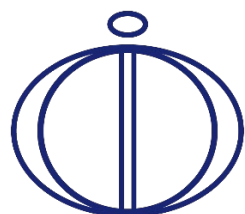


Diffusion Module (DICTRA) Example Macros

Thermo-Calc Version 2021b



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[exa1](#)

One-phase problem. Homogenization of a binary Fe-Ni alloy. In this example it is assumed there is initially a linear Ni-concentration profile.

[exa2a](#)

One-phase problem. Homogenization of a binary Fe-Ni alloy. A Ni rich and a Ni lean alloy are put together and initially there is a step profile.

[exa2b](#)

One-phase problem. Homogenization of a binary Fe-Ni alloy. This example is identical to a2a but instead it uses implicit time integration instead of the trapezoidal method for solving the PDEs.

[exa3](#)

One-phase problem. Uphill diffusion in an Fe-Si-C alloy. This is an example to simulate uphill diffusion in a ternary single phase austenite matrix due to the classical Darken experiment published by L.S. Darken: Trans. Aime, v.180 (1949), pp. 430-438.

[exa4](#)

One-phase problem. Carburization of binary Fe-C alloy: Comparison to an analytical erf solution. This is a simple binary simulation with a single phase region. The numerical simulation is compared with an analytical erf solution. For this reason a special database erf.tdb is created where the diffusion coefficient is set to a concentration independent value.

[exa5](#)

One-phase problem. Carburization of a binary Fe-0.15 wt% C alloy. A mixture of 40% N₂ and 60% cracked methanol is used as carrier gas. The carburizing "carbon potential" in the gas is 0.85 wt%. A surface reaction controls the flux of C at the surface.

[exa6](#)

One-phase problem. Diffusion through a tube wall. A simple example about diffusion through a tube wall. The tube material is an Fe-0.6Mn-0.7Si-0.05C alloy. On the inside wall a carbon activity of 0.9 is maintained whereas on the outside the C-activity is very low. This example demonstrates the use of the command SET-FIRST-INTERFACE as well as the MIXED boundary conditions.

[exa7](#)

One phase example. Homogenization heat treatment. The initial segregation profile is created from a Scheil calculation (see macro create_initial_profile.TCM). The command INPUT_SCHEIL_PROFILE in the DICTRA MONITOR performs most of the set up. Only time and temperature must be entered after the INPUT_SCHEIL_PROFILE command is executed.

[exb1a](#)

Moving boundary problem. Austenite to ferrite transformation in a binary Fe-C alloy. This example calculates a ferrite(BCC)/austenite(FCC) transformation in a binary Fe-C alloy. The initial state is an austenite of 2mm thickness. The composition of the austenite is Fe-0.15wt%C.

[exb1b](#)

Moving boundary problem. Austenite to ferrite transformation in a binary Fe-C alloy. This is the same example as in exb1a but now the problem is with ferrite as an inactive phase adjacent to the initial austenite.

[exb1c](#)

Moving boundary problem. Austenite to ferrite transformation in a binary Fe-C alloy. This is the same example as in exb1a and exb1b but now the simulation starts at a higher temperature and assumes a gradual cooling down to 1050 K.

[exb2](#)

Moving boundary problem. Cementite dissolution in an Fe-Cr-C alloy. This example calculates the dissolution of a spherical cementite particle in an austenite matrix. This case is from Z.-K. Liu, L. Håglund, B. Jähnsson and J. Ågren: Metall. Trans.A, v.22A (1991), pp. 1745-1752.

[exb3](#)

Moving boundary example. Dissolution of 23-carbide in an austenitic matrix. This example calculates the dissolution of an M23C6 particle in an austenite matrix. A film of ferrite is allowed to nucleate around the carbide during the precipitation.

[exb4a](#)

Moving boundary problem. Solidification path of an Fe-18%Cr-8%Ni alloy: Eutectic reaction. This example demonstrates the solidification path of an Fe-18%Cr-8%Ni alloy. A eutectic reaction is assumed, LIQUID -> BCC + FCC. Hence the BCC and FCC regions should be on separate sides of the liquid region. Comparison is made with both a Scheil-Gulliver simulation and equilibrium solidification conditions, both done in Thermo-Calc.

[exb4b](#)

Moving boundary problem. Solidification path of an Fe-18%Cr-8%Ni alloy: Peritectic reaction. This example is the same as exb4a but now a peritectic reaction is assumed: LIQUID + BCC -> FCC. Hence the FCC region should appear in between the LIQUID and the BCC. Comparison is made with both a Scheil-Gulliver simulation and equilibrium solidification conditions, both done in Thermo-Calc.

[exb4c](#)

Moving boundary problem. Solidification path of an Fe-18%Cr-8%Ni alloy. This example is the same as exb4b but now the diffusivity data is amended for the LIQUID and a high value for the diffusivity is used to simulate a case where it is assumed that the composition in the LIQUID is always homogeneous. This example is less realistic than exb4b. Comparison is made with both a Scheil-Gulliver simulation and equilibrium solidification conditions, both done in Thermo-Calc.

[exb4d](#)

Moving boundary problem. Solidification path of an Fe-18%Cr-8%Ni alloy. This example is the same as exb4b but instead of controlling the temperature the amount of heat extracted is given. Comparison is made with both a Scheil-Gulliver simulation and equilibrium solidification conditions, both done in Thermo-Calc.

[exb5](#)

Moving boundary problem. Ternary diffusion couple of Fe-Ni-Cr alloys. This example demonstrates the evaluation of a ternary Fe-Cr-Ni diffusion couple. A thin slice of alpha phase (38%Cr, 0%Ni) is clamped between two thicker slices of gamma phase (27%Cr, 20%Ni). The assembly is subsequently heat treated at 1373 K. This example corresponds to diffusion couple A in M. Kajihara, C.-B. Lim and M. Kikuchi: ISIJ International 33 (1993), pp. 498-507. See also M. Kajihara and M. Kikuchi: Acta Metall.Mater. 41 (1993), pp.2045-2059.

[exb6](#)

Moving boundary problem. Microsegregation of phosphorus. This example illustrates the effect of microsegregation of phosphorus during peritectic solidification in steel.

[exb7](#)

This example modifies the database interactively, which is not yet supported by GES6. Therefore, we enforce the use of GES5.

exc1

Cell calculation. Carbon cannon in ferrite/austenite: Fe-C system, 2-cell calculation This example simulates what happens to a ferrite plate that has inherited the carbon content of its parent austenite. The ferrite plate formed is embedded in an austenite matrix. This setup corresponds to a proposed mechanism for formation of Widmannst tten ferrite or for the ferrite phase of the bainite structure. It is assumed that the phase boundary between ferrite and austenite is immobile, this is achieved in the simulation by putting the ferrite and the austenite in two different cells. See also M. Hillert, L. H glund and J.  gren: Acta Metall. Mater. 41 (1993), pp.1951-1957.

exc2

Cell calculation. Cementite dissolution in an Fe-Cr-C alloy: Three particle sizes and three different cells This example calculates the dissolution of cementite particles in an austenite matrix. This example is the same as exc1 but instead there are three particle sizes. A total of six particles are considered using three different cells. This is to represent some size distribution among the cementite particles. See also Z.-K. Liu, L. H glund, B. J nsson and J.  gren: Metall.Trans.A, v. 22A (1991), pp. 1745-1752.

exd1a

Diffusion in dispersed systems. Carburization of Ni-25Cr alloy: Dispersed system model This example is about carburization of a Ni-25Cr alloy. In this case the M3C2 and M7C3 carbides are entered as spheroid phases in a FCC matrix. This simulation can be run with either the DISPERSED SYSTEM MODEL or the HOMOGENIZATION MODEL. In this example the DISPERSED SYSTEM MODEL is used, which requires that the default HOMOGENIZATION MODEL is disabled. With the DISPERSED SYSTEM MODEL the command ENTER_LABYRINTH_FUNCTION is used to take into account the impeding effect of dispersed phases on long-range diffusion. For the HOMOGENIZATION MODEL the command ENTER_HOMOGENIZATION_FUNCTION should be used. This case is from A. Engstr m, L. H glund and J.  gren: Metall.Trans.A v. 25A (1994), pp. 1127-1134.

exd1b

Diffusion in dispersed systems. Carburization of Ni-25Cr alloy: Homogenization model This example is about carburization of a Ni-25Cr alloy. In this case the M3C2 and M7C3 carbides are entered as spheroid phases in a FCC matrix. This case is from A. Engstr m, L. H glund and J.  gren: Metall.Trans. A, v.25A (1994), pp. 1127-1134. This simulation can be run with the DISPERSED SYSTEM MODEL or HOMOGENIZATION MODEL. The default HOMOGENIZATION MODEL is used and then ENTER_HOMOGENIZATION_FUNCTION should be used instead of ENTER_LABYRINTH_FUNCTION.

exd2a

Diffusion in dispersed systems. Diffusion couple of Fe-Cr-Ni alloys: Dispersed system model This example calculates the interdiffusion in a diffusion couple between a two-phase (FCC+BCC) and a single-phase (FCC) Fe-Ni-Cr alloy. This case is from A. Engstr m: Scand. J. Met., v. 24, 1995, pp.12-20. This simulation can be run with either the DISPERSED SYSTEM MODEL or the HOMOGENIZATION MODEL. In this example the DISPERSED SYSTEM MODEL is used, which requires that the default HOMOGENIZATION MODEL is disabled. With the DISPERSED SYSTEM MODEL the command ENTER_LABYRINTH_FUNCTION is used to take into account the impeding effect of dispersed phases on long-range diffusion. For the HOMOGENIZATION MODEL the command ENTER_HOMOGENIZATION_FUNCTION should be used.

exd2b

Diffusion in dispersed systems. Diffusion couple of Fe-Cr-Ni alloys: Homogenization model This example calculates the interdiffusion in a diffusion couple between a two-phase (FCC+BCC) and a single-phase (FCC) Fe-Ni-Cr alloy. This case is from A. Engstr m: Scand. J. Met., v. 24, 1995, pp.12-20. This simulation can be run with either the DISPERSED SYSTEM MODEL or the HOMOGENIZATION MODEL. Here the default HOMOGENIZATION MODEL is used and then ENTER_HOMOGENIZATION_FUNCTION should be used instead of ENTER_LABYRINTH_FUNCTION.

exd3

Diffusion in dispersed systems. Diffusion couple of Fe-Cr-Ni alloys: Homogenization model This example uses the homogenization model. It is taken from H. Larsson and A. Engstr m, Acta. Mater. v.54 (2006), pp. 2431-2439. Experimental data from A. Engstr m, Scand J Metall, v.243 (1995), p.12. The homogenization model can be used for multiphase simulations like the dispersed system model, but unlike the dispersed system model there is no need to have a single continuous matrix phase and, furthermore, there is no need to limit the size of time-steps. The set-up is performed in the same manner as for the dispersed system model, which means that a certain phase is entered as the matrix phase and the other phases are entered as spheroidal, but the choice of matrix phase will not affect the simulation.

exe1

Cooperative growth. Growth of pearlite in an Fe-Mn-C alloy An example of pearlite growth in an Fe-0.50wt%C-0.91wt%Mn steel.

exf1

Coarsening problem. Coarsening of M6C precipitate in an Fe-Mo-C alloy This example calculates the Ostwald-ripening of a spherical M6C carbide in an austenite matrix.

exg1

Kinetic data example. Checking mobilities and diffusivities in an Fe-Ni alloy This is an example file to check the mobilities and diffusivities in an Fe-Ni alloy.

exg2

Kinetic data example. Optimization of mobilities in Ni-Al fcc alloys A file for reading thermodynamic data and setting up the kinetic parameters that are needed for an optimization of the FCC phase in the binary Ni-Al system. See also A. Engstr m and J.  gren: ("Assessment of Diffusional Mobilities in Face-Centered Cubic Ni-Cr-Al Alloys" in Z. Metallkunde, Feb. 1996).

exh1

Deviation from local equilibrium. Ferrite/austenite diffusion couple with interface mobility This example calculates the growth of ferrite into austenite with a limited interface mobility. this is done by adding a Gibbs-energy contribution to the ferrite using the SET-SURFACE-ENERGY command.

exh2

Deviation from local equilibrium. Ferrite/austenite para-equilibrium in an Fe-Ni-C alloy This example calculates the growth of ferrite into austenite in an Fe-2.02%Ni-0.0885%C alloy using the para-equilibrium model. The results are compared with experimental information from Hutchinson, C. R., A. Fuchsmann, and Yves Brechet. "The diffusional formation of ferrite from austenite in Fe-C-Ni alloys." Metall. Mat. Trans. A 35.4 (2004): 1211-1221.

exh3

Deviation from local equilibrium. Diffusion induced by a temperature gradient (thermomigration) This calculation shows how a temperature gradient induces diffusion.

exi1

Diffusion in complex phases. Diffusion in a system with B2 ordering This example shows diffusion in a system with B2 ordering. The datafile AlFeNi-data.TDB contains both a thermodynamic and kinetic description for the ordered and disordered BCC.

exi2

Diffusion in complex phases. Diffusion of carbon in cementite This example demonstrates the use of the model for calculation of diffusion through a stoichiometric phase. The flux of a component in the stoichiometric phase is assumed to be proportional to the difference in chemical potential at each side of the stoichiometric phase multiplied with the mobility for the component in the phase. The mobility is assessed from experimental information and is basically the tracer diffusivity for the component.

exi3a

Diffusion in complex phases. Diffusion in iron oxide (FeO) This example shows the oxidation of an iron sample and the consequent growth of an oxide layer.

exi3b Diffusion in complex phases. Diffusion in iron oxide (FeO) with a grain boundary contribution This example shows the oxidation of an iron sample and consequent growth of an oxide layer using the grain boundary diffusion contribution model.



One-Phase Problems



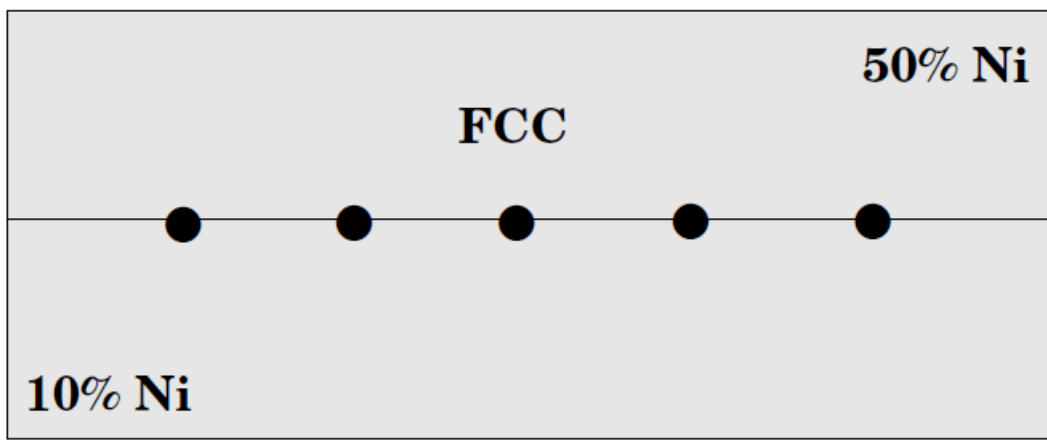


Example exa1

Homogenization of a binary Fe-Ni alloy: Linear Concentration Profile

Simple homogenization of a binary Fe-Ni alloy. It is assumed there is initially a linear Ni-concentration profile.

T = 1400 K



Results

exal-setup

SYS:About

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

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Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exal\setup.DCM.test"
SYS: @@
SYS: @@ One-phase problem.
SYS: @@ Homogenization of a binary Fe-Ni alloy.
SYS: @@ In this example it is assumed there is initially a linear
SYS: @@ Ni-concentration profile.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: goto_module
MODULE NAME: data
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA          /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ USE THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: @@
TDB_TCFE11: switch_database
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2
TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
```

```

SNOB1      = SGTE Noble Metal Alloys Database v1.2
STBC2      = SGTE Thermal Barrier Coating TDB v2.2
STBC1      = SGTE Thermal Barrier Coating TDB v1.1
SALT1      = SGTE Molten Salts Database v1.3
SNUX6      = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2      = TC Semi-Conductors v2.1
SLAG4      = Fe-containing Slag v4.1
SLAG3      = Fe-containing Slag v3.2
SLAG2      = Fe-containing Slag v2.2
SLAG1      = Fe-containing Slag v1.2
TCOX11     = Metal Oxide Solutions v11.0
TCOX10     = Metal Oxide Solutions v10.1
TCOX9      = Metal Oxide Solutions v9.0
TCOX8      = Metal Oxide Solutions v8.0
TCOX7      = Metal Oxide Solutions v7.0
TCOX6      = Metal Oxide Solutions v6.0
TCOX5      = Metal Oxide Solutions v5.1
TCOX4      = Metal Oxide Solutions v4.1
ION3       = Ionic Solutions v3.0
ION2       = Ionic Solutions v2.6
ION1       = Ionic Solutions v1.5
NOX2       = NPL Oxide Solutions Database v2.1
TCNOBL1    = Noble Metals Alloys v1.0
TCNOBL2    = Noble Metals Alloys v2.0
TCSLD4     = Solder Alloys v4.0
TCSLD3     = Solder Alloys v3.3
TCSLD2     = Solder Alloys v2.0
TCSLD1     = Solder Alloys v1.1
TCSI1      = Ultrapure Silicon v1.2
TCMP2      = Materials Processing v2.5
TCES1      = Combustion/Sintering v1.1
TSCS1      = Super Conductor v1.0
TCFC1      = SOFC Database v1.0
TCNF2      = Nuclear Fuels v2.1b
NUMT2      = Nuclear Materials v2.1
NUOX4      = Nuclear Oxides v4.2
NUTO1      = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1      = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19     = IRSN NUCLEA-19
NUCL15     = IRSN NUCLEA-15_4
NUCL10     = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19     = IRSN Mephista-19
MEPH15     = IRSN Mephista-15_1
MEPH11     = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3      = Aqueous Solution v3.0
TCAQ2      = Aqueous Solution v2.7
AQ52       = TGG Aqueous Solution Database v2.6
GCE2       = TGG Geochemical/Environmental TDB v2.3
FEDEMO    = Iron Demo Database v4.0
ALDEMO    = Aluminum Demo Database v4.0
NIDEMO    = Nickel Demo Database v2.0
CUDEMO    = Copper Demo Database v1.0
SLDEMO    = Solder Demo Database v1.0
OXDEMO    = Oxide Demo Database v3.0
SUBDEMO   = Substance Demo Database v1.0
PAQ2       = Public Aqueous Soln (SIT) TDB v2.5
PG35       = PG35 Binary Semi-Conductors TDB v1.3
PURE5     = SGTE Unary (Pure Elements) TDB v5.2
MOB2       = Alloys Mobility v2.7
MOB1       = Alloys Mobility v1.3
MOBFE1     = Steels/Fe-Alloys Mobility v1.1
MOBFE2     = Steels/Fe-Alloys Mobility v2.0
MOBFE3     = Steels/Fe-Alloys Mobility v3.0
MOBFE4     = Steels/Fe-Alloys Mobility v4.0
MOBFE5     = Steels/Fe-Alloys Mobility v5.0
MOBFE6     = Steels/Fe-Alloys Mobility v6.0
MOBN15     = Ni-Alloys Mobility v5.1
MOBN14     = Ni-Alloys Mobility v4.1
MOBN13     = Ni-Alloys Mobility v3.2
MOBN12     = Ni-Alloys Mobility v2.4
MOBN11     = Ni-Alloys Mobility v1.10
MOBAL6     = Al-Alloys Mobility v6.0
MOBAL5     = Al-Alloys Mobility v5.0
MOBAL4     = Al-Alloys Mobility v4.0
MOBAL3     = Al-Alloys Mobility v3.0
MOBAL2     = Al-Alloys Mobility v2.0
MOBAL1     = Al-Alloys Mobility v1.0
MOBCU1     = Cu-Alloys Mobility v1.0
MOBCU2     = Cu-Alloys Mobility v2.0
MOBCU3     = Cu-Alloys Mobility v3.0
MOBCU4     = Cu-Alloys Mobility v4.0
MOBHEA1    = High Entropy Alloys Mobility v1.0
MOBHEA2    = High Entropy Alloys Mobility v2.0
MOBMG2     = Mg-Alloys Mobility v2.0
MOBMG1     = Mg-Alloys Mobility v1.0
MOBSI1     = Si-Alloys Mobility v1.0
MOBSLD1    = Solder-Alloys Mobility v1.1
MOBT14     = Ti-Alloys Mobility v4.0
MOBT13     = Ti-Alloys Mobility v3.1
MOBT12     = Ti-Alloys Mobility v2.0
MOBT11     = Ti-Alloys Mobility v1.0
MALDEMO   = Al-Alloys Mobility demo database v2.0
MFEDEMO   = Fe-Alloys Mobility demo database v2.0
MNIDEMO   = Ni-Alloys Mobility demo database v1.0
MCUDEMO   = Cu-Alloys Mobility demo database v1.0
USER      = User defined Database

```

```

DATABASE NAME /TCFE11/: fedemo
Current database: Iron Demo Database v4.0

VA                      /- DEFINED
TDB_FEDEMO: @@
TDB_FEDEMO: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_FEDEMO: @@
TDE_FEDEMO: define_system
ELEMENTS: fe ni           NI  DEFINED
TDB_FEDEMO: @@
TDB_FEDEMO: @@
TDB_FEDEMO: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDE_FEDEMO: @@
TDB_FEDEMO: reject
ELEMENTS, SPECIES, PHASES, CONSTITUENT OR SYSTEM: /PHASES/: phase
PHASES: *

```

```

LIQUID:L           BCC_A2          LAVES_PHASE_C14
CBCC_A12         CUB_A13        FCC_A1
HCP_A3   REJECTED

TDB_FEDEMO: @@
TDB_FEDEMO: @@ RESTORE THE THERMODYNAMIC DATA FOR THE FCC PHASE
TDB_FEDEMO: @@
TDB_FEDEMO: restore
ELEMENTS, SPECIES, PHASES OR CONSTITUENTS: /ELEMENTS/: phase
PHASES: fcc
FCC_A1 RESTORED
TDB_FEDEMO: @@
TDB_FEDEMO: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_FEDEMO: @@
TDB_FEDEMO: get_data
09:20:45,366 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
    volumes'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
-OK-
TDB_FEDEMO: @@
TDB_FEDEMO: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_FEDEMO: @@ SWITCH TO THE MOBILITY DATABASE TO RETRIEVE DATA
TDB_FEDEMO: @@
TDB_FEDEMO: append_database
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2
TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
SNOB1 = SGTE Noble Metal Alloys Database v1.2
STBC2 = SGTE Thermal Barrier Coating TDB v2.2
STBC1 = SGTE Thermal Barrier Coating TDB v1.1
SALT1 = SGTE Molten Salts Database v1.3
SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2 = TC Semi-Conductors v2.1
SLAG4 = Fe-containing Slag v4.1
SLAG3 = Fe-containing Slag v3.2
SLAG2 = Fe-containing Slag v2.2
SLAG1 = Fe-containing Slag v1.2
TCOX11 = Metal Oxide Solutions v11.0
TCOX10 = Metal Oxide Solutions v10.1
TCOX9 = Metal Oxide Solutions v9.0
TCOX8 = Metal Oxide Solutions v8.0
TCOX7 = Metal Oxide Solutions v7.0

```

```

TCOX6 = Metal Oxide Solutions v6.0
TCOX5 = Metal Oxide Solutions v5.1
TCOX4 = Metal Oxide Solutions v4.1
ION3 = Ionic Solutions v3.0
ION2 = Ionic Solutions v2.6
ION1 = Ionic Solutions v1.5
NOX2 = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4 = Solder Alloys v4.0
TCSLD3 = Solder Alloys v3.3
TCSLD2 = Solder Alloys v2.0
TCSLD1 = Solder Alloys v1.1
TCSI1 = Ultrapure Silicon v1.2
TCMP2 = Materials Processing v2.5
TCE51 = Combustion/Sintering v1.1
TCS51 = Super Conductor v1.0
TCFC1 = SOFC Database v1.0
TCNF2 = Nuclear Fuels v2.1b
NUMT2 = Nuclear Materials v2.1
NUOX4 = Nuclear Oxides v4.2
NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephista-19
MEPH15 = IRSN Mephista-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQ52 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SLDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.2
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3
MOBFE1 = Steels/Fe-Alloys Mobility v1.1
MOBFE2 = Steels/Fe-Alloys Mobility v2.0
MOBFE3 = Steels/Fe-Alloys Mobility v3.0
MOBFE4 = Steels/Fe-Alloys Mobility v4.0
MOBFE5 = Steels/Fe-Alloys Mobility v5.0
MOBFE6 = Steels/Fe-Alloys Mobility v6.0
MOBN15 = Ni-Alloys Mobility v5.1
MOBN14 = Ni-Alloys Mobility v4.1
MOBN13 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBS11 = Si-Alloys Mobility v1.0
MOBSD1 = Solder-Alloys Mobility v1.1
MOBT14 = Ti-Alloys Mobility v4.0
MOBT13 = Ti-Alloys Mobility v3.1
MOBT12 = Ti-Alloys Mobility v2.0
MOBT11 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

```

```

DATABASE NAME /FEDEMO/: mfedemo
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: define_system
ELEMENTS: fe ni
    FE           NI   DEFINED
APP: reject
ELEMENTS, SPECIES, PHASES, CONSTITUENT OR SYSTEM: /PHASES/: phase
PHASES: *
    BCC_A2          FCC_A1  REJECTED
APP:
APP: restore
ELEMENTS, SPECIES, PHASES OR CONSTITUENTS: /ELEMENTS/: phase
PHASES: fcc
    FCC_A1  RESTORED
APP:
APP: get_data
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
-OK-
APP:
APP: @@
```

```

APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: goto_module
MODULE NAME: dictra_monitor
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set_condition
GLOBAL OR BOUNDARY CONDITION /GLOBAL/: global
VARIABLE : T
LOW TIME LIMIT /0/: 0
T (TIME,X)= 1400;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
DIC>
DIC> @@
DIC> @@ START BY ENTERING A REGION
DIC> @@
DIC> enter_region
REGION NAME : austenite
DIC>
DIC> @@
DIC> @@ ENTER A GRID INTO THE REGION
DIC> @@
DIC> @@ FOR SIMPLICITY, AN EQUIDISTANT GRID IS USED
DIC> @@
DIC> enter_grid_coordinates
REGION NAME : /AUSTENITE/: austenite
WIDTH OF REGION /1/: 1e-4
TYPE /LINEAR/: AUTO
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE active PHASE INTO THE REGION
DIC> @@
DIC> enter_phase_in_region
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /AUSTENITE/: austenite
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC> @@
DIC> @@ ENTER AN INITIAL Ni COMPOSITION INTO THE PHASE. A LINEAR
DIC> @@ VARIATION IN THE REGION IS ASSUMED.
DIC> @@
DIC> enter_compositions
REGION NAME : /AUSTENITE/: austenite
PHASE NAME: /FCC_A1/: fcc#1
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: weight_percent
PROFILE FOR /NI/: ni
TYPE /LINEAR/: linear
VALUE OF FIRST POINT : 10
VALUE OF LAST POINT : /10/: 50
DIC>
DIC> @@
DIC> @@ THE BOUNDARY CONDITION IS A CLOSED SYSTEM AS WE DO NOT SPECIFY
DIC> @@ ANYTHING ELSE
DIC> @@
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set_simulation_time
END TIME FOR INTEGRATION /.1/: 1E6
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /100000/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO FILE AND EXIT
DIC> @@
DIC> save_workspaces exa1 Y
DIC>
DIC> set_interactive
--OK--
DIC>

```

exa1-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa1\run.DCM.test"
DIC>
DIC>
DIC> @@ exa1_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE a1
DIC> @@
DIC> goto module
MODULE NAME: dictra_monitor
TIME STEP AT TIME 0.00000E+00
DIC>
DIC> read_workspaces exa1
OK
DIC>
DIC> @@
DIC> @@ Start the simulation
DIC> @@
DIC> simulate_reaction
Region: AUSTENITE
linear grid 75
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.1001000E-03 DT = 0.1000000E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.40010010 DT = 0.40000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 241.69569 DT = 241.29559 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 724.28686 DT = 482.59117 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 1689.4692 DT = 965.18235 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 3619.8339 DT = 1930.3647 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 7480.5633 DT = 3860.7294 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 15202.022 DT = 7721.4588 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 30644.940 DT = 15442.918 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 61530.775 DT = 30885.835 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 123302.44 DT = 61771.670 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399617
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 223302.44 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 323302.44 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 423302.44 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600385 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 523302.44 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 623302.44 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600385 NI = .290319863399615
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 723302.44 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 823302.44 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600384 NI = .290319863399616
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 923302.44 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600385 NI = .290319863399615
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 1000000.0 DT = 76697.555 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .709680136600385 NI = .290319863399615
TOTAL SIZE OF SYSTEM: 1E-04 [m]
MUST SAVE WORKSPACE ON FILE
```

WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 0.10010000E-03
DELETING TIME-RECORD FOR TIME 0.40010010
DELETING TIME-RECORD FOR TIME 241.69569
DELETING TIME-RECORD FOR TIME 724.28686
DELETING TIME-RECORD FOR TIME 1689.4692
DELETING TIME-RECORD FOR TIME 3619.8339
DELETING TIME-RECORD FOR TIME 7480.5633
DELETING TIME-RECORD FOR TIME 15202.022
DELETING TIME-RECORD FOR TIME 30644.940
DELETING TIME-RECORD FOR TIME 61530.775
DELETING TIME-RECORD FOR TIME 123302.44
DELETING TIME-RECORD FOR TIME 223302.44
DELETING TIME-RECORD FOR TIME 323302.44
DELETING TIME-RECORD FOR TIME 423302.44
DELETING TIME-RECORD FOR TIME 523302.44
DELETING TIME-RECORD FOR TIME 623302.44
DELETING TIME-RECORD FOR TIME 723302.44
DELETING TIME-RECORD FOR TIME 823302.44

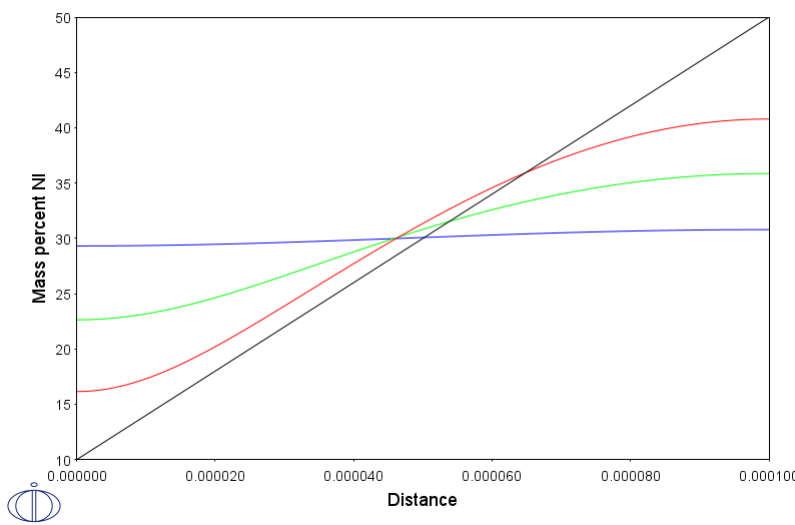
KEEPING TIME-RECORD FOR TIME 923302.44
AND FOR TIME 1000000.0
WORKSPACE RECLAIMED

TIMESTEP AT 1000000.00 SELECTED

DIC>
DIC> @@
DIC> @@ THE SIMULATION IS FINISHED
DIC> @@
DIC>
DIC> set_interactive
--OK--
DIC>

exal-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exal\plot.DCM.test"
DIC>
DIC>
DIC> @@ exal_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE a1
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> goto_module
MODULE NAME: dictra_monitor
TIME STEP AT TIME 1.00000E+06
DIC> read_workspaces exal
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post_processor
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1:
POST-1: @@
POST-1: @@ PLOT SOME CONCENTRATION PROFILES
POST-1: @@
POST-1: set_diagram_axis
AXIS (X, Y OR Z) : x
VARIABLE : distance
INFO: Distance is set as independent variable
DISTANCE : /GLOBAL/: global
POST-1:
POST-1: set_diagram_axis
AXIS (X, Y OR Z) : y
VARIABLE : weight-percent
FOR COMPONENT : ni
POST-1:
POST-1: set_plot_condition
CONDITION /TIME/: time
VALUE(S) /LAST/: 0 1e5 3e5 1e6
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot_diagram
2021.05.13.09.22.44
Time=0,100000,300000,1000000
CELL #1
```



```
POST-1:
POST-1:
POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1:
POST-1: set_interactive
--OK---
POST-1:
```

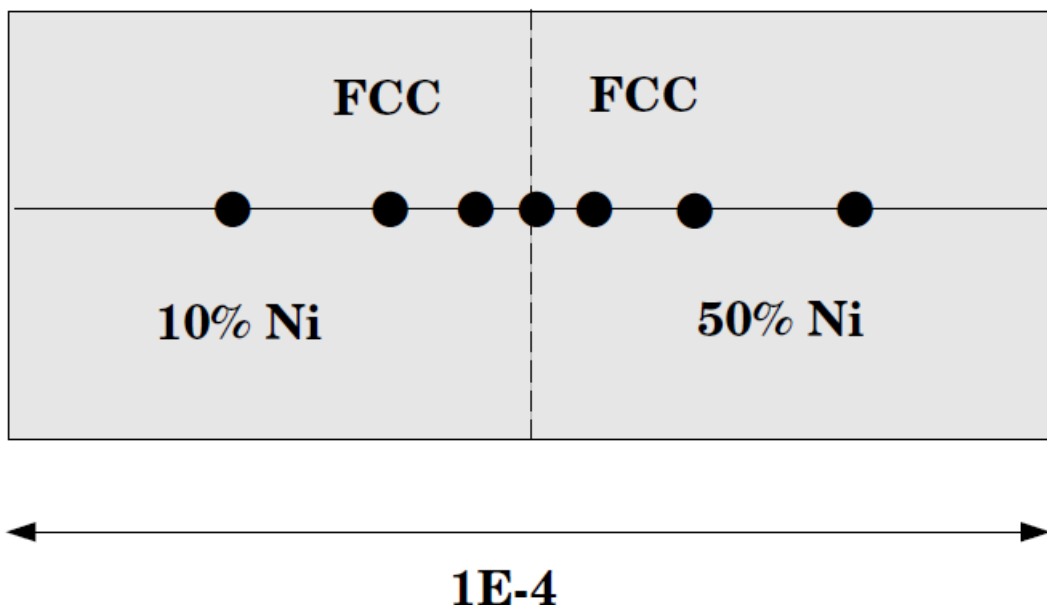


Example exa2a

Homogenization of a binary Fe-Ni alloy: Step-profile

Simple homogenization of a binary Fe-Ni alloy. A Ni rich and a Ni lean alloy are put together and initially there is a step profile.

$$T = 1400 \text{ K}$$



exa2a-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa2a\setup.DCM.test"
SYS: @@
SYS: @@ One-phase problem.
SYS: @@ Homogenization of a binary Fe-Ni alloy.
SYS: @@ A Ni rich and a Ni lean alloy are put together and initially
SYS: @@ there is a step profile.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exa2a_setup.DCM
SYS:
SYS: @@
SYS: @@ IN exa1 ALL THE COMMANDS WERE WRITTEN IN FULL BUT NOW ABBREVIATED
SYS: @@ COMMANDS ARE USED
SYS: @@
SYS:
SYS: @@
SYS: @@ FIRST DEFINE A LOG-FILE FOR THIS EXAMPLE
SYS: @@
SYS: set_log_file setup
Heading:
SYS: @@
SYS: @@ NOW GO TO THE DATABASE MODULE
SYS: @@
SYS: go da
... the command in full is GOTO_MODULE
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /- DEFINED
TDB_FCC_A1  REJECTED
TDB_TCFE11: @@
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: @@
TDB_TCFE11: sw FEDEMO
... the command in full is SWITCH_DATABASE
Current database: Iron Demo Database v4.0

VA           /- DEFINED
TDB_FEDEMO: @@
TDB_FEDEMO: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_FEDEMO: @@
TDB_FEDEMO: def-system fe ni
... the command in full is DEFINE_SYSTEM
FE          NI  DEFINED
TDB_FEDEMO: @@
TDB_FEDEMO: @@
TDB_FEDEMO: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_FEDEMO: @@
TDB_FEDEMO: rej ph * all
... the command in full is REJECT
LIQUID:L      BCC_A2          LAVES_PHASE_C14
CBCC_A12     CUB_A13         FCC_A1
HCP_A3       REJECTED
TDB_FEDEMO: res ph fcc
... the command in full is RESTORE
FCC_A1      RESTORED
TDB_FEDEMO: @@
TDB_FEDEMO: @@
TDB_FEDEMO: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_FEDEMO: @@
TDB_FEDEMO: get
... the command in full is GET_DATA
09:24:02,549 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
... the command in full is AMEND_PHASE_DESCRIPTION
PARAMETERS ...
FUNCTIONS ...

List of references for assessed data

'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
    volumes'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
-OK-
TDB_FEDEMO: @@
TDB_FEDEMO: @@
TDB_FEDEMO: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_FEDEMO: @@ SWITCH TO THE MOBILITY DATABASE TO RETRIEVE DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app
... the command in full is APPEND_DATABASE
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9  = Steels/Fe-Alloys v9.3
TCFE8  = Steels/Fe-Alloys v8.2
```

TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2
TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
SNOB1 = SGTE Noble Metal Alloys Database v1.2
STBC2 = SGTE Thermal Barrier Coating TDB v2.2
STBC1 = SGTE Thermal Barrier Coating TDB v1.1
SALT1 = SGTE Molten Salts Database v1.3
SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2 = TC Semi-Conductors v2.1
SLAG4 = Fe-containing Slag v4.1
SLAG3 = Fe-containing Slag v3.2
SLAG2 = Fe-containing Slag v2.2
SLAG1 = Fe-containing Slag v1.2
TCOX11 = Metal Oxide Solutions v11.0
TCOX10 = Metal Oxide Solutions v10.1
TCOX9 = Metal Oxide Solutions v9.0
TCOX8 = Metal Oxide Solutions v8.0
TCOX7 = Metal Oxide Solutions v7.0
TCOX6 = Metal Oxide Solutions v6.0
TCOX5 = Metal Oxide Solutions v5.1
TCOX4 = Metal Oxide Solutions v4.1
ION3 = Ionic Solutions v3.0
ION2 = Ionic Solutions v2.6
ION1 = Ionic Solutions v1.5
NOX2 = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4 = Solder Alloys v4.0
TCSLD3 = Solder Alloys v3.3
TCSLD2 = Solder Alloys v2.0
TCSLD1 = Solder Alloys v1.1
TCSI1 = Ultrapure Silicon v1.2
TCMP2 = Materials Processing v2.5
TCES1 = Combustion/Sintering v1.1
TCSR1 = Super Conductor v1.0
TFCF1 = SOFC Database v1.0
TCNF2 = Nuclear Fuels v2.1b
NUMT2 = Nuclear Materials v2.1
NUOX4 = Nuclear Oxides v4.2
NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephisto-19
MEPH15 = IRSN Mephisto-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQ52 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SLDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.2
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3

```

MOBFE1 = Steels/Fe-Alloys Mobility v1.1
MOBFE2 = Steels/Fe-Alloys Mobility v2.0
MOBFE3 = Steels/Fe-Alloys Mobility v3.0
MOBFE4 = Steels/Fe-Alloys Mobility v4.0
MOBFE5 = Steels/Fe-Alloys Mobility v5.0
MOBFE6 = Steels/Fe-Alloys Mobility v6.0
MOBN15 = Ni-Alloys Mobility v5.1
MOBNI4 = Ni-Alloys Mobility v4.1
MOBN13 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBSI1 = Si-Alloys Mobility v1.0
MOBSDL1 = Solder-Alloys Mobility v1.1
MOBT14 = Ti-Alloys Mobility v4.0
MOBT13 = Ti-Alloys Mobility v3.1
MOBT12 = Ti-Alloys Mobility v2.0
MOBT11 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

```

DATABASE NAME /FEDEMO/: mfedemo

Current database: Fe-Alloys Mobility demo database v2.0

```

VA DEFINED
APP: def-sys fe ni
... the command in full is DEFINE_SYSTEM
FE NI DEFINED
APP: rej ph * all
... the command in full is REJECT
BCC_A2 FCC_A1 REJECTED
APP: res ph fcc
... the command in full is RESTORE
FCC_A1 RESTORED
APP: get
... the command in full is GET_DATA
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

```

List of references for assessed data

```

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
-OK-
APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: go d-m
... the command in full is GOTO_MODULE
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond
... the command in full is SET_CONDITION
GLOBAL OR BOUNDARY CONDITION /GLOBAL/:
glob
VARIABLE : T
LOW TIME LIMIT /0/: 0
T(TIME,X)= 1400;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
DIC>
DIC> @@
DIC> @@ ENTER A REGION
DIC> @@
DIC> enter-region
REGION NAME : austenite
DIC>
DIC> @@
DIC> @@ ENTER A GRID INTO THE REGION.
DIC> @@ IN THIS CASE WE WANT SEVERAL POINTS IN THE MIDDLE OF THE REGION,
DIC> @@ SO A DOUBLE GEOMETRIC GRID IS CONSTRUCTED.
DIC> @@
DIC> enter-grid
... the command in full is ENTER_GRID_COORDINATES
REGION NAME : /AUSTENITE/:
austenite
WIDTH OF REGION /1/: 1e-4
TYPE /LINEAR/: double
NUMBER OF POINTS /50/: 60
VALUE OF R IN THE GEOMETRICAL SERIE FOR LOWER PART OF REGION: 0.9
VALUE OF R IN THE GEOMETRICAL SERIE FOR UPPER PART OF REGION: 1.11
DIC>
DIC> @@
DIC> @@ ENTER THE active PHASES INTO THE REGIONS
DIC> @@
DIC> enter-phase
... the command in full is ENTER_PHASE_IN_REGION
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUSTENITE/:
austenite
PHASE TYPE /MATRIX/: matrix

```


exa2a-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa2a\run.DCM.test"
... the command in full is MACRO_FILE_OPEN
DIC>
DIC>
DIC> @@ exa2a_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE a2a
DIC> @@
DIC>
DIC> @@
DIC> @@ LET US DEFINE A LOG-FILE FOR THIS EXAMPLE
DIC> @@
DIC> @@set-log-file run
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AN READ SETUP FROM FILE
DIC> @@
DIC> go d-m
... the command in full is GOTO_MODULE
TIME STEP AT TIME 0.00000E+00
DIC>
DIC> read exa2a
... the command in full is READ_WORKSPACES
... the command in full is DEFINE_COMPONENTS
... the command in full is SELECT_EQUILIBRIUM
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> simulate
... the command in full is SIMULATEREACTION
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
... the command in full is SET_NUMERICAL_LIMITS
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
2 GRIDPOINT(S) ADDED TO CELL #1 REGION: AUSTENITE
TIME = 0.10000000E-06 DT = 0.10000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.10010000E-03 DT = 0.10000000E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.40010010 DT = 0.40000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 10.408313 DT = 10.008213 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 30.424739 DT = 20.016426 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 70.457590 DT = 40.032851 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 150.52329 DT = 80.065703 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 310.65470 DT = 160.13141 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 1271.4431 DT = 640.52562 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 2552.4944 DT = 1281.0512 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 5114.5969 DT = 2562.1025 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 10238.802 DT = 5124.2050 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 20487.212 DT = 10248.410 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 40984.032 DT = 20496.820 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 81977.671 DT = 40993.640 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 163964.95 DT = 81987.280 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406168 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 263964.95 DT = 100000.00 SUM OF SQUARES = 0.0000000
```

```

U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 363964.95 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 463964.95 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .291111754593831
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 563964.95 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .291111754593831
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 663964.95 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406168 NI = .291111754593833
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 763964.95 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406167 NI = .291111754593833
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 863964.95 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406167 NI = .291111754593833
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 963964.95 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406168 NI = .291111754593832
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 1000000.0 DT = 36035.049 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406168 NI = .291111754593832
TOTAL SIZE OF SYSTEM: 1E-04 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 0.10010000E-03
DELETING TIME-RECORD FOR TIME 0.40010010
DELETING TIME-RECORD FOR TIME 10.408313
DELETING TIME-RECORD FOR TIME 30.424739
DELETING TIME-RECORD FOR TIME 70.457590
DELETING TIME-RECORD FOR TIME 150.52329
DELETING TIME-RECORD FOR TIME 310.65470
DELETING TIME-RECORD FOR TIME 630.91751
DELETING TIME-RECORD FOR TIME 1271.4431
DELETING TIME-RECORD FOR TIME 2552.4944
DELETING TIME-RECORD FOR TIME 5114.5969
DELETING TIME-RECORD FOR TIME 10238.802
DELETING TIME-RECORD FOR TIME 20487.212
DELETING TIME-RECORD FOR TIME 40984.032
DELETING TIME-RECORD FOR TIME 81977.671
DELETING TIME-RECORD FOR TIME 163964.95
DELETING TIME-RECORD FOR TIME 263964.95
DELETING TIME-RECORD FOR TIME 363964.95
DELETING TIME-RECORD FOR TIME 463964.95
DELETING TIME-RECORD FOR TIME 563964.95
DELETING TIME-RECORD FOR TIME 663964.95
DELETING TIME-RECORD FOR TIME 763964.95
DELETING TIME-RECORD FOR TIME 863964.95

KEEPING TIME-RECORD FOR TIME 963964.95
AND FOR TIME 1000000.0
WORKSPACE RECLAIMED

```

```

TIMESTEP AT 1000000.00 SELECTED

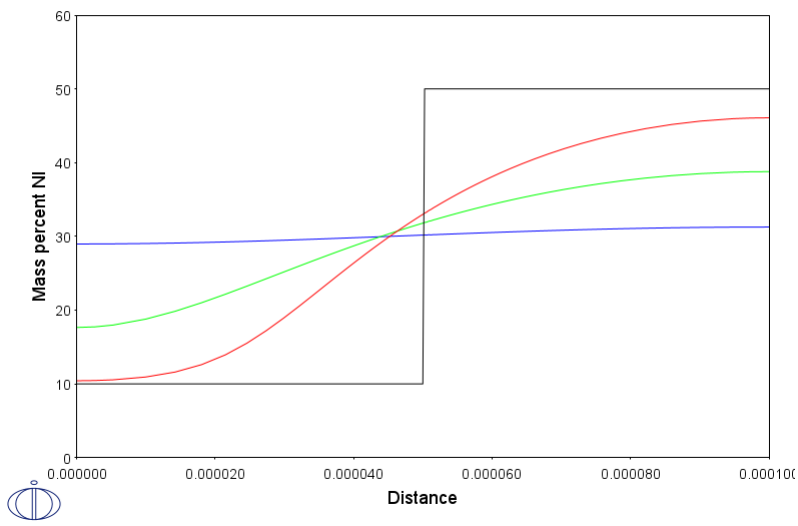
DIC>
DIC> @@
DIC> @@ THE SIMULATION IS FINISHED
DIC> @@
DIC>
DIC> set-inter
... the command in full is SET_INTERACTIVE
--OK--
DIC>

```

exa2a-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa2a\plot.DCM.test"
... the command in full is MACRO_FILE_OPEN
DIC>
DIC>
DIC> @@ exa2a_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE a2a
DIC> @@
DIC>
DIC> @@
DIC> @@ LET US DEFINE A LOG-FILE FOR THIS EXAMPLE
DIC> @@
DIC> set-log-file plot
AMBIGUOUS COMMAND, USE HELP
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
... the command in full is GOTO_MODULE
TIME STEP AT TIME 1.00000E+06
DIC>
DIC> read exa2a
... the command in full is READ_WORKSPACES
... the command in full is DEFINE_COMPONENTS
... the command in full is SELECT_EQUILIBRIUM
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
... the command in full is POST_PROCESSOR
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1:
POST-1: @@
POST-1: @@ PLOT SOME NI-CONCENTRATION PROFILES
POST-1: @@
POST-1: s-d-a
... the command in full is SET_DIAGRAM_AXIS
AXIS (X, Y OR Z) : x
VARIABLE : dist
INFO: Distance is set as independent variable
... the command in full is SET_INDEPENDENT_VARIABLE
DISTANCE : /GLOBAL/: glo
POST-1:
POST-1: s-d-a
... the command in full is SET_DIAGRAM_AXIS
AXIS (X, Y OR Z) : y
VARIABLE : weight-percent
FOR COMPONENT : ni
POST-1:
POST-1: s-p-c
... the command in full is SET_PLOT_CONDITION
CONDITION /TIME/: time
VALUE(S) /LAST/: 0 1e5 3e5 1e6
POST-1:
POST-1: @@
POST-1: @@ SET SCALING ON Y-AXIS BEFORE PLOTTING
POST-1: @@
POST-1: s-s-s
... the command in full is SET_SCALING_STATUS
AXIS (X, Y OR Z) : y
AUTOMATIC SCALING (Y OR N) /N/: n
MIN VALUE : 0
MAX VALUE : 60
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
... the command in full is MAKE_EXPERIMENTAL_DATAFILE
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
... the command in full is PLOT_DIAGRAM
```

2021.05.13.09.26.05
Time = 0,100000,300000,1000000
CELL #1



POST-1:
POST-1:
POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1:
POST-1: set-inter
... the command in full is SET_INTERACTIVE_MODE
--OK--
POST-1:

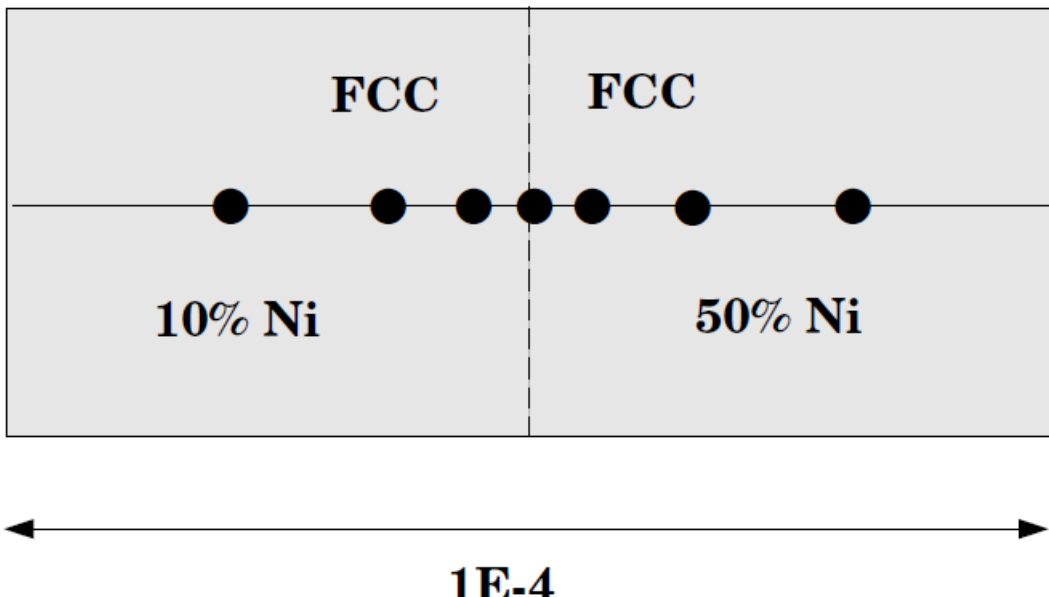


Example exa2b

Homogenization of a binary Fe-Ni alloy

Simple homogenization of a binary Fe-Ni alloy. We have put together a Ni rich and a Ni lean alloy. This example is identical to exa2a. However, in this example implicit time integration is used instead of the trapezoidal method for solving the PDEs.

T = 1400 K



exa2b-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa2b\setup.DCM.test"
SYS: @@
SYS: @@ One-phase problem.
SYS: @@ Homogenization of a binary Fe-Ni alloy.
SYS: @@ This example is identical to a2a but instead it uses implicit time
SYS: @@ integration instead of the trapezoidal method for solving the PDEs.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exa2b_setup.DCM
SYS:
SYS: @@
SYS: @@ FIRST DEFINE A LOG-FILE FOR THIS EXAMPLE
SYS: @@
SYS: set_log_file setup
Heading:
SYS: @@
SYS: @@ THEN GO TO THE DATABASE MODULE
SYS: @@
SYS: go da
... the command in full is GOTO_MODULE
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA                               /- DEFINED
DICTRA_FCC_A1   REJECTED
TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ USE THE TCFE DATABASE FOR THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw fedemo
... the command in full is SWITCH_DATABASE
Current database: Iron Demo Database v4.0

VA                               /- DEFINED
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_FEDEMO: @@
TDB_FEDEMO: def-system fe ni
... the command in full is DEFINE_SYSTEM
FE                           NI  DEFINED
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_FEDEMO: @@
TDB_FEDEMO: rej ph * all
... the command in full is REJECT
LIQUID:L           BCC_A2           LAVES_PHASE_C14
CBCC_A12          CUB_A13          FCC_A1
HCP_A3            REJECTED
TDB_FEDEMO: res ph fcc
... the command in full is RESTORE
FCC_A1             RESTORED
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_FEDEMO: @@
TDB_FEDEMO: get
... the command in full is GET_DATA
09:27:23,792 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
... the command in full is AMEND_PHASE_DESCRIPTION
PARAMETERS ...
FUNCTIONS ...

List of references for assessed data

'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
    volumes'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
-OK-
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_FEDEMO: @@ SWITCH TO THE MOBILITY DATABASE TO RETRIEVE DATA.
TDB_FEDEMO: @@
TDB_FEDEMO: app
... the command in full is APPEND_DATABASE
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
```

TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2
TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
SNOB1 = SGTE Noble Metal Alloys Database v1.2
STBC2 = SGTE Thermal Barrier Coating TDB v2.2
STBC1 = SGTE Thermal Barrier Coating TDB v1.1
SALT1 = SGTE Molten Salts Database v1.3
SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2 = TC Semi-Conductors v2.1
SLAG4 = Fe-containing Slag v4.1
SLAG3 = Fe-containing Slag v3.2
SLAG2 = Fe-containing Slag v2.2
SLAG1 = Fe-containing Slag v1.2
TCOX11 = Metal Oxide Solutions v11.0
TCOX10 = Metal Oxide Solutions v10.1
TCOX9 = Metal Oxide Solutions v9.0
TCOX8 = Metal Oxide Solutions v8.0
TCOX7 = Metal Oxide Solutions v7.0
TCOX6 = Metal Oxide Solutions v6.0
TCOX5 = Metal Oxide Solutions v5.1
TCOX4 = Metal Oxide Solutions v4.1
ION3 = Ionic Solutions v3.0
ION2 = Ionic Solutions v2.6
ION1 = Ionic Solutions v1.5
NOX2 = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4 = Solder Alloys v4.0
TCSLD3 = Solder Alloys v3.3
TCSLD2 = Solder Alloys v2.0
TCSLD1 = Solder Alloys v1.1
TCSI1 = Ultrapure Silicon v1.2
TCMP2 = Materials Processing v2.5
TCES1 = Combustion/Sintering v1.1
TCSC1 = Super Conductor v1.0
TCFC1 = SOFC Database v1.0
TCNF2 = Nuclear Fuels v2.1b
NUMT2 = Nuclear Materials v2.1
NUOX4 = Nuclear Oxides v4.2
NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephista-19
MEPH15 = IRSN Mephista-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQSO2 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SDDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.2
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3
MOBFE1 = Steels/Fe-Alloys Mobility v1.1
MOBFE2 = Steels/Fe-Alloys Mobility v2.0
MOBFE3 = Steels/Fe-Alloys Mobility v3.0
MOBFE4 = Steels/Fe-Alloys Mobility v4.0
MOBFE5 = Steels/Fe-Alloys Mobility v5.0

```

MOBFE6 = Steels/Fe-Alloys Mobility v6.0
MOBN15 = Ni-Alloys Mobility v5.1
MOBN14 = Ni-Alloys Mobility v4.1
MOBN13 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBSI1 = Si-Alloys Mobility v1.0
MOBSDL1 = Solder-Alloys Mobility v1.1
MOBT14 = Ti-Alloys Mobility v4.0
MOBT13 = Ti-Alloys Mobility v3.1
MOBT12 = Ti-Alloys Mobility v2.0
MOBT11 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

```

DATABASE NAME /FEDEMO/: mfedemo

Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys fe ni
... the command in full is DEFINE_SYSTEM

FE DEFINED
APP: rej ph * all
... the command in full is REJECT

BCC_A2 FCC_A1 REJECTED
APP: res ph fcc
... the command in full is RESTORE

FCC_A1 RESTORED
APP: get
... the command in full is GET_DATA

ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'

-OK-

APP:

APP: @@

APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP

APP: @@

APP: go d-m

... the command in full is GOTO_MODULE

NO TIME STEP DEFINED

DIC>

DIC> @@

DIC> @@ ENTER THE GLOBAL CONDITION T

DIC> @@

DIC> set-cond

... the command in full is SET_CONDITION

GLOBAL OR BOUNDARY CONDITION /GLOBAL/: glob

VARIABLE : T

LOW TIME LIMIT /0/: 0

T(TIME,X)= 1400;

HIGH TIME LIMIT /*/: *

ANY MORE RANGES /N/: N

DIC>

DIC> @@

DIC> @@ ENTER A REGION

DIC> @@

DIC> enter-region

REGION NAME : austenite

DIC>

DIC> @@

DIC> @@ ENTER A GRID INTO THE REGION.

DIC> @@ AS IN EXAMPLE a2a WE WANT SEVERAL POINTS IN THE MIDDLE OF THE REGION,

DIC> @@ SO A DOUBLE GEOMETRIC GRID IS ALSO CONSTRUCTED IN THIS EXAMPLE.

DIC> @@

DIC> enter-grid

... the command in full is ENTER_GRID_COORDINATES

REGION NAME : /AUSTENITE/: austenite

WIDTH OF REGION /1/: 1e-4

TYPE /LINEAR/: double

NUMBER OF POINTS /50/: 60

VALUE OF R IN THE GEOMETRICAL SERIE FOR LOWER PART OF REGION: 0.9

VALUE OF R IN THE GEOMETRICAL SERIE FOR UPPER PART OF REGION: 1.11

DIC>

DIC> @@

DIC> @@ ENTER ACTIVE PHASES INTO THE REGIONS

DIC> @@

DIC> enter-phase

... the command in full is ENTER_PHASE_IN_REGION

ACTIVE OR INACTIVE PHASE /ACTIVE/: act

REGION NAME : /AUSTENITE/: austenite

PHASE TYPE /MATRIX/: matrix

PHASE NAME: /NONE/: fcc#1

DIC>

DIC> @@

DIC> @@ ENTER THE INITIAL Ni COMPOSITION INTO THE PHASE. READ DATA FROM

DIC> @@ THE FILE ni.dat WHICH CONTAINS THE Ni-PROFILE.

exa2b-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa2b\run.DCM.test"
... the command in full is MACRO_FILE_OPEN
DIC>
DIC>
DIC> @@ exa2b_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE a2b
DIC> @@
DIC> @@
DIC> @@
DIC> @@ LET US DEFINE A LOG-FILE FOR THIS EXAMPLE
DIC> @@
DIC> set-log-file run
AMBIGUOUS COMMAND, USE HELP
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ SETUP FROM FILE
DIC> @@
DIC> go d-m
... the command in full is GOTO_MODULE
TIME STEP AT TIME 0.00000E+00
DIC> read exa2b
... the command in full is READ_WORKSPACES
... the command in full is DEFINE_COMPONENTS
... the command in full is SELECT_EQUILIBRIUM
OK
DIC>
DIC> @@
DIC> @@ Start the simulation
DIC> @@
DIC> simulate
... the command in full is SIMULATE_REACTION
... the command in full is SET_NUMERICAL_LIMITS
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
2 GRIDPOINT(S) ADDED TO CELL #1 REGION: AUSTENITE
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.1001000E-03 DT = 0.1000000E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.40010010 DT = 0.40000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 10.403455 DT = 10.003355 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 30.410165 DT = 20.006710 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 70.423585 DT = 40.013420 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 150.45043 DT = 80.026840 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 310.50411 DT = 160.05368 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 630.61147 DT = 320.10736 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 1270.8262 DT = 640.21472 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 2551.2556 DT = 1280.4294 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 5112.1145 DT = 2560.8589 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 10233.832 DT = 5121.7178 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 20477.268 DT = 10243.436 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .70888824540617 NI = .29111175459383
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 40964.139 DT = 20486.871 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406171 NI = .291111754593829
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 81937.881 DT = 40973.742 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406172 NI = .291111754593828
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 163885.37 DT = 81947.484 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406173 NI = .291111754593827
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 263885.37 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406169 NI = .291111754593831
```

```

TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep          0 seconds
TIME = 363885.37 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406171 NI = .291111754593829
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep          0 seconds
TIME = 463885.37 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406166 NI = .291111754593834
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep          0 seconds
TIME = 563885.37 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406164 NI = .291111754593836
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep          0 seconds
TIME = 663885.37 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406165 NI = .291111754593835
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep          0 seconds
TIME = 763885.37 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406162 NI = .291111754593838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep          0 seconds
TIME = 863885.37 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406163 NI = .291111754593836
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep          0 seconds
TIME = 963885.37 DT = 100000.00 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: FE = .708888245406167 NI = .291111754593833
TOTAL SIZE OF SYSTEM: 1E-04 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME    0.0000000
DELETING TIME-RECORD FOR TIME   0.10000000E-06
DELETING TIME-RECORD FOR TIME   0.10010000E-03
DELETING TIME-RECORD FOR TIME   0.40010010
DELETING TIME-RECORD FOR TIME   10.403455
DELETING TIME-RECORD FOR TIME   30.410165
DELETING TIME-RECORD FOR TIME   70.423585
DELETING TIME-RECORD FOR TIME   150.45043
DELETING TIME-RECORD FOR TIME   310.50411
DELETING TIME-RECORD FOR TIME   630.61147
DELETING TIME-RECORD FOR TIME   1270.8262
DELETING TIME-RECORD FOR TIME   2551.2556
DELETING TIME-RECORD FOR TIME   5112.1145
DELETING TIME-RECORD FOR TIME   10233.832
DELETING TIME-RECORD FOR TIME   20477.268
DELETING TIME-RECORD FOR TIME   40964.139
DELETING TIME-RECORD FOR TIME   81937.881
DELETING TIME-RECORD FOR TIME   163885.37
DELETING TIME-RECORD FOR TIME   263885.37
DELETING TIME-RECORD FOR TIME   363885.37
DELETING TIME-RECORD FOR TIME   463885.37
DELETING TIME-RECORD FOR TIME   563885.37
DELETING TIME-RECORD FOR TIME   663885.37
DELETING TIME-RECORD FOR TIME   763885.37
DELETING TIME-RECORD FOR TIME   863885.37

KEEPING TIME-RECORD FOR TIME    963885.37
AND FOR TIME                   1000000.0
WORKSPACE RECLAIMED

TIMESTEP AT      1000000.00      SELECTED
```

