ ,  $^R\#$ ,  $^L\#$  and/or  $^N$ ). #' are unsigned integers.  $^$  is the caret character and does not mean a control character. Between an # (in an operator) and a numeric number (as a part of the text), there must always be a comma sign (, ) or the number is not plotted in the text, because the # with the number is interpreted as another incorrect #. In some of the commands, #=0 resets the option to previous (or default) value.

#### STRING OPERATION CODES IN THE DIGLIB SOFTWARE

For the PostScript hardcopy fonts, these operation codes are not valid; see <u>"LaTeX</u> Formatting Codes" on page 14:

Operator	Operation
^S#	Set character size to size #
^G	Set font to Greek
^F#	Set font to font number #
^U#	Move up # units
^D#	Move down # units
^R#	Move right # units
^L#	Move left # units
^N	Do not move, remain at current

### **EXAMPLES**

STRING Alpha1 ^Ga^F0^D0^S8,1^S0^U0

#### is set to $\alpha_1$

STRING M23C6 M^D0^S8,23^S0^U0C^D0^S8,6^S0^U0

### is set to M<sub>23</sub>C<sub>6</sub>

STRING ACA2CO3 ^Ga^F0^D0^S8Ca^D0^S4,2^S0^U0^S8C0^D0^S4,3^S0^U0

is set to Ca<sub>2</sub>CO<sub>3</sub>

## **SYMBOLSIZE**

SYMBOLSIZE redefines the current symbol size setting. The symbol size has an initial default value.



<u>"Example 3 – Using Strings and Various Line Types" on page 34, "Example 7 – Color Codes" on page 46 and "Example 8 - DIGLIB Symbols" on page 51</u>

Syntax	SYMBOLSIZE \$\$
	<pre>\$\$ must be an unsigned real.</pre>

## TEXT

TEXT outputs, at the current position, the text following the keyword TEXT or the text in the string string name that is defined with <u>"STRING" on page 27</u>.

Syntax	TEXT text <b>Or</b> ~string_name
	It is possible to mix ^ operators and previously defined string names using the $\sim$ operator.

## **Examples of DATAPLOT Files and Outputs**

Once your Thermo-Calc software is installed you can open the Examples and PDF documentation (Manuals) from the main menu:

- To open the Manuals folder select Help → Manuals Folder. Note for Linux openSUSE users: This link is not functional. To access the folder, navigate to the default location based on your installation type
- To open the Examples based on whether you are in Console Mode or Graphical Mode, select File → Examples Files or Help → Examples Files.

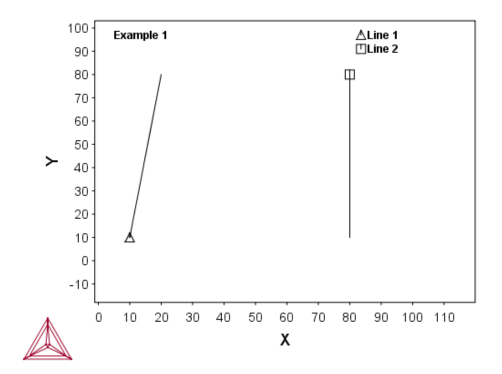
### In this section:

Example 1 – Draw Lines and Symbols	31
Example 2 – Draw Polygons and Symbols	32
Example 3 – Using Strings and Various Line Types	34
Example 4 – Draw Curves Defined by Functions	36
Example 5 – Use Included Files for Predefined Symbols	38
Example 6 – Triangular Diagrams for Ternary Systems	40
Example 7 – Color Codes	46
Example 8 - DIGLIB Symbols	51
Example 9 - DIGLIB Fonts and Lines	59

## **Example 1 – Draw Lines and Symbols**

The macro text below produces the example output. You can copy the text into your own EXP file, or use the example file DATAPLOT 1 - Draw Lines and Symbols.EXP available with your installation. To open the folder: From the Thermo-Calc menu, select Help → Manuals Folder.

### **Output Example**



### Macro Text

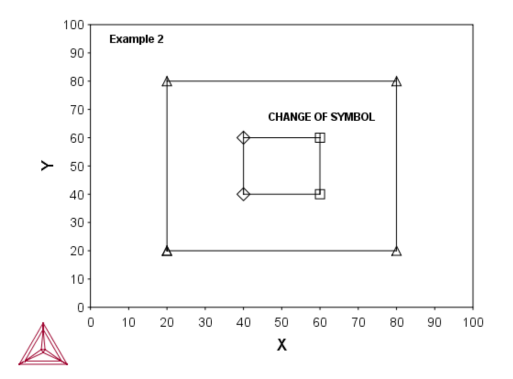
DATAPLOT Example 1 PROLOG 1 EXAMPLE 1 0<X<100, 0<Y<100 XSCALE 0.00000 100 XTYPE LINEAR YTYPE LINEAR XLENGTH 11.5000 YLENGTH 11.5000 TITLE EXAMPLE 1 XTEXT X YTEXT Y

```
DATASET 1 Two lines started with two symbols
ATTRIBUTE CENTER
0.05 0.95 N'Example 1
0.7 0.95 NS'Line 1
0.7 0.90 NS2'Line 2
10 10 S1
20 80 D
80 80 S2
80 10 D
50 60
```

## **Example 2 – Draw Polygons and Symbols**

The macro text below produces the example output. You can copy the text into your own EXP file, or use the example file DATAPLOT 2 - Draw Polygons and Symbols.EXP available with your installation. To open the folder: From the Thermo-Calc menu, select Help → Manuals Folder.

#### **Output Example**



#### Macro Text

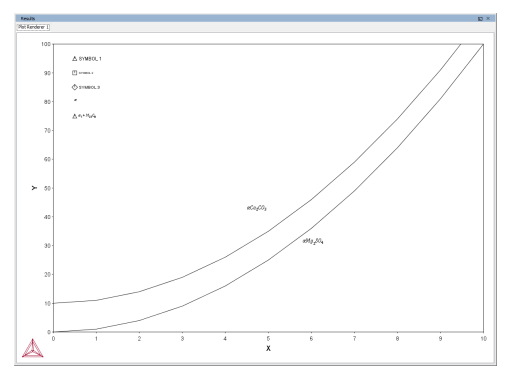
\$DATAPLOT Example 2 PROLOG 2 EXAMPLE 2 0<X<100, 0<Y<100 XSCALE 0.00000 100 YSCALE 0.00000 100 XTYPE LINEAR YTYPE LINEAR XLENGTH 11.5000 YLENGTH 11.5000 TITLE EXAMPLE 2 XTEXT X YTEXT Y DATASET 2 Two ploygons with three types of symbols ATTRIBUTE CENTER CLIP OFF 0.05 0.95 N'Example 2 1.1 0.95 NS1'SYMBOL 1 1.1 0.90 NS2'SYMBOL 2 1.1 0.85 NS3'SYMBOL 3 BLOCK X=C1; Y=C2; GOC=C3, DSWA 40 40 M 40 60 60 60 S2'CHANGE OF SYMBOL 60 40 40 40 SO BLOCKEND BLOCK X=C1\*100; Y=C2\*100; GOC=C3,DSWA 0.2 0.2 MS1 0.2 0.8 0.8 0.8 0.8 0.2 0.2 0.2 BLOCKEND \$Draw an additional polygon: DRAWLINE 20,10 30,15 40,30 50,80

DRAWLINE 50,80 60,60 90,30

## **Example 3 – Using Strings and Various Line Types**

The macro text below produces the example output. You can copy the text into your own EXP file, or use the example file DATAPLOT 3 - Using String and Line Types.EXP available with your installation. To open the folder: From the Thermo-Calc menu, select Help → Manuals Folder.

#### **Output Example**



#### Macro Text

\$DATAPLOT Example 3
PROLOG 3 EXAMPLE 3 0<X<10, 0<Y<100
XSCALE 0.00000 10
YSCALE 0.00000 100
XTYPE LINEAR
YTYPE LINEAR
XLENGTH 11.5000
YLENGTH 11.5000
TITLE EXAMPLE 3
XTEXT X</pre>

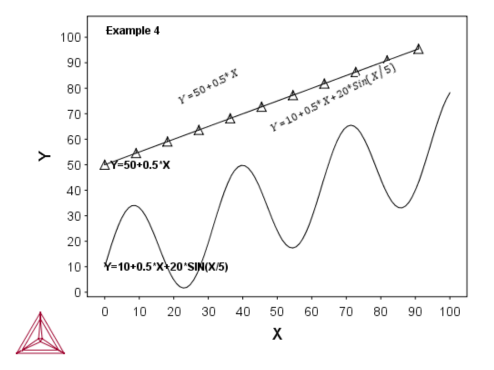
```
YTEXT Y
DATASET 3 Draw curves; plot formatted texts and symbols
$Define some strings:
STRING BCC \latex \alpha
STRING BCC1 \latex \alpha 1 + M 23C 8
STRING M23C6 M 23C 6
STRING ACA2CO3 \latex \alpha Ca_2CO_3
STRING AMG2SO4 \latex \alpha Mg_2SO_4
ATTRIBUTE CENTER
CLIP OFF
FONT 2
0.05 0.95 NS1'SYMBOL 1
CHARSIZE 0.2
0.05 0.90 NS2' SYMBOL 2
CHARSIZE 0.3
0.05 0.85 NS3'SYMBOL 3
0.05 0.80 N' ~BCC
0.05 0.75 NS1'~BCC1
SYMBOLSIZE 0.4
CHARSIZE 0.4
0.45 0.42 N'~ACA2CO3
0.58 0.30 N'~AMG2SO4
CLIP ON
LINETYPE 1
BLOCK X=C1; Y=C1*C1; GOC=C2, DWA
0 M
1
2
3
4
5
6
7
8
9
```

```
10
BLOCKEND
LINETYPE 2
BLOCK X=C1; Y=C1*C1+10; GOC=C2, DWA
0 M
1
2
3
4
5
6
7
8
9
10
BLOCKEND
```

## **Example 4 – Draw Curves Defined by Functions**

The macro text below produces the example output. You can copy the text into your own EXP file, or use the example file DATAPLOT 4 – Curves and Functions.EXP available with your installation. To open the folder: From the Thermo-Calc menu, select Help  $\rightarrow$  Manuals Folder.

### **Output Example**



#### Macro Text

\$DATAPLOT Example 4 PROLOG 4 EXAMPLE 4 0<X<100, 0<Y<100 XSCALE 0.00000 100 YSCALE 0.00000 100 XTYPE LINEAR YTYPE LINEAR XLENGTH 11.5000 YLENGTH 11.5000 TITLE EXAMPLE 4 XTEXT X YTEXT Y DATASET 4 Plot two functions as lines: ATTRIBUTE CENTER 0.05 0.95 N'Example 4 \$ Draw two lines defined by FUNCTIONS: FUNCTION Y=10+0.5\*X+20\*Sin(X/5); 0 100 100; DWA; FUNCTION Y=50+0.5\*X; 0 100 10; DS1WA; \$ Write functions beside the lines: \$ Note the real rotation angle (27 degree) can be seen \$ only on the PostScript hardcopy 0.25 0.68 N'^RO27Y=50+0.5\*X 0.25 0.45 N'^RO27Y=10+0.5\*X+20\*Sin(X/5)

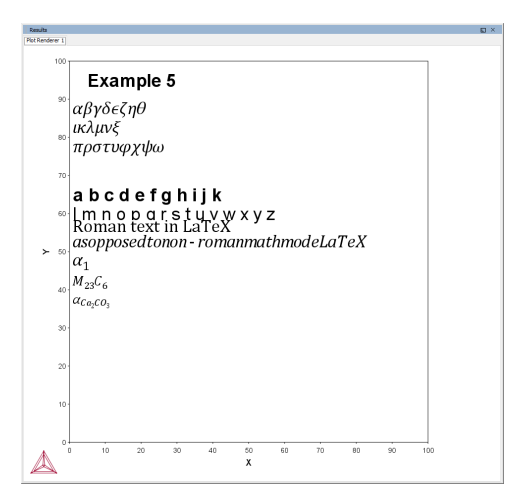
## **Example 5 – Use Included Files for Predefined Symbols**

The macro text below produces the example output. You can copy the text into your own EXP file, or use the example file DATAPLOT 5 – Predefined Symbols.EXP available with your installation. To open the folder: From the Thermo-Calc menu, select Help  $\rightarrow$  Manuals Folder.

### **Output Example**

8\$B

In the plot result, the hard-to-read LaTeX line is intended to highlight how to type readable text using LaTeX.



#### Macro Text

```
$DATAPLOT Example 5 Symbols
PROLOG 5 EXAMPLE 5 0<X<100, 0<Y<100
XSCALE 0.00000 100
YSCALE 0.00000 100
XTYPE LINEAR
YTYPE LINEAR
XLENGTH 11.5000
YLENGTH 11.5000
TITLE EXAMPLE 5
XTEXT X
YTEXT Y
DATASET 5 Write equations and strings with different properties
STRING TEST1A \latex \alpha \beta \gamma \delta \epsilon \zeta \eta \theta
STRING TEST1B \latex \iota \kappa \lambda \mu \nu \xi \omicron
```

```
STRING TESTIC \latex \pi \rho \sigma \tau \upsilon \phi \chi \psi \omega
STRING TEST2A a b c d e f g h i j k
STRING TEST2B l m n o p q r s t u v w x y z
STRING BCC1 \latex \alpha 1
STRING ACA2CO3 \latex \alpha {Ca 2CO 3}
ATTRIBUTE CENTER
CHARSIZE 1
0.05 0.95 N'Example 5
CLIP OFF
0.01 0.85 N'~TEST1A
0.01 0.80 N'~TEST1B
0.01 0.75 N'~TEST1C
FONT 1
0.01 0.65 N'~TEST2A
FONT 2
0.01 0.60 N'~TEST2B
0.01 0.55 N'\latex \textrm{Roman text in LaTeX}
0.01 0.50 N'\latex as opposed to non-roman math mode LaTeX
0.01 0.45 N'~BCC1
CHARSIZE 0.8
0.01 0.40 N'\latex M {23}C 6
0.01 0.35 N'~ACA2CO3
```

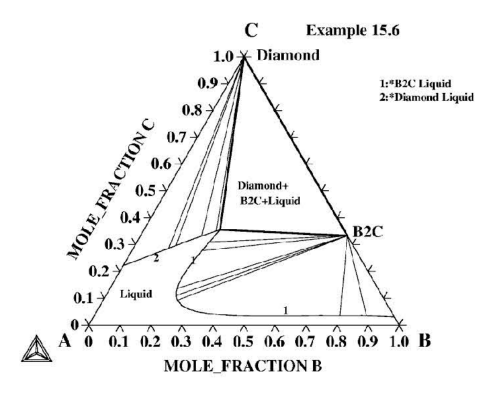
## **Example 6 – Triangular Diagrams for Ternary Systems**

The macro text below produces the example output. You can copy the text into your own EXP file, or use the example file DATAPLOT 6 – Ternary Plots.EXP available with your installation. To open the folder: From the Thermo-Calc menu, select Help  $\rightarrow$  Manuals Folder.

### **Output Example**



In the DATAPLOT language, this type of diagram is only available for Console Mode.



### Macro Text

\$DATAPLOT Example 6 - Ternary Phase Diagram

PROLOG 6 EXAMPLE 6 0<X<0.969224, 0<Y<1.00000

XSCALE 0.00000 0.969224

YSCALE 0.307492E-01 1.00000

**XTYPE LINEAR** 

YTYPE LINEAR

XLENGTH 11.5000

YLENGTH 11.5000

TITLE A-B-C at T=1000 K

**XTEXT MOLE\_FRACTION B** 

YTEXT MOLE\_FRACTION C

DIAGRAM\_TYPE TRIANGULAR YES YES

DATASET 6 Plot a ternary phase diagram

CLIP OFF

0.80 0.90 N'Example 6

0.86 0.31 N'B2C

0.52 0.97 N'Diamond

CHARSIZE 0.25

1.4E+01 1.10E+01 MVA'1:\*B2C Liquid

1.4E+01 1.05E+01 MVA'2:\*Diamond Liquid

0.10 0.10 N'Liquid

0.48 0.45 N'Diamond+

0.48 0.40 N' B2C+Liquid

5.80E-01 5.40E-02 MWA' 1

1.90E-01 2.40E-01 MWA' 1

0.65E-01 2.50E-01 MWA' 2

CHARSIZE 0.45

-0.10 -0.05 N'A

1.06 -0.05 N'B

0.48 1.05 N'C

CHARSIZE 0.35

\$\$ Calculated A-B-C Phase Equilibrium Data:

\$ PHASE REGION FOR:

\$F0 LIQUID

\$E DIAMOND\_A4

\$F0 B2C

\$ INVARIANT EQUILIBRIUM

COLOR 2

BLOCK X=C1; Y=C2; GOC=C3,WAD;

2.4555855989E-01 3.5568857193E-01 M

0.00000000E+00 9.9999523163E-01

2.4555855989E-01 3.5568857193E-01 M

6.66666666653E-01 3.3333334327E-01

0.00000000E+00 9.9999523163E-01 M

6.66666666653E-01 3.3333334327E-01

COLOR 1

BLOCKEND

\$ PHASE REGION FOR:

\$E LIQUID

\$F0 B2C

BLOCK X=C1; Y=C2; GOC=C3,WAD;

\$ PLOTTED COLUMNS ARE : X(LIQUID,B) and X(LIQUID,C)

2.2030337155E-01 1.2340000272E-01 M

2.2632879019E-01 1.1058768630E-01

2.3371633887E-01 9.9345825613E-02

2.4253317714E-01 8.9345827699E-02

2.6429468393E-01 7.2744041681E-02

2.8429466486E-01 6.2814079225E-02

2.9617273808E-01 5.8319382370E-02

3.2811737061E-01 4.9470417202E-02

3.6353862286E-01 4.3130427599E-02

3.9895987511E-01 3.8979098201E-02

4.5209178329E-01 3.5266116261E-02

5.2293431759E-01 3.3152002841E-02

6.1148744822E-01 3.3077053726E-02

6.4690870047E-01 3.3490389585E-02

6.8232995272E-01 3.4017231315E-02

- 7.3546189070E-01 3.4814555198E-02
- 7.5317251682E-01 3.5033416003E-02
- 8.0630439520E-01 3.5373892635E-02 8.5943627357E-01 3.4983776510E-02
- 9.1256815195E-01 3.3575300127E-02
- 9.6747112274E-01 3.0857827514E-02
- 9.6922445297E-01 3.0749246478E-02
- 2.2030337155E-01 1.2340000272E-01 M
- 2.1294665337E-01 1.5308913589E-01
- 2.1171525121E-01 1.8851040304E-01
- 2.1532440186E-01 2.2393165529E-01
- 2.2180187702E-01 2.5935292244E-01
- 2.2992117703E-01 2.9477417469E-01
- 2.3888295889E-01 3.3019542694E-01
- 2.4555855989E-01 3.5568857193E-01
- \$ PLOTTED COLUMNS ARE : X(B2C,B) and X(B2C,C)
- 6.66666666653E-01 3.3333334327E-01 M
- 6.66666666653E-01 3.3333334327E-01
- \$ TIELINES
- COLOR 3
- 6.66666666653E-01 3.3333334327E-01 M
- 8.7775242329E-01 3.4625384957E-02
- 6.66666666653E-01 3.3333334327E-01 M
- 7.9250496626E-01 3.5342670977E-02
- 6.66666666653E-01 3.3333334327E-01 M
- 2.4555824697E-01 3.5568737984E-01
- 6.66666666653E-01 3.3333334327E-01 M

2.3944084346E-01 9.2542596161E-02

6.66666666653E-01 3.3333334327E-01 M

2.3359020054E-01 3.0954307318E-01

6.66666666653E-01 3.3333334327E-01 M

2.2585247457E-01 2.7766343951E-01

6.66666666653E-01 3.3333334327E-01 M

2.1618695557E-01 1.3621240854E-01

2.2632879019E-01 1.1058768630E-01 M

6.66666666653E-01 3.3333334327E-01

COLOR 1

BLOCKEND

\$ PHASE REGION FOR:

\$F0 DIAMOND A4

\$E B2C

BLOCK X=C1; Y=C2; GOC=C3,WAD;

\$ PLOTTED COLUMNS ARE : X(DIAMOND\_A4,B) and X(DIAMOND\_A4,C)

0.00000000E+00 9.9999523163E-01 M

0.00000000E+00 9.9999976158E-01

BLOCKEND

\$ PHASE REGION FOR:

\$E LIQUID

\$F0 DIAMOND\_A4

BLOCK X=C1; Y=C2; GOC=C3,WAD;

\$ PLOTTED COLUMNS ARE : X(LIQUID,B) and X(LIQUID,C)

2.4555855989E-01 3.5568857193E-01 M

2.0263540745E-01 3.3483061194E-01

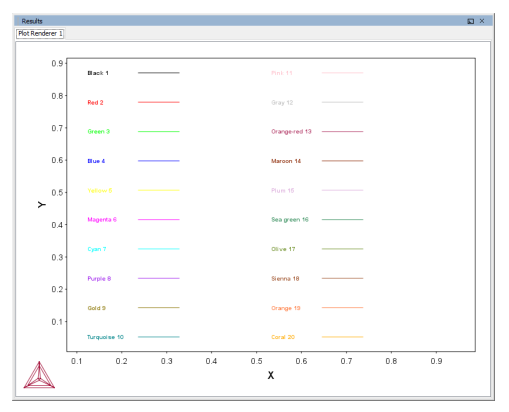
1.6243904829E-01 3.1375369430E-01

- 1.2243904918E-01 2.9153180122E-01
- 8.2439050078E-02 2.6854267716E-01
- 4.2439054698E-02 2.4548016489E-01
- 2.4390530307E-03 2.2313812375E-01
- 2.4999999937E-07 2.2181616724E-01
- \$ TIELINES
- COLOR 3
- 0.000000000E+00 9.9999511242E-01 M
- 2.3529186845E-01 3.5085919499E-01
- 0.00000000E+00 9.9999469519E-01 M
- 1.9826586545E-01 3.3260950446E-01
- 0.00000000E+00 9.9999409914E-01 M
- 1.3240024447E-01 2.9716023803E-01
- 0.00000000E+00 9.9999397993E-01 M
- 1.1439975351E-01 2.8695335984E-01
- BLOCKEND

## **Example 7 – Color Codes**

The macro text below produces the example output. You can copy the text into your own EXP file, or use the example file DATAPLOT 7 – Color Codes.EXP available with your installation. To open the folder: From the Thermo-Calc menu, select Help  $\rightarrow$  Manuals Folder.