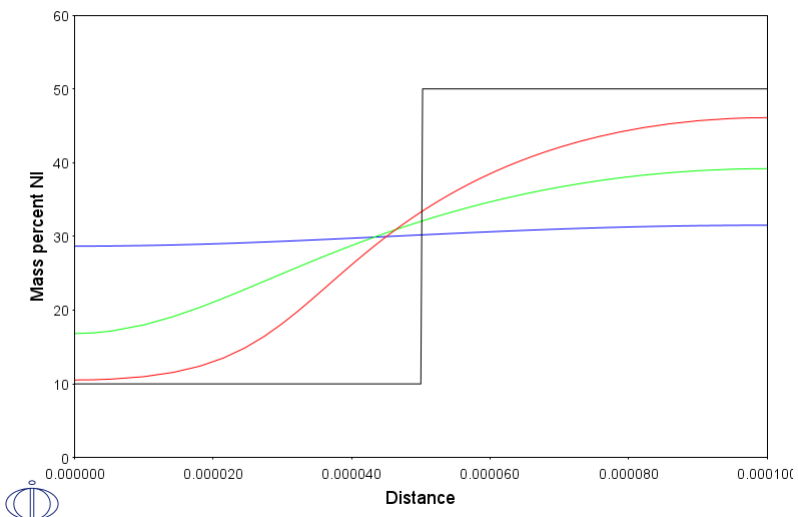
```

DIC>
DIC> @@
DIC> @@ THE SIMULATION IS FINISHED
DIC> @@
DIC>
DIC> set-inter
... the command in full is SET_INTERACTIVE
--OK--
DIC>
```

exa2b-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa2b\plot.DCM.test"
... the command in full is MACRO_FILE_OPEN
DIC>
DIC>
DIC> @@ exa2b_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE a2b
DIC> @@
DIC> @@
DIC> @@
DIC> @@ LET US DEFINE A LOG-FILE FOR THIS EXAMPLE
DIC> @@
DIC> set-log-file plot
AMBIGUOUS COMMAND, USE HELP
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
... the command in full is GOTO_MODULE
TIME STEP AT TIME 1.00000E+06
DIC> read exa2b
... the command in full is READ_WORKSPACES
... the command in full is DEFINE_COMPONENTS
... the command in full is SELECT_EQUILIBRIUM
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
... the command in full is POST_PROCESSOR
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1:
POST-1: @@
POST-1: @@ PLOT SOME CONCENTRATION PROFILES
POST-1: @@
POST-1: s-d-a
... the command in full is SET_DIAGRAM_AXIS
AXIS (X, Y OR Z) : x
VARIABLE : dist
INFO: Distance is set as independent variable
... the command in full is SET_INDEPENDENT_VARIABLE
DISTANCE : /GLOBAL/: glo
POST-1:
POST-1: s-d-a
... the command in full is SET_DIAGRAM_AXIS
AXIS (X, Y OR Z) : y
VARIABLE : w-p
FOR COMPONENT : ni
POST-1:
POST-1: s-p-c
... the command in full is SET_PLOT_CONDITION
CONDITION /TIME/: time
VALUE(S) /LAST/: 0 1e5 3e5 1e6
POST-1:
POST-1: @@
POST-1: @@ SET SCALING ON Y-AXIS BEFORE PLOTTING
POST-1: @@
POST-1: s-s-s
... the command in full is SET_SCALING_STATUS
AXIS (X, Y OR Z) : y
AUTOMATIC SCALING (Y OR N) /N/: n
MIN VALUE : 0
MAX VALUE : 60
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
... the command in full is MAKE_EXPERIMENTAL_DATAFILE
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
... the command in full is PLOT_DIAGRAM
```

2021.05.13.09.29.25
Time = 0,100000,300000,1000000
CELL #1



POST-1:
POST-1:
POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1:
POST-1: set-inter
... the command in full is SET_INTERACTIVE_MODE
--OK--
POST-1:



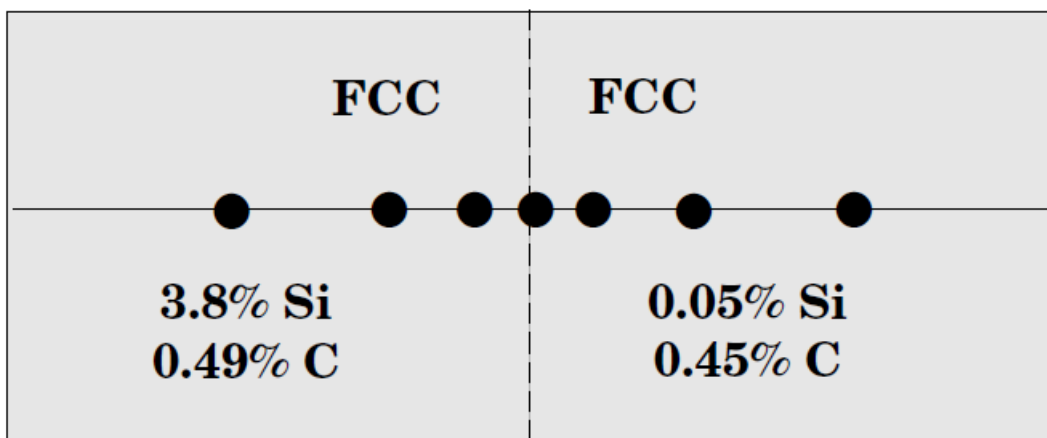
Example exa3

Uphill diffusion in an Fe-Si-C alloy

Simulation of uphill diffusion in a ternary single phase austenite matrix due to the classical darken experiment published by L.S. Darken (Trans. Aime, v.180 (1949), pp. 430-438).

In this example, two pieces of austenite (3.80 wt%Si, 0.49 wt%C) and (0.05 wt%Si, 0.45 wt%C) are put together and are subsequently annealed at 1050 C for 13 days. As both pieces are austenite they must be entered into the same region. This is done by giving the compositions of Si and C in each gridpoint individually. These data are then stored on file.

$$T = 1323 \text{ K}$$



exa3-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa3\setup.DCM.test"
SYS: @@
SYS: @@ One-phase problem.
SYS: @@ Uphill diffusion in an Fe-Si-C alloy
SYS: @@ This is an example to simulate uphill diffusion in a ternary single
SYS: @@ phase austenite matrix due to the classical Darken experiment published
SYS: @@ by L.S. Darken: Trans. Aime, v.180 (1949), pp. 430-438.
SYS: @@
SYS: @@ In this example two pieces of austenite (3.80 wt%Si, 0.49 wt%C) and
SYS: @@ (0.05 wt%Si, 0.45 wt%C) are put together and are subsequently annealed
SYS: @@ at 1050C for 13 days. As both pieces are austenite they must be entered
SYS: @@ into the same region. This is done by individually giving the compositions
SYS: @@ of Si and C in each grid point. These data are then stored to file.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ darken_setup.DCM
SYS:
SYS: @@
SYS: @@ Note that LOG-FILES used previously in examples a2a and a2b are
SYS: @@ no longer used.
SYS: @@
SYS: @@
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA          /- DEFINED
DICTRA_FCC_A1 REJECTED

TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ USE A TCFE DATABASE FOR THE THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw tcfef9
Current database: Steels/Fe-Alloys v9.3

VA          /- DEFINED
L12_FCC      B2_BCC           DICTRA_FCC_A1
REJECTED

TDB_TCFE9: def-sys fe si c
FE           SI               C
DEFINED

TDB_TCFE9: rej ph * all
GAS:G          LIQUID:L        BCC_A2
FCC_A1         HCP_A3         CBCC_A12
CUB_A13        DIAMOND_FCC_A4 GRAPHITE
CEMENTITE      M23C6          M7C3
M5C2          KSI_CARBIDE    FE4N_LP1
FECN_CHI       LAVES_PHASE_C14 M3SI
CR3SI         FE2SI          FESI2_H
FESI2_L        MSI            M5SI3
AL4C3         FE8SI2C       SIC
AL5FE4        MP_B31         M2P_C22
M203C:I      REJECTED

TDB_TCFE9: res ph fcc
FCC_A1        RESTORED

TDB_TCFE9: get
09:31:17,315 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
FUNCTIONS .....

List of references for assessed data

'A. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.-G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
  volumes'
'X.-G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'P. Gustafsson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'B. Hallstedt, unpublished work (2016); C-Fe-Mn Epsilon martensite.'
'J. Lacaze and B. Sundman, Metall. Mater. Trans. A, 22A (1991) 2211-2223;
  Fe-Si and Fe-Si-C'
'J. Miettinen and B. Hallstedt, CALPHAD, 22 (1998) 231-256; Fe-Si and Fe
  -Si-C'

-OK-
TDB_TCFE9:
TDB_TCFE9: @@
TDB_TCFE9: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_TCFE9: @@
TDB_TCFE9: app mobfe4
Current database: Steels/Fe-Alloys Mobility v4.0

VA DEFINED
B2_BCC REJECTED
APP: def-sys fe si c
FE           SI               C
DEFINED
APP: rej ph * all
```

```

BCC_A2          CEMENTITE          FCC_A1
FE4N_LP1        HCP_A3           LIQUID:L
REJECTED
APP: res ph fcc
FCC_A1 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'D. Bergner et al., Defect and Diffusion Forum 66-69(1989)409. Impurity
diffusion of Si in fcc Fe.'
-OK-
APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 1323; * N
DIC>
DIC> @@
DIC> @@ ENTER THE REGION austenite
DIC> @@
DIC> enter-region
REGION NAME : austenite
DIC>
DIC> @@
DIC> @@ ENTER THE GRID
DIC> @@ NOTE THAT GRID POINT DISTANCES ARE SMALLEST AROUND THE MIDDLE
DIC> @@
DIC> enter-grid
REGION NAME : /AUSTENITE/: austenite
WIDTH OF REGION /1/: 50E-3
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER THE PHASE INTO A REGION (BOTH PIECES ARE AUSTENITIC)
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /AUSTENITE/: austenite
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC> @@
DIC> @@ ENTER COMPOSITIONS INTO THE PHASE
DIC> @@
DIC> enter-composition
REGION NAME : /AUSTENITE/: austenite
PHASE NAME: /FCC_A1/: fcc#1
DEPENDENT COMPONENT ? /SI/: FE
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: C func 0.49-0.04*hs(x-25e-3);
PROFILE FOR /SI/: SI func 3.80-3.75*hs(x-25e-3);
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME AND OTHER SIMULATION PARAMETERS
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /1/: 1e10
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /1E+09/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC> save exa3 Y
DIC>
DIC> set-inter
--OK--
DIC>
```

exa3-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa3\run.DCM.test"
DIC>
DIC>
DIC> @@ darken_run.DCM
DIC>
DIC>
DIC> @@ ENTER THE DICTRA MONITOR
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC>
DIC> @@
DIC> @@ READ SETUP FROM FILE AND START SIMULATION
DIC> @@
DIC> read exa3
OK
DIC>
DIC> sim
Region: AUSTENITE
double geometric
coarse at outer boundaries dense at 0.25047E-01
lower part 0.80000      22
upper part 1.2500       22
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: C = .0215351122808224 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
U-FRACTION IN SYSTEM: C = .0215351122808224 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808224 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          1 seconds
TIME = 0.1001000E-03 DT = 0.1000000E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808224 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 0.40010010 DT = 0.40000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808224 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 21.165600 DT = 20.765500 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808224 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 62.696601 DT = 41.531000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808224 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 145.75860 DT = 83.062001 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808224 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 311.88260 DT = 166.12400 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808223 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 644.13060 DT = 332.24800 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808223 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 1308.6266 DT = 664.49600 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808225 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 2637.6186 DT = 1328.9920 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .021535112280823 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 5295.6026 DT = 2657.9840 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808237 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 10611.571 DT = 5315.9680 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808241 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 21243.507 DT = 10631.936 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808231 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 42507.379 DT = 21263.872 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .02153511228082 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 85035.123 DT = 42527.744 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808147 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          0 seconds
TIME = 170090.61 DT = 85055.489 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808123 FE = .96291921367461
SI = .0370807863253901
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep          1 seconds
```

```

TIME = 340201.59 DT = 170110.98 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122808354 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 680423.54 DT = 340221.95 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122809501 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 1360867.5 DT = 680443.91 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122812752 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 2721755.3 DT = 1360887.8 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122812752 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 5443530.9 DT = 2721775.6 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122811047 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 10887082. DT = 5443551.3 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122807851 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 21774185. DT = 10887103. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122805233 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 43548390. DT = 21774205. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122804133 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 87096800. DT = 43548410. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122803482 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.17419362E+09 DT = 87096820. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122803353 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 1 seconds
TIME = 0.34838726E+09 DT = 0.17419364E+09 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .021535112280406 FE = .96291921367461
SI = .0370807863253899
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.69677454E+09 DT = 0.34838728E+09 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .021535112280591 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.13935491E+10 DT = 0.69677456E+09 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122806838 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.23935491E+10 DT = 0.10000000E+10 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122809351 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.33935491E+10 DT = 0.10000000E+10 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .021535112281113 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.43935491E+10 DT = 0.10000000E+10 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122816696 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.53935491E+10 DT = 0.10000000E+10 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122822483 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.63935491E+10 DT = 0.10000000E+10 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122828043 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.73935491E+10 DT = 0.10000000E+10 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122833152 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.83935491E+10 DT = 0.10000000E+10 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122836845 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 0 seconds
TIME = 0.93935491E+10 DT = 0.10000000E+10 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122841004 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
CPU time used in timestep 1 seconds
TIME = 0.10000000E+11 DT = 0.60645090E+09 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0215351122842448 FE = .96291921367461
SI = .03708078632539
TOTAL SIZE OF SYSTEM: .05 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 0.10010000E-03
DELETING TIME-RECORD FOR TIME 0.40010010

```

```
DELETING TIME-RECORD FOR TIME      21.165600
DELETING TIME-RECORD FOR TIME      62.696601
DELETING TIME-RECORD FOR TIME     145.75860
DELETING TIME-RECORD FOR TIME     311.88260
DELETING TIME-RECORD FOR TIME     644.13060
DELETING TIME-RECORD FOR TIME    1308.6266
DELETING TIME-RECORD FOR TIME    2637.6186
DELETING TIME-RECORD FOR TIME    5295.6026
DELETING TIME-RECORD FOR TIME   10611.571
DELETING TIME-RECORD FOR TIME   21243.507
DELETING TIME-RECORD FOR TIME   42507.379
DELETING TIME-RECORD FOR TIME   85035.123
DELETING TIME-RECORD FOR TIME 170090.61
DELETING TIME-RECORD FOR TIME 340201.59
DELETING TIME-RECORD FOR TIME 680423.54
DELETING TIME-RECORD FOR TIME 1360867.5
DELETING TIME-RECORD FOR TIME 2721755.3
DELETING TIME-RECORD FOR TIME 5443530.9
DELETING TIME-RECORD FOR TIME 10887082.
DELETING TIME-RECORD FOR TIME 21774185.
DELETING TIME-RECORD FOR TIME 43548390.
DELETING TIME-RECORD FOR TIME 87096800.
DELETING TIME-RECORD FOR TIME 0.17419362E+09
DELETING TIME-RECORD FOR TIME 0.34838726E+09
DELETING TIME-RECORD FOR TIME 0.69677454E+09
DELETING TIME-RECORD FOR TIME 0.13935491E+10
DELETING TIME-RECORD FOR TIME 0.23935491E+10
DELETING TIME-RECORD FOR TIME 0.33935491E+10
DELETING TIME-RECORD FOR TIME 0.43935491E+10
DELETING TIME-RECORD FOR TIME 0.53935491E+10
DELETING TIME-RECORD FOR TIME 0.63935491E+10
DELETING TIME-RECORD FOR TIME 0.73935491E+10
DELETING TIME-RECORD FOR TIME 0.83935491E+10
```

```
KEEPING TIME-RECORD FOR TIME 0.93935491E+10
AND FOR TIME 0.10000000E+11
WORKSPACE RECLAIMED
```

```
TIMESTEP AT 0.100000000E+11 SELECTED
```

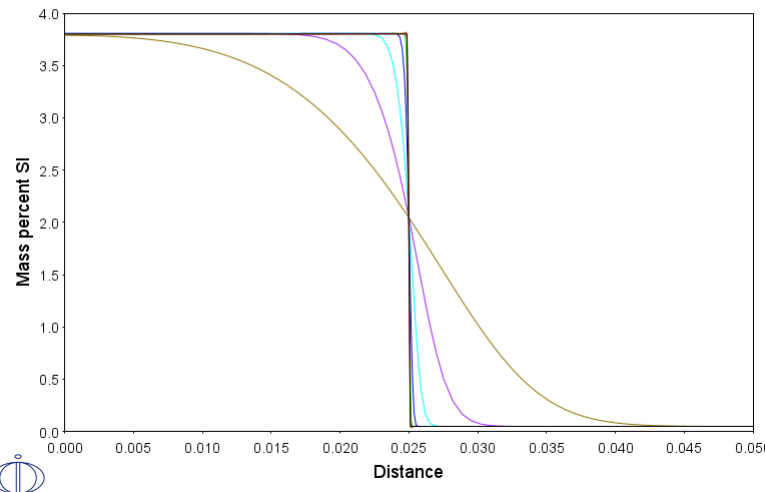
```
DIC>
DIC> set-inter
--OK--
DIC>
```

exa3-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa3\plot.DCM.test"
DIC>
DIC>
DIC> @@ darken_plot.DCM
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MODULE AND SPECIFY THE STORE-RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 1.00000E+10
DIC> read exa3
OK
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ PLOT THE CONCENTRATION PROFILE FOR Si AT TIMES 0, 1E5, 1123200, 1E7,
POST-1: @@ 1E8, 1E9 AND 1E10 S
POST-1: @@
POST-1: @@ SET DISTANCE IN SYSTEM AS X-AXIS, WEIGHT-% SI ON Y-AXIS AND SPECIFY
POST-1: @@ FOR WHICH SIMULATION TIMES TO PLOT THE PROFILES.
POST-1: @@
POST-1: set-diagram-axis x distance global
INFO: Distance is set as independent variable
POST-1: set-diagram-axis y weight-percent si
POST-1: set-plot-condition time 0 1E5 1123200 1e7 1E8 1E9 1E10
POST-1:
POST-1: @@
POST-1: @@ PLOT THE DIAGRAM
POST-1: @@
POST-1: set-title
TITLE : Figure a3.1
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Figure a3.1

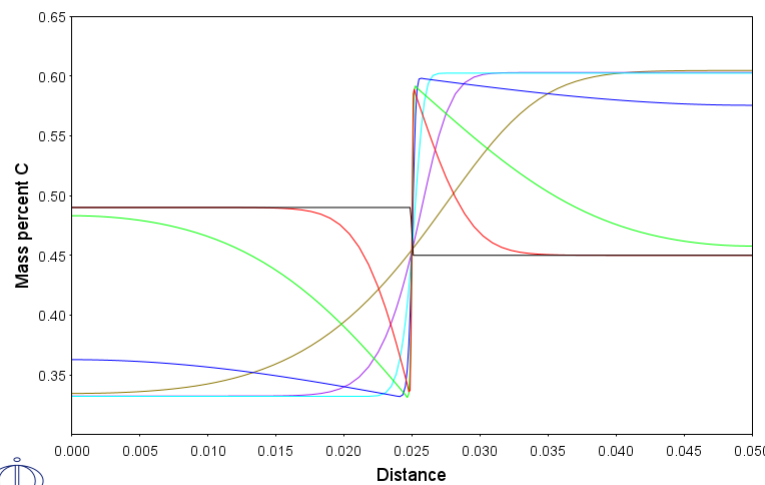
2021.05.13.09.35.49
Time=0,100000,1123200,1E+07,1E+08,1E+09,1E+10
CELL #1



```
POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1: @@
POST-1: @@ PLOT THE CONCENTRATION PROFILES FOR C
POST-1: @@
POST-1: @@ WE ONLY NEED TO CHANGE THE Y-AXIS
POST-1: @@
POST-1: set-diagram-axis y w-p c
POST-1: set-title Figure a3.2
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Figure a3.2

2021.05.13.09.35.51
 Time = 0,100000,1123200,1E+07,1E+08,1E+09,1E+10
 CELL #1

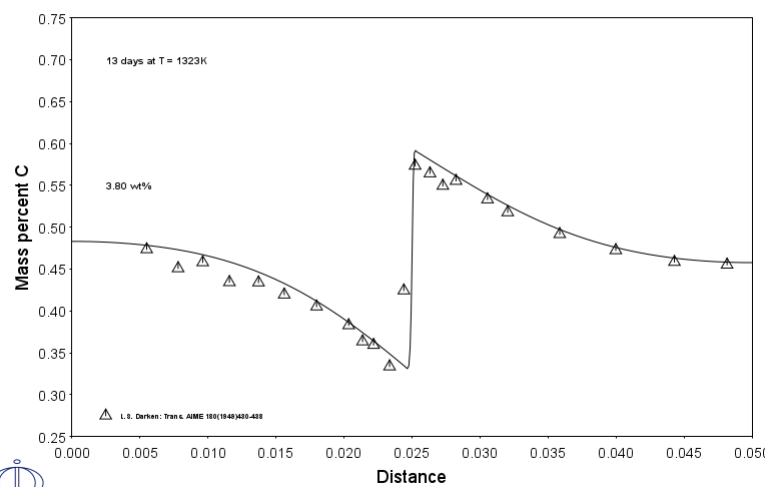


```

POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1: @@ 
POST-1: @@ COMPARE WITH DARKEN'S EXPERIMENTS
POST-1: @@
POST-1: append_experimental_data yes exa3.exp 0; 1
POST-1:
POST-1: set-plot-condition time 1123200
POST-1:
POST-1: s-s-s
AXIS (X, Y OR Z) : y
AUTOMATIC SCALING (Y OR N) /N/: n
MIN VALUE : 0.25
MAX VALUE : 0.75
POST-1:
POST-1: set-title Figure a3.3
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
  
```

Figure a3.3

2021.05.13.09.35.53
 Time = 1123200
 CELL #1



```

POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1: set-inter
--OK--
POST-1:
  
```

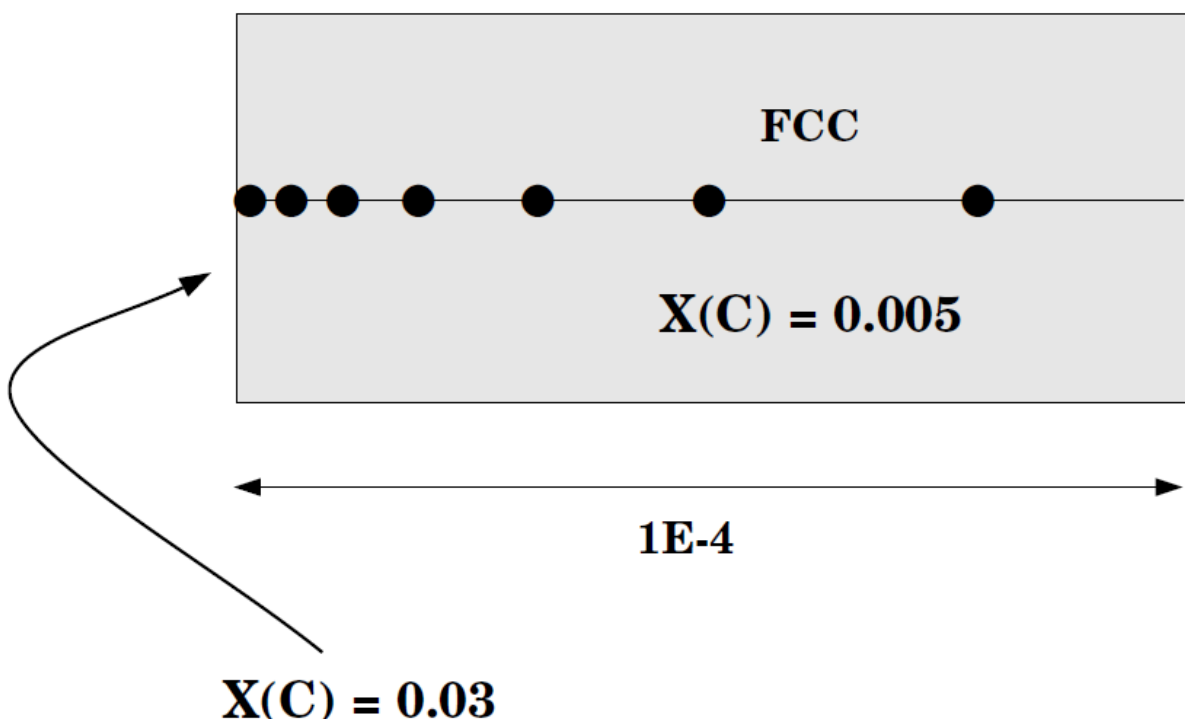


Example exa4

Carburization of a binary Fe-C alloy: Comparison to analytical erf solution

This is a simple binary simulation with one single phase region. It compares a numerical simulation with an analytical erf-solution. For this reason a special database is created (*erf.tdb*) where the diffusion coefficient is set to a concentration independent value.

T = 1200 K



exa4-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa4\setup.DCM.test"
SYS: @@
SYS: @@ One-phase problem.
SYS: @@ Carburization of binary Fe-C alloy: Comparison to an analytical erf solution
SYS: @@ This is a simple binary simulation with a single phase region.
SYS: @@ The numerical simulation is compared with an analytical erf solution.
SYS: @@ For this reason a special database erf.tdb is created where the
SYS: @@ diffusion coefficient is set to a concentration independent value.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exa4_setup.DCM
SYS:
SYS:
SYS: @@
SYS: @@ READ THE DATA FROM THE DATABASES
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: SW FEDEMO
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-system fe,c
 FE C DEFINED
TDB_FEDEMO: rej-ph *
 GAS:G LIQUID:L BCC_A2
 LAVES_PHASE_C14 CBCC_A12 CEMENTITE
 CUB_A13 DIAMOND_FCC_A4 FCC_A1
 GRAPHITE HCP_A3 KSI_CARBIDE
 M23C6 M5C2 M7C3
REJECTED
TDB_FEDEMO: rest-ph fcc
 FCC_A1 RESTORED
TDB_FEDEMO: get
09:37:14,437 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES
ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
-OK-
TDB_FEDEMO:
TDB_FEDEMO: append user exa4.TDB
Current database: User defined Database
This database does not support the DATABASE_INFORMATION command

VA DEFINED
09:37:20,624 INFO USER_790547025_12, number of lines read: 29
09:37:31,884 INFO Parsing of USER_790547025_12 completed in 11331 ms
TDB_APP: def-system fe,c
 FE C DEFINED
TDB_APP: rej-ph *
 FCC_A1 REJECTED
TDB_APP: rest-ph fcc
 FCC_A1 RESTORED
TDB_APP: get
ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS
-OK-
TDB_APP:
TDB_APP: @@
TDB_APP: @@ GO TO THE DICTRA MODULE AND SET UP THE SYSTEM
TDB_APP: @@
TDB_APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 1200; * N
DIC>
DIC> @@
DIC> @@ ENTER THE REGION steel
DIC> @@
DIC> enter-region
REGION NAME : steel
DIC>
DIC> @@

```

DIC> @@ ENTER THE GRID
DIC> @@ CARBON ENTERS THE SYSTEM FROM THE LOWER BOUNDARY AND CONSEQUENTLY
DIC> @@ MORE POINTS ARE REQUIRED AT THAT BOUNDARY. THIS IS WHY A GEOMETRIC
DIC> @@ GRID IS USED.
DIC> @@
DIC> enter-grid
REGION NAME : /STEEL/: steel
WIDTH OF REGION /1/: 1E-4
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER THE PHASE INTO THE REGION
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /STEEL/: steel
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITION IN THE FCC PHASE
DIC> @@
DIC> enter-composition
REGION NAME : /STEEL/: steel
PHASE NAME: /FCC_A1/: fcc#1
COMPOSITION TYPE /MOLE_FRACTION/: mole-fraction
PROFILE FOR /C/: c
TYPE /LINEAR/: linear
VALUE OF FIRST POINT : 0.005
VALUE OF LAST POINT : /5E-3/: 0.005
DIC>
DIC>
DIC> @@
DIC> @@ SET A FIXED COMPOSITION AS THE BOUNDARY VALUE
DIC> @@
DIC> set-condition
GLOBAL OR BOUNDARY CONDITION /GLOBAL/: boundary
BOUNDARY /LOWER/: lower
CONDITION TYPE /CLOSED_SYSTEM/: state-variable-value
State variable expression #1 : /N=1/: n=1
State variable expression #2 : x(c)=0.03
DIC>
DIC>
DIC> @@
DIC> @@ SET A SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 100
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /10/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC>
DIC>
DIC> save exa4 Y
DIC>
DIC> set-inter
--OK--
DIC>

```

exa4-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa4\run.DCM.test"
DIC>
DIC>
DIC> @@ exa4_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE a4
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exa4
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim
Region: STEEL
single geometric dense at 0.0000
1.1388 96
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: C = .0050251256281407 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
U-FRACTION IN SYSTEM: C = .0050251256281407 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00502599895421297 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.26001020E-05 DT = 0.25001020E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00502674234772398 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.76003060E-05 DT = 0.50002040E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00502762855271361 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.17600714E-04 DT = 0.10000408E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00502880490898696 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.37601530E-04 DT = 0.20000816E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00503042289514107 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.77603162E-04 DT = 0.40001632E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00503268130112881 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.15760643E-03 DT = 0.80003264E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00503585491527872 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.31761295E-03 DT = 0.16000653E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00504032903043467 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.63762601E-03 DT = 0.32001306E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00504664654067874 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 0.12776521E-02 DT = 0.64002611E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00505557392100181 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.25577043E-02 DT = 0.12800522E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00506819425567398 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.51178088E-02 DT = 0.25601044E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00508603865206295 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.10238018E-01 DT = 0.51202089E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00511127199989312 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.20478435E-01 DT = 0.10240418E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0051469556180961 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.40959271E-01 DT = 0.20480836E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00519741865567008 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.81920942E-01 DT = 0.40961671E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0052687830573381 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.16384428 DT = 0.81923342E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00536970755217346 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.32769097 DT = 0.16384468 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00551243555924854 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.65538434 DT = 0.32769337 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00571428313781583 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 1.3107711 DT = 0.65538674 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00599973850521004 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 2.6215446 DT = 1.3107735 SUM OF SQUARES = 0.0000000
```

```

U-FRACTION IN SYSTEM: C = .00640343298842195 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 5.2430915 DT = 2.6215470 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00697433052866847 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 10.486185 DT = 5.2430939 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00778159421638383 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 20.486185 DT = 10.000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00887736257293564 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 30.486185 DT = 10.000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00972196664325369 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 40.486185 DT = 10.000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .010436351986536 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 50.486185 DT = 10.000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0110669328932657 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 60.486185 DT = 10.000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0116377078880251 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 70.486185 DT = 10.000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0121630223717782 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 90.486185 DT = 10.000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0131119785059757 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 100.00000 DT = 9.5138146 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .013526298212787 FE = 1
TOTAL SIZE OF SYSTEM: 1E-04 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 0.26001020E-05
DELETING TIME-RECORD FOR TIME 0.76003060E-05
DELETING TIME-RECORD FOR TIME 0.17600714E-04
DELETING TIME-RECORD FOR TIME 0.37601530E-04
DELETING TIME-RECORD FOR TIME 0.77603162E-04
DELETING TIME-RECORD FOR TIME 0.15760643E-03
DELETING TIME-RECORD FOR TIME 0.31761295E-03
DELETING TIME-RECORD FOR TIME 0.63762601E-03
DELETING TIME-RECORD FOR TIME 0.12776521E-02
DELETING TIME-RECORD FOR TIME 0.25577043E-02
DELETING TIME-RECORD FOR TIME 0.51178088E-02
DELETING TIME-RECORD FOR TIME 0.10238018E-01
DELETING TIME-RECORD FOR TIME 0.20478435E-01
DELETING TIME-RECORD FOR TIME 0.40959271E-01
DELETING TIME-RECORD FOR TIME 0.81920942E-01
DELETING TIME-RECORD FOR TIME 0.16384428
DELETING TIME-RECORD FOR TIME 0.32769097
DELETING TIME-RECORD FOR TIME 0.65538434
DELETING TIME-RECORD FOR TIME 1.3107711
DELETING TIME-RECORD FOR TIME 2.6215446
DELETING TIME-RECORD FOR TIME 5.2430915
DELETING TIME-RECORD FOR TIME 10.486185
DELETING TIME-RECORD FOR TIME 20.486185
DELETING TIME-RECORD FOR TIME 30.486185
DELETING TIME-RECORD FOR TIME 40.486185
DELETING TIME-RECORD FOR TIME 50.486185
DELETING TIME-RECORD FOR TIME 60.486185
DELETING TIME-RECORD FOR TIME 70.486185
DELETING TIME-RECORD FOR TIME 80.486185

KEEPING TIME-RECORD FOR TIME 90.486185
AND FOR TIME 100.00000
WORKSPACE RECLAIMED

```

```

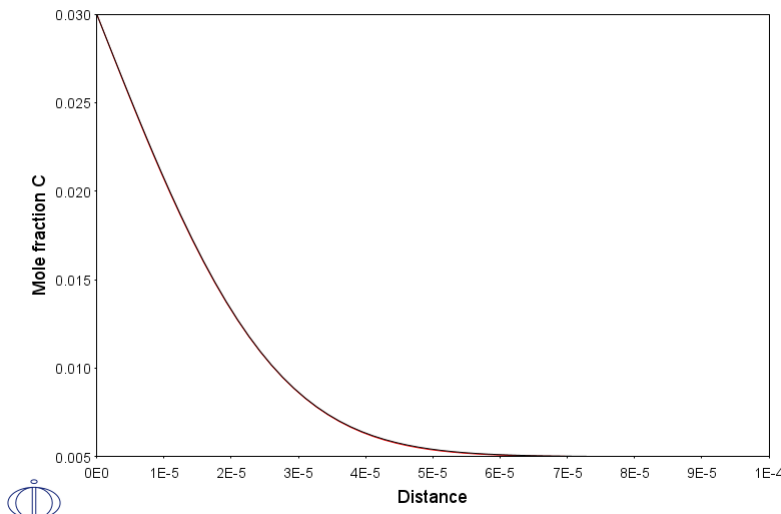
TIMESTEP AT 100.000000 SELECTED

DIC>
DIC> @@
DIC> @@ THE SIMULATION IS FINISHED
DIC> @@
DIC>
DIC> set-inter
--OK--
DIC>

```

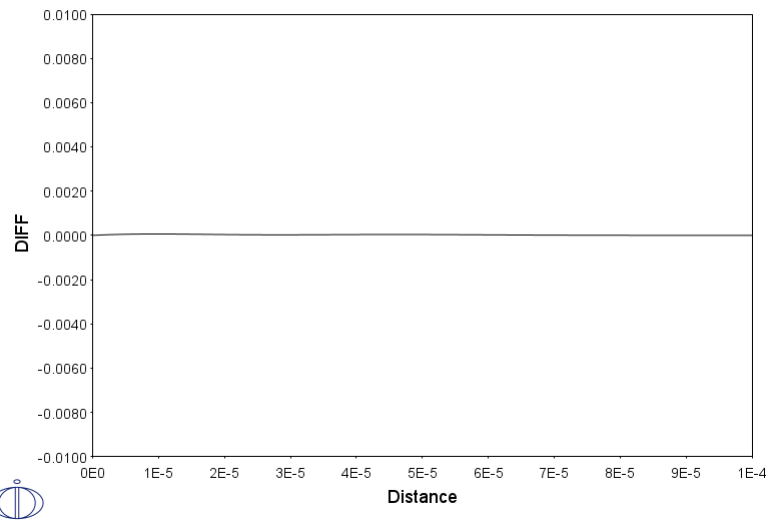
exa4-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa4\plot.DCM.test"
DIC>
DIC>
DIC> @@ exa4_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE exa4
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 1.00000E+02
DIC> read exa4
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ PLOT A COMPOSITION PROFILE
POST-1: @@
POST-1: s-d-a x distance global
INFO: Distance is set as independent variable
POST-1: s-d-a y x(c)
POST-1: s-p-c time 25
POST-1: @@
POST-1: @@
POST-1: @@ ENTER THE ANALYTICAL SOLUTION, CALLED ERFSOL
POST-1: @@
POST-1: enter-symbol
Function or table /FUNCTION/: function
NAME: erfsol
FUNCTION: 0.03-0.025*erf(gd/sqrt(4*dc(fcc,c,c,fe)*25));
POST-1:
POST-1: @@
POST-1: @@ COMPARE THE ANALYTICAL AND NUMERICAL SOLUTIONS
POST-1: @@
POST-1: enter-symbol
Function or table /FUNCTION/: table
NAME: aaa
Variable(s) x(c) erfsol
POST-1:
POST-1: s-d-a y aaa
COLUMN NUMBER /*/: 1 2
POST-1:
POST-1: set-axis-text
AXIS (X, Y OR Z) : y
AUTOMATIC AXIS TEXT (Y OR N) /N/: n
AXIS TEXT : Mole fraction C
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.09.39.35
Time = 25
CELL #1
```



```
POST-1:
POST-1:
POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1:
POST-1: @@
POST-1: @@ PLOT THE DIFFERENCE
POST-1: @@
POST-1: enter func diff=x(c)-erfsol;
POST-1: s-d-a y diff
POST-1: s-s-s y n -1e-2 1e-2
POST-1:
```

```
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
POST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot  
2021.05.13.09.39.36  
Time = 25  
CELL #1
```



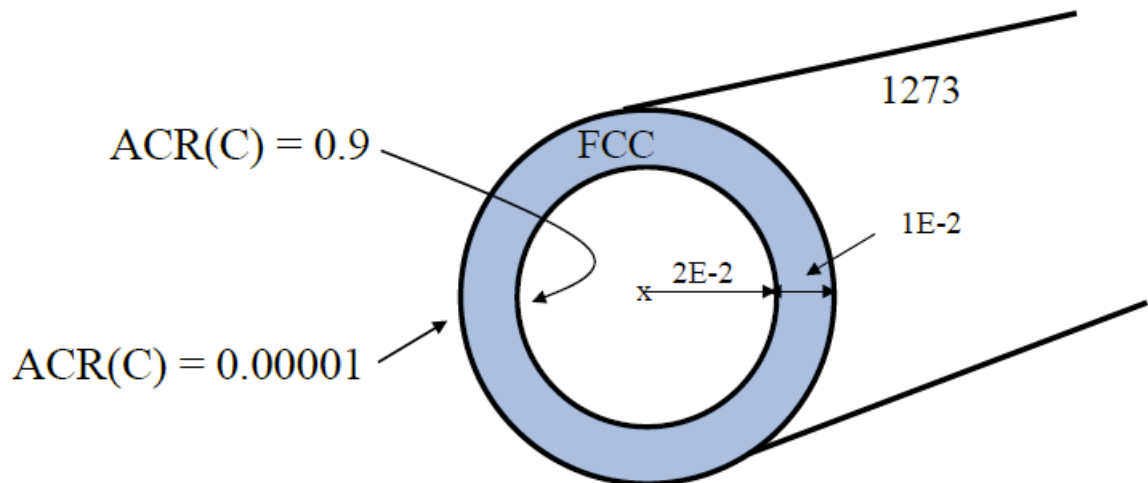
```
POST-1:  
POST-1:  
POST-1:  
POST-1:@?<Hit_return_to_continue>  
POST-1:  
POST-1: set-interactive  
--OK--  
POST-1:
```



Example exa6

Diffusion through a tube wall: Boundary conditions result in a gradient in C-activity

A simple example of diffusion through a tube wall. The tube-material is an Fe-0.6%Mn-0.7%Si-0.05%C alloy. On the inside wall a carbon activity of 0.9 is maintained whereas on the outside the C-activity is very low. This example demonstrates the use of the command SET-FIRST-INTERFACE as well as the use of MIXED boundary conditions.



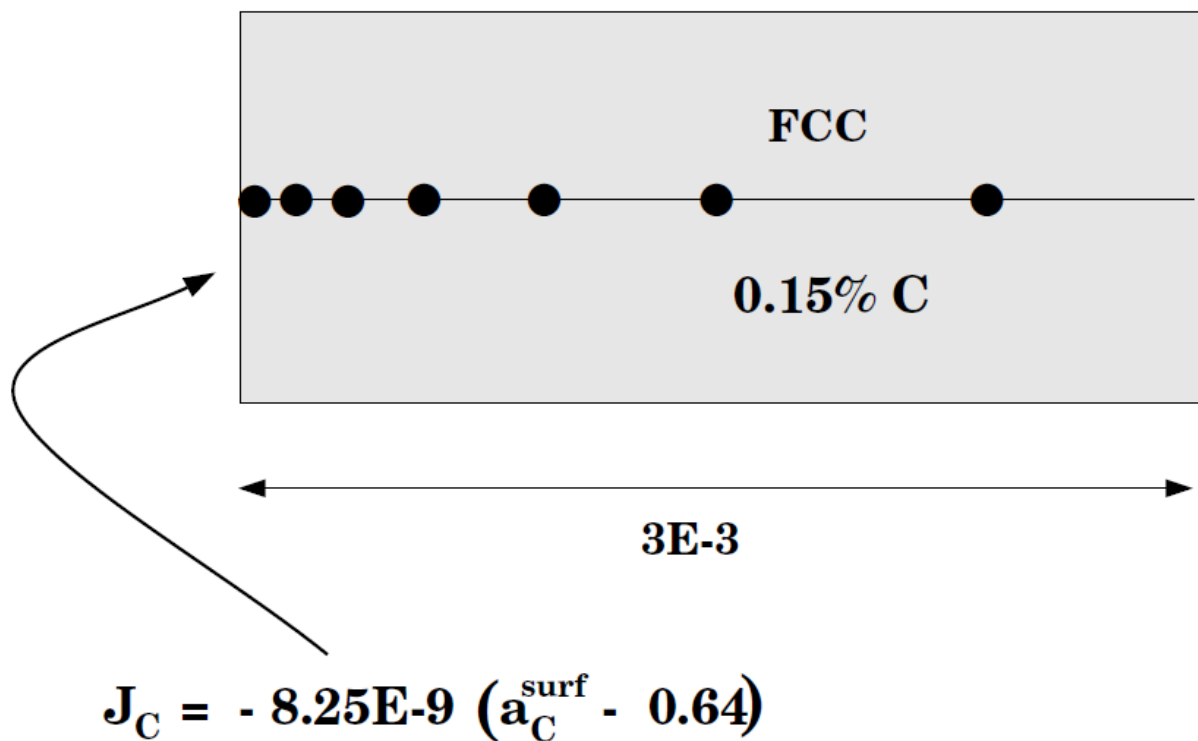


Example exa5

Carburization of a binary Fe-0.15 wt% C alloy: A surface reaction controls the flux of C at the surface

A mixture of 40% N₂ and 60% cracked methanol is used as carrier gas. The carburizing "carbon potential" in the gas is 0.85 wt%.

$$T = 1173 \text{ K}$$



exa5-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa5\setup.DCM.test"
SYS: i>_@@
NO SUCH COMMAND, USE HELP
SYS: @@ One-phase problem.
SYS: @@ Carburization of a binary Fe-0.15 wt% C alloy.
SYS: @@ A mixture of 40% N2 and 60% cracked methanol is used as carrier gas.
SYS: @@ The carburizing "carbon potential" in the gas is 0.85 wt%.
SYS: @@ A surface reaction controls the flux of C at the surface.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exa5_setup.DCM
SYS:
SYS: @@
SYS: @@ GO TO THE DATABASES AND READ THE THERMODYNAMIC AND KINETIC DATA
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: sw FEDEMO
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-sys fe,c
FE C DEFINED
TDB_FEDEMO: rej-ph *
GAS:G LIQUID:L BCC_A2
LAVES_PHASE_C14 CBCC_A12 CEMENTITE
CUB_A13 DIAMOND_FCC_A4 FCC_A1
GRAPHITE HCP_A3 KSI_CARBIDE
M23C6 M5C2 M7C3
REJECTED
TDB_FEDEMO: rest-ph fcc graphite
FCC_A1 GRAPHITE RESTORED
TDB_FEDEMO: get
09:40:54,298 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES
ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
'B. Uhrenius, Int. J. Refract. Met. Hard Mater. 12 (1994) 121 -127; Molar
volumes'
-OK-
TDB_FEDEMO:@?
TDB_FEDEMO: append
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2

TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
SNOB1 = SGTE Noble Metal Alloys Database v1.2
STBC2 = SGTE Thermal Barrier Coating TDB v2.2
STBC1 = SGTE Thermal Barrier Coating TDB v1.1
SALT1 = SGTE Molten Salts Database v1.3
SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2 = TC Semi-Conductors v2.1
SLAG4 = Fe-containing Slag v4.1
SLAG3 = Fe-containing Slag v3.2
SLAG2 = Fe-containing Slag v2.2
SLAG1 = Fe-containing Slag v1.2
TCOX11 = Metal Oxide Solutions v11.0
TCOX10 = Metal Oxide Solutions v10.1
TCOX9 = Metal Oxide Solutions v9.0
TCOX8 = Metal Oxide Solutions v8.0
TCOX7 = Metal Oxide Solutions v7.0
TCOX6 = Metal Oxide Solutions v6.0
TCOX5 = Metal Oxide Solutions v5.1
TCOX4 = Metal Oxide Solutions v4.1
ION3 = Ionic Solutions v3.0
ION2 = Ionic Solutions v2.6
ION1 = Ionic Solutions v1.5
NOX2 = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4 = Solder Alloys v4.0
TCSLD3 = Solder Alloys v3.3
TCSLD2 = Solder Alloys v2.0
TCSLD1 = Solder Alloys v1.1
TCSI1 = Ultrapure Silicon v1.2
TCMP2 = Materials Processing v2.5
TCES1 = Combustion/Sintering v1.1
TCSC1 = Super Conductor v1.0
TCFC1 = SOFC Database v1.0
TCNF2 = Nuclear Fuels v2.1b
NUMT2 = Nuclear Materials v2.1
NUOX4 = Nuclear Oxides v4.2
NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephista-19
MEPH15 = IRSN Mephista-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQ52 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SDDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.2
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3
MOBFE1 = Steels/Fe-Alloys Mobility v1.1
MOBFE2 = Steels/Fe-Alloys Mobility v2.0
MOBFE3 = Steels/Fe-Alloys Mobility v3.0
MOBFE4 = Steels/Fe-Alloys Mobility v4.0
MOBFE5 = Steels/Fe-Alloys Mobility v5.0
MOBFE6 = Steels/Fe-Alloys Mobility v6.0
MOBN15 = Ni-Alloys Mobility v5.1
MOBN14 = Ni-Alloys Mobility v4.1
MOBN13 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBS11 = Si-Alloys Mobility v1.0
MOBSDL1 = Solder-Alloys Mobility v1.1
MOBTI4 = Ti-Alloys Mobility v4.0
MOBTI3 = Ti-Alloys Mobility v3.1
MOBTI2 = Ti-Alloys Mobility v2.0
MOBTI1 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0

```

MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER     = User defined Database

DATABASE NAME /FEDEMO/: mfedemo
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys fe,c
FE C DEFINED
APP: rej-ph *
BCC_A2 FCC_A1 REJECTED
APP: rest-ph fcc
FCC_A1 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
-OK-
APP:@?
APP: @@
APP: @@ GO TO THE DICTRA MONITOR TO SET UP THE INITIAL STATE OF THE SPECIMEN
APP: @@
APP: go d-m
NO TIME STEP DEFINED
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC>
DIC> set-cond glob T 0 1173; * N
DIC>
DIC> @@
DIC> SELECT A REFERENCE STATE FOR THE C ACTIVITY
DIC> @@
DIC> set-ref-state
Component: c
Reference state: graph
Temperature /*/: *
Pressure /100000/: 1e5
DIC>
DIC> @@
DIC> ENTER A REGION, GRID, PHASE AND COMPOSITION
DIC> @@
DIC> enter-region
REGION NAME : steel
DIC>
DIC> enter-grid
REGION NAME : /STEEL/: steel
WIDTH OF REGION /1/: 3E-3
TYPE /LINEAR/: AUTO
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /STEEL/: steel
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC> enter-composition
REGION NAME : /STEEL/: steel
PHASE NAME: /FCC_A1/: fcc#1
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: c
TYPE /LINEAR/: lin
VALUE OF FIRST POINT : 0.15
VALUE OF LAST POINT : /0.15/: 0.15
DIC>
DIC> @@
DIC> @@ NOW SET THE BOUNDARY CONDITIONS. WE ARE ONLY INTERESTED IN THE
DIC> @@ SURFACE REGION, FOR EXAMPLE IT IS SUFFICIENT TO SET CONDITIONS AT THE
DIC> @@ LOWER BOUNDARY.
DIC> @@
DIC>
DIC> @@
DIC> @@ Specify the activity flux function which controls the uptake of C.
DIC> @@
DIC> @@ The functions f and g and the parameter N has to be specified.
DIC> @@ k k
DIC> @@
DIC> @@ N
DIC> @@ J V = f (variables)*(ACTIVITY -g (variables)) (1)
DIC> @@ k m k k
DIC> @@
DIC> @@ f and g in equation 1 is the mass-transfer coefficient and
DIC> @@ k k
DIC> @@ the activity of k in the gas, respectively. ACTIVITY in eq. 1 means
DIC> @@ the actual activity of species k at the surface.
DIC> @@
DIC>
DIC> @@
DIC> @@ The main carburizing reaction for our atmosphere is:
DIC> @@
DIC> @@ CO + H -> C + H O (I)
DIC> @@ 2 <- - 2
DIC> @@
DIC> @@ Following Sproge and Å...gren (J. Heat Treating, v6, no 1, 1988 pp. 9-19)
DIC> @@ we calculate the mass-transfer coefficient for carbon, f in
DIC> @@ eq. 1 above by means of eq. 3, 4 and 12 in Sproge and Å...gren's paper.
DIC> @@
DIC> @@
DIC> @@ A * K * P * sqrt( P )
DIC> @@ I CO H
DIC> @@
DIC> @@ f = ----- / gamma (2)
DIC> @@ a + B * K * P * sqrt( P )

```

```

DIC> @@ C I CO H
DIC> @@
DIC> @@
DIC> @@ K is the equilibrium constant for reaction (I)
DIC> @@
DIC> @@ A and B are constants defined in Sproge and Å...gren's paper. gamma
DIC> @@ is the activity coefficient for carbon in the steel.
DIC> @@
DIC> @@ Assume a constant value for P * sqrt( P ) = 0.14
DIC> @@ CO H
DIC> @@
DIC> @@ The carbon activity in the gas is controlled by the partial
DIC> @@ pressure of water as can be understood from reaction (I).
DIC> @@
DIC> @@ Assume that the carbon activity, a of the gas is 0.64
DIC> @@ C
DIC> @@ which corresponds to a carburizing "carbon potential" of 0.85 wt%.
DIC> @@
DIC> @@ In this way we may calculate f to 8.25E-9 mol/s.
DIC> @@
DIC>
DIC> set-cond
GLOBAL OR BOUNDARY CONDITION /GLOBAL/: bound
BOUNDARY /LOWER/: lower
CONDITION TYPE /CLOSED_SYSTEM/: activity_flux_function
ENTER THE EXPRESSION AS:
N
J_V = f (variables)*(ACTIVITY -g (variables))
k_m_k
FLUX OF FCC_A1,C
LOW TIME LIMIT /0/: 0
f(T,P,TIME)= -8.25E-9;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
N /1/: 1
LOW TIME LIMIT /0/: 0
g(T,P,TIME)= 0.64;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
DIC>
DIC> @@
DIC> @@ SPECIFY A SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 18000
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /1800/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO FILE
DIC> @@
DIC> Save exa5.Y
DIC>
DIC> set-inter
--OK--
DIC>

```

exa5-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa5\run.DCM.test"
DIC>
DIC>
DIC> @@ exa5_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE a5
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC> read exa5
OK
DIC>
DIC> @@
DIC> @@ Start the simulation
DIC> @@
DIC> sim
Region: STEEL
single geometric dense at 0.0000
1.2176 101
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: C = .00698495916383109 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
U-FRACTION IN SYSTEM: C = .00698495916383109 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916398349 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.24659766E-05 DT = 0.23659766E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916758908 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.71979297E-05 DT = 0.47319532E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495917479988 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.16661836E-04 DT = 0.94639063E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495918922068 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.35589649E-04 DT = 0.18927813E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495921806013 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.73445274E-04 DT = 0.37855625E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495927573319 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.14915652E-03 DT = 0.75711250E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495939106305 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.30057903E-03 DT = 0.15142250E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0069849596216772 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.60342403E-03 DT = 0.30284500E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0069849600827771 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.24204940E-02 DT = 0.12113800E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698496284726693 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.48432541E-02 DT = 0.24227600E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698496652968263 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.96887741E-02 DT = 0.48455200E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698497388635673 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.19379814E-01 DT = 0.96910401E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0069849885766828 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 1 seconds
TIME = 0.38761894E-01 DT = 0.19382080E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698501789240893 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.77526054E-01 DT = 0.38764160E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698507634097207 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.15505437 DT = 0.77528320E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698519272377668 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.31011102 DT = 0.15505664 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698542404641377 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.62022430 DT = 0.31011328 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698588265674895 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 1.2404509 DT = 0.62022656 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698678864751835 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
```

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TIME = 2.4809040 DT = 1.2404531 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698856957978377 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 4.9618102 DT = 2.4809063 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0069920463860113 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 9.9236228 DT = 4.9618125 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00699876996632461 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 1 seconds
TIME = 19.847248 DT = 9.9236250 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00701160601590707 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 39.694498 DT = 19.847250 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0070356937636148 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 79.388998 DT = 39.694500 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00707989462116312 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 158.77800 DT = 79.389000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00715874757109912 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 317.55600 DT = 158.77800 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00729474138750393 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 635.11200 DT = 317.55600 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00752054375503505 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 1270.2240 DT = 635.11200 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00788106518443646 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 1 seconds
TIME = 2540.4480 DT = 1270.2240 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00843611780537564 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 4340.4480 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00904710226999639 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 6140.4480 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00955134567260921 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 7940.4480 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00999113675764907 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 9740.4480 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0103863764716757 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 11540.448 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0107484090267321 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 1 seconds
TIME = 13340.448 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0110844421240418 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 15140.448 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0113993948509824 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 16940.448 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0116968050295672 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 18000.000 DT = 1059.5520 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0118647499013374 FE = 1
TOTAL SIZE OF SYSTEM: .003 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 0.24659766E-05
DELETING TIME-RECORD FOR TIME 0.71979297E-05
DELETING TIME-RECORD FOR TIME 0.16661836E-04
DELETING TIME-RECORD FOR TIME 0.35589649E-04
DELETING TIME-RECORD FOR TIME 0.73445274E-04
DELETING TIME-RECORD FOR TIME 0.14915652E-03
DELETING TIME-RECORD FOR TIME 0.30057903E-03
DELETING TIME-RECORD FOR TIME 0.60342403E-03
DELETING TIME-RECORD FOR TIME 0.12091140E-02
DELETING TIME-RECORD FOR TIME 0.24204940E-02
DELETING TIME-RECORD FOR TIME 0.48432541E-02
DELETING TIME-RECORD FOR TIME 0.96887741E-02
DELETING TIME-RECORD FOR TIME 0.19379814E-01
DELETING TIME-RECORD FOR TIME 0.38761894E-01
DELETING TIME-RECORD FOR TIME 0.77526054E-01
DELETING TIME-RECORD FOR TIME 0.15505437
DELETING TIME-RECORD FOR TIME 0.31011102
DELETING TIME-RECORD FOR TIME 0.62022430
DELETING TIME-RECORD FOR TIME 1.2404509
DELETING TIME-RECORD FOR TIME 2.4809040
DELETING TIME-RECORD FOR TIME 4.9618102
DELETING TIME-RECORD FOR TIME 9.9236228
DELETING TIME-RECORD FOR TIME 19.847248
DELETING TIME-RECORD FOR TIME 39.694498
DELETING TIME-RECORD FOR TIME 79.388998
DELETING TIME-RECORD FOR TIME 158.77800
DELETING TIME-RECORD FOR TIME 317.55600
DELETING TIME-RECORD FOR TIME 635.11200
DELETING TIME-RECORD FOR TIME 1270.2240
DELETING TIME-RECORD FOR TIME 2540.4480
DELETING TIME-RECORD FOR TIME 4340.4480
DELETING TIME-RECORD FOR TIME 6140.4480

```

DELETING TIME-RECORD FOR TIME 7940.4480
DELETING TIME-RECORD FOR TIME 9740.4480
DELETING TIME-RECORD FOR TIME 11540.448
DELETING TIME-RECORD FOR TIME 13340.448
DELETING TIME-RECORD FOR TIME 15140.448

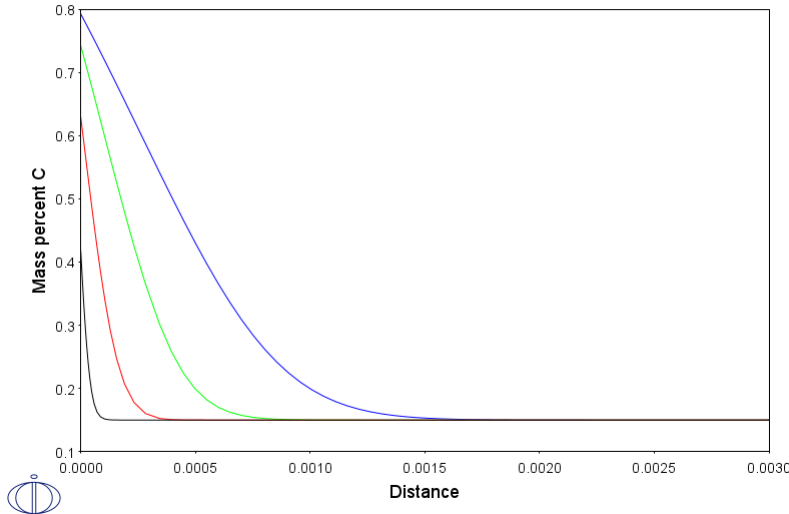
KEEPING TIME-RECORD FOR TIME 16940.448
AND FOR TIME 18000.000
WORKSPACE RECLAIMED

TIMESTEP AT 18000.0000 SELECTED

DIC>
DIC> set-inter
--OK--
DIC>

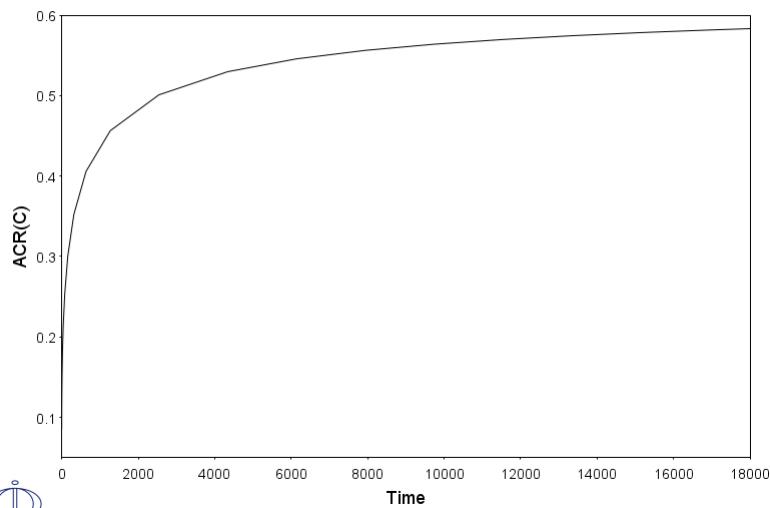
exa5-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa5\plot.DCM.test"
DIC>
DIC>
DIC> @@ exa5_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE a5
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 1.80000E+04
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC> read exa5
OK
DIC>
DIC> @@
DIC> @@ ENTER THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ PLOT SOME DIFFERENT CONCENTRATION PROFILES
POST-1: @@
POST-1: s-d-a x dist glo
INFO: Distance is set as independent variable
POST-1: s-d-a y w-p c
POST-1: s-p-c time 100 1000 5000 18000
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.09.42.54
Time = 100,1000,5000,18000
CELL #1
```

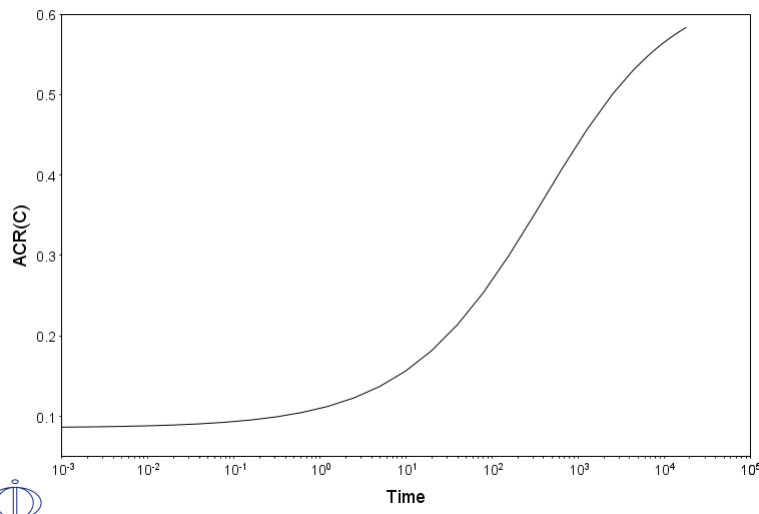


```
POST-1:
POST-1:
POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1:
POST-1: @@
POST-1: @@ PLOT THE VARIATION OF THE C ACTIVITY AT THE SURFACE
POST-1: @@
POST-1: s-d-a y acr(c)
POST-1:
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1:
POST-1: s-p-c
CONDITION /TIME/: interface
INTERFACE : first
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

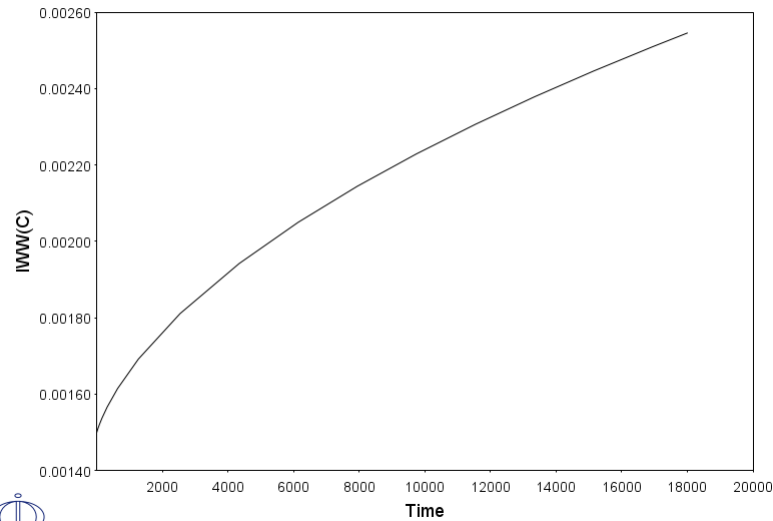
2021.05.13.09.42.58
"FIRST" INTERFACE OF SYSTEM
CELL #1



POST-1:
POST-1:
POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1:
POST-1: @@
POST-1: @@ USE A LOGARITHMIC SCALE ON THE X-AXIS
POST-1: @@
POST-1: set-axis-type
AXIS (X, Y OR Z) : x
AXIS TYPE /LINEAR/: logarithmic
POST-1:
POST-1: s-s-s x n 0.001 2e4
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.09.43.00
"FIRST" INTERFACE OF SYSTEM
CELL #1



POST-1:
POST-1:
POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1:
POST-1: @@
POST-1: @@ PLOT THE AVERAGE WEIGHT FRACTION OF C IN THE SPECIMEN
POST-1: @@
POST-1: s-d-a y iww(c)
POST-1:
POST-1: set-ax-ty
AXIS (X, Y OR Z) : x
AXIS TYPE /LINEAR/: linear
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot



```
POST-1:  
POST-1:  
POST-1:  
POST-1:@?<Hit_return_to_continue>  
POST-1:  
POST-1:  
POST-1:  
POST-1: set-inter  
--OK--  
POST-1:
```

exa6-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa6\setup.DCM.test"
SYS: @@
SYS: @@ One-phase problem.
SYS: @@ Diffusion through a tube wall.
SYS: @@ A simple example about diffusion through a tube wall.
SYS: @@ The tube material is an Fe-0.6%Mn-0.7%Si-0.05%C alloy. On
SYS: @@ the inside wall a carbon activity of 0.9 is maintained whereas on
SYS: @@ the outside the C-activity is very low. This example demonstrates
SYS: @@ the use of the command SET-FIRST-INTERFACE as well as the MIXED
SYS: @@ boundary conditions.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ setup.DCM
SYS:
SYS:
SYS: @@
SYS: @@ GO TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: @@
TDB_TCFE11: @@ USE THE TCFE DATABASE FOR THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw tcfef9
Current database: Steels/Fe-Alloys v9.3

VA           /- DEFINED
L12_FCC      B2_BCC          DICTRA_FCC_A1
REJECTED
TDB_TCFE9: def-sys fe si mn c
FE           SI              MN
C DEFINED
TDB_TCFE9: rej ph * all
GAS:G         LIQUID:L        BCC_A2
FCC_A1        HCP_A3          CBCC_A12
CUB_A13       DIAMOND_FCC_A4 GRAPHITE
CEMENTITE     M23C6           M7C3
M5C2          KSI_CARBIDE    FE4N_LP1
FEFCN_CHI     LAVES_PHASE_C14 M3SI
MNN9SI2       MN11SI19       MN6SI
G_PHASE       CR3SI          FE2SI
FESI2_H       FESI2_L        MSI
M5SSI3        AL4C3          FE8SI2C
SIC           MN5SIC         CUZN_EPSILON
AL5FE4        MP_B31        M2P_C22
FLUORITE_C1:I ZRO2_TETR:I   M2O3C:I
M2O3H:I      REJECTED
TDB_TCFE9: res ph fcc,grap
FCC_A1        GRAPHITE RESTORED
TDB_TCFE9: get
09:44:19,861 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
FUNCTIONS .....

List of references for assessed data

'A. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'B. Uhrenius, Int. J. Refract. Met. Hard Mater. 12 (1994) 121-127; Molar
  volumes'
'X.-G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
  volumes'
'X.-G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'B. Hallstedt, unpublished work (2016); C-Fe-Mn Epsilon martensite.'
'D. Djurovic, B. Hallstedt, J. von Appen, R. Dronskowsk, CALPHAD, 34, 279
  -85(2010); Mn-C'
'J. Lacaze and B. Sundman, Metall. Mater. Trans. A, 22A (1991) 2211-2223;
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'W. Huang, CALPHAD, 13 (1989) 243-252; TRITA-MAC 388 (rev 1989); FE-MN'
'W. Zheng, J. Alloys Compd., 632 (2015) 661-675'
'D. Djurovic, B. Hallstedt, J. von Appen, R. Dronskowsk, CALPHAD,
  submitted, 2011; Fe-Mn-C'
'J. Miettinen and B. Hallstedt, CALPHAD, 22 (1998) 231-256; Fe-Si and Fe
  -Si-C'
'A. Forsberg and J. Agren, J. Phase Equilib., 14 (1993) 354-363; Fe-Mn-Si'
-OK-
TDB_TCFE9: @@
TDB_TCFE9: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE KINETIC DATA
TDB_TCFE9: @@
TDB_TCFE9: app
Use one of these databases
TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
```

```

TCFE9   = Steels/Fe-Alloys v9.3
TCFE8   = Steels/Fe-Alloys v8.2
TCFE7   = Steels/Fe-Alloys v7.0
TCFE6   = Steels/Fe-Alloys v6.2
TCFE5   = Steels/Fe-Alloys v5.0
TCFE4   = Steels/Fe-Alloys v4.1
TCFE3   = Steels/Fe-Alloys v3.1
TCFE2   = Steels/Fe-Alloys v2.1
TCFE1   = Steels/Fe-Alloys v1.0
FEDAT   = TCS/TT Steels Database v1.0
TCNI11  = Ni-Alloys v11.0
TCNI10  = Ni-Alloys v10.0
TCNI9   = Ni-Alloys v9.1
TCNI8   = Ni-Alloys v8.2
TCNI7   = Ni-Alloys v7.2
TCNI6   = Ni-Alloys v6.1
TCNI5   = Ni-Alloys v5.1
TCNI4   = Ni-Alloys v4.0
TCNI1   = Ni-Alloys v1.3
TCAL8   = Al-Alloys v8.0
TCAL7   = Al-Alloys v7.1
TCAL6   = Al-Alloys v6.0
TCAL5   = Al-Alloys v5.1
TCAL4   = Al-Alloys v4.0
TCAL3   = Al-Alloys v3.0
TCAL2   = Al-Alloys v2.1
TCAL1   = Al-Alloys v1.2
TCMG6   = Mg-Alloys v6.1
TCMG5   = Mg-Alloys v5.1
TCMG4   = Mg-Alloys v4.0
TCMG3   = Mg-Alloys v3.0
TCMG2   = Mg-Alloys v2.0
TCMG1   = Mg-Alloys v1.1
TCTI3   = Ti-Alloys v3.0
TCTI2   = Ti-Alloys v2.2
TCTI1   = Ti-Alloys v1.0
TCCU4   = Cu-Alloys v4.0
TCCU3   = Cu-Alloys v3.1
TCCU2   = Cu-Alloys v2.0
TCCU1   = Cu-Alloys v1.0
TCCC1   = Cemented carbide v1.0
TCHEA5  = High Entropy Alloy v5.0
TCHEA4  = High Entropy Alloy v4.2
TCHEA3  = High Entropy Alloy v3.1
TCHEA2  = High Entropy Alloy v2.1.1
TCHEA1  = High Entropy Alloy v1.0
SSOL7   = SGTE Alloy Solutions Database v7.0
SSOL6   = SGTE Alloy Solutions Database v6.0
SSOL5   = SGTE Alloy Solutions Database v5.0
SSOL4   = SGTE Alloy Solutions Database v4.9g
SSOL2   = SGTE Alloy Solutions Database v2.1
SSUB6   = SGTE Substances Database v6.0
SSUB5   = SGTE Substances Database v5.2
SSUB4   = SGTE Substances Database v4.1
SSUB3   = SGTE Substances Database v3.3
SSUB2   = SGTE Substances Database v2.2
SNOB3   = SGTE Noble Metal Alloys Database v3.1
SNOB2   = SGTE Noble Metal Alloys Database v2.1
SNOB1   = SGTE Noble Metal Alloys Database v1.2
STBC2   = SGTE Thermal Barrier Coating TDB v2.2
STBC1   = SGTE Thermal Barrier Coating TDB v1.1
SALT1   = SGTE Molten Salts Database v1.3
SNUX6   = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2   = TC Semi-Conductors v2.1
SLAG4   = Fe-containing Slag v4.1
SLAG3   = Fe-containing Slag v3.2
SLAG2   = Fe-containing Slag v2.2
SLAG1   = Fe-containing Slag v1.2
TCOX11  = Metal Oxide Solutions v11.0
TCOX10  = Metal Oxide Solutions v10.1
TCOX9   = Metal Oxide Solutions v9.0
TCOX8   = Metal Oxide Solutions v8.0
TCOX7   = Metal Oxide Solutions v7.0
TCOX6   = Metal Oxide Solutions v6.0
TCOX5   = Metal Oxide Solutions v5.1
TCOX4   = Metal Oxide Solutions v4.1
ION3    = Ionic Solutions v3.0
ION2    = Ionic Solutions v2.6
ION1    = Ionic Solutions v1.5
NOX2   = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4  = Solder Alloys v4.0
TCSLD3  = Solder Alloys v3.3
TCSLD2  = Solder Alloys v2.0
TCSLD1  = Solder Alloys v1.1
TCSI1   = Ultrapure Silicon v1.2
TCMP2   = Materials Processing v2.5
TCES1   = Combustion/Sintering v1.1
TCS1   = Super Conductor v1.0
TCFC1   = SOFC Database v1.0
TCNF2   = Nuclear Fuels v2.1b
NUMT2   = Nuclear Materials v2.1
NUOX4   = Nuclear Oxides v4.2
NUTO1   = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1   = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19  = IRSN NUCLEA-19
NUCL15  = IRSN NUCLEA-15_4
NUCL10  = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19  = IRSN Mephista-19
MEPH15  = IRSN Mephista-15_1
MEPH11  = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3   = Aqueous Solution v3.0
TCAQ2   = Aqueous Solution v2.7
AQ52   = TGG Aqueous Solution Database v2.6
GCE2   = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SLDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2   = Public Aqueous Soln (SIT) TDB v2.5
PG35   = PG35 Binary Semi-Conductors TDB v1.3
PURE5  = SGTE Unary (Pure Elements) TDB v5.2

```

```

MOB2      = Alloys Mobility v2.7
MOB1      = Alloys Mobility v1.3
MOBFE1   = Steels/Fe-Alloys Mobility v1.1
MOBFE2   = Steels/Fe-Alloys Mobility v2.0
MOBFE3   = Steels/Fe-Alloys Mobility v3.0
MOBFE4   = Steels/Fe-Alloys Mobility v4.0
MOBFE5   = Steels/Fe-Alloys Mobility v5.0
MOBFE6   = Steels/Fe-Alloys Mobility v6.0
MOBN15   = Ni-Alloys Mobility v5.1
MOBN14   = Ni-Alloys Mobility v4.1
MOBN13   = Ni-Alloys Mobility v3.2
MOBN12   = Ni-Alloys Mobility v2.4
MOBN11   = Ni-Alloys Mobility v1.10
MOBAL6   = Al-Alloys Mobility v6.0
MOBAL5   = Al-Alloys Mobility v5.0
MOBAL4   = Al-Alloys Mobility v4.0
MOBAL3   = Al-Alloys Mobility v3.0
MOBAL2   = Al-Alloys Mobility v2.0
MOBAL1   = Al-Alloys Mobility v1.0
MOBCU1   = Cu-Alloys Mobility v1.0
MOBCU2   = Cu-Alloys Mobility v2.0
MOBCU3   = Cu-Alloys Mobility v3.0
MOBCU4   = Cu-Alloys Mobility v4.0
MOBHEA1  = High Entropy Alloys Mobility v1.0
MOBHEA2  = High Entropy Alloys Mobility v2.0
MOBMG2   = Mg-Alloys Mobility v2.0
MOBMG1   = Mg-Alloys Mobility v1.0
MOBSI1   = Si-Alloys Mobility v1.0
MOBSLD1  = Solder-Alloys Mobility v1.1
MOBT14   = Ti-Alloys Mobility v4.0
MOBT13   = Ti-Alloys Mobility v3.1
MOBT12   = Ti-Alloys Mobility v2.0
MOBT11   = Ti-Alloys Mobility v1.0
MALDEMO  = Al-Alloys demo database v2.0
MFEDEMO  = Fe-Alloys Mobility demo database v2.0
MNIDEMO  = Ni-Alloys Mobility demo database v1.0
MCUDEMO  = Cu-Alloys Mobility demo database v1.0
USER     = User defined Database

```

DATABASE NAME /TCFE9: mobfe4
 Current database: Steels/Fe-Alloys Mobility v4.0

```

VA DEFINED
B2_BCC REJECTED
APP: def-sys fe si mn c
  FE                      SI                      MN
  C DEFINED
APP: rej ph * all
  BCC_A2                  CEMENTITE             FCC_A1
  FE4N_LP1                HCP_A3                 LIQUID:L
  REJECTED
APP: res ph fcc
  FCC_A1 RESTORED
APP: get
  ELEMENTS .....
  SPECIES .....
  PHASES .....
  PARAMETERS .....
  FUNCTIONS .....

```

List of references for assessed data

```

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'Y. Liu, L. Zhang, et al. CALPHAD 33(2009)614-23; Fe-Mn-C (fcc)'
'M. Yin, et al. (2010) unpublished work.'
'W. Zheng, et al., J. Alloys and Compounds, 632 (2015) 661-675; Fe-Mn-Si
(fcc)'
'D. Bergner et al., Defect and Diffusion Forum 66-69(1989)409. Impurity
diffusion of Si in fcc Fe.'

```

-OK-

```

APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR WHERE THE PROBLEM IS SET UP
APP: @@
APP: go d-m

```

```

NO TIME STEP DEFINED
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE

```

```

DIC>
DIC> @@
DIC> @@ ENTER GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 1273; * N
DIC>

```

```

DIC> @@
DIC> @@ SET THE REFERENCE STATE FOR CARBON
DIC> @@
DIC> set-ref C grap * 101325
DIC>

```

```

DIC> @@
DIC> @@ ENTER A REGION
DIC> @@
DIC> enter-region aus
DIC>

```

```

DIC> @@
DIC> @@ ENTER A DOUBLE GEOMETRIC GRID INTO THE REGION
DIC> @@
DIC> enter-grid

```

```

REGION NAME : /AUS/: aus

```

```

WIDTH OF REGION /1/: 1e-2

```

```

TYPE /LINEAR/: AUTO

```

```

DIC>
DIC> @@
DIC> @@ SET THE GEOMETRY (1 = CYLINDER)
DIC> @@
DIC> enter-geo

```

```

GEOMETRICAL EXPONENT /0/: 1

```

```

DIC>

```

```

DIC> @@
DIC> @@ SET THE FIRST INTERFACE => TUBE

```

```

DIC> @@

```

```

DIC> set-first-interface
COORDINATE FOR FIRST INTERFACE /0/: 2e-2
DIC>
DIC> @@
DIC> @@ ENTER AN active PHASE IN THE REGION
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUS/: aus
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc_a1#1
DIC>
DIC> @@
DIC> @@ ENTER INITIAL COMPOSITIONS INTO THE PHASE
DIC> @@
DIC> enter-composition
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1/: fcc#1
DEPENDENT COMPONENT ? /SI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: si lin 0.7 0.7
PROFILE FOR /MN/: mn lin 0.6 0.6
PROFILE FOR /SI/: c lin 5e-2 5e-2
DIC>
DIC> @@
DIC> @@ SET THE BOUNDARY CONDITIONS ON BOTH THE LOWER AND UPPER PART OF THE REGION
DIC> @@
DIC> @@ USE MIXED CONDITIONS: AN ACTIVITY CONDITION FOR C AND CLOSED
DIC> @@ SYSTEMS FOR MN AND SI.
DIC> @@
DIC> set-cond
GLOBAL OR BOUNDARY CONDITION /GLOBAL/: boundary
BOUNDARY /LOWER/: lower
CONDITION TYPE /CLOSED_SYSTEM/: mixed
Dependent substitutional element:FE
Dependent interstitial element:VA
TYPE OF CONDITION FOR COMPONENT C /ZERO_FLUX/: activity
LOW TIME LIMIT /0/: 0
ACR(C) (TIME)= 0.9;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
TYPE OF CONDITION FOR COMPONENT MN /ZERO_FLUX/: zero_flux
TYPE OF CONDITION FOR COMPONENT SI /ZERO_FLUX/: zero_flux
DIC>
DIC> set-cond
GLOBAL OR BOUNDARY CONDITION /GLOBAL/: boundary
BOUNDARY /UPPER/: upper
CONDITION TYPE /CLOSED_SYSTEM/: mixed
Dependent substitutional element:FE
Dependent interstitial element:VA
TYPE OF CONDITION FOR COMPONENT C /ZERO_FLUX/: activity
LOW TIME LIMIT /0/: 0
ACR(C) (TIME)= 1e-5;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
TYPE OF CONDITION FOR COMPONENT MN /ZERO_FLUX/: zero_flux
TYPE OF CONDITION FOR COMPONENT SI /ZERO_FLUX/: zero_flux
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 1e9
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /100000000/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC>
DIC> @@
DIC> @@ SAVE TO FILE
DIC> @@
DIC> save exa6 y
DIC>
DIC> set-inter
--OK--
DIC>

```

exa6-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa6\run.DCM.test"
DIC>
DIC>
DIC> @@
DIC> FILE FOR RUNNING exa6
DIC> @@
DIC>
DIC> @@
DIC> ENTER THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC> read exa6
OK
DIC>
DIC> @@
DIC> Start the simulation
DIC> @@
DIC> simulate
Region: AUS
double geometric
dense at outer boundaries, coarse at 0.50000E-02
lower part 1.2500 22
upper part 0.80000 22
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: C = .00115488575879621 FE = .490055682684517
MN = .00302988813183616 SI = .00691442924890042
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
U-FRACTION IN SYSTEM: C = .00115488575879621 FE = .490055682684517
MN = .00302988813183616 SI = .00691442924890042
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00116822817821891 FE = .490055682684517
MN = .00302988813183616 SI = .00691442924890042
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 0.10010000E-03 DT = 0.1000000E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00116829188694328 FE = .490055682684517
MN = .00302988813183616 SI = .00691442924890042
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 3.0557610 DT = 3.0556609 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00118959189435653 FE = .490055682684521
MN = .00302988813183664 SI = .00691442924889593
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 9.1670828 DT = 6.1113218 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00121111672154579 FE = .490055682684524
MN = .00302988813183702 SI = .00691442924889251
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 21.389726 DT = 12.222644 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00123890626455181 FE = .490055682684528
MN = .00302988813183746 SI = .00691442924888853
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 45.835013 DT = 24.445287 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00127670303015352 FE = .490055682684532
MN = .00302988813183797 SI = .00691442924888392
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 94.725588 DT = 48.890574 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00132921753226138 FE = .490055682684536
MN = .00302988813183855 SI = .00691442924887889
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 192.50674 DT = 97.781148 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00140291029963708 FE = .49005568268454
MN = .00302988813183909 SI = .00691442924887446
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 388.06903 DT = 195.56230 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00150683893865951 FE = .490055682684541
MN = .00302988813183934 SI = .00691442924887331
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 779.19363 DT = 391.12459 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00165382101622263 FE = .490055682684533
MN = .00302988813183861 SI = .00691442924888217
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 1561.4428 DT = 782.24919 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00186208912002595 FE = .49005568268452
MN = .00302988813183733 SI = .00691442924889648
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 3125.9412 DT = 1564.4984 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00215763950853838 FE = .490055682684545
MN = .00302988813184034 SI = .00691442924886802
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 6254.9379 DT = 3128.9967 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00257762247234064 FE = .490055682684614
MN = .00302988813184873 SI = .00691442924879132
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 12512.931 DT = 6257.9935 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00317612553687887 FE = .490055682684701
MN = .00302988813186053 SI = .00691442924869196
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 25028.918 DT = 12515.987 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00403212708748458 FE = .490055682684788
MN = .00302988813187388 SI = .00691442924859185
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 50060.892 DT = 25031.974 SUM OF SQUARES = 0.0000000
```

```

U-FRACTION IN SYSTEM: C = .00526185946385771 FE = .490055682684867
MN = .00302988813188714 SI = .00691442924849985
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 100124.84 DT = 50063.948 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00703878462392717 FE = .490055682684956
MN = .00302988813189985 SI = .00691442924839735
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 200252.74 DT = 100127.90 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00961468135205799 FE = .490055682685084
MN = .0030298881319146 SI = .00691442924825484
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds

```

output ignored...

```

... output resumed

CPU time used in timestep 1 seconds
TIME = 1602043.3 DT = 801023.17 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .016516584267494 FE = .490055682687091
MN = .00302988813214933 SI = .00691442924601361
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 307117.5 DT = 1469074.2 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165253458723486 FE = .490055682689399
MN = .00302988813240262 SI = .00691442924345169
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 5521426.5 DT = 2450309.1 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165254570895416 FE = .490055682693946
MN = .00302988813287881 SI = .00691442923842824
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 9715746.1 DT = 4194319.5 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165253215591159 FE = .490055682703382
MN = .00302988813383215 SI = .00691442922803932
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 17813876. DT = 8098130.0 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165250052909427 FE = .490055682726195
MN = .00302988813607049 SI = .00691442920298776
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 34010136. DT = 16196260. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165246276464085 FE = .490055682784398
MN = .00302988814169422 SI = .00691442913916063
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 66402656. DT = 32392520. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165243580713323 FE = .490055682920989
MN = .00302988815540701 SI = .00691442898885753
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 0.13118770E+09 DT = 64785040. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165244582014679 FE = .490055683192306
MN = .00302988818558275 SI = .00691442868736454
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 0.23061023E+09 DT = 99422537. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165263917143557 FE = .490055683555767
MN = .00302988823056496 SI = .00691442827892098
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 0.33012543E+09 DT = 99515198. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165303267851654 FE = .49005568386721
MN = .00302988827178368 SI = .0069144279262599
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 0.42970616E+09 DT = 99580728. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165336326908907 FE = .490055684142115
MN = .00302988830951066 SI = .00691442761362775
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 0.52934108E+09 DT = 99634921. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165366300962674 FE = .490055684390274
MN = .0030298883443561 SI = .00691442733062356
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 0.62902222E+09 DT = 99681143. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .016539438424926 FE = .490055684617852
MN = .00302988837682077 SI = .00691442707058051
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 0.72874408E+09 DT = 99721854. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165421143527838 FE = .490055684829008
MN = .00302988840729343 SI = .0069144268289518
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 0.82850228E+09 DT = 99758206. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .016544689475698 FE = .490055685026688
MN = .00302988843607474 SI = .0069144266024909
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 0.92829328E+09 DT = 99791002. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165471833610484 FE = .490055685213062
MN = .00302988846340022 SI = .00691442638879089
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 1 seconds
TIME = 0.98979354E+09 DT = 61500257. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165498694311618 FE = .490055685322987
MN = .00302988847959189 SI = .006914426267451
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
CPU time used in timestep 0 seconds
TIME = 0.10000000E+10 DT = 10206459. SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0165516055116173 FE = .490055685340902
MN = .00302988848223566 SI = .0069144262421159
TOTAL SIZE OF SYSTEM: .00314159265359 [m^2]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000

```

```
DELETING TIME-RECORD FOR TIME 0.1000000E-06
DELETING TIME-RECORD FOR TIME 0.1001000E-03
DELETING TIME-RECORD FOR TIME 3.0557610
DELETING TIME-RECORD FOR TIME 9.1670828
DELETING TIME-RECORD FOR TIME 21.389726
DELETING TIME-RECORD FOR TIME 45.835013
DELETING TIME-RECORD FOR TIME 94.725588
DELETING TIME-RECORD FOR TIME 192.50674
DELETING TIME-RECORD FOR TIME 388.06903
DELETING TIME-RECORD FOR TIME 779.19363
DELETING TIME-RECORD FOR TIME 1561.4428
DELETING TIME-RECORD FOR TIME 3125.9412
DELETING TIME-RECORD FOR TIME 6254.9379
DELETING TIME-RECORD FOR TIME 12512.931
DELETING TIME-RECORD FOR TIME 25028.918
DELETING TIME-RECORD FOR TIME 50060.892
DELETING TIME-RECORD FOR TIME 100124.84
DELETING TIME-RECORD FOR TIME 200252.74
DELETING TIME-RECORD FOR TIME 400508.53
DELETING TIME-RECORD FOR TIME 801020.11
DELETING TIME-RECORD FOR TIME 1602043.3
DELETING TIME-RECORD FOR TIME 3071117.5
DELETING TIME-RECORD FOR TIME 5521426.5
DELETING TIME-RECORD FOR TIME 9715746.1
DELETING TIME-RECORD FOR TIME 17813876.
DELETING TIME-RECORD FOR TIME 34010136.
DELETING TIME-RECORD FOR TIME 66402656.
DELETING TIME-RECORD FOR TIME 0.13118770E+09
DELETING TIME-RECORD FOR TIME 0.23061023E+09
DELETING TIME-RECORD FOR TIME 0.33012543E+09
DELETING TIME-RECORD FOR TIME 0.42970616E+09
DELETING TIME-RECORD FOR TIME 0.52934108E+09
DELETING TIME-RECORD FOR TIME 0.62902222E+09
DELETING TIME-RECORD FOR TIME 0.72874408E+09
DELETING TIME-RECORD FOR TIME 0.82850228E+09
DELETING TIME-RECORD FOR TIME 0.92829328E+09
```

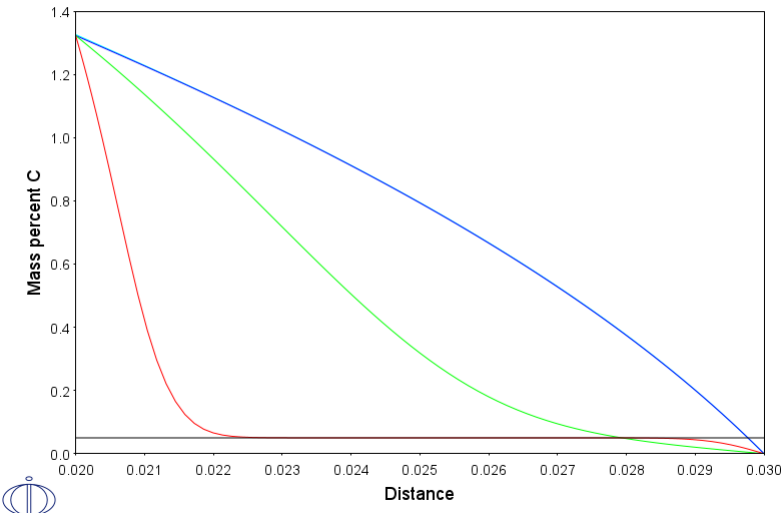
```
KEEPING TIME-RECORD FOR TIME 0.98979354E+09
AND FOR TIME 0.10000000E+10
WORKSPACE RECLAIMED
```

```
TIMESTEP AT 0.10000000E+10 SELECTED
```

```
DIC>
DIC> @@
DIC> @@ THE SIMULATION IS FINISHED
DIC> @@
DIC>
DIC> set-inter
--OK---
DIC>
```

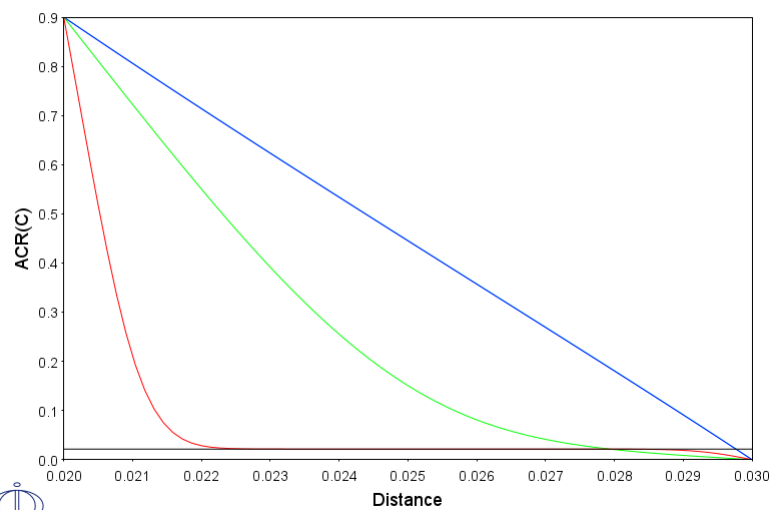
exa6-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa6\plot.DCM.test"
DIC>
DIC>
DIC> @@ exa6_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE exa6
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 1.00000E+09
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC> read exa6
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1:
POST-1: @@
POST-1: @@ PLOT THE CONCENTRATION OF C AT DIFFERENT TIMES
POST-1: @@
POST-1: s-d-a x distance global
INFO: Distance is set as independent variable
POST-1: s-d-a y w-p c
POST-1: s-p-c time 0,1e4,2e5,1e7,1e9
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.09.46.39
Time=0,10000,20000,1E+07,1E+09
CELL #1
```

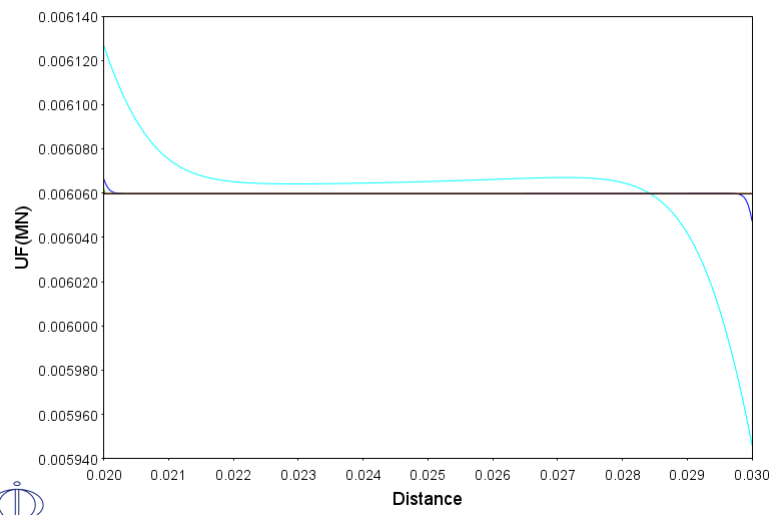


```
POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: @@
POST-1: @@ PLOT THE ACTIVITY OF C
POST-1: @@
POST-1:
POST-1: s-d-a y acr(c)
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

2021.05.13.09.46.49
Time = 0,10000,200000,1E+07,1E+09
CELL #1

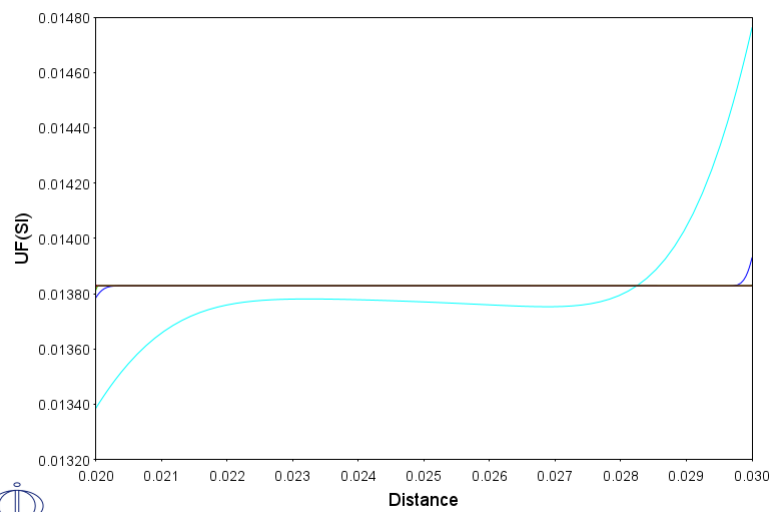


POST-1:
POST-1:
POST-1:
POST-1: Hit RETURN to continue
POST-1:
POST-1: @@
POST-1: @@ LET US LOOK AT THE MN AND SI PROFILES
POST-1: @@
POST-1: @@ WE PLOT THE U-FRACTION OF MN AND SI WHICH WILL BE INDEPENDENT
POST-1: @@ OF THE C-CONCENTRATION.
POST-1: @@
POST-1: s-d-a y u-f mn
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.09.46.51
Time = 0,10000,200000,1E+07,1E+09
CELL #1

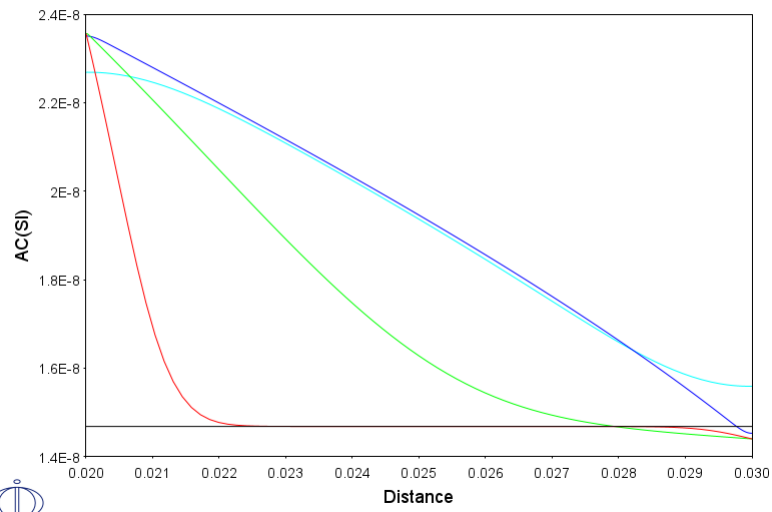


POST-1:
POST-1:
POST-1:
POST-1:
POST-1: Hit RETURN to continue
POST-1:
POST-1: s-d-a y u-f si
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

2021.05.13.09.46.54
Time = 0,10000,200000,1E+07,1E+09
CELL #1



```
POST-1:  
POST-1:  
POST-1:  
POST-1:  
POST-1:Hit RETURN to continue  
POST-1:  
POST-1: @@  
POST-1: @@ FINALLY, LOOK AT THE ACTIVITY PROFILES OF SI  
POST-1: @@  
POST-1: s-d-a y ac(si)  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
POST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot  
2021.05.13.09.47.01  
Time = 0,10000,200000,1E+07,1E+09  
CELL #1
```



```
POST-1:  
POST-1:  
POST-1:  
POST-1:  
POST-1:  
POST-1:Hit RETURN to continue  
POST-1:  
POST-1: set-inter  
--OK--  
POST-1:
```

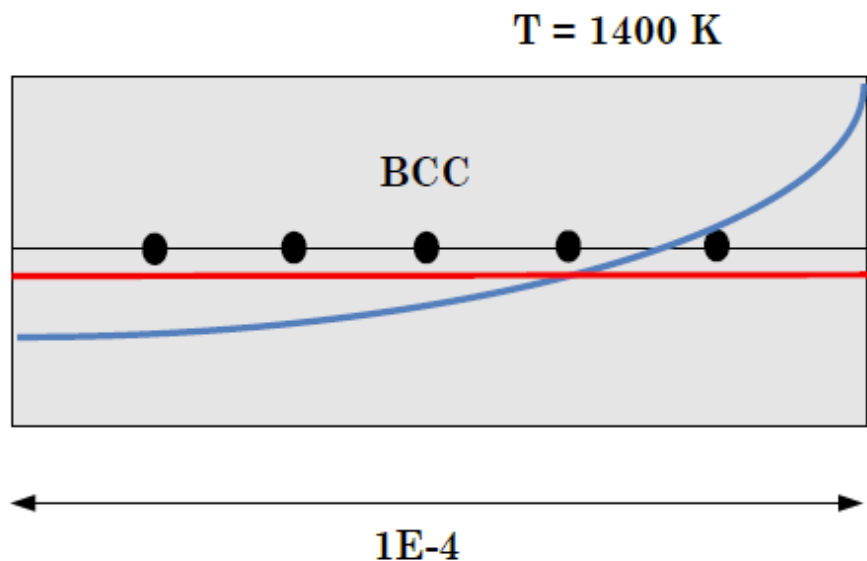


Example exa7

Homogenization heat-treatment

(Initial profile imported from Scheil simulation)

The initial segregation profile is created from a Scheil calculation (see macro `create_initial_profile.TCM`). The command `INPUT_SCHEIL_PROFILE` in the DICTRA monitor performs most of the setup. Only time and temperature must be entered after the `INPUT_SCHEIL_PROFILE` command is executed.



exa7-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa7\setup.DCM.test"
SYS: @@ One phase example.
SYS: @@ Homogenization heat treatment
SYS: @@ The initial segregation profile is created from a Scheil
SYS: @@ calculation (see macro create_initial_profile.TCM). The command
SYS: @@ INPUT_SCHEIL_PROFILE in the DICTRA MONITOR performs most of the
SYS: @@ set up. Only time and temperature must be entered after the
SYS: @@ INPUT_SCHEIL_PROFILE command is executed.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ In this example only a single phase, ferrite, is entered in the simulation
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: sw FEDEMO
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-sys fe cr ni mn
FE CR NI
MN DEFINED
TDB_FEDEMO: rej ph *
LIQUID:L BCC_A2 LAVES_PHASE_C14
Cbcc_A12 CHI_A12 CUB_A13
DIAMOND_FCC_A4 FCC_A1 HCP_A3
SIGMA REJECTED
TDB_FEDEMO: rest ph bcc
bcc_A2 RESTORED
TDB_FEDEMO: get
09:48:19,008 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES
ELEMENTS
SPECIES
PHASES
PARAMETERS
FUNCTIONS

List of references for assessed data

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'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
 volumes'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'J-O. Andersson and B. Sundman, CALPHAD, 11 (1987) 83-92; TRITA 0270
 (1986); CR-Fe'
'B.J. Lee, Metall. Trans. A, 24A (1993) 1919-1933; Cr-Mn, Fe-Cr -Mn'
'B.J. Lee, CALPHAD, 17 (1993) 251-268; revision of Fe-Cr and Fe -Ni liquid'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); CR -NI'
'W. Huang, CALPHAD, 13 (1989) 243-252; TRITA-MAC 388 (rev 1989); FE-MN'
'L.J. Zhang, Int.J. Mater. Res., 100(2) 160-175 (2009), Fe-Mn-Ni'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
'Shuhong Liu, unpublished work (2010), Mn-Ni, Al-Mn-Ni, Mn-Ni-Zn, Al-Cu-Fe-Mg
 -Mn-Si'
-OK-
TDB_FEDEMO:
TDB_FEDEMO: app MFEDEMO
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys fe cr ni mn
FE CR NI
MN DEFINED
APP: rej ph *
BCC_A2 FCC_A1 REJECTED
APP: rest ph bcc
bcc_A2 RESTORED
APP: get
ELEMENTS
SPECIES
PHASES
PARAMETERS
FUNCTIONS

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: ISIJ International, 35(1995)1415-1421; Cr, Fe and Ni
 diffusion bcc Cr-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
 in bcc Fe'
'Assessed from data presented in Landolt-Bornstein, Vol. 26, ed. H.
 Mehrer, Springer (1990); Impurity diff of Mn in bcc Fe.''
-OK-
APP:
APP: go dict-mon
NO TIME STEP DEFINED
DIC>
DIC>
DIC> @@ THE INPUT_SCHEIL_PROFILE COMMAND PERFORMS MOST OF THE SET UP
DIC> input_scheil_profile
INFO: SCHEIL_REGION CREATED

```
FILE NAME /XF.TXT/: segregation_profile.TXT
ENTER WIDTH OF REGION /1/: 100e-6
INFO: LINEAR GRID IN SCHEIL_REGION ENTERED WITH 100 GRID POINTS
ENTER MAIN SOLID SOLUTION PHASE
PHASE NAME: bcc#
INFO: COMPOSITION PROFILE ENTERED IN REGION
SHOULD MORE PHASES BE ENTERED IN THE REGION /NO/: n
INFO: TO COMPLETE SETUP, ENTER TEMPERATURE AND
SIMULATION TIME
DIC>
DIC>
DIC> @@ ENTER THE HEAT TREATMENT TEMPERATURE
DIC> s-cond
GLOBAL OR BOUNDARY CONDITION /GLOBAL/: glob
VARIABLE : t
LOW TIME LIMIT /0/: 0 1473; * n
DIC>
DIC>
DIC> @@ ENTER A SIMULATION TIME
DIC> se-si-ti
END TIME FOR INTEGRATION /1/: 3600
AUTOMATIC TIMESTEP CONTROL /YES/: y
MAX TIMESTEP DURING INTEGRATION /360/: 360
INITIAL TIMESTEP : /1E-07/: 1e-7
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/: 1e-9
DIC>
DIC>
DIC> save exa7 y
DIC>
DIC> set-inter
--OK--
DIC>
```

exa7-run

DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa7\run.DCM.test"
DIC> go dict-mon
TIME STEP AT TIME 0.00000E+00
DIC>
DIC> read exa7
OK
DIC>
DIC> sim
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: CR = .180420497424242 FE = .796498787878788
MN = .009990168782828 NI = .0130905459141414
TOTAL SIZE OF SYSTEM: 1E-04 [m]
U-FRACTION IN SYSTEM: CR = .180420497424242 FE = .796498787878788
MN = .009990168782828 NI = .0130905459141414
TOTAL SIZE OF SYSTEM: 1E-04 [m]
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424242 FE = .796498787878788
MN = .009990168782828 NI = .0130905459141414
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.1001000E-03 DT = 0.1000000E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424243 FE = .796498787878788
MN = .009990168782828 NI = .0130905459141414
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.41908397 DT = 0.41898387 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424242 FE = .796498787878788
MN = .009990168782828 NI = .0130905459141414
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 1.2570517 DT = 0.83796774 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424242 FE = .796498787878788
MN = .0099901687828279 NI = .0130905459141418
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 2.9329872 DT = 1.6759355 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424243 FE = .7964987878788
MN = .0099901687828289 NI = .0130905459141397
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 6.2848581 DT = 3.3518710 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424257 FE = .796498787878809
MN = .00999016878284567 NI = .0130905459140889
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 12.988600 DT = 6.7037419 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424302 FE = .7964987878887
MN = .0099901687828762 NI = .0130905459139341
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 26.396084 DT = 13.407484 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424318 FE = .7964987878861
MN = .00999016878290129 NI = .0130905459139196
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 53.211052 DT = 26.814968 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424435 FE = .7964987878771
MN = .00999016878299364 NI = .0130905459138002
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 106.84099 DT = 53.629935 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424574 FE = .79649878787836
MN = .00999016878336976 NI = .0130905459136696
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 214.10086 DT = 107.25987 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424432 FE = .7964987878782
MN = .00999016878339539 NI = .0130905459139731
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 428.62060 DT = 214.51974 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .18042049742409 FE = .79649878787776
MN = .0099901687835091 NI = .0130905459146418
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 788.62060 DT = 360.00000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497424077 FE = .7964987877897
MN = .00999016878346275 NI = .013090545914564
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 1148.6206 DT = 360.00000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497423612 FE = .7964987877388
MN = .0099901687833258 NI = .0130905459156742
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 1508.6206 DT = 360.00000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497423264 FE = .7964987877025
MN = .0099901687831708 NI = .0130905459165403
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 1868.6206 DT = 360.00000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497422964 FE = .7964987875229
MN = .00999016878345041 NI = .0130905459183568
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 2228.6206 DT = 360.00000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497422684 FE = .7964987872687
MN = .00999016878399131 NI = .0130905459206371
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 2588.6206 DT = 360.00000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497422471 FE = .7964987870924
MN = .00999016878437158 NI = .0130905459222343
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 2948.6206 DT = 360.00000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497422315 FE = .796498787869698
MN = .00999016878464259 NI = .0130905459233444
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds

```
TIME = 3308.6206 DT = 360.00000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497422201 FE = .796498787868845
MN = .00999016878483191 NI = .013090545924121
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 3600.0000 DT = 291.37940 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .180420497422136 FE = .796498787868424
MN = .00999016878491989 NI = .0130905459245201
TOTAL SIZE OF SYSTEM: 1E-04 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 0.10010000E-03
DELETING TIME-RECORD FOR TIME 0.41908397
DELETING TIME-RECORD FOR TIME 1.2570517
DELETING TIME-RECORD FOR TIME 2.9329872
DELETING TIME-RECORD FOR TIME 6.2848581
DELETING TIME-RECORD FOR TIME 12.988600
DELETING TIME-RECORD FOR TIME 26.396084
DELETING TIME-RECORD FOR TIME 53.211052
DELETING TIME-RECORD FOR TIME 106.84099
DELETING TIME-RECORD FOR TIME 214.10086
DELETING TIME-RECORD FOR TIME 428.62060
DELETING TIME-RECORD FOR TIME 788.62060
DELETING TIME-RECORD FOR TIME 1148.6206
DELETING TIME-RECORD FOR TIME 1508.6206
DELETING TIME-RECORD FOR TIME 1868.6206
DELETING TIME-RECORD FOR TIME 2228.6206
DELETING TIME-RECORD FOR TIME 2588.6206
DELETING TIME-RECORD FOR TIME 2948.6206

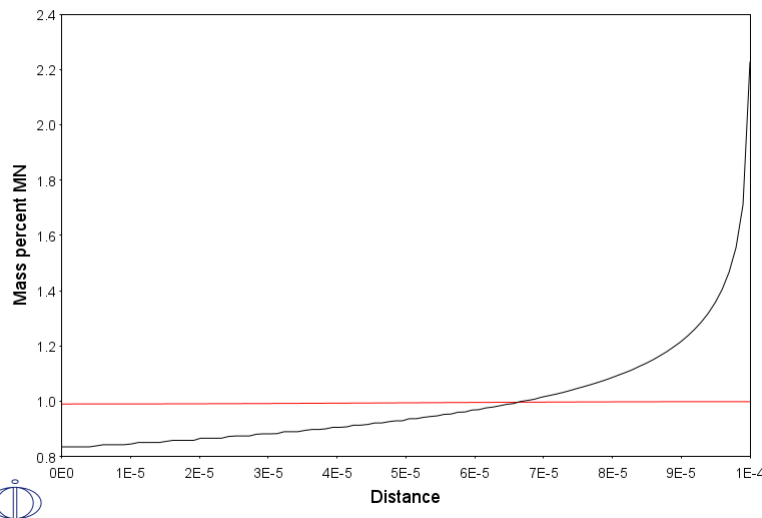
KEEPING TIME-RECORD FOR TIME 3308.6206
AND FOR TIME 3600.0000
WORKSPACE RECLAIMED
```

TIMESTEP AT 3600.00000 SELECTED

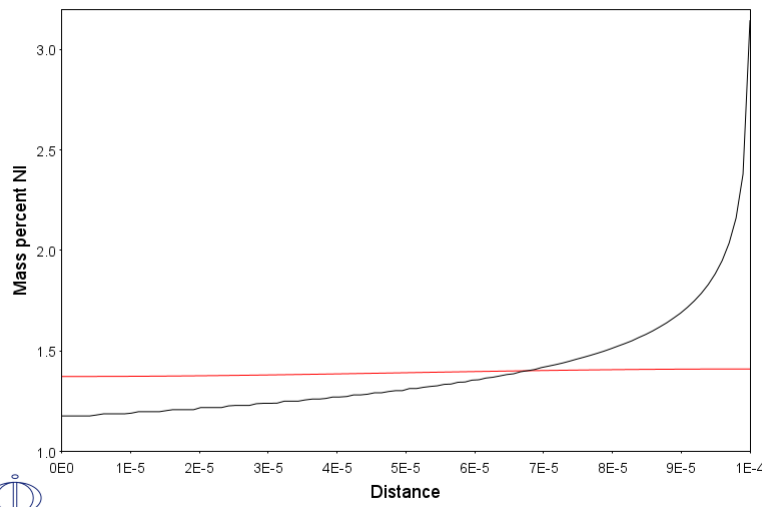
```
DIC>
DIC> set-inter
--OK--
DIC>
```

exa7-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exa7\plot.DCM.test"
DIC> go dict-mon
TIME STEP AT TIME 3.60000E+03
DIC>
DIC> read exa7
OK
DIC>
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: s-p-c time 0,3600
POST-1: s-d-a x d g
INFO: Distance is set as independent variable
POST-1: s-d-a y w-p mn
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.09.50.19
Time = 0,3600
CELL #1
```



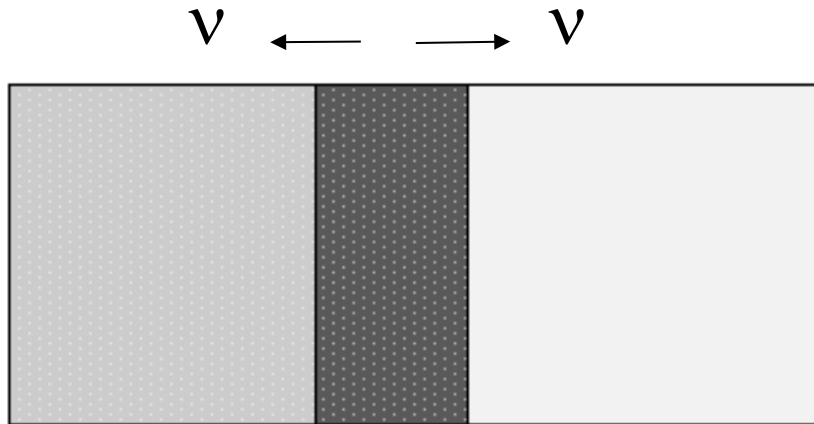
```
POST-1:
POST-1: @@ Hit enter for the next plot
POST-1:@?
POST-1:
POST-1: s-d-a y w-p ni
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.09.50.20
Time = 0,3600
CELL #1
```



```
POST-1:
POST-1: set-inter
--OK---
POST-1:
```



Moving Boundary Problems



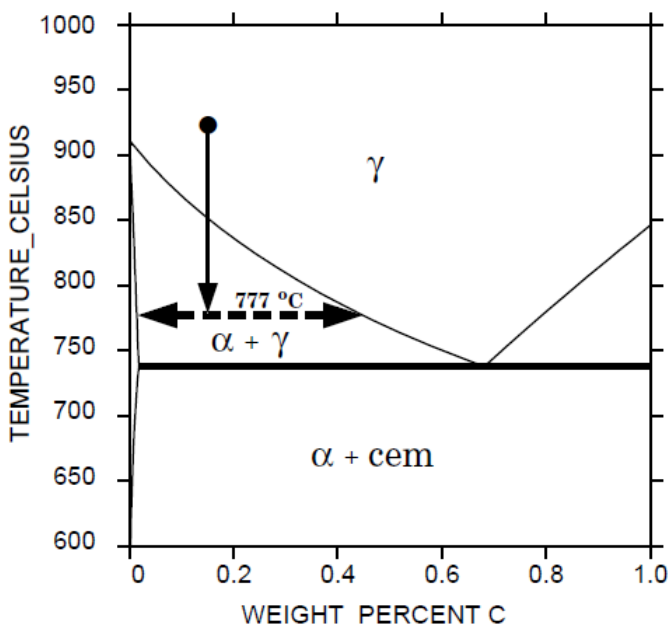


Example exb1a

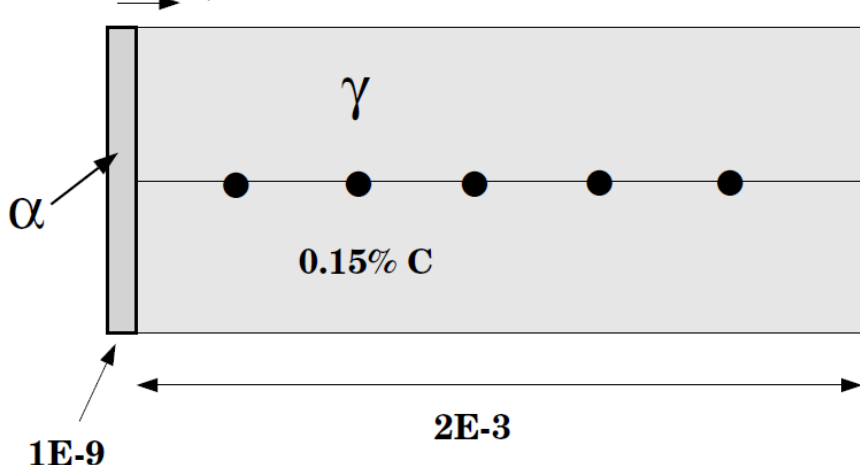
γ to α transformation in a binary Fe-C alloy

This example calculates a ferrite (BCC)/austenite (FCC) transformation in a binary Fe-C alloy. The initial state is an austenite of 2 mm thickness. The composition of the austenite is Fe-0.15wt%C. After austenitization the specimen has been quenched down to 1050K. The system is assumed closed, no boundary conditions are set (a closed system is the default). Ferrite is expected to grow into the austenite. For this reason you start with a thin region with ferrite adjacent to the austenite.

Fe - C Phase diagram



$T = 1050 \text{ K}$



exbla-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exbla\setup.DCM.test"
SYS: @@
SYS: @@ Moving boundary problem.
SYS: @@ Austenite to ferrite transformation in a binary Fe-C alloy
SYS: @@ This example calculates a ferrite(BCC)/austenite(FCC)transformation
SYS: @@ in a binary Fe-C alloy. The initial state is an austenite of 2mm
SYS: @@ thickness. The composition of the austenite is Fe-0.15wt%C.
SYS: @@
SYS: @@ After austenitization the specimen is quenched down to 1050K.
SYS: @@ The system is assumed closed, so no boundary conditions are set
SYS: @@ (a closed system is the default). Ferrite is expected to grow
SYS: @@ into the austenite, which is why we start with a thin
SYS: @@ region with ferrite adjacent to the austenite.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exbla_setup.DCM
SYS:
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /- DEFINED
DICTRA_FCC_A1  REJECTED
TDB_TCFE11: @@
TDB_TCFE11: @@
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE THE DATA
TDB_TCFE11: @@
TDB_TCFE11: sw FEDEMO
Current database: Iron Demo Database v4.0

VA           /- DEFINED
TDB_FEDEMO: @@
TDB_FEDEMO: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_FEDEMO: @@
TDB_FEDEMO: def-sys fe c
FE           C  DEFINED
TDB_FEDEMO: @@
TDB_FEDEMO: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_FEDEMO: @@
TDB_FEDEMO: rej ph * all
GAS:G          LIQUID:L          BCC_A2
LAVES_PHASE_C14  Cbcc_A12        CEMENTITE
CUB_A13         DIAMOND_FCC_A4   FCC_A1
GRAPHITE        HCP_A3          KSI_CARBIDE
M23C6          M5C2            M7C3
REJECTED
TDB_FEDEMO: res ph fcc bcc
FCC_A1          BCC_A2    RESTORED
TDB_FEDEMO: @@
TDB_FEDEMO: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_FEDEMO: @@
TDB_FEDEMO: get
09:51:37,261 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'P. Franke, estimated parameter within SGTE, 2007; Fe-C, Ni-C, Mo-C, C-Mn'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'

-OK-
TDB_FEDEMO: @@
TDB_FEDEMO: @@
TDB_FEDEMO: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE DATA
TDB_FEDEMO: @@
TDB_FEDEMO: append
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
```

TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2
TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
SNOB1 = SGTE Noble Metal Alloys Database v1.2
STBC2 = SGTE Thermal Barrier Coating TDB v2.2
STBC1 = SGTE Thermal Barrier Coating TDB v1.1
SALT1 = SGTE Molten Salts Database v1.3
SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2 = TC Semi-Conductors v2.1
SLAG4 = Fe-containing Slag v4.1
SLAG3 = Fe-containing Slag v3.2
SLAG2 = Fe-containing Slag v2.2
SLAG1 = Fe-containing Slag v1.2
TCOX11 = Metal Oxide Solutions v11.0
TCOX10 = Metal Oxide Solutions v10.1
TCOX9 = Metal Oxide Solutions v9.0
TCOX8 = Metal Oxide Solutions v8.0
TCOX7 = Metal Oxide Solutions v7.0
TCOX6 = Metal Oxide Solutions v6.0
TCOX5 = Metal Oxide Solutions v5.1
TCOX4 = Metal Oxide Solutions v4.1
ION3 = Ionic Solutions v3.0
ION2 = Ionic Solutions v2.6
ION1 = Ionic Solutions v1.5
NOX2 = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4 = Solder Alloys v4.0
TCSLD3 = Solder Alloys v3.3
TCSLD2 = Solder Alloys v2.0
TCSLD1 = Solder Alloys v1.1
TCSI1 = Ultrapure Silicon v1.2
TCMP2 = Materials Processing v2.5
TCE51 = Combustion/Sintering v1.1
TCSC1 = Super Conductor v1.0
TCFC1 = SOFC Database v1.0
TCNF2 = Nuclear Fuels v2.1b
NUMT2 = Nuclear Materials v2.1
NUOX4 = Nuclear Oxides v4.2
NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephista-19
MEPH15 = IRSN Mephista-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQ52 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SLDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.2
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3
MOBFE1 = Steels/Fe-Alloys Mobility v1.1
MOBFE2 = Steels/Fe-Alloys Mobility v2.0
MOBFE3 = Steels/Fe-Alloys Mobility v3.0
MOBFE4 = Steels/Fe-Alloys Mobility v4.0
MOBFE5 = Steels/Fe-Alloys Mobility v5.0
MOBFE6 = Steels/Fe-Alloys Mobility v6.0

```

MOBN15 = Ni-Alloys Mobility v5.1
MOBN14 = Ni-Alloys Mobility v4.1
MOBN13 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBS11 = Si-Alloys Mobility v1.0
MOBSDL1 = Solder-Alloys Mobility v1.1
MOBT14 = Ti-Alloys Mobility v4.0
MOBT13 = Ti-Alloys Mobility v3.1
MOBT12 = Ti-Alloys Mobility v2.0
MOBT11 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

```

DATABASE NAME /FEDEMO/: MFEDEMO

Current database: Fe-Alloys Mobility demo database v2.0

```

VA DEFINED
APP: def-sys fe c
    FE C DEFINED
APP: rej ph * all
    BCC_A2 FCC_A1 REJECTED
APP: res ph fcc bcc
    FCC_A1 BCC_A2 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

```

List of references for assessed data

```

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Z. Metallkunde 85(1994)498-501; C and N diffusion in bcc Cr
-Fe-Ni'
'B. Jonson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'
-OK-

```

```

APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER GLOBAL CONDITION T
DIC> @@
DIC> set-condition global T 0 1050; * N
DIC>
DIC> @@
DIC> @@ START BY ENTERING THE REGIONS ferrite AND austenite WHERE WE
DIC> @@ PUT THE BCC AND FCC PHASE, RESPECTIVELY. THE FERRITE REGION IS
DIC> @@ ASSUMED INITIALLY TO BE VERY THIN, 1E-9 METERS.
DIC> @@
DIC> enter-region
REGION NAME : ferrite
DIC>
DIC> enter-region
REGION NAME : austenite
ATTACH TO REGION NAMED /FERRITE/:
ATTACHED TO THE RIGHT OF FERRITE /YES/:
DIC>
DIC> @@
DIC> @@ ENTER GRIDS INTO THE REGIONS
DIC> @@
DIC> enter-grid
REGION NAME : /FERRITE/: ferrite
WIDTH OF REGION /1/: 1e-9
TYPE /LINEAR/: AUTO
DIC>
DIC> enter-grid austenite
WIDTH OF REGION /1/: 20e-4
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER active PHASES INTO THE REGIONS
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /FERRITE/: ferrite
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: bcc
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /AUSTENITE/: austenite
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC>
DIC>

```

```

DIC> @@
DIC> @@ ENTER AN INITIAL COMPOSITION INTO BCC
DIC> @@
DIC> enter-composition
REGION NAME : /FERRITE/: ferrite
PHASE NAME: /BCC_A2/: bcc
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: C
TYPE /LINEAR/: linear
VALUE OF FIRST POINT : 0.01
VALUE OF LAST POINT : /1E-2/: 0.01
DIC>
DIC> @@
DIC> @@ ENTER AN INITIAL COMPOSITION INTO FCC
DIC> @@
DIC> enter-composition
REGION NAME : /AUSTENITE/: austenite
PHASE NAME: /FCC_A1/: fcc#1
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: C
TYPE /LINEAR/: linear
VALUE OF FIRST POINT : 0.15
VALUE OF LAST POINT : /0.15/: 0.15
DIC>
DIC> @@
DIC> @@ THE BOUNDARY CONDITION IS A CLOSED SYSTEM AS WE DO NOT SPECIFY
DIC> @@ ANYTHING ELSE
DIC> @@
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 1e9
AUTOMATIC TIMESTEP CONTROL /YES/: YES
MAX TIMESTEP DURING INTEGRATION /100000000/: 1e8
INITIAL TIMESTEP : /1E-07/: 1E-5
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/: 1E-5
DIC>
DIC> @@
DIC> @@ IMPLICIT (1) TIME INTEGRATION IS USED INSTEAD OF THE MORE ACCURATE
DIC> @@ (BUT LESS STABLE) TRAPETZOIDAL METHOD WHICH IS THE DEFAULT.
DIC> @@
DIC> s-s-c
NS01A PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA TIMESTEPS IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/:
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/:
DEGREE OF IMPURITY WHEN INTEGRATING PDES (AUTO, 0 -> 0.5 -> 1) : /AUTO/: 1.0
MAX TIMESTEP CHANGE PER TIMESTEP : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/:
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exbla Y
DIC>
DIC> set-inter
---OK---
DIC>

```

exbla-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exbla\run.DCM.test"
DIC>
DIC>
DIC> @@ exbla_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE bla
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exbla
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> simulate
Region: FERRITE
single geometric dense at 0.10000E-08
1.0000 24
Region: AUSTENITE
single geometric dense at 0.0000
1.1762 99
Trying old scheme 3
U-FRACTION IN SYSTEM: C = .00698495590385911 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
U-FRACTION IN SYSTEM: C = .00698495590385911 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
3.981486698395194E-005 3.982285206223790E-005 1.590378050717499E-012 8.422113835746911E-020 TIME = 0.10000000E-04 DT = 0.10000000E-04 SUM OF SQUARES = 0.84221138E-19
04 DT = 0.10000000E-04 SUM OF SQUARES = 0.84221138E-19
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.29175547E-02 AND 0.29175547E-02
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.30175547E-07
U-FRACTION IN SYSTEM: C = .00698495590212123 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
13 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

CPU time used in timestep 0 seconds
3.499840016754033E-006 3.500511904249956E-006 1.189295167153537E-007 4.111386278498581E-008 2.962711489812290E-009 TIME = 0.2530748984847927E-012 1.388215646533131E-016 6.302028868602742E-024 TIME = 0.30000000E-04 DT = 0.20000000E-04 SUM OF SQUARES = 0.63020289E-23
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.11112117E-03 AND 0.11112117E-03
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.32397971E-07
U-FRACTION IN SYSTEM: C = .00698495800112155 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
13 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

CPU time used in timestep 0 seconds
8.791374341914706E-008 8.798286131288386E-008 8.680197148511708E-012 8.237998777204505E-016 5.570658332104316E-024 TIME = 0.70000000E-04 DT = 0.40000000E-04 SUM OF SQUARES = 0.55706583E-23
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.81460777E-04 AND 0.81460777E-04
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.35656402E-07
U-FRACTION IN SYSTEM: C = .00698495800112151 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
CPU time used in timestep 0 seconds
3.282053178498482E-006 3.283518217738399E-006 3.522797165763882E-006 1.817586682189996E-007 4.865296022642763E-008 TIME = 3.905763819835909E-011 5.263908083351613E-015 6.537906182240236E-022 TIME = 0.15000000E-03 DT = 0.80000000E-04 SUM OF SQUARES = 0.65379062E-21
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.58646939E-04 AND 0.58646939E-04
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.40348157E-07
U-FRACTION IN SYSTEM: C = .00698495794642423 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
CPU time used in timestep 0 seconds
2 GRIDPOINT(S) ADDED TO CELL #1 REGION: FERRITE
4.713654652390555E-007 4.723390763359801E-007 4.586676384660827E-012 4.399046976856031E-017 TIME = 0.31000000E-03 DT = 0.16000000E-03 SUM OF SQUARES = 0.43990470E-16
03 DT = 0.16000000E-03 SUM OF SQUARES = 0.43990470E-16
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.51923334E-04 AND 0.51923334E-04
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.48655891E-07
U-FRACTION IN SYSTEM: C = .00698495386454072 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
CPU time used in timestep 0 seconds
4.378612637278334E-006 4.381488086560821E-006 1.153721994970485E-009 2.848526252439885E-013 1.743908466437677E-020 TIME = 0.63000000E-03 DT = 0.32000000E-03 SUM OF SQUARES = 0.17439085E-19
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.35629005E-04 AND 0.35629005E-04
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.60057172E-07
U-FRACTION IN SYSTEM: C = .006984948587741 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
CPU time used in timestep 0 seconds
2.098301746880194E-006 2.099772516707655E-006 6.174131548456762E-010 1.696049180678346E-013 1.285332330763806E-020 TIME = 0.12700000E-02 DT = 0.64000000E-03 SUM OF SQUARES = 0.12853323E-19

output ignored...