### **Output Example**



#### Macro Text

\$DATAPLOT Example 7 - DIGLIB Colors PROLOG 1 DIGLIB Colors XSCALE 0.0 1.0 YSCALE 0.0 1.0 XTYPE LINEAR YTYPE LINEAR XLENGTH 15.5000 YLENGTH 11.5000 TITLE DIGLIB Colors XTEXT X YTEXT Y DATASET 1 DIGLIB Colors: \$ DIGLIB Colors: CHARSIZE 0.3 SYMBOLSIZE 0.5 COLOR 1

0.05 0.95 N' Black 1 BLOCK X=C1; Y=C2; GOC=C3,N 0.175 0.95 M 0.275 0.95 D BLOCKEND COLOR 2 0.05 0.85 N' Red 2 BLOCK X=C1; Y=C2; GOC=C3,N 0.175 0.85 M 0.275 0.85 D BLOCKEND COLOR 3 0.05 0.75 N' Green 3 BLOCK X=C1; Y=C2; GOC=C3,N 0.175 0.75 M 0.275 0.75 D BLOCKEND COLOR 4 0.05 0.65 N' Blue 4 BLOCK X=C1; Y=C2; GOC=C3,N 0.175 0.65 M 0.275 0.65 D BLOCKEND COLOR 5 0.05 0.55 N' Yellow 5 BLOCK X=C1; Y=C2; GOC=C3,N 0.175 0.55 M 0.275 0.55 D BLOCKEND COLOR 6 0.05 0.45 N' Magenta 6 BLOCK X=C1; Y=C2; GOC=C3,N 0.175 0.45 M 0.275 0.45 D BLOCKEND

```
COLOR 7
0.05 0.35 N' Cyan 7
BLOCK X=C1; Y=C2; GOC=C3,N
0.175 0.35 M
0.275 0.35 D
BLOCKEND
COLOR 8
0.05 0.25 N' Purple 8
BLOCK X=C1; Y=C2; GOC=C3,N
0.175 0.25 M
0.275 0.25 D
BLOCKEND
COLOR 9
0.05 0.15 N' Gold 9
BLOCK X=C1; Y=C2; GOC=C3,N
0.175 0.15 M
0.275 0.15 D
BLOCKEND
COLOR 10
0.05 0.05 N' Turquoise 10
BLOCK X=C1; Y=C2; GOC=C3,N
0.175 0.05 M
0.275 0.05 D
BLOCKEND
COLOR 11
0.5 0.95 N' Pink 11
BLOCK X=C1; Y=C2; GOC=C3,N
0.625 0.95 M
0.725 0.95 D
BLOCKEND
COLOR 12
0.5 0.85 N' Gray 12
BLOCK X=C1; Y=C2; GOC=C3,N
0.625 0.85 M
0.725 0.85 D
```

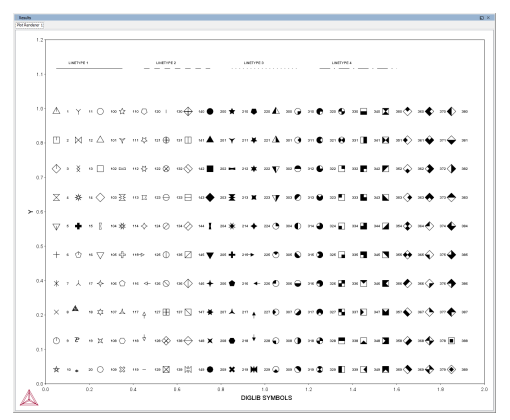
```
BLOCKEND
COLOR 13
0.5 0.75 N' Orange-red 13
BLOCK X=C1; Y=C2; GOC=C3,N
0.625 0.75 M
0.725 0.75 D
BLOCKEND
COLOR 14
0.5 0.65 N' Maroon 14
BLOCK X=C1; Y=C2; GOC=C3,N
0.625 0.65 M
0.725 0.65 D
BLOCKEND
COLOR 15
0.5 0.55 N' Plum 15
BLOCK X=C1; Y=C2; GOC=C3,N
0.625 0.55 M
0.725 0.55 D
BLOCKEND
COLOR 16
0.5 0.45 N' Sea green 16
BLOCK X=C1; Y=C2; GOC=C3,N
0.625 0.45 M
0.725 0.45 D
BLOCKEND
COLOR 17
0.5 0.35 N' Olive 17
BLOCK X=C1; Y=C2; GOC=C3,N
0.625 0.35 M
0.725 0.35 D
BLOCKEND
COLOR 18
0.5 0.25 N' Sienna 18
BLOCK X=C1; Y=C2; GOC=C3,N
0.625 0.25 M
```

0.725 0.25 D BLOCKEND COLOR 19 0.5 0.15 N' Orange 19 BLOCK X=C1; Y=C2; GOC=C3,N 0.625 0.15 M 0.725 0.15 D BLOCKEND COLOR 20 0.5 0.05 N' Coral 20 BLOCK X=C1; Y=C2; GOC=C3,N 0.625 0.05 M 0.725 0.05 D BLOCKEND

## **Example 8 - DIGLIB Symbols**

The macro text below produces the example output. You can copy the text into your own EXP file, or use the example file DATAPLOT 8 – DIGLIB Symbols.EXP available with your installation. To open the folder: From the Thermo-Calc menu, select Help  $\rightarrow$  Manuals Folder.

### **Output Example**



A default symbol is the current symbol in the run of the software (it is usually the No 1 symbol if the POST-processor is switched on for the first time). S (i.e., # is not specified) means that the current symbol is plotted. SO (i.e., #=0) means that no symbol is plotted.

### Macro Text

```
$DATAPLOT DIGLIB Symbols 2
PROLOG 1 DIGLIB Symbols 2 0<X<2.0, 0<Y<1.2
XSCALE 0.0 2.0
YSCALE 0.0 1.2
XTYPE LINEAR
YTYPE LINEAR
XLENGTH 15.5000
YLENGTH 11.5000
TITLE DIGLIB Symbols
XTEXT DIGLIB SYMBOLS
DATASET 1 Various DIGLIB Symbols:
$ DIGLIB Symbols:</pre>
```

```
CHARSIZE 0.3
SYMBOLSIZE 0.5
0.05 0.95 WS1' 1
0.05 0.85 WS2' 2
0.05 0.75 WS3' 3
0.05 0.65 WS4' 4
0.05 0.55 WS5' 5
0.05 0.45 WS6' 6
0.05 0.35 WS7' 7
0.05 0.25 WS8' 8
0.05 0.15 WS9' 9
0.05 0.05 WS10' 10
0.15 0.95 WS11' 11
0.15 0.85 WS12' 12
0.15 0.75 WS13' 13
0.15 0.65 WS14' 14
0.15 0.55 WS15' 15
0.15 0.45 WS16' 16
0.15 0.35 WS17' 17
0.15 0.25 WS18' 18
0.15 0.15 WS19' 19
0.15 0.05 WS20' 20
0.25 0.95 WS100' 100
0.25 0.85 WS101' 101
0.25 0.75 WS102' 102
0.25 0.65 WS103' 103
0.25 0.55 WS104' 104
0.25 0.45 WS105' 105
0.25 0.35 WS106' 106
0.25 0.25 WS107' 107
0.25 0.15 WS108' 108
0.25 0.05 WS109' 109
0.35 0.95 WS110' 110
0.35 0.85 WS111' 111
0.35 0.75 WS112' 112
```

0.35	0.65	WS113'	113
0.35	0.55	WS114'	114
0.35	0.45	WS115'	115
0.35	0.35	WS116'	116
0.35	0.25	WS117'	117
0.35	0.15	WS118'	118
0.35	0.05	WS119'	119
0.45	0.95	WS120'	120
0.45	0.85	WS121'	121
0.45	0.75	WS122'	122
0.45	0.65	WS123'	123
0.45	0.55	WS124'	124
0.45	0.45	WS125'	125
0.45	0.35	WS126'	126
0.45	0.25	WS127'	127
0.45	0.15	WS128'	128
0.45	0.05	WS129'	129
0.55	0.95	WS130'	130
0.55	0.85	WS131'	131
0.55	0.75	WS132'	132
0.55	0.65	WS133'	133
0.55	0.55	WS134'	134
0.55	0.45	WS135'	135
0.55	0.35	WS136'	136
0.55	0.25	WS137'	137
0.55	0.15	WS138'	138
0.55	0.05	WS139'	139
0.65	0.95	WS140'	140
0.65	0.85	WS141'	141
0.65	0.75	WS142'	142
0.65	0.65	WS143'	143
0.65	0.55	WS144'	144
0.65	0.45	WS145'	145
0.65	0.35	WS146'	146
0.65	0.25	WS147'	147

0.65	0.15	WS148'	148
0.65	0.05	WS149'	149
0.75	0.95	WS200'	200
0.75	0.85	WS201'	201
0.75	0.75	WS202'	202
0.75	0.65	WS203'	203
0.75	0.55	WS204'	204
0.75	0.45	WS205'	205
0.75	0.35	WS206'	206
0.75	0.25	WS207'	207
0.75	0.15	WS208'	208
0.75	0.05	WS209'	209
0.85	0.95	WS210'	210
0.85	0.85	WS211'	211
0.85	0.75	WS212'	212
0.85	0.65	WS213'	213
0.85	0.55	WS214'	214
0.85	0.45	WS215'	215
0.85	0.35	WS216'	216
0.85	0.25	WS217'	217
0.85	0.15	WS218'	218
0.85	0.05	WS219'	219
0.95	0.95	WS220'	220
0.95	0.85	WS221'	221
0.95	0.75	WS222'	222
0.95	0.65	WS223'	223
0.95	0.55	WS224'	224
0.95	0.45	WS225'	225
0.95	0.35	WS226'	226
0.95	0.25	WS227'	227
0.95	0.15	WS228'	228
0.95	0.05	WS229'	229
1.05	0.95	WS300'	300
1.05	0.85	WS301'	301
1.05	0.75	WS302'	302

1.05	0.65	WS303'	303
1.05	0.55	WS304'	304
1.05	0.45	WS305'	305
1.05	0.35	WS306'	306
1.05	0.25	WS307'	307
1.05	0.15	WS308'	308
1.05	0.05	WS309'	309
1.15	0.95	WS310'	310
1.15	0.85	WS311'	311
1.15	0.75	WS312'	312
1.15	0.65	WS313'	313
1.15	0.55	WS314'	314
1.15	0.45	WS315'	315
1.15	0.35	WS316'	316
1.15	0.25	WS317'	317
1.15	0.15	WS318'	318
1.15	0.05	WS319'	319
1.25	0.95	WS320'	320
1.25	0.85	WS321'	321
1.25	0.75	WS322'	322
1.25	0.65	WS323'	323
1.25	0.55	WS324'	324
1.25	0.45	WS325'	325
1.25	0.35	WS326'	326
1.25	0.25	WS327'	327
1.25	0.15	WS328'	328
1.25	0.05	WS329'	329
1.35	0.95	WS330'	330
1.35	0.85	WS331'	331
1.35	0.75	WS332'	332
1.35	0.65	WS333'	333
1.35	0.55	WS334'	334
1.35	0.45	WS335'	335
1.35	0.35	WS336'	336
1.35	0.25	WS337'	337

1.35	0.15	WS338'	338
1.35	0.05	WS339'	339
1.45	0.95	WS340'	340
1.45	0.85	WS341'	341
1.45	0.75	WS342'	342
1.45	0.65	WS343'	343
1.45	0.55	WS344'	344
1.45	0.45	WS345'	345
1.45	0.35	WS346'	346
1.45	0.25	WS347'	347
1.45	0.15	WS348'	348
1.45	0.05	WS349'	349
1.55	0.95	WS350'	350
1.55	0.85	WS351'	351
1.55	0.75	WS352'	352
1.55	0.65	WS353'	353
1.55	0.55	WS354'	354
1.55	0.45	WS355'	355
1.55	0.35	WS356'	356
1.55	0.25	WS357'	357
1.55	0.15	WS358'	358
1.55	0.05	WS359'	359
1.65	0.95	WS360'	360
1.65	0.85	WS361'	361
1.65	0.75	WS362'	362
1.65	0.65	WS363'	363
1.65	0.55	WS364'	364
1.65	0.45	WS365'	365
1.65	0.35	WS366'	366
1.65	0.25	WS367'	367
1.65	0.15	WS368'	368
1.65	0.05	WS369'	369
1.75	0.95	WS370'	370
1.75	0.85	WS371'	371
1.75	0.75	WS372'	372

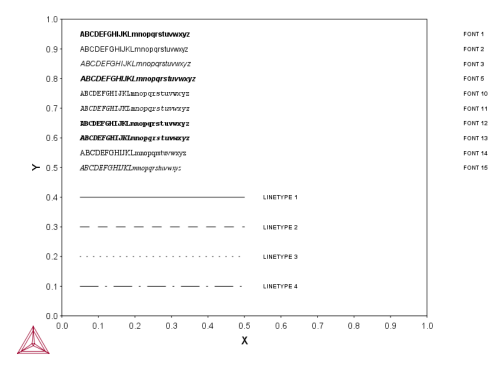
1.75 0.65 WS373' 373 1.75 0.55 WS374' 374 1.75 0.45 WS375' 375 1.75 0.35 WS376' 376 1.75 0.25 WS377' 377 1.75 0.15 WS378' 378 1.75 0.05 WS379' 379 1.85 0.95 WS380' 380 1.85 0.85 WS381' 381 1.85 0.75 WS382' 382 1.85 0.65 WS383' 383 1.85 0.55 WS384' 384 1.85 0.45 WS385' 385 1.85 0.35 WS386' 386 1.85 0.25 WS387' 387 1.85 0.15 WS388' 388 1.85 0.05 WS389' 389 \$ DIGLIB Line Types: LINETYPE 1 BLOCK X=C1; Y=C2; GOC=C3, DWA 0.05 1.10 M 0.35 1.10 BLOCKEND LINETYPE 2 BLOCK X=C1; Y=C2; GOC=C3,DWA 0.45 1.10 M 0.75 1.10 BLOCKEND LINETYPE 3 BLOCK X=C1; Y=C2; GOC=C3,DWA 0.85 1.10 M 1.15 1.10 BLOCKEND LINETYPE 4 BLOCK X=C1; Y=C2; GOC=C3, DWA

```
1.25 1.10 M
1.60 1.10
BLOCKEND
0.10 1.12 W' LINETYPE 1
0.50 1.12 W' LINETYPE 2
0.90 1.12 W' LINETYPE 3
1.30 1.12 W' LINETYPE 4
```

## **Example 9 - DIGLIB Fonts and Lines**

The macro text below produces the example output. You can copy the text into your own EXP file, or use the example file DATAPLOT 9 – DIGLIB Fonts.EXP available with your installation. To open the folder: From the Thermo-Calc menu, select Help  $\rightarrow$  Manuals Folder.

#### **Output Example**



#### Macro Text

\$DATAPLOT Example 9 - DIGLIB Fonts
PROLOG 1 DIGLIB Fonts & Line Types 0<X<1, 0<Y<1
XSCALE 0.0 1.0
YSCALE 0.0 1.0</pre>

XTYPE LINEAR YTYPE LINEAR XLENGTH 11.5000 YLENGTH 11.5000 TITLE DIGLIB Fonts XTEXT X YTEXT Y DATASET 1 Various DIGLIB Fonts and Line Types: \$ DIGLIB Fonts: CHARSIZE .35 FONT 1 0.05 0.95 N'ABCDEFGHIJKLmnopqrstuvwxyz FONT 2 0.05 0.90 N'ABCDEFGHIJKLmnopqrstuvwxyz FONT 3 0.05 0.85 N'ABCDEFGHIJKLmnopqrstuvwxyz FONT 5 0.05 0.80 N'ABCDEFGHIJKLmnopqrstuvwxyz FONT 10 0.05 0.75 N'ABCDEFGHIJKLmnopqrstuvwxyz FONT 11 0.05 0.70 N'ABCDEFGHIJKLmnopqrstuvwxyz FONT 12 0.05 0.65 N'ABCDEFGHIJKLmnopqrstuvwxyz FONT 13 0.05 0.60 N'ABCDEFGHIJKLmnopqrstuvwxyz FONT 14 0.05 0.55 N'ABCDEFGHIJKLmnopqrstuvwxyz FONT 15 0.05 0.50 N'ABCDEFGHIJKLmnopqrstuvwxyz FONT 1 CLIP OFF CHARSIZE .30 1.10 0.95 N'FONT 1 1.10 0.90 N'FONT 2

1.10 0.85 N'FONT 3 1.10 0.80 N'FONT 5 1.10 0.75 N'FONT 10 1.10 0.70 N'FONT 11 1.10 0.65 N'FONT 12 1.10 0.60 N'FONT 13 1.10 0.55 N'FONT 14 1.10 0.50 N'FONT 15 CLIP ON \$ DIGLIB Line Types: LINETYPE 1 BLOCK X=C1; Y=C2; GOC=C3,DWA 0.05 0.40 M 0.50 0.40 BLOCKEND \$DRAWLINE 0.05,0.40 0.5,0.40 LINETYPE 2 BLOCK X=C1; Y=C2; GOC=C3,DWA 0.05 0.30 M 0.50 0.30 BLOCKEND \$DRAWLINE 0.05,0.30 0.5,0.30 LINETYPE 3 BLOCK X=C1; Y=C2; GOC=C3,DWA 0.05 0.20 M 0.50 0.20 BLOCKEND \$DRAWLINE 0.05,0.20 0.5,0.20 LINETYPE 4 BLOCK X=C1; Y=C2; GOC=C3,DWA 0.05 0.10 M 0.50 0.10 BLOCKEND \$DRAWLINE 0.05,0.10 0.5,0.10 0.55 0.40 N' LINETYPE 1

0.55 0.30 N' LINETYPE 2 0.55 0.20 N' LINETYPE 3 0.55 0.10 N' LINETYPE 4

# **Database Manager User Guide**

Thermo-Calc Version 2018b





## Introduction to the Database Manager Guide

### In this section:

Thermodynamic and Kinetic/Mobility Databases	3
Initialization of the DATA Module	3
Initiation Parameter Examples	4

## **Thermodynamic and Kinetic/Mobility Databases**

This guide describes how to define and construct a thermodynamic or kinetic database. It includes details about the data structures and formats of the Thermo-Calc databases as well as the Diffusion Module (DICTRA) database-extensions.

The Thermo-Calc databases are created and maintained in the Thermo-Calc Database Format (TDB) which is an international standard for the CALPHAD-type thermodynamic calculations and kinetic simulations. This document gives a full description of the TDB format (with the Diffusion Module (DICTRA) database extension) and is a comprehensive guide about how to create and manage a Thermo-Calc (and Diffusion Module (DICTRA)) database.