... output resumed

TIME = 0.57592186E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.15364008E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.19926141E-15 AND -0.19926141E-15
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.13794590E-02
U-FRACTION IN SYSTEM: C = .00698584273647938 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

CPU time used in timestep 0 seconds
4.535237501308182E-005 4.535773928372776E-005 6.377119607700181E-022 TIME = 0.67592186E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.63771196E-21
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.13492716E-15 AND 0.13492716E-15
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.13794725E-02
U-FRACTION IN SYSTEM: C = .00698570514621074 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
20 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FERRITE

CPU time used in timestep 0 seconds
1.970327668730446E-005 1.970567086562696E-005 5.024070003177980E-019 TIME = 0.77592186E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.50240700E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.85417341E-16 AND -0.85417341E-16
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.13794639E-02
U-FRACTION IN SYSTEM: C = .00698579224946741 FE = 1
```

```

TOTAL SIZE OF SYSTEM: .002000001 [m]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

CPU time used in timestep          0 seconds
9.869293459165407E-006      9.870366152308391E-006      1.893331983544551E-
022     TIME = 0.87592186E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.18933320E-21
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.70327802E-16 AND 0.70327802E-16
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.13794710E-02
U-FRACTION IN SYSTEM: C = .00698572053357829 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
10 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FERRITE

CPU time used in timestep          0 seconds
1.668288167588449E-005      1.668402994762454E-005      2.250176276744608E-
028     TIME = 0.97592186E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.22501763E-27
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.13134925E-15 AND -0.13134925E-15
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.13794578E-02
U-FRACTION IN SYSTEM: C = .00698585447532275 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

CPU time used in timestep          0 seconds
4.795129124478679E-005      4.795492713364641E-005      1.073766223439627E-
019     TIME = 0.10000000E+10 DT = 24078140. SUM OF SQUARES = 0.10737662E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.21081044E-15 AND 0.21081044E-15
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.13794629E-02
U-FRACTION IN SYSTEM: C = .00698580271423688 FE = 1
TOTAL SIZE OF SYSTEM: .002000001 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME    0.0000000
DELETING TIME-RECORD FOR TIME    0.10000000E-04
DELETING TIME-RECORD FOR TIME    0.30000000E-04
DELETING TIME-RECORD FOR TIME    0.70000000E-04
DELETING TIME-RECORD FOR TIME    0.15000000E-03
DELETING TIME-RECORD FOR TIME    0.31000000E-03
DELETING TIME-RECORD FOR TIME    0.63000000E-03
DELETING TIME-RECORD FOR TIME    0.12700000E-02
DELETING TIME-RECORD FOR TIME    0.25500000E-02
DELETING TIME-RECORD FOR TIME    0.51100000E-02
DELETING TIME-RECORD FOR TIME    0.10230000E-01
DELETING TIME-RECORD FOR TIME    0.20470000E-01
DELETING TIME-RECORD FOR TIME    0.40950000E-01
DELETING TIME-RECORD FOR TIME    0.81910000E-01
DELETING TIME-RECORD FOR TIME    0.16383000
DELETING TIME-RECORD FOR TIME    0.32767000
DELETING TIME-RECORD FOR TIME    0.65535000
DELETING TIME-RECORD FOR TIME    1.3107100
DELETING TIME-RECORD FOR TIME    2.6214300
DELETING TIME-RECORD FOR TIME    5.2428700
DELETING TIME-RECORD FOR TIME    10.485750
DELETING TIME-RECORD FOR TIME    20.971510
DELETING TIME-RECORD FOR TIME    41.943030
DELETING TIME-RECORD FOR TIME    83.886070
DELETING TIME-RECORD FOR TIME    167.77215
DELETING TIME-RECORD FOR TIME    335.54431
DELETING TIME-RECORD FOR TIME    671.08863
DELETING TIME-RECORD FOR TIME    1342.1773
DELETING TIME-RECORD FOR TIME    2684.3546
DELETING TIME-RECORD FOR TIME    5368.7091
DELETING TIME-RECORD FOR TIME    10737.418
DELETING TIME-RECORD FOR TIME    21474.836
DELETING TIME-RECORD FOR TIME    42949.673
DELETING TIME-RECORD FOR TIME    85899.346
DELETING TIME-RECORD FOR TIME    171798.69
DELETING TIME-RECORD FOR TIME    343597.38
DELETING TIME-RECORD FOR TIME    687194.77
DELETING TIME-RECORD FOR TIME    1374389.5
DELETING TIME-RECORD FOR TIME    2748779.1
DELETING TIME-RECORD FOR TIME    5497558.1
DELETING TIME-RECORD FOR TIME    10995116.
DELETING TIME-RECORD FOR TIME    21990233.
DELETING TIME-RECORD FOR TIME    43980465.
DELETING TIME-RECORD FOR TIME    87960930.
DELETING TIME-RECORD FOR TIME    0.17592186E+09
DELETING TIME-RECORD FOR TIME    0.27592186E+09
DELETING TIME-RECORD FOR TIME    0.37592186E+09
DELETING TIME-RECORD FOR TIME    0.47592186E+09
DELETING TIME-RECORD FOR TIME    0.57592186E+09
DELETING TIME-RECORD FOR TIME    0.67592186E+09
DELETING TIME-RECORD FOR TIME    0.77592186E+09
DELETING TIME-RECORD FOR TIME    0.87592186E+09

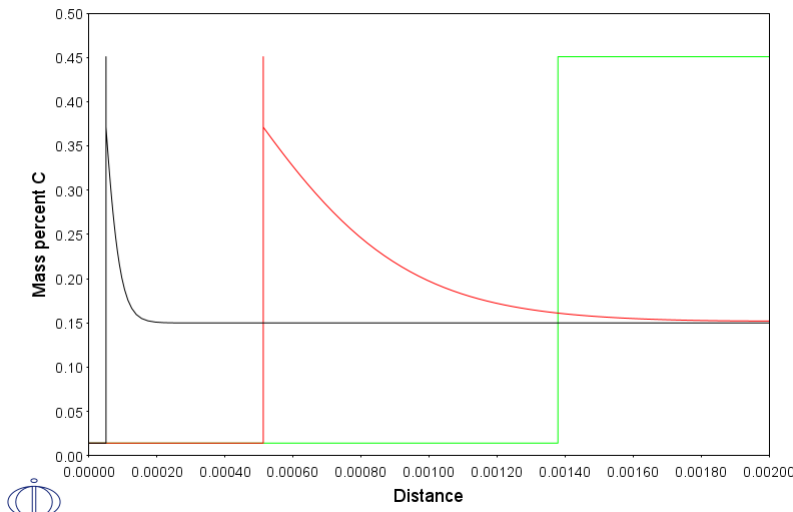
KEEPING TIME-RECORD FOR TIME    0.97592186E+09
AND FOR TIME                  0.10000000E+10
WORKSPACE RECLAIMED

TIMESTEP AT      0.100000000E+10 SELECTED

DIC>
DIC>
DIC>
DIC> @@
DIC> @@ THE SIMULATION IS FINISHED
DIC> @@
DIC>
DIC> set-inter
--OK--
DIC>
```

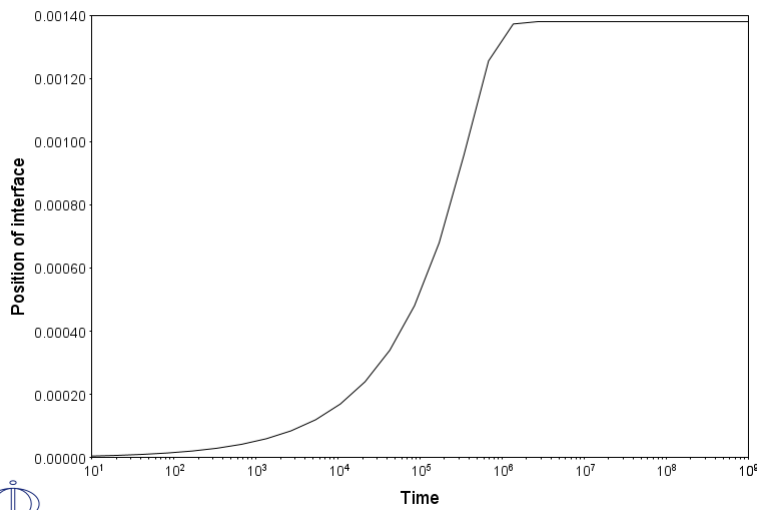
exbla-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exbla\plot.DCM.test"
DIC>
DIC>
DIC> @@ exbla_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE bia
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 1.00000E+09
DIC> read exbla
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1:
POST-1: @@
POST-1: @@ PLOT THE CARBON CONCENTRATIONS AT DIFFERENT TIMES
POST-1: @@
POST-1: s-d-a x distance global
INFO: Distance is set as independent variable
POST-1: s-d-a y w-p c
POST-1: s-p-c time le3,1e5,1e9
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.09.53.40
Time=1000,100000,1E+09
CELL #1
```



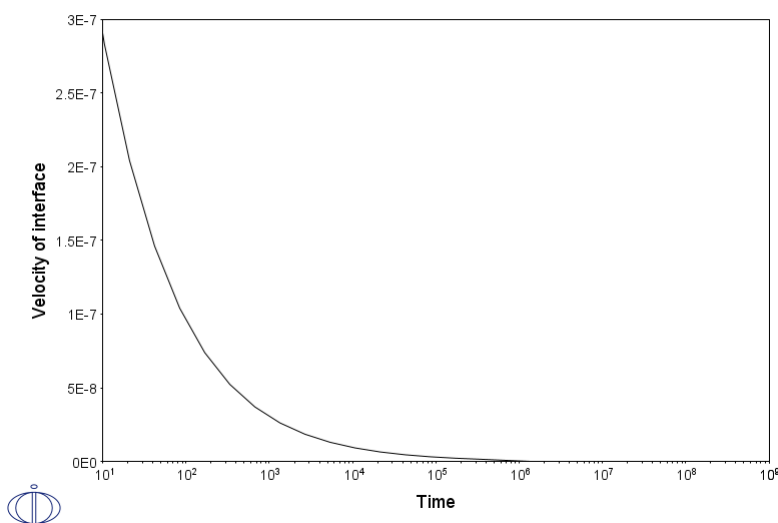
```
POST-1:
POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: @@
POST-1: @@ PLOT THE POSITION OF THE BCC/FCC INTERPHASE
POST-1: @@
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-d-a y
VARIABLE : pos
INTERFACE : aus
UPPER OR LOWER INTERFACE OF REGION AUSTENITE#1 /LOWER/: lower
POST-1:
POST-1: set_axis_type
AXIS (X, Y OR Z) : x
AXIS TYPE /LINEAR/: log
POST-1:
POST-1: s-s-s
AXIS (X, Y OR Z) : x
AUTOMATIC SCALING (Y OR N) /N/: n
MIN VALUE : 10
MAX VALUE : 1e9
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
    OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

2021.05.13.09.53.42
LOWER INTERFACE OF REGION "AUSTENITE#1"
CELL #1



POST-1:
POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: @@
POST-1: @@ PLOT THE VELOCITY OF THE BCC/FCC INTERPHASE
POST-1: @@
POST-1: s-d-a
AXIS (X, Y OR Z) : y
VARIABLE : velocity
INTERFACE : aus
UPPER OR LOWER INTERFACE OF REGION AUSTENITE#1 /LOWER/: lower
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

2021.05.13.09.53.42
LOWER INTERFACE OF REGION "AUSTENITE#1"
CELL #1



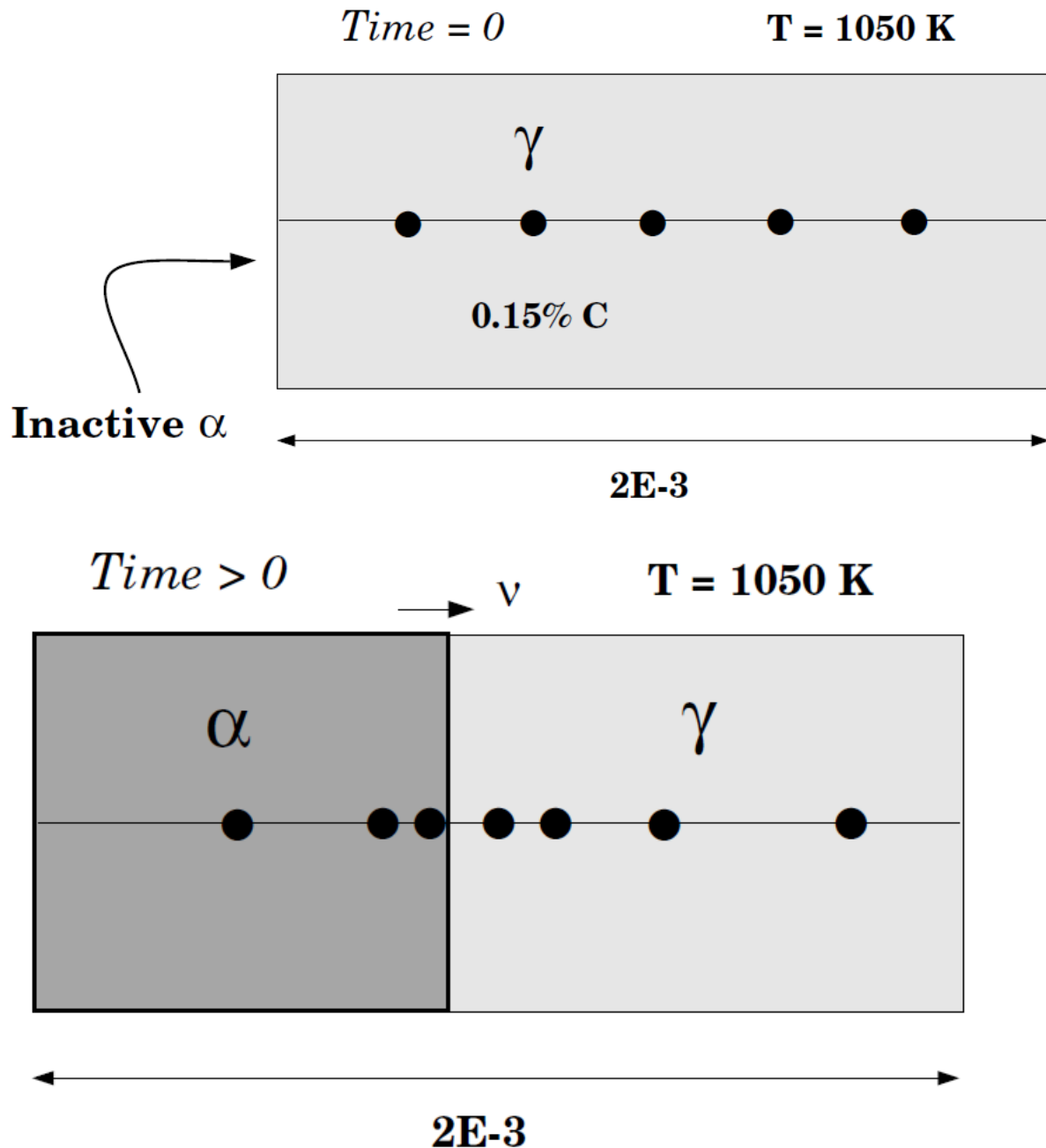
POST-1:
POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: set-inter
--OK--
POST-1:



Example exb1b

γ to α transformation in a binary Fe-C alloy: Inactive α

This is the same example as in exb1a but now the problem is with ferrite as an inactive phase adjacent to the initial austenite.



exblb-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exblb\setup.DCM.test"
SYS: @@
SYS: @@ Moving boundary problem.
SYS: @@ Austenite to ferrite transformation in a binary Fe-C alloy
SYS: @@ This is the same example as in exbla but now the problem is with
SYS: @@ ferrite as an inactive phase adjacent to the initial austenite.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exblb_setup.DCM
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA                                /- DEFINED
DICTRA_FCC_A1   REJECTED
TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: @@
TDB_TCFE11: sw FEDEMO
Current database: Iron Demo Database v4.0

VA                                /- DEFINED
TDB_FEDEMO: @@
TDB_FEDEMO: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_FEDEMO: @@
TDB_FEDEMO: def-sys fe c
FE                               C DEFINED
TDB_FEDEMO: @@
TDB_FEDEMO: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_FEDEMO: @@
TDB_FEDEMO: rej ph * all
GAS:G                         LIQUID:L          BCC_A2
LAVES_PHASE_C14      CBCC_A12        CEMENTITE
CUB_A13             DIAMOND_FCC_A4    FCC_A1
GRAPHITE           HCP_A3           KSI_CARBIDE
M23C6              M5C2            M7C3
REJECTED
TDB_FEDEMO: res ph fcc bcc
FCC_A1                  BCC_A2   RESTORED
TDB_FEDEMO: @@
TDB_FEDEMO: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_FEDEMO: @@
TDB_FEDEMO: get
09:55:00,953 INFO  *** Invoking Gibbs Energy System v6 ***
REINITIATING GES ....
ELEMENTS ....
SPECIES ....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'P. Franke, estimated parameter within SGTE, 2007; Fe-C, Ni-C, Mo-C, C-Mn'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
-OK-
TDB_FEDEMO: @@
TDB_FEDEMO: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE THE DATA.
TDB_FEDEMO: @@
TDB_FEDEMO: app
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
```

TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2
TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
SNOB1 = SGTE Noble Metal Alloys Database v1.2
STBC2 = SGTE Thermal Barrier Coating TDB v2.2
STBC1 = SGTE Thermal Barrier Coating TDB v1.1
SALT1 = SGTE Molten Salts Database v1.3
SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2 = TC Semi-Conductors v2.1
SLAG4 = Fe-containing Slag v4.1
SLAG3 = Fe-containing Slag v3.2
SLAG2 = Fe-containing Slag v2.2
SLAG1 = Fe-containing Slag v1.2
TCOX11 = Metal Oxide Solutions v11.0
TCOX10 = Metal Oxide Solutions v10.1
TCOX9 = Metal Oxide Solutions v9.0
TCOX8 = Metal Oxide Solutions v8.0
TCOX7 = Metal Oxide Solutions v7.0
TCOX6 = Metal Oxide Solutions v6.0
TCOX5 = Metal Oxide Solutions v5.1
TCOX4 = Metal Oxide Solutions v4.1
ION3 = Ionic Solutions v3.0
ION2 = Ionic Solutions v2.6
ION1 = Ionic Solutions v1.5
NOX2 = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4 = Solder Alloys v4.0
TCSLD3 = Solder Alloys v3.3
TCSLD2 = Solder Alloys v2.0
TCSLD1 = Solder Alloys v1.1
TCSI1 = Ultrapure Silicon v1.2
TCMP2 = Materials Processing v2.5
TCES1 = Combustion/Sintering v1.1
TCSC1 = Super Conductor v1.0
TCFC1 = SOFC Database v1.0
TCNF2 = Nuclear Fuels v2.1b
NUMT2 = Nuclear Materials v2.1
NUOX4 = Nuclear Oxides v4.2
NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephista-19
MEPH15 = IRSN Mephista-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQ52 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SLDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.2
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3
MOBF61 = Steels/Fe-Alloys Mobility v1.1
MOBF62 = Steels/Fe-Alloys Mobility v2.0
MOBF63 = Steels/Fe-Alloys Mobility v3.0
MOBF64 = Steels/Fe-Alloys Mobility v4.0
MOBF65 = Steels/Fe-Alloys Mobility v5.0
MOBF66 = Steels/Fe-Alloys Mobility v6.0
MOBN15 = Ni-Alloys Mobility v5.1
MOBN14 = Ni-Alloys Mobility v4.1
MOBN13 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0

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MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBSI1 = Si-Alloys Mobility v1.0
MOBSDL1 = Solder-Alloys Mobility v1.1
MOBT14 = Ti-Alloys Mobility v4.0
MOBT13 = Ti-Alloys Mobility v3.1
MOBT12 = Ti-Alloys Mobility v2.0
MOBT11 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

DATABASE NAME /FEDEMO/: MFEDEMO
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys fe c
FE C DEFINED
APP: rej ph * all FCC_A1 REJECTED
BCC_A2 FCC_A2 RESTORED
APP: res ph fcc bcc
FCC A1 BCC_A2 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Z. Metallkunde 85(1994)498-501; C and N diffusion in bcc Cr
-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'

-OK-
APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 1050; * N
DIC>
DIC> @@
DIC> @@ START BY ENTERING THE REGION austenite WHERE WE PUT THE fcc PHASE
DIC> @@
DIC> enter-region
REGION NAME : austenite
DIC>
DIC> @@
DIC> @@ ENTER THE GRID INTO THE REGION
DIC> @@
DIC> enter-grid
REGION NAME : /AUSTENITE/: austenite
WIDTH OF REGION /1/: 20e-4
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER THE active PHASES INTO THE REGIONS
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /AUSTENITE/: austenite
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC> @@
DIC> @@ ENTER THE inactive PHASES INTO THE REGIONS
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inactive
ATTACH TO REGION NAMED /AUSTENITE/: austenite
ATTACHED TO THE RIGHT OF AUSTENITE /YES/: no
PHASE NAME: /NONE/: bcc
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-5
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITION FOR FCC
DIC> @@
DIC> enter-composition
REGION NAME : /AUSTENITE/: austenite
PHASE NAME: /FCC_A1/: fcc#1
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: C
TYPE /LINEAR/: lin
VALUE OF FIRST POINT : 0.15
VALUE OF LAST POINT : /0.15/: 0.15
DIC>
DIC> @@
DIC> @@ THE BOUNDARY CONDITION IS A CLOSED SYSTEM AS WE DO NOT SPECIFY
DIC> @@ ANYTHING ELSE

```

```
DIC> @@  
DIC> @@  
DIC> @@ SET THE SIMULATION TIME  
DIC> @@  
DIC> set-simulation-time  
END TIME FOR INTEGRATION /1/: 1e9  
AUTOMATIC TIMESTEP CONTROL /YES/:  
MAX TIMESTEP DURING INTEGRATION /100000000/:  
INITIAL TIMESTEP : /1E-07/:  
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:  
DIC>  
DIC>  
DIC>  
DIC> @@  
DIC> @@ IMPLICIT (1) TIME INTEGRATION IS USED INSTEAD OF THE MORE ACCURATE  
DIC> @@ (BUT LESS STABLE) TRAPEZOIDAL METHOD WHICH IS THE DEFAULT.  
DIC> @@  
DIC> s-s-c  
NSO1A PRINT CONTROL : /0/:  
FLUX CORRECTION FACTOR : /1/:  
NUMBER OF DELTA TIMESTEPS IN CALLING MULDIF: /2/:  
CHECK INTERFACE POSITION /AUTO/:  
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:  
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:  
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/:  
DEGREE OF IMPURITY WHEN INTEGRATING PDES (AUTO, 0 -> 0.5 -> 1): /AUTO/: 1.0  
MAX TIMESTEP CHANGE PER TIMESTEP : /2/:  
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:  
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:  
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/:  
DIC> @@  
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT  
DIC> @@  
DIC> save exb1b Y
```

This file contains results from a previous SIMULATEREACTION command.
The SAVE command will save the current status of the program but destroy
the results from the previous simulation.

```
DIC>  
DIC> set-inter  
--OK---  
DIC>
```

exb1b-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb1b\run.DCM.test"
DIC>
DIC>
DIC> @@ exb1b_run.DCM
DIC>
DIC> @@ FILE FOR RUNNING EXAMPLE b1b
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exb1b
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim
Region: AUSTENITE
single geometric dense at 0.0000
1.2284 102
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
INFO: PHASE BCC_A2 IS SCHEDULED TO APPEAR
REGION STATUS CHANGE, ITERATING: TIME= 0.50000000E-07
REGION STATUS CHANGE, ITERATING: TIME= 0.25000000E-07
TIME = 0.25000000E-07 DT = 0.25000000E-07
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE

KEEPING TIME-RECORD FOR TIME 0.000000
AND FOR TIME 0.25000000E-07
WORKSPACE RECLAIMED
WIDTH OF NEW REGION R_BCC_A2 /1E-06/:
Trying old scheme 3
START VALUE(S) FOR INTERFACE #2 R_BCC_A2/AUSTENITE, CELL #1
-----
VELOCITY /1/:
25 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

Trying old scheme 3
U-FRACTION IN SYSTEM: C = .00698156310125388 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
U-FRACTION IN SYSTEM: C = .00698156310125388 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
1 GRIDPOINT(S) ADDED TO CELL #1 REGION: R_BCC_A2
2.641796457615056E-003 2.641820164613401E-003 0.237813210027546 2.896595830854927E-003 2.863309993054893E-003
2.465854445577573E-003 3.038053080121225E-003 2.558808984270720E-003 2.415833955535861E-003
2.363564686115593E-003 3.038053080121225E-003 2.415833955535861E-003 2.496303890482957E-003
2.376837928709497E-003 2.369135059341974E-003 2.366895698823924E-003 2.361896017028972E-003
2.360225243301796E-003 2.369135059341974E-003 2.361896017028972E-003 2.359389068072184E-003
2.358662180339569E-003 2.369135059341974E-003 2.359389068072184E-003 2.361278860714089E-003
2.358761592036592E-003 2.358659683976153E-003 2.358709289015734E-003 2.358624346087840E-003
2.358591707433402E-003 2.358824346542741E-003 2.358624346087840E-003 2.358620293471447E-003
2.358624346087840E-003 2.358620293471447E-003 2.358624346087840E-003 2.358624346087840E-003

      ERROR RETURN FROM NS01A BECAUSE 5 CALLS OF CALFUN FAILED TO IMPROVE THE RESIDUALS
*** ERROR 1890 IN DCNS01: ERROR RETURN FROM NS01A
2.641796457615056E-003 2.641796461150068E-003 2.641796457615056E-003 0.237508579819959 2.641796457615056E-003
003 2.896591057258121E-003 2.641796461150068E-003 2.641796457615056E-003 2.896591057258121E-003 2.641796457615056E-003
003 0.237508579819959 2.641796457615056E-003 2.896591057258121E-003 2.775733791491543E-003 2.641796457615056E-003
003 3.038163915825988E-003 2.641796457615056E-003 2.775733791491543E-003 2.641796457615056E-003 2.641796457615056E-003
003 2.558805763213206E-003 2.558805763213206E-003 2.363552975740587E-003 2.363552975740587E-003 2.363552975740587E-003
003 5.738025859324140E-003 2.363552975740587E-003 2.558805763213206E-003 2.558805763213206E-003 2.363552975817769E-003
003 2.363552975740587E-003

output ignored...

... output resumed

CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.62828906E-16 AND 0.62828906E-16
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13796467E-02
U-FRACTION IN SYSTEM: C = .00698392110790924 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
5 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_BCC_A2

CPU time used in timestep 0 seconds
2.334103381700427E-006 2.334487099558977E-006 4.999449696713977E-006
029 TIME = 0.60500275E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.49994497E-28
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.13337965E-16 AND -0.13337965E-16
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13796454E-02
U-FRACTION IN SYSTEM: C = .00698393470913857 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

CPU time used in timestep 0 seconds
4.150258752341024E-005 4.150602239630179E-005 1.600696493093292E-005
018 TIME = 0.70500275E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.16006965E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.18516576E-16 AND 0.18516576E-16
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13796473E-02
U-FRACTION IN SYSTEM: C = -.00698391582708524 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_BCC_A2

CPU time used in timestep 0 seconds
2.685801124862139E-005 2.686184732485364E-005 3.633434088767752E-005
017 TIME = 0.80500275E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.36334341E-16
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.72645873E-17 AND -0.72645873E-17
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13796465E-02
```

U-FRACTION IN SYSTEM: C = .00698392323505816 FE = 1
 TOTAL SIZE OF SYSTEM: .002 [m]
 1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

CPU time used in timestep 0 seconds
 6.865812362918548E-005 6.866052985034949E-005 1.475550048511266E-005
 017 TIME = 0.90500275E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.14755500E-16
 CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.33508870E-16 AND 0.33508870E-16
 POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13796499E-02
 U-FRACTION IN SYSTEM: C = .00698389064794 FE = 1
 TOTAL SIZE OF SYSTEM: .002 [m]
 1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_BCC_A2

CPU time used in timestep 0 seconds
 1.236458183132819E-005 1.236929231472599E-005 2.095296479852855E-005
 025 TIME = 0.10000000E+10 DT = 94997254. SUM OF SQUARES = 0.20952965E-24
 CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.15597347E-16 AND 0.15597347E-16
 POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13796514E-02
 U-FRACTION IN SYSTEM: C = .00698387395528536 FE = 1
 TOTAL SIZE OF SYSTEM: .002 [m]
 MUST SAVE WORKSPACE ON FILE
 WORKSPACE SAVED ON FILE
 RECLAIMING WORKSPACE
 DELETING TIME-RECORD FOR TIME 0.0000000
 DELETING TIME-RECORD FOR TIME 0.25000000E-07
 DELETING TIME-RECORD FOR TIME 0.12500000E-06
 DELETING TIME-RECORD FOR TIME 0.60937151E-05
 DELETING TIME-RECORD FOR TIME 0.18031145E-04
 DELETING TIME-RECORD FOR TIME 0.41906006E-04
 DELETING TIME-RECORD FOR TIME 0.89655726E-04
 DELETING TIME-RECORD FOR TIME 0.18515517E-03
 DELETING TIME-RECORD FOR TIME 0.37615405E-03
 DELETING TIME-RECORD FOR TIME 0.75815182E-03
 DELETING TIME-RECORD FOR TIME 0.15221473E-02
 DELETING TIME-RECORD FOR TIME 0.30501384E-02
 DELETING TIME-RECORD FOR TIME 0.61061205E-02
 DELETING TIME-RECORD FOR TIME 0.12218085E-01
 DELETING TIME-RECORD FOR TIME 0.24442013E-01
 DELETING TIME-RECORD FOR TIME 0.48889870E-01
 DELETING TIME-RECORD FOR TIME 0.97785584E-01
 DELETING TIME-RECORD FOR TIME 0.19557701
 DELETING TIME-RECORD FOR TIME 0.39115987
 DELETING TIME-RECORD FOR TIME 0.78232558
 DELETING TIME-RECORD FOR TIME 1.5646570
 DELETING TIME-RECORD FOR TIME 3.1293199
 DELETING TIME-RECORD FOR TIME 6.2586455
 DELETING TIME-RECORD FOR TIME 12.517297
 DELETING TIME-RECORD FOR TIME 25.034600
 DELETING TIME-RECORD FOR TIME 50.069205
 DELETING TIME-RECORD FOR TIME 100.13842
 DELETING TIME-RECORD FOR TIME 200.27684
 DELETING TIME-RECORD FOR TIME 400.55368
 DELETING TIME-RECORD FOR TIME 801.10737
 DELETING TIME-RECORD FOR TIME 1602.2148
 DELETING TIME-RECORD FOR TIME 3204.4295
 DELETING TIME-RECORD FOR TIME 6408.8590
 DELETING TIME-RECORD FOR TIME 12817.718
 DELETING TIME-RECORD FOR TIME 25635.436
 DELETING TIME-RECORD FOR TIME 51270.872
 DELETING TIME-RECORD FOR TIME 102541.74
 DELETING TIME-RECORD FOR TIME 205083.49
 DELETING TIME-RECORD FOR TIME 410166.98
 DELETING TIME-RECORD FOR TIME 820333.96
 DELETING TIME-RECORD FOR TIME 1640667.9
 DELETING TIME-RECORD FOR TIME 3281335.8
 DELETING TIME-RECORD FOR TIME 6562671.6
 DELETING TIME-RECORD FOR TIME 13125343.
 DELETING TIME-RECORD FOR TIME 26250687.
 DELETING TIME-RECORD FOR TIME 52501373.
 DELETING TIME-RECORD FOR TIME 0.10500275E+09
 DELETING TIME-RECORD FOR TIME 0.20500275E+09
 DELETING TIME-RECORD FOR TIME 0.30500275E+09
 DELETING TIME-RECORD FOR TIME 0.40500275E+09
 DELETING TIME-RECORD FOR TIME 0.50500275E+09
 DELETING TIME-RECORD FOR TIME 0.60500275E+09
 DELETING TIME-RECORD FOR TIME 0.70500275E+09
 DELETING TIME-RECORD FOR TIME 0.80500275E+09

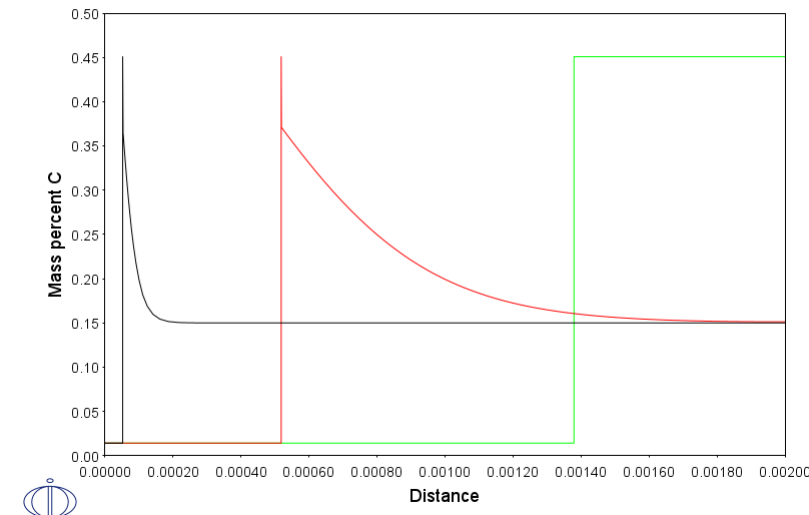
KEEPING TIME-RECORD FOR TIME 0.90500275E+09
 AND FOR TIME 0.10000000E+10
 WORKSPACE RECLAIMED

TIMESTEP AT 0.100000000E+10 SELECTED

DIC>
DIC>
DIC>
DIC> @@
DIC> @@ THE SIMULATION IS FINISHED
DIC> @@
DIC>
DIC> set-inter
--OK--
DIC>

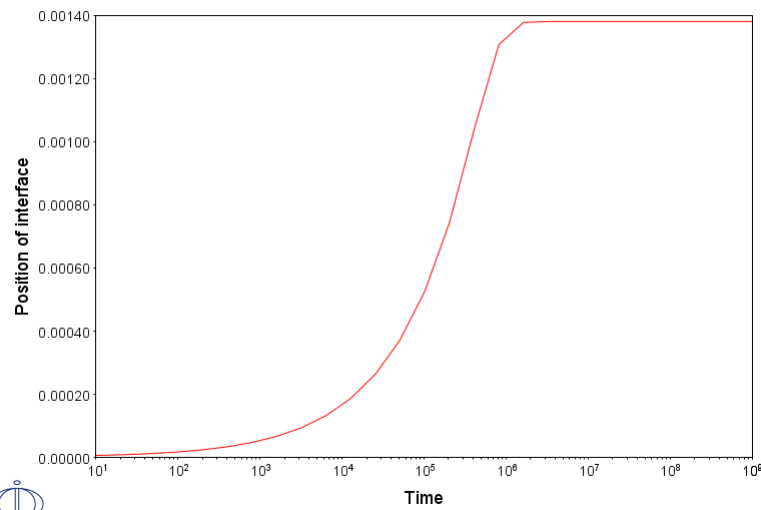
exb1b-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb1b\plot.DCM.test"
DIC>
DIC>
DIC> @@ exb1b_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE b1b
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 1.00000E+09
DIC> read exb1b
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1:
POST-1: @@
POST-1: @@ PLOT THE CARBON CONCENTRATIONS AT DIFFERENT TIMES
POST-1: @@
POST-1: s-d-a y w-p c
POST-1: s-d-a x dist glob
INFO: Distance is set as independent variable
POST-1: s-p-c time 1e3,1e5,1e9
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.09.57.08
Time=1000,100000,1E+09
CELL #1
```



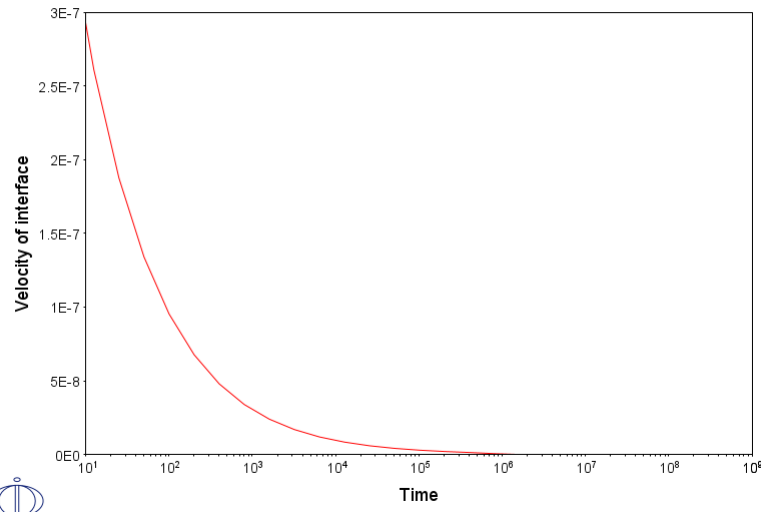
```
POST-1:
POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: @@
POST-1: @@ PLOT THE POSITION OF THE BCC/FCC INTERPHASE
POST-1: @@
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-d-a y
VARIABLE : pos
INTERFACE : aus
UPPER OR LOWER INTERFACE OF REGION AUSTENITE#1 /LOWER/: lower
POST-1:
POST-1: set_axis_type
AXIS (X, Y OR Z) : x
AXIS TYPE /LINEAR/: log
POST-1:
POST-1: s-s-s
AXIS (X, Y OR Z) : x
AUTOMATIC SCALING (Y OR N) /N/: n
MIN VALUE : 10
MAX VALUE : 1e9
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...          OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

2021.05.13.09.57.10
LOWER INTERFACE OF REGION "AUSTENITE#1"
CELL #1



```
POST-1:  
POST-1:  
POST-1:  
POST-1:  
POST-1:Hit RETURN to continue  
POST-1:  
POST-1: @@  
POST-1: @@ PLOT THE VELOCITY OF THE BCC/FCC INTERPHASE  
POST-1: @@  
POST-1: s-d-a  
AXIS (X, Y OR Z) : y  
VARIABLE : velocity  
INTERFACE : aus  
UPPER OR LOWER INTERFACE OF REGION AUSTENITE#1 /LOWER/: lower  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ...          OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

2021.05.13.09.57.12
LOWER INTERFACE OF REGION "AUSTENITE#1"
CELL #1



```
POST-1:  
POST-1:  
POST-1:  
POST-1:  
POST-1:Hit RETURN to continue  
POST-1:  
POST-1: set-inter  
--OK--  
POST-1:
```



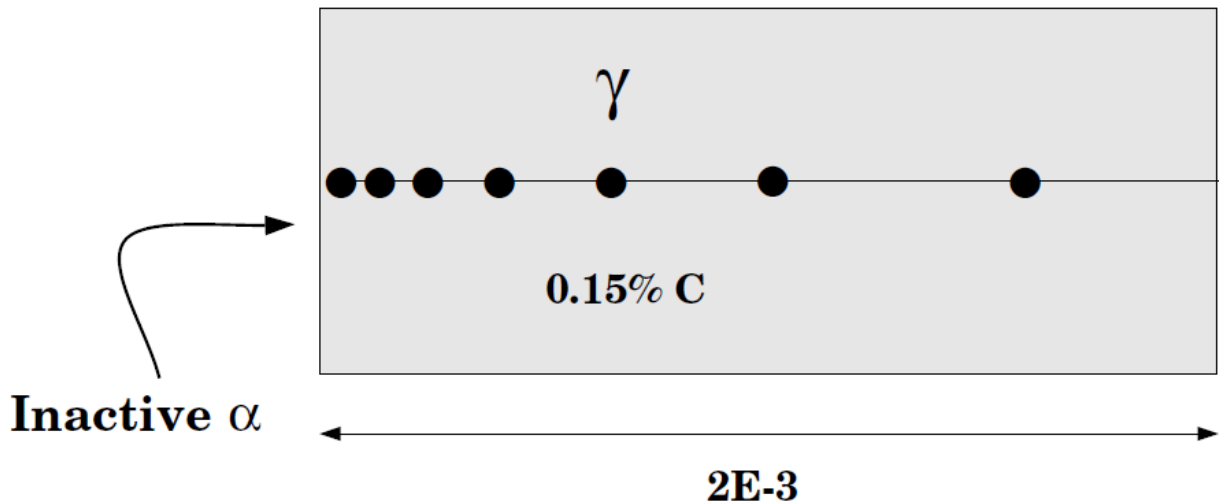
Example exb1c

γ to α transformation in a binary Fe-C alloy: Gradual cool down to 1050 K

This is the same example as in exb1a and exb1b but now the simulation starts at a higher temperature and assumes a gradual cooling down to 1050 K. When 1050 K is reached, the temperature is kept constant and thus has an isothermal transformation. As in exb1b, ferrite is in an inactive phase adjacent to the initial austenite.

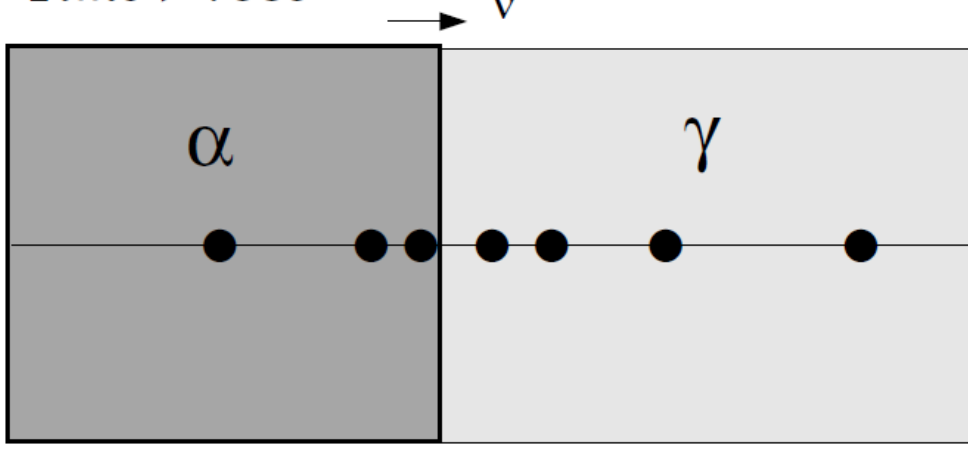
Time = 0

T = 1173 - Time * 0.1667



Time > 738s

T = 1050 K



2E-3

exblc-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exblc\setup.DCM.test"
SYS: @@
SYS: @@ Moving boundary problem.
SYS: @@ Austenite to ferrite transformation in a binary Fe-C alloy
SYS: @@ This is the same example as in exbla and exlbl but now the
SYS: @@ simulation starts at a higher temperature and assumes a gradual cooling
SYS: @@ down to 1050 K.
SYS: @@
SYS: @@ When 1050 K is reached, the temperature is kept constant and thus has an
SYS: @@ isothermal transformation. As in exlbl ferrite is an inactive
SYS: @@ phase adjacent to the initial austenite.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /* DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: @@
TDB_TCFE11: sw FEDEMO
Current database: Iron Demo Database v4.0

VA           /* DEFINED
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_FEDEMO: @@
TDB_FEDEMO: def-sys fe c
FE           C DEFINED
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_FEDEMO: @@
TDB_FEDEMO: rej ph * all
GAS:G          LIQUID:L          BCC_A2
LAVES_PHASE_C14    Cbcc_A12        CEMENTITE
CUB_A13         DIAMOND_FCC_A4   FCC_A1
GRAPHITE        HCP_A3          KSI_CARBIDE
M23C6          M5C2            M7C3
REJECTED
TDB_FEDEMO: res ph fcc bcc
FCC_A1          BCC_A2 RESTORED
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_FEDEMO: @@
TDB_FEDEMO: get
09:58:29,922 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'P. Franke, estimated parameter within SGTE, 2007; Fe-C, Ni-C, Mo-C, C-Mn'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'

-OK-
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
```

TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2
TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
SNOB1 = SGTE Noble Metal Alloys Database v1.2
STBC2 = SGTE Thermal Barrier Coating TDB v2.2
STBC1 = SGTE Thermal Barrier Coating TDB v1.1
SALT1 = SGTE Molten Salts Database v1.3
SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2 = TC Semi-Conductors v2.1
SLAG4 = Fe-containing Slag v4.1
SLAG3 = Fe-containing Slag v3.2
SLAG2 = Fe-containing Slag v2.2
SLAG1 = Fe-containing Slag v1.2
TCOX11 = Metal Oxide Solutions v11.0
TCOX10 = Metal Oxide Solutions v10.1
TCOX9 = Metal Oxide Solutions v9.0
TCOX8 = Metal Oxide Solutions v8.0
TCOX7 = Metal Oxide Solutions v7.0
TCOX6 = Metal Oxide Solutions v6.0
TCOX5 = Metal Oxide Solutions v5.1
TCOX4 = Metal Oxide Solutions v4.1
ION3 = Ionic Solutions v3.0
ION2 = Ionic Solutions v2.6
ION1 = Ionic Solutions v1.5
NOX2 = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4 = Solder Alloys v4.0
TCSLD3 = Solder Alloys v3.3
TCSLD2 = Solder Alloys v2.0
TCSLD1 = Solder Alloys v1.1
TCSI1 = Ultrapure Silicon v1.2
TCPMP2 = Materials Processing v2.5
TCES1 = Combustion/Sintering v1.1
TCSC1 = Super Conductor v1.0
TCFC1 = SOFC Database v1.0
TCNF2 = Nuclear Fuels v2.1b
NUMT2 = Nuclear Materials v2.1
NUOX4 = Nuclear Oxides v4.2
NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephista-19
MEPH15 = IRSN Mephista-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQ52 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SLDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.2
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3
MOBFE1 = Steels/Fe-Alloys Mobility v1.1
MOBFE2 = Steels/Fe-Alloys Mobility v2.0
MOBFE3 = Steels/Fe-Alloys Mobility v3.0
MOBFE4 = Steels/Fe-Alloys Mobility v4.0
MOBFE5 = Steels/Fe-Alloys Mobility v5.0
MOBFE6 = Steels/Fe-Alloys Mobility v6.0
MOBN15 = Ni-Alloys Mobility v5.1
MOBN14 = Ni-Alloys Mobility v4.1
MOBN13 = Ni-Alloys Mobility v3.2

```

MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBS11 = Si-Alloys Mobility v1.0
MOBSD1 = Solder-Alloys Mobility v1.1
MOBTI4 = Ti-Alloys Mobility v4.0
MOBTI3 = Ti-Alloys Mobility v3.1
MOBTI2 = Ti-Alloys Mobility v2.0
MOBTI1 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

```

DATABASE NAME /FEDEMO/: MFEDEMO

Current database: Fe-Alloys Mobility demo database v2.0

```

VA DEFINED
APP: def-sys fe c
FE C DEFINED
APP: rej ph * all FCC_A1 REJECTED
BCC_A2 FCC_A1 RESTORED
APP: res ph fcc bcc
FCC_A1 BCC_A2 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
FUNCTIONS .....

```

List of references for assessed data

```

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Z. Metallkunde 85(1994)498-501; C and N diffusion in bcc Cr
-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'

```

-OK-

```

APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER GLOBAL CONDITION T
DIC> @@
DIC> @@ ASSUME THAT THE COOLING RATE IS 10K/MINUTE DOWN TO 1050K
DIC> @@
DIC> set-cond
GLOBAL OR BOUNDARY CONDITION /GLOBAL/: glob
VARIABLE : T
LOW TIME LIMIT /0/: 0
T(TIME,X)= 1173-time*0.1667;
HIGH TIME LIMIT /*/: 738
ANY MORE RANGES /N/: y
T(TIME,X)= 1050;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
DIC>
DIC> @@
DIC> @@ START BY ENTERING THE REGION austenite WHERE WE PUT THE fcc PHASE
DIC> @@
DIC> enter-region
REGION NAME : austenite
DIC>
DIC> @@
DIC> @@ ENTER THE GRID INTO THE REGION
DIC> @@
DIC> enter-grid
REGION NAME : /AUSTENITE/: austenite
WIDTH OF REGION /1/: 20e-4
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER THE active PHASES INTO THE REGIONS
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUSTENITE/: austenite
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC> @@
DIC> @@ ENTER THE inactive PHASES INTO THE REGIONS
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /AUSTENITE/: austenite
ATTACHED TO THE RIGHT OF AUSTENITE /YES/: no
PHASE NAME: /NONE/: bcc
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-5
CONDITION TYPE /CLOSED_SYSTEM/: closed

```

```
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITION FOR FCC
DIC> @@
DIC> enter-composition
REGION NAME : /AUSTENITE/: austenite
PHASE NAME: /FCC_A1/: fcc#1
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: C
TYPE /LINEAR/: lin
VALUE OF FIRST POINT : 0.15
VALUE OF LAST POINT : /0.15/: 0.15
DIC>
DIC> @@
DIC> @@ THE BOUNDARY CONDITION IS A CLOSED SYSTEM AS WE DO NOT SPECIFY
DIC> @@ ANYTHING ELSE.
DIC> @@
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 738
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /73.8/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC>
DIC> @@
DIC> @@ IMPLICIT (1) TIME INTEGRATION IS USED INSTEAD OF THE MORE ACCURATE
DIC> @@ (BUT LESS STABLE) TRAPEZOIDAL METHOD WHICH IS THE DEFAULT.
DIC> @@
DIC> s-s-c
NS01A PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA Timesteps IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/:
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/:
DEGREE OF IMPURITY WHEN INTEGRATING PDES (AUTO, 0 -> 0.5 -> 1): /AUTO/: 1.0
MAX TIMESTEP CHANGE PER Timestep : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/:
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exblc Y
DIC>
DIC> set-inter
--OK--
DIC>
```

exb1c-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb1c\run.DCM.test"
DIC>
DIC>
DIC> @@ exb1c_run.DCM
DIC>
DIC> @@ FILE FOR RUNNING EXAMPLE exb1c
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exb1c
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim
Region: AUSTENITE
single geometric dense at 0.0000
1.2084 101
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.24660033E-05 DT = 0.23660033E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.71980098E-05 DT = 0.47320065E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.16662023E-04 DT = 0.94640131E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.35590049E-04 DT = 0.18928026E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.73446101E-04 DT = 0.37856052E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.14915821E-03 DT = 0.75712105E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.30058242E-03 DT = 0.15142421E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.60343083E-03 DT = 0.30284842E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.12091277E-02 DT = 0.60569684E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.24205213E-02 DT = 0.12113937E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.48433087E-02 DT = 0.24227873E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.9688834E-02 DT = 0.48455747E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.19380033E-01 DT = 0.96911494E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.38762332E-01 DT = 0.19382299E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383108 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.77526929E-01 DT = 0.38764598E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383109 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.15505612 DT = 0.77529195E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383109 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.31011451 DT = 0.15505839 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383109 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 0.62023129 DT = 0.31011678 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383109 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 1.2404649 DT = 0.62023356 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383109 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 2.4809320 DT = 1.2404671 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383109 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 4.9618662 DT = 2.4809342 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383109 FE = 1
```

```

TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 9.9237347 DT = 4.9618685 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0069849591638311 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 19.847472 DT = 9.9237370 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383107 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 39.694946 DT = 19.847474 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383105 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 76.594946 DT = 36.900000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00698495916383098 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds

output ignored...

... output resumed

3.603227678061542E-005 3.604114340896829E-005 5.596528604264911E-003
033 TIME = 0.43290458E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.55965286E-32
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.67782292E-16 AND 0.67782292E-16
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13794411E-02
U-FRACTION IN SYSTEM: C = .00698601770553276 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
5 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_BCC_A2

CPU time used in timestep 0 seconds
1.346741670350485E-005 1.346841105784895E-005 1.360472343084249E-018
018 TIME = 0.53290458E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.13604723E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.11351159E-15 AND -0.11351159E-15
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13794298E-02
U-FRACTION IN SYSTEM: C = .00698613345760261 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

CPU time used in timestep 0 seconds
1.337421555810630E-005 1.337587500157234E-005 1.528234819010838E-027
027 TIME = 0.63290458E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.15282348E-26
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.68073223E-16 AND 0.68073223E-16
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13794366E-02
U-FRACTION IN SYSTEM: C = .00698606404075852 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
4 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_BCC_A2

CPU time used in timestep 0 seconds
7.843152332779303E-006 7.843914423743844E-006 5.293114704408065E-032
032 TIME = 0.73290458E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.52931147E-31
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.70606021E-16 AND -0.70606021E-16
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13794295E-02
U-FRACTION IN SYSTEM: C = .00698613604039153 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
4.483924980437298E-006 4.484522648424397E-006 1.136218712246799E-021
021 TIME = 0.83290458E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.11362187E-20
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.34633176E-16 AND 0.34633176E-16
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13794330E-02
U-FRACTION IN SYSTEM: C = .00698610072363056 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
2 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_BCC_A2

CPU time used in timestep 0 seconds
2.726925031826145E-006 2.724639367223028E-006 2.225652791982499E-030
030 TIME = 0.93290458E+09 DT = 0.10000000E+09 SUM OF SQUARES = 0.22256528E-29
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.42037398E-16 AND 0.42037398E-16
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13794372E-02
U-FRACTION IN SYSTEM: C = .00698605785650249 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
2 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_BCC_A2

CPU time used in timestep 0 seconds
3.315675851048835E-004 3.315981840536485E-004 3.966614047191010E-028
028 TIME = 0.10000000E+10 DT = 67095423. SUM OF SQUARES = 0.39666140E-27
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.48086084E-16 AND -0.48086084E-16
POSITION OF INTERFACE R_BCC_A2 / AUSTENITE IS 0.13794340E-02
U-FRACTION IN SYSTEM: C = .00698609075687852 FE = 1
TOTAL SIZE OF SYSTEM: .002 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 738.00000
DELETING TIME-RECORD FOR TIME 769.68675
DELETING TIME-RECORD FOR TIME 833.06026
DELETING TIME-RECORD FOR TIME 959.80726
DELETING TIME-RECORD FOR TIME 1213.3013
DELETING TIME-RECORD FOR TIME 1720.2893
DELETING TIME-RECORD FOR TIME 2734.2654
DELETING TIME-RECORD FOR TIME 4762.2175
DELETING TIME-RECORD FOR TIME 8818.1217
DELETING TIME-RECORD FOR TIME 16929.930
DELETING TIME-RECORD FOR TIME 33153.547
DELETING TIME-RECORD FOR TIME 65600.781
DELETING TIME-RECORD FOR TIME 130495.25
DELETING TIME-RECORD FOR TIME 260284.19
DELETING TIME-RECORD FOR TIME 519862.06
DELETING TIME-RECORD FOR TIME 1039017.8
DELETING TIME-RECORD FOR TIME 2077329.3
DELETING TIME-RECORD FOR TIME 4153952.3
DELETING TIME-RECORD FOR TIME 8307198.2
DELETING TIME-RECORD FOR TIME 16613690.
DELETING TIME-RECORD FOR TIME 33226674.
DELETING TIME-RECORD FOR TIME 66452642.
DELETING TIME-RECORD FOR TIME 0.13290458E+09
DELETING TIME-RECORD FOR TIME 0.23290458E+09
DELETING TIME-RECORD FOR TIME 0.33290458E+09
DELETING TIME-RECORD FOR TIME 0.43290458E+09
DELETING TIME-RECORD FOR TIME 0.53290458E+09
DELETING TIME-RECORD FOR TIME 0.63290458E+09
DELETING TIME-RECORD FOR TIME 0.73290458E+09

```

DELETING TIME-RECORD FOR TIME 0.83290458E+09

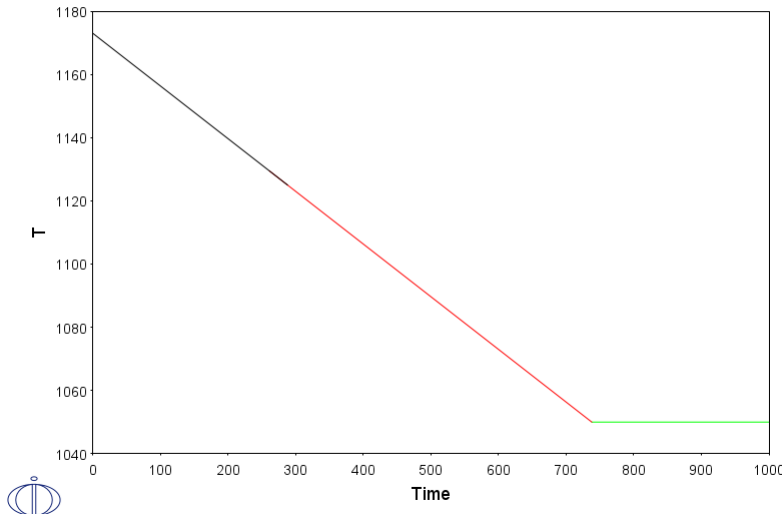
KEEPING TIME-RECORD FOR TIME 0.93290458E+09
AND FOR TIME 0.10000000E+10
WORKSPACE RECLAIMED

TIMESTEP AT 0.10000000E+10 SELECTED

DIC>
DIC>
DIC>
DIC>
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ THE SIMULATION IS FINISHED
DIC> @@
DIC>
DIC> set-inter
--OK---
DIC>

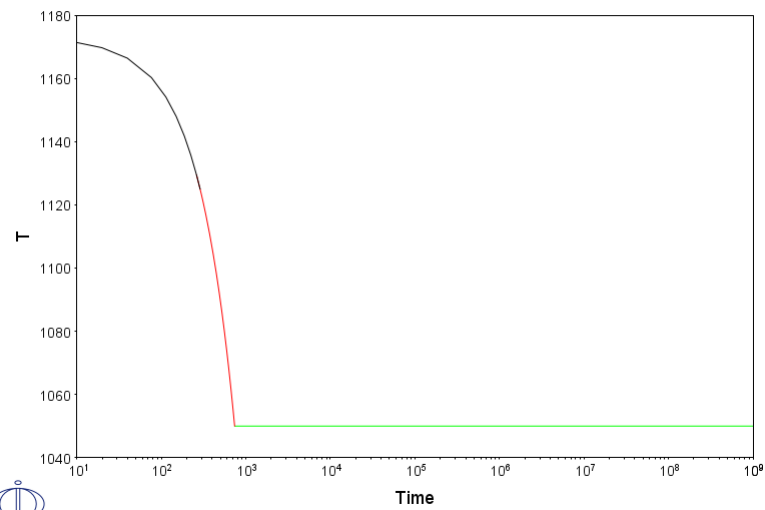
exb1c-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb1c\plot.DCM.test"
DIC>
DIC>
DIC> @@ exb1c_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE b1c
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 1.00000E+09
DIC> read exb1c
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1:
POST-1: @@
POST-1: @@ PLOT TEMPERATURE VS. TIME
POST-1: @@
POST-1: s-d-a y t
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-p-c
CONDITION /TIME/: interface
INTERFACE : austenite
UPPER OR LOWER INTERFACE OF REGION AUSTENITE#1 /LOWER/: lower
POST-1: s-s-s x n 0 1000
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.10.00.31
LOWER INTERFACE OF REGION "AUSTENITE#1"
CELL #1
```



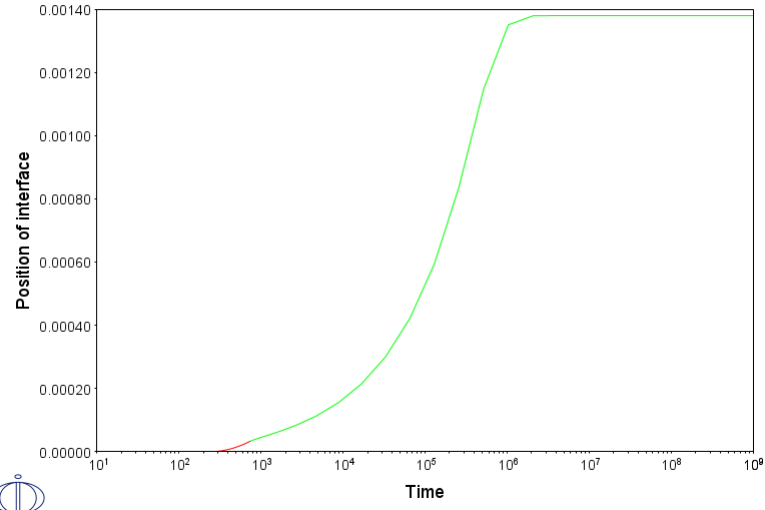
```
POST-1:
POST-1:
POST-1:
POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1:
POST-1: @@
POST-1: @@ PLOT VS. LOG TIME
POST-1: @@
POST-1: set-axis-type x log
POST-1: s-s-s x n 10 1e9
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

2021.05.13.10.00.33
LOWER INTERFACE OF REGION "AUSTENITE#1"
CELL #1



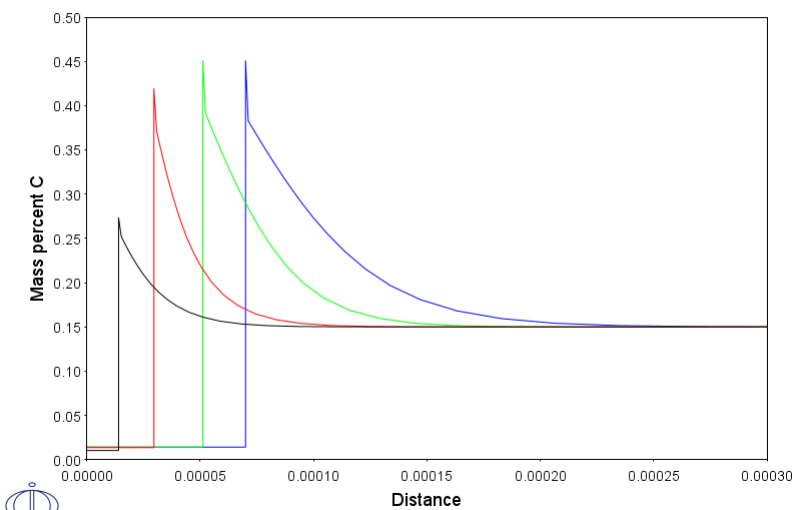
```
POST-1:  
POST-1:  
POST-1:  
POST-1:@?<Hit_return_to_continue>  
POST-1:  
POST-1: @@  
POST-1: @@ PLOT THE POSITION OF THE BCC/FCC INTERPHASE  
POST-1: @@  
POST-1: s-d-a  
AXIS (X, Y OR Z) : y  
VARIABLE : position  
INTERFACE : austenite  
UPPER OR LOWER INTERFACE OF REGION AUSTENITE#1 /LOWER/: lower  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ... OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