Usually, a Thermo-Calc or Diffusion Module (DICTRA) database is constructed as a single text file with an extension of TDB, for example, PAQ2setup.TDB. For some databases (especially large databases used with older versions of Thermo-Calc or Diffusion Module (DICTRA)) a database may consist of one primary-definition file (\*\*\*setup.TDB) and several additional-definition TDB files (such as \*\*\*param.TDB, \*\*\*funct.TDB, \*\*\*refer.TDB, etc. These are built for a variety of defined parameters, functions, references and so forth, and are linked to the \*\*\*setup.TDB file through the correct use of the "TYPE\_DEFINITION " on page 21 and/or "FTP\_FILE" on page 24 commands inside the \*\*\*setup.TDB file.

With Thermo-Calc, Precipitation Module (TC-PRISMA) and Thermo-Calc Software Development Kits (SDKs), you can code everything into a single **\*\*\*setup.TDB** file for a database at any scale.

## **Initialization of the DATA Module**

The following explains how a database initiation file (or a database directory file) is constructed to work with the DATA module. When you start, DATA looks for this file that has information about the available predefined databases.

The *database initiation file* is called:

- Windows: TC\_INITD.TDB (or TC\_INITD)
- Linux/UNIX: initd.tdb

You can have your own file, but it is more efficient to define an environment variable with the default name as above, and then during installation, translate it to a common initiation file name. The common file is automatically copied by the Thermo-Calc installation script to the following directories:

- Windows: \DATA\ under the directory defined by the TCPATH parameter
- Linux/UNIX:/data/ under the directory defined by the TC\_DATA parameter.

The database manager can find the file on a local computer (independent installations) or a connected server (server installations).

When editing or modifying the definitions of initiation parameters (including short-names, paths and subdirectories, database definition file names, and instructive database descriptions) for the available databases in the database initiation file a specific format is used with the fields:

- The first field gives the abbreviated name for the database (to a maximum of five characters)
- The second field specifies a path and filename (maximum 78 characters) containing the database path definition and the actual database definition (*setup*) file name, where the extension is either TDB (which stands for original textual TDB file) or TDC (which is encrypted from the original TDB file). There are differences between platforms with respect to entering parameters as listed in the table.
- The third field (maximum 60 characters) details an optional full name (and version/subversion numbers) for the brief database description.

### **Initiation Parameter Examples**

8

Parameter	Windows	Linux/Unix
The setup file name must consist of a main part and an extension of a maximum of three characters:	For example, SSOL5SETUP.TDB	For example, ssol5setup.tdb
Database path definition	Use a back-slash $\$	Use a forward-slash /
All databases should be normally located in subdirectories that are under the directory defined by the:	TCPATH parameter or under its subdirectory \DATA\	TC_DATA parameter or under its subdirectory /data/

Each entry record (i.e. for a single database) must finish with an exclamation mark !. There should be no spaces or commas within the first and second fields since these characters are taken as field separators, while spaces or commas can be used in the third field. You can write each database entry on one or two lines (up to the!), it is recommended that the third field is added as follows:

- Windows: on the same line as the first and second fields.
- Linux/UNIX: on the following line.

The following are examples of an initiation file defining two databases named SSOL6 and TCFE8.

#### WINDOWS

```
SSOL6 TCPATH\DATA\SSOL6\SSOL6SETUP.TDC SGTE Solutions Database
version 6.0 !
TCFE8 TCPATH\DATA\TCFE8\TCFE8SETUP.TDC TCS Steels/Fe-Alloys
Database version 8.0 !
```

### LINUX/UNIX

ssol5 TC\_DATA/data/ssol5/ssol5setup.tdc
 SGTE Solutions Database version 5.1 !
Tcfe8 TC\_DATA/data/tcfe8/tcfe8setup.tdc
 TCS Steels/Fe-Alloys Database version 8.0 !

## The Database Definition File and Keywords

### In this section:

Database Definition File Syntax
ELEMENT
SPECIES
PHASE
Examples of Recommended Suffixes
GIBBS Phase-type Codes
Additional clarification
CONSTITUENT
ADD_CONSTITUENT
COMPOUND_PHASE
ALLOTROPIC_PHASE
TEMPERATURE_LIMITS
DEFINE_SYSTEM_DEFAULT
DEFAULT_COMMAND
DATABASE_INFORMATION
TYPE_DEFINITION
FTP_FILE
FUNCTION
PARAMETER
GIBBS Parameter Name
OPTIONS
TABLE
ASSESSED_SYSTEMS
REFERENCE_FILE

LIST_OF_REFERENCE	
ADD_REFERENCE	40
CASE and ENDCASE	
VERSION_DATA	

## **Database Definition File Syntax**

The database definition (\*.TDB) file (normally named \*\*\*setup.TDB) consists of a set of keyword codes each followed by one or several parameters (arguments).

- A complete keyword entry must end with an exclamation mark (!).
- A single keyword entry can be up to 2000 characters long. However, the maximum length of a line in an \*.TDB file is 78 characters--it may be necessary to continue the keyword parameter (arguments) on several lines. The ! must be at the end of the last line.
- It is recommended to always have at least one empty space at the beginning of each continuation line for the keyword parameters (arguments); otherwise, the DATA module can misunderstand the parameters (or arguments), or issue an error message.
- The keyword and its various parameters (arguments) are separated by a space or a comma.
- A dollar sign (\$) in the first position of the line indicates that the line is a comment line, which is ignored by the DATA module.

When the database is selected the DATA module only reads the database definition file (\*\*\*setup.TDB) once, from beginning to end. The DATA module checks continuously when reading the definition file. This implies that (nearly) everything must be declared or defined before it is used in any other way.

For example, if the GRAPHITE phase is to be included in the database definition, the element C (carbon) and the phase GRAPHITE must be defined before declaring that carbon dissolves in graphite. This definition order is necessary to build the internal data structure acceptable by the DATA module (during its consistency checking).

This topic gives a description of the available keywords and the appropriate arguments. A basic knowledge of the Gibbs Energy System (GIBBS) module is assumed.

This syntax is used:

KEYWORD [arg.1]\*# [arg.2]\*## {optional arg.3}!

The keywords are written in full length but can be abbreviated as long as the abbreviation is unique. A keyword may have syntax consisting of several arguments and optional arguments. The number, # or ##, in the notation, [...]\*# or [...]\*##, indicates an argument with a maximum length of # ASCII characters.

# Arguments within square brackets $[\ldots]$ must always be given, but are optional when enclosed in curly brackets $\{\ldots\}$ .

### **ELEMENT**

8\$B

ELEMENT [element name]\*2 [ref. state]\*24 [mass] [H298] [S298] !

The *element name* (maximum two characters) is the one found in the periodic chart but there are no naming restrictions. However, the GIBBS module only recognizes UPPER-case element names (if the Upper Case Mode is selected by the GIBBS command <u>"REINITIATE" on page 121</u>), which means that lower-case (if defined in a database) is automatically converted to UPPER-case by the DATA/GIBBS module.

The elements are automatically entered as species using the same names of the elements. If, for example, the species corresponding to FE needs to be named FE1, you can define the species as FE1, which results in an element named FE and a species named FE1. Vacancies (VA) and electrons (denoted either as /- in gaseous, liquid or solid phases, or ZE in an aqueous solution phase), need to be entered as special elements for correct handling by the DATA module.

The *reference (ref.) state* (maximum 24 characters) is the stable phase (at 298.15 K and 1 bar) that should contain this element and be used as the reference state for all element thermodynamic data. The *mass*, given in gram per mole, is used in various calculation programs and should always be given the correct value. *H298* and *S298* denote the enthalpy and entropy difference between 0 and 298.15 K for the element in SI units. If these are unknown, the values can be set to zero. All this information (reference state, H298 and S298) define the SER (Stable Element Reference state).

### **EXAMPLES**

ELEMENT /	/ –	ELECTRON_GAS	0.0	0.0	0.0	!		
ELEMENT \	VA	VACUUM	0.0	0.0	0.0	!		
ELEMENT 2	ΖE	UNIT_CHARGE	0.000	0000	0001 (	0.0	0.0	!
ELEMENT A	AL	FCC_A1	26.98	3154	4577.2	296	28.321	15 !
ELEMENT (	С	GRAPHITE	12.01	L1	1054.0		5.74	!
ELEMENT H	FΕ	BCC_A2	55.84	17	4489	2	7.28	!
ELEMENT (	0	1/2_MOLE_02(G)	15.99	994	4341	10	2.5158	3!
ELEMENT 1	ΤI	HCP_A3	47.88	3	4810	3	0.648	!
ELEMENT 2	ZR	HCP_A3	91.22	24	5566.2	7 3	9.181	!

ELEMENT ZY DUMMY 1 1 !

### **SPECIES**

SPECIES [species name]\*24 [stoichiometric formula] !

This keyword defines species in the data structure. Every *species name* (maximum 24 characters) must be unique. The species are built from the predefined set of elements in the stoichiometric formula. If an undefined element is referenced, DATA displays an error message and the data structure is probably damaged.

The species names do not have to be the same as the stoichiometry formula, although in general this is recommended. The elements are automatically entered as species using the same names of the elements.



You can define a species name as a mixture of UPPER-case and lower-case letters in a database, but the DATA module automatically converts all lower-case to UPPER-case because the GIBBS module only recognizes UPPER-case species names.

When naming the species in a database, use special characters (such as +, -, \_, / and .) in species names. Avoid using other special characters (such as ( and )).

The *stoichiometric formula* is written with a simplified chemical notation, in which the chemical elements should always be given in UPPER-case and in any preferred order, and the stoichiometric coefficients are written in either real numerical factors or integer digits.

It is important that the numerical factor of 1 is not left out. Subgroups are not allowed in a stoichiometry formula; however, while specifying the stoichiometry formula for a specific species in a database, you can specify it in a way that some elements (always together with the corresponding partial stoichiometric coefficients) are repeated (as in the examples below).

### **EXAMPLES**

8

SPECIES	AL203	AL203 !	
SPECIES	Silica	SI102 !	
SPECIES	NaSb_60H	NA1SB106H6	!
SPECIES	FE+2	FE/+2 !	
SPECIES	SB-3	SB/-3 !	

SPECIES	AlCl2/3	AL.33333CL.666667 !
SPECIES	AL1CL1H2O2	AL1CL1H2O2 !
SPECIES	AlCl3_3H2O	AL1CL3H6O3 !
SPECIES	AlO2H2Cl.H6O3	AL102H2CL1H603 !
SPECIES	AlCl2-OH.3H2O	AL1CL201H1H603 !
SPECIES	AlCl2OH.3Water	AL101H1CL2H6O3 !

### PHASE

```
PHASE [phase name]*24 [data-type code]*8 [numb. subl.] [sites in
subl. 1] [sites in subl. 2] etc... {auxiliary text string} !
```

This keyword defines a phase and its properties (except for what species are allowed to enter it and for its parameters).

The *phase name* (maximum 24 characters) must be unique; otherwise the DATA module sees it as an attempt to redefine a previously defined phase. This causes DATA to display an error message and ignore the rest of the line. A phase name can be suffixed by an underscore (\_and letters to identify the physical state(s) or structure type(s) of the phase.

### **Examples of Recommended Suffixes**

Suffix	Definition
ABC_S	The ABC phase in solid state.
ABC_S2	The ABC phase in solid state 2.
ABC_S3	The ABC phase in solid state 3.
ABC_LT	The ABC phase in solid state at low temperatures.
ABC_HT	The ABC phase in solid state at high temperatures.
ABC_L	The ABC phase in liquid state.
ABC_LIQ	The ABC phase in liquid state.
FCC_A1	The FCC phase in disordered structure type A1.
FCC_L12	The FCC phase in ordered structure type L12.

The *phase name* can also be attached with a colon sign (:) and a letter for a legal GIBBS phase-type code (e.g. IONIC\_LIQ:Y and GAS:G).

### **GIBBS** Phase-type Codes

Code	Definition
G	Bit set for a gaseous mixture phase.
A	Bit set for an aqueous solution phase.
Y	Bit set for an ionic liquid solution phase (specially treated by the Ionic Two-Sublattice Liquid Model).
L	Bit set for a liquid solution phase (but not A (aqueous) or Y (ionic liquid)).
I	Bit set for a phase with charged species (but not G (gaseous), A (aqueous) or Y (ionic liquid)).
F	Bit set for an ordered FCC or HCP solution phase using the <i>Four Substitutional-Sublattice Ordering Model</i> (additionally, such a phase can also have interstitial sublattices).
В	Bit set for an ordered BCC solution phase using the <i>Four Substitutional-Sublattice Ordering Model</i> (additionally, such a phase can also have interstitial sublattices).



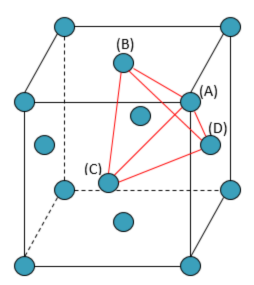
Other invalid characters (e.g. M or P) are eventually treated, together with the colon (:) as a part of a phase name.

A G phase (gaseous mixture) or an A phase (aqueous solution) is usually treated as a substitutional phase without sublattice, and an L phase (ordinary liquid solution) is normally (but not always) modelled as a substitutional phase without sublattice, too.

For ordered FCC or HCP phases, these four substitutional sublattices represent four corners of the regular tetrahedron on these lattices, all of which are the nearest neighbours, as shown.

### FCC UNIT CELL EXAMPLE

An FCC unit cell with the lattice positions indicated that correspond to the G (FCC,A:B:C:D) end member. All lattice positions are equivalent for a four substitutional-sublattice ordering model.



A *Normal 4-Sublattice Model* requires that all the G parameters for each of the endmembers with the same elements but distributed on different sites be given separately. However, as these corners are identical lattice points, the phase-type option F means that the G parameters need be given only once. The possible permutations are handled automatically.

### **Additional clarification**

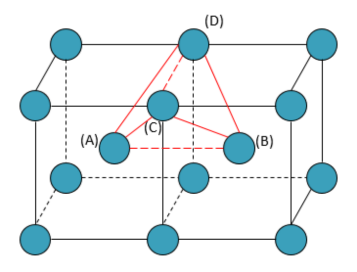
An A-B binary solution phase (with the element A located on one sublattice site and B on the other three sublattice sites) treated by the Normal 4-Sublattice Model has to have four G parameters for four end-members, i.e.

- G(phase,A:B:B:B)
- G(phase,B:A:B:B)
- G(phase,B:B:A:B, and
- G(phase,B:B:B:A)

This is because in the general case these G parameters can be different from each other. But for the FCC and HCP orderings, they are identical and thus all G parameters of such endmembers need to be given only once, and the possible permutations are then automatically handled by the GIBBS module. Also, only one of the identical permutations is listed; in this example, G (phase, A:B:B:B) where it is alphabetically the first in the list of permutations. This significantly simplifies the usage of this model (*Four Substitutional-Sublattice Ordering Model*) in multicomponent alloys. For ordered BCC phases, the phase-type option B means the same thing but it is more complicated since the 4-substitutional-sublattice ordering phase represents an irregular tetrahedron with two pairs of sites that are next nearest neighbours as shown:

### BCC UNIT CELL EXAMPLE

Two BCC unit cells with the lattice positions indicated that correspond to the G (BCC,A:B:C:D,0) end member. Lattice positions (A) and (B) are equivalent, as are lattice positions (C) and (B) for a four substitutional-sublattice ordering model.



For an end member described by the parameter G (phase, A:B:C:D) A and B are next nearest neighbours, as are C and D. And the nearest neighbours of A (or B) are C and D. Thus, for an A-B binary solution phase (with the element A located on two sublattice sites and B on two sublattice sites) treated by the Normal 4-Sublattice Model, the end-member described by the G (phase, A:A:B:B) term has four nearest neighbour bonds between A and B atoms, whereas the end-member described by the G (phase, A:B:A:B) term has two nearest neighbour bonds between A and B atoms and three next nearest neighbour bonds.

The first end-member (described by the G(phase, A:A:B:B) term) represents B2-ordering and the second (described by the G(phase, A:B:A:B) term) stands for B32-ordering. There are two permutations of the G(phase, A:A:B:B) term and four permutations of the G (phase, A:B:A:B) term, automatically conducted in the *Four Substitutional-Sublattice Ordering Model*. If you enter the unary, binary, ternary and quaternary parameters you are dealing with 1, 6, 21, 55 parameters for BCC:B, 1, 5, 15, 35 parameters for FCC:F and 1, 16, 81, 256 parameters for phases without F/B. An additional feature with the phase-type options F and B is that a composition set that represents the solution phase has a suffix (indicating what ordering the phase has) that is automatically added to its phase name in some listings of equilibrium calculations (when performing either single-point or stepping or mapping calculations, and when plotting the calculated property diagrams or phase diagrams).

Solution Phase	Suffix for Disordered Phase	Suffix for Ordered Phase
FCC PHASE	FCC A1	FCC_L12
		FCC_L10
	BCC_A2	BCC_B2
BCC PHASE		BCC_B32
DCC PRASE		BCC_D03
		BCC_L21
HCP PHASE		HCP_D019
NCF FRASE	HCP_A3	HCP_B19

Such suffix indications can be:

If you want to convert an existing database TDB-file to use the F/B feature, add the phase-type code to the corresponding phase name in the PHASE and CONSTITUENT commands in the TDB-file, then when running the LIST\_DATA command in the GES module. The created database file is in this less verbose format.

The *data-type code* consists of 1 to 8 characters where each character must stand for an action, which is to be coupled to this phase. The keyword <u>"TYPE\_DEFINITION " on page 21</u>, described below, must be used in the current database to specify what action should be taken by DATA for each character code.

The data entries [numb. subl.] [sites in subl. 1] [sites in subl. 2] etc., specify the total number of sublattices (always as an integer digit) and the sites (i.e. stoichiometric coefficients) of each of the sublattices (given in either integer digits or real numerical factors) for the phase.

Optionally, an *auxiliary text string* (maximum 78 characters) can be given after the last [sites in sublattice #] but before the exclamation mark !. This string displays in connection with the phase name in some listings within the DATA module.

### EXAMPLES

```
PHASE GAS:G % 1 1.0 !
PHASE LIQUID:L %ZCDQ 2 1.0 1.0
> Metallic liquid solution, modelled by CEF Model. !
PHASE IONIC-LIQ:Y %ZCDQ 2 1.0 1.0
> Ionic liquid solution, modelled by Ionic Two-Sublattice Model. !
PHASE SPINEL:I
                 %ZA 4 1 2 2 4
> Complex Spinel Solution, by CEF model with ionic constraints. !
PHASE M23C6 % 3 20.0 3.0 6.0 !
              %&A 2 1 1
PHASE FCC A1
> Disordered FCC phase; also as MX carbides/nitrides. !
PHASE FCC L10 %&AX 3 0.75 0.25 1
> Ordered FCC phase, modelled by 2-Sublattice Model for Ordering.
PHASE FCC L12:F % AX 5 0.25 0.25 0.25 1.0
> Ordered FCC phase, modelled by 4-Sublattice Model for Ordering.
PHASE AQUEOUS: A %HIJMR 1 1.0
> Aqueous Solution: using the Complete Revised HKF Model. !
```

### CONSTITUENT

8

CONSTITUENT [phase name]\*24 [constituent description]\*2000 !

This keyword (and the <u>"ADD\_CONSTITUENT" on the next page</u> keyword for large solution phase) defines the phase-constitution as a list of constituents (for a substitutional phase with no sublattice) or of constituent arrays (for a sublattice phase).

The *phase name* (maximum 24 characters) must be a predefined phase (i.e. already through the <u>"PHASE" on page 11</u> keyword).

It is important that if a phase bears a legal phase-type (among G, A, Y, L, I, F and B) in its phase definition (already by the PHASE keyword; such as GAS:G, SLAG:L, LIQUID:L, IONIC\_LIQ:Y, SPINEL:I, FCC\_L12:F, HCP\_D021:F, BCC\_B2:B, AQUEOUS:A), such a valid phase-type code must also always be attached to the phase name in the CONSTITUENT keyword (and the ADD\_CONSTITUENT keyword). Specifying the phase name in UPPER-case is recommended. You can define a phase name as a mixture of UPPER-case and lower-case letters in a database, but the DATA module automatically converts all lower-case to UPPER-case because the GIBBS module only recognizes UPPER-case phase names.

The *constituent description* (maximum 2000 characters) is a list of the species that enter a phase. The list starts with a colon (:), indicating the start of the sub-list of species for the first sublattice, and different sublattices are separated by colons. The complete sequence ends with a final colon.

Optionally, each sublattice may specify which species are considered to be major constituents. This is done by adding a percent sign (%) directly to the species name. The start values on the site fractions of the major constituents should sum to 0.99 on a specific sublattice. Thus, the minor constituents (i.e. those without a %) add up to 0.01. A maximum of 2000 characters can be coded in the constituent description, continuing in sequent lines. If the phase has a constituent description longer than 2000 characters, the rest can be coded in one or several ADD\_CONSTITUENT keywords.