2021.05.13.10.00.34
LOWER INTERFACE OF REGION "AUSTENITE#1"
CELL #1



```
POST-1:  
POST-1:  
POST-1:  
POST-1:  
POST-1:@?<Hit_return_to_continue>  
POST-1:  
POST-1: @@  
POST-1: @@ PLOT THE CARBON CONCENTRATION VS. DISTANCE  
POST-1: @@  
POST-1: s-d-a y w-p c  
POST-1: s-d-a x dis glob  
INFO: Distance is set as independent variable  
POST-1: s-p-c time 500,700,1200,2000  
POST-1: set-axis-type x lin  
POST-1: s-s-s x n 0 3e-4  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ... OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

2021.05.13.10.00.35
Time = 500,700,1200,2000
CELL #1



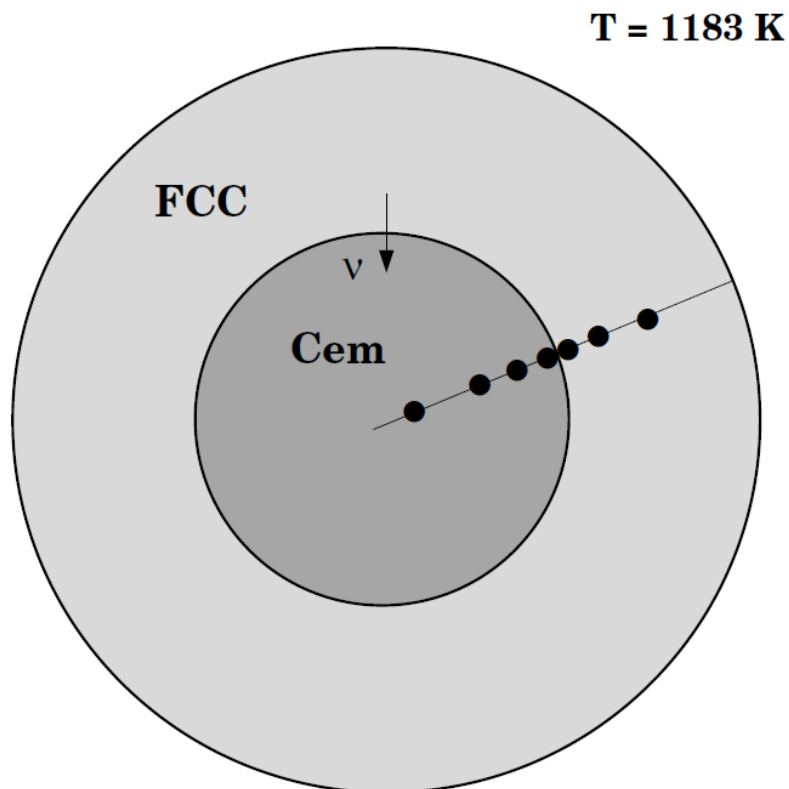
POST-1:
POST-1:
POST-1:
POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1: set-inter
--OK---
POST-1:



Example exb2

Cementite dissolution in an Fe-Cr-C alloy

This example calculates the dissolution of a spherical cementite particle in an austenite matrix. This case is from Z.-K. Liu, L. Höglund, B. Jönsson and J. Ågren (Metall. Trans.A, v.22A, 1991, pp. 1745-1752). In order to achieve the correct average composition in the calculation it is necessary to take into account the fact that the calculation is set up using the volume fraction of the phases. To calculate the initial state at the heat treatment temperature we need first to determine the state at the normalizing temperature. To calculate the volume fraction of the phases we need to enter a number of functions that calculate these quantities.



Koermann, T. Hickel, J. Neugebauer, CALPHAD, 34, 129 -33(2010); Fe-C'
 'J. Bratberg, Z. Metallkd., 96 (2005) 335-344; Fe-Cr-Mo-C'
 'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
 'R. Naraghi, Thermo-Calc Software AB, Sweden, 2016; FCC Fe-Cr-C and C-Cr-Ni'
 -OK-

TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE THE MOBILITY DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app mobfe4
 Current database: Steels/Fe-Alloys Mobility v4.0

```

VA DEFINED
B2_BCC REJECTED
APP: def-sp fe cr c
FE CR C
DEFINED
APP: rej ph * all
BCC_A2 CEMENTITE FCC_A1
FE4N_LP1 HCP_A3 LIQUID:L
REJECTED
APP: res ph fcc cementite
FCC_A1 CEMENTITE RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
FUNCTIONS .....

```

List of references for assessed data

```

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Z. Metallkunde 85(1994)502-509; C diffusion in fcc Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'This parameter has been estimated'
-OK-
APP:
APP: @@
APP: @@ ENTER THE POLY-3 MONITOR
APP: @@
APP: go p-3

```

POLY version 3.32

POLY: @@
POLY: @@ SET THE CONDITIONS AT THE NORMALIZING TEMPERATURE

```

POLY: @@
POLY: set-cond T=1008,P=101325,N=1
POLY: set-cond X(CR)=0.0206,X(C)=0.0391
POLY:
POLY:
POLY: @@
POLY: @@ ENTER FUNCTIONS TO DETERMINE THE VOLUME-FRACTIONS
POLY: @@
POLY:
POLY: @@ Radius of the cementite particle
POLY: ent-symb var rcem=0.5255e-6;
POLY:
POLY: @@ total number of moles of substitutional components
POLY: ent-symb func nstot=n(fe)+n(cr);
POLY:
POLY: @@ number of moles of substitutional components in cementite
POLY: ent-symb func nscem=n(cem,fe)+n(cem,cr);
POLY:
POLY: @@ volume fraction (U-fraction) of cementite
POLY: ent-symb func vfcem=nscem/nstot;
POLY:
POLY: @@ total radius of the system
POLY: ent-symb func rtot=rcem/vfcem**^(1/3);
POLY:
POLY: @@ radius of the surrounding austenite matrix
POLY: ent-symb func rmat=rtot-rcem;
POLY:
POLY:
POLY: @@
POLY: @@ COMPUTE THE EQUILIBRIUM
POLY: @@
POLY: compute-eq
Using global minimization procedure
Calculated 4113 grid points in 4 s
Found the set of lowest grid points in 0 s
Calculated POLY solution 0 s, total time 4 s
POLY:
POLY:
POLY: @@
POLY: @@ SHOW THE COMPUTED VALUES TO BE USED IN THE DICTRA CALCULATION
POLY: @@
POLY: show rmat
RMAT=5.3924863E-7
POLY: show w(cem,cr),w(bcc,cr),w(bcc,c)
W(CEMENTITE,CR)=0.12581645
W(BCC_A2,CR)=4.4332285E-3
W(BCC_A2,C)=1.510215E-4
POLY:
POLY: ent var wmatcr=w(bcc,cr);
POLY: ent var wmatc=w(bcc,c);
POLY: ent var wcemcr=w(cem,cr);
POLY:
POLY: @@
POLY: @@ ENTER THE DICTRA MONITOR
POLY: @@
POLY: go d-m
NO TIME STEP DEFINED
*** ENTERING BCC_A2 AS A DIFFUSION NONE PHASE
DIC>
DIC> @@

```

```

DIC> @@ ENTER GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob t 0 1183; * n
DIC> @@
DIC> @@ ENTER THE REGIONS carb AND aus
DIC> @@
DIC> enter-region
REGION NAME : carb
DIC>
DIC> enter-region
REGION NAME : aus
ATTACH TO REGION NAMED /CARB/:
ATTACHED TO THE RIGHT OF CARB /YES/:
DIC> @@
DIC> @@ ENTER GEOMTRICAL GRIDS INTO THE REGIONS
DIC> @@
DIC> @@
DIC> @@ THE INITIAL SIZE OF THE CEMENTITE PARTICLE IS ASSUMED TO BE KNOWN
DIC> @@ (IN THIS CASE THE VALUE IS TAKEN FROM LIU ET AL. WHO ESTIMATED THE
DIC> @@ AVERAGE INITIAL DIAMETER OF THE PARTICLES TO 1.051E-6 METERS).
DIC> @@
DIC> enter-grid
REGION NAME : /CARB/: carb
WIDTH OF REGION /1/: rcm
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ THE SIZE OF THE FCC REGION CAN BE CALCULATED FROM A MASS BALANCE
DIC> @@ AFTER ESTIMATING THE INITIAL COMPOSITIONS IN THE TWO PHASES
DIC> @@
DIC> enter-grid
REGION NAME : /AUS/: aus
WIDTH OF REGION /1/: rmat
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER PHASES INTO REGIONS
DIC> @@
DIC> enter-phase act carb matrix cementite
DIC> enter-phase act aus matrix fcc#1
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITIONS IN THE PHASES
DIC> @@
DIC> enter-composition
REGION NAME : /CARB/: carb
PHASE NAME: /CEMENTITE/: cementite
COMPOSITION TYPE /MOLE_FRACTION/: weig-fraction
PROFILE FOR /CR/: cr lin wcemcr wcemcr
DIC>
DIC> enter-composition
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1#1/: fcc#1
DEPENDENT COMPONENT ? /FE/: fe
COMPOSITION TYPE /MOLE_FRACTION/: weig-fraction
PROFILE FOR /C/: CR lin wmatcr wmatcr
PROFILE FOR /CR/: C lin wmatc wmatc
DIC>
DIC> @@
DIC> @@ SET TO A SPHERICAL GEOMETRY
DIC> @@
DIC> enter-geo
GEOOMETRICAL EXPONENT /0/: 2
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME AND OTHER SIMULATION PARAMETERS
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /1/: 10000
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /1000/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exb2 Y
DIC>
DIC> set-inter
--OK---
DIC>

```

exb2-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb2\run.DCM.test"
DIC>
DIC>
DIC> @@ exb2_run.DCM
DIC>
DIC>
DIC> @@ READ THE SET UP FROM FILE AND START THE SIMULATION
DIC> @@
DIC>
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
*** ENTERING BCC_A2 AS A DIFFUSION NONE PHASE
DIC> read exb2
OK
DIC> sim
Region: CARB
single geometric dense at 0.52550E-06
0.85084 96
Region: AUS
single geometric dense at 0.0000
1.0072 63
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
Trying old scheme 4
GENERATING STARTING VALUES FOR CELL # 1 INTERFACE # 2
DETERMINING INITIAL EQUILIBRIUM VALUES
CALCULATING STARTING VALUES: 9 EQUILIBRIUM CALCULATIONS DONE 6 OUT OF 9
04
U-FRACTION IN SYSTEM: C = .0406910188418179 CR = .0214382349908298
FE = .978561765139677
TOTAL SIZE OF SYSTEM: 5.0562655986E-18 [m^3]
U-FRACTION IN SYSTEM: C = .0406910188418179 CR = .0214382349908298
FE = .978561765139677
TOTAL SIZE OF SYSTEM: 5.0562655986E-18 [m^3]
0.175650510569693 0.175691773365777 0.175650565704887 2.957532598305099E-003 7.754437973902466E-
005 5.789560110638669E-006 8.570927948261712E-007 3.747773449054368E-007 3.453239602567087E-
007 1.349945520965789E-007 1.350781378777377E-007 1.346761780040160E-007 1.361219741567956E-
007 1.343486158187189E-007 1.340757896497654E-007 1.335307279086690E-007 1.344901657987451E-
007 1.324431149502718E-007 1.302779707752355E-007 1.259882989092764E-007 1.269270676242141E-
007 1.175749112140888E-007 1.014430318623444E-007 7.21795337839238E-008 7.299680976072143E-
008 2.72813525993651E-008 7.241610557431686E-011 4.089073975800423E-014 7.514683315310822E-
019 TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.75146833E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.39436721E-02 AND -0.39436721E-02
POSITION OF INTERFACE CARB / AUS IS 0.52510563E-06
U-FRACTION IN SYSTEM: C = .040729015482772 CR = .0214383281162587
FE = .978561672014248
TOTAL SIZE OF SYSTEM: 5.0562655986E-18 [m^3]
15 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARB

CPU time used in timestep 1 seconds
2.248325209321158E-005 2.248775172894889E-005 2.248323619911258E-005 1.887681971785783E-009 1.633889346556181E-
009 1.019554384038724E-009 6.882195845839138E-010 6.885275294802593E-010 2.216755169621960E-
010 6.151372771543853E-014 2.128400055073508E-017 TIME = 0.3000000E-06 DT = 0.2000000E-
06 SUM OF SQUARES = 0.21284001E-16
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.22443437E-04 AND -0.22443437E-04
POSITION OF INTERFACE CARB / AUS IS 0.52510114E-06
U-FRACTION IN SYSTEM: C = .04072976539935 CR = .0214383291134799
FE = .978561671017027
TOTAL SIZE OF SYSTEM: 5.0562655986E-18 [m^3]
CPU time used in timestep 0 seconds
6.026886171006244E-004 6.026583061736450E-004 6.026723276374160E-004 8.498836241942660E-005 3.515902363183597E-
005 2.728069475198861E-007 4.04929651693932E-010 6.350486825359254E-015 2.88154926615981E-
018 TIME = 0.7000000E-06 DT = 0.4000000E-06 SUM OF SQUARES = 0.28815493E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.46375286E-05 AND -0.46375286E-05
POSITION OF INTERFACE CARB / AUS IS 0.52509929E-06
U-FRACTION IN SYSTEM: C = .0407315536943019 CR = .0214383294273154
FE = .978561670703191
TOTAL SIZE OF SYSTEM: 5.0562655986E-18 [m^3]
CPU time used in timestep 0 seconds
2.365053654994453E-004 2.365034011193425E-004 2.364987464518160E-004 7.190707106424921E-005 1.444190616366121E-
007 4.667910968230499E-011 2.104087464936592E-014 2.043436109102669E-017 TIME = 0.1500000E-05 DT = 0.8000000E-
06 SUM OF SQUARES = 0.20434361E-16
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.40628788E-05 AND -0.40628788E-05
POSITION OF INTERFACE CARB / AUS IS 0.52509604E-06
U-FRACTION IN SYSTEM: C = .0407314707249636 CR = .0214383297893152
FE = .978561670341192
TOTAL SIZE OF SYSTEM: 5.0562655986E-18 [m^3]

output ignored...

... output resumed

2.625780106432001E-009 5.088645634293734E-014 1.23711967778825E-
018 TIME = 8301.5634 DT = 1000.0000 SUM OF SQUARES = 0.12371197E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.10714303E-10 AND -0.10714303E-10
POSITION OF INTERFACE CARB / AUS IS 0.24630884E-06
U-FRACTION IN SYSTEM: C = .0407634546383461 CR = .021438390755656
FE = .978561609374851
TOTAL SIZE OF SYSTEM: 5.0562655986E-18 [m^3]
CPU time used in timestep 1 seconds
8.020743552166865E-009 8.075927620319851E-009 8.037806988719298E-009 2.656775571773380E-009 9.146648175368995E-
010 6.108854557489235E-015 5.066615456766682E-
018 TIME = 9301.5634 DT = 1000.0000 SUM OF SQUARES = 0.50666155E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.10120591E-10 AND -0.10120591E-10
POSITION OF INTERFACE CARB / AUS IS 0.23618825E-06
U-FRACTION IN SYSTEM: C = .0407634719546566 CR = .0214383896298978
FE = .978561610500609
TOTAL SIZE OF SYSTEM: 5.0562655986E-18 [m^3]
CPU time used in timestep 1 seconds
7.715606069317417E-007 7.711461084002090E-007 7.708252905711602E-007 6.733834081453556E-008 2.579683533447208E-
012 1.675208725553256E-017 TIME = 10000.000 DT = 698.43661 SUM OF SQUARES = 0.16752087E-16
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.95749100E-11 AND -0.95749100E-11
POSITION OF INTERFACE CARB / AUS IS 0.22950079E-06
U-FRACTION IN SYSTEM: C = .0407634715905315 CR = .021438388875538
FE = .978561611242953
TOTAL SIZE OF SYSTEM: 5.0562655986E-18 [m^3]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
```

RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 0.30000000E-06
DELETING TIME-RECORD FOR TIME 0.70000000E-06
DELETING TIME-RECORD FOR TIME 0.15000000E-05
DELETING TIME-RECORD FOR TIME 0.31000000E-05
DELETING TIME-RECORD FOR TIME 0.63000000E-05
DELETING TIME-RECORD FOR TIME 0.12700000E-04
DELETING TIME-RECORD FOR TIME 0.25500000E-04
DELETING TIME-RECORD FOR TIME 0.51100000E-04
DELETING TIME-RECORD FOR TIME 0.99418841E-04
DELETING TIME-RECORD FOR TIME 0.17900385E-03
DELETING TIME-RECORD FOR TIME 0.30071417E-03
DELETING TIME-RECORD FOR TIME 0.47831765E-03
DELETING TIME-RECORD FOR TIME 0.72934329E-03
DELETING TIME-RECORD FOR TIME 0.10758312E-02
DELETING TIME-RECORD FOR TIME 0.15428938E-02
DELETING TIME-RECORD FOR TIME 0.21467122E-02
DELETING TIME-RECORD FOR TIME 0.29017133E-02
DELETING TIME-RECORD FOR TIME 0.38194432E-02
DELETING TIME-RECORD FOR TIME 0.49085135E-02
DELETING TIME-RECORD FOR TIME 0.61737016E-02
DELETING TIME-RECORD FOR TIME 0.76140450E-02
DELETING TIME-RECORD FOR TIME 0.92225149E-02
DELETING TIME-RECORD FOR TIME 0.10986151E-01
DELETING TIME-RECORD FOR TIME 0.12887549E-01
DELETING TIME-RECORD FOR TIME 0.14913628E-01
DELETING TIME-RECORD FOR TIME 0.17051229E-01
DELETING TIME-RECORD FOR TIME 0.19290844E-01
DELETING TIME-RECORD FOR TIME 0.21633465E-01
DELETING TIME-RECORD FOR TIME 0.24076421E-01
DELETING TIME-RECORD FOR TIME 0.26619478E-01
DELETING TIME-RECORD FOR TIME 0.29264709E-01
DELETING TIME-RECORD FOR TIME 0.32010278E-01
DELETING TIME-RECORD FOR TIME 0.34862213E-01
DELETING TIME-RECORD FOR TIME 0.37829432E-01
DELETING TIME-RECORD FOR TIME 0.40921190E-01
DELETING TIME-RECORD FOR TIME 0.44135912E-01
DELETING TIME-RECORD FOR TIME 0.47480207E-01
DELETING TIME-RECORD FOR TIME 0.50960807E-01
DELETING TIME-RECORD FOR TIME 0.54592579E-01
DELETING TIME-RECORD FOR TIME 0.58382957E-01
DELETING TIME-RECORD FOR TIME 0.62356658E-01
DELETING TIME-RECORD FOR TIME 0.66520761E-01
DELETING TIME-RECORD FOR TIME 0.68801534E-01
DELETING TIME-RECORD FOR TIME 0.71495808E-01
DELETING TIME-RECORD FOR TIME 0.74611177E-01
DELETING TIME-RECORD FOR TIME 0.78157596E-01
DELETING TIME-RECORD FOR TIME 0.82113712E-01
DELETING TIME-RECORD FOR TIME 0.86489284E-01
DELETING TIME-RECORD FOR TIME 0.91274871E-01
DELETING TIME-RECORD FOR TIME 0.96471057E-01
DELETING TIME-RECORD FOR TIME 0.10205818
DELETING TIME-RECORD FOR TIME 0.10804601
DELETING TIME-RECORD FOR TIME 0.11444776
DELETING TIME-RECORD FOR TIME 0.12132749
DELETING TIME-RECORD FOR TIME 0.12873099
DELETING TIME-RECORD FOR TIME 0.13674515
DELETING TIME-RECORD FOR TIME 0.14543216
DELETING TIME-RECORD FOR TIME 0.15494171
DELETING TIME-RECORD FOR TIME 0.16560505
DELETING TIME-RECORD FOR TIME 0.17785673
DELETING TIME-RECORD FOR TIME 0.19238367
DELETING TIME-RECORD FOR TIME 0.21033391
DELETING TIME-RECORD FOR TIME 0.23438452
DELETING TIME-RECORD FOR TIME 0.27062133
DELETING TIME-RECORD FOR TIME 0.33531949
DELETING TIME-RECORD FOR TIME 0.41255614
DELETING TIME-RECORD FOR TIME 0.50834602
DELETING TIME-RECORD FOR TIME 0.63458019
DELETING TIME-RECORD FOR TIME 0.81556830
DELETING TIME-RECORD FOR TIME 0.91469957
DELETING TIME-RECORD FOR TIME 1.0981843
DELETING TIME-RECORD FOR TIME 1.4390940
DELETING TIME-RECORD FOR TIME 1.6570157
DELETING TIME-RECORD FOR TIME 2.0571409
DELETING TIME-RECORD FOR TIME 2.8315093
DELETING TIME-RECORD FOR TIME 4.3109112
DELETING TIME-RECORD FOR TIME 7.1209759
DELETING TIME-RECORD FOR TIME 12.538328
DELETING TIME-RECORD FOR TIME 23.254969
DELETING TIME-RECORD FOR TIME 44.688251
DELETING TIME-RECORD FOR TIME 87.554816
DELETING TIME-RECORD FOR TIME 173.28794
DELETING TIME-RECORD FOR TIME 344.75420
DELETING TIME-RECORD FOR TIME 687.68671
DELETING TIME-RECORD FOR TIME 1373.5517
DELETING TIME-RECORD FOR TIME 2301.5634
DELETING TIME-RECORD FOR TIME 3301.5634
DELETING TIME-RECORD FOR TIME 4301.5634
DELETING TIME-RECORD FOR TIME 5301.5634
DELETING TIME-RECORD FOR TIME 6301.5634
DELETING TIME-RECORD FOR TIME 7301.5634
DELETING TIME-RECORD FOR TIME 8301.5634

KEEPING TIME-RECORD FOR TIME 9301.5634
AND FOR TIME 10000.000
WORKSPACE RECLAIMED

TIMESTEP AT 10000.0000 SELECTED

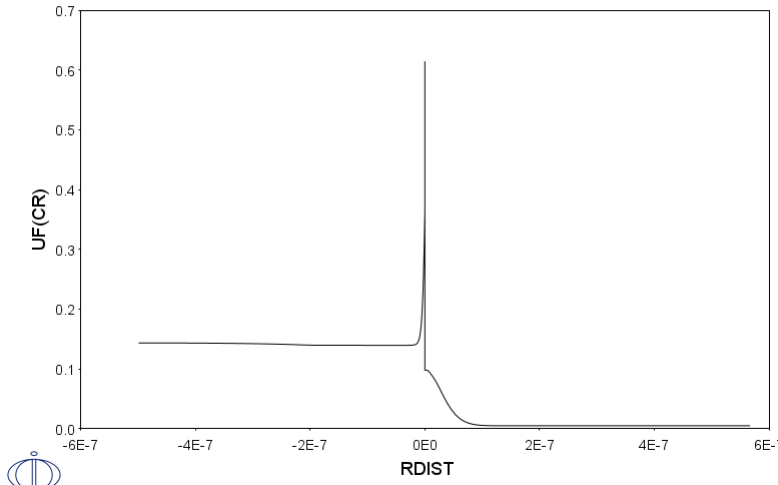
DIC>
DIC> set-inter
--OK--
DIC>

exb2-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb2\plot.DCM.test"
DIC>
DIC>
DIC> @@ exb2_plot.DCM
DIC>
DIC> @@
DIC> FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE b2
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 1.00000E+04
*** ENTERING BCC_A2 AS A DIFFUSION NONE PHASE
DIC> read exb2
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ LET US PLOT CHROMIUM CONCENTRATION PROFILES
POST-1: @@ WE THEN SET THE DISTANCE AS X-AXIS (NOTE THAT DISTANCE IS AUTOMATICALLY
POST-1: @@ SET AS THE INDEPENDENT VARIABLE) AND U-FRACTION CARBON AS Y-AXIS
POST-1: @@ REMEMBER THAT THE PLOT CONDITION ALSO MUST BE SET.
POST-1: @@
POST-1: @@ NOTICE THAT ALL DISTANCES IN THE DATA FILE ARE GIVEN RELATIVE TO THE
POST-1: @@ CEM/FCC INTERFACE. FOR THIS REASON AN OFFSET MUST BE GIVEN TO THE
POST-1: @@ DATA ACCORDING TO THE ACTUAL PARTICLE RADIUS AT THE SPECIFIED TIME.
POST-1: @@
POST-1: enter-symb
Function or table /FUNCTION/: func
NAME: rdist
FUNCTION: gd-poi(carb,u);
POST-1:
POST-1: s-d-a x rdist
POST-1:
POST-1: s-i-v
VARIABLE /TIME/: dist
DISTANCE : /GLOBAL/: glo
POST-1:
POST-1: s-d-a y uf(cr)
POST-1:
POST-1: s-p-c time 10
POST-1:
POST-1: @@
POST-1: @@ SET THE TITLE ON THE PLOT
POST-1: @@
POST-1: set-title Figure b2.1
POST-1:
POST-1: @@
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Figure b2.1

2021.05.13.10.06.59
Time = 10
CELL #1



```
POST-1:
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: @@
POST-1: @@ INCLUDE EXPERIMENTAL DATA POINTS ON THE PLOT FOR COMPARISION
POST-1: @@
POST-1: @@ FIRST LIST DATASETS
POST-1: @@
POST-1: app y exb2.exp
PROLOGUE NUMBER: /0/: 0
```

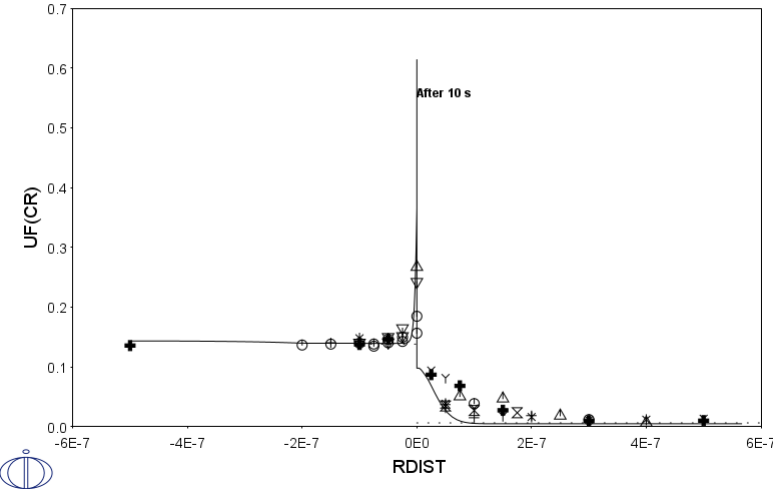
```

DATASET NUMBER(s): /-1/: -1
DATASET 1 CONCENTRATION PROFILE T=10S
DATASET 2 CONCENTRATION PROFILE T=100S
DATASET 3 CONCENTRATION PROFILE T=1000S
DATASET 4 CONCENTRATION PROFILE T=10000S
DATASET 5 VOLUME FRACTION CEMENTITE VS. TIME
DATASET 6 MEAN PARTICLE DIAMETER VS. TIME
POST-1:
POST-1: @@
POST-1: @@ SELECT THE PROPER DATASET
POST-1: @@
POST-1: app y exb2.exp
PROLOGUE NUMBER: /0/: 0
DATASET NUMBER(s): /-1/: 1
POST-1:
POST-1: set-title Figure b2.2
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

```

Figure b2.2

2021.05.13.10.07.02
Time = 10
CELL #1



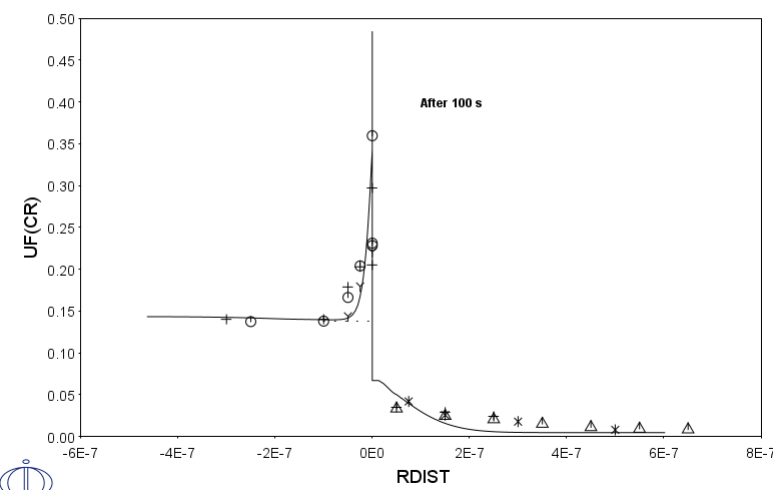
```

POST-1:
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: @@
POST-1: @@ PLOT ALSO FOR 100, 1000 AND 10000 seconds
POST-1: @@
POST-1:
POST-1: s-p-c time 100
POST-1:
POST-1: app y exb2.exp
PROLOGUE NUMBER: /0/: 0
DATASET NUMBER(s): /-1/: 2
POST-1:
POST-1: set-title Figure b2.3
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

```

Figure b2.3

2021.05.13.10.07.05
Time = 100
CELL #1



```

POST-1:
POST-1:

```

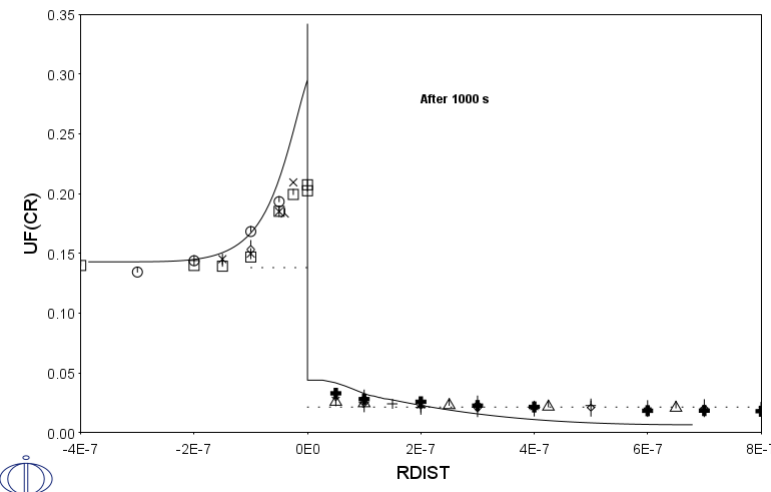
```

POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1:
POST-1: s-p-c time 1000
POST-1:
POST-1: app y exb2.exp
PROLOGUE NUMBER: /0/: 0
DATASET NUMBER(s): /-1/: 3
POST-1:
POST-1: set-title Figure b2.4
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

```

Figure b2.4

2021.05.13.10.07.09
Time = 1000
CELL #1



After 1000 s

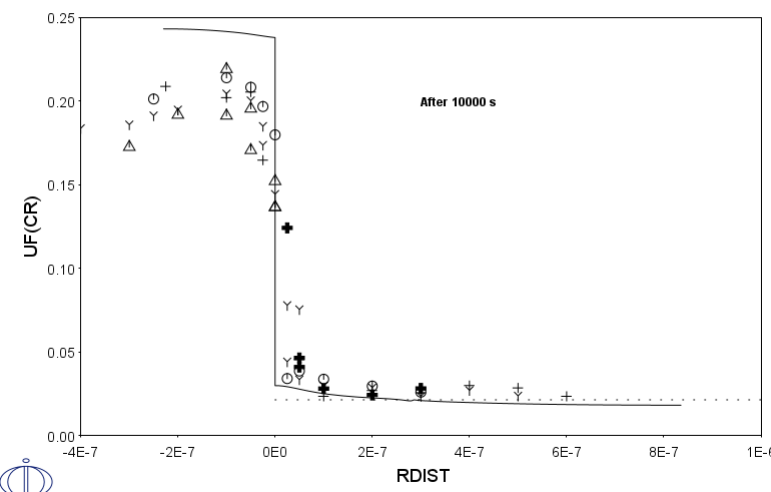
```

POST-1:
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1:
POST-1: s-p-c time 10000
POST-1:
POST-1: app y exb2.exp
PROLOGUE NUMBER: /0/: 0
DATASET NUMBER(s): /-1/: 4
POST-1:
POST-1: set-title Figure b2.5
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

```

Figure b2.5

2021.05.13.10.07.11
Time = 10000
CELL #1



After 10000 s

```

POST-1:
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: @@ ALSO PLOT HOW THE VOLUME FRACTION OF CEMENTITE VARIES
POST-1: @@ WITH TIME
POST-1: @@

```

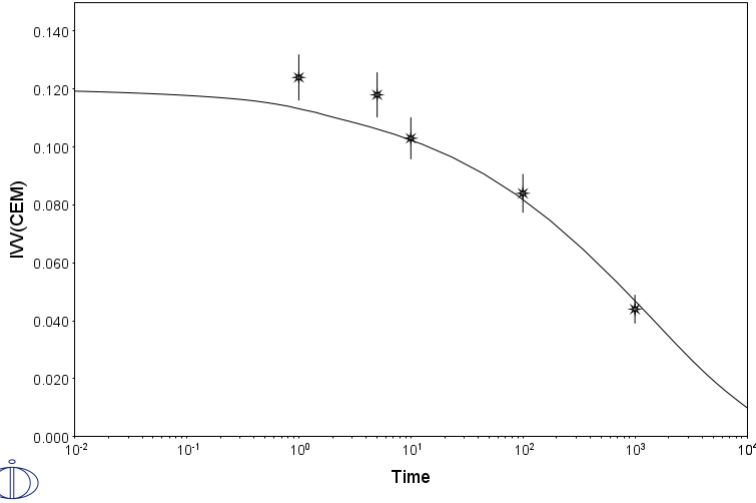
```

POST-1: s-d-a y ivv(cem)
POST-1: s-s-s y n 0 .15
POST-1:
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: set-axis-type x log
POST-1: s-s-s x n .01 10000
POST-1:
POST-1: s-p-c integral
POST-1:
POST-1: app y exb2.exp
PROLOGUE NUMBER: /0/: 0
DATASET NUMBER(s): /-1/: 5
POST-1:
POST-1: set-title Figure b2.6
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

```

Figure b2.6

2021.05.13.10.07.15
CELL #1



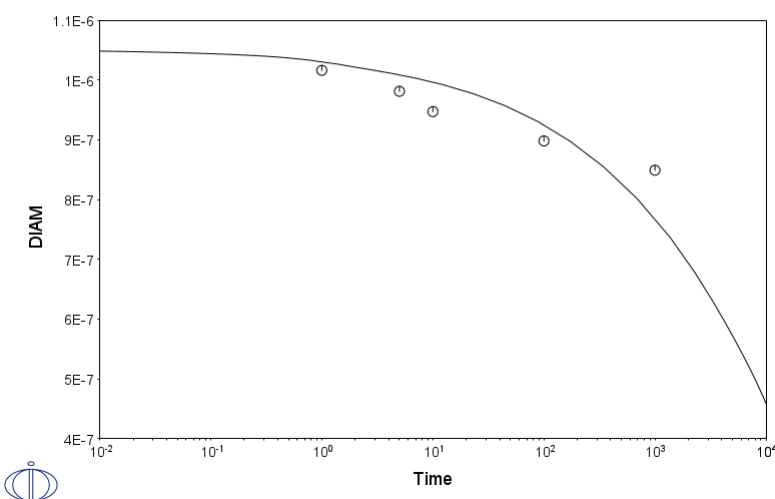
```

POST-1:
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: @@
POST-1: @@ ALSO PLOT HOW THE DIAMETER OF CEMENTITE VARIES WITH TIME
POST-1: @@
POST-1: enter func diam=2*poi(carb,u);
POST-1: s-d-a y diam
POST-1:
POST-1: s-p-c interface carb upper
POST-1:
POST-1: app y exb2.exp
PROLOGUE NUMBER: /0/: 0
DATASET NUMBER(s): /-1/: 6
POST-1:
POST-1: set-title Figure b2.7
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

```

Figure b2.7

2021.05.13.10.07.18
UPPER INTERFACE OF REGION "CARB#1"
CELL #1



POST-1:

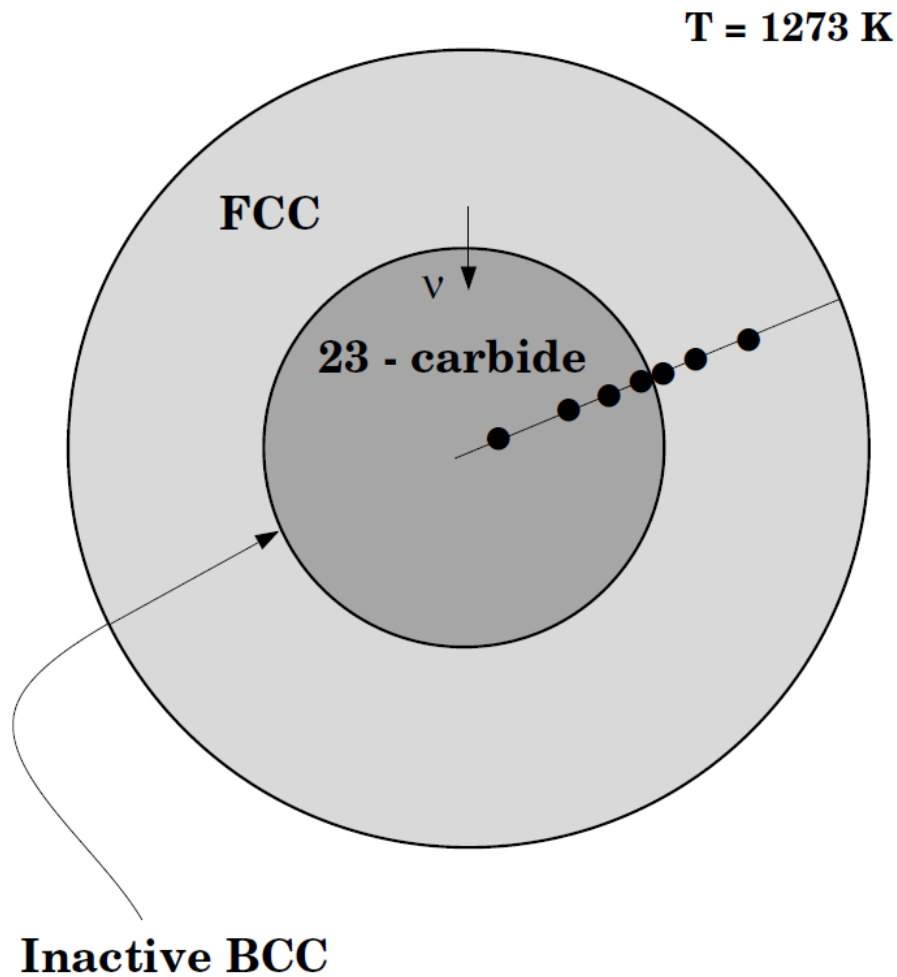
```
POST-1:  
POST-1:  
POST-1:  
POST-1:@?<_hit_return_to_continue_>  
POST-1:  
POST-1: set-inter  
--OK---  
POST-1:
```



Example exb3

Dissolution of 23-carbide in an austenitic matrix

This example calculates the dissolution of an M₂₃C₆ particle in an austenite matrix. A film of ferrite is allowed to nucleate around the carbide during the precipitation.



exb3-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb3\setup.DCM.test"
SYS:
SYS: @@ Moving boundary example.
SYS: @@ Dissolution of 23-carbide in an austenitic matrix
SYS: @@ This example calculates the dissolution of an M23C6 particle in an
SYS: @@ austenite matrix. A film of ferrite is allowed to nucleate around the
SYS: @@ carbide during the precipitation.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASES
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: @@
TDB_TCFE11: @@ USE A DATABASE FOR THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw fedemo
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-sys fe c cr
 C CR
 DEFINED
TDB_FEDEMO: rej ph *
 GAS:G LIQUID:L BCC_A2
 LAVES_PHASE_C14 CBCC_A12 CEMENTITE
 CHI_A12 CUB_A13 DIAMOND_FCC_A4
 FCC_A1 GRAPHITE HCP_A3
 KSI_CARBIDE M23C6 M3C2
 M5C2 M7C3 SIGMA
REJECTED
TDB_FEDEMO: res ph fcc bcc m23
 FCC_A1 BCC_A2 M23C6
 RESTORED
TDB_FEDEMO: get
10:08:36,053 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES
ELEMENTS
SPECIES
PHASES
Creating a new composition set FCC_A1#2
PARAMETERS
FUNCTIONS

List of references for assessed data

'J-O. Andersson, CALPHAD, 11 (1987) 271-276; TRITA 0314; C-CR'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'P. Franke, estimated parameter within SGTE, 2007; Fe-C, Ni-C, Mo-C, C-Mn'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
 volumes'
'A.V. Khvan, B. Hallstedt, C. Broeckmann, CALPHAD, 46, 24 -33(2014); Cr-Fe
 -C'
'J-O. Andersson, Metall. Trans. A, 19A (1988) 627-636 TRITA 0207 (1986); C
 -CR-FE'
'J-O. Andersson and B. Sundman, CALPHAD, 11 (1987) 83-92; TRITA 0270
(1986); CR-FE'
'J. Bratberg, Z. Metallkd., 96 (2005) 335-344; Fe-Cr-Mo-C'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'R. Naraghi, Thermo-Calc Software AB, Sweden, 2016; FCC Fe-Cr-C and C-Cr-Ni'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'D. Djurovic, B. Hallstedt, J. von Appen, R. Dronskowski, Calphad 35.4
(2011) 479-491; Fe-Mn-C'
-OK-
TDB_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE THE MOBILITY DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app mfedemo
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys c cr fe
 C CR
 DEFINED
APP: rej ph *
 BCC_A2 FCC_A1 REJECTED
APP: res ph fcc bcc
 FCC_A1 BCC_A2 RESTORED
APP: get
ELEMENTS
SPECIES
PHASES
Creating a new composition set FCC_A1#3
PARAMETERS
FUNCTIONS

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Z. Metallkunde 85(1994)502-509; C diffusion in fcc Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Z. Metallkunde 85(1994)498-501; C and N diffusion in bcc Cr
-Fe-Ni'
'B. Jonsson: ISIJ International, 35(1995)1415-1421; Cr, Fe and Ni
diffusion bcc Cr-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'

OK

APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR
APP: @@
APP: go d-m
NO TIME STEP DEFINED
*** ENTERING M23C6 AS A DIFFUSION NONE PHASE

DIC>
DIC>
DIC> @@ THE MOBILITY DATABASE LACKS KINETIC DATA FOR THE M23-CARBIDE
DIC> @@ SO AN ESTIMATE FOR THE MOBILITIES IN THIS PHASE ARE ENTERED.
DIC> ent-mob-est M23 c
M[M23,C](T)= 0;
DIC>
DIC> ent-mob-est M23 cr
M[M23,CR](T)= 3e-11*exp(-278000/8.3145/T);
DIC>
DIC> ent-mob-est M23 fe
M[M23,FE](T)= 1e-11*exp(-275000/8.3145/T);
DIC>
DIC> @@
DIC> @@ ENTER GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 1273; * N
DIC>
DIC> @@
DIC> @@ ENTER THE REGIONS carbide AND matrix
DIC> @@
DIC> enter-region carbide
DIC> enter-region matrix
ATTACH TO REGION NAMED /CARBIDE/:
ATTACHED TO THE RIGHT OF CARBIDE /YES/:
DIC> @@
DIC> @@ ASSUME SOME REASONABLE SIZE OF THE CARBIDE PARTICLE
DIC> @@
DIC> enter-grid carbide 5.0000000E-7 AUTO
DIC> @@
DIC> @@ THE SIZE OF THE FCC REGION WE CAN CALCULATE FROM A MASS BALANCE
DIC> @@ AFTER ESTIMATING THE INITIAL COMPOSITIONS IN THE TWO PHASES.
DIC> @@
DIC> enter-grid matrix 5.55859755E-7 AUTO
DIC> @@
DIC> @@ ENTER PHASES INTO THE REGION MATRIX. BOUNDARY CONDITIONS ARE GIVEN
DIC> @@ IF THE INACTIVE PHASE bcc IS NUCLEATED
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /CARBIDE/: matrix
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /MATRIX/: matrix
ATTACHED TO THE RIGHT OF MATRIX /YES/: no
PHASE NAME: /NONE/: bcc#1
DEPENDENT COMPONENT ? /FE/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC> @@
DIC> @@ ENTER THE PHASE INTO THE REGION carbide
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /CARBIDE/: carbide
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: m23c6
DIC>
DIC> @@
DIC> @@ ENTER COMPOSITIONS INTO THE PHASES
DIC> @@
DIC> enter-composition
REGION NAME : /CARBIDE/: carbide
PHASE NAME: /M23C6/: m23c6
DEPENDENT COMPONENT ? /FE/: fe
COMPOSITION TYPE /MOLE_FRACTION/: mole-fraction
PROFILE FOR /CR/: cr lin 0.55079807 0.55079807
DIC>
DIC>
DIC> enter-composition
REGION NAME : /MATRIX/: matrix
PHASE NAME: /FCC_A1#1/: fcc#1
DEPENDENT COMPONENT ? /FE/: fe
COMPOSITION TYPE /MOLE_FRACTION/: mole-fraction
PROFILE FOR /C/: cr lin 8.5203899E-2 8.5203899E-2
PROFILE FOR /CR/: c lin 1.8072433E-4 1.8072433E-4
DIC>
DIC> @@
DIC> @@ SET TO A SPHERICAL GEOMETRY
DIC> @@
DIC> enter-geo
GEOOMETRICAL EXPONENT /0/: 2
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME AND OTHER SIMULATION PARAMETERS

```
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /1/: 8000
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /800/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exb3 Y
DIC>
DIC>
DIC>
DIC> set-inter
--OK---
DIC>
```

exb3-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb3\run.DCM.test"
DIC>
DIC>
DIC> @@ exb3_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE b3
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE SET UP FROM FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exb3
OK
DIC>
DIC> @@
DIC> @@ WHEN THE FERRITE NUCLEATES WE USE DEFAULT VALUES
DIC> @@ AS STARTING VALUES FOR THE WIDTH OF THE NEW REGION
DIC> @@ AND THE VELOCITY OF THE INTERFACES
DIC> @@
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim yes
Region: CARBIDE
single geometric dense at 0.50000E-06
0.83296 92
Region: MATRIX
single geometric dense at 0.0000
1.0051 62
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
Trying old scheme 4
GENERATING STARTING VALUES FOR CELL # 1 INTERFACE # 2
DETERMINING INITIAL EQUILIBRIUM VALUES
CALCULATING STARTING VALUES: 9 EQUILIBRIUM CALCULATIONS
U-FRACTION IN SYSTEM: C = .0278637912207471 CR = .149918318671311
FE = .850081681459196
TOTAL SIZE OF SYSTEM: 4.93068569406E-18 [m^3]
U-FRACTION IN SYSTEM: C = .0278637912207471 CR = .149918318671311
FE = .850081681459196
TOTAL SIZE OF SYSTEM: 4.93068569406E-18 [m^3]
0.56566300795141 0.565797019192576 0.565663812089353 1.015925208843506E-002 3.012654895297374E-
004 4.77345260290525E-005 2.99573903808362E-006 7.21536208790236E-007 8.262638058237697E-
007 4.246506052259695E-008 4.119324775544597E-008 3.974271203488054E-008 4.355497133381697E-
008 3.870625294979825E-008 3.673410109463311E-008 3.294771045521285E-008 3.668595706394525E-
008 2.600355364026559E-008 1.460917538560299E-008 1.65819065504747E-009 4.335933841029224E-
012 1.279149124343428E-015 2.899183627817856E-017 TIME = 0.10000000E-06 DT = 0.10000000E-
06 SUM OF SQUARES = 0.28991836E-16
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.17310735E-02 AND -0.17310735E-02
POSITION OF INTERFACE CARBIDE / MATRIX IS 0.49982689E-06
U-FRACTION IN SYSTEM: C = .0278769048021911 CR = .149918499752463
FE = .850081500378044
TOTAL SIZE OF SYSTEM: 4.93068569406E-18 [m^3]
16 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARBIDE

CPU time used in timestep 2 seconds
1.096405342624566E-004 1.096626445880325E-004 1.096405350852248E-004 5.301312550514082E-008 2.238795244595802E-
008 6.726062328394531E-011 8.232621487274427E-014 3.182654756339016E-019 TIME = 0.30000000E-06 DT = 0.20000000E-
06 SUM OF SQUARES = 0.31826548E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.29934902E-05 AND -0.29934902E-05
POSITION OF INTERFACE CARBIDE / MATRIX IS 0.49982629E-06
U-FRACTION IN SYSTEM: C = .0278774003819474 CR = .149918500361418
FE = .850081499769089
TOTAL SIZE OF SYSTEM: 4.93068569406E-18 [m^3]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: MATRIX

CPU time used in timestep 1 seconds
1.217834959640360E-004 1.217822143862585E-004 1.217736602730391E-004 6.952193690456773E-007 2.897691539868494E-
009 5.94476007713682E-012 3.496734461402032E-016 2.032526948286751E-022 TIME = 0.70000000E-06 DT = 0.40000000E-
06 SUM OF SQUARES = 0.20325269E-21
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.28086480E-05 AND -0.28086480E-05
POSITION OF INTERFACE CARBIDE / MATRIX IS 0.49982517E-06
U-FRACTION IN SYSTEM: C = .027877419496498 CR = .149918501468152
FE = .850081498662355
TOTAL SIZE OF SYSTEM: 4.93068569406E-18 [m^3]

output ignored...

... output resumed

8.227990151515691E-012 6.644349202219934E-017 TIME = 5409.6764 DT = 800.00000 SUM OF SQUARES = 0.66443492E-16
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.72469007E-12 AND -0.72469007E-12
POSITION OF INTERFACE CARBIDE / MATRIX IS 0.39298838E-06
U-FRACTION IN SYSTEM: C = .0278315366128045 CR = .149829818102637
FE = .85017018202787
TOTAL SIZE OF SYSTEM: 4.93068569406E-18 [m^3]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: MATRIX

CPU time used in timestep 1 seconds
7.28957649320801E-008 7.291652649070389E-008 7.282475060639085E-008 4.660101478085030E-008 3.690335781027564E-
008 1.927578251022090E-008 9.373787665480115E-010 1.604800356814237E-015 2.936178286076429E-
019 TIME = 6209.6764 DT = 800.00000 SUM OF SQUARES = 0.29361783E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.28917572E-12 AND -0.28917572E-12
POSITION OF INTERFACE CARBIDE / MATRIX IS 0.392975704E-06
U-FRACTION IN SYSTEM: C = .02783153774494937 CR = .14982981860118
FE = .850170181529327
TOTAL SIZE OF SYSTEM: 4.93068569406E-18 [m^3]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: MATRIX

CPU time used in timestep 1 seconds
2.512054726275328E-008 2.512540223109571E-008 2.507787701776461E-008 1.662810811079077E-008 1.517974844775034E-
008 1.244982451205507E-008 7.683070933640646E-009 7.681065394535538E-009 1.617461485851593E-
009 2.967270075043390E-015 4.940354918402674E-
```

020 TIME = 7009.6764 DT = 800.00000 SUM OF SQUARES = 0.49403549E-19
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.39242850E-13 AND -0.39242850E-13
 POSITION OF INTERFACE CARBIDE / MATRIX IS 0.39272565E-06
 U-FRACTION IN SYSTEM: C = .0278315402086449 CR = .149829818985228
 FE = .850170181145279
 TOTAL SIZE OF SYSTEM: 4.93068569406E-18 [m^3]
 1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: MATRIX

CPU time used in timestep 1 seconds					
8.558014626556451E-009	8.558398232896329E-009	8.532476480695151E-009	5.576393278647864E-009	5.511701269025334E-	
009 5.365919512143457E-009	5.097369656807043E-009	5.098502113828289E-009	4.564634566841469E-		
009 3.600267924469804E-009	2.023954154531248E-009	2.023825894613357E-009	2.325233958152578E-		
010 7.941002790464720E-018	TIME = 7809.6764 DT = 800.00000	SUM OF SQUARES = 0.79410028E-17			
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.10417307E-12 AND 0.10417307E-12					
POSITION OF INTERFACE CARBIDE / MATRIX IS 0.39280899E-06					
U-FRACTION IN SYSTEM: C = .0278315407879869 CR = .149829819000751					
FE = .850170181129756					
TOTAL SIZE OF SYSTEM: 4.93068569406E-18 [m^3]					
30 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARBIDE					
CPU time used in timestep 1 seconds					
7.410550559270847E-008	7.410335722342407E-008	7.444093059196022E-008	2.903266432359087E-008	2.866492353727829E-	
008 2.751852345689572E-008	2.565458543205420E-008	2.564001639465827E-008	2.186294335122158E-		
008 1.537322331289984E-008	5.737901025146476E-009	1.307140972294803E-			
016 TIME = 8000.0000 DT = 190.32356	SUM OF SQUARES = 0.13071410E-15				
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.41375431E-12 AND 0.41375431E-12					
POSITION OF INTERFACE CARBIDE / MATRIX IS 0.39288774E-06					
U-FRACTION IN SYSTEM: C = .0278315407310841 CR = .149829819005946					
FE = .850170181124561					
TOTAL SIZE OF SYSTEM: 4.93068569406E-18 [m^3]					
MUST SAVE WORKSPACE ON FILE					
WORKSPACE SAVED ON FILE					
RECLAIMING WORKSPACE					
DELETING TIME-RECORD FOR TIME 45.242773					
DELETING TIME-RECORD FOR TIME 67.499169					
DELETING TIME-RECORD FOR TIME 67.499179					
DELETING TIME-RECORD FOR TIME 67.499199					
DELETING TIME-RECORD FOR TIME 67.499239					
DELETING TIME-RECORD FOR TIME 67.499319					
DELETING TIME-RECORD FOR TIME 67.499479					
DELETING TIME-RECORD FOR TIME 67.499799					
DELETING TIME-RECORD FOR TIME 67.500439					
DELETING TIME-RECORD FOR TIME 67.501719					
DELETING TIME-RECORD FOR TIME 67.504279					
DELETING TIME-RECORD FOR TIME 67.509399					
DELETING TIME-RECORD FOR TIME 67.519639					
DELETING TIME-RECORD FOR TIME 67.540119					
DELETING TIME-RECORD FOR TIME 67.581079					
DELETING TIME-RECORD FOR TIME 67.662999					
DELETING TIME-RECORD FOR TIME 67.826839					
DELETING TIME-RECORD FOR TIME 68.154519					
DELETING TIME-RECORD FOR TIME 68.809879					
DELETING TIME-RECORD FOR TIME 70.120599					
DELETING TIME-RECORD FOR TIME 72.742039					
DELETING TIME-RECORD FOR TIME 77.984919					
DELETING TIME-RECORD FOR TIME 88.470679					
DELETING TIME-RECORD FOR TIME 109.44220					
DELETING TIME-RECORD FOR TIME 151.38524					
DELETING TIME-RECORD FOR TIME 235.27132					
DELETING TIME-RECORD FOR TIME 403.04348					
DELETING TIME-RECORD FOR TIME 738.58780					
DELETING TIME-RECORD FOR TIME 1409.6764					
DELETING TIME-RECORD FOR TIME 2209.6764					
DELETING TIME-RECORD FOR TIME 3009.6764					
DELETING TIME-RECORD FOR TIME 3809.6764					
DELETING TIME-RECORD FOR TIME 4609.6764					
DELETING TIME-RECORD FOR TIME 5409.6764					
DELETING TIME-RECORD FOR TIME 6209.6764					
DELETING TIME-RECORD FOR TIME 7009.6764					
KEEPING TIME-RECORD FOR TIME 7809.6764					
AND FOR TIME 8000.0000					
WORKSPACE RECLAIMED					
TIMESTEP AT 8000.00000 SELECTED					

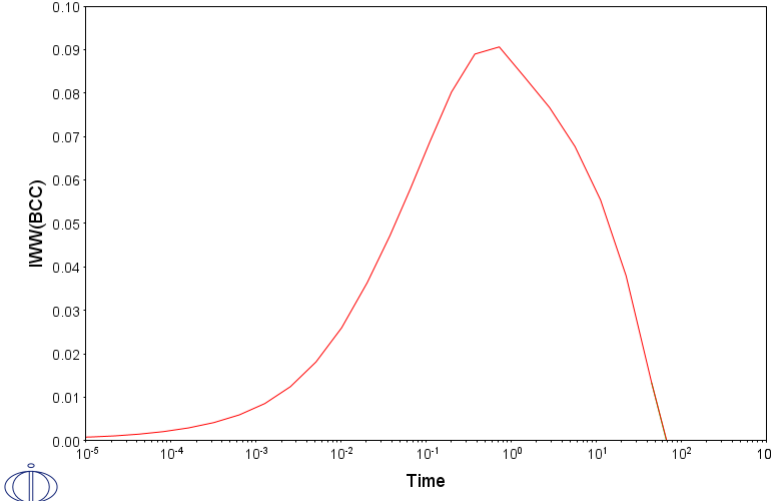
DIC>
DIC> set-inter
--OK--
DIC>

exb3-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb3\plot.DCM.test"
DIC>
DIC>
DIC> @@ exb3_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE b3
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MODULE AND SPECIFY THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 8.00000E+03
DIC> read exb3
OK
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ LET US SEE HOW THE AMOUNT OF FERRITE VARIED DURING THE
POST-1: @@ SIMULATION
POST-1: @@
POST-1: s-d-a y iww(bcc)
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-ax-typ x log
POST-1: s-s-s x n 1E-5 1E3
POST-1: s-s-s y n 0 0.1
POST-1:
POST-1: set-tit Figure b3.1
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Figure b3.1

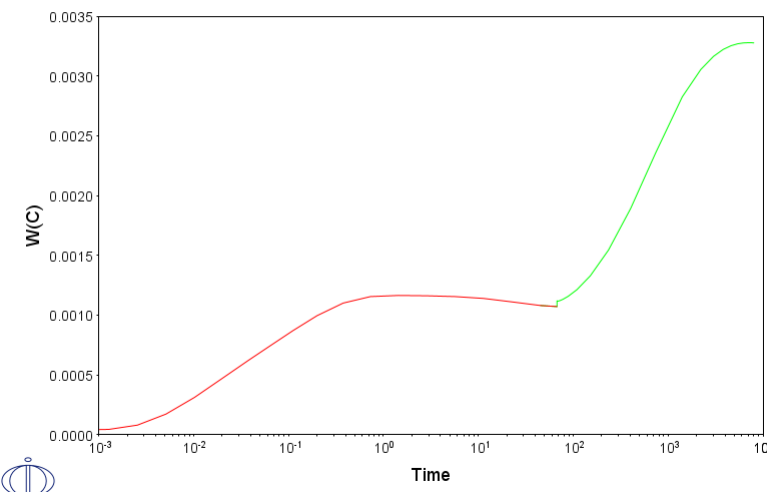
2021.05.13.10.17.02
CELL #1



```
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: @@
POST-1: @@ NOW LOOK AT THE ALLOYING ELEMENTS AT THE UPPER BOUND OF THE SYSTEM
POST-1: @@
POST-1: s-d-a y w(c)
POST-1: s-s-s x n 1E-3 1E4
POST-1: s-p-c interface last
POST-1:
POST-1: set-tit Figure b3.2
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Figure b3.2

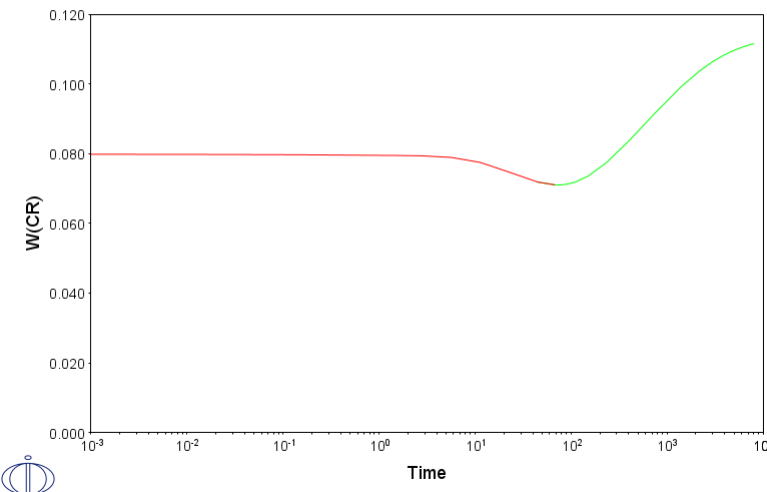
2021.05.13.10.17.04
UPPER INTERFACE OF REGION "LAST"
CELL #1



```
POST-1:  
POST-1:  
POST-1:@?<_hit_return_to_continue_>  
POST-1:  
POST-1: s-d-a y w(cr)  
POST-1: s-s-s y n 0 0.12  
POST-1:  
POST-1: set-tit Figure b3.3  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ...          OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Figure b3.3

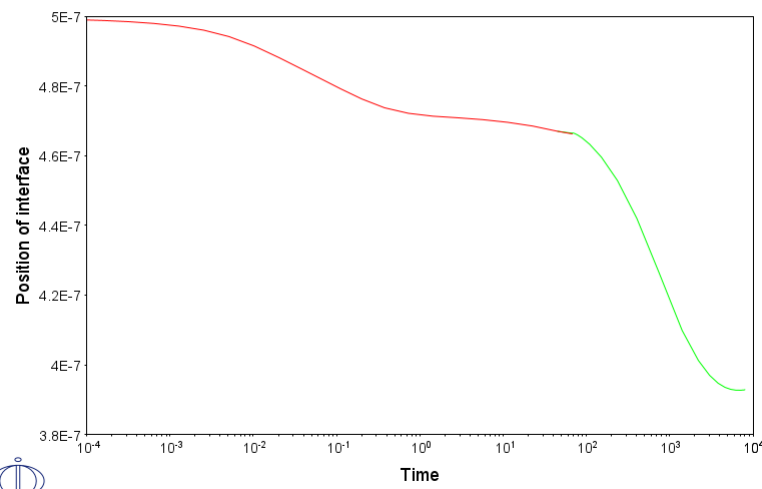
2021.05.13.10.17.06
UPPER INTERFACE OF REGION "LAST"
CELL #1



```
POST-1:  
POST-1:  
POST-1:@?<_hit_return_to_continue_>  
POST-1:  
POST-1: @@  
POST-1: @@ AND FINALLY LOOK AT THE CHANGE OF RADIUS OF THE M23-CARBIDE  
POST-1: @@  
POST-1: s-d-a y position carbide upper  
POST-1: s-s-s x n 1E-4 1E4  
POST-1:  
POST-1: set-tit Figure b3.4  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ...          OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Figure b3.4

2021.05.13.10.17.08
UPPER INTERFACE OF REGION "CARBIDE#1"
CELL #1



```
POST-1:  
POST-1:  
POST-1:@?<_hit_return_to_continue_>  
POST-1:  
POST-1:  
POST-1: set-inter  
---OK---  
POST-1:
```



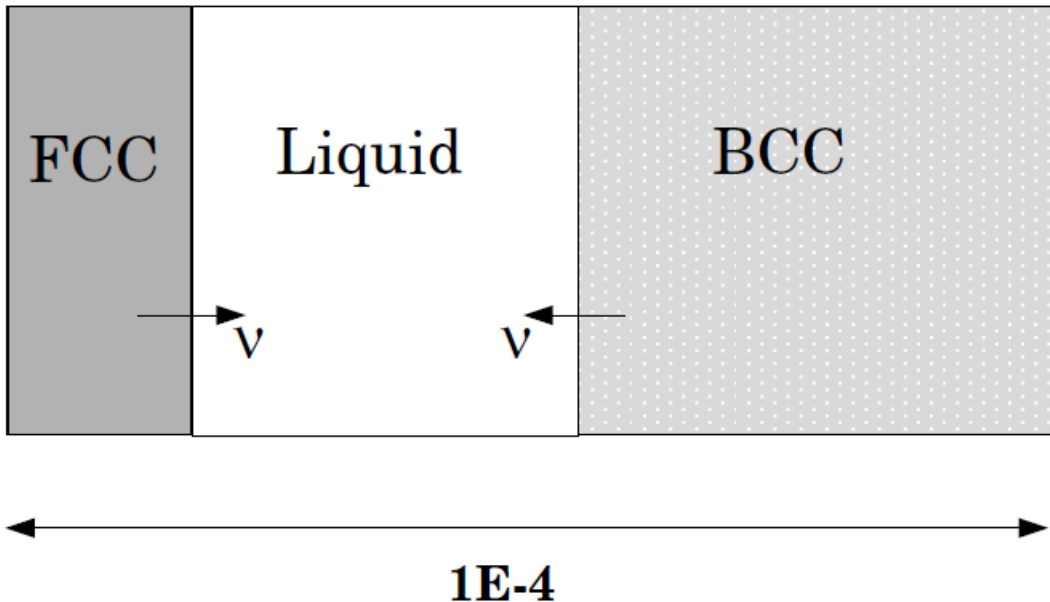
Example exb4a

Solidification path of a Fe-18%Cr-8%Ni alloy: Eutectic reaction

This example demonstrates the solidification path of an Fe-18%Cr-8%Ni alloy. A eutectic reaction is assumed, LIQUID \rightarrow BCC + FCC. Hence the BCC and FCC regions should be on separate sides of the liquid region. Comparison is made with both a Scheil-Gulliver simulation and equilibrium solidification conditions, both made with Thermo-Calc.

$$Time > 0$$

$$T = 1900 - 1 * \text{Time} \text{ K}$$



exb4a-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4a\setup.DCM.test"
SYS: @@
SYS: @@ Moving boundary problem.
SYS: @@ Solidification path of an Fe-18%Cr-8%Ni alloy: Eutectic reaction
SYS: @@ This example demonstrates the solidification path of an Fe-18%Cr-8%Ni
SYS: @@ alloy. A eutectic reaction is assumed, LIQUID -> BCC + FCC. Hence the
SYS: @@ BCC and FCC regions should be on separate sides of the liquid region.
SYS: @@ Comparison is made with both a Scheil-Gulliver simulation and equilibrium
SYS: @@ solidification conditions, both done in Thermo-Calc.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exb4a_setup.DCM
SYS:
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:-
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: sw tcfe9
Current database: Steels/Fe-Alloys v9.3

VA           /- DEFINED
L12_FCC      B2_BCC          DICTRA_FCC_A1
REJECTED
TDB_TCFE9:-
TDB_TCFE9: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_TCFE9: def-sys fe ni cr
FE           NI              CR
DEFINED
TDB_TCFE9:-
TDB_TCFE9: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_TCFE9: rej ph /all
LIQUID:L      BCC_A2          FCC_A1
HCP_A3        CBC_A12         CUB_A13
SIGMA         CHI_A12         LAVES_PHASE_C14
CR3SI         NBNI3           NISTI
CRZN17        BETA1           GAMMA
AL5FE4        FLUORITE_C1:I  ZRO2_TETR:I
M2O3C:I       CENI2           CENI5
REJECTED
TDB_TCFE9: res ph fcc liq bcc
FCC_A1        LIQUID:L        BCC_A2
RESTORED
TDB_TCFE9:-
TDB_TCFE9: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_TCFE9: get
10:18:30,719 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
FUNCTIONS .....