#### **EXAMPLES**

```
CONSTITUENT BCC_A2 :FE

CONSTITUENT IONIC-LIQ:Y :FE+2 : SB-3: !

CONSTITUENT M23C6 :CR FE :FE CR W MO : C: !

CONSTITUENT AQUEOUS:A :H20% AG+1 AGF AGCL AGCL2-1 AGI3-2 AGS04-1

AGC2H4+1

AGN2H6+1 AGC2N2-1 AGC2H4NO2 AL+3 ALF3 ALO2-1, ... : !

CONSTITUENT SPINEL:I : AL+3% CR+3 FE+2% FE+3 MG+2% NI+2

: AL+3% CA+2 CR+3 FE+2 FE+3 MG+2% NI+2 VA

: FE+2 MG+2 VA%

: N-3 O-2% :!
```

## ADD\_CONSTITUENT

ADD CONSTITUENT [phase name]\*24 [constituent description]\*2000 !

This keyword adds more constituents to a phase that already has some constituents. Its syntax is the same as for the <u>"CONSTITUENT" on the previous page</u> keyword. This keyword can be used several times, if the phase is very large, e.g. a gaseous mixture or a complex aqueous solution. This is useful when there are so many constituents in a phase that the 2000 characters available for the constituent description list is not enough.



Constituents are not necessary on all sublattices. In the second example below, no addition is made to the first sublattice.

#### **EXAMPLES**

ADD\_CONSTITUENT GAS :S1 S2 S3 ... : ! ADD\_CONSTITUENT IM-PHASE : :CR:W ... : ! ADD\_CONSTITUENT AQUEOUS:A :CUCL+1 CUCL2 CUCL2-1 CUCL3-2 CUOH+1 CUO2H2 CUO3H3-1 CUO4H4-2 CU2OH+3 CU2O2H2+2 CU3O4H4+2 NIO2H2 NIO3H3-1 NIO4H4-2 NI2OH+3 NI4O4H4+4 ZNOH+1 ZNO2H2 ZNO3H3-1 ZNO4H4-2 ... : !

### **COMPOUND\_PHASE**

COMPOUND PHASE [phase name]\*24 [data-type code]\*8 [constituent] !

The keyword is a compact way to simultaneously define a species, a *compound phase* (maximum 24 characters) and its phase-constituent. It is useful for stoichiometric phases with constant compositions. The species name and stoichiometric formula must be identical, i.e. being the given *constituent*. The phase has this species as its only constituent. This keyword allows the database definition file for a large substance database to be more compact; it is a combination of the <u>"SPECIES" on page 10</u>, <u>"PHASE" on page 11</u> and <u>"CONSTITUENT" on page 16</u> keywords.

#### **EXAMPLES**

COMPOUND\_PHASE AL2O3 % AL2O3 ! COMPOUND\_PHASE MAGNETITE %MF FE3O4 ! COMPOUND\_PHASE QUARTZ % SIO2 !

### ALLOTROPIC\_PHASE

ALLOTROPIC\_PHASE [phase name]\*24 [data-type code]\*8 [constituent] !

This keyword does the same as the <u>"COMPOUND\_PHASE" above</u> keyword for entering an allotropic phase (maximum 24 characters), but does not enter the constituent as a species to the data structure. Use this if the species is already defined.

#### **EXAMPLES**

ALLOTROPIC\_PHASE BETHA-AL2O3 % AL2O3 ! ALLOTROPIC\_PHASE CRISTOBALITE % SIO2 ! ALLOTROPIC\_PHASE TRIDYMITE % SIO2 !

## **TEMPERATURE\_LIMITS**

TEMPERATURE LIMITS [lower limit] [upper limit] !

This keyword sets the default upper and lower temperature limits used by the GIBBS module for Gibbs energy parameters and functions. It can be used only once in one database definition file and all its sequential files.

### EXAMPLE

TEMPERATURE\_LIMITS 500.0 1800.0 !

## **DEFINE\_SYSTEM\_DEFAULT**

DEFINE\_SYSTEM\_DEFAULT [keyword] {G-ref. type index} !

This keyword sets the default value to ELEMENT or SPECIES in the DATA command <u>"DEFINE</u> <u>SYSTEM" on page 15</u> in the *Thermo-Calc Console Mode Command Reference*.

For a substance database, it can be appropriate to have ELEMENT as a default value whereas a large solution database can benefit from having SPECIES as a default value. A proper default value is useful for beginners. An advanced user is more likely to use the DATA commands DEFINE\_ELEMENT and DEFINE\_SPECIES to override the default value.

*{G-ref. type index}* is an integer indicating the reference state type for an element when entering and listing data in the GIBBS module. The following lists legal numbers and the corresponding meaning (the reference state type for an element):

Number	Definition
1	symbol: G
2	symbol: H298
3	symbol: H0

### EXAMPLE

DEFINE SYSTEM DEFAULT element 2 !

## DEFAULT\_COMMAND

DEFAULT\_COMMAND [secondary keyword and parameters] !

This keyword specifies commands to be executed by the DATA module at database initialization. The syntax of the available command is currently not the same as the user available DATA commands but the actions are similar. The available *secondary keyword and parameters* in syntax are:

DEFINE\_SYSTEM\_ELEMENT [element names] DEFINE\_SYSTEM\_SPECIES [species names] DEFINE\_SYSTEM\_CONSTITUENT [phase] [sublattice] [species] REJECT\_SYSTEM\_ELEMENT [element names] REJECT\_SYSTEM\_SPECIES [species names] REJECT\_SYSTEM\_CONSTITUENT [phase] [sublattice] [species] REJECT\_PHASE [phase names] RESTORE PHASE [phase names]

#### **EXAMPLES**

DEFAULT\_COMMAND DEFINE\_SYSTEM\_ELEMENT FE VA ! DEFAULT\_COMMAND REJECT\_SYSTEM\_CONSTITUENT LIQUID 2 C ! DEFAULT\_COMMAND REJECT\_PHASE LIQUID ! DEFAULT\_COMMAND RESTOR\_PHASE GAS !

### DATABASE\_INFORMATION

DATABASE INFORMATION [text]\*10000 !

This keyword defines a text for the detailed description of the current database. The text can be listed with the DATA command DATBASE\_INFORMATION. An apostrophe (') can be used in the text to indicate a new line; and two apostrophes, ('') can be used in the text to indicate a new line plus an empty line.



The continuous *text* length (each line with max 78 characters) is 10,000 characters.

#### **EXAMPLE**

```
DATABASE_INFORMATION This is the XXX-Alloy Solution Database '
in the A-B-C-D-..... System. '
Developed by TCS, released in May 2001. ''
... more ... !
```

## **TYPE\_DEFINITION**

```
TYPE_DEFINITION [data-type code]*1 [secondary keyword with
parameters] !
```

This keyword couples phases to an action performed by the DATA module when the DATA command <u>"GET\_DATA" on page 16</u> is executed.

### SECONDARY KEYWORDS

The secondary keywords and associated parameters in syntax are:

```
SEQ [filename]
RND# [filename]
GES [valid GIBBS command with parameters]
POLY3 [valid POLY command with parameters]
TDB [valid DATA command with parameters]
IF [conditional statement] THEN [keyword with parameters]
AFTER [valid GIBBS command with parameters]
```

The secondary keyword SEQ specifies a sequential file that stores parameters belonging to the phases using the associated data type code (which is defined by this keyword). A special case where the filename is given as an asterisk (\*) implies that the database definition file also acts as a sequential data storage file. This case makes it possible to have a single file for a small database, which is especially suited for personal databases.

The secondary keyword RND should be concatenated with a positive integer # to indicate the type of the random file. Currently, there are these types of random files:

- RNDO, the default, is used for complete Gibbs energy expressions (GO parameters), where the search field is the unabbreviated parameter name.
- RND1 is designated for functions, where the function name is used as the search field.
- RND2 is reserved for binary interaction parameters, where its search field is also the unabbreviated parameter name without any interaction order notation.



Ternary and higher order interaction parameters must be specified on a sequential file. Moreover, the internal structures of these random files are subject to changes with different versions of DATA, and with implementations of DATA on various computer systems. For more information, see the FORTRAN program TDBSORT, which is available from Thermo-Calc Software AB.

The secondary keywords GES, POLY3, or TDB specifies a modification of, or addition to, phases having the associated data type code, such as magnetic contribution, another excess model, or any other valid GIBBS/POLY/DATA command that applies to a certain phase. By implementing this as a call to the interactive GIBBS/POLY/DATA module, flexibility is achieved. If a new type of addition is implemented in a GIBBS/POLY/DATA module, it can be immediately used in the database definition file without reprogramming the DATA module.



In several examples below, the use of the at (@) sign indicates any phase to which the relevant type definition (e.g. A, B, 4, or E) applies.

The secondary keywords IF and THEN allow specification of a conditional statement structured with respect to the phase constitution that controls the execution of a following type-definition (keyword with parameters) string. See the last four examples.

The secondary keyword AFTER is similar to the GES keyword except the defined GIBBS command is executed after all parameters are entered. The reason for this is that the command has no effect unless there is a parameter. Following the AFTER keyword, a GIBBS command must be given and it is executed after entering the parameters of the phase.

The data-type code (always as one string) can be any normal or special character, e.g. 0, 5, A, F, M, &, &, and so forth, and is referred in the definition keywords <u>"PHASE" on page 11</u>, <u>"COMPOUND\_PHASE" on page 18</u> and <u>"ALLOTROPIC\_PHASE" on page 18</u> for various phases.

A phase can have several ADDITIONAL parts of different types (that are enforced by certain TYPE\_DEFINITIONS which call the GIBBS command AMEND\_PHASE\_DESCRIPTION for describing various contributions to Gibbs energy). However, the DATA/GIBBS module automatically deletes any early-defined ADDITIONAL part(s) of the same type.

The DATA module can selectively retrieve functions that are necessary for a defined system from a database that has functions stored in its setup file or SEQ sequential function file, while all other functions irrelevant for the defined system are ignored and are not saved in associated GIBBS and POLY workspaces.

### **EXAMPLES**

```
TYPE_DEF % SEQ TCPATH\DATA\[DATABASE]\PARAMETERS.TDB !

TYPE_DEF I SEQ TCPATH\DATA\[DATABASE]\INTERACTION-PARAMS.TDB !

TYPE_DEF G RND0 TCPATH\DATA\[DATABASE]\GZERO-PARAMS.TDB !

TYPE_DEF F RND1 TCPATH\DATA\[DATABASE]\FUNCTIONS.TDB !

TYPE_DEF & RND2 TCPATH\DATA\[DATABASE]\BINARY-INTERACTIONS.TDB !

TYPE_DEF A GES AM_PH_DES @ MAGNETIC -1 0.40 !

TYPE DEF B GES AM PH DES @ MAGNETIC -3 0.28 !
```

```
TYPE DEF 4 GES AM PH DES @ EXCESS MODEL REDLICH-KISTER KOHLER !
TYPE DEF 5 GES AM PH DES AQUEOUS EXCESS MODEL HKF !
TYPE DEF 6 GES AM PH DES AQUEOUS HKF ELECTROSTATIC !
TYPE DEF 7 GES AM PH DES AQUEOUS STATUS 02084000,,, !
TYPE DEF 8 GES AM PH DES AQUEOUS MAJOR CONST 1 H2O !
TYPE DEF E AFTER AM PH DES LIQUID EXCESS MIXED-EXCESS
                        A B LEGENDRE C A POLYNOM , , , !
TYPE DEF T AFTER AM PH DES LIQUID TERN-EXT TOOP-KOHLER B A C ,,,,
TYPE DEF Q AFTER AM PH DES LIQUID TERN-EXT KOHLER FE CR NI !
TYPE DEF C IF (PD AND PT AND SN) THEN TDB RESTORE PHASE BCT A5 !
TYPE DEF D IF (PD AND (PT OR SN)) THEN TDB REJECT PHASE BCC A2 !
TYPE DEF E IF ((NB OR TI OR V) AND (C OR N)) THEN
          GES AM PH DES @ COMP SET ,, CR NB TI V: C N: !
TYPE DEF F IF (ALO3/2 OR CRO3/2 OR FEO OR MNO OR SIO2) THEN
         GES AM PH DES LIQUID COMP SET ,, ALN%, ALO3/2%, CRO3/2%,
            FEO%, FEO3/2, MNO3/2%, MNS%, SIO2%, TIO2% : !
TYPE DEF R GES AM PH DES FE LIQUID FRACTION LIMITS Fe 0 0.6
        Ag 0 0.01 Al 0 0.05 Ca 0 0.05 Co 0 0.01 Cr 0 0.01
        Cu 0 0.02 Mg 0 0.05 Mn 0 0.05 Mo 0 0.05 Nb 0 0.05
        Ni 0 0.05 Pb 0 0.05 Si 0 0.10 Sn 0 0.02 Ti 0 0.05
        U 0 0.01 V 0 0.02 W 0 0.02 Zr 0 0.03
        B 0 0.01 C 0 0.01 H 0 0.01 N 0 0.01 O 0 0.01
        P 0 0.01 S 0 0.01 !
```

#### **ORDERED PHASE RESTORES DISORDER CONTRIBUTION**

When an ordered phase with disordered contribution is restored, the disordered contribution is automatically restored as well. You can also restore the disordered phase *without* restoring the ordered phase.

However the DIS\_PART TYPE\_DEFINITION must always be set on the ordered phase. It is important that the disordered phase is declared *before* the ordered phase. Otherwise it causes an error when DIS\_PART TYPE\_DEFINITION is executed and then either the ordered or disordered phase is rejected.

#### EXAMPLE

8

```
TYPE_DEFINITION & GES A_P_D FCC_A1 MAGNETIC -3.0 2.80000E-01 !
PHASE FCC_A1 %& 2 1 1 !
CONSTITUENT FCC_A1 :CR,NI% : C%,VA : !
$ THIS PHASE HAS A DISORDERED CONTRIBUTION FROM FCC_A1
TYPE_DEFINITION ' GES AMEND_PHASE_DESCRIPTION FCC_L12 DIS_PART
FCC_A1,,,!
TYPE_DEFINITION ( GES A_P_D FCC_L12 MAGNETIC -3.0 2.80000E-01
!
PHASE FCC_L12 %'( 3 .75 .25 1 !
CONSTITUENT FCC L12 :CR,NI% : CR,NI% : C,VA% : !
```

For example a GES command such as the following only affects phases defined above or at the phase with this TYPE\_DEFINITION in the TDB file:

TYPE DEFINITION ( GES A P D FCC L12 DIS PART FCC A1,,, !

For example a TDB command such as the following only affects phases defined *below* the phase with this TYPE\_DEFINITION in the TDB file:

TYPE\_DEFINITION ( TDB RESTORE\_PHASE FCC A1 !

### **FTP\_FILE**

FTP FILE [filename] !

FTP\_FILE is a special function random file and the function names correspond to the record numbers where these record names and the functions are stored. The FTP\_FILE decreases search time for the associated database in the DATA module. The file is used for large substance databases along with a SEQ sequential or RND0 random file for storage of G0 parameters referring the functions named FxxxxT that are stored in the FTP file. The integer number xxxx is a search code used by DATA when such files are processed.



No modification of this file type is allowed.

### EXAMPLE

```
FTP FILE TCPATH\DATA\[DATABASE]\FTP-FILE.DATA !
```

## **FUNCTION**

8

GIBBS can use predefined functions in the expression (TP-Function) of a Gibbs energy parameter or in other functions. This is often used when several parameters (or functions) have a common sub-expression, such as for metastable modifications of elements. This keyword can appear in both files for database definition and sequential storage, but not in FTP files. A valid *function name* can have up to 8 characters.

A function always starts with a lowest temperature limit of its applicability, followed by one or more (up to 10) expressions (TP-Functions) that are coded as mathematical relations of constants, functions of stable variables (T and P) and other entered functions (normally with a # suffix, e.g. +3\*GHSERAL#).

### "ENTER\_PARAMETER" on page 137 in the Thermo-Calc Console Mode Command Reference

The expression is a FORTRAN-like expression and operators +, -, \*, = and \*\* can be used (\*\* only with integer powers). Unary-functions LN or LOG (both for natural logarithm) and EXP (for exponential) can also be used. Each expression (TP-Function) should end with a semicolon (;), and be followed by its upper applicable temperature limit and a continuation indicator (Y to continue with the next expression or N to end the function's expression). If there is no continuation after a specific expression (TP-Function), the reference index can optionally be given after the N indicator.

A complete/valid function entry can be written in several continuation lines if the function's expression (TP-Function) is too long or if there is more than one applicable expression (TP-Function), as the maximum length of each line is 78 characters.

It is recommended to always have at least one empty space at the beginning of each continuation line. Otherwise, the DATA module may misunderstand the expression or issue some error messages when reading the function entry. Avoid entering functions like the one below. FUNCTION GHSERXY 298.15
-1000+1058\*T-38.9\*T\*LOG(T)+GFUNXY#; 6000 N !

Such a function is read by the DATA module as

1000+1058\*T-38.9\*T\*LOG(T)+GFUNXY#

rather than

-1000+1058\*T-38.9\*T\*LOG(T)+GFUNXY#

This is because the DATA module concatenates all lines and removes extra spaces before trying to enter the function in the GIBBS workspace. Thus, the – sign is taken as a delimiter between 298.15 and 1000, and the function incorrectly becomes:

```
FUNCTION GHSERXY 298.15 1000+1058*T-38.9*T*LOG(T)+GFUNXY#; 6000 N !
```

Avoid this mistake by giving at least one empty space as the first character of a new line, such as

FUNCTION GHSERXY 298.15
-1000+1058\*T-38.9\*T\*LOG(T)+GFUNXY#; 6000 N !

which is read correctly as

FUNCTION GHSERXY 298.15 -1000+1058\*T-38.9\*T\*LOG(T)+GFUNXY#; 6000 N !

The lowest-temperature limit (in Kelvin) for the applicability of the (first) TP-Function in a function is normally set by default as 298.15 K, in most cases. However, you can set another limit when it is applicable (according to experimental data and assessments).

An upper-temperature limit (in Kelvin; followed by a Y or N sign) for the applicability of each TP-Function in a function must be given after the semicolon (;) immediately following the specific TP-Function. The highest-temperature limit (in Kelvin) for the applicability of the current function is always followed by the N sign. If a negative number is given as the lowest-temperature limit, it assumes there are breakpoints in pressure for this function. In these cases, it is interpreted as the lowest-pressure limit (in Pascal), and the other limits in the current function is also taken as pressure limit values (in Pascal).

The temperature/pressure limits for the functions are checked during calculations. An indicator is set if the actual temperature/pressure condition is below the lowest temperature/pressure limit or above the highest temperature/pressure limit. In these cases, an extrapolation is done using the TP-Function valid in the nearest temperature/pressure range.

The optional reference index {*Ref. Index*} is an integer number indicating where to find the particular function in a special reference file. The references are listed when doing the GET\_ DATA command in the DATA module. They can also be listed in the GIBBS module with the command LIST\_DATA with the option R.

For accounting for the reference indices, also see the keyword <u>"REFERENCE</u> FILE" on page 38.

The reference index field can also be an abbreviation (such as REF: 250,  $REF_002$ , or REF-SGTE) which denotes the original reference. In this case, the reference cannot be obtained when issuing the DATA command GET\_DATA or the GIBBS command LIST\_DATA (with the option R).

However, the references directly coded in the database definition file (\*\*\*setup.TDB) starting with a letter can be shown when issuing the DATA command GET\_DATA or the GIBBS command LIST\_DATA (with the option N or R). Normally, such references must be located after the LIST\_OF\_REFERENCE keyword. It is recommended to use reference code names such as REF001, REF018, etc. The reference list, which is generated by the GIBBS command <u>"LIST\_DATA" on page 115</u> <file> with the N or R option, is also possible to be directly read by the DATA module.

The DATA module can selectively retrieve functions which are necessary for a defined system from a database that has functions stored in its setup file or SEQ sequential function file, while all other functions irrelevant for the defined system are ignored and are not saved in associated GIBBS and POLY workspaces. Previously, this can only be done for large databases that have functions stored in RND1 random or FTP function files.

### **EXAMPLES**

FUNCTION GFREE 298.15 1000+GFUNXY#; 6000 N !
FUNCTION GFUNXY 298.15 -1000+200\*T+30\*T\*LOG(T); 6000 N 505 !
FUNCTION G0\_CAO 298.15 -663538.11+352.67749\*T-57.7533\*T\*LN(T)
+5.3895E-03\*T\*\*2-8.879385E-07\*T\*\*3+575530\*T\*\*(-1);
1400.00 Y -625196.99+78.896993\*T-20.40145\*T\*LN(T)
-1.112923E-02\*T\*\*2+5.1896733E-07\*T\*\*3-6917350\*T\*\*(-1);
2900.00 Y -499226.55-490.37695\*T+51.95912\*T\*LN(T)
-2.961051E-02\*T\*\*2+1.4033905E-06\*T\*\*3-48114685\*T\*\*(-1);
3172.00 Y -587711.89+375.04117-62.76\*T\*LN(T);
6000.00 N REF020 !