List of references for assessed data

'A. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'X.-G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'X.-G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
    volumes'
'J. Brillo and I. Egry, Int. J. Thermophysics, 24, 1155-1170'
'B.-J. Lee, CALPHAD, 17 (1993) 251-268; revision of Fe-Cr and Fe-Ni liquid'
'J-O. Andersson and B. Sundman, CALPHAD, 11 (1987) 83-92; TRITA 0270
    (1986); CR-FE'
'B.-J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'A. Dinsdale and T. Chart, MTDS NPL, Unpublished work (1986); CR-NI'
'A. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
-OK-
TDB_TCFE9:-
TDB_TCFE9: @@
TDB_TCFE9: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_TCFE9: @@ SWITCH TO A MOBILITY DATABASE AND APPEND THE DATA
TDB_TCFE9: @@
TDB_TCFE9: app
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
```

FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2
TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
SNOB1 = SGTE Noble Metal Alloys Database v1.2
STBC2 = SGTE Thermal Barrier Coating TDB v2.2
STBC1 = SGTE Thermal Barrier Coating TDB v1.1
SALT1 = SGTE Molten Salts Database v1.3
SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2 = TC Semi-Conductors v2.1
SLAG4 = Fe-containing Slag v4.1
SLAG3 = Fe-containing Slag v3.2
SLAG2 = Fe-containing Slag v2.2
SLAG1 = Fe-containing Slag v1.2
TCOX11 = Metal Oxide Solutions v11.0
TCOX10 = Metal Oxide Solutions v10.1
TCOX9 = Metal Oxide Solutions v9.0
TCOX8 = Metal Oxide Solutions v8.0
TCOX7 = Metal Oxide Solutions v7.0
TCOX6 = Metal Oxide Solutions v6.0
TCOX5 = Metal Oxide Solutions v5.1
TCOX4 = Metal Oxide Solutions v4.1
ION3 = Ionic Solutions v3.0
ION2 = Ionic Solutions v2.6
ION1 = Ionic Solutions v1.5
NOX2 = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4 = Solder Alloys v4.0
TCSLD3 = Solder Alloys v3.3
TCSLD2 = Solder Alloys v2.0
TCSLD1 = Solder Alloys v1.1
TCSI1 = Ultrapure Silicon v1.2
TCMP2 = Materials Processing v2.5
TCES1 = Combustion/Sintering v1.1
TCSR1 = Super Conductor v1.0
TCFC1 = SOFC Database v1.0
TCNF2 = Nuclear Fuels v2.1b
NUMT2 = Nuclear Materials v2.1
NUOX4 = Nuclear Oxides v4.2
NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephista-19
MEPH15 = IRSN Mephista-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQSO2 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SLDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.0
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3
MOBFE1 = Steels/Fe-Alloys Mobility v1.1
MOBFE2 = Steels/Fe-Alloys Mobility v2.0
MOBFE3 = Steels/Fe-Alloys Mobility v3.0
MOBFE4 = Steels/Fe-Alloys Mobility v4.0
MOBFE5 = Steels/Fe-Alloys Mobility v5.0
MOBFE6 = Steels/Fe-Alloys Mobility v6.0
MOBNI5 = Ni-Alloys Mobility v5.1

```

MOBNI4 = Ni-Alloys Mobility v4.1
MOBNI3 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBSI1 = Si-Alloys Mobility v1.0
MOBSLD1 = Solder-Alloys Mobility v1.1
MOBT14 = Ti-Alloys Mobility v4.0
MOBT13 = Ti-Alloys Mobility v3.1
MOBT12 = Ti-Alloys Mobility v2.0
MOBT11 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

DATABASE NAME /TCFE9/: mobfe4
Current database: Steels/Fe-Alloys Mobility v4.0

VA DEFINED
B2_BCC REJECTED
APP: def-sys fe ni cr
    FE                      NI                      CR
    DEFINED
APP: rej ph /all          FCC_A1                  HCP_A3
    BCC_A2                  LIQUID:L               BCC_A2
    LIQUID:L REJECTED
APP: res ph fcc liq bcc
    FCC_A1                  LIQUID:L
    RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Ni diffusion fcc Cr-Ni'
'B. Jonsson: Z. Metallkunde 86(1995)686-692; Cr, Fe and Ni diffusion fcc
    Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
    -Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
'B. Jonsson: ISIJ International, 35(1995)1415-1421; Cr, Fe and Ni
    diffusion bcc Cr-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
    in bcc Fe'
'No assessed or estimated parameters'
'Estimation by using the modified Sutherland equation from Chen et al.,
    Phil. Magazine 94 (2014) 1552.'
'L. Zhang et al., Acta Mater., 58(2010)3664.'

-OK-
APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER GLOBAL CONDITION T
DIC> @@
DIC> @@ LOWER THE TEMPERATURE TO A RATE OF 1 K/s
DIC> @@
DIC> set-cond glob T 0 1900-1*TIME; * N
DIC>
DIC> @@
DIC> @@ ENTER A REGION CALLED smalta
DIC> @@
DIC> enter-region smalta
DIC>
DIC> @@
DIC> @@ ENTER A DOUBLE GEOMETRIC GRID INTO THE REGION
DIC> @@
DIC> enter-grid
REGION NAME : /SMALTA/: smalta
WIDTH OF REGION /1/: 1e-4
TYPE /LINEAR/: AUTO
DIC>
DIC>
DIC> @@
DIC> @@ ENTER active PHASES INTO THE REGION
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /SMALTA/: smalta
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: liq
DIC>
DIC> @@
DIC> @@ ENTER inactive PHASES INTO THE REGION: ONE PHASE ON EACH SIDE OF THE LIQUID
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /SMALTA/: smalta

```

```

ATTACHED TO THE RIGHT OF SMALTA /YES/: no
PHASE NAME: /NONE/: fcc#1
DEPENDENT COMPONENT ? /NI/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-5
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /SMALTA/: smalta
ATTACHED TO THE RIGHT OF SMALTA /YES/: yes
PHASE NAME: /NONE/: bcc#1
DEPENDENT COMPONENT ? /NI/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-5
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC> @@
DIC> @@ ENTER A START COMPOSITION FOR THE LIQUID
DIC> @@
DIC> enter-composition
REGION NAME : /SMALTA/: smalta
PHASE NAME: /LIQUID/: liq
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /CR/: cr lin 18 18
PROFILE FOR /NI/: ni lin 8 8
DIC>
DIC> @@
DIC> @@ THE BOUNDARY CONDITION IS A CLOSED SYSTEM (THE DEFAULT) AS WE DO NOT SPECIFY
DIC> @@ ANYTHING ELSE.
DIC> @@
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /1/: 200
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /20/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC>
DIC> @@
DIC> @@ CHECK THE INTERFACE POSITION. THIS IS TO MAKE SURE THAT THE
DIC> @@ LIQUID REGION DOES NOT SHRINK TOO MUCH DURING A TIMESTEP.
DIC> @@ IN ADDITION THE TIMESTEP IS CONTROLLED BY THE PHASE INTERFACE
DIC> @@ DISPLACEMENT DURING THE SIMULATION.
DIC> @@
DIC> s-s-c
NS01A PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA TIMESTEPS IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/: AUTO
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/:
DEGREE OF IMPLICITY WHEN INTEGRATING PDEs (AUTO, 0 -> 0.5 -> 1) : /AUTO/:
MAX TIMESTEP CHANGE PER TIMESTEP : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/:
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exb4a Y
DIC>
DIC> set-inter
--OK---
DIC>
```

exb4a-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4a\run.DCM.test"
DIC>
DIC>
DIC> @@ exb4a_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE b4a
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE SET UP FROM FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exb4a
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim yes
Region: SMALTA
double geometric
dense at outer boundaries, coarse at 0.50000E-04
lower part 1.2275 22
upper part 0.81465 22
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
INFO: TIMESTEP IS CONTROLLED BY INTERFACE POSITION
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
TIME = 0.10000000E-06 DT = 0.10000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 0.26028659E-05 DT = 0.25028659E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.76085978E-05 DT = 0.50057319E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.17620062E-04 DT = 0.10011464E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.37642989E-04 DT = 0.20022928E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 0.77688844E-04 DT = 0.40045855E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.15778055E-03 DT = 0.80091710E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.31796397E-03 DT = 0.16018342E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.63833082E-03 DT = 0.32036684E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.12790645E-02 DT = 0.64073368E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.25605319E-02 DT = 0.12814674E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.51234666E-02 DT = 0.25629347E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.10249336E-01 DT = 0.51258694E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.20501075E-01 DT = 0.10251739E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.41004553E-01 DT = 0.20503478E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.82011508E-01 DT = 0.41006956E-01 SUM OF SQUARES = 0.0000000
```

F-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882258

TOTAL SIZE OF SYSTEM: 1E-04 [m]

CPU time used in timestep 0 seconds
TIME = 0.16402542 DT = 0.82013911E-01 SUM OF SQUARES = 0.0000000

U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882258

TOTAL SIZE OF SYSTEM: 1E-04 [m]

CPU time used in timestep 0 seconds

output ignored...

... output resumed

013	4.451016681584280E-018	TIME = 178.38332	DT = 1.3107200	SUM OF SQUARES = 0.44510167E-17	4.398999059913720E-
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.59985813E-07 AND -0.59985813E-07					
POSITION OF INTERFACE R_FCC_A1 / R_BCC_A2 IS 0.70798215E-05					
U-FRACTION IN SYSTEM: CR = -.191500837210103 FE = .733100607091574 NI = .0753985556983228					
TOTAL SIZE OF SYSTEM: 1E-04 [m]					
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_BCC_A2					
CPU time used in timestep 1 seconds					
1.087065393704081E-005	1.087060191036813E-005	1.086454200977664E-005	3.773771610984094E-007	3.512334048357269E-	
011 3.747162386535219E-016	6.343584820225726E-				
021 TIME = 181.00476 DT = 2.6214400 SUM OF SQUARES = 0.63435848E-20					
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.43957921E-07 AND -0.43957921E-07					
POSITION OF INTERFACE R_FCC_A1 / R_BCC_A2 IS 0.69645885E-05					
U-FRACTION IN SYSTEM: CR = -.191500837128086 FE = .733100607278848 NI = .0753985555930666					
TOTAL SIZE OF SYSTEM: 1E-04 [m]					
2 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_BCC_A2					
CPU time used in timestep 2 seconds					
7.96841032302261E-006	7.968559090050239E-006	7.965071815030743E-006	9.965452851655026E-007	1.848642455401581E-	
007 1.454590664712276E-012	9.973971029481365E-				
017 TIME = 186.24764 DT = 5.2428800 SUM OF SQUARES = 0.99739710E-16					
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.17721381E-07 AND -0.17721381E-07					
POSITION OF INTERFACE R_FCC_A1 / R_BCC_A2 IS 0.68716774E-05					
U-FRACTION IN SYSTEM: CR = .191500837051579 FE = .733100607519551 NI = .0753985554288702					
TOTAL SIZE OF SYSTEM: 1E-04 [m]					
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_BCC_A2					
CPU time used in timestep 2 seconds					
5.760483174316431E-006	5.760637276761246E-006	5.758982848124157E-006	2.368951635922415E-006	1.576724015433373E-	
006 3.73510118696255E-007	4.317210487061832E-011	4.720019209951113E-015	6.408809900060585E-		
019 TIME = 196.73340 DT = 10.485760 SUM OF SQUARES = 0.64088099E-18					
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.21802006E-07 AND 0.21802006E-07					
POSITION OF INTERFACE R_FCC_A1 / R_BCC_A2 IS 0.71002880E-05					
U-FRACTION IN SYSTEM: CR = .191500837049122 FE = .733100607565332 NI = .075398555385546					
TOTAL SIZE OF SYSTEM: 1E-04 [m]					
CPU time used in timestep 2 seconds					
1.322183101525621E-005	1.322204636875977E-005	1.321670781935714E-005	7.855597921282976E-008	2.238377524146627E-	
011 5.880537982680392E-017	TIME = 200.00000 DT = 3.2666030	SUM OF SQUARES = 0.58805380E-16			
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.13886077E-07 AND 0.13886077E-07					
POSITION OF INTERFACE R_FCC_A1 / R_BCC_A2 IS 0.71456483E-05					
U-FRACTION IN SYSTEM: CR = -.191500837043234 FE = .733100607594293 NI = .0753985553624729					
TOTAL SIZE OF SYSTEM: 1E-04 [m]					
MUST SAVE WORKSPACE ON FILE					
WORKSPACE SAVED ON FILE					
RECLAIMING WORKSPACE					
DELETING TIME-RECORD FOR TIME 175.56749					
DELETING TIME-RECORD FOR TIME 175.76189					
DELETING TIME-RECORD FOR TIME 175.76190					
DELETING TIME-RECORD FOR TIME 175.76192					
DELETING TIME-RECORD FOR TIME 175.76196					
DELETING TIME-RECORD FOR TIME 175.76204					
DELETING TIME-RECORD FOR TIME 175.76220					
DELETING TIME-RECORD FOR TIME 175.76252					
DELETING TIME-RECORD FOR TIME 175.76316					
DELETING TIME-RECORD FOR TIME 175.76444					
DELETING TIME-RECORD FOR TIME 175.76700					
DELETING TIME-RECORD FOR TIME 175.77212					
DELETING TIME-RECORD FOR TIME 175.78236					
DELETING TIME-RECORD FOR TIME 175.80284					
DELETING TIME-RECORD FOR TIME 175.84380					
DELETING TIME-RECORD FOR TIME 175.92572					
DELETING TIME-RECORD FOR TIME 176.08956					
DELETING TIME-RECORD FOR TIME 176.41724					
DELETING TIME-RECORD FOR TIME 177.07260					
DELETING TIME-RECORD FOR TIME 178.38332					
DELETING TIME-RECORD FOR TIME 181.00476					
DELETING TIME-RECORD FOR TIME 186.24764					
KEEPING TIME-RECORD FOR TIME 196.73340					
AND FOR TIME 200.00000					
WORKSPACE RECLAIMED					
TIMESTEP AT 200.000000 SELECTED					
DIC>					
DIC> @@					
DIC> @@ THE SIMULATION IS FINISHED					

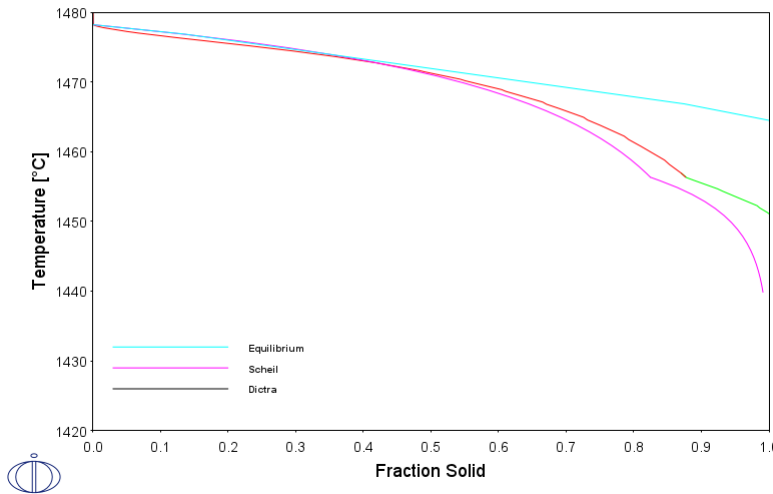
```
DIC> @@  
DIC>  
DIC> set-inter  
--OK---  
DIC>
```

exb4a-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4a\plot.DCM.test"
DIC>
DIC>
DIC> @@ exb4a_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE b4a
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 2.00000E+02
DIC> read exb4a
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: set-title Fe-18%Cr-8%Ni
POST-1:
POST-1: @@
POST-1: @@ PLOT THE FRACTION OF SOLID AND COMPARE WITH A SCHEIL-GULLIVER SIMULATION
POST-1: @@ AND EQUILIBRIUM SOLIDIFICATION (DATA ON FILE exb4.exp)
POST-1: @@
POST-1: enter function fs=1-ivv(liquid);
POST-1: s-d-a x fs
POST-1: s-s-s x n 0 1
POST-1: set-axis-text
AXIS (X, Y OR Z) : x
AUTOMATIC AXIS TEXT (Y OR N) /N/: n
AXIS TEXT : Fraction Solid
POST-1:
POST-1: s-d-a y t-c
POST-1: s-s-s y n 1420 1480
POST-1:
POST-1: s-p-c interface smalta lower
POST-1:
POST-1: app y exb4a.exp 0; 1
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...orking ...          OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Fe-18%Cr-8%Ni

2021.05.13.10.24.20
LOWER INTERFACE OF REGION "SMALTA#1"
CELL #1



```
POST-1:
POST-1:
POST-1: set-inter
--OK---
POST-1:
```



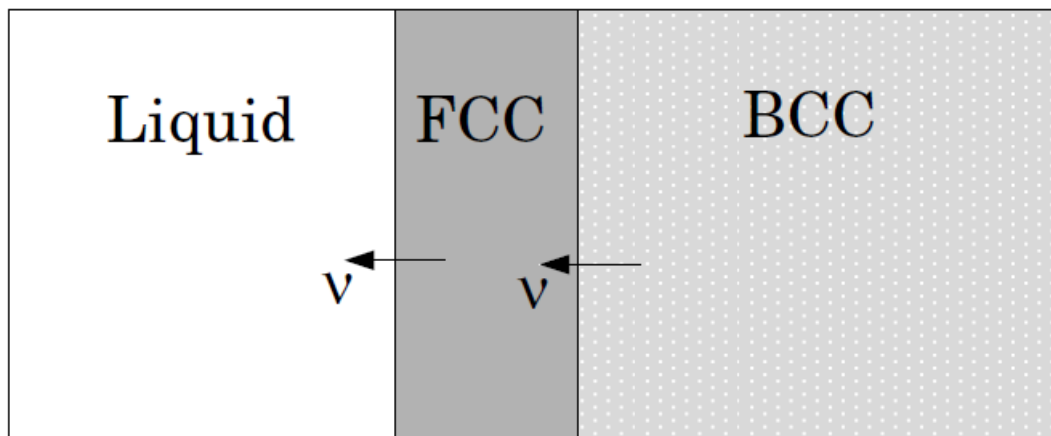
Example exb4b

Solidification path of an Fe-18%Cr-8%Ni alloy: Peritectic reaction

This example is the same as exb4a but now a peritectic reaction is assumed, LIQUID + BCC \rightarrow FCC. Hence the FCC region should appear in between the LIQUID and the BCC. Comparison is made with both a Scheil-Gulliver simulation and equilibrium solidification conditions, both made with Thermo-Calc.

Time > 0

$$T = 1900 - 1 * \text{Time} \text{ K}$$



← →
1E-4

exb4b-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4b\setup.DCM.test"

SYS: @@
SYS: @@ Moving boundary problem.
SYS: @@ Solidification path of an Fe-18%Cr-8%Ni alloy: Peritectic reaction
SYS: @@ This example is the same as exb4a but now a peritectic reaction is assumed:
SYS: @@ LIQUID + BCC -> FCC. Hence the FCC region should appear in between the LIQUID
SYS: @@ and the BCC. Comparison is made with both a Scheil-Gulliver simulation and
SYS: @@ equilibrium solidification conditions, both done in Thermo-Calc.

SYS: -----

NO SUCH COMMAND, USE HELP

SYS:
SYS: @@ exb4b_setup.DCM
SYS:
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da

THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED

DICTRA_FCC_A1 REJECTED

TDB_TCFE11:
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: sw fedemo

Current database: Iron Demo Database v4.0

VA /- DEFINED

TDB_FEDEMO:
TDB_FEDEMO: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_FEDEMO: def-sys fe ni cr

FE NI CR

DEFINED

TDB_FEDEMO:
TDB_FEDEMO: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_FEDEMO: rej ph /all

LIQUID:L BCC_A2 LAVES_PHASE_C14
CBCC_A12 CHI_A12 CUB_A13
FCC_A1 HCP_A3 SIGMA

REJECTED

TDB_FEDEMO: res ph fcc liq bcc
FCC_A1 LIQUID:L BCC_A2
RESTORED

TDB_FEDEMO:
TDB_FEDEMO: @@ RETRIEVE DATA FROM THE DATABASE FILE

TDB_FEDEMO: get

10:25:37,630 INFO *** Invoking Gibbs Energy System v6 ***

REINITIATING GES

ELEMENTS

SPECIES

PHASES

PARAMETERS ...

FUNCTIONS

List of references for assessed data

'A.T. Dinsdale, SGT Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'M. Ghasemi, Thermo-Calc Software AB, Sweden, 2020; Unifying liquid volume
database'
'M. Ghasemi, Thermo-Calc Software AB, Sweden, 2019; Viscosity of metallic
liquid'
'M Ghasemi, Thermo-Calc Software AB: assessing the surface tension of
metallic liquid'
'Y. Tang, in TCCU2, Thermo-Calc Software AB, Stockholm, Sweden'
'M. J. Assael, J. Phys. Chem. Ref. Data 35 (2006) 285; Fe, Al'
'M. J. Assael, High Temp. High Press., 41 (2012); Sb, Pb, Bi, Ni, Ag'
'B.J. Lee, CALPHAD, 17 (1993) 251-268; revision of Fe-Cr and Fe -Ni liquid'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'J-O. Andersson and B. Sundman, CALPHAD, 11 (1987) 83-92; TRITA 0270
(1986); CR-FE'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); CR -NI'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'

-OK

TDB_FEDEMO:
TDB_FEDEMO: @@

TDB_FEDEMO: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE AND APPEND THE DATA.

TDB_FEDEMO: @@

TDB_FEDEMO: app

Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0

FEDAT = TCS/TT Steels Database v1.0
 TCNI11 = Ni-Alloys v11.0
 TCNI10 = Ni-Alloys v10.0
 TCNI9 = Ni-Alloys v9.1
 TCNI8 = Ni-Alloys v8.2
 TCNI7 = Ni-Alloys v7.2
 TCNI6 = Ni-Alloys v6.1
 TCNI5 = Ni-Alloys v5.1
 TCNI4 = Ni-Alloys v4.0
 TCNI1 = Ni-Alloys v1.3
 TCAL8 = Al-Alloys v8.0
 TCAL7 = Al-Alloys v7.1
 TCAL6 = Al-Alloys v6.0
 TCAL5 = Al-Alloys v5.1
 TCAL4 = Al-Alloys v4.0
 TCAL3 = Al-Alloys v3.0
 TCAL2 = Al-Alloys v2.1
 TCAL1 = Al-Alloys v1.2
 TCMG6 = Mg-Alloys v6.1
 TCMG5 = Mg-Alloys v5.1
 TCMG4 = Mg-Alloys v4.0
 TCMG3 = Mg-Alloys v3.0
 TCMG2 = Mg-Alloys v2.0
 TCMG1 = Mg-Alloys v1.1
 TCTI3 = Ti-Alloys v3.0
 TCTI2 = Ti-Alloys v2.2
 TCTI1 = Ti-Alloys v1.0
 TCCU4 = Cu-Alloys v4.0
 TCCU3 = Cu-Alloys v3.1
 TCCU2 = Cu-Alloys v2.0
 TCCU1 = Cu-Alloys v1.0
 TCCC1 = Cemented carbide v1.0
 TCHEA5 = High Entropy Alloy v5.0
 TCHEA4 = High Entropy Alloy v4.2
 TCHEA3 = High Entropy Alloy v3.1
 TCHEA2 = High Entropy Alloy v2.1.1
 TCHEA1 = High Entropy Alloy v1.0
 SSOL7 = SGTE Alloy Solutions Database v7.0
 SSOL6 = SGTE Alloy Solutions Database v6.0
 SSOL5 = SGTE Alloy Solutions Database v5.0
 SSOL4 = SGTE Alloy Solutions Database v4.9g
 SSOL2 = SGTE Alloy Solutions Database v2.1
 SSUB6 = SGTE Substances Database v6.0
 SSUB5 = SGTE Substances Database v5.2
 SSUB4 = SGTE Substances Database v4.1
 SSUB3 = SGTE Substances Database v3.3
 SSUB2 = SGTE Substances Database v2.2
 SNOB3 = SGTE Noble Metal Alloys Database v3.1
 SNOB2 = SGTE Noble Metal Alloys Database v2.1
 SNOB1 = SGTE Noble Metal Alloys Database v1.2
 STBC2 = SGTE Thermal Barrier Coating TDB v2.2
 STBC1 = SGTE Thermal Barrier Coating TDB v1.1
 SALT1 = SGTE Molten Salts Database v1.3
 SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
 SEMC2 = TC Semi-Conductors v2.1
 SLAG4 = Fe-containing Slag v4.1
 SLAG3 = Fe-containing Slag v3.2
 SLAG2 = Fe-containing Slag v2.2
 SLAG1 = Fe-containing Slag v1.2
 TCOX11 = Metal Oxide Solutions v11.0
 TCOX10 = Metal Oxide Solutions v10.1
 TCOX9 = Metal Oxide Solutions v9.0
 TCOX8 = Metal Oxide Solutions v8.0
 TCOX7 = Metal Oxide Solutions v7.0
 TCOX6 = Metal Oxide Solutions v6.0
 TCOX5 = Metal Oxide Solutions v5.1
 TCOX4 = Metal Oxide Solutions v4.1
 ION3 = Ionic Solutions v3.0
 ION2 = Ionic Solutions v2.6
 ION1 = Ionic Solutions v1.5
 NOX2 = NPL Oxide Solutions Database v2.1
 TCNOBL1 = Noble Metals Alloys v1.0
 TCNOBL2 = Noble Metals Alloys v2.0
 TCSLD4 = Solder Alloys v4.0
 TCSLD3 = Solder Alloys v3.3
 TCSLD2 = Solder Alloys v2.0
 TCSLD1 = Solder Alloys v1.1
 TCSI1 = Ultrapure Silicon v1.2
 TCMP2 = Materials Processing v2.5
 TCES1 = Combustion/Sintering v1.1
 TCSC1 = Super Conductor v1.0
 TCFCl = SOFC Database v1.0
 TCNF2 = Nuclear Fuels v2.1b
 NUMT2 = Nuclear Materials v2.1
 NUOX4 = Nuclear Oxides v4.2
 NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
 NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
 NUCL19 = IRSN NUCLEA-19
 NUCL15 = IRSN NUCLEA-15_4
 NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
 MEPH19 = IRSN Mephista-19
 MEPH15 = IRSN Mephista-15_1
 MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
 TCAQ3 = Aqueous Solution v3.0
 TCAQ2 = Aqueous Solution v2.7
 AQS2 = TGG Aqueous Solution Database v2.6
 GCE2 = TGG Geochemical/Environmental TDB v2.3
 FEDEMO = Iron Demo Database v4.0
 ALDEMO = Aluminum Demo Database v4.0
 NI DEMO = Nickel Demo Database v2.0
 CUDEMO = Copper Demo Database v1.0
 SLDEMO = Solder Demo Database v1.0
 OXDEMO = Oxide Demo Database v3.0
 SUBDEMO = Substance Demo Database v1.0
 PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
 PG35 = PG35 Binary Semi-Conductors TDB v1.3
 PURE5 = SGTE Unary (Pure Elements) TDB v5.2
 MOB2 = Alloys Mobility v2.7
 MOB1 = Alloys Mobility v1.3
 MOBFE1 = Steels/Fe-Alloys Mobility v1.1
 MOBFE2 = Steels/Fe-Alloys Mobility v2.0
 MOBFE3 = Steels/Fe-Alloys Mobility v3.0
 MOBFE4 = Steels/Fe-Alloys Mobility v4.0
 MOBFE5 = Steels/Fe-Alloys Mobility v5.0
 MOBFE6 = Steels/Fe-Alloys Mobility v6.0
 MOBNI5 = Ni-Alloys Mobility v5.1


```

ATTACHED TO THE RIGHT OF SMALTA /YES/: yes
PHASE NAME: /NONE/: fcc#1
DEPENDENT COMPONENT ? /NI/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-5
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /SMALTA/: smalta
ATTACHED TO THE RIGHT OF SMALTA /YES/: yes
PHASE NAME: /NONE/: bcc#1
DEPENDENT COMPONENT ? /NI/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-5
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC> @@
DIC> @@ ENTER THE START COMPOSITION FOR THE LIQUID
DIC> @@
DIC> enter-composition
REGION NAME : /SMALTA/: smalta
PHASE NAME: /LIQUID/: liq
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /CR/: cr lin 18 18
PROFILE FOR /NI/: ni lin 8 8
DIC>
DIC> @@
DIC> @@ THE BOUNDARY CONDITION IS A CLOSED SYSTEM (DEFAULT) AS WE DO NOT SPECIFY
DIC> @@ ANYTHING ELSE.
DIC> @@
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 200
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /20/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ CHECK THE INTERFACE POSITION. THIS IS TO MAKE SURE THAT THE LIQUID REGION
DIC> @@ DOES NOT SHRINK TOO MUCH DURING A TIMESTEP. IN ADDITION, THE TIMESTEP IS
DIC> @@ CONTROLLED BY THE PHASE INTERFACE DISPLACEMENT DURING THE SIMULATION.
DIC> @@
DIC> s-s-c
NS01A PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA TIMESTEPS IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/: yes
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/:
DEGREE OF IMPLICITY WHEN INTEGRATING PDEs (AUTO, 0 -> 0.5 -> 1): /AUTO/:
MAX TIMESTEP CHANGE PER TIMESTEP : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/: @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exb4b Y
DIC>
DIC> set-inter
--OK--
DIC>

```

exb4b-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4b\run.DCM.test"
DIC>
DIC> @@ exb4b_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE b4b
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE SET UP FROM FILE
DIC> @@
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exb4b
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim yes
Region: SMALTA
single geometric dense at 0.10000E-03
0.93810          83
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           1 seconds
TIME = 0.27420944E-05 DT = 0.26420944E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.80262833E-05 DT = 0.52841888E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           1 seconds
TIME = 0.18594661E-04 DT = 0.10568378E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.39731416E-04 DT = 0.21136755E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.82004927E-04 DT = 0.42273511E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.16655195E-03 DT = 0.845457022E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.33564599E-03 DT = 0.16909404E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.67383408E-03 DT = 0.33818809E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.13502102E-02 DT = 0.67637617E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.27029626E-02 DT = 0.13527523E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.54084673E-02 DT = 0.27055047E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.10819477E-01 DT = 0.54110094E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.21641495E-01 DT = 0.10822019E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882253
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.43285533E-01 DT = 0.21644038E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882252
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
TIME = 0.86573608E-01 DT = 0.43288075E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219291
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep           0 seconds
```

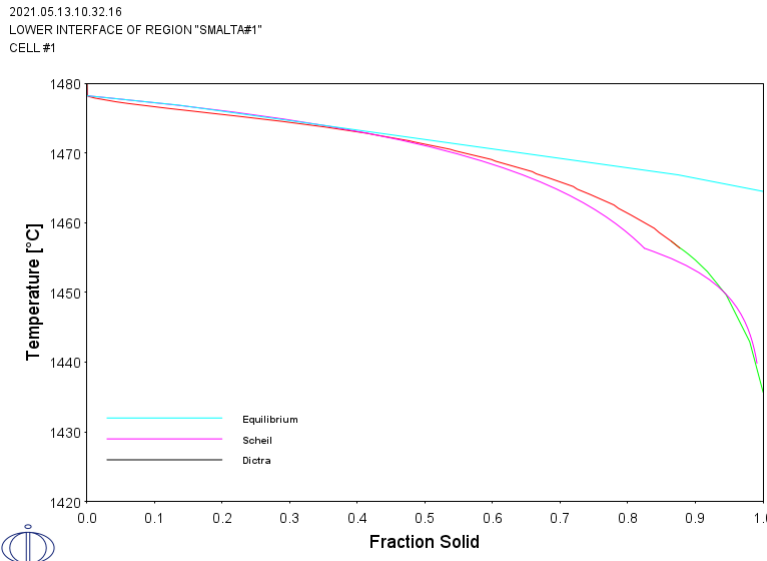

--OK--

DIC>

exb4b-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4b\plot.DCM.test"
DIC>
DIC>
DIC> @@ exb4b_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE b4b
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 2.00000E+02
DIC> read exb4b
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: set-title Fe-18%Cr-8%Ni
POST-1:
POST-1: @@
POST-1: @@ PLOT THE FRACTION OF SOLID AND COMPARE WITH A SCHEIL-GULLIVER
POST-1: @@ SIMULATION AND EQUILIBRIUM SOLIDIFICATION (DATA ON FILE exb4.exp)
POST-1: @@
POST-1: enter func fs=1-ivv(liquid);
POST-1: s-d-a x fs
POST-1: s-s-s x n 0 1
POST-1: s-ax-te x n Fraction Solid
POST-1: 
POST-1: s-d-a y t-c
POST-1: s-s-s y n 1420 1480
POST-1: 
POST-1: s-p-c interf smalta lower
POST-1: 
POST-1: app y exb4b.exp 0; 1
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...orking ...          OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Fe-18%Cr-8%Ni



```
POST-1:
POST-1:
POST-1:
POST-1: set-inter
--OK---
POST-1:
```



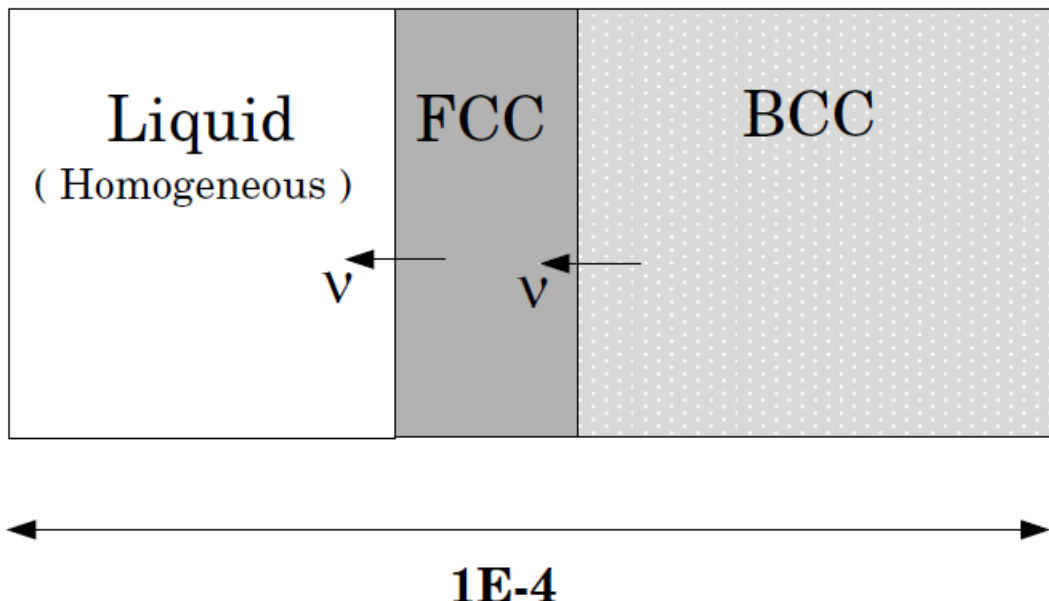
Example exb4c

Solidification path of an Fe-18%Cr-8%Ni alloy: Peritectic reaction, homogeneous liquid

This example is the same as exb4b but now the diffusivity data is amended for the LIQUID and a very high value for the diffusivity is used in order to simulate a case where we assume that the composition in the LIQUID is always homogeneous. This case should be considered less realistic than exb4b. Comparison is made with both a Scheil-Gulliver simulation and equilibrium solidification conditions, both made with Thermo-Calc.

Time > 0

$$T = 1900 - 1 * \text{Time} \text{ K}$$



exb4c-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4c\setup.DCM.test"
SYS: @@
SYS: @@ Moving boundary problem.
SYS: @@ Solidification path of an Fe-18%Cr-8%Ni alloy
SYS: @@ This example is the same as exb4b but now the diffusivity data is amended
SYS: @@ for the LIQUID and a high value for the diffusivity is used to simulate a
SYS: @@ case where it is assumed that the composition in the LIQUID is always
SYS: @@ homogeneous. This example is less realistic than exb4b.
SYS: @@ Comparison is made with both a Scheil-Gulliver simulation and equilibrium
SYS: @@ solidification conditions, both done in Thermo-Calc.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exb4c_setup.DCM
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: sw tcfef9
Current database: Steels/Fe-Alloys v9.3

VA           /- DEFINED
L12_FCC      B2_BCC          DICTRA_FCC_A1
REJECTED
TDB_TCFE9:
TDB_TCFE9: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_TCFE9: def-sys fe ni cr
FE           NI             CR
DEFINED
TDB_TCFE9:
TDB_TCFE9: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_TCFE9: rej ph /all
LIQUID:L      BCC_A2          FCC_A1
HCP_A3        CBC_A13         CUB_A13
SIGMA         CHI_A12         LAVES_PHASE_C14
CR3SI         NBNI3          NI3TI
CRZN17        BETA1          GAMMA
AL5FE4        FLUORITE_C1:I ZRO2_TETR:I
M2O3C:I       CENI2          CENI5
REJECTED
TDB_TCFE9: res ph fcc liq bcc
FCC_A1        LIQUID:L       BCC_A2
RESTORED
TDB_TCFE9:
TDB_TCFE9: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_TCFE9: get
10:34:09,382 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
FUNCTIONS .....

List of references for assessed data

'A. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'X.-G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'X.-G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
    volumes'
'J. Brillo and I. Egry, Int. J. Thermophysics, 24, 1155-1170'
'B.-J. Lee, CALPHAD, 17 (1993) 251-268; revision of Fe-Cr and Fe-Ni liquid'
'J-O. Andersson and B. Sundman, CALPHAD, 11 (1987) 83-92; TRITA 0270
    (1986); CR-FE'
'B.-J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'A. Dinsdale and T. Chart, MTDS NPL, Unpublished work (1986); CR-NI'
'A. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
-OK-
TDB_TCFE9:
TDB_TCFE9: @@
TDB_TCFE9: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_TCFE9: @@ SWITCH TO A MOBILITY DATABASE AND APPEND THE DATA.
TDB_TCFE9: @@
TDB_TCFE9: app
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
```

FEDAT = TCS/TT Steels Database v1.0
 TCNI11 = Ni-Alloys v11.0
 TCNI10 = Ni-Alloys v10.0
 TCNI9 = Ni-Alloys v9.1
 TCNI8 = Ni-Alloys v8.2
 TCNI7 = Ni-Alloys v7.2
 TCNI6 = Ni-Alloys v6.1
 TCNI5 = Ni-Alloys v5.1
 TCNI4 = Ni-Alloys v4.0
 TCNI1 = Ni-Alloys v1.3
 TCAL8 = Al-Alloys v8.0
 TCAL7 = Al-Alloys v7.1
 TCAL6 = Al-Alloys v6.0
 TCAL5 = Al-Alloys v5.1
 TCAL4 = Al-Alloys v4.0
 TCAL3 = Al-Alloys v3.0
 TCAL2 = Al-Alloys v2.1
 TCAL1 = Al-Alloys v1.2
 TCMG6 = Mg-Alloys v6.1
 TCMG5 = Mg-Alloys v5.1
 TCMG4 = Mg-Alloys v4.0
 TCMG3 = Mg-Alloys v3.0
 TCMG2 = Mg-Alloys v2.0
 TCMG1 = Mg-Alloys v1.1
 TCTI3 = Ti-Alloys v3.0
 TCTI2 = Ti-Alloys v2.2
 TCTI1 = Ti-Alloys v1.0
 TCCU4 = Cu-Alloys v4.0
 TCCU3 = Cu-Alloys v3.1
 TCCU2 = Cu-Alloys v2.0
 TCCU1 = Cu-Alloys v1.0
 TCCC1 = Cemented carbide v1.0
 TCHEA5 = High Entropy Alloy v5.0
 TCHEA4 = High Entropy Alloy v4.2
 TCHEA3 = High Entropy Alloy v3.1
 TCHEA2 = High Entropy Alloy v2.1.1
 TCHEA1 = High Entropy Alloy v1.0
 SSOL7 = SGTE Alloy Solutions Database v7.0
 SSOL6 = SGTE Alloy Solutions Database v6.0
 SSOL5 = SGTE Alloy Solutions Database v5.0
 SSOL4 = SGTE Alloy Solutions Database v4.9g
 SSOL2 = SGTE Alloy Solutions Database v2.1
 SSUB6 = SGTE Substances Database v6.0
 SSUB5 = SGTE Substances Database v5.2
 SSUB4 = SGTE Substances Database v4.1
 SSUB3 = SGTE Substances Database v3.3
 SSUB2 = SGTE Substances Database v2.2
 SNOB3 = SGTE Noble Metal Alloys Database v3.1
 SNOB2 = SGTE Noble Metal Alloys Database v2.1
 SNOB1 = SGTE Noble Metal Alloys Database v1.2
 STBC2 = SGTE Thermal Barrier Coating TDB v2.2
 STBC1 = SGTE Thermal Barrier Coating TDB v1.1
 SALT1 = SGTE Molten Salts Database v1.3
 SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
 SEMC2 = TC Semi-Conductors v2.1
 SLAG4 = Fe-containing Slag v4.1
 SLAG3 = Fe-containing Slag v3.2
 SLAG2 = Fe-containing Slag v2.2
 SLAG1 = Fe-containing Slag v1.2
 TCOX11 = Metal Oxide Solutions v11.0
 TCOX10 = Metal Oxide Solutions v10.1
 TCOX9 = Metal Oxide Solutions v9.0
 TCOX8 = Metal Oxide Solutions v8.0
 TCOX7 = Metal Oxide Solutions v7.0
 TCOX6 = Metal Oxide Solutions v6.0
 TCOX5 = Metal Oxide Solutions v5.1
 TCOX4 = Metal Oxide Solutions v4.1
 ION3 = Ionic Solutions v3.0
 ION2 = Ionic Solutions v2.6
 ION1 = Ionic Solutions v1.5
 NOX2 = NPL Oxide Solutions Database v2.1
 TCNOBL1 = Noble Metals Alloys v1.0
 TCNOBL2 = Noble Metals Alloys v2.0
 TCSLD4 = Solder Alloys v4.0
 TCSLD3 = Solder Alloys v3.3
 TCSLD2 = Solder Alloys v2.0
 TCSLD1 = Solder Alloys v1.1
 TCSI1 = Ultrapure Silicon v1.2
 TCMP2 = Materials Processing v2.5
 TCES1 = Combustion/Sintering v1.1
 TCSC1 = Super Conductor v1.0
 TCFCl = SOFC Database v1.0
 TCNF2 = Nuclear Fuels v2.1b
 NUMT2 = Nuclear Materials v2.1
 NUOX4 = Nuclear Oxides v4.2
 NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
 NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
 NUCL19 = IRSN NUCLEA-19
 NUCL15 = IRSN NUCLEA-15_4
 NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
 MEPH19 = IRSN Mephista-19
 MEPH15 = IRSN Mephista-15_1
 MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
 TCAQ3 = Aqueous Solution v3.0
 TCAQ2 = Aqueous Solution v2.7
 AQS2 = TGG Aqueous Solution Database v2.6
 GCE2 = TGG Geochemical/Environmental TDB v2.3
 FEDEMO = Iron Demo Database v4.0
 ALDEMO = Aluminum Demo Database v4.0
 NI DEMO = Nickel Demo Database v2.0
 CUDEMO = Copper Demo Database v1.0
 SLDEMO = Solder Demo Database v1.0
 OXDEMO = Oxide Demo Database v3.0
 SUBDEMO = Substance Demo Database v1.0
 PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
 PG35 = PG35 Binary Semi-Conductors TDB v1.3
 PURE5 = SGTE Unary (Pure Elements) TDB v5.2
 MOB2 = Alloys Mobility v2.7
 MOB1 = Alloys Mobility v1.3
 MOBFE1 = Steels/Fe-Alloys Mobility v1.1
 MOBFE2 = Steels/Fe-Alloys Mobility v2.0
 MOBFE3 = Steels/Fe-Alloys Mobility v3.0
 MOBFE4 = Steels/Fe-Alloys Mobility v4.0
 MOBFE5 = Steels/Fe-Alloys Mobility v5.0
 MOBFE6 = Steels/Fe-Alloys Mobility v6.0
 MOBNI5 = Ni-Alloys Mobility v5.1

```

MOBNI4 = Ni-Alloys Mobility v4.1
MOBNI3 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBSI1 = Si-Alloys Mobility v1.0
MOBSLD1 = Solder-Alloys Mobility v1.1
MOBT14 = Ti-Alloys Mobility v4.0
MOBT13 = Ti-Alloys Mobility v3.1
MOBT12 = Ti-Alloys Mobility v2.0
MOBT11 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

```

```

DATABASE NAME /TCFE9/: mobfe4
Current database: Steels/Fe-Alloys Mobility v4.0

```

```

VA DEFINED
B2_BCC REJECTED
APP: def-sys fe ni cr
    FE                      NI                      CR
    DEFINED
APP: rej ph /all          FCC_A1                  HCP_A3
    BCC_A2                  LIQUID:L
    LIQUID:L REJECTED
APP: res ph fcc liq bcc
    FCC_A1                  LIQUID:L
    BCC_A2
    RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

```

List of references for assessed data

```

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Ni diffusion fcc Cr-Ni'
'B. Jonsson: Z. Metallkunde 86(1995)686-692; Cr, Fe and Ni diffusion fcc
    Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
    -Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
'B. Jonsson: ISIJ International, 35(1995)1415-1421; Cr, Fe and Ni
    diffusion bcc Cr-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
    in bcc Fe'
'No assessed or estimated parameters'
'Estimation by using the modified Sutherland equation from Chen et al.,
    Phil. Magazine 94 (2014) 1552.'
'L. Zhang et al., Acta Mater., 58(2010)3664.'

```

-OK-

```

APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: go d-m
NO TIME STEP DEFINED

```

```

DIC>
DIC> @@
DIC> @@ LIST THE MOBILITIES IN THE LIQUID
DIC> @@
DIC> list-mobility-data
Sorry, LIST-DATA disabled for this database

```

```

DIC>
DIC>
DIC> liquid
NO SUCH COMMAND, USE HELP
DIC>
DIC>
DIC> @@
DIC> @@ AMEND THE DIFFUSIVITY DATA IN THE LIQUID
DIC> @@
DIC> @@ CHANGE TO A DIFFUSIVITY THAT IS 1000 TIMES HIGHER THAN THE
DIC> @@ VALUE IN THE MOBILITY DATABASE. THIS SHOULD BE ENOUGH IN ORDER TO
DIC> @@ ASSUME THAT THE COMPOSITION IN THE LIQUID IS AT ALL TIMES HOMOGENEOUS.
DIC> @@

```

```

DIC> amend_mobility_data
PARAMETER:
*** ERROR, PLEASE RE-ENTER EACH PART SEPARATELY

```

```

IDENTIFIER: dq
PHASE NAME: liquid&cr
CONSTITUENT: cr
INTERACTING CONSTITUENT:

```

```

DQ(LIQUID&CR#1,CR;0) = Sorry, database encrypted
DO YOU WANT TO CHANGE THE NUMBER OF RANGES /NO/: yes
I AM SORRY BUT YOU MUST THEN REENTER ALL RANGES

```

```

DQ(LIQUID&CR#1,CR;0) =

```

```

LOW TEMPERATURE LIMIT /298.15/: 298.15

```

```

FUNCTION: +R*T*LN(1E-06);

```

```

HIGH TEMPERATURE LIMIT /6000/: 6000

```

```

ANY MORE RANGES /N/: no

```

```

*** WARNING, NO MAGNETIC CONTRIBUTION DEFINED

```

```

DIC>
DIC> amend_mobility_data
PARAMETER: dq(liquid&cr,fe;0)
DQ(LIQUID&CR#1,FE;0) = Sorry, database encrypted
DO YOU WANT TO CHANGE THE NUMBER OF RANGES /NO/: y
I AM SORRY BUT YOU MUST THEN REENTER ALL RANGES
DQ(LIQUID&CR#1,FE;0) =
LOW TEMPERATURE LIMIT /298.15/: 298.15 +R*T*LN(1E-06); 6000 n

*** WARNING, NO MAGNETIC CONTRIBUTION DEFINED

DIC>
DIC> am-mob dq(liquid&cr,ni;0)
DQ(LIQUID&CR#1,NI;0) = Sorry, database encrypted
DO YOU WANT TO CHANGE THE NUMBER OF RANGES /NO/: y
I AM SORRY BUT YOU MUST THEN REENTER ALL RANGES
DQ(LIQUID&CR#1,NI;0) =
LOW TEMPERATURE LIMIT /298.15/: 298.15 +R*T*LN(1E-06); 6000 n

*** WARNING, NO MAGNETIC CONTRIBUTION DEFINED

DIC>
DIC> am-mob dq(liquid&ni,cr;0)
DQ(LIQUID&NI#1,CR;0) = Sorry, database encrypted
DO YOU WANT TO CHANGE THE NUMBER OF RANGES /NO/: y
I AM SORRY BUT YOU MUST THEN REENTER ALL RANGES
DQ(LIQUID&NI#1,CR;0) =
LOW TEMPERATURE LIMIT /298.15/: 298.15 +R*T*LN(1E-06); 6000 n

*** WARNING, NO MAGNETIC CONTRIBUTION DEFINED

DIC>
DIC> am-mob dq(liquid&ni,ni;0)
DQ(LIQUID&NI#1,NI;0) = Sorry, database encrypted
DO YOU WANT TO CHANGE THE NUMBER OF RANGES /NO/: y
I AM SORRY BUT YOU MUST THEN REENTER ALL RANGES
DQ(LIQUID&NI#1,NI;0) =
LOW TEMPERATURE LIMIT /298.15/: 298.15 +R*T*LN(1E-06); 6000 n

*** WARNING, NO MAGNETIC CONTRIBUTION DEFINED

DIC>
DIC> am-mob dq(liquid&fe,cr;0)
DQ(LIQUID&FE#1,CR;0) = Sorry, database encrypted
DO YOU WANT TO CHANGE THE NUMBER OF RANGES /NO/: y
I AM SORRY BUT YOU MUST THEN REENTER ALL RANGES
DQ(LIQUID&FE#1,CR;0) =
LOW TEMPERATURE LIMIT /298.15/: 298.15 +R*T*LN(1E-06); 6000 n

*** WARNING, NO MAGNETIC CONTRIBUTION DEFINED

DIC>
DIC> am-mob dq(liquid&fe,fe;0)
DQ(LIQUID&FE#1,FE;0) = Sorry, database encrypted
DO YOU WANT TO CHANGE THE NUMBER OF RANGES /NO/: y
I AM SORRY BUT YOU MUST THEN REENTER ALL RANGES
DQ(LIQUID&FE#1,FE;0) =
LOW TEMPERATURE LIMIT /298.15/: 298.15 +R*T*LN(1E-06); 6000 n

*** WARNING, NO MAGNETIC CONTRIBUTION DEFINED

DIC>
DIC> li-mob
AMBIGUOUS COMMAND, USE HELP
DIC>
DIC>
DIC> liquid
NO SUCH COMMAND, USE HELP
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> @@ LOWER THE TEMPERATURE TO A RATE OF 1 K/s
DIC> set-cond glob T 0 1900-1*TIME; * N
DIC>
DIC> @@
DIC> @@ ENTER A REGION CALLED smalta
DIC> @@
DIC> enter-region smalta
DIC>
DIC> @@
DIC> @@ ENTER A GEOMETRIC GRID INTO THE REGION
DIC> @@
DIC> enter-grid

```

```

REGION NAME : /SMALTA/: smalta
WIDTH OF REGION /1/: 1e-4
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER active PHASES INTO THE REGION
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /SMALTA/: smalta
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: liq
DIC>
DIC> @@
DIC> @@ ENTER inactive PHASES INTO THE REGION, BOTH PHASES ON THE SAME SIDE
DIC> @@ OF THE LIQUID REGION IN ORDER TO GET A PERITECTIC REACTION.
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /SMALTA/: smalta
ATTACHED TO THE RIGHT OF SMALTA /YES/: yes
PHASE NAME: /NONE/: fcc#1
DEPENDENT COMPONENT ? /NI/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-5
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /SMALTA/: smalta
ATTACHED TO THE RIGHT OF SMALTA /YES/: yes
PHASE NAME: /NONE/: bcc#1
DEPENDENT COMPONENT ? /NI/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-5
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC> @@
DIC> @@ ENTER THE START COMPOSITION FOR THE LIQUID
DIC> @@
DIC> enter-composition
REGION NAME : /SMALTA/: smalta
PHASE NAME: /LIQUID/: liq
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /CR/: cr lin 18 18
PROFILE FOR /NI/: ni lin 8 8
10:34:33,022 INFO Preparing system for use: MOBFE4_MODIFIED_162089485851912
10:34:35,739 INFO Preparing phase for use: LIQUID
DIC>
DIC> @@
DIC> @@ THE BOUNDARY CONDITION IS A CLOSED SYSTEM (DEFAULT) AS WE DO NOT SPECIFY
DIC> @@ ANYTHING ELSE
DIC> @@
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 200
AUTOMATIC TIMESTEP CONTROL /YES/: yes
MAX TIMESTEP DURING INTEGRATION /20/: 1
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ CHECK THE INTERFACE POSITION. THIS IS TO MAKE SURE THAT THE LIQUID
DIC> @@ REGION DOES NOT SHRINK TOO MUCH DURING A TIMESTEP. IN ADDITION THE TIMESTEP
DIC> @@ IS CONTROLLED BY THE PHASE INTERFACE DISPLACEMENT DURING THE SIMULATION.
DIC> @@
DIC> s-s-c
NSOIA PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA Timesteps IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/: yes
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/:
DEGREE OF IMPLICITY WHEN INTEGRATING PDEs (AUTO, 0 -> 0.5 -> 1): /AUTO/:
MAX TIMESTEP CHANGE PER TIMESTEP : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/: @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exb4c Y
DIC>
DIC> set-inter
--OK---
DIC>

```

exb4c-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4c\run.DCM.test"
DIC>
DIC> @@ exb4c_run.DCM
DIC>
DIC> @@
DIC> FILE FOR RUNNING EXAMPLE b4b
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE SET UP FROM FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exb4c
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim yes
Region: SMALTA
single geometric dense at 0.10000E-03
0.99375 62
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
10:35:34,423 INFO Preparing phase for use: BCC_A2
10:35:35,276 INFO Preparing phase for use: FCC_A1
TIME = 0.10000000E-06 DT = 0.10000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 2 seconds
TIME = 0.10773101E-05 DT = 0.97731005E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 0.30319302E-05 DT = 0.19546201E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.694411704E-05 DT = 0.39092402E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 0.14759651E-04 DT = 0.78184804E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.30396612E-04 DT = 0.15636961E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 0.61670533E-04 DT = 0.31273922E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.12421838E-03 DT = 0.62547843E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.24931406E-03 DT = 0.12509569E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.49950544E-03 DT = 0.25019137E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.99988818E-03 DT = 0.50038275E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.20006537E-02 DT = 0.10007655E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992482 FE = .733068011219293
NI = .0754116207882257
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.40021847E-02 DT = 0.20015310E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992482 FE = .733068011219293
NI = .0754116207882259
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.80052466E-02 DT = 0.40030620E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992478 FE = .733068011219298
NI = .0754116207882241
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.16011371E-01 DT = 0.80061240E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992479 FE = .733068011219296
NI = .0754116207882191
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.32023619E-01 DT = 0.16012248E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992493 FE = .733068011219288
NI = .0754116207882191
```

```

TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.64048114E-01 DT = 0.32024496E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992497 FE = .733068011219264
NI = .0754116207882391
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.12809711 DT = 0.64048992E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992473 FE = .733068011219282
NI = .0754116207882449
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.25619509 DT = 0.12809798 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .191520367992311 FE = .733068011219435
NI = .0754116207882534

```

output ignored...

... output resumed

```

1.699448762847264E-008 3.196946048987361E-009 6.365914350669771E-010 8.137206590427579E-011 4.822931506030856E-
015 3.642429876836610E-016 TIME = 199.16001 DT = 1.0000000 SUM OF SQUARES = 0.36424299E-15
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.10042491E-06 AND -0.10042491E-06
POSITION OF INTERFACE SMALTA / R_FCC_A1 IS 0.46634068E-06
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.11494123E-07 AND 0.11494123E-07
POSITION OF INTERFACE R_FCC_A1 / R_BCC_A2 IS 0.10974304E-04
U-FRACTION IN SYSTEM: CR = .191209953993244 FE = .733434883964446
NI = .0753551620423105
TOTAL SIZE OF SYSTEM: 1E-04 [m]
33 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_FCC_A1

CPU time used in timestep 3 seconds
3.727050292694413E-007 3.732636074917638E-007 3.727007199184641E-007 3.726201750127047E-007 3.725913590737159E-
007 1.890188648825606E-008 6.899777276244959E-010 9.886941641614766E-013 3.184161732385594E-
016 TIME = 200.00000 DT = 0.83999467 SUM OF SQUARES = 0.31841617E-15
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.85596223E-07 AND -0.85596223E-07
POSITION OF INTERFACE SMALTA / R_FCC_A1 IS 0.39444031E-06
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.13575622E-07 AND 0.13575622E-07
POSITION OF INTERFACE R_FCC_A1 / R_BCC_A2 IS 0.10985707E-04
U-FRACTION IN SYSTEM: CR = .191209945953057 FE = .733434891748842
NI = .0753551622981016
TOTAL SIZE OF SYSTEM: 1E-04 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 170.32213
DELETING TIME-RECORD FOR TIME 170.48228
DELETING TIME-RECORD FOR TIME 170.48229
DELETING TIME-RECORD FOR TIME 170.48229
DELETING TIME-RECORD FOR TIME 170.48229
DELETING TIME-RECORD FOR TIME 170.48230
DELETING TIME-RECORD FOR TIME 170.48231
DELETING TIME-RECORD FOR TIME 170.48233
DELETING TIME-RECORD FOR TIME 170.48239
DELETING TIME-RECORD FOR TIME 170.48249
DELETING TIME-RECORD FOR TIME 170.48269
DELETING TIME-RECORD FOR TIME 170.48310
DELETING TIME-RECORD FOR TIME 170.48392
DELETING TIME-RECORD FOR TIME 170.48556
DELETING TIME-RECORD FOR TIME 170.48884
DELETING TIME-RECORD FOR TIME 170.49539
DELETING TIME-RECORD FOR TIME 170.50850
DELETING TIME-RECORD FOR TIME 170.53471
DELETING TIME-RECORD FOR TIME 170.58714
DELETING TIME-RECORD FOR TIME 170.69200
DELETING TIME-RECORD FOR TIME 170.90171
DELETING TIME-RECORD FOR TIME 171.32114
DELETING TIME-RECORD FOR TIME 172.16001
DELETING TIME-RECORD FOR TIME 173.16001
DELETING TIME-RECORD FOR TIME 174.16001
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DELETING TIME-RECORD FOR TIME 193.16001
DELETING TIME-RECORD FOR TIME 194.16001
DELETING TIME-RECORD FOR TIME 195.16001
DELETING TIME-RECORD FOR TIME 196.16001
DELETING TIME-RECORD FOR TIME 197.16001
DELETING TIME-RECORD FOR TIME 198.16001
KEEPING TIME-RECORD FOR TIME 199.16001
AND FOR TIME 200.00000
WORKSPACE RECLAIMED
TIMESTEP AT 200.000000 SELECTED

```

DIC>
DIC>
DIC>
DIC>
DIC>
DIC>
DIC>
DIC>

```
DIC>
DIC> @@  
DIC> @@ THE SIMULATION IS FINISHED  
DIC> @@  
DIC>  
DIC> set-inter  
--OK---  
DIC>
```

exb4c-plot

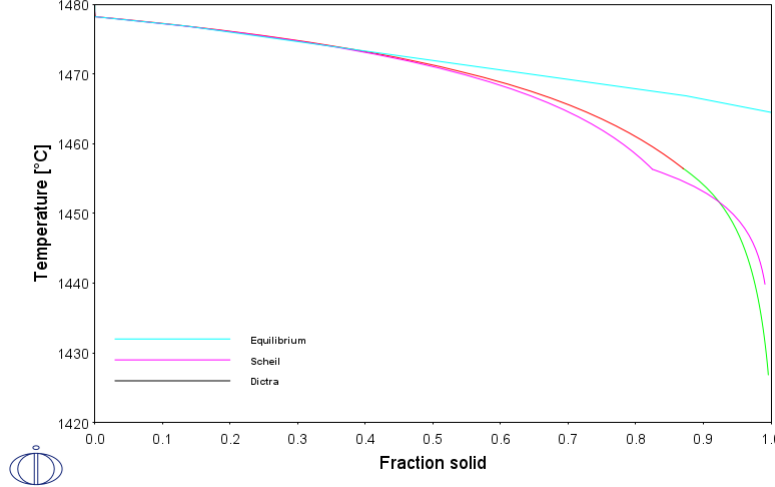
```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4c\plot.DCM.test"
DIC>
DIC>
DIC> @@ exb4c_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE b4c
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 2.00000E+02
DIC> read exb4c
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: set-title Fe-18%Cr-8%Ni
POST-1:
POST-1: @@
POST-1: @@ PLOT THE FRACTION OF SOLID AND COMPARE WITH A SCHEIL-GULLIVER
POST-1: @@ SIMULATION AND EQUILIBRIUM SOLIDIFICATION (DATA ON FILE exb4.exp).
POST-1: @@ IN THIS CASE WE CAN SEE THAT ALL THREE LINES INITIALLY FALL
POST-1: @@ ON THE SAME LINE.
POST-1: @@
POST-1: enter func fs=1-ivv(liquid);
POST-1: s-d-a x fs
POST-1: s-s-s x n 0 1
POST-1: s-ax-te x n Fraction solid
POST-1:
POST-1: s-d-a y t-c
POST-1: s-s-s y n 1420 1480
POST-1:
POST-1: s-p-c interf smalta lower
POST-1:
POST-1: app y exb4c.exp 0; 1
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Fe-18%Cr-8%Ni

2021.05.13.10.40.29

LOWER INTERFACE OF REGION "SMALTA#1"

CELL #1



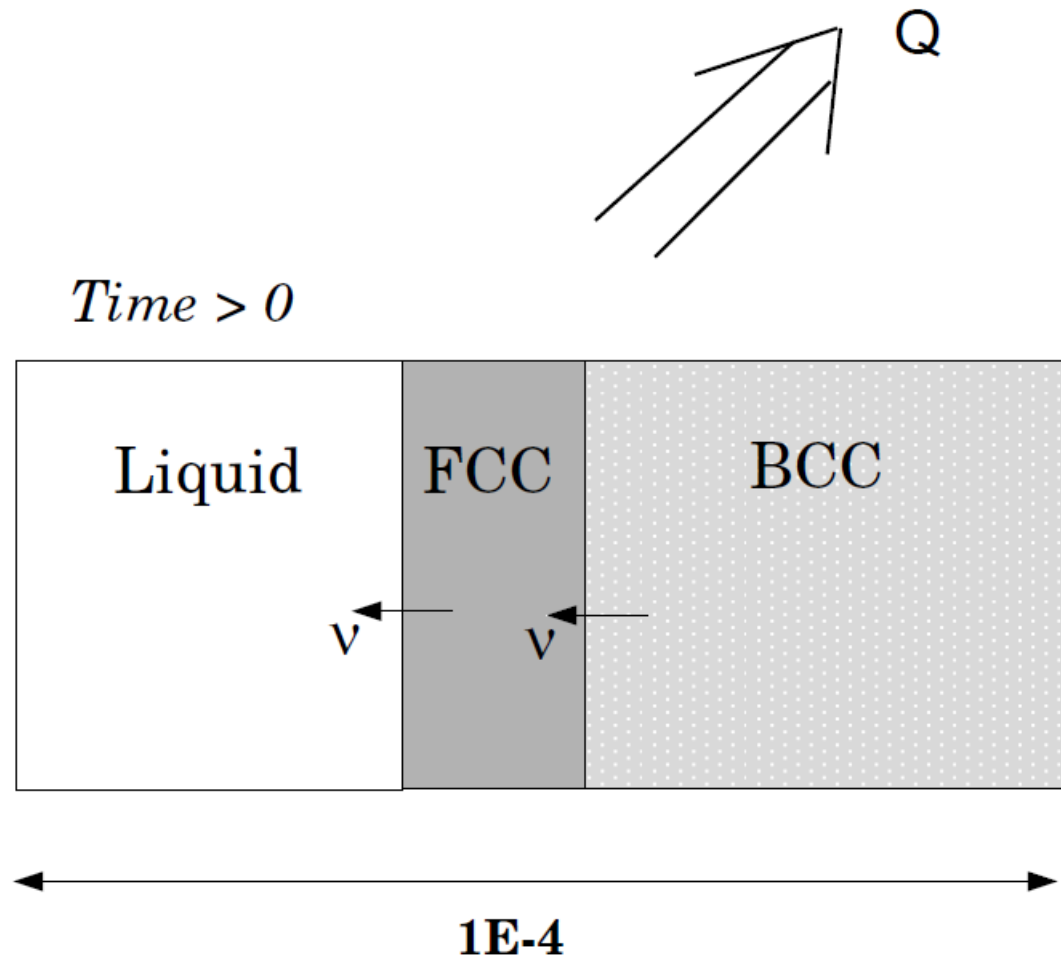
```
POST-1:
POST-1:
POST-1:
POST-1: set-inter
--OK--
POST-1:
```



Example exb4d

Solidification path of an Fe-18%Cr-8%Ni alloy: Peritectic reaction, heat-flux controls the temperature

This example is the same as exb4b but instead of controlling the temperature the amount heat extracted is given. Comparison is made with both a Scheil-Gulliver simulation and equilibrium solidification conditions, both made with Thermo-Calc.



exb4d-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4d\setup.DCM.test"
SYS: @@
SYS: @@ Moving boundary problem.
SYS: @@ Solidification path of an Fe-18%Cr-8%Ni alloy
SYS: @@ This example is the same as exb4b but instead of controlling the temperature
SYS: @@ the amount of heat extracted is given. Comparison is made with both a
SYS: @@ Scheil-Gulliver simulation and equilibrium solidification conditions,
SYS: @@ both done in Thermo-Calc.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exb4d_setup.DCM
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA          /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: sw fedemo
Current database: Iron Demo Database v4.0

VA          /- DEFINED
TDB_FEDEMO: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_FEDEMO: def-sys fe ni cr
FE          NI          CR
DEFINED
TDB_FEDEMO:
TDB_FEDEMO: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_FEDEMO: rej ph /all
LIQUID:L      BCC_A2      LAVES_PHASE_C14
CBCC_A12     CHI_A12     CUB_A13
FCC_A1       HCP_A3      SIGMA
REJECTED
TDB_FEDEMO: res ph fcc liq bcc
FCC_A1       LIQUID:L    BCC_A2
RESTORED
TDB_FEDEMO:
TDB_FEDEMO: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_FEDEMO: get
10:41:55,012 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES ....
ELEMENTS ....
SPECIES ....
PHASES .....
PARAMETERS ...
FUNCTIONS ...

List of references for assessed data

'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'M. Ghasemi, Thermo-Calc Software AB, Sweden, 2020; Unifying liquid volume
database'
'M. Ghasemi, Thermo-Calc Software AB, Sweden, 2019; Viscosity of metallic
liquid'
'M. Ghasemi, Thermo-Calc Software AB: assessing the surface tension of
metallic liquid'
'Y. Tang, in TCCU2, Thermo-Calc Software AB, Stockholm, Sweden'
'M. J. Assael, J. Phys. Chem. Ref. Data 35 (2006) 285; Fe, Al'
'M. J. Assael, High Temp. High Press., 41 (2012); Sb, Pb, Bi, Ni, Ag'
'B.J. Lee, CALPHAD, 17 (1993) 251-268; revision of Fe-Cr and Fe -Ni liquid'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'J-O. Andersson and B. Sundman, CALPHAD, 11 (1987) 83-92; TRITA 0270
(1986); CR-FE'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); CR -NI'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
-OK-
TDB_FEDEMO: @@
TDB_FEDEMO: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_FEDEMO: @@ SWITCH TO THE MOBILITY DATABASE AND APPEND THE DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
```

TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2
TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
SNOB1 = SGTE Noble Metal Alloys Database v1.2
STBC2 = SGTE Thermal Barrier Coating TDB v2.2
STBC1 = SGTE Thermal Barrier Coating TDB v1.1
SALT1 = SGTE Molten Salts Database v1.3
SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2 = TC Semi-Conductors v2.1
SLAG4 = Fe-containing Slag v4.1
SLAG3 = Fe-containing Slag v3.2
SLAG2 = Fe-containing Slag v2.2
SLAG1 = Fe-containing Slag v1.2
TCOX11 = Metal Oxide Solutions v11.0
TCOX10 = Metal Oxide Solutions v10.1
TCOX9 = Metal Oxide Solutions v9.0
TCOX8 = Metal Oxide Solutions v8.0
TCOX7 = Metal Oxide Solutions v7.0
TCOX6 = Metal Oxide Solutions v6.0
TCOX5 = Metal Oxide Solutions v5.1
TCOX4 = Metal Oxide Solutions v4.1
ION3 = Ionic Solutions v3.0
ION2 = Ionic Solutions v2.6
ION1 = Ionic Solutions v1.5
NOX2 = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4 = Solder Alloys v4.0
TCSLD3 = Solder Alloys v3.3
TCSLD2 = Solder Alloys v2.0
TCSLD1 = Solder Alloys v1.1
TCSI1 = Ultrapure Silicon v1.2
TCPMP2 = Materials Processing v2.5
TCES1 = Combustion/Sintering v1.1
TCSC1 = Super Conductor v1.0
TCFC1 = SOFC Database v1.0
TCNF2 = Nuclear Fuels v2.1b
NUMT2 = Nuclear Materials v2.1
NUOX4 = Nuclear Oxides v4.2
NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephista-19
MEPH15 = IRSN Mephista-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQ52 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SLDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.2
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3
MOBF1 = Steels/Fe-Alloys Mobility v1.1
MOBF2 = Steels/Fe-Alloys Mobility v2.0
MOBF3 = Steels/Fe-Alloys Mobility v3.0
MOBF4 = Steels/Fe-Alloys Mobility v4.0
MOBF5 = Steels/Fe-Alloys Mobility v5.0
MOBF6 = Steels/Fe-Alloys Mobility v6.0
MOBN15 = Ni-Alloys Mobility v5.1
MOBN14 = Ni-Alloys Mobility v4.1

```

MOBN13 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBSI1 = Si-Alloys Mobility v1.0
MOBSLD1 = Solder-Alloys Mobility v1.1
MOBTI4 = Ti-Alloys Mobility v4.0
MOBTI3 = Ti-Alloys Mobility v3.1
MOBTI2 = Ti-Alloys Mobility v2.0
MOBTI1 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

```

```

DATABASE NAME /FEDEMO/: mobfe2
Current database: Steels/Fe-Alloys Mobility v2.0
TCS Steel Mobility Database Version 2.0 from 2011-12-09.

```

VA DEFINED

*** WARNING: This database cannot be used with GES6, temporarily reverting to G
ESS5

```

APP: def-sys fe ni cr
    FE           NI          CR
    DEFINED
APP: rej ph /all
    BCC_A2      FCC_A1      HCP_A3
    LIQUID:L   REJECTED
APP: res ph fcc liq bcc
    FCC_A1      LIQUID:L   BCC_A2
    RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

```

List of references for assessed data

```

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Ni diffusion fcc Cr-Ni'
'B. Jonsson: Z. Metallkunde 86(1995)686-692; Cr, Fe and Ni diffusion fcc
Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
'B. Jonsson: ISIJ International, 35(1995)1415-1421; Cr, Fe and Ni
diffusion bcc Cr-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'

```

-OK-

```

APP: @@
APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: go d-m
NO TIME STEP DEFINED

```

```

DIC> @@
DIC> @@ EXTRACT HEAT 91.19 J/mole/s
DIC> @@
DIC> set-cond glob Q 0 91.19; * N
DIC>
DIC> @@

```

```

DIC> @@ ENTER AN INITIAL TEMPERATURE
DIC> @@
DIC> set-initial-temp 1900
DIC>

```

```

DIC> @@
DIC> @@ ENTER A REGION CALLED smalta
DIC> @@
DIC> enter-region smalta
DIC>
DIC> @@
DIC> @@ ENTER A GEOMETRIC GRID INTO THE REGION
DIC> @@
DIC> enter-grid

```

```

REGION NAME : /SMALTA/: smalta
WIDTH OF REGION /1/: 1e-4
TYPE /LINEAR/: AUTO

```

```

DIC>
DIC>
DIC> @@
DIC> @@ ENTER active PHASES INTO THE REGION
DIC> @@
DIC> enter-phase

```

```

ACTIVE OR INACTIVE PHASE /ACTIVE/: act

```

```

REGION NAME : /SMALTA/: smalta
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: liq

```

```

DIC>
DIC> @@
DIC> @@ ENTER inactive PHASES INTO THE REGION, BOTH PHASES ON THE
DIC> @@ SAME SIDE OF THE LIQUID REGION TO GET A PERITECTIC REACTION.
DIC> @@

```

```

DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /SMALTA/: smalta
ATTACHED TO THE RIGHT OF SMALTA /YES/: yes
PHASE NAME: /NONE/: fcc#1
DEPENDENT COMPONENT ? /NI/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-3
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /SMALTA/: smalta
ATTACHED TO THE RIGHT OF SMALTA /YES/: yes
PHASE NAME: /NONE/: bcc#1
DEPENDENT COMPONENT ? /NI/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-3
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC> @@
DIC> @@ ENTER A START COMPOSITION FOR THE LIQUID
DIC> @@
DIC> enter-composition
REGION NAME : /SMALTA/: smalta
PHASE NAME: /LIQUID/: liq
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /CR/: cr lin 18 18
PROFILE FOR /NI/: ni lin 8 8
DIC>
DIC> @@
DIC> @@ THE BOUNDARY CONDITION IS A CLOSED SYSTEM (DEFAULT) AS WE DO NOT SPECIFY
DIC> @@ ANYTHING ELSE
DIC> @@
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 200
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /20/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ CHECK THE INTERFACE POSITION. THIS IS TO MAKE SURE THAT THE LIQUID REGION
DIC> @@ DOES NOT SHRINK TOO MUCH DURING A TIMESTEP. IN ADDITION THE TIMESTEP IS
DIC> @@ CONTROLLED BY THE PHASE INTERFACE DISPLACEMENT DURING THE SIMULATION.
DIC> @@
DIC> s-s-c
NS01A PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA TIMESTEPS IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/: yes
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/: no
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/:
DEGREE OF IMPLICITY WHEN INTEGRATING PDEs (AUTO, 0 -> 0.5 -> 1): /AUTO/:
MAX TIMESTEP CHANGE PER TIMESTEP : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/: @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exb4d Y
DIC>
DIC> set-inter
---OK---
DIC>
```

exb4d-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4d\run.DCM.test"
DIC>
DIC> @@ exb4d_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE b4b
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE SET UP FROM FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exb4d
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim
Region: SMALTA
single geometric dense at 0.10000E-03
0.92197 87
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
TEMPERATURE: 1900. ENTHALPY: 0.7394E+05
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
TEMPERATURE: 1900. ENTHALPY: 0.7394E+05
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
3.863918610051234E-007 3.864691456616007E-007 3.837208091757153E-016 3.821106605297053E-025 TIME = 0.10000000E-006 DT = 0.10000000E-006 SUM OF SQUARES = 0.38211066E-24
TEMPERATURE: 1900. ENTHALPY: 0.7394E+05
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
9.508205000173971E-018 TIME = 0.26002472E-05 DT = 0.25002472E-05 SUM OF SQUARES = 0.95082050E-17
TEMPERATURE: 1900. ENTHALPY: 0.7394E+05
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882255
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
3.803283213789877E-017 TIME = 0.76007416E-05 DT = 0.50004944E-05 SUM OF SQUARES = 0.38032832E-16
TEMPERATURE: 1900. ENTHALPY: 0.7394E+05
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
1.521313188418321E-016 1.521313188418321E-016 1.521313188418321E-016 9.309554162466689E-020 1.431856013280617E-016
016 3.399417495083793E-017 3.399417495083793E-017 1.089145661025413E-019 1.089145661025413E-
019 1.089145661025413E-019 1.089145661025413E-019 2.673309085540699E-017 1.089153455095704E-
019 1.089153455095704E-019 1.089153455095704E-019 1.092199812185641E-
018 1.089145661025413E-019 TIME = 0.17601730E-04 DT = 0.10000989E-04 SUM OF SQUARES = 0.10891457E-18
TEMPERATURE: 1900. ENTHALPY: 0.7394E+05
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
1.666578045201216E-016 1.666578045201216E-016 1.666578045201216E-016 1.352478211635110E-012 1.666578045201216E-
016 7.573827019817665E-030 7.573827019817665E-030 7.573827019817665E-030 7.573827019817665E-
030 1.666578045201216E-016 5.562266575976920E-030 5.562266575976920E-030 5.562266575976920E-
030 5.562266575976920E-030 5.562266575976920E-030 5.562266575976920E-030 1.041609024951602E-
017 1.546768164561797E-029 TIME = 0.37603708E-04 DT = 0.20001978E-04 SUM OF SQUARES = 0.15467682E-28

ERROR RETURN FROM NS01A BECAUSE A NEARBY STATIONARY POINT OF F(X) IS PREDICTED
*** ERROR 1890 IN DCNS01: ERROR RETURN FROM NS01A
1.521313188418321E-016 1.521313188418321E-016 1.521313188418321E-016 9.309554162466689E-020 1.431856013280617E-
016 3.399417495083793E-017 3.399417495083793E-017 1.089145661025413E-019 1.089145661025413E-
019 1.089145661025413E-019 1.089145661025413E-019 2.673309085540699E-017 1.089153455095704E-
019 1.089153455095704E-019 1.089153455095704E-019 1.092199812185641E-
018 1.089145661025413E-019 TIME = 0.17601730E-04 DT = 0.10000989E-04 SUM OF SQUARES = 0.10891457E-18
TEMPERATURE: 1900. ENTHALPY: 0.7394E+05
U-FRACTION IN SYSTEM: CR = .191520367992483 FE = .733068011219292
NI = .0754116207882254
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
1.666578045201216E-016 1.666578045201216E-016 1.666578045201216E-016 1.352478211635110E-012 1.666578045201216E-
016 7.573827019817665E-030 7.573827019817665E-030 7.573827019817665E-030 7.573827019817665E-
030 1.666578045201216E-016 5.562266575976920E-030 5.562266575976920E-030 5.562266575976920E-
030 5.562266575976920E-030 5.562266575976920E-030 5.562266575976920E-030 1.041609024951602E-
017 1.546768164561797E-029 TIME = 0.37603708E-04 DT = 0.20001978E-04 SUM OF SQUARES = 0.15467682E-28

ERROR RETURN FROM NS01A BECAUSE A NEARBY STATIONARY POINT OF F(X) IS PREDICTED
*** ERROR 1890 IN DCNS01: ERROR RETURN FROM NS01A
1.666578045201216E-016 1.666578045201216E-016 1.666578045201216E-016 1.352478211635110E-012 1.666578045201216E-
016 7.573827019817665E-030 7.573827019817665E-030 7.573827019817665E-030 7.573827019817665E-
030 1.666578045201216E-016 5.562266575976920E-030 5.562266575976920E-030 5.562266575976920E-
030 5.562266575976920E-030 5.562266575976920E-030 5.562266575976920E-030 1.041609024951602E-
017 1.546768164561797E-029 TIME = 0.37603708E-04 DT = 0.20001978E-04 SUM OF SQUARES = 0.15467682E-28

output ignored...

... output resumed

TOTAL SIZE OF SYSTEM: 1E-04 [m]
6 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_FCC_A1

CPU time used in timestep 1 seconds
1.010611839641213E-007 1.010624112174118E-007 1.010620821966687E-007 1.010407815978725E-007 7.802524850104379E-
009 1.010731516713628E-013 8.491417896581680E-013 8.491417896581680E-013 8.491417896581680E-013
019 TIME = 197.90144 DT = 0.39375000 SUM OF SQUARES = 0.84914179E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.80357395E-07 AND 0.80357395E-07
POSITION OF INTERFACE R_FCC_A1 / R_BCC_A2 IS 0.53376600E-05
TEMPERATURE: 1702. ENTHALPY: 0.5587E+05
U-FRACTION IN SYSTEM: CR = .19142649256958 FE = .73317849065027
NI = .0753950167801494
TOTAL SIZE OF SYSTEM: 1E-04 [m]
14 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: R_FCC_A1

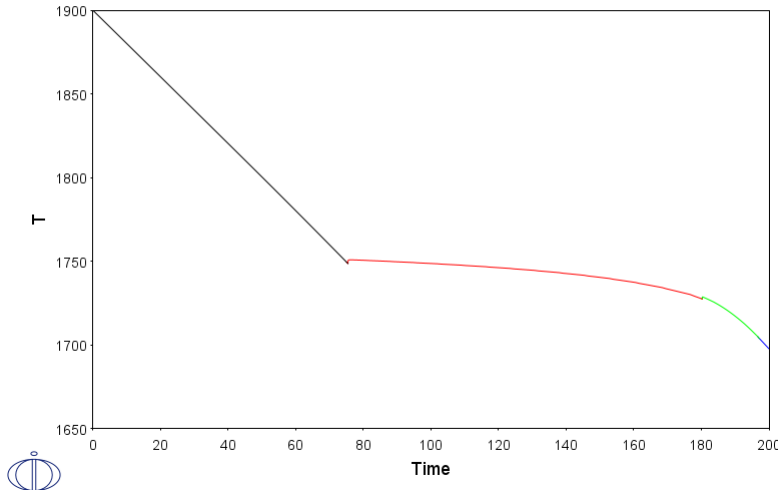
CPU time used in timestep 0 seconds
5.895191095916032E-007 5.896964303258992E-007 5.902402434406712E-007 5.893737193956263E-007 3.967197178262157E-
007 3.583238933502586E-007 2.901878750191593E-007 2.902122464542427E-007 1.672868017032316E-
007 2.285303868088257E-008 1.345973070721034E-014 6.74659362445023E-
018 TIME = 198.68894 DT = 0.78750000 SUM OF SQUARES = 0.67465936E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.89888651E-07 AND 0.89888651E-07
POSITION OF INTERFACE R_FCC_A1 / R_BCC_A2 IS 0.54084473E-05
TEMPERATURE: 1701. ENTHALPY: 0.5580E+05
U-FRACTION IN SYSTEM: CR = .191426492569597 FE = .733178490650262
NI = .0753950167801402
TOTAL SIZE OF SYSTEM: 1E-04 [m]
```


exb4d-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb4d\plot.DCM.test"
DIC>
DIC>
DIC> @@ exb4d_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE b4b
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 2.00000E+02
DIC> read exb4d
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: set-title Fe-18%Cr-8%Ni
POST-1:
POST-1: @@
POST-1: @@ PLOT THE FRACTION OF SOLID AND COMPARE WITH SCHEIL-GULLIVER
POST-1: @@ SIMULATION AND EQUILIBRIUM SOLIDIFICATION (DATA ON FILE exb4.exp)
POST-1: @@
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-d-a y T
POST-1: s-p-c inter first
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...orking ...          OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Fe-18%Cr-8%Ni

2021.05.13.10.47.39
"FIRST" INTERFACE OF SYSTEM
CELL #1



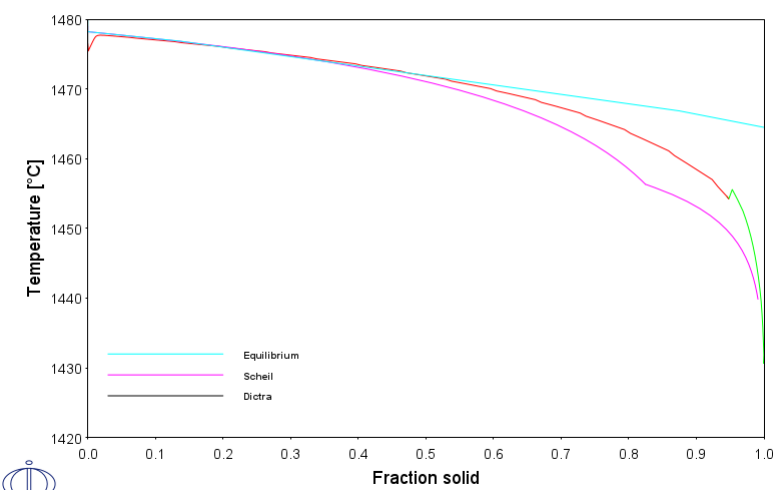
```
POST-1:
POST-1:Hit RETURN to continue
POST-1: enter func fs=1-ivv(liquid);
POST-1: s-d-a x fs
POST-1: s-s-s x n 0 1
POST-1: s-ax-te x n Fraction solid
POST-1:
POST-1: s-d-a y t-c
POST-1: s-s-s y n 1420 1480
POST-1:
POST-1: s-p-c interf smalta lower
POST-1:
POST-1: app y exb4d.exp 0; 1
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...orking ...          OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Fe-18%Cr-8%Ni

2021.05.13.10.47.42

LOWER INTERFACE OF REGION "SMALTA#1"

CELL #1



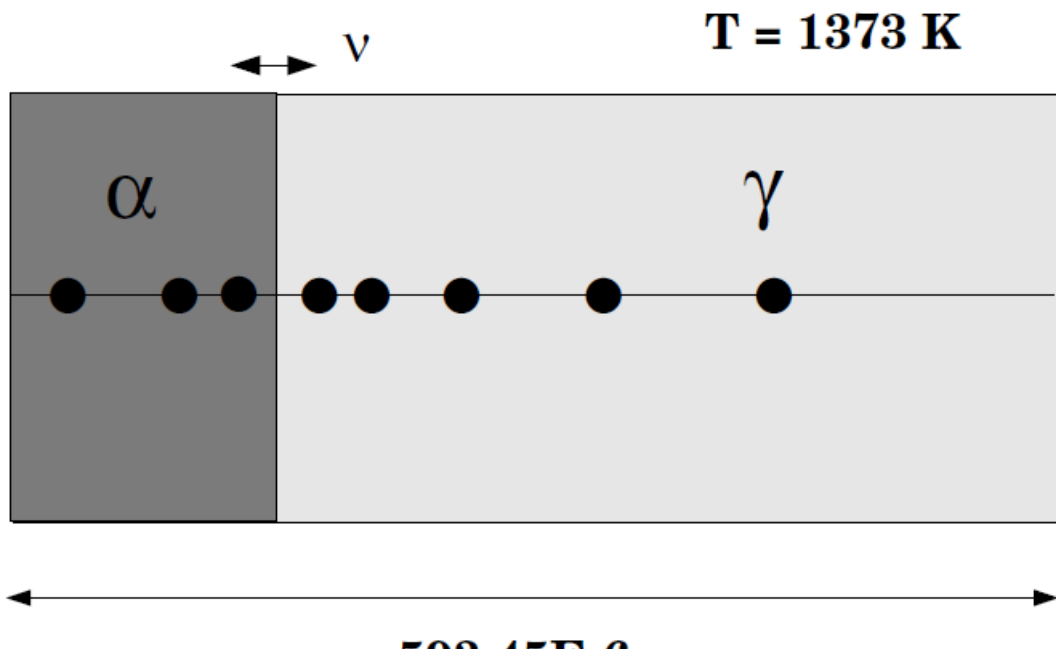
POST-1:
POST-1:
POST-1:
POST-1: set-inter
--OK---
POST-1:



Example exb5

$\gamma/\alpha/\gamma$ diffusion couple of Fe-Ni-Cr alloys

This example demonstrates the evaluation of a ternary Fe-Cr-Ni diffusion couple. A thin slice of α phase (38%Cr, 0%Ni) is clamped between two thicker slices of γ phase (27%Cr, 20%Ni). The assembly is subsequently heat treated at 1373K. This setup corresponds to diffusion couple A in M. Kajihara, C.-B. Lim and M. Kikuchi: ISIJ International 33 (1993), pp. 498-507. See also M. Kajihara and M. Kikuchi: Acta Metall.Mater. 41 (1993), pp.2045-2059.



TCMG4 = Mg-Alloys v4.0
 TCMG3 = Mg-Alloys v3.0
 TCMG2 = Mg-Alloys v2.0
 TCMG1 = Mg-Alloys v1.1
 TCTI3 = Ti-Alloys v3.0
 TCTI2 = Ti-Alloys v2.2
 TCTI1 = Ti-Alloys v1.0
 TCCU4 = Cu-Alloys v4.0
 TCCU3 = Cu-Alloys v3.1
 TCCU2 = Cu-Alloys v2.0
 TCCU1 = Cu-Alloys v1.0
 TCCC1 = Cemented carbide v1.0
 TCHEA5 = High Entropy Alloy v5.0
 TCHEA4 = High Entropy Alloy v4.2
 TCHEA3 = High Entropy Alloy v3.1
 TCHEA2 = High Entropy Alloy v2.1.1
 TCHEA1 = High Entropy Alloy v1.0
 SSOL7 = SGTE Alloy Solutions Database v7.0
 SSOL6 = SGTE Alloy Solutions Database v6.0
 SSOL5 = SGTE Alloy Solutions Database v5.0
 SSOL4 = SGTE Alloy Solutions Database v4.9g
 SSOL2 = SGTE Alloy Solutions Database v2.1
 SSUB6 = SGTE Substances Database v6.0
 SSUB5 = SGTE Substances Database v5.2
 SSUB4 = SGTE Substances Database v4.1
 SSUB3 = SGTE Substances Database v3.3
 SSUB2 = SGTE Substances Database v2.2
 SNOB3 = SGTE Noble Metal Alloys Database v3.1
 SNOB2 = SGTE Noble Metal Alloys Database v2.1
 SNOB1 = SGTE Noble Metal Alloys Database v1.2
 STBC2 = SGTE Thermal Barrier Coating TDB v2.2
 STBC1 = SGTE Thermal Barrier Coating TDB v1.1
 SALT1 = SGTE Molten Salts Database v1.3
 SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
 SEMC2 = TC Semi-Conductors v2.1
 SLAG4 = Fe-containing Slag v4.1
 SLAG3 = Fe-containing Slag v3.2
 SLAG2 = Fe-containing Slag v2.2
 SLAG1 = Fe-containing Slag v1.2
 TCOX11 = Metal Oxide Solutions v11.0
 TCOX10 = Metal Oxide Solutions v10.1
 TCOX9 = Metal Oxide Solutions v9.0
 TCOX8 = Metal Oxide Solutions v8.0
 TCOX7 = Metal Oxide Solutions v7.0
 TCOX6 = Metal Oxide Solutions v6.0
 TCOX5 = Metal Oxide Solutions v5.1
 TCOX4 = Metal Oxide Solutions v4.1
 ION3 = Ionic Solutions v3.0
 ION2 = Ionic Solutions v2.6
 ION1 = Ionic Solutions v1.5
 NOX2 = NPL Oxide Solutions Database v2.1
 TCNOBL1 = Noble Metals Alloys v1.0
 TCNOBL2 = Noble Metals Alloys v2.0
 TCSLD4 = Solder Alloys v4.0
 TCSLD3 = Solder Alloys v3.3
 TCSLD2 = Solder Alloys v2.0
 TCSLD1 = Solder Alloys v1.1
 TCSI1 = Ultrapure Silicon v1.2
 TCM2 = Materials Processing v2.5
 TCES1 = Combustion/Sintering v1.1
 TCSC1 = Super Conductor v1.0
 TCFC1 = SOFC Database v1.0
 TCNF2 = Nuclear Fuels v2.1b
 NUMT2 = Nuclear Materials v2.1
 NUOX4 = Nuclear Oxides v4.2
 NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
 NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
 NUCL19 = IRSN NUCLEA-19
 NUCL15 = IRSN NUCLEA-15_4
 NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
 MEPH19 = IRSN Mephista-19
 MEPH15 = IRSN Mephista-15_1
 MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
 TCAQ3 = Aqueous Solution v3.0
 TCAQ2 = Aqueous Solution v2.7
 AQS2 = TGG Aqueous Solution Database v2.6
 GCE2 = TGG Geochemical/Environmental TDB v2.3
 FEDEMO = Iron Demo Database v4.0
 ALDEMO = Aluminum Demo Database v4.0
 NI DEMO = Nickel Demo Database v2.0
 CUDEMO = Copper Demo Database v1.0
 SLDEMO = Solder Demo Database v1.0
 OXDEMO = Oxide Demo Database v3.0
 SUBDEMO = Substance Demo Database v1.0
 PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
 PG35 = PG35 Binary Semi-Conductors TDB v1.3
 PURE5 = SGTE Unary (Pure Elements) TDB v5.3
 MOB2 = Alloys Mobility v2.7
 MOB1 = Alloys Mobility v1.3
 MOBFE1 = Steels/Fe-Alloys Mobility v1.1
 MOBFE2 = Steels/Fe-Alloys Mobility v2.0
 MOBFE3 = Steels/Fe-Alloys Mobility v3.0
 MOBFE4 = Steels/Fe-Alloys Mobility v4.0
 MOBFE5 = Steels/Fe-Alloys Mobility v5.0
 MOBFE6 = Steels/Fe-Alloys Mobility v6.0
 MOBN15 = Ni-Alloys Mobility v5.1
 MOBN14 = Ni-Alloys Mobility v4.1
 MOBN13 = Ni-Alloys Mobility v3.2
 MOBN12 = Ni-Alloys Mobility v2.4
 MOBN11 = Ni-Alloys Mobility v1.10
 MOBAL6 = Al-Alloys Mobility v6.0
 MOBAL5 = Al-Alloys Mobility v5.0
 MOBAL4 = Al-Alloys Mobility v4.0
 MOBAL3 = Al-Alloys Mobility v3.0
 MOBAL2 = Al-Alloys Mobility v2.0
 MOBAL1 = Al-Alloys Mobility v1.0
 MOBCU1 = Cu-Alloys Mobility v1.0
 MOBCU2 = Cu-Alloys Mobility v2.0
 MOBCU3 = Cu-Alloys Mobility v3.0
 MOBCU4 = Cu-Alloys Mobility v4.0
 MOBHEA1 = High Entropy Alloys Mobility v1.0
 MOBHEA2 = High Entropy Alloys Mobility v2.0
 MOBMG2 = Mg-Alloys Mobility v2.0
 MOBMG1 = Mg-Alloys Mobility v1.0
 MOBS11 = Si-Alloys Mobility v1.0
 MOBSDL1 = Solder-Alloys Mobility v1.1

```

MOBTI4 = Ti-Alloys Mobility v4.0
MOBTI3 = Ti-Alloys Mobility v3.1
MOBTI2 = Ti-Alloys Mobility v2.0
MOBTI1 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

DATABASE NAME /FEDEMO/: mfedemo
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys cr fe ni
CR FE NI
DEFINED
APP: rej-ph /all
BCC_A2 FCC_A1 REJECTED
APP: res-ph bcc,fcc
BCC_A2 FCC_A1 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Ni diffusion fcc Cr-Ni'
'B. Jonsson: Z. Metallkunde 86(1995)686-692; Cr, Fe and Ni diffusion fcc
'Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
'-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
'B. Jonsson: ISIJ International, 35(1995)1415-1421; Cr, Fe and Ni
diffusion bcc Cr-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'

-OK-
APP:
APP: @@ GO TO THE DICTRA MODULE TO SET UP THE SIMULATION
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ SET THE GLOBAL CONDITIONS
DIC> @@
DIC> set-cond glob T 0 1373; * N
DIC>
DIC> @@
DIC> @@ ENTER TWO REGIONS, ONE FOR EACH PHASE
DIC> @@
DIC> enter-region alpha
DIC> enter-region gamma
ATTACH TO REGION NAMED /ALPHA/:
ATTACHED TO THE RIGHT OF ALPHA /YES/:
DIC> @@
DIC> @@ ENTER THE GRID SIZE AND SPACINGS
DIC> @@
DIC> enter-grid alpha 93.45E-6 AUTO
DIC> enter-grid gamma 500.0E-6 AUTO
DIC>
DIC> @@
DIC> @@ SPECIFY WHICH PHASE GOES INTO WHICH REGION
DIC> @@
DIC> enter-phase act alpha matrix bcc
DIC> enter-phase act gamma matrix fcc
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITIONS IN THE PHASES
DIC> @@ IT IS IMPORTANT NOT TO PUT 0%NI IN PHASE BCC,
DIC> @@ ENTER SOME SMALL VALUE INSTEAD
DIC> @@
DIC> enter-composition
REGION NAME : /ALPHA/: alpha
PHASE NAME: /BCC_A2/: bcc
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-f
PROFILE FOR /CR/: cr lin .38 .38
PROFILE FOR /NI/: ni lin 1e-5 1e-5
DIC>
DIC> enter-composition
REGION NAME : /GAMMA/: gamma
PHASE NAME: /FCC_A1/: fcc
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-f
PROFILE FOR /CR/: cr lin .27 .27
PROFILE FOR /NI/: ni lin .28 .28
DIC>
DIC> @@
DIC> @@ SPECIFY THE SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 36E5
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /360000/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC>
DIC> SAVE exb5 Y
DIC>
DIC> set-inter
--OK--
DIC>
```


exb5-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb5\run.DCM.test"
DIC>
DIC>
DIC> @@ exb5_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE b5
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE SET UP FROM FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exb5
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> simulate
Region: ALPHA
single geometric dense at 0.93450E-04
0.80000 36
Region: GAMMA
single geometric dense at 0.0000
1.2500 103
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
Trying old scheme 4
GENERATING STARTING VALUES FOR CELL # 1 INTERFACE # 2
DETERMINING INITIAL EQUILIBRIUM VALUES
CALCULATING STARTING VALUES: 9 EQUILIBRIUM CALCULATIONS *** ERROR 1611 IN QTHISS: TOO MANY ITERATIONS
Give the command INFO TROUBLE for help
DONE 6 OUT OF 9 *** ERROR 1611 IN QTHISS: TOO MANY ITERATIONS
Give the command INFO TROUBLE for help
DONE 9 OUT OF 9 try 1 failed
try 2 failed
try 3 failed
DETERMINED ACTIVITIES ACR(NI) 2.00457405352E-04
UNABLE TO OBTAIN GOOD STARTING VALUE USING THE OLD SCHEME
USE NEW SCHEME /YES:
Trying new scheme
GENERATING STARTING VALUES FOR CELL # 1 INTERFACE # 2
DETERMINING INITIAL EQUILIBRIUM VALUES
CALCULATING STARTING VALUES: 18 EQUILIBRIUM CALCULATIONS DONE 1 OUT OF 18
04
U-FRACTION IN SYSTEM: CR = .305280432605602 FE = .471672082221692
NI = .223047485172706
TOTAL SIZE OF SYSTEM: 5.9345E-04 [m]
U-FRACTION IN SYSTEM: CR = .305280432605602 FE = .471672082221692
NI = .223047485172706
TOTAL SIZE OF SYSTEM: 5.9345E-04 [m]
3.503057562886359E-002 3.503838978365914E-002 3.503070722821699E-002 6.010860849728436E-004 4.996097709884086E-
004 3.843186467579116E-004 2.950169743173567E-004 2.951542157074272E-004 1.282237850461803E-
004 1.381556889258480E-007 1.911573577860582E-009 4.796670199127128E-011 2.303329573479827E-
015 9.076446564795043E-022 TIME = 0.10000000E-06 DT = 0.10000000E-06 SUM OF SQUARES = 0.90764466E-21
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.16001447 AND 0.16001447
POSITION OF INTERFACE ALPHA / GAMMA IS 0.93466001E-04
U-FRACTION IN SYSTEM: CR = .305280432536186 FE = .471672082122531
TOTAL SIZE OF SYSTEM: 5.9345E-04 [m]
26 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: GAMMA

CPU time used in timestep 0 seconds
9.795982825923318E-004 9.797942120145652E-004 9.795987026603873E-004 1.691347619213920E-012 1.336248725813137E-
012 1.296755508755182E-012 1.534999389686798E-012 6.680078113294400E-012 1.211461410126802E-
012 1.208868038883532E-012 1.209218294227699E-012 6.030069421951742E-012 1.204231833024221E-
012 1.202965670449502E-012 1.202807770706800E-012 5.949304371358198E-012 1.202017608479717E-
012 1.201382005677497E-012 1.200629566641394E-012 5.941375535485801E-012 1.199444110338236E-
012 1.197808658376458E-012 1.194411670664971E-012 5.934987258752826E-012 1.188089753613608E-
012 1.175555861730537E-012 1.150438012336459E-012

output ignored...

... output resumed

5.916111599434281E-012 5.615139793007579E-018 TIME = 2202011.3 DT = 360000.00 SUM OF SQUARES = 0.56151398E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.76087561E-11 AND -0.76087561E-11
POSITION OF INTERFACE ALPHA / GAMMA IS 0.95399428E-04
U-FRACTION IN SYSTEM: CR = .30528044496888 FE = .471672025958996
NI = .223047529072116
TOTAL SIZE OF SYSTEM: 5.9345E-04 [m]
7 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: GAMMA

CPU time used in timestep 1 seconds
1.857515243005923E-006 1.860328097861410E-006 1.852882361726696E-006 1.312188501911818E-008 2.815003807071727E-
009 2.721627042567782E-013 6.082124334976564E-022 TIME = 2562011.3 DT = 360000.00 SUM OF SQUARES = 0.60821243E-21
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.64452738E-11 AND -0.64452738E-11
POSITION OF INTERFACE ALPHA / GAMMA IS 0.93079129E-04
U-FRACTION IN SYSTEM: CR = .305280445340601 FE = .47167202530499
NI = .223047529354409
TOTAL SIZE OF SYSTEM: 5.9345E-04 [m]
6 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: GAMMA

CPU time used in timestep 1 seconds
6.450201781477898E-007 6.464378036012519E-007 6.423207643428668E-007 3.408122513600546E-009 6.667770584835788E-
010 3.267199212234195E-015 1.058076701748795E-022 TIME = 2922011.3 DT = 360000.00 SUM OF SQUARES = 0.10580767E-21
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.58181917E-11 AND -0.58181917E-11
POSITION OF INTERFACE ALPHA / GAMMA IS 0.90984580E-04
U-FRACTION IN SYSTEM: CR = .305280445518116 FE = .471672025224492
NI = .223047529257392
TOTAL SIZE OF SYSTEM: 5.9345E-04 [m]
7 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: GAMMA

CPU time used in timestep 1 seconds
3.680113606276320E-007 3.689840658726437E-007 3.659972733632945E-007 1.727065188767271E-009 3.148537793874792E-
```

```

010      1.466472825752511E-015      3.291013268494610E-
023      TIME = 3282011.3      DT = 360000.00      SUM OF SQUARES = 0.32910133E-22
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.53569964E-11 AND -0.53569964E-11
POSITION OF INTERFACE ALPHA / GAMMA IS 0.89056061E-04
U-FRACTION IN SYSTEM: CR = .305280446098058 FE = .47167202353897
NI = .223047530362973
TOTAL SIZE OF SYSTEM: 5.9345E-04 [m]
5 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: GAMMA

CPU time used in timestep           1 seconds
4.266368522437976E-009      4.268910195100429E-009      4.285225593790484E-009      1.878542591846417E-009      8.600711256692142E-
010      4.296300014532382E-016      2.000400010021238E-
024      TIME = 3600000.0      DT = 317988.70      SUM OF SQUARES = 0.20004000E-23
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.49339128E-11 AND -0.49339128E-11
POSITION OF INTERFACE ALPHA / GAMMA IS 0.87487133E-04
U-FRACTION IN SYSTEM: CR = .305280446350209 FE = .471672022910764
NI = .223047530739028
TOTAL SIZE OF SYSTEM: 5.9345E-04 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 0.93355784E-05
DELETING TIME-RECORD FOR TIME 0.27806735E-04
DELETING TIME-RECORD FOR TIME 0.64749049E-04
DELETING TIME-RECORD FOR TIME 0.13863368E-03
DELETING TIME-RECORD FOR TIME 0.28640293E-03
DELETING TIME-RECORD FOR TIME 0.58194144E-03
DELETING TIME-RECORD FOR TIME 0.11730185E-02
DELETING TIME-RECORD FOR TIME 0.23551725E-02
DELETING TIME-RECORD FOR TIME 0.47194806E-02
DELETING TIME-RECORD FOR TIME 0.94480967E-02
DELETING TIME-RECORD FOR TIME 0.18905329E-01
DELETING TIME-RECORD FOR TIME 0.37819794E-01
DELETING TIME-RECORD FOR TIME 0.75648723E-01
DELETING TIME-RECORD FOR TIME 0.15130658
DELETING TIME-RECORD FOR TIME 0.19274610
DELETING TIME-RECORD FOR TIME 0.21544345
DELETING TIME-RECORD FOR TIME 0.26083817
DELETING TIME-RECORD FOR TIME 0.35162760
DELETING TIME-RECORD FOR TIME 0.40135502
DELETING TIME-RECORD FOR TIME 0.50080986
DELETING TIME-RECORD FOR TIME 0.69971953
DELETING TIME-RECORD FOR TIME 0.91386552
DELETING TIME-RECORD FOR TIME 1.3421575
DELETING TIME-RECORD FOR TIME 2.0765292
DELETING TIME-RECORD FOR TIME 3.2890531
DELETING TIME-RECORD FOR TIME 5.5909634
DELETING TIME-RECORD FOR TIME 10.194784
DELETING TIME-RECORD FOR TIME 13.853758
DELETING TIME-RECORD FOR TIME 21.171706
DELETING TIME-RECORD FOR TIME 35.807602
DELETING TIME-RECORD FOR TIME 65.079394
DELETING TIME-RECORD FOR TIME 89.577418
DELETING TIME-RECORD FOR TIME 138.57347
DELETING TIME-RECORD FOR TIME 236.56556
DELETING TIME-RECORD FOR TIME 319.77937
DELETING TIME-RECORD FOR TIME 486.20698
DELETING TIME-RECORD FOR TIME 761.28227
DELETING TIME-RECORD FOR TIME 1232.2011
DELETING TIME-RECORD FOR TIME 2093.8574
DELETING TIME-RECORD FOR TIME 3662.1611
DELETING TIME-RECORD FOR TIME 6798.7685
DELETING TIME-RECORD FOR TIME 13071.983
DELETING TIME-RECORD FOR TIME 25618.413
DELETING TIME-RECORD FOR TIME 50711.272
DELETING TIME-RECORD FOR TIME 100896.99
DELETING TIME-RECORD FOR TIME 201268.43
DELETING TIME-RECORD FOR TIME 402011.30
DELETING TIME-RECORD FOR TIME 762011.30
DELETING TIME-RECORD FOR TIME 1122011.3
DELETING TIME-RECORD FOR TIME 1482011.3
DELETING TIME-RECORD FOR TIME 1842011.3
DELETING TIME-RECORD FOR TIME 2202011.3
DELETING TIME-RECORD FOR TIME 2562011.3
DELETING TIME-RECORD FOR TIME 2922011.3

KEEPING TIME-RECORD FOR TIME 3282011.3
AND FOR TIME 3600000.0
WORKSPACE RECLAIMED

TIMESTEP AT 3600000.00 SELECTED

DIC> set-inter
--OK--
DIC>

```

exb5-plot

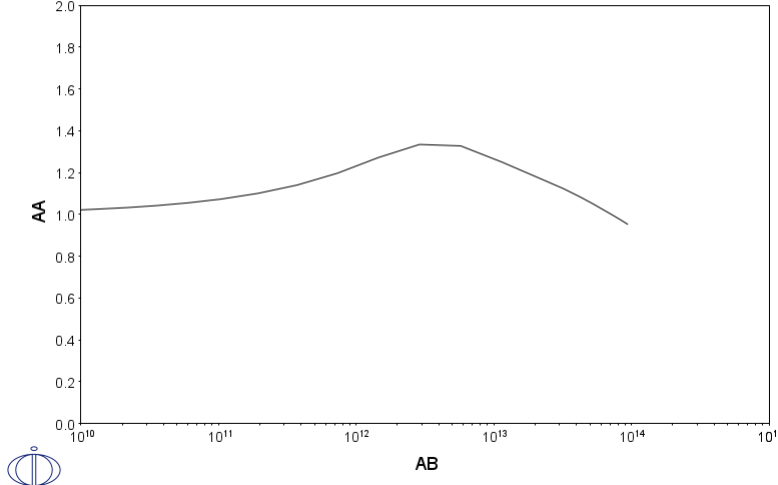
```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb5\plot.DCM.test"
DIC>
DIC>
DIC> @@ exb5_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE b5
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 3.60000E+06
DIC> read exb5
OK
DIC>
DIC> @@
DIC> @@ ENTER THE POST PROCESSOR, PLOT SOME QUANTITIES AND COMPARE WITH EXPERIMENTS
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: set-title Diffusion Couple A
POST-1:
POST-1: @@
POST-1: @@ WE ARE INTERESTED IN THE POSITION OF THE UPPER INTERFACE OF REGION ALPHA
POST-1: @@
POST-1: s-p-c interf alpha upper
POST-1:
POST-1: @@
POST-1: @@ 10 IS THE INITIAL THICKNESS USED FOR NORMALIZATION
POST-1: @@
POST-1: enter func l0=186.9e-6;
POST-1: enter func aa=2*poi(alpha,u)/l0;
POST-1: enter func ab=time/10**2;
POST-1: s-i-v time
POST-1:
POST-1: s-d-a x ab
POST-1: s-s-s x n 1e10 1e15
POST-1: s-ax-ty x log
POST-1:
POST-1: s-d-a y aa
POST-1: s-s-s y n 0 2
POST-1:
POST-1: app y exb5.exp
PROLOGUE NUMBER: /0/: 0
DATASET NUMBER(s): /-1/: 7
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Diffusion Couple A

2021.05.13.10.51.54

UPPER INTERFACE OF REGION "ALPHA#1"

CELL #1

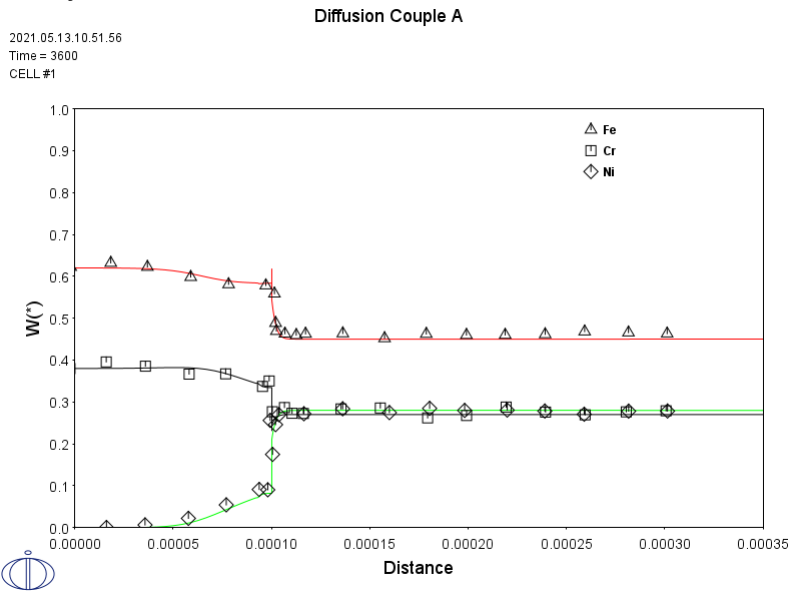


```
POST-1:
POST-1:
POST-1:
POST-1: Hit RETURN to continue
POST-1:
POST-1: @@
POST-1: @@ PLOT THE CONCENTRATION PROFILES FOR DIFFERENT ANNEALING TIMES
POST-1: @@
POST-1: s-d-a x dist glo
INFO: Distance is set as independent variable
POST-1: s-ax-ty x lin
POST-1: s-s-s x n 0 350e-6
POST-1:
POST-1: s-d-a y w(*)
POST-1: s-s-s y n 0 1
POST-1:
POST-1: s-p-c time 3600
POST-1:
```

```

POST-1: app y exb5.exp 0; 1
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

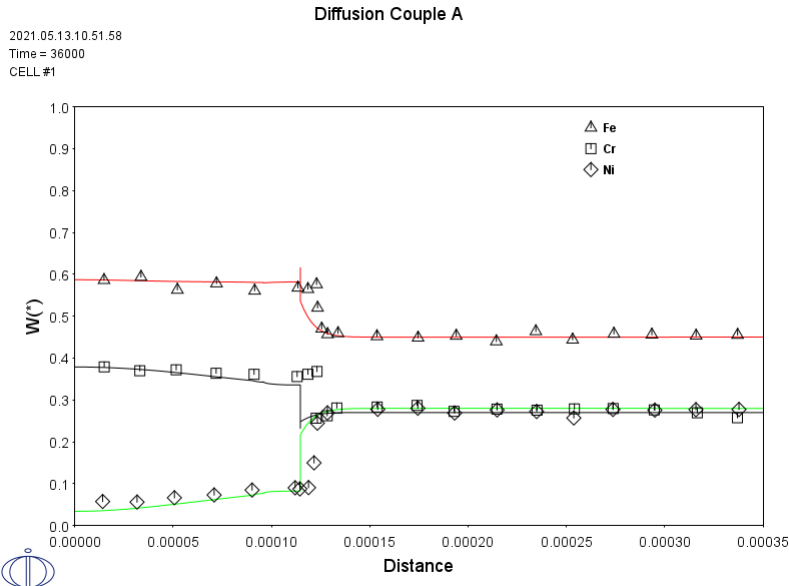
```



```

POST-1:
POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: s-p-c time 36000
POST-1: app y exb5.exp 0; 2
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

```



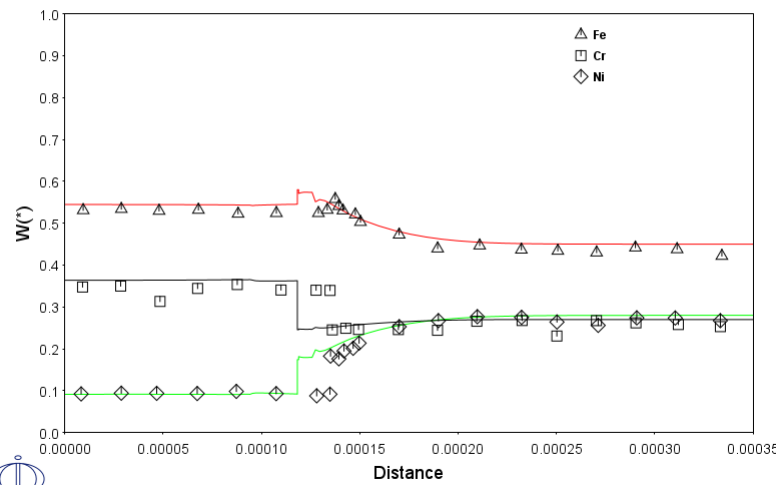
```

POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: s-p-c time 360000
POST-1: app y exb5.exp 0; 3
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

```

Diffusion Couple A

2021.05.13.10.52.00
Time = 360000
CELL #1

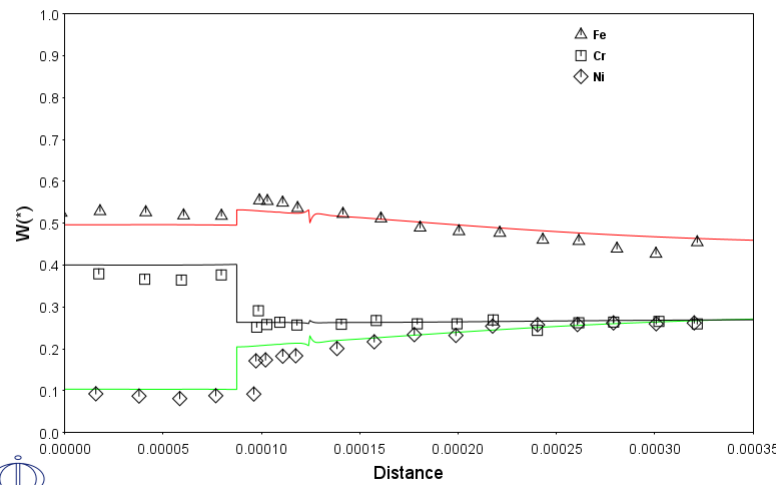


POST-1:

```
POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: s-p-c time 3600000
POST-1: app y exb5.exp 0; 4
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Diffusion Couple A

2021.05.13.10.52.01
Time = 3600000
CELL #1

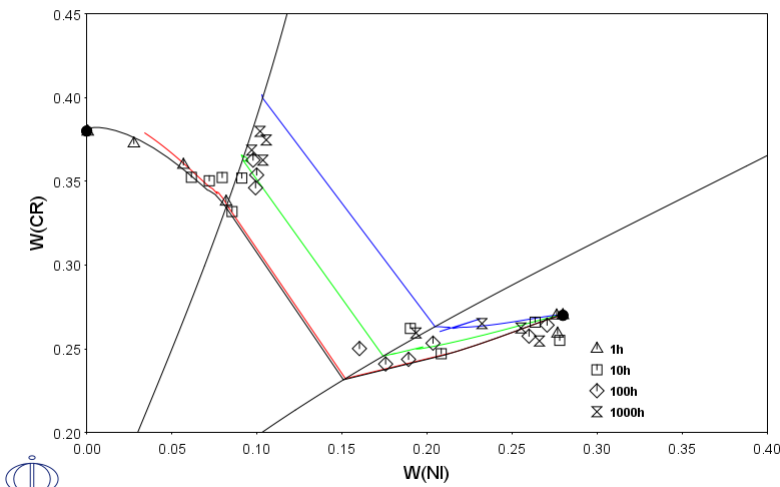


POST-1:

```
POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: @@ 
POST-1: @@ FINALLY PLOT DIFFUSION PATHS.
POST-1: @@
POST-1: s-d-a x w(ni)
POST-1: s-s-s x n .00 .40
POST-1:
POST-1: s-d-a y w(cr)
POST-1: s-s-s y n .20 .45
POST-1:
POST-1: s-i-v dist glob
POST-1:
POST-1: s-p-c time 3600,36000,360000,3600000
POST-1:
POST-1: app y exb5.exp 0; 5 6
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Diffusion Couple A

2021.05.13.10.52.03
Time = 3600,36000,360000,3600000
CELL #1



POST-1:
POST-1:
POST-1:
POST-1: set-inter
--OK---
POST-1:



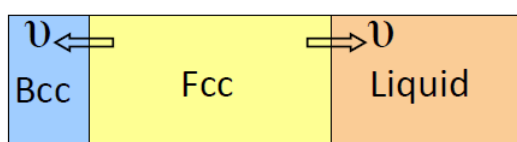
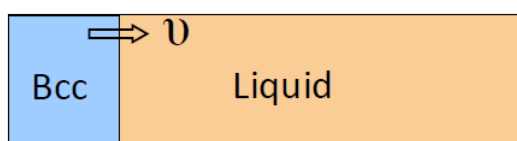
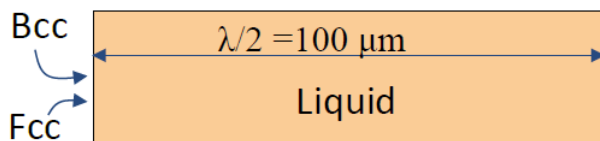
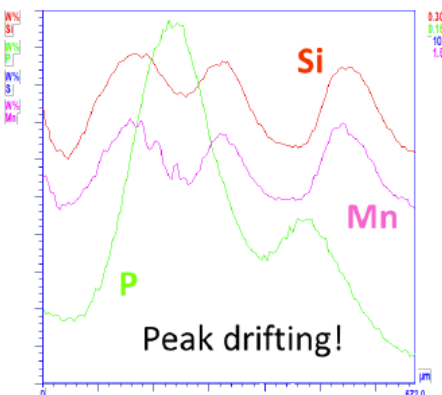
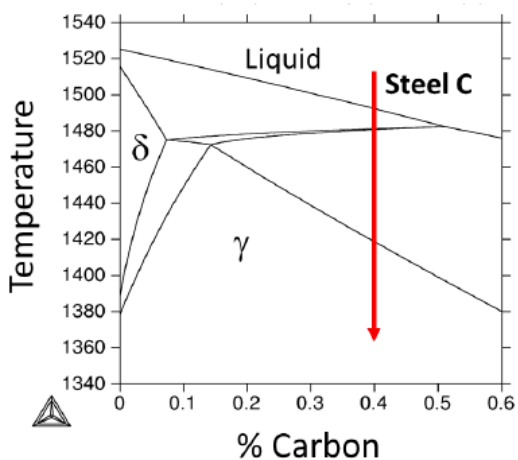
Example exb6

Micro-segregation of phosphorus

This example illustrates the effect of microsegregation of phosphorus during peritectic solidification in steel.

Steel C:

Fe - 0.8% Mn - 0.7% Si - 0.03% P - 0.4% C



exb6-setup

SYS:About

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb6\setup.DCM.test"
SYS: @@
SYS: @@ Moving boundary problem.
SYS: @@ Microsegregation of phosphorus
SYS: @@ This example illustrates the effect of microsegregation
SYS: @@ of phosphorus during peritectic solidification in steel.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:
TDB_TCFE11: @@ USE A TCFE DATABASE FOR THERMODYNAMIC DATA
TDB_TCFE11: sw tcfef9
Current database: Steels/Fe-Alloys v9.3

VA           /- DEFINED
L12_FCC      B2_BCC          DICTRA_FCC_A1
REJECTED
TDB_TCFE9:
TDB_TCFE9: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_TCFE9: def-sys fe c si mn p
FE           C               SI
MN           P   DEFINED
TDB_TCFE9:
TDB_TCFE9: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_TCFE9: rej ph /all
GAS:G         LIQUID:L        BCC_A2
FCC_A1        HCP_A3          CBCC_A12
CUB_A13       DIAMOND_FCC_A4 RED_P
WHITE_P        GRAPHITE        CEMENTITE
M23C6         M7C3            M5C2
KSI_CARBIDE   FB4N_LP1       FECN_CHI
LAVES_PHASE_C14 M3SI           MN9Si2
MN11Si19      MN6Si           G_PHASE
CR3Si         FE2Si          FESi2_H
FESi2_L        MSI             M5Si3
AL4C3         FE8Si2C        SIC
MN5SiC        CUZN_EPSILON    ALSFE4
MP_B31        CU3P_D021       M2P_C22
M3P_D0E        MN3P_D0E       FENBP
FESi4P4       SIP             SIP2
FLUORITE_C1:I ZRO2_TETR:I   M2O3C:I
M2O3H:I      REJECTED
TDB_TCFE9: res ph fcc liq bcc
FCC_A1        LIQUID:L        BCC_A2
RESTORED
TDB_TCFE9:
TDB_TCFE9: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_TCFE9: get
10:53:21,592 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
FUNCTIONS .....

List of references for assessed data

'A. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'B. Uhrenius, Int. J. Refract. Met. Hard Mater. 12 (1994) 121-127; Molar
' volumes'
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'L. Kjellqvist, Thermo-Calc Software AB, Sweden, 2012; Molar volumes'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'P. Franke, Estimated parameter within SGTE, 2007; Fe-C, Ni-C, Mo-C, C-Mn'
'B. Hallstedt, unpublished work (2016); C-Fe-Mn Epsilon martensite.'
'D. Djurovic, B. Hallstedt, J. von Appen, R. Dronskowski, CALPHAD, 34, 279
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' 1989); C-FE-MN'
'J.-H. Shim, C.-S. Oh, and D.N. Lee, Z. Metallkd., 2000, 91, p 114-120; Fe
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'J. Grobner, H.L. Lukas and F. Aldinger, CALPHAD, 20 (1996) 247-254; Si-C
' and Al-Si-C'
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'Fe-Si and Fe-Si-C'
'W. Huang, CALPHAD, 13 (1989) 243-252; TRITA-MAC 388 (rev 1989); FE-MN'
'J.-H. Shim, C.-S. Oh and D.N. Lee, J. Korean Inst. Met. Mater., 34 (1996)
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'Cr-Fe-P'
'J. Miettinen and B. Hallstedt, CALPHAD, 22 (1998) 231-256; Fe-Si and Fe
'-Si-C'
'J. Miettinen, G. Vassilev, J. Phase Equilib. Diffus., 35(5), 2014, 587-594;
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. Mn-P and Fe-Mn-P'
 'W. Zheng, J. Alloys Compd., 632 (2015) 661-675'
 'S.-M. Liang and R. Schmid-Fetzer, J. Phase Equilib., 2014, 35, 24-35'
 'D. Djurovic, B. Hallstedt, J. von Appen, R. Dronskowski, CALPHAD,
 submitted, 2011; Fe-Mn-C'
 'R. Naraghi, Thermo-Calc Software AB, Sweden, 2014; Revision of the liquid
 C-Fe-Si description'
 'W. Zheng et al., J. Iron Steel Res. Int. 24(2) (2017) 190-197'
 'A. Forsberg and J. Agren, J. Phase Equilib., 14 (1993) 354-363; Fe-Mn-Si'
 'J. Miettinen, G. Vassilev, J. Phase Equilib. Diffus., 37(5) 2016, 283-290;
 Fe-P-Si'
 -OK-

TDB_TCFE9: @@

TDB_TCFE9: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.

TDB_TCFE9: @@ SWITCH TO A MOBILITY DATABASE AND APPEND THE DATA.

TDB_TCFE9: @@

TDB_TCFE9: app

Use one of these databases

TCFE11	= Steels/Fe-Alloys v11.0
TCFE10	= Steels/Fe-Alloys v10.1
TCFE9	= Steels/Fe-Alloys v9.3
TCFE8	= Steels/Fe-Alloys v8.2
TCFE7	= Steels/Fe-Alloys v7.0
TCFE6	= Steels/Fe-Alloys v6.2
TCFE5	= Steels/Fe-Alloys v5.0
TCFE4	= Steels/Fe-Alloys v4.1
TCFE3	= Steels/Fe-Alloys v3.1
TCFE2	= Steels/Fe-Alloys v2.1
TCFE1	= Steels/Fe-Alloys v1.0
FEDAT	= TCS/TT Steels Database v1.0
TCNI11	= Ni-Alloys v11.0
TCNI10	= Ni-Alloys v10.0
TCNI9	= Ni-Alloys v9.1
TCNI8	= Ni-Alloys v8.2
TCNI7	= Ni-Alloys v7.2
TCNI6	= Ni-Alloys v6.1
TCNI5	= Ni-Alloys v5.1
TCNI4	= Ni-Alloys v4.0
TCNI1	= Ni-Alloys v1.3
TCAL8	= Al-Alloys v8.0
TCAL7	= Al-Alloys v7.1
TCAL6	= Al-Alloys v6.0
TCAL5	= Al-Alloys v5.1
TCAL4	= Al-Alloys v4.0
TCAL3	= Al-Alloys v3.0
TCAL2	= Al-Alloys v2.1
TCAL1	= Al-Alloys v1.2
TCMG6	= Mg-Alloys v6.1
TCMG5	= Mg-Alloys v5.1
TCMG4	= Mg-Alloys v4.0
TCMG3	= Mg-Alloys v3.0
TCMG2	= Mg-Alloys v2.0
TCMG1	= Mg-Alloys v1.1
TCTI3	= Ti-Alloys v3.0
TCTI2	= Ti-Alloys v2.2
TCTI1	= Ti-Alloys v1.0
TCCU4	= Cu-Alloys v4.0
TCCU3	= Cu-Alloys v3.1
TCCU2	= Cu-Alloys v2.0
TCCU1	= Cu-Alloys v1.0
TCCC1	= Cemented carbide v1.0
TCHEA5	= High Entropy Alloy v5.0
TCHEA4	= High Entropy Alloy v4.2
TCHEA3	= High Entropy Alloy v3.1
TCHEA2	= High Entropy Alloy v2.1.1
TCHEA1	= High Entropy Alloy v1.0
SSOL7	= SGTE Alloy Solutions Database v7.0
SSOL6	= SGTE Alloy Solutions Database v6.0
SSOL5	= SGTE Alloy Solutions Database v5.0
SSOL4	= SGTE Alloy Solutions Database v4.9g
SSOL2	= SGTE Alloy Solutions Database v2.1
SSUB6	= SGTE Substances Database v6.0
SSUB5	= SGTE Substances Database v5.2
SSUB4	= SGTE Substances Database v4.1
SSUB3	= SGTE Substances Database v3.3
SSUB2	= SGTE Substances Database v2.2
SNOB3	= SGTE Noble Metal Alloys Database v3.1
SNOB2	= SGTE Noble Metal Alloys Database v2.1
SNOB1	= SGTE Noble Metal Alloys Database v1.2
STBC2	= SGTE Thermal Barrier Coating TDB v2.2
STBC1	= SGTE Thermal Barrier Coating TDB v1.1
SALT1	= SGTE Molten Salts Database v1.3
SNUX6	= SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2	= TC Semi-Conductors v2.1
SLAG4	= Fe-containing Slag v4.1
SLAG3	= Fe-containing Slag v3.2
SLAG2	= Fe-containing Slag v2.2
SLAG1	= Fe-containing Slag v1.2
TCOX11	= Metal Oxide Solutions v11.0
TCOX10	= Metal Oxide Solutions v10.1
TCOX9	= Metal Oxide Solutions v9.0
TCOX8	= Metal Oxide Solutions v8.0
TCOX7	= Metal Oxide Solutions v7.0
TCOX6	= Metal Oxide Solutions v6.0
TCOX5	= Metal Oxide Solutions v5.1
TCOX4	= Metal Oxide Solutions v4.1
ION3	= Ionic Solutions v3.0
ION2	= Ionic Solutions v2.6
ION1	= Ionic Solutions v1.5
NOX2	= NPL Oxide Solutions Database v2.1
TCNOBL1	= Noble Metals Alloys v1.0
TCNOBL2	= Noble Metals Alloys v2.0
TCSLD4	= Solder Alloys v4.0
TCSLD3	= Solder Alloys v3.3
TCSLD2	= Solder Alloys v2.0
TCSLD1	= Solder Alloys v1.1
TCSI1	= Ultrapure Silicon v1.2
TCMP2	= Materials Processing v2.5
TCES1	= Combustion/Sintering v1.1
TCSC1	= Super Conductor v1.0
TCFC1	= SOFC Database v1.0
TCNF2	= Nuclear Fuels v2.1b
NUMT2	= Nuclear Materials v2.1
NUOX4	= Nuclear Oxides v4.2