### PARAMETER

PARAMETER [GIBBS parameter name] [lowest temp. limit]

This keyword can appear in both files for database definition and sequential storage, but not in FTP files. After the keyword, a valid *GIBBS parameter name* should be given.

It is used to define standard Gibbs energies (i.e. the G parameters for Gibbs energy of formations) of all valid end-members of various stoichiometric and solution phases, and excess Gibbs energies (i.e. the L parameters for Gibbs energy of interactions) of all binary, ternary, quaternary or higher-order interactions in various solution phases; both standard Gibbs energies and excess energies can also have parameters for contributions from PT-depended volume variations (i.e. the VO, VA, VB, VC and VK parameters for molar volume, thermal expansivity, bulk modulus, isothermal compressibility and high-pressure fitting parameter), magnetic ordering (i.e. the TC and BM parameters for Curie temperature and Bohr magneton number) and hypothetical electrostatic interactions (i.e. BM parameter for Born functions  $\omega$ Pr,Tr of aqueous solute species).

The general form of a parameter is:

```
<identifier>(<phase>, <constituent array>; <digit>) <xxx>
<expression> <yyy> <keyword Y or N> <zzz> !
```

### **GENERAL FORM**

Name	Description
identifier	The parameter type.
phase	The phase name (maximum 24 characters).
constituent array	The specific constituent array in the phase.
digit	The degree of composition-dependent interaction contribution (an integer number from 0 to 9), that is only for excess energy (L), Curie temperature (TC) and Bohr magneton number (BMAGN), as well as for volume-related parameters (V0 or VA or VB or VC or VK); if it is valued as zero, or if it is for the standard Gibbs energy (G) for which the degree is always zero, it can be omitted.

Name	Description
expression	The mathematical relation to describe the parameter.
xxx and YYY	The low and high temperature limits respectively for the applicable temperature range of the parameter expression
keyword Y or N	The indicator on if there is continuation for the parameter expression or not
ZZZ	The reference index/number for the assessment of this parameter;
!	The exclamation mark is used to indicate that the current parameter definition is ended

### **GIBBS Parameter Name**

The GIBBS parameter name has a general form of:

```
<identifier>(<phase>,<constituent array>;<digit>)
```

### NAME EXAMPLES

Parameter	Definition
G(GAS,C102)	The Gibbs energy of formation of a CO2 molecule in gas.
G(FCC, FE:VA)	The Gibbs energy of formation of fcc Fe with interstitials.
L(LIQ,Fe,Cr;0)	The regular solution parameter for Fe and Cr in liquid.
L(LIQ,Fe,Cr;1)	The sub-regular solution parameter.
TC(BCC,Fe:Va)	The Curie temperature of bcc Fe.
BMAGN(BCC, Fe:Va)	The Bohr magneton number parameter of bcc Fe.

The GIBBS parameter name consists of several parts. The first is a *type-identifier*. The following type-identifiers are legal:

Type- Identifier	Definition
G	Standard energy parameter (Gibbs energy of formation)

Type- Identifier	Definition
L	Excess energy parameter (Gibbs energy of interaction)
тс	Curie temperature for magnetic ordering
BMAGN or BM	Bohr magneton number for magnetic ordering (or Born function $\omega$ Pr,Tr for aqueous solute species).
V0	Molar volume at 298.15 K and 1 bar (a numeric value only)
VA	$\int\limits_{1}^{T}lpha(T)dT$ Integrated thermal expansivity $^{298.15}$
VB	Bulk modulus at 1 bar
VC	Isothermal compressibility
VK	High-pressure fitting parameter

You can also use G for interaction parameters; and on output list (performed by the GIBBS command LIST\_PARAMETER or LIST\_PHASE\_DATA) the type-identifier L is always used for interaction parameters. Note that the type-identifier BM is also used for Born functions  $\omega$ Pr,Tr of aqueous solute species.

The identifier must be followed by an opening parenthesis, a phase name, a comma and a constituent array. Optionally, the constituent array can be followed by a semicolon and a digit. The parameter name is terminated by a closing parenthesis.

It is important that if a phase bears a legal phase-type (among G, A, Y, L, I, F and B) in its phase definition (already by the PHASE keyword; such as GAS:G, LIQUID:L, SLAG:L, IONIC\_LIQ:Y, SPINEL:I, FCC\_L12:F, HCP\_D021:F, BCC\_
B2:B, AQUEOUS:A), such a valid phase-type code should not be attached to the phase name in the PARAMETER keyword.

8

Specifying the phase name in UPPER-case is recommended. You can define a phase name as a mixture of UPPER-case and lower-case letters in a database, but the DATA module automatically converts all lower-case to UPPER-case because the GIBBS module only recognizes UPPER-case phase names.

The *constituent array* consists of a list of constituent names. Interaction parameters have two or more constituents from the same sublattice separated by a comma. If the phase has sublattices, at least one constituent in each sublattice must be specified. The constituents in different sublattices must be given in sublattice order and are separated by a colon.

After the component array, a sub-index digit can be specified after a semicolon. This digit must be in the range 0 to 9. The interpretation of the sub-index depends on the excess energy model used for the phase. If no semicolon and digit are given, the sub-index value is assumed to be as zero.

The excess energy parameters, e.g. the regular/subregular (binary) parameter or ternary parameters, are multiplied with two or more fractions of the constituents from the same sublattice of the solution phase. These additional constituents must be given as interacting constituents.

Be careful about the sign of odd terms, for example, L(BCC, B, A:VA; 1) is treated as L(BCC, A, B:VA; 1), i.e. it is always put into alphabetical order.



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Solution phases with sublattices may have interacting constituents in each sublattice.

You can use an asterisk (\*) to denote that the excess interaction parameter is independent of the constituents of a specific sublattice. For example,  $L(FCC_L12, AL, NI:*)$  means that the interaction parameter is for the binary interaction between constituents AL and NI on the first sublattice in the FCC\_L12 solution phase, while it is independent of all constituents on the second sublattice. A interaction parameter in the list of constituents is always added to the Gibbs energy and the asterisk (\*) is calculated with the term of  $[1-\sum y(specified constituents)]$ , which implies that in an A-B binary system the following three L parameters are identical (but in higher-order systems, they are different):

- L(phase, A, B) is multiplied with X(A) \*X(B)
- L(phase, A, \*) is multiplied with X(A) \* (1-X(A))
- L(phase, B, \*) is multiplied with X(B) \* (1-X(B))

A parameter always starts with a lowest temperature limit of its applicability, followed by one or more (up to 10) expressions (TP-Functions) coded as mathematical relations of constants, functions of stable variables (T and P) and entered functions (normally with a # suffix, e.g. +3\*GSERAL#).

The expression is a FORTRAN-like expression and operators +, -, \*, = and \*\* can be used (\*\* only with integer powers). Unary-functions LN or LOG (both for natural logarithm) and EXP (for exponential) can also be used. Each expression (TP-Function) should ends with a semicolon (;) and be followed by its upper applicable temperature limit and a continuation indicator (Y to continue with the next expression, or N to end the parameter's expression). If there is no continuation after a specific expression (TP-Function), the reference index can be optionally given after the N indicator.

A complete/valid parameter entry can be written in several continuation lines if the parameter's expression (TP-Function) is too long or if there is more than one applicable expression (TP-Function), as the maximum length of each line is 78 characters.

It is recommended to always have at least one empty space at the beginning of each continuation line. Avoid entering parameters such as:

```
PARAMETR G(LIQUID,A,B) 298.15
-2000+4568*T+2*GFUNAB#; 6000 N !
```

**B** 

Such a parameter is read by the DATA module as 2000+4568\*T+2\*GFUNAB#, rather than as – 2000 +4568\*T+2\*GFUNAB#.

Avoid this mistake by giving at least one empty space as the first character of a new line, such as

```
PARAMETR G(LIQUID,A,B) 298.15
-2000+4568*T+2*GFUNAB#; 6000 N !
```

The lowest-temperature limit (in Kelvin) for the applicability of the (first) TP-Function in a parameter is normally set by default as 298.15 K, in most cases; however, you can set another limit when it is applicable (according to experimental data and assessments). An upper-temperature limit (in Kelvin; followed by a Y or N sign) for the applicability of each TP-Function in a parameter must be given after the semicolon (;) immediately following the specific TP-Function; and the highest-temperature limit (in Kelvin) for the applicability of the current parameter is always followed by the N sign. If a negative number is given as the lowest-temperature limit, it is assumed that there are breakpoints in pressure for this parameter. In such cases, it is interpreted as the lowest-pressure limit (in Pascal), and the other limits in the current parameter are also taken as pressure limit values (in Pascal).

The temperature/pressure limits for the parameters are checked during calculations. An indicator is set if the actual temperature/pressure condition is below the lowest temperature/pressure limit or above the highest temperature/pressure limit. In these cases, an extrapolation is done using the TP-Function valid in the nearest temperature/pressure range.

The optional reference index {*Ref. Index*} is an integer number indicating where to find the particular parameter in a special reference file. The references are listed when doing the GET\_DATA command in the DATA module. These can also be listed in the GIBBS module with the command LIST\_DATA and the option R or N.



For accounting the reference indices, also see the keyword <u>"REFERENCE\_FILE"</u> on page 38.

The reference index field can also be an abbreviation (such as REF:250,  $REF_002$ , or REF-SGTE) denoting the original reference. In this case, the reference cannot be obtained when issuing the DATA command GET\_DATA or the GIBBS command LIST\_DATA (with the option R or N).

However, the references directly coded in the database definition file (\*\*\*setup.TDB) that starts with a letter can be shown when issuing the DATA command GET\_DATA or the GIBBS command LIST\_DATA (with the option R or N). Normally, such references must be located after the LIST\_OF\_REFERENCE keyword. It is recommended to use reference code names such as REF001, REF018, etc. The reference list, which is generated by the GIBBS command LIST\_DATA <file> with the N option, is thus also possible to be directly read by the DATA module.

#### **EXAMPLES**

```
PARAMETER G(BCC, FE:VA) 298.15 1000+200*T+...; 6000 N 91DIN !
PARAMETER TC (BCC, FE:VA)
                           298.15 +1043; 6000 N 91DIN !
PARAMETER BMAGN(BCC, FE:VA) 298.15 +2.22; 6000 N 91DIN !
PARAMETER G(SIGMA, FE:CR:CR;0) 298.15 1000+200*T+...; 6000 N 101 !
PARAMETER G(LIQUID, AL; 0) 298.15 +11005.553-11.840873*T
        +7.9401E-20*T**7+GHSERAL#;
933.60 Y +10481.974-11.252014*T+1.234264E+28*T**(-9)+GHSERAL#;
   2900.00 N REF:283 !
PARAMETER G(BCC A2, PB:C) 298.15 UN ASS#; 300 N REF:0 !
PARAMETER G(BCC A2,NI:C;0) 298.15 +GHSERNI#+3*GHSERCC#
                                    +400000-100*T; 6000 N REF071 !
PARAMETER G(BCC A2, MN:VA) 298.15 +GMNBCC#; 6000 N REF285 !
PARAMETER BM(AQUEOUS, OH-1) 298.15 +Z0002PW0#; 1600 N 155 !
PARAMETER L(BCC,FE,CO:VA;0) 298.15 1000+200*T+...; 6000 N !
PARAMETER L(BCC,FE,CO:VA;1) 298.15 1000+200*T+...; 6000 N !
PARAMETER L(BCC, FE, CO:VA; 2) 298.15 1000+200*T+...; 6000 N !
```

PARAM TC(BCC\_A2,CO,MO:VA;O) 298.15 -3700; 6000 N R454 !
PARAM TC(BCC\_A2,CO,MO:VA;1) 298.15 +2300; 6000 N R454 !
PARAM BMAGN(BCC\_A2,CO,MO:VA;O) 298.15 -3.445; 6000 N R454 !
PARAM VO(BCC\_A2,CR,FE:VA;O) 298.15 +2ERO#; 6000 N REFO6V !
PARAM VO(BCC\_A2,CR,FE:VA;1) 298.15 -1.10524097E-7; 6000 N REFO6V !
PARAM VO(BCC\_A2,CR,FE:VA;2) 298.15 +1.40024130E-7; 6000 N REFO6V !
PARAM VA(BCC\_A2,CR,FE:VA;0) 298.15 -6.49444634E-6\*DELTAT#; 6000 N
REFO6V !
PARAM VA(BCC\_A2,CR,FE:VA;1) 298.15 +2.91269321E-5\*DELTAT#; 6000 N
REFO6V !

### **OPTIONS**

OPTIONS /[alloy name]([composition limitations for all alloying elements]) !

This keyword defines an alloy in a database. An alloy has a name, a major component and a number of alloying elements. The purpose for defining an alloy is to be able to tell you about applicable composition limits of the current database in applications to that particular type of alloy. It is possible to have several alloys in the same database. The alloys are given after the keyword in the database.

The alloy name must be proceeded by a slash (/) and terminated by the opening parenthesis, with no spaces are allowed in between.

- The alloy name is maximum 8 characters.
- After the parenthesis, follows the major element and a parenthesis with its minimum mass and minimum mole percent given inside.
- Then, the alloying element names, each with its maximum mass and mole percent are given within parenthesis.
- There must a space between definitions for each alloying element.
- The alloy definition is terminated by a closing parenthesis, and the whole OPTIONS keyword by the exclamation mark (!).

#### **EXAMPLE**

OPTIONS /SSteel(Fe(60,60) CR(30,30) NI(15,15) SI(1,1) N(.1,1)) !

## TABLE

8

It is recommended to always have at least one empty space at the beginning of each continuation line. Otherwise, the DATA module may issue some error messages when reading the table entry.

```
TABLE [name]*8 [start temp] [end temp] [delta temp] [table values]
!
```

This keyword can appear in both files for database definition and sequential storage, but not in FTP files. It makes a table of Gibbs energy as a function of temperature where the values are given from the start temperature to the end temperature, at a step of the delta temperature.

### EXAMPLE

```
TABLE DEMTAB 1000.0 1500.0 100.0 -2912.9008 -2834.2416 -2755.5824
-2677.7600 -2600.7744 -2524.2072 !
```

## ASSESSED\_SYSTEMS

ASSESSED\_SYSTEM [descriptions on special treatments for specific assessed systems]\*8000 !

This keyword can be included in the database definition file (the \*\*\*setup.TDB file). A maximum of 8000 characters after the keyword (to describe some special options when the DATA, GIBBS and POLY models deal with the existing systems with assessed data) is allowed until the exclamation mark (!).

Several ASSESSED\_SYSTEMS keywords can be used in the same TDB file in order to have more lists of assessed systems. There is no limit to the number of the entries in the same TDB file.

The assessed systems in the database, and the special treatment options, are typed after the keyword. The elements (always in UPPER CASE) in each assessed system must be in alphabetical order and be separated by a hyphen, such as C-FE for the Fe-C binary system, and C-CR-FE for the Fe-Cr-C ternary system. A space must be between each assessed system. Information on assessed binary, ternary or higher-order systems may also be given in this way.



A ternary system like C-CR-FE does not imply that the binary C-CR, C-FE and CR-FE are assessed. There is no way to indicate partially assessed systems.

- There is a field to give some descriptive information for a specific system, with various options about how to:
- Reject or restore phase(s) from the current database in the DATA module;
- Set major constituent(s) in the first composition set and to set a second composition set for a specific phase available in the current database in the GIBBS module; and
- Calculate this specific system in the POLY module.
- This is the facility used by the BIN (binary phase diagrams) and TERN (ternary phase diagrams) modules in the Thermo-Calc software/database package.

The descriptive information must immediately follow the specific system name, and must be enclosed within parenthesis (and), and the left parenthesis must follow directly after the system, such as :

```
AL-NI(TDB +L12 ;G5 C-S:L12/NI:AL:VA ;P3 STP:.8/1200/1
STP:.2/600/1)
```

The syntax TDB means that the commands to the DATA module proceed, and +L12 in the example means that the phase called L12 should be restored (it has been rejected by default).

The directive ; G5 means that the following are commands to the GIBBS module.  $c_s$ : means creating a second composition set, after the colon follows the phase name and after the slash the major constituents.

After the directive ; P3 follows commands to the POLY module. STP: means setting a start point with the value of the X-axis first (composition for the second element in a binary system), the slash separates the Y-axis value (temperature), and possibly one or more directions (-1, 1, -2 or 2).

A summary of the allowed syntax is:

TDB accepts

- +phase and -phase for restore/reject.
- ; G5 accepts
- MAJ:phase/constituent-array for major constituents of the first composition set.
- C S:phase/constituent-array for a second composition set.
- ; P3 accepts
- TMM:lt/ht for the low-/high-temperature limits (lt and ht; for instance TMM:500/4000) suitable for calculating phase diagrams and property diagrams of a binary system.
- \* for a default start point which is set as:

- for a binary system: at the composition  $x(2^{nd} \text{ element}) = .1234$ , temperature T=1100 K and with the default directions; or
- for a ternary system: at the compositions  $x (2^{nd} element) = .1234$  and  $x (3^{rd} element) = .1234$  and with the default directions.
- STP:x/t/d1/d2/d3 for a specific start point in a binary system which is set as at the composition X (2<sup>nd</sup> element) =x and temperature T=t (in K), and with the directions d1, d2 and/or d3.
- STP:x1/x2/d1/d2/d3 for a specific start point in a ternary system which is set as at the compositions X(2<sup>nd</sup> element)=x1 and X(3<sup>rd</sup> element)=x2 and with the directions d1, d2 and/or d3.
- The direction(s) can be defined as -1, 1, -2 or 2. If no direction is specified, all default directions are used (meaning no ADD command is enforced in the POLY module).
- If only one start point is specified, the direction(s) may be omitted; if more than one start points are specified, at least one direction for each start point must be given for all start points.

Each entry for a specific binary or ternary sub-system can be written in one or several lines (each line with 78 characters).

### **OTHER EXAMPLES**

```
ASSESSED_SYSTEMS

AL-NI(TDB +L12 +BCC_B2 ;G5 C_S:L12/NI:AL:VA

;P3 STP:.8/1000/1 STP:.45/700/1 STP:.7/700/1)

AL-PB(TDB -HCP -BCC

;G5 MAJ:LIQ/AL MAJ:FCC/AL:VA C-S:LIQ/PB C-S:FCC/PB:VA ;P3

*)

CR-FE(;G5 C-S:BCC/CR:VA ;P3 STP:.6/1200/1/-2/2)

AG-CU(;G5 MAJ:FCC/AG:VA C_S:FCC/CU:VA ;P3 STP:.3/1000)

C-NB(;P3 STP:.9/1100/1)

C-SI(;P3 *)

CO-CR(;G5 MAJ:FCC/CO:VA C_S:FCC/CR:VA ;P3 STP:.1/1100)

CR-FE(TDB -HCP ;G5 C_S:BCC/CR:VA ;P3 STP:.6/1200/1/-2/2)

CR-NI(;P3 *)

CR-W(;G5 MAJ:BCC/W:VA C_S:BCC/CR:VA

;P3 TMM:500/4000 STP:.3/700/1 STP:.3/1800/2 )
```

```
CU-FE(TDB -HCP ;G5 MAJ:LIQ/CU MAJ:FCC/FE:VA C_S:FCC/CU:VA
    ;P3 STP:.9/1400)
FE-N(TDB +FE4N ;P3 *)
FE-O(TDB -LIQUID +IONIC ;G5 C_S:ION_LIQ/FE+2:O-2 MAJ:ION_
LIQ/FE+2:VA
    ;P3 STP:.2/2000/1 )
FE-S(TDB -LIQUID +IONIC ;G5 C_S:ION_LIQ/FE+2:S MAJ:ION_LIQ/FE+2:S-
2 ;P3 *)
AL-MG-SI(;P3 *)
C-CR-FE ;G5 MAJ:BCC/FE:VA C_S:BCC/CR:VA ;P3 *) !
```



The semicolon (;) is a part of the ;G5 and ;P3 directives. A long descriptive information can be written in more than one line, such as for the AL-NI, AL-PB, CU-FE and FE-O systems shown above.

The directive ; P3  $\star$  is needed if the default start point should be used. If there is no P3 directive, the BIN or TERN module generates some 20 different start points in order to cover all possible compositions and temperatures (for a binary system) or all possible compositions (for a ternary system under any specific temperature).

### **REFERENCE\_FILE**

REFERENCE FILE [file name] !

This keyword takes a reference file name as its argument. This reference file (that must be edited in a restrictive way and be saved as a blocked file with a fixed line-length for each line in the file, as described below) contains a complete list of the references for the various parameters (and sometimes functions) in the database. The file must have a fixed record structure: each reference entry with one or several records, and each record with exactly 78 characters written in one single blocked line; and no empty space is allowed at the beginning of all lines in the file. If there are more than one record entered for a reference entry, all the continuation lines must start with an ampersand ( $\alpha$ ). The line number of the first record for a specific reference entry is then accounted as the unique integer for that specific reference, which is referred when a parameter or function calls this integer as the optional *{Ref. Index}*.



Also see keyword <u>"PARAMETER" on page 27</u> or <u>"FUNCTION" on page 25</u> about specifying a reference index.

#### EXAMPLE

/-1<G> T.C.R.A.S. Class 1

```
AG1.64TE1
               THERMODATA 01/93
&28/01/93
&SILVER 1.64-TELLURIDE. Solid Standard State.
AG1
                HULTGREN SELECTED VAL. SGTE **
&AT.WEIGHT 107.870, STANDARD STATE: CODATA KEY VALUE.MPT=1234.93K.
&--U.D. 30/10/85.
AG1<G>
               T.C.R.A.S Class: 1
AG1/+1<G> T.C.R.A.S Class: 1
AG1BR1
                N.P.L.
                                              SGTE **
&Tfusion uncertain and heat vaporization estimated.
                THERMODATA 01/93
AG1BR1<G>
&28/01/93
&Gaseous Standard State.
AG1BR103
                BARIN & KNACKE.SUPPL.REF:62,* SGTE **
&AGO3BR SILVER OXYTRIBROMIDE
```

In the example, the unique integers of related references (for assessed elements, species, phases, interactions, etc.) are:

1/-1<G> 2AG1.64TE1 5 AG1 8AG1<G> 9AG1/+1<G> 10AG1BR1 12AG1BR1<G> 15AG1BR103

### LIST\_OF\_REFERENCE

```
LIST_OF_REFERENCE
NUMBER SOURCE
[REFxxx] '[Detailed reference]'
......!
```

This keyword starts a reference list that is directly coded in the database definition file (\*\*\*setup.TDB). Its argument begins on the following line, and normally has an explanation line (NUMBER SOURCE) that is followed by various reference codes. Each reference code

may occupy one or more lines (each line with maximum 78 characters), but must have a reference code name (that starts with a letter) and the detailed reference information (that is written within two single-quotation marks, ''). It is recommended to use reference code names such as REF001, REF018, etc. A maximum of 400,000 characters after the keyword is allowed until the exclamation mark (!).

Such a reference list can be shown when issuing the DATA command <u>"GET\_DATA" on</u> page 16 or the GIBBS command <u>"LIST\_DATA" on page 115</u> (with the R option). The reference list, which is generated by the GIBBS command LIST\_DATA <file> with the N option, has this structure, and is thus possible to be directly read by the DATA module.

#### **EXAMPLE**

```
LIST_OF_REFERENCES

NUMBER SOURCE

REF283 'Alan Dinsdale, SGTE Data for Pure Elements,

Calphad Vol 15(1991) p 317-425,

also in NPL Report DMA(A)195 Rev. August 1990'

REF224 'P-Y Chevalier, Thermochimica Acta, 130 (1988) p 33-41; AG-

SI'

!
```

### ADD\_REFERENCE

```
ADD_REFERENCE
[REFxxx] '[Detailed reference]'
......!
```

Use this keyword to have an unlimited number of references. The use of this is the same as that of <u>"LIST\_OF\_REFERENCE" on the previous page</u>, and is used after it. Several ADD\_ REFERENCES sections can be used in a single database if there are too many references.

It is recommended to always start with a so-called *empty reference* (such as the DUMPO reference in the following example) as the first entry in each of the ADD\_REFERENCE sections to make sure that all references are listed appropriately as retrieving data by issuing the "GET\_DATA" on page 16 command.

#### EXAMPLE

```
ADD_REFERENCE
DUMP0 'Empty reference 0'
```

```
REF4 'J-O Andersson, B. Sundman, CALPHAD Vol 11, (1987), p 83-
92
          TRITA 0270 (1986); CR-FE'
   REF5 'J-O Andersson, Met. Trans A, Vol 19A, (1988) p 627-636
          TRITA 0207 (1986); C-CR-FE'
   REFS 'Pingfang Shi (2006), TCS PTERN Public Ternary Alloys
Database,
          v1.2; Modified L0(BCC, Fe, C) and L0(BCC, Cr, C) parameters
          at high temperatures.'
   . . . . . . .
!
ADD REFERENCE
   DUMP1 'Empty reference 1'
   REF275 'A. Fernandez Guillermet, Z. Metallkde. Vol 79(1988)
p.524-536,
          TRITA-MAC 362 (1988); C-CO-NI AND C-CO-FE-NI'
   REF393
          'K. Frisk, Metall. Trans. Vol 21A (1990) p 2477-2488,
          TRITA 0409 (1989); CR-FE-N'
   REF1096 'P. Gustafson, Metall. Trans. 19A(1988) p 2547-2554,
          TRITA-MAC 348, (1987); C-CR-FE-W'
   . . . . . . .
!
```

### **CASE and ENDCASE**

```
CASE [ELEMENT/SPECIE/PHASE] !
IF (boolean algebra on element, species or phase names) THEN
[GIBBS/POLY/DATA command] !
ENDCASE !
```

This keyword takes as its argument, a definition on which type of the following Boolean algebra operates. A simple Boolean algebra using AND and OR with a maximum of four levels of parentheses works. The CASE construction must end with the ENDCASE keyword. This makes it possible to have additional GIBBS or POLY commands executed depending on the user selection of elements, species or phases. The DATA commands that can be given as secondary keyword to "DEFAULT COMMAND " on page 19 can also be executed.

#### EXAMPLE 1

```
CASE ELEMENT !
IF((CR OR TI OR V) AND N)
THEN GES AM_PH_DES @ C_S ,, CR MO TI V:C N: !
ENDCASE !
```

#### EXAMPLE 2

```
CASE ELEMENT !
IF(O) THEN TDB DEFINE_SYSTEM_ELEMENT /- !
ENDCASE !
```

#### EXAMPLE 3

```
CASE ELEMENT !
IF(AL AND FE)
THEN TDB DEF_SYS_ELEMENT VA !
ENDCASE !
```

### **VERSION\_DATA**

```
VERSION DATE [string] *78 !
```

The string is denoted as the version/revision date and database manager of the database. Nowadays, this directive is used mainly for the purpose of keeping the development and revision history mostly updated.

#### EXAMPLE

```
<code>VERSION_DATE</code> Last update and adjustment: Database Manager, 2015-09-25 !
```

# Working with the Mobility Databases

### In this section:

About the Diffusion Module (DICTRA) and the Databases	. 44
Extensions to Database Definition File Syntax	. 45
PARAMETER	. 45
DIFFUSION	46
ZEROVOLUME_SPECIES	47

# About the Diffusion Module (DICTRA) and the Databases



The following is applicable to Console Mode.

A Diffusion Module (DICTRA) installation shares directories with Thermo-Calc, including the \DATA\ (or /data/) area and the database initiation file (TC\_INITD.TDB or initd.tdb).

When you want to switch or add self-generated databases/data-sets or any new database, use the predefined database list in the database initiation file as part of the Thermo-Calc installation packages.

You can add comment lines in the database initiation file, which must start with a \$ sign. These lines are ignored by the DATA module. This is also applicable if the database manager or user wants to temporarily disable a database in the predefined database list. If there are too many databases in the \DATA\ area that the DATA module cannot handle properly, the \$ sign can be used to temporarily comment about the uncommon databases. However, this does not often happen.

If Thermo-Calc is used with a Windows platform, such additional database initiation files can be located at any directory of any driver, on either a local computer or connected server. The NEW\_DICTORY\_FILE command displays an **Open** window to access a database initiation file if the file name or its path is not given on the same line of the NEW\_DICTORY\_FILE command, or if it is incomplete or incorrect, so that the path (in the Look in box) and database initiation file name (in the File name box) can be appropriately selected. However, if Thermo-Calc is run on a Linux/UNIX platform, these files must be located in the current working directory (where Thermo-Calc is started).

In an additional database initiation file, the first database entry may need to have the same path definition structure as in the ordinary database initiation file. Copy the entry line(s) for common databases from the original file to a database initiation file. The entries of the additional databases follow. Similar to the standard databases predefined in the database initiation file, all databases should normally be located in subdirectories under the directory as defined by the TCPATH parameter (Windows) or the TC\_DATA parameter (Linux/UNIX), or under its subdirectory  $\Delta TAA$  or /data/.

### WINDOWS DATABASE INITIATION FILE EXAMPLE

For a Windows platform, the following example is an additional database initiation file, called MYINITD1.TDB:

```
$
$ DATABASES TCC (Additional TCC Databases)
PURE5 TCPATH\DATA\PURE5\PURE5SETU.TDB SGTE Pure Elements
Database, version 5 !
AD1
     TCPATH\DATA\ADD1\AD1SETUP.TDB TCS ADD1 Solution
Database, version 1 !
     TCPATH\ADDDATA\ADD2\AD2SETUP.TDB TCS ADD2 Solution Database
AD2
1
$AD20 TCPATH\ADDDATA\ADD20ld\AD2SETUP.TDB TCS ADD2 Database
(old) !
AD3
     TCPATH\DATA\NEWDATA\ADD3\AD3SETUP.TDB TCS ADD3 Solution
Database !
AD4
     TCPATH\DATA\NEWDATA\MYPROJ1\ADD4\AD4SETUP.TDB
     MYPROJECT1 ADD4 Solution Database !
$
$ DATABASES DIC (Additional DICTRA Databases)
DCAD1 TCPATH\DICDATA\DCADD1\DCAD1SET.TDB
          TCS DCADD1 Mobility Database !
```

### **Extensions to Database Definition File Syntax**

Software packages for simulation of diffusional phase transformations, such as the Diffusion Module (DICTRA) and Precipitation Module (TC-PRISMA), need both thermodynamic data and kinetic data (i.e. diffusivities or mobilities). Naturally, the handling and storage of kinetic data also benefits from the use of some kind of database management. Thus, the TDB database definition file syntax has been extended to incorporate some new keywords needed for storing kinetic data - <u>"PARAMETER" on page 27</u>, <u>"DIFFUSION" on the next page</u>, and <u>"ZEROVOLUME\_SPECIES " on page 47</u>.

### PARAMETER

This keyword allows you to enter all types of normal GIBBS parameters for thermodynamic data as well as five special extensions suitable for kinetic data used in the Diffusion Module (DICTRA).

Valid extensions to special GIBBS parameter names are:

- MQ: Activation enthalpy for mobility equation.
- MF: Pre-exponential factor for mobility equation.
- DQ: Activation enthalpy for diffusivity equation.
- DF: Pre-exponential factor for diffusivity equation.
- vs: Volume counted per mole of volume carrying species.

#### **EXAMPLES**

```
PARAMETER MQ(BCC,FE:VA) 298.15 1000+200*T+...; 6000 N !
PARAMETER MF(BCC,CO:VA) 298.15 1000+200*T+...; 6000 N !
PARAMETER DQ(FCC,FE:VA) 298.15 1043+...; 6000 N 10 !
PARAMETER DF(FCC,CR:C ) 298.15 1000+200*T+...; 6000 N 10 !
PARAMETER VS(FCC) 298.15 1000+200*T+...; 6000 N 11 !
```

### DIFFUSION

DIFFUSION [model keyword] [phase name] [additional parameter(s)] !

This keyword specifies the type of diffusion model to use for a phase if the default model is not desired. The default model calculates the full diffusion matrix. A diffusivity is calculated from the different mobilities and the thermodynamic factors. The former ones are calculated as:

M = exp ( $\Sigma$ MF/RT) exp ( $\Sigma$ MQ/RT) / RT

where  $\Sigma$  stands for a weighted summation of the different MF's and MQ's plus possibly a Redlich-Kister term.

#### VALID MODEL KEYWORDS

Keyword	Description
NONE	No diffusion in this phase

Keyword	Description
DILUTE	Constitution list of dependent species in each sublattice must be given as an additional parameter. Only the diagonal terms in the diffusion matrix are calculated. D = exp ( $\Sigma$ DF/RT) exp ( $\Sigma$ DQ/RT).
SIMPLE	Constitution list of dependent species in each sublattice must be given as additional parameter. Only the diagonal terms in the diffusion matrix are calculated. D = $\Sigma$ DF + $\Sigma$ DQ.
MAGNETIC	The so-called ALPHA and ALPHA2 parameters must be given as additional parameters. ALPHA is for the substitutional magnetic model and ALPHA2 for the interstitial one. By appending an $\&$ sign and a species name after the alpha keyword one can supply individual values for the different species. The full diffusion matrix is calculated.

#### **EXAMPLES**

```
DIFFUSION NONE SIGMA !
DIFFUSION DILUTE CEMENTITE : FE : C : !
DIFFUSION MAGNETIC BCC_A2 ALPHA=0.3 ALPHA2&C=1.8 ALPHA2&N=0.6 !
```

# **ZEROVOLUME\_SPECIES**

ZEROVOLUME SPECIES [list of species] !

In the Diffusion Module (DICTRA), the assumption that the volume is carried by the substitutional elements only is applied. The interstitial elements are assumed to have zero molar volumes. This keyword uses a list of which species are to be considered as zero volume ones for an argument.



There is no underscore (\_) between Zero and Volume.

#### **EXAMPLE**

ZEROVOLUME\_SPECIES VA C N !

# **Database Definition File Examples**

### In this section:

Example 1: A Steel Database	. 49
Example 2: A Custom Database for the Sb-Sn Binary System	. 50
Example 3: A Public Database for the Fe-Cr-C Ternary System	.53

# **Example 1: A Steel Database**

```
TEMP-LIM 500.0 2000.0 !
Ś
$ELEMENT, NAME, REF.STATE, ATOMIC-MASS, H0, S0 !
ELEMENT VA VACUUM 0.0 0.0 0.0 !
ELEMENT C GRAPHITE 12.011 0.0 0.0 !
ELEMENT V BCC
                        50.9415 0.0 0.0 !
ELEMENT CR BCC-PARAMAGNETIC 51.996 0.0 0.0 !
ELEMENT FE FCC-PARAMAGNETIC 55.847 0.0 0.0 !
ELEMENT NI FCC-PARAMAGNETIC 58.69 0.0 0.0 !
                95.94 0.0 0.0 !
ELEMENT MO BCC
ELEMENT W BCC 183.85 0.0 0.0 !
Ś
$PHASE, NAME, TYPE, NR-OF-SUBL, SITES-IN-EACH-SUBL. !
PHASE BCC
           B1M 2 1.0 3.0 !
PHASE FCC
             F2M 2 1.0 1.0 !
PHASE HCP
            0 2 2.0 1.0 !
PHASE LIQUID 3 2 1.0 1.0 !
PHASE CEMENTITE 4 2 3.0 1.0 !
PHASE M23C6 4 2 23.0 6.0 !
PHASE M7C3
            4 2 7.0 3.0 !
             4 4 2.0 2.0 2.0 1.0 !
PHASE M6C
PHASE SIGMA 0 3 10.0 4.0 16.0 !
PHASE MU-PHASE 0 3 7.0 2.0 4.0 !
PHASE R-PHASE 0 3 27.0 14.0 12.0 !
PHASE GRAPHITE 4 1 1.0 !
$
$CONSTITUENT, PHASE-NAME : CONSTITUENTS !
CONSTITUENT BCC :V CR FE NI MO W:VA C: !
CONSTITUENT FCC :V CR FE NI MO W:VA C: !
CONSTITUENT HCP :CR FE NI:VA C N: !
CONSTITUENT LIQUID :C V CR FE NI MO W VA:VA C: !
CONSTITUENT CEMENTITE :CR FE:C: !
CONSTITUENT M23C6 :CR FE:C: !
CONSTITUENT M7C3 :CR FE:C: !
CONSTITUENT M6C :FE:W:FE W:C: !
```

```
CONSTITUENT SIGMA :FE:V CR MO:FE V CR MO: !
CONSTITUENT MU-PHASE :FE:MO W:FE MO W: !
CONSTITUENT R-PHASE :FE:MO :FE MO: !
CONSTITUENT GRAPHITE :C: !
Ś
$TYPE DEFINITIIONS:
TYPE-DEFINITION 0 SEQ TCPATH\DATA\METDATA\TC-THEREST.TDB !
TYPE-DEFINITION 1 SEQ TCPATH\DATA\METDATA\TC-BCC.TDB !
TYPE-DEFINITION 2 SEQ TCPATH\DATA\METDATA\TC-FCC.TDB !
TYPE-DEFINITION 3 SEQ TCPATH\DATA\METDATA\TC-LIQUID.TDB !
TYPE-DEFINITION 4 SEQ TCPATH\DATA\METDATA\TC-CARBIDES.TDB !
TYPE-DEFINITION M SEO TCPATH\DATA\METDATA\TC-CURIE-BOHR.TDB !
TYPE-DEFINITION B GES AM-PH BCC MAGNETIC -1 .4 !
TYPE-DEFINITION F GES AM-PH FCC MAGNETIC -3 .28 !
Ś
$DEFAULT COMMANDS:
DEFAULT-COMMAND DEF ELEMENT VA !
DEFAULT-COMMAND REJ SYS-CONST LIQUID 1 VA !
$DATABASE INFORMATION:
DATABASE-INFO The following binary and ternary systems are available: '
  FE-CR-NI by Hertzman'
  FE-MO
            Fernandez'
 FE-CR-C
            Andersson'
  FE-W-C
           Gustafson'
  FE-W Andersson & Gustafson' !
```

# Example 2: A Custom Database for the Sb-Sn Binary System

```
      $ELEMENT, NAME, REF.STATE, ATOMIC-MASS, H0, S0 !

      ELEM VA VACUUM
      0.0
      0.0
      0.0 !

      ELEM MG HCP(A3)
      24.305
      0.0
      0.0 !

      ELEM SB RHOMBOHEDRAL(A7)
      121.75
      0.0
      0.0 !

      ELEM SN BCT(A5)
      118.69
      0.0
      0.1 !

      ELEM /- ELECTRON-GAS
      0
      0
      !
```

```
$SPECIES, NAME, STOICHIOMETRIC-FORMULA !
SPECIE MG1 MG1!
SPECIE MG2 MG2!
SPECIE MG2+ MG/+2!
SPECIE SB1 SB1!
SPECIE SB2 SB2!
SPECIE SB4 SB4!
SPECIE SB3- SB/-3!
SPECIE SB5- SB/-5!
SPECIE SN1 SN1!
SPECIE SN4- SN/-4!
Ś
$PHASE, NAME, TYPE, NR-OF-SUBL, SITES-IN-EACH-SUBL. !
PHASE BCT
           Z 1 1.0!
PHASE HCP
              Z 1 1.0!
PHASE RHOMBO
              Z 1 1.0!
PHASE GAS:G Z 1 1.0!
PHASE LIQUID:L Z 1 1.0!
PHASE IONICLIQ:Y Z 2 1 1!
PHASE SPLIQ:Y Z 2 1 1!
PHASE BMG3SB2:I Z 2 3 2!
PHASE AMG3SB2:I Z 2 3 2!
PHASE MG2SN:I Z 2 2 1!
PHASE SBSN Z 2 1 1!
PHASE SB2SN3 Z 2 2 3!
$
$CONSTITUENT, PHASE-NAME : CONSTITUENTS !
CONSTITUENT RHOMBO :SB SN:!
CONSTITUENT HCP :MG SN:!
CONSTITUENT BCT :SB SN:!
CONSTITUENT GAS:G :MG1 MG2 SB1 SB2 SB4 SN1:
 > Gas phase, using the Ideal EOS and Mixing Model. !
CONSTITUENT LIQUID:L :SB SN:!
CONSTITUENT IONICLIQ:Y :MG2+:SB SB3- SN SN4- VA:
 > This is the Ionic Liquid Solution Phase. !
CONSTITUENT SPLIQ:Y :MG2+:SB SB3- SN SN4- VA:!
CONSTITUENT BMG3SB2:I :MG2+:SB3- SB5- VA SN4-:!
```

```
CONSTITUENT AMG3SB2:I :MG2+:SB3- VA SN4-:!
CONSTITUENT MG2SN:I :MG2+ VA:SB3- SN4-:!
CONSTITUENT SBSN :SB SN:SB SN:!
CONSTITUENT SB2SN3 :SB:SN:!
Ś
$DEFAULT COMMANDS:
DEFAULT-COM DEF-ELEM VA /-!
DEFAULT-COM REJ-PHASE LIQUID!
DEFAULT-COM REJ-PHASE SPLIQ!
$
$TYPE DEFINITIIONS:
TYPE-DEFINITION Z SEQ * !
Ś
$DATABASE INFORMATION:
DATABASE_INFO The Sb-Sn system with isentropic temperatures!
$
$VERSION DATE:
VERSION DATE Last update 1986-05-18 11:39:49 !
$
Ś
$ HERE COMES THE THERMODYNAMIC DATA (expressed in functions & parameters):
Ś
FUNCTION MGLIQUID 298.15 -4630.90976+192.994374*T-34.0888057*T*LOG(T)
   -36544605.6*T**(-2); 6000 N!
$
FUNCTION MGSOLID 298.15 -8367.34+143.677876*T-26.1849785*T*LOG(T)
   +4.858E-4*T**2-1.393669E-6*T**3+78950*T**(-1);
   923.00 Y -13804.4772 +202.909445*T-34.0888057*T*LOG(T)
   -3.65446056E7*T**(-2) +1.06753982E28*T**(-9); 6000 N!
$
FUNCTION SBLIQUID 298.15 9071.98+146.800*T-31.38*T*LOG(T)
  -2.441646E8*T**(-2); 6000 N!
$
. . . . .
..... <more>
$
FUNCTION LFCT 298.15 -17325.6+5.03600*T; 6000 N!
```

```
FUNCTION GFCTSBSN 298.15 LFCT+SBSOLID+SNSOLID+2948.291+3721.286;
   6000 N!
FUNCTION ISB 298.15 15000; 6000 N!
FUNCTION ISN 298.15 47199.9-95.6270*T; 6000 N!
Ś
. . . . .
..... <more>
Ś
PARAMETER G(RHOMBO, SB; 0) 298.15 SBSOLID; 6000 N!
PARAMETER G(RHOMBO, SN; 0) 298.15 2035+SNSOLID; 6000 N!
$
PARAMETER G(HCP, MG; 0) 298.15 MGSOLID; 6000 N!
PARAMETER G(HCP, SN; 0) 298.15 32000+SNSOLID; 6000 N!
PARAMETER G(HCP, MG, SN; 0) 298.15 -69566-9.23183*T; 6000 N!
Ś
PARAMETER G(BCT, SN;0) 298.15 SNSOLID; 6000 N!
PARAMETER G(BCT, SB;0) 298.15 1000+SBSOLID; 6000 N!
PARAMETER G(BCT,SB,SN;0) 298.15 0.5*ISB+0.5*ISN; 6000 N!
PARAMETER G(BCT, SB, SN;1) 298.15 0.5*ISB-0.5*ISN; 6000 N!
Ś
PARAMETER G(IONICLIQ, MG2+:SB3-;0) 298.15 -204389-4.98506*T
   -2.75637E9*T**(-2)+3*MGLIQUID+2*SBLIQUID; 6000 N!
PARAMETER G(IONICLIQ, MG2+:SN4-;0) 298.15 -98639.5+881.073*T
   -174.523*T*LOG(T)-1.79808E9*T**(-2); 6000 N!
PARAMETER G(IONICLIQ, MG2+:SB;0) 298.15 SBLIQUID; 6000 N!
$
. . . . . .
..... <more>
Ś
```

# **Example 3: A Public Database for the Fe-Cr-C Ternary System**

```
$ Revision history:
$ Created as PDEMO by Pingfang Shi on 2004-10-05
$ Renamed to DFeCrC by Pingfang Shi on 2006-10-25
```

```
$
$ FURTHER MODIFICATIONS:
$ pfs: /20041005 (PDEMO)
Ś
      * Retrieved all definitions from PTERN for the Fe-Cr-C ternary !
$ pfs: /20061025 (DFeCrC)
      * Ignore ELEMENT /- definition.
Ś
$
      * Modify the L(BCC_A2,FE:C,VA;0) [and L(BCC_A2,CR:C,VA;0)]
$
        parameters, in order to avoid BCC-appearance (Fe-C)
        at temperatures higher than 3900 K.
Ś
          $L(BCC A2,CR:C,VA;0) 298.15 -190*T;
$
                                               6000 N REF1 !
          L(BCC A2,CR:C,VA;0) 298.15 -190*T; 3000 Y 0; 6000 N REF1 !
Ś
          $L(BCC A2,FE:C,VA;0) 298.15 -190*T;
$
                                          6000 N REF3 !
Ś
          L(BCC A2,FE:C,VA;0) 298.15 -190*T; 3900 Y 0; 6000 N REF3 !
$
      * Add Cr into HCP_A3 phase, and
        add all necessary GO/L parameters (according to SSOL2/TCFE3):
Ś
          G(HCP A3, CR:VA;0)
$
          TC(HCP A3, CR:VA;0)
$
$
          BMAGN(HCP A3, CR:VA;0)
$
          G(HCP A3, FE:C;0)
$
          G(HCP_A3,CR:C;0)
          L(HCP A3, CR:C, VA; 0)
$
$
          L(HCP A3, CR, FE:VA;0)
Ś
      * Add "TMM:300/3000" limits to all three binary joins in the
          ASSESSED SYSTEM section.
Ś
$
      * Change the reference indices, e.g., 101 to REF1, in order to
Ś
          always get the reference list correctly.
$
      * Re-arrange database format slightly.
Ś
      * Updated the DATABASE INFO section.
$-----
$
$ DATABASE INFORMATION:
$
DATABASE INFO
                                        DFeCrC '
```

TCS Demo Fe-Cr-C Alloy Solutions Database'

(only for demonstration purposes in TCC-Demo/TCW-Demo)'

\*\*\*\*\*

(Version 1.2, Oct. 2006) ''

Copyright @ 2004-2012: Thermo-Calc Software AB, Sweden '' The DFeCrC database (previously called PDEMO) is the TCS Demo Fe-Cr-C ' Alloy Solutions Database, which is especially designed for primarily ' demonstrating the uses of: '

 $^{\star}$  TCC-Demo software: BIN, TERN, SCHEIL and some other modules; '

\* TCW-Demo software: '

"Binary" module calculations of phase diagrams and property diagrams' in available binary systems; '

"Ternary" module calculations of phase diagrams (e.g., isothermal' sections, monovariant lines involving liquid, liquidus' surface projections) and property diagrams in available' ternary systems; '

"Scheil" module simulations of alloy solidifications; and/or ' "Equilibrium" calculation routines for phase diagrams and property' diagrams in available binary or ternary systems. '

It can also be used for other types of calculations/simulations in both' TCC and TCW software, and in their application programming interfaces.'' It includes critically-assessed data for all possible three binary joins' (i.e. Cr-C, Fe-Cr and Fe-Cr) and one complete ternary alloy solutions ' within the Fe-Cr-C system. However, due to the fact that experimental' data for the Fe-C and Cr-C binary joins and Fe-Cr-C ternary system at ' extremely-high temperatures are not sufficient, the available assessed' data can not be applied to temperature conditions higher than 4000 K.'' Included thermodynamic data are available for various stoichiometric ' and solution phases, e.g. liquid mixture, various alloy solutions and ' intermetallic phases. But the gaseous mixture phase is excluded in this' DEMO version.''

The DFeCrC database can be used not only in calculating various types ' of phase diagrams (binary phase diagrams, ternary isothermal sections,' ternary monovariant lines involving liquid, ternary liquidus surface ' projections, ternary isopleth sections, etc.) and property diagrams ' (the easiest ways are through the BIN and TERN modules), but also in ' simulating alloy solidification processes (with the SCHEIL module) of'

```
Fe-based or Cr-based alloys (but limited within the Fe-Cr-C ternary '
system in this DEMO version). Many unique features of the TCC/TCW '
software can be demonstrated using this and other specially-designed '
DEMO databases. ''
However, this DFeCrC database (similar to other DEMO and/or PUBLIC TC '
databases) is provided within the TCC-Demo/TCW-Demo software only for '
the purposes of demonstration, testing and evaluation. For R&D projects'
and teaching activities, you are highly encouraged to only use the FULL'
versions of the TCC and/or TCW software together with some commercial '
databases that are appropriate for your specific applications. Please '
contact us for all kinds of details. ''
Release History: Version 1.1 with minor improvements, Oct. 2004 '
               Version 1.2 with minor modifications, Oct. 2006 ''
Edited by: Dr. Pingfang Shi (Thermo-Calc Software, Oct. 2006). ''
1
$ -----
```

\$VERSION\_DATE Last update and adjustment: Pingfang Shi, 2004-10-05 !
VERSION\_DATE Last update and adjustment: Pingfang Shi, 2006-10-25 !

```
$
```

```
$ Definition of Elements in the Database System:
$-----
$ELEM NAME STABLE_ELEMENT_REF ATOMIC MASS H298-H0 S298 !
$-----
$ELEMENT /- ELECTRON GAS
                      0.0000E+00 0.0000E+00 0.0000E+00!
ELEMENT VA VACUUM
                     0.0000E+00 0.0000E+00 0.0000E+00!
ELEMENT C GRAPHITE
                     1.2011E+01 1.0540E+03 5.7400E+00!
ELEMENT CR BCC A2
                     5.1996E+01 4.0500E+03 2.3560E+01!
ELEMENT FE BCC A2
                     5.5847E+01 4.4890E+03 2.7280E+01!
Ś
```

```
$
$ TYPE DEFINITIONS for phase descriptions:
$-----
$... For magnetic contributions:
TYPE_DEFINITION A GES AMEND_PHASE_DES @ MAGNETIC -1.0 4.00000E-01 !
TYPE DEFINITION B GES AMEND PHASE DES @ MAGNETIC -3.0 2.80000E-01 !
$... For Cr/Fe-C in FCC phase and Cr/Fe-C in HCP phase:
TYPE DEFINITION C IF(C) THEN GES AMEND PHASE DES 0 C S ,, :C: !
Ś
$ Default Commands:
$-----
$DEFINE SYSTEM DEFAULT SPECIES 2 !
DEFINE SYSTEM DEFAULT ELEMENT 2 !
DEFAULT COMMAND DEF SYS ELEMENT VA !
$
$ FUNCTIONS for various phases:
$-----
Ś
FUNCT GHSERCC 298.15 -17368.441+170.73*T-24.3*T*LN(T)
   -4.723E-04*T**2+2562600*T** (-1) -2.643E+08*T** (-2)
   +1.2E+10*T**(-3); 6000 N!
FUNCT GHSERCR 298.15 -8856.94+157.48*T-26.908*T*LN(T)
   +.00189435*T**2-1.47721E-06*T**3+139250*T**(-1); 2180 Y
    -34869.344+344.18*T-50*T*LN(T)-2.88526E+32*T**(-9); 6000 N!
FUNCT GHSERFE 298.15 +1225.7+124.134*T-23.5143*T*LN(T)
   -.00439752*T**2-5.8927E-08*T**3+77359*T**(-1); 1811.00 Y
   -25383.581+299.31255*T-46*T*LN(T)+2.29603E+31*T**(-9); 6000 N!
FUNCT GFELIQ 298.15 +GHSERFE#+12040.17-6.55843*T
```

-3.6751551E-21\*T\*\*7; 1811 Y -10839.7+291.302\*T-46\*T\*LN(T); 6000 N! FUNCT GFEFCC 298.15 +GHSERFE#-1462.4+8.282\*T -1.15\*T\*LN(T)+6.4E-04\*T\*\*2; 1811 Y -27098.266+300.25256\*T-46\*T\*LN(T)+2.78854E+31\*T\*\*(-9); 6000 N! FUNCT GFECEM 298.15 +GPCEM1#-10745+706.04\*T-120.6\*T\*LN(T); 6000 N! FUNCT GFEM23C6 298.15 +7.666667\*GFECEM#-1.6666667\*GHSERCC# +66920-40\*T; 6000 N! FUNCT GCRFCC 298.15 +GHSERCR#+7284+.163\*T; 6000 N! FUNCT GCRM23C6 298.15 -521983+3622.24\*T-620.965\*T\*LN(T) -.126431\*T\*\*2; 6000 N! FUNCT GCRM3C2 298.15 -100823.8+530.66989\*T-89.6694\*T\*LN(T)

-.0301188\*T\*\*2; 6000 N! FUNCT GCRM7C3 298.15 -201690+1103.128\*T-190.177\*T\*LN(T) -.0578207\*T\*\*2; 6000 N!

FUNCT	GPCLIQ	298.15	+YCLIQ#*EXP(ZCLIQ#); 6	1 000	Ν!	
FUNCT	ACLIQ	298.15	+2.32E-05*T+2.85E-09*T**2	; 60	000	N!
FUNCT	BCLIQ	298.15	+1+3.2E-10*P;	60	000	N!
FUNCT	CCLIQ	298.15	1.6E-10;	60	000	N!
FUNCT	DCLIQ	298.15	+1*LN(BCLIQ#);	60	000	N!
FUNCT	ECLIQ	298.15	+1*LN(CCLIQ#);	60	000	N!
FUNCT	VCLIQ	298.15	+7.626E-06*EXP(ACLIQ#);	60	000	N!
FUNCT	XCLIQ	298.15	+1*EXP(.5*DCLIQ#)-1;	60	000	N!
FUNCT	YCLIQ	298.15	+VCLIQ#*EXP(-ECLIQ#);	60	000	N!
FUNCT	ZCLIQ	298.15	+1*LN(XCLIQ#);	60	000	N!

FUNCT	GPCGRA	298.15	+YCGRA#*EXP(ZCGRA#);	5000	N!	
FUNCT	ACGRA	298.15	+2.32E-05*T+2.85E-09*T**2	2; (	6000	N!
FUNCT	BCGRA	298.15	+1+3.6E-10*P;	(	6000	N!
FUNCT	CCGRA	298.15	3.3E-10;	(	6000	N!
FUNCT	DCGRA	298.15	+1*LN(BCGRA#);	(	6000	N!
FUNCT	ECGRA	298.15	+1*LN(CCGRA#);	(	6000	N!
FUNCT	VCGRA	298.15	+5.259E-06*EXP(ACGRA#);	(	6000	N!
FUNCT	XCGRA	298.15	+1*EXP(.9166667*DCGRA#)-1	L; (	6000	N!
FUNCT	YCGRA	298.15	+VCGRA#*EXP(-ECGRA#);	(	6000	N!

```
FUNCT ZCGRA
              298.15 +1*LN(XCGRA#);
                                                6000 N!
FUNCT GPCFCC
              298.15 +YCFCC#*EXP(ZFEFCC#); 6000 N!
FUNCT ACFCC
              298.15 +1.44E-04*T;
                                             6000 N!
FUNCT VCFCC
              298.15 +1.031E-05*EXP(ACFCC#); 6000 N!
FUNCT YCFCC
              298.15 +VCFCC#*EXP(-EFEFCC#); 6000 N!
FUNCT GPCRLIQ 298.15 +YCRLIQ#*EXP(ZCRLIQ#); 6000 N!
FUNCT ACRLIQ
              298.15 +1.7E-05*T+9.2E-09*T**2; 6000 N!
FUNCT BCRLIQ
              298.15 +1+4.65E-11*P;
                                             6000 N!
FUNCT CCRLIQ
              298.15 3.72E-11;
                                             6000 N!
FUNCT DCRLIO
              298.15 +1*LN(BCRLIO#);
                                            6000 N!
FUNCT ECRLIO
              298.15 +1*LN(CCRLIO#);
                                            6000 N!
FUNCT VCRLIQ
              298.15 +7.653E-06*EXP(ACRLIQ#); 6000 N!
FUNCT XCRLIQ
              298.15 +1*EXP(.8*DCRLIQ#)-1;
                                            6000 N!
FUNCT YCRLIQ
              298.15 +VCRLIQ#*EXP(-ECRLIQ#); 6000 N!
FUNCT ZCRLIQ
              298.15 +1*LN(XCRLIQ#);
                                             6000 N!
FUNCT GPCRBCC 298.15 +YCRBCC#*EXP(ZCRBCC#); 6000 N!
FUNCT ACRBCC
              298.15 +1.7E-05*T+9.2E-09*T**2; 6000 N!
FUNCT BCRBCC
              298.15 +1+2.6E-11*P;
                                             6000 N!
FUNCT CCRBCC
              298.15 2.08E-11;
                                             6000 N!
FUNCT DCRBCC
              298.15 +1*LN(BCRBCC#);
                                            6000 N!
FUNCT ECRBCC
              298.15 +1*LN(CCRBCC#);
                                            6000 N!
FUNCT VCRBCC
              298.15 +7.188E-06*EXP(ACRBCC#); 6000 N!
              298.15 +1*EXP(.8*DCRBCC#)-1; 6000 N!
FUNCT XCRBCC
FUNCT YCRBCC
              298.15 +VCRBCC#*EXP(-ECRBCC#); 6000 N!
FUNCT ZCRBCC
              298.15 +1*LN(XCRBCC#);
                                            6000 N!
FUNCT GPFELIQ 298.15 +YFELIQ#*EXP(ZFELIQ#); 6000 N!
FUNCT AFELIQ
              298.15 +1.135E-04*T;
                                                            6000 N!
              298.15 +1+4.98009787E-12*P+3.20078924E-14*T*P; 6000 N!
FUNCT BFELIQ
FUNCT CFELIQ
              298.15 +4.22534787E-12+2.71569924E-14*T;
                                                            6000 N!
FUNCT DFELIQ 298.15 +1*LN(BFELIQ#);
                                                            6000 N!
FUNCT EFELIO 298.15 +1*LN(CFELIO#);
                                                            6000 N!
FUNCT VFELIQ 298.15 +6.46677E-06*EXP(AFELIQ#);
                                                            6000 N!
FUNCT XFELIQ 298.15 +1*EXP(.8484467*DFELIQ#)-1;
                                                           6000 N!
```

```
FUNCT YFELIQ 298.15 +VFELIQ#*EXP(-EFELIQ#);
                                                            6000 N!
FUNCT ZFELIQ 298.15 +1*LN(XFELIQ#);
                                                            6000 N!
FUNCT GPFEFCC 298.15 +YFEFCC#*EXP(ZFEFCC#); 6000 N!
FUNCT AFEFCC 298.15 +7.3097E-05*T;
                                                            6000 N!
FUNCT BFEFCC 298.15 +1+3.25236341E-11*P+3.36607808E-16*T*P; 6000 N!
FUNCT CFEFCC 298.15 +2.62285341E-11+2.71455808E-16*T;
                                                           6000 N!
FUNCT DFEFCC 298.15 +1*LN(BFEFCC#);
                                                           6000 N!
FUNCT EFEFCC 298.15 +1*LN(CFEFCC#);
                                                            6000 N!
FUNCT VFEFCC
             298.15 +6.688726E-06*EXP(AFEFCC#);
                                                            6000 N!
FUNCT XFEFCC 298.15 +1*EXP(.8064454*DFEFCC#)-1;
                                                            6000 N!
FUNCT YFEFCC 298.15 +VFEFCC#*EXP(-EFEFCC#);
                                                            6000 N!
FUNCT ZFEFCC 298.15 +1*LN(XFEFCC#);
                                                            6000 N!
```

FUNCT	GPFEBCC	298.15	+YFEBCC#*EXP(ZFEBCC#); 6000 N!	
FUNCT	AFEBCC	298.15	+2.3987E-05*T+1.2845E-08*T**2;	6000 N!
FUNCT	BFEBCC	298.15	+1+2.80599565E-11*P+3.06481523E-16*T*P;	6000 N!
FUNCT	CFEBCC	298.15	+2.20949565E-11+2.41329523E-16*T;	6000 N!
FUNCT	DFEBCC	298.15	+1*LN(BFEBCC#);	6000 N!
FUNCT	EFEBCC	298.15	+1*LN(CFEBCC#);	6000 N!
FUNCT	VFEBCC	298.15	+7.042095E-06*EXP(AFEBCC#);	6000 N!
FUNCT	XFEBCC	298.15	+1*EXP(.7874195*DFEBCC#)-1;	6000 N!
FUNCT	YFEBCC	298.15	+VFEBCC#*EXP(-EFEBCC#);	6000 N!
FUNCT	ZFEBCC	298.15	+1*LN(XFEBCC#);	6000 N!

```
FUNCT GPFEHCP 298.15 +YFEHCP#*EXP(ZFEHCP#); 6000 N!
FUNCT AFEHCP 298.15 +7.3646E-5*T;
                                                            6000 N!
FUNCT BFEHCP 298.15 +1+32.5236341E-12*P+3.36607808E-16*P*T; 6000 N!
FUNCT CFEHCP 298.15 +26.2285341E-12+2.71455808E-16*T;
                                                          6000 N!
FUNCT DFEHCP 298.15 +LOG(BFEHCP#);
                                                           6000 N!
FUNCT EFEHCP 298.15 +LOG(CFEHCP#);
                                                            6000 N!
FUNCT VFEHCP 298.15 +6.59121E-6*EXP(AFEHCP#);
                                                           6000 N!
FUNCT XFEHCP 298.15 +EXP(0.8064454*DFEHCP#)-1;
                                                           6000 N!
FUNCT YFEHCP 298.15 +VFEHCP#*EXP(-1*EFEHCP#);
                                                           6000 N!
FUNCT ZFEHCP 298.15 +LOG(XFEHCP#);
                                                            6000 N!
```

FUNCT GPCEM1 298.15 +VCEM1#\*P; 6000 N!

```
FUNCT ACEM1 298.15 -1.36E-05*T+4E-08*T**2;
                                      6000 N!
                                      6000 N!
FUNCT VCEM1 298.15 +2.339E-05*EXP(ACEM1#);
FUNCT GPSIG1 298.15 +1.09E-04*P;
                                 6000 N!
FUNCT GPSIG2 298.15 +1.117E-04*P;
                                6000 N!
$
$ Define the various Phase and their Constituents, and
$ Assign parameters to phases:
$-----
$PHASE NAME:TYPE MARKCODE #SUBL SITES IN EACH SUBL. !
$-----
PHASE LIQUID:L % 1 1.0
 > This is metallic liquid solution phase, with C species !
CONST LIQUID:L : C, CR, FE : !
PARAM G(LIQUID,C;0) 298.15 +GHSERCC#+GPCLIO#
                         +117369-24.63*T; 6000 N REF0 !
PARAM G(LIQUID,CR;0) 298.15 +GHSERCR#+GPCRLIQ#
              +24339.955-11.420225*T+2.37615E-21*T**7; 2180 Y
                        +GHSERCR#+GPCRLIQ#
              +18409.36-8.563683*T+2.88526E+32*T**(-9); 6000 N REF0 !
PARAM G(LIQUID, FE; 0) 298.15 +GFELIQ#+GPFELIQ#; 6000 N REF0 !
PARAM L(LIQUID,C,CR;0) 298.15 -90526-25.9116*T; 6000 N REF1 !
                                      6000 N REF1 !
PARAM L(LIQUID, C, CR; 1) 298.15 +80000;
PARAM L(LIQUID, C, CR; 2) 298.15 +80000;
                                      6000 N REF1 !
PARAM L(LIQUID, C, FE; 0) 298.15 -124320+28.5*T; 6000 N REF3 !
PARAM L(LIQUID,C,FE;1) 298.15 +19300;
                                     6000 N REF3 !
PARAM L(LIQUID, C, FE; 2) 298.15 +49260-19*T; 6000 N REF3 !
PARAM L(LIQUID,CR,FE;0) 298.15 -14550+6.65*T; 6000 N REF4 !
PARAM L(LIQUID,C,CR,FE;0) 298.15 -496063; 6000 N REF2 !
PARAM L(LIQUID, C, CR, FE; 1) 298.15 +57990; 6000 N REF2 !
PARAM L(LIQUID, C, CR, FE; 2) 298.15 +61404; 6000 N REF2 !
```

```
$PHASE FCC A1 %BC 2 1 1 !
$ Note the C TYPE DEF for 2nd FCC composition-set (MC) is not necessary
Ś
  for the (Fe,Cr)-C system.
PHASE FCC A1 %B 2 1 1 !
CONST FCC A1 : CR, FE%
            : C,VA% : !
PARAM G(FCC A1, CR:VA; 0) 298.15 +GCRFCC#+GPCRBCC#; 6000 N REF0 !
PARAM TC(FCC A1,CR:VA;0) 298.15 -1109; 6000 N REF0 !
PARAM BMAGN(FCC A1,CR:VA;0) 298.15 -2.46; 6000 N REF0 !
PARAM G(FCC A1, FE:VA; 0) 298.15 +GFEFCC#+GPFEFCC#; 6000 N REF0 !
PARAM TC(FCC A1,FE:VA;0) 298.15 -201; 6000 N REF0 !
PARAM BMAGN(FCC A1, FE:VA; 0) 298.15 -2.1; 6000 N REF0 !
PARAM G(FCC A1, CR:C;0) 298.15 +GHSERCR#+GHSERCC#
                            +1200-1.94*T; 6000 N REF2 !
PARAM G(FCC A1, FE:C;0) 298.15 +GFEFCC#+GHSERCC#+GPCFCC#
                            +77207-15.877*T; 6000 N REF3 !
PARAM TC(FCC A1,FE:C;0) 298.15 -201; 6000 N REF3 !
PARAM BMAGN(FCC_A1,FE:C;0) 298.15 -2.1; 6000 N REF3 !
PARAM L(FCC A1, CR:C, VA; 0) 298.15 -11977+6.8194*T; 6000 N REF2 !
PARAM L(FCC A1, FE:C, VA; 0) 298.15 -34671; 6000 N REF3 !
PARAM L(FCC A1, CR, FE:VA; 0) 298.15 +10833-7.477*T; 6000 N REF4 !
PARAM L(FCC A1,CR,FE:VA;1) 298.15 +1410; 6000 N REF4 !
PARAM L(FCC A1,CR,FE:C;0) 298.15 -74319+3.2353*T; 6000 N REF2 !
PHASE BCC A2 %A 2 1 3 !
CONST BCC_A2 : CR%, FE%
            : C,VA% : !
PARAM G(BCC A2, CR:VA;0) 298.15 +GHSERCR#+GPCRBCC#; 6000 N REF0 !
```

PARAM TC(BCC A2,CR:VA;0) 298.15 -311.5; 6000 N REF0 !

```
PARAM BMAGN(BCC A2,CR:VA;0) 298.15 -.01; 6000 N REF0 !
PARAM G(BCC A2,FE:VA;0) 298.15 +GHSERFE#+GPFEBCC#; 6000 N REF0 !
PARAM TC(BCC A2,FE:VA;0) 298.15 1043; 6000 N REF0 !
PARAM BMAGN (BCC A2, FE:VA; 0) 298.15 2.22; 6000 N REF0 !
PARAM G(BCC A2,CR:C;0) 298.15 +GHSERCR#+3*GHSERCC#+GPCRBCC#+3*GPCGRA#
                               +416000; 6000 N REF1 !
PARAM TC(BCC A2,CR:C;0) 298.15 -311.5; 6000 N REF1 !
PARAM BMAGN(BCC A2,CR:C;0) 298.15 -.008; 6000 N REF1 !
PARAM G(BCC A2, FE:C;0) 298.15 +GHSERFE#+3*GHSERCC#+GPFEBCC#+3*GPCGRA#
                               +322050+75.667*T; 6000 N REF3 !
PARAM TC(BCC A2,FE:C;0) 298.15 1043; 6000 N REF3 !
PARAM BMAGN(BCC A2, FE:C; 0) 298.15 2.22; 6000 N REF3 !
$ PF-20061025: Modify the L(BCC A2,FE:C,VA;0) [and L(BCC A2,CR:C,VA;0)]
        parametera, in order to avoid BCC-appearance (Fe-C)
Ś
         at temperatures higher than 3900 K.
Ś
$PARAM L(BCC A2,CR:C,VA;0) 298.15 -190*T; 6000 N REF1 !
PARAM L(BCC A2, CR:C, VA; 0) 298.15 -190*T; 3000 Y 0; 6000 N REFS !
$PARAM L(BCC_A2,FE:C,VA;0) 298.15 -190*T; 6000 N REF3 !
PARAM L(BCC A2,FE:C,VA;0) 298.15 -190*T; 3900 Y 0; 6000 N REFS !
Ś
PARAM L(BCC A2,CR,FE:VA;0) 298.15 +20500-9.68*T; 6000 N REF4 !
PARAM TC (BCC A2, CR, FE:VA; 0) 298.15 1650; 6000 N REF4 !
PARAM TC (BCC A2, CR, FE:VA; 1) 298.15 550; 6000 N REF4 !
PARAM BMAGN(BCC A2, CR, FE:VA;0) 298.15 -.85; 6000 N REF4 !
PARAM L(BCC A2, CR, FE:C; 0) 298.15 -1250000+667.7*T; 6000 N REF2 !
PARAM TC(BCC A2,CR,FE:C;0) 298.15 1650; 6000 N REF5 !
PARAM TC (BCC A2, CR, FE:C;1) 298.15 550; 6000 N REF5 !
PARAM BMAGN(BCC A2,CR,FE:C;0) 298.15 -.85; 6000 N REF5 !
$PHASE HCP A3 %BC 2 1 .5 !
$ Note the C TYPE DEF for 2nd HCP composition-set (M2C) is not necessary
$ for the (Fe,Cr)-C system.
PHASE HCP A3 %B 2 1 .5 !
```

```
CONST HCP_A3 : CR,FE,
            : VA%,C : !
PARAM G(HCP_A3,CR:VA;0) 298.15 +GHSERCR#+4438; 6000 N REF1 !
PARAM TC(HCP A3,CR:VA;0) 298.15 -1109; 6000 N REF1 !
PARAM BMAGN(HCP A3, CR:VA; 0) 298.15 -2.46; 6000 N REF1 !
PARAM G(HCP A3, FE:VA;0) 298.15 +GHSERFE#+GPFEHCP#
            -3705.78+12.591*T-1.15*T*LN(T)+6.4E-04*T**2; 1811 Y
                           +GHSERFE#+GPFEHCP#
            -3957.199+5.24951*T+4.9251E+30*T**(-9); 6000 N REF0 !
PARAM G(HCP A3, CR:C;0) 298.15 +GHSERCR#+.5*GHSERCC#
            -18504+9.4173*T-2.4997*T*LN(T)+.001386*T**2; 6000 N REF1 !
PARAM G(HCP A3, FE:C; 0) 298.15 +GFEFCC#+.5*GHSERCC#+GPCFCC#
                           +52905-11.9075*T; 6000 N REF3 !
PARAM L(HCP A3, CR:C, VA; 0) 298.15 +4165; 6000 N REF1 !
PARAM L(HCP A3, FE:C, VA; 0) 298.15 -22126; 6000 N REF3 !
$PARAM L(HCP A3, FE:C, VA; 0) 298.15 -17335; 6000 N TCFE3 !
PARAM L(HCP A3, CR, FE:VA; 0) 298.15 +10833-7.477*T; 6000 N REF4 !
PHASE SIGMA % 3 8 4 18 !
CONST SIGMA : FE
           : CR
           : CR,FE : !
PARAM G(SIGMA, FE:CR:CR;0) 298.15 +8*GFEFCC#+22*GHSERCR#
                          +92300-95.96*T+GPSIG1#; 6000 N REF4 !
PARAM G(SIGMA, FE:CR:FE;0) 298.15 +8*GFEFCC#+4*GHSERCR#+18*GHSERFE#
                          +117300-95.96*T+GPSIG2#; 6000 N REF4 !
PHASE CEMENTITE % 2 3 1 !
CONST CEMENTITE : CR, FE%
              : C : !
PARAM G(CEMENTITE, CR:C;0) 298.15 +3*GHSERCR#+GHSERCC#
```

```
-48000-9.2888*T; 6000 N REF2 !
PARAM G(CEMENTITE, FE:C;0) 298.15 +GFECEM#; 6000 N REF3 !
PARAM L(CEMENTITE, CR, FE:C; 0) 298.15 +25278-17.5*T; 6000 N REF2 !
PHASE M3C2 % 2 3 2 !
CONST M3C2 : CR
         : C : !
PARAM G(M3C2,CR:C;0) 298.15 +GCRM3C2#; 6000 N REF2 !
PHASE M7C3 % 2 7 3 !
CONST M7C3 : CR%,FE
         : C : !
PARAM G(M7C3,CR:C;0) 298.15 +GCRM7C3#;
                                     6000 N REF2 !
PARAM G(M7C3, FE:C;0) 298.15 +7*GHSERFE#+3*GHSERCC#
                        +75000-48.2168*T; 6000 N REF2 !
PARAM L(M7C3,CR,FE:C;0) 298.15 -4520-10*T; 6000 N REF2 !
PHASE M23C6 % 3 20 3 6 !
CONST M23C6 : CR%, FE%
          : CR%, FE%
          : C : !
PARAM G(M23C6,CR:CR:C;0) 298.15 +GCRM23C6#; 6000 N REF5 !
PARAM G(M23C6,FE:CR:C;0) 298.15 +.1304348*GCRM23C6#
                          +.8695652*GFEM23C6#; 6000 N REF5 !
PARAM G(M23C6,CR:FE:C;0) 298.15 +.8695652*GCRM23C6#
                          +.1304348*GFEM23C6#; 6000 N REF5 !
PARAM G(M23C6, FE:FE:C;0) 298.15 +GFEM23C6#;
                                         6000 N REF5 !
PARAM L(M23C6,CR,FE:CR:C;0) 298.15 -205342+141.6667*T; 6000 N REF2 !
```

```
PARAM L(M23C6,CR,FE:FE:C;0) 298.15 -205342+141.6667*T; 6000 N REF2 !
PHASE GRAPHITE % 1 1.0 !
CONST GRAPHITE : C : !
PARAM G(GRAPHITE,C;0) 298.15 +GHSERCC#+GPCGRA#; 6000 N REF0 !
$
$ ASSESSED SYSTEMS information:
$
 * necessary for the BIN and TERN modules in TCC
Ś
               the Binary Phase Diagram module in TCW
Ś
             and the Equilibrium calculation routine in TCW
$-----
ASSESSED_SYSTEM
$... Binary ...
 C-FE(;P3 TMM:300/4000 *)
 C-CR(;P3 TMM:300/4000 *)
 CR-FE(;G5 MAJ:BCC/FE:VA C S:BCC/CR:VA
     ;P3 TMM:300/3000 STP:.6/1200/1/-2/2)
$ CR-FE(TDB -HCP ;G5 MAJ:BCC/FE:VA C S:BCC/CR:VA
    ;P3 TMM:300/3000 STP:.6/1200/1/-2/2)
$
Ś
$... Ternary ...
 C-CR-FE(;G5 MAJ:BCC/FE:VA C S:BCC/CR:VA
      ;P3 STP:.4/.5/1 STP:0.1/0.5/1)
!
Ś
$
$ LIST OF REFERENCE:
Ś
$-----
LIST OF REFERENCES
NUMBER SOURCE
```

```
REF0 'Alan Dinsdale, SGTE Data for Pure Elements, Calphad Vol 15(1991)
        p 317-425, also in NPL Report DMA(A)195 Rev. August 1990'
REF1 'J-O Andersson, Calphad Vol 11 (1987) p 271-276, TRITA 0314; C-CR'
REF2 'Byeong-Joo Lee, unpublished revision (1991); C-Cr-Fe-Ni'
REF3 'P. Gustafson, Scan. J. Metall. vol 14, (1985) p 259-267
        TRITA 0237 (1984); C-FE'
REF4 'J-O Andersson, B. Sundman, CALPHAD Vol 11, (1987), p 83-92
        TRITA 0270 (1986); CR-FE'
REF5 'J-O Andersson, Met. Trans A, Vol 19A, (1988) p 627-636
        TRITA 0207 (1986); C-CR-FE'
REFS 'Pingfang Shi (2006), TCS PTERN Public Ternary Alloys Database,
        v1.2; Modified L0(BCC,Fe,C) and L0(BCC,Cr,C) parameters
        at high temperatures.'
```

```
!
```

# The Database Checker

### In this section:

About the Database Checker	69
Database Checker Revision History	69

# **About the Database Checker**

The Thermo-Calc software package includes a program to check that the syntax of unencrypted database files is correct. The program applies the syntax rules set out in the *Database Manager User Guide* and reports errors and issues warnings. This program is intended for advanced users who develop and manage databases.



Thermo-Calc accepts deviations from these syntax rules. This means that a database can work even if the Database Checker reports errors and warnings. For example, an error is reported if an abbreviated phase name is found, but phase name abbreviations are accepted by Thermo-Calc and its add-on modules.

The executable **DatabaseChecker** file is found in the Thermo-Calc home directory. The program can also be launched by selecting **Tools Database Checker** from the main menu.



<u>"Database Checker Revision History" below</u> for releases notes that may assist database developers.

# **Database Checker Revision History**

The DatabaseChecker application is standalone and need not be installed separately, just download the containing zip file, open it and follow the instructions in the README.TXT file. The program requires that Java is installed on the target machine. Please note that large amounts of RAM may be required when checking large databases.

### **RELEASE NOTES VERSION 1.4**

- The DatabaseChecker user interface has been modified to allow checking of several database files in one batch. Instead of selecting an individual database file, the user now selects a directory that contains databases. The application then shows a tree structure with all database files in the chosen directory and its subdirectories. The user may then select any number of these files for processing.
- A check of parameters whose interaction order is 3 has been introduced. For such parameters, there must either exist one parameter with the interaction degree 0 or three parameters with the interaction degrees 0, 1 and 2.
- To facilitate optimization of databases, the tool now generates information messages about functions and parameters that will always return 0.

### RELEASE NOTES, VERSION 1.3.2

• The tool now detects conflicts between identical parameters even if one of them have been defined with the G prefix and the other with the L prefix.

### **RELEASE NOTES, VERSION 1.3**

A bug prevented integers bigger than 32 bits from being parsed. This has been corrected.

The tool now checks that the species and number of sublattices used in parameter definitions match the phase definition and generates an error message for every mismatch. Ionic liquid parameters are currently exempted from this check as not to break the widespread use of defining only one constituent for ionic liquid parameters.

Type definitions and case statements are now parsed for errors. However, only a subset of all possible GES, POLY and TDB commands are accepted and all other commands will result in an error message. The accepted commands are:

- GES AMEND\_PHASE\_DESCRIPTION
- COMPOSITION\_SETS
- MAJOR\_CONSTITUENTS
- DISORDERED\_PART
- MAGNETIC\_ORDERING
- STATUS\_BITS
- DEFAULT\_STABLE
- FRACTION\_LIMITS
- EXCESS\_MODEL
- REDLICH-KISTER\_MUGGIANU
- HKF
- HKF\_ELECTROSTATIC
- DEBYE\_HUCKEL
- GES CHANGE\_STATUS
- TDB DEFINE\_SYSTEM\_ELEMENTS
- TDB RESTORE\_PHASES
- TDB REJECT\_PHASES

Some improvements to warning and error messages have been made to facilitate finding errors.

#### **RELEASE NOTES, VERSION 1.2**

- The tool now uses parallel processing of database files on multi-core systems. This significantly improves performance when large databases are checked.
- It is no longer required that inter-dependent database statements are defined in a specific order in the database file (e.g., species need not any longer be defined before the elements on which they depend). However, for the sake of readability, it is still advised that the database statements occur in a natural order in the file.
- Because of the parallel processing of database statements, multiple definition errors (cases where one name has been used to define more than one entity in the database e.g, two identically named species) may now be generated for any of the violating occurences and not only for the last occurence in the file.

### **RELEASE NOTES, VERSION 1.1**

- Phase names were not properly recognized in parameter definitions if there was one or more whitespace characters between the opening parenthesis and the phase name. This has been corrected.
- The error message Could not parse parameter, <species name> has been given without a sublattice number but is defined in more than one sublattice is now displayed only for phases where the general diffusion model is used. Note: this implementation is temporary and may be subject to changes later.
- The meaning of the information message The phase <phase name> expects the following parameters: [<parameter names>] was unclear to users. It is now longer shown for the parameter names T and P. For all other names, the message is now displayed as an error message with the following text: The phase <phase name> contains references to the following undefined functions: [<function names>].
- The tool now looks for function names that are abbreviations of other function names and warns about this, since abbreviated names may cause problems during the execution of macro files that have been auto-generated from GES.
- The tools also warns for functions whose names are in the format V<integer>, since this may cause conflicts with names that are used by PARROT.
- Parameter names are now shown in a format that is less cryptic and more resembles the format used in the database. However, the format of diffusion parameters may still differ somewhat from what is used in the database.

### **RELEASE NOTES, VERSION 1.0**

• The tool parses a TDB file and produces a log with error, warning and information messages about the state of the file

- The message log can be exported and saved as a separate file for later analysis
- The tool is based on the database command definitions given in TC\_Database\_ ManagersGuide and DICTRA27\_UsersGuide. All commands in these documents are supported. Some undocumented DIFFUSION commands are not supported.
- Databases that span several files are correctly handled, also blocked files
- The tool is in some aspects more strict than required by the guides (see Additional Information below)
- The tool performs the following checks:
- Syntax control of database statements: ERROR
- Line size: ERROR or WARNING depending on context
- Multiply defined elements, species or phases: ERROR
- Multiply defined functions and parameters: ERROR
- Multiply defined assessed system: ERROR
- Multiply defined references: ERROR
- References entries that are never used: WARNING
- References to non-existent type definitions: ERROR
- Type definitions that are never used: WARNING
- Unused functions and parameters: WARNING
- Functions and parameters with undefined references: WARNING
- References to undefined elements, species or phases: ERROR
- Dependencies to external names from a phase: INFORMATION. (All parameters of a phase are evaluated with respect to their dependencies to other parameters and functions and all unresolved names are listed. Typically, this should only yield dependencies on T and possibly P)

#### **ADDITIONAL INFORMATION**

In addition to the command definitions given in the user guides listed above, the following rules are also applied:

- Names are case-sensitive, meaning that e.g., the defined element MN cannot be referred to under the name Mn;
- Phase names cannot be abbreviated. When referred to in parameter descriptions their full names must be used.