```

NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephista-19
MEPH15 = IRSN Mephista-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQ82 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SLDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.2
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3
MOBFE1 = Steels/Fe-Alloys Mobility v1.1
MOBFE2 = Steels/Fe-Alloys Mobility v2.0
MOBFE3 = Steels/Fe-Alloys Mobility v3.0
MOBFE4 = Steels/Fe-Alloys Mobility v4.0
MOBFE5 = Steels/Fe-Alloys Mobility v5.0
MOBFE6 = Steels/Fe-Alloys Mobility v6.0
MOBN15 = Ni-Alloys Mobility v5.1
MOBN14 = Ni-Alloys Mobility v4.1
MOBN13 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBS11 = Si-Alloys Mobility v1.0
MOBSLD1 = Solder-Alloys Mobility v1.1
MOBT14 = Ti-Alloys Mobility v4.0
MOBT13 = Ti-Alloys Mobility v3.1
MOBT12 = Ti-Alloys Mobility v2.0
MOBT11 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

```

```

DATABASE NAME /TCFE9/: mobfe4
Current database: Steels/Fe-Alloys Mobility v4.0

```

```

VA DEFINED
B2_BCC REJECTED
APP: def-sys fe c si mn p
    FE          C           SI
    MN          P  DEFINED
APP: rej ph /all
    BCC_A2      CEMENTITE   FCC_A1
    FE4N_LP1    HCP_A3     LIQUID:L
    REJECTED
APP: res ph fcc liq bcc
    FCC_A1      LIQUID:L   BCC_A2
    RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

```

List of references for assessed data

```

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
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diffusion of Si in fcc Fe.'
'B. Jonsson: Z. Metallkunde 85(1994)498-501; C and N diffusion in bcc Cr
-Fe-Ni'
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'Assessed from data presented in Landolt-Bornstein, Vol. 26, ed. H.
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'Assessed from data presented in Landolt-Bornstein, Vol. 26, ed. H.
Mehrer, Springer (1990); Impurity diff of P in bcc Fe.'
'No assessed or estimated parameters'
'Estimation by using the modified Sutherland equation from Chen et al.,
Phil. Magazine 94 (2014) 1552.'
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diffusion of Cr in fcc Cu.'
'Y. Tang et al., CALPHAD, 49(2015)58.'

```

-OK-

```

APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC:  @@ ENTER THE GLOBAL CONDITION T
DIC: @@
DIC:  @@ LOWER THE TEMPERATURE TO A RATE OF 0.2 K/s
DIC: @@
DIC: set-cond glob T 0 1780-0.2*TIME; * N
DIC:>
DIC:> @@
DIC:  @@ ENTER A REGION CALLED smalta
DIC: @@
DIC: enter-region smalta
DIC:>
DIC:> @@
DIC:  @@ ENTER A DOUBLE GEOMETRIC GRID INTO THE REGION
DIC: @@
DIC: enter-grid
REGION NAME : /SMALTA/: smalta
WIDTH OF REGION /1/: 1e-4
TYPE /LINEAR/: AUTO
DIC:>
DIC:> @@
DIC:  @@ ENTER active PHASES INTO THE REGION
DIC: @@
DIC: enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /SMALTA/: smalta
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: liq
DIC:>
DIC:> @@
DIC:  @@ ENTER inactive PHASES INTO THE REGION, BOTH PHASES ON THE SAME SIDE
DIC: @@ OF THE LIQUID REGION TO GET A PERITECTIC REACTION.
DIC: @@
DIC: enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /SMALTA/: smalta
ATTACHED TO THE RIGHT OF SMALTA /YES/: no
PHASE NAME: /NONE/: fcc#1
DEPENDENT COMPONENT ? /SI/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-5
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC>
DIC: enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: inact
ATTACH TO REGION NAMED /SMALTA/: smalta
ATTACHED TO THE RIGHT OF SMALTA /YES/: no
PHASE NAME: /NONE/: bcc#1
DEPENDENT COMPONENT ? /SI/: fe
REQUIRED DRIVING FORCE FOR PRECIPITATION: /1E-05/: 1e-5
CONDITION TYPE /CLOSED_SYSTEM/: closed
DIC:>
DIC:> @@
DIC:  @@ ENTER A START COMPOSITION FOR THE LIQUID
DIC: @@
DIC: enter-composition
REGION NAME : /SMALTA/: smalta
PHASE NAME: /LIQUID/: liq
DEPENDENT COMPONENT ? /SI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: c lin 0.4 0.4
PROFILE FOR /MN/: si lin 0.7 0.7
PROFILE FOR /P/: mn lin 0.8 0.8
PROFILE FOR /SI/: p lin 0.03 0.03
DIC:>
DIC:> @@
DIC:  @@ THE BOUNDARY CONDITION IS A CLOSED SYSTEM (DEFAULT) AS WE DO NOT SPECIFY
DIC:  @@ ANYTHING ELSE
DIC:>
DIC:> @@
DIC:  @@ SET THE SIMULATION TIME
DIC:>
DIC: set-simulation-time
END TIME FOR INTEGRATION /.1/: 3000
AUTOMATIC TIMESTEP CONTROL /YES/: yes
MAX TIMESTEP DURING INTEGRATION /300/: 15
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC:>
DIC:>
DIC:>
DIC:>
DIC:  @@ CHECK THE INTERFACE POSITION. THIS IS TO MAKE SURE THAT THE LIQUID REGION
DIC:  @@ DOES NOT SHRINK TOO MUCH DURING A TIMESTEP. IN ADDITION THE TIMESTEP IS
DIC:  @@ CONTROLLED BY THE PHASE INTERFACE DISPLACEMENT DURING THE SIMULATION.
DIC:>
DIC: s-s-c
NS01A PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA Timesteps IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/: yes
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/:
DEGREE OF IMPLICITY WHEN INTEGRATING PDEs (AUTO, 0 -> 0.5 -> 1) : /AUTO/:
MAX TIMESTEP CHANGE PER TIMESTEP : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/:
DIC:>
DIC:> @@
DIC:  @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC:>

```

```
DIC> save exb6 Y
DIC>
DIC> set-inter
--OK---
DIC>
```

exb6-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb6\run.DCM.test"
DIC>
DIC>
DIC> @@ exb6_run.DCM
DIC>
DIC> @@
DIC> @@ READ THE SET UP FROM FILE AND START THE SIMULATION
DIC> @@
DIC>
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exb6
OK
DIC> sim
Region: SMALTA
single geometric dense at 0.0000
1.0649 83
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 0.22705714E-05 DT = 0.21705714E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.66117143E-05 DT = 0.43411429E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 0.1529400E-04 DT = 0.86822858E-05 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.32658572E-04 DT = 0.17364572E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.67387715E-04 DT = 0.34729143E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237327E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.13684600E-03 DT = 0.69458286E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.27576257E-03 DT = 0.13891657E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 0.55359572E-03 DT = 0.27783314E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669964 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.11092620E-02 DT = 0.55566629E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.22205946E-02 DT = 0.11113326E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669964 P = 5.39133528237327E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.44432597E-02 DT = 0.22226652E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669964 P = 5.39133528237327E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.88885901E-02 DT = 0.44453303E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669964 P = 5.39133528237327E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.17779251E-01 DT = 0.88906606E-02 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237327E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
```

```

TIME = 0.35560572E-01 DT = 0.17781321E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .018537587104735 FE = .977481861209078
MN = .00810568126669962 P = 5.39133528237327E-04
SI = .0138733239959838
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.71123215E-01 DT = 0.35562643E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .018537587104735 FE = .977481861209078
MN = .00810568126669963 P = 5.39133528237328E-04
SI = .0138733239959839
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 1 seconds
TIME = 0.14224850 DT = 0.71125285E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047351 FE = .977481861209078
MN = .00810568126669965 P = 5.39133528237335E-04
SI = .0138733239959839
TOTAL SIZE OF SYSTEM: 1E-04 [m]
CPU time used in timestep 0 seconds
TIME = 0.28449907 DT = 0.14225057 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0185375871047352 FE = .977481861209078
MN = .0081056812666997 P = 5.39133528237319E-04

```

output ignored...

... output resumed

```

DELETING TIME-RECORD FOR TIME 925.22043
DELETING TIME-RECORD FOR TIME 940.22043
DELETING TIME-RECORD FOR TIME 955.22043
DELETING TIME-RECORD FOR TIME 970.22043
DELETING TIME-RECORD FOR TIME 985.22043
DELETING TIME-RECORD FOR TIME 1000.2204
DELETING TIME-RECORD FOR TIME 1015.2204
DELETING TIME-RECORD FOR TIME 1030.2204
DELETING TIME-RECORD FOR TIME 1045.2204
DELETING TIME-RECORD FOR TIME 1060.2204
DELETING TIME-RECORD FOR TIME 1075.2204
DELETING TIME-RECORD FOR TIME 1090.2204
DELETING TIME-RECORD FOR TIME 1105.2204
DELETING TIME-RECORD FOR TIME 1120.2204
DELETING TIME-RECORD FOR TIME 1135.2204
DELETING TIME-RECORD FOR TIME 1150.2204
DELETING TIME-RECORD FOR TIME 1165.2204
DELETING TIME-RECORD FOR TIME 1180.2204
DELETING TIME-RECORD FOR TIME 1195.2204
DELETING TIME-RECORD FOR TIME 1210.2204
DELETING TIME-RECORD FOR TIME 1225.2204
DELETING TIME-RECORD FOR TIME 1240.2204
DELETING TIME-RECORD FOR TIME 1255.2204
DELETING TIME-RECORD FOR TIME 1270.2204
DELETING TIME-RECORD FOR TIME 1285.2204
DELETING TIME-RECORD FOR TIME 1300.2204
DELETING TIME-RECORD FOR TIME 1315.2204
DELETING TIME-RECORD FOR TIME 1330.2204
DELETING TIME-RECORD FOR TIME 1345.2204
DELETING TIME-RECORD FOR TIME 1360.2204
DELETING TIME-RECORD FOR TIME 1375.2204
DELETING TIME-RECORD FOR TIME 1390.2204
DELETING TIME-RECORD FOR TIME 1405.2204
DELETING TIME-RECORD FOR TIME 1420.2204
DELETING TIME-RECORD FOR TIME 1435.2204
DELETING TIME-RECORD FOR TIME 1450.2204
DELETING TIME-RECORD FOR TIME 1465.2204
DELETING TIME-RECORD FOR TIME 1480.2204
DELETING TIME-RECORD FOR TIME 1495.2204
DELETING TIME-RECORD FOR TIME 1510.2204
DELETING TIME-RECORD FOR TIME 1525.2204
DELETING TIME-RECORD FOR TIME 1540.2204
DELETING TIME-RECORD FOR TIME 1555.2204
DELETING TIME-RECORD FOR TIME 1570.2204
DELETING TIME-RECORD FOR TIME 1585.2204
DELETING TIME-RECORD FOR TIME 1600.2204
DELETING TIME-RECORD FOR TIME 1615.2204
DELETING TIME-RECORD FOR TIME 1630.2204
DELETING TIME-RECORD FOR TIME 1645.2204
DELETING TIME-RECORD FOR TIME 1660.2204
DELETING TIME-RECORD FOR TIME 1675.2204
DELETING TIME-RECORD FOR TIME 1690.2204
DELETING TIME-RECORD FOR TIME 1705.2204
DELETING TIME-RECORD FOR TIME 1720.2204
DELETING TIME-RECORD FOR TIME 1735.2204
DELETING TIME-RECORD FOR TIME 1750.2204
DELETING TIME-RECORD FOR TIME 1765.2204
DELETING TIME-RECORD FOR TIME 1780.2204
DELETING TIME-RECORD FOR TIME 1795.2204
DELETING TIME-RECORD FOR TIME 1810.2204
DELETING TIME-RECORD FOR TIME 1825.2204
DELETING TIME-RECORD FOR TIME 1840.2204
DELETING TIME-RECORD FOR TIME 1855.2204
DELETING TIME-RECORD FOR TIME 1870.2204
DELETING TIME-RECORD FOR TIME 1885.2204
DELETING TIME-RECORD FOR TIME 1900.2204
DELETING TIME-RECORD FOR TIME 1915.2204
DELETING TIME-RECORD FOR TIME 1930.2204
DELETING TIME-RECORD FOR TIME 1945.2204
DELETING TIME-RECORD FOR TIME 1960.2204
DELETING TIME-RECORD FOR TIME 1975.2204
DELETING TIME-RECORD FOR TIME 1990.2204
DELETING TIME-RECORD FOR TIME 2005.2204
DELETING TIME-RECORD FOR TIME 2020.2204
DELETING TIME-RECORD FOR TIME 2035.2204
DELETING TIME-RECORD FOR TIME 2050.2204
DELETING TIME-RECORD FOR TIME 2065.2204
DELETING TIME-RECORD FOR TIME 2080.2204
DELETING TIME-RECORD FOR TIME 2095.2204
DELETING TIME-RECORD FOR TIME 2110.2204
DELETING TIME-RECORD FOR TIME 2125.2204
DELETING TIME-RECORD FOR TIME 2140.2204
DELETING TIME-RECORD FOR TIME 2155.2204
DELETING TIME-RECORD FOR TIME 2170.2204
DELETING TIME-RECORD FOR TIME 2185.2204
DELETING TIME-RECORD FOR TIME 2200.2204
DELETING TIME-RECORD FOR TIME 2215.2204

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DELETING TIME-RECORD FOR TIME 2230.2204
DELETING TIME-RECORD FOR TIME 2245.2204
DELETING TIME-RECORD FOR TIME 2260.2204
DELETING TIME-RECORD FOR TIME 2275.2204
DELETING TIME-RECORD FOR TIME 2290.2204
DELETING TIME-RECORD FOR TIME 2305.2204
DELETING TIME-RECORD FOR TIME 2320.2204
DELETING TIME-RECORD FOR TIME 2335.2204
DELETING TIME-RECORD FOR TIME 2350.2204
DELETING TIME-RECORD FOR TIME 2365.2204
DELETING TIME-RECORD FOR TIME 2380.2204
DELETING TIME-RECORD FOR TIME 2395.2204
DELETING TIME-RECORD FOR TIME 2410.2204
DELETING TIME-RECORD FOR TIME 2425.2204
DELETING TIME-RECORD FOR TIME 2440.2204
DELETING TIME-RECORD FOR TIME 2455.2204
DELETING TIME-RECORD FOR TIME 2470.2204
DELETING TIME-RECORD FOR TIME 2485.2204
DELETING TIME-RECORD FOR TIME 2500.2204
DELETING TIME-RECORD FOR TIME 2515.2204
DELETING TIME-RECORD FOR TIME 2530.2204
DELETING TIME-RECORD FOR TIME 2545.2204
DELETING TIME-RECORD FOR TIME 2560.2204
DELETING TIME-RECORD FOR TIME 2575.2204
DELETING TIME-RECORD FOR TIME 2590.2204
DELETING TIME-RECORD FOR TIME 2605.2204
DELETING TIME-RECORD FOR TIME 2620.2204
DELETING TIME-RECORD FOR TIME 2635.2204
DELETING TIME-RECORD FOR TIME 2650.2204
DELETING TIME-RECORD FOR TIME 2665.2204
DELETING TIME-RECORD FOR TIME 2680.2204
DELETING TIME-RECORD FOR TIME 2695.2204
DELETING TIME-RECORD FOR TIME 2710.2204
DELETING TIME-RECORD FOR TIME 2725.2204
DELETING TIME-RECORD FOR TIME 2740.2204
DELETING TIME-RECORD FOR TIME 2755.2204
DELETING TIME-RECORD FOR TIME 2770.2204
DELETING TIME-RECORD FOR TIME 2785.2204
DELETING TIME-RECORD FOR TIME 2800.2204
DELETING TIME-RECORD FOR TIME 2815.2204
DELETING TIME-RECORD FOR TIME 2830.2204
DELETING TIME-RECORD FOR TIME 2845.2204
DELETING TIME-RECORD FOR TIME 2860.2204
DELETING TIME-RECORD FOR TIME 2875.2204
DELETING TIME-RECORD FOR TIME 2890.2204
DELETING TIME-RECORD FOR TIME 2905.2204
DELETING TIME-RECORD FOR TIME 2920.2204
DELETING TIME-RECORD FOR TIME 2935.2204
DELETING TIME-RECORD FOR TIME 2950.2204
DELETING TIME-RECORD FOR TIME 2965.2204
DELETING TIME-RECORD FOR TIME 2980.2204

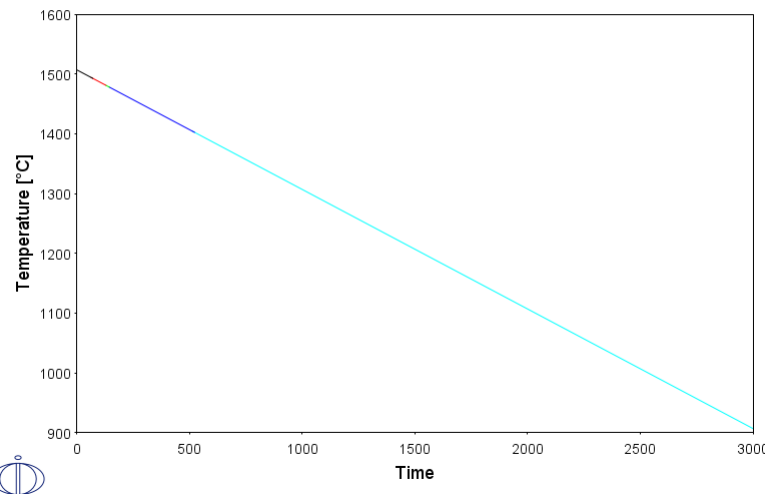
KEEPING TIME-RECORD FOR TIME 2995.2204
AND FOR TIME 3000.0000
WORKSPACE RECLAIMED

TIMESTEP AT 3000.00000 SELECTED

DIC>
DIC> set-inter
---OK---
DIC>

exb6-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb6\plot.DCM.test"
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 3.00000E+03
DIC> read exb6
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: set-title Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P
POST-1:
POST-1: s-d-a y t-c
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-p-c interf first
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...orking ...          OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P
2021.05.13.11.30.52
"FIRST" INTERFACE OF SYSTEM
CELL #1
```



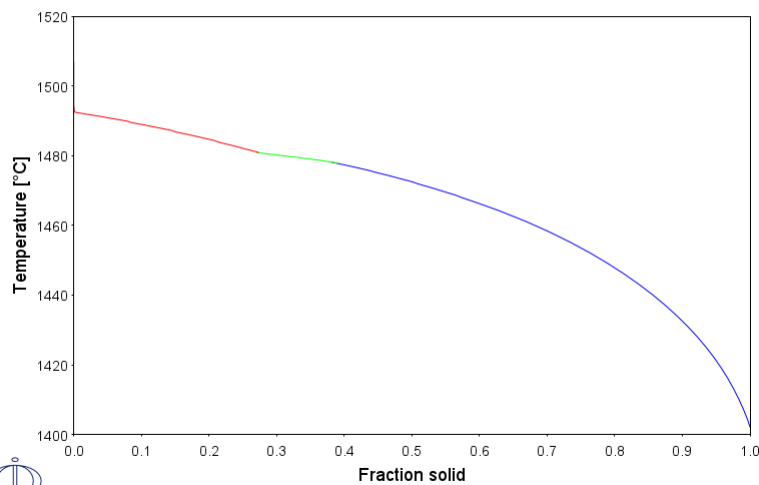
```
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: @@
POST-1: @@ PLOT THE FRACTION OF SOLID
POST-1: @@
POST-1: enter func fs=1-ivv(liq);
POST-1: s-d-a x fs
POST-1: s-s-s x n 0 1
POST-1: s-ax-te x n Fraction solid
POST-1:
POST-1: s-d-a y t-c
POST-1:
POST-1:
POST-1: s-p-c interf smalta lower
POST-1:
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...orking ...          OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

2021.05.13.11.30.55

LOWER INTERFACE OF REGION "SMALTA#1"

CELL #1

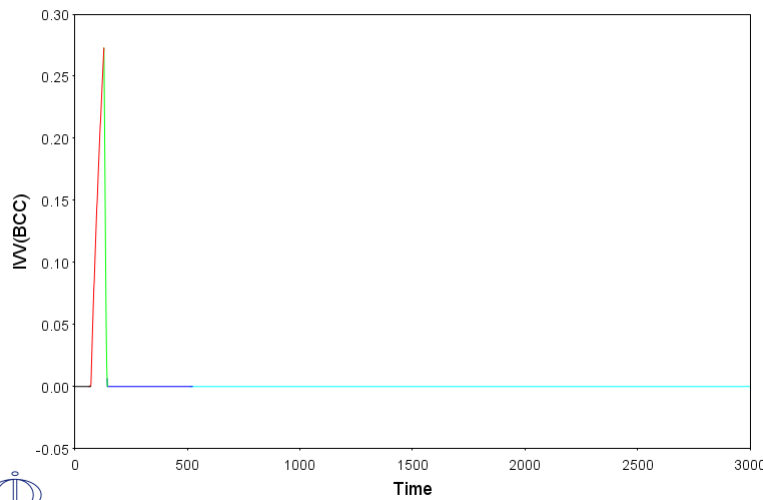


```
POST-1:  
POST-1:Hit RETURN to continue  
POST-1: s-d-a y ivv(bcc)  
POST-1: s-d-a x time  
INFO: Time is set as independent variable  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ...orking ... OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

2021.05.13.11.31.00

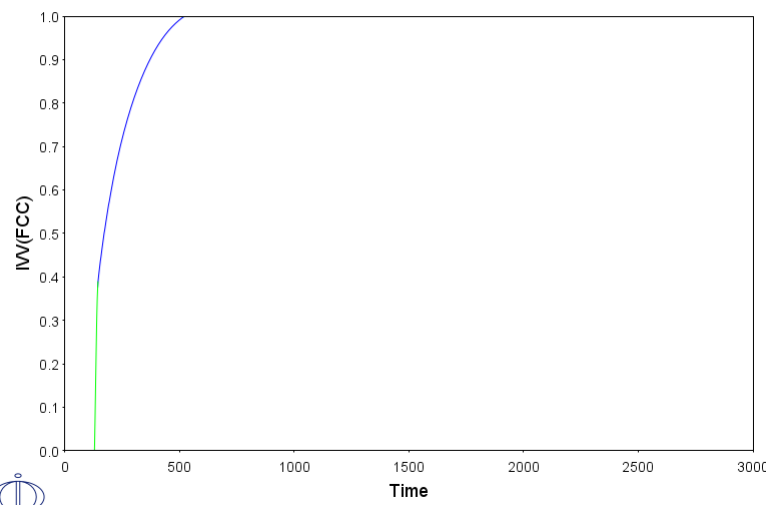
CELL #1



```
POST-1:  
POST-1:Hit RETURN to continue  
POST-1:  
POST-1: s-d-a y ivv(fcc)  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ...orking ... OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

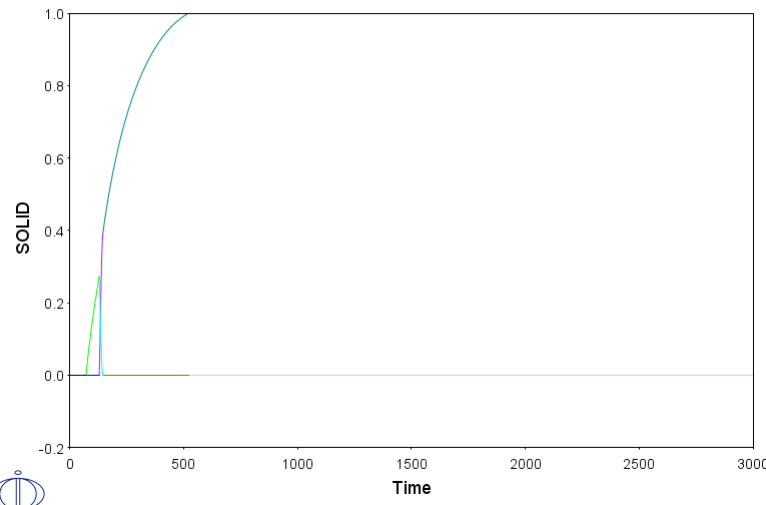
2021.05.13.11.31.05
CELL #1



```
POST-1:  
POST-1:Hit RETURN to continue  
POST-1: ent table solid  
Variable(s) ivv(bcc) ivv(fcc)  
POST-1:  
POST-1: s-d-a y solid  
COLUMN NUMBER /*/  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ...orking ... OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

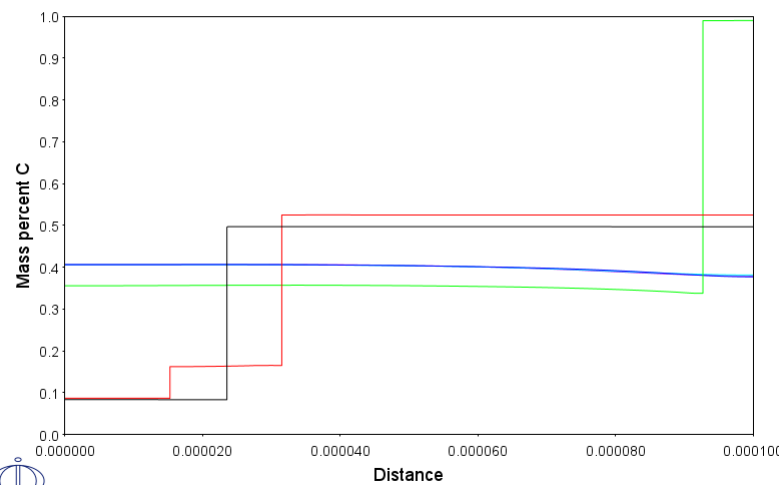
2021.05.13.11.31.12
CELL #1



```
POST-1:  
POST-1:Hit RETURN to continue  
POST-1: s-d-a y w-p c  
POST-1: s-d-a x dis gl  
INFO: Distance is set as independent variable  
POST-1: s-p-c time 120,135,400,700,1500,3000  
POST-1:  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ...orking ...OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

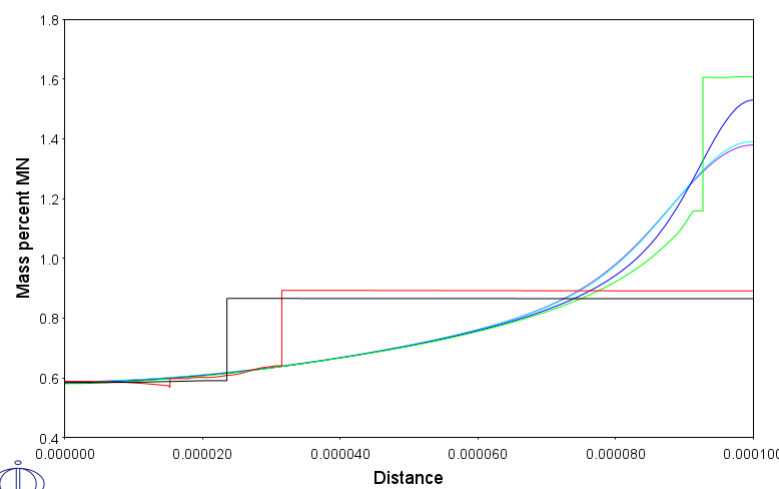
2021.05.13.11.31.15
Time = 120,135,400,700,1500,3000
CELL #1



```
POST-1:  
POST-1: Hit RETURN to continue  
POST-1: s-d-a y w-p mn  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ...orking ...OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

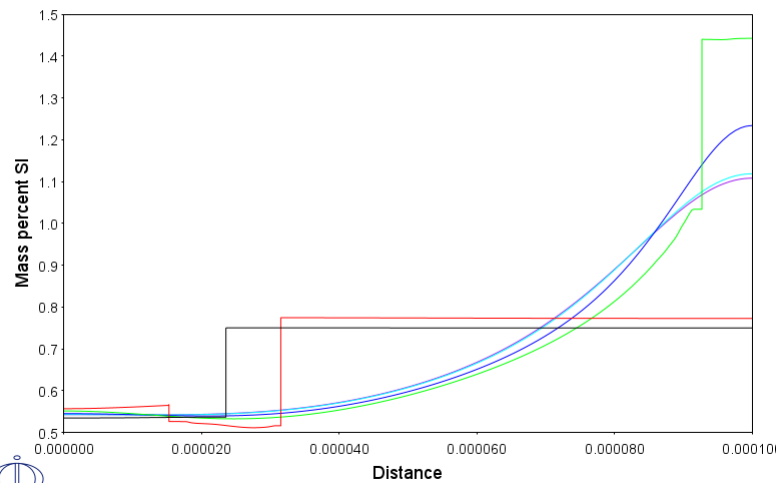
2021.05.13.11.31.18
Time = 120,135,400,700,1500,3000
CELL #1



```
POST-1:  
POST-1: Hit RETURN to continue  
POST-1: s-d-a y w-p si  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ...orking ...OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

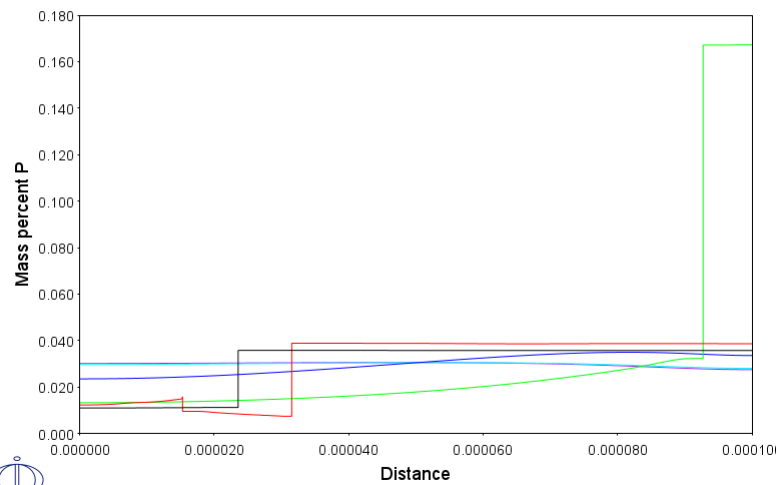
2021.05.13.11.31.20
Time = 120,135,400,700,1500,3000
CELL #1



```
POST-1:  
POST-1: Hit RETURN to continue  
POST-1: s-d-a y w-p p  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ...orking ...OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

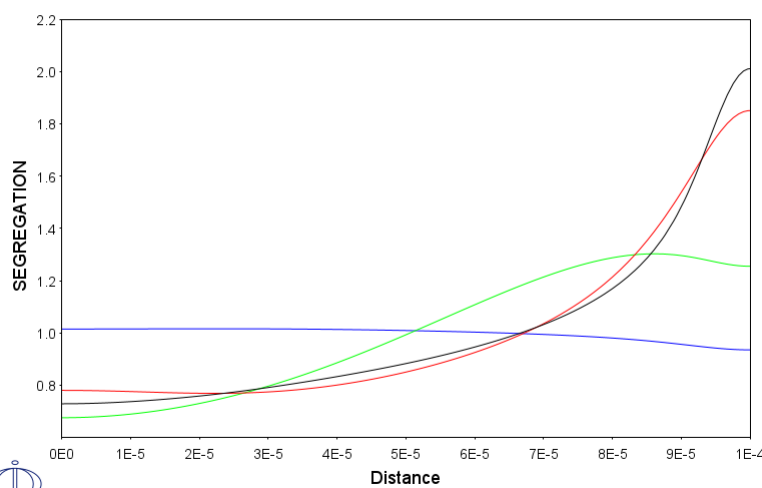
2021.05.13.11.31.22
Time = 120,135,400,700,1500,3000
CELL #1



```
POST-1:  
POST-1: Hit RETURN to continue  
POST-1:  
POST-1: ent function mnn  
FUNCTION: w(mn)/0.008  
&  
POST-1: ent function sin  
FUNCTION: w(si)/0.007  
&  
POST-1: ent function pn  
FUNCTION: w(p)/0.0003  
&  
POST-1: ent function cn  
FUNCTION: w(c)/0.004  
&  
POST-1: ent tabel segregation  
Variable(s) mnn sin pn cn  
POST-1:  
POST-1:  
POST-1: s-d-a y segregation  
COLUMN NUMBER /*:  
POST-1:  
POST-1:  
POST-1: s-p-c time 610  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
POST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

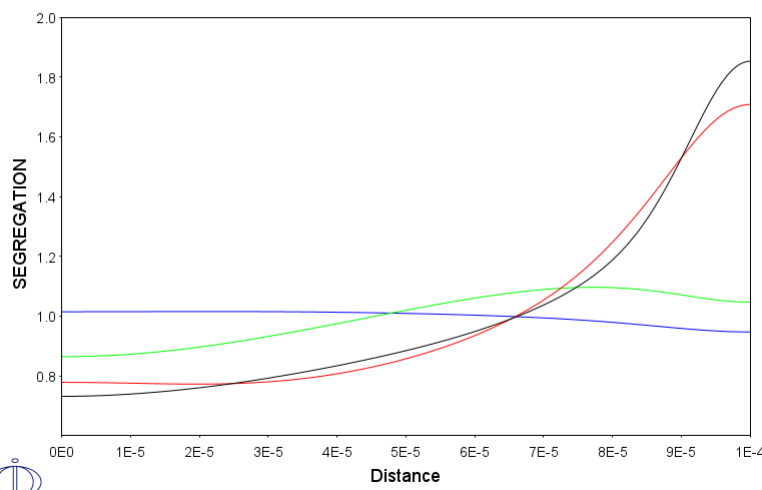
2021.05.13.11.31.25
Time = 610
CELL #1



POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: s-p-c time 800
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

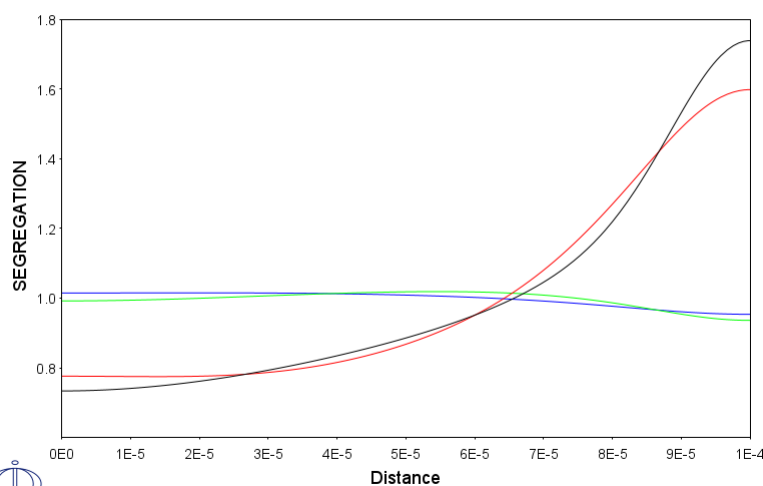
2021.05.13.11.31.27
Time = 800
CELL #1



POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: s-p-c time 1500
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

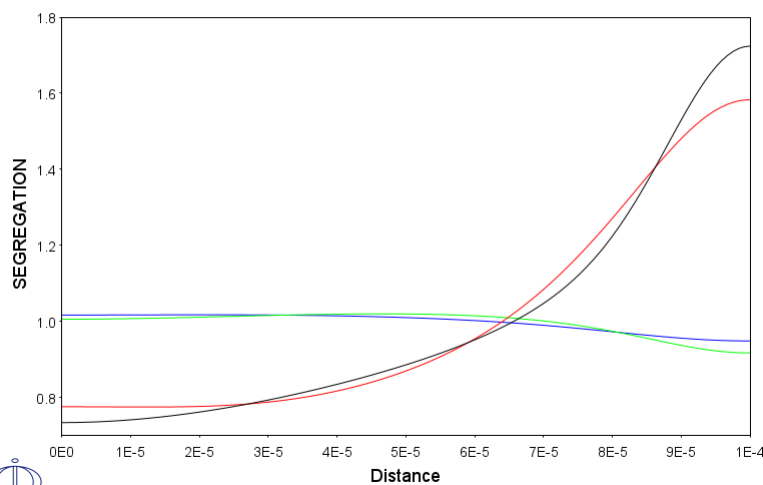
2021.05.13.11.31.33
Time = 1500
CELL #1



```
POST-1:  
POST-1:Hit RETURN to continue  
POST-1:  
POST-1: s-p-c time 3000  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
POST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

Fe-0.4%C-0.7%Si-0.8%Mn-0.03%P

2021.05.13.11.31.44
Time = 3000
CELL #1



```
POST-1:  
POST-1:Hit RETURN to continue  
POST-1: set-inter  
--OK--  
POST-1:
```

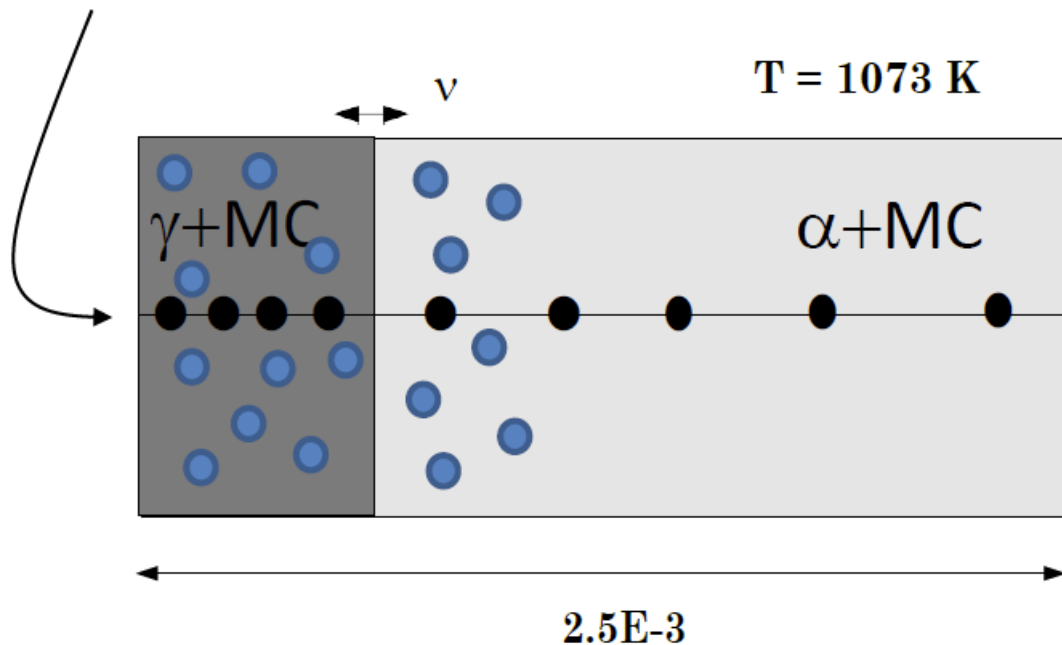


Example exb7

Moving boundary problem with multiple phases on each side of the boundary

This example shows how to enter dispersed phases on either side of a phase interface. The particular case shows how the kinetics of a ferrite to austenite transformation is affected by simultaneous precipitation of niobium carbide. The transformation is caused by carburization.

$$\text{ACR}(\text{C}) = 1$$



exb7-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb7\setup.DCM"
SYS: @@ This example modifies the database interactively, which is not
SYS: @@ yet supported by GES6. Therefore, we enforce the use of GES5.
SYS: set-ges-version 5
SYS:
SYS: @@
SYS: @@ Moving boundary problem.
SYS: @@ Moving boundary problem with multiple phases on each side
SYS: @@ This example shows how to enter dispersed phases on either side
SYS: @@ of a phase interface. The particular case shows how
SYS: @@ the kinetics of a ferrite to austenite transformation is
SYS: @@ affected by simultaneous precipitation of niobium carbide.
SYS: @@ The transformation is caused by carburization.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ USE A TCFE DATABASE FOR THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw tcfe9
Current database: Steels/Fe-Alloys v9.3

VA /- DEFINED
L12_FCC B2_BCC DICTRA_FCC_A1
REJECTED
TDB_TCFE9:
TDB_TCFE9: @@
TDB_TCFE9: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_TCFE9: @@
TDB_TCFE9: def-species fe c nb
 FE C NB
 DEFINED
TDB_TCFE9:
TDB_TCFE9: @@
TDB_TCFE9: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_TCFE9: @@
TDB_TCFE9: rej ph * all
GAS:G LIQUID:L BCC_A2
FCC_A1 HCP_A3 CBCC_A12
CUB_A13 DIAMOND_FCC_A4 GRAPHITE
CEMENTITE M23C6 M7C3
M6C M5C2 KS1_CARBIDE
Z_PHASE FE4N_LP1 FECN_CHI
SIGMA MU_PHASE LAVES_PHASE_C14
G_PHASE CR3SI NBNI3
AL5FE4 REJECTED
TDB_TCFE9: res ph fcc bcc grap
 FCC_A1 BCC_A2 GRAPHITE
RESTORED
TDB_TCFE9:
TDB_TCFE9: @@
TDB_TCFE9: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_TCFE9: @@
TDB_TCFE9: get
REINITIATING GES
ELEMENTS
SPECIES
PHASES
Creating a new composition set FCC_A1#2
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'A. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'B. Uhrenius, Int. J. Refract. Met. Hard Mater. 12 (1994) 121-127; Molar volumes'
'X.-G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar volumes'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'P. Franke, Estimated parameter within SGTE, 2007; Fe-C, Ni-C, Mo-C, C-Mn'
'P. Gustafsson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.-G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'B. Hallstedt, unpublished work (2016); C-Fe-Mn Epsilon martensite.'
'A.V. Khvan, B. Hallstedt, CALPHAD, 40, 10-15(2013); Fe-Nb, Nb-C, Nb-N, Fe -Mn-Nb, Fe-Nb-C, Fe-Nb-N, Nb-C-N, Fe-Mn-Nb-C, Fe-Mn-Nb-N'
'B.-J. Lee, Metall. Mater. Trans. A, 32A (2001) 2423-39'
'S. Canderyd, Report IM-2005-109, Stockholm, Sweden; Fe-Nb-C'
'W. Huang, Z. Metallkd., 6 (1990) 397-404; Fe-Nb-C'
-OK-
TDB_TCFE9: @@
TDB_TCFE9: @@ NOW APPEND A SSUB DATABASE FROM WHICH WE READ THE THERMODYNAMIC
TDB_TCFE9: @@ DESCRIPTION OF NIOBIUM CARBIDE
TDB_TCFE9: @@

```

TDB_TCFE9:
TDB_TCFE9: app SSUB6
Current database: SGTE Substances Database v6.0

VA DEFINED
APP: def-sys fe c nb
FE C NB
    DEFINED
APP: rej ph *
GAS:G C0_749NB1_S C0_877NB1_S
CO_98NB1_S C_S C_L
DIAMOND CIFE3_S C1NB1_S
C1NB2_S C60_S FE_S
FE_S2 FE_S3 FE_L
FE2NB1_S NB_S NB_L
REJECTED

APP: rest ph c1nb1_s
C1NB1_S RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

C1NB1 I. BARIN 3rd. Edition
C1NB1
Data taken from BARIN 3rd. Ed. (1995)
-OK-
APP:
APP:
APP: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
APP: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE THE DATA
APP: @@
APP: app mobfe4
Current database: Steels/Fe-Alloys Mobility v4.0

VA DEFINED
B2_BCC REJECTED
APP: def-sys fe c nb
FE C NB
    DEFINED
APP: rej ph * all
BCC_A2 CEMENTITE FCC_A1
FE4N_LP1 HCP_A3 LIQUID:L
REJECTED

APP: res ph fcc bcc
FCC_A1 BCC_A2 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'J. Geise and Ch. Herzig, Z. Metallkd. 76(1985)622.; Impurity diffusion of
Nb in fcc Fe.'
'Assessed from data presented by B. B. Yu and R. F. Davis J. Phys. Chem.
Solids 40(1979)997.; C Self-Diff in NbC.'
'B. Jonsson: Z. Metallkunde 85(1994)498-501; C and N diffusion in bcc Cr
-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'
'R. F. Peart, Acta Metall. 10(1962)519.; Impurity diffusion of Fe in bcc
Nb.'
'Assessed from data presented in Landolt-Bornstein, Vol. 26, ed. H.
Mehrer, Springer (1990); Impurity diff of Nb in bcc Fe.'
'R. E. Einziger et al., Phys. Rev. B 17(1978)440.; self-diffusion of Nb in
bcc Nb.'

-OK-
APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: go d-m
NO TIME STEP DEFINED
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
*** ENTERING C1NB1_S AS A DIFFUSION NONE PHASE
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 1073.15; * N
DIC>
DIC> @@
DIC> @@ ENTER REGIONS ferr AND aus
DIC> @@
DIC> enter-region
REGION NAME : ferr
DIC>
DIC> ent-reg
REGION NAME : aus
ATTACH TO REGION NAMED /FERR/: ferr
ATTACHED TO THE RIGHT OF FERR /YES/: n
DIC>
DIC> @@
DIC> @@ ENTER GRIDS INTO REGIONS
DIC> @@
DIC> enter-grid
REGION NAME : /AUS/: ferr
WIDTH OF REGION /1/: 2.499999e-3
TYPE /LINEAR/: AUTO
DIC>
DIC>

```

```

DIC> enter-grid
REGION NAME : /AUS/: aus
WIDTH OF REGION /1/: 1e-9
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER active PHASES INTO REGIONS
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /AUS/: ferr
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: bcc
DIC>
DIC>
DIC> en-ph
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUS/: ferr
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC>
DIC> en-ph
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUS/: aus
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: c1nb1_s
INFO: EQUILIBRIUM COMPOSITION AND FRACTION OF SPHEROID PHASES USED AS DEFAULT
DIC>
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /AUS/: aus
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC>
DIC> en-ph
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUS/: aus
PHASE TYPE /MATRIX/: sph
PHASE NAME: /NONE/: c1nb1_s
INFO: EQUILIBRIUM COMPOSITION AND FRACTION OF SPHEROID PHASES USED AS DEFAULT
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITIONS OF THE PHASES
DIC> @@
DIC> enter-composition
REGION NAME : /AUS/: ferr
PHASE NAME: /BCC_A2/: bcc
DEPENDENT COMPONENT ? /NB/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: C
TYPE /LINEAR/: lin
VALUE OF FIRST POINT : 1e-3
VALUE OF LAST POINT : /1E-3/: 1e-3
PROFILE FOR /NB/: nb
TYPE /LINEAR/: lin
VALUE OF FIRST POINT : 0.28
VALUE OF LAST POINT : /0.28/: 0.28
DIC>
DIC> en-co
REGION NAME : /AUS/: ferr
PHASE NAME: /BCC_A2/: c1nb1_s
USE EQUILIBRIUM VALUE /Y/: y
DIC>
DIC> enter-composition
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1#1/: fcc#1
DEPENDENT COMPONENT ? /NB/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: C
TYPE /LINEAR/: lin
VALUE OF FIRST POINT : 0.89
VALUE OF LAST POINT : /0.89/: 0.89
PROFILE FOR /NB/: nb
TYPE /LINEAR/: lin
VALUE OF FIRST POINT : 0.28
VALUE OF LAST POINT : /0.28/: 0.28
DIC>
DIC> en-co
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1#1/: c1nb1_s
USE EQUILIBRIUM VALUE /Y/: y
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /1/: 32400
AUTOMATIC TIMESTEP CONTROL /YES/: y
MAX TIMESTEP DURING INTEGRATION /3240/: 3240
INITIAL TIMESTEP : /1E-07/: 1e-8
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/: 1e-15
DIC>
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ SET THE REFERENCE PHASE OF CARBON AS GRAPHITE
DIC> @@
DIC> s-ref
Component: c
Reference state: grap
Temperature /*/: *
Pressure /100000/: 1e5
DIC>
DIC> @@
DIC> @@ SET THE BOUNDARY CONDITION.
DIC> @@ THE CARBON ACTIVITY IS THE ONE ON THE BOUNDARY
DIC> @@
DIC> s-cond
GLOBAL OR BOUNDARY CONDITION /GLOBAL/: bound
BOUNDARY /LOWER/: low
CONDITION TYPE /CLOSED_SYSTEM/: mix

```

```
Dependent substitutional element:FE
Dependent interstitial element:VA
TYPE OF CONDITION FOR COMPONENT C /ZERO_FLUX/: act
LOW TIME LIMIT /0/: 0
ACR(C) (TIME)= 1.0;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
TYPE OF CONDITION FOR COMPONENT NB /ZERO_FLUX/: zero
DIC>
DIC> @@
DIC> @@ ENABLE THE HOMOGENIZATION MODEL
DIC> @@
DIC> ho y y
INFO: HOMOGENIZATION MODEL ENABLED
DIC>
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exb7 Y
DIC>
DIC>
DIC> set-inter
--OK--
DIC>
```

exb7-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb7\run.DCM"
DIC> @@
DIC> @@ READ THE SET UP FROM FILE AND START THE SIMULATION
DIC> @@
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
*** ENTERING C1NB1_S AS A DIFFUSION NONE PHASE
DIC> read exb7
OK
DIC> sim yes
Region: AUS
double geometric
dense at outer boundaries, coarse at 0.50000E-09
lower part 1.0800 16
upper part 0.92593 16
Region: FERR
single geometric dense at 0.0000
1.0800 114
DEGREE OF IMPLICITY SET TO EULER BACKWARD
STARTING SIMULATION USING HOMOGENIZATION MODEL
-----
WARNING: ELEMENT C
IS BOTH INTERSTITIAL AND SUBSTITUTIONAL
AND RESULTS MUST BE INTERPRETED WITH CARE
INFO: PHASE WITH LIMITED SOLUBILITY OF ELEMENT(S) EXIST
AFallback PHASE ZZDICTRA GHOST WILL BE DEFINED
ALONG WITH THE FOLLOWING PARAMETERS:
G(ZZDICTRA_GHOST,C;0)-H298(GRAPHITE,C;0)
G(ZZDICTRA_GHOST,FE;0)-H298(BCC_A2,FE;0)
G(ZZDICTRA_GHOST,NB;0)-H298(BCC_A2,NB;0)
L(ZZDICTRA_GHOST,C,FE;0)
L(ZZDICTRA_GHOST,C,NB;0)
L(ZZDICTRA_GHOST,FE,NB;0)
INFO: FCC_A1#2 is stable but not entered in the simulation
WARNING:C1NB1_S HAS NO VOLUME FRACTION, CREATING ONE
WARNING:C1NB1_S HAS NO VOLUME FRACTION, CREATING ONE
Starting time-step t0= 0.000000 dt= 0.1000000E-07
Error 408
Error 408
Starting time-step t0= 0.48828125E-11 dt= 0.48828125E-11
Starting time-step t0= 0.97656250E-11 dt= 0.97656250E-11
Starting time-step t0= 0.19531250E-10 dt= 0.19531250E-10
Starting time-step t0= 0.39062500E-10 dt= 0.39062500E-10
Starting time-step t0= 0.78125000E-10 dt= 0.78125000E-10
Starting time-step t0= 0.15625000E-09 dt= 0.15625000E-09
Starting time-step t0= 0.31250000E-09 dt= 0.15625000E-09
Starting time-step t0= 0.46875000E-09 dt= 0.31250000E-09
Starting time-step t0= 0.78125000E-09 dt= 0.31250000E-09
Starting time-step t0= 0.10937500E-08 dt= 0.31250000E-09
Starting time-step t0= 0.14062500E-08 dt= 0.31250000E-09
Starting time-step t0= 0.17187500E-08 dt= 0.62500000E-09
Starting time-step t0= 0.23437500E-08 dt= 0.62500000E-09
Starting time-step t0= 0.29687500E-08 dt= 0.12500000E-08
Starting time-step t0= 0.42187500E-08 dt= 0.25000000E-08
Starting time-step t0= 0.67187500E-08 dt= 0.50000000E-08
Starting time-step t0= 0.11718750E-07 dt= 0.10000000E-07
Starting time-step t0= 0.21718750E-07 dt= 0.10000000E-07
Starting time-step t0= 0.31718750E-07 dt= 0.10000000E-07
Starting time-step t0= 0.41718750E-07 dt= 0.10000000E-07
Starting time-step t0= 0.51718750E-07 dt= 0.20000000E-07
Starting time-step t0= 0.71718750E-07 dt= 0.20000000E-07
Starting time-step t0= 0.91718750E-07 dt= 0.20000000E-07
Starting time-step t0= 0.11171875E-06 dt= 0.40000000E-07
Starting time-step t0= 0.15171875E-06 dt= 0.40000000E-07
Starting time-step t0= 0.19171875E-06 dt= 0.80000000E-07
Starting time-step t0= 0.27171875E-06 dt= 0.80000000E-07
Starting time-step t0= 0.35171875E-06 dt= 0.16000000E-06
Starting time-step t0= 0.51171875E-06 dt= 0.32000000E-06
Starting time-step t0= 0.83171875E-06 dt= 0.64000000E-06
Starting time-step t0= 0.14717187E-05 dt= 0.12800000E-05
Starting time-step t0= 0.27517187E-05 dt= 0.25600000E-05
Starting time-step t0= 0.53117187E-05 dt= 0.51200000E-05
Starting time-step t0= 0.10431719E-04 dt= 0.10240000E-04
Starting time-step t0= 0.20671719E-04 dt= 0.20480000E-04
Starting time-step t0= 0.41151719E-04 dt= 0.40960000E-04
Starting time-step t0= 0.82111719E-04 dt= 0.81920000E-04
Starting time-step t0= 0.16403172E-03 dt= 0.16384000E-03
Starting time-step t0= 0.32787172E-03 dt= 0.32768000E-03
Starting time-step t0= 0.65555172E-03 dt= 0.65536000E-03
Starting time-step t0= 0.13109117E-02 dt= 0.13107200E-02
Starting time-step t0= 0.26216317E-02 dt= 0.26214400E-02
Starting time-step t0= 0.52430717E-02 dt= 0.52428800E-02
Starting time-step t0= 0.10485952E-01 dt= 0.10485760E-01
Starting time-step t0= 0.20971712E-01 dt= 0.20971520E-01
Starting time-step t0= 0.41943232E-01 dt= 0.41943040E-01
Starting time-step t0= 0.83886272E-01 dt= 0.41943040E-01
Starting time-step t0= 0.12582931 dt= 0.41943040E-01
Starting time-step t0= 0.16777235 dt= 0.83886080E-01
Starting time-step t0= 0.18874387 dt= 0.20971520E-01
Error 408
Error 408
Starting time-step t0= 0.18907155 dt= 0.32768000E-03
Starting time-step t0= 0.18939923 dt= 0.32768000E-03
Starting time-step t0= 0.18972691 dt= 0.32768000E-03
Starting time-step t0= 0.19005459 dt= 0.32768000E-03
Starting time-step t0= 0.19038227 dt= 0.32768000E-03
Starting time-step t0= 0.19070995 dt= 0.32768000E-03
Starting time-step t0= 0.19103763 dt= 0.32768000E-03
Starting time-step t0= 0.19136531 dt= 0.32768000E-03
Starting time-step t0= 0.19169299 dt= 0.32768000E-03
Starting time-step t0= 0.19202067 dt= 0.65536000E-03
Starting time-step t0= 0.19267603 dt= 0.65536000E-03
Starting time-step t0= 0.19333139 dt= 0.65536000E-03
Starting time-step t0= 0.19398675 dt= 0.65536000E-03
Starting time-step t0= 0.19464211 dt= 0.65536000E-03
Starting time-step t0= 0.19529747 dt= 0.65536000E-03
Starting time-step t0= 0.19595283 dt= 0.13107200E-02
Starting time-step t0= 0.19726355 dt= 0.13107200E-02
```

```

Starting time-step t0= 0.19857427      dt= 0.13107200E-02
Starting time-step t0= 0.19988499      dt= 0.13107200E-02
Starting time-step t0= 0.20119571      dt= 0.13107200E-02
Starting time-step t0= 0.20250643      dt= 0.26214400E-02
Starting time-step t0= 0.20512787      dt= 0.26214400E-02
Starting time-step t0= 0.20774931      dt= 0.52428800E-02
Starting time-step t0= 0.21299219      dt= 0.52428800E-02
Starting time-step t0= 0.21823507      dt= 0.52428800E-02
Starting time-step t0= 0.22347795      dt= 0.10485760E-01
Starting time-step t0= 0.23396371      dt= 0.10485760E-01
Starting time-step t0= 0.24444947      dt= 0.10485760E-01
Starting time-step t0= 0.25493523      dt= 0.10485760E-01
Starting time-step t0= 0.26542099      dt= 0.10485760E-01
Starting time-step t0= 0.27590675      dt= 0.20971520E-01
Starting time-step t0= 0.29687827      dt= 0.20971520E-01
Starting time-step t0= 0.31784979      dt= 0.20971520E-01
Starting time-step t0= 0.33882131      dt= 0.20971520E-01
Starting time-step t0= 0.35979283      dt= 0.20971520E-01
Starting time-step t0= 0.38076435      dt= 0.20971520E-01
Starting time-step t0= 0.40173587      dt= 0.20971520E-01
Starting time-step t0= 0.42270739      dt= 0.20971520E-01
Starting time-step t0= 0.44367891      dt= 0.20971520E-01
Starting time-step t0= 0.46465043      dt= 0.20971520E-01
Starting time-step t0= 0.48562195      dt= 0.20971520E-01
Starting time-step t0= 0.50659347      dt= 0.41943040E-01

```

output ignored...

... output resumed

```

DELETING TIME-RECORD FOR TIME    26647.717
DELETING TIME-RECORD FOR TIME    26690.666
DELETING TIME-RECORD FOR TIME    26733.616
DELETING TIME-RECORD FOR TIME    26776.566
DELETING TIME-RECORD FOR TIME    26830.253
DELETING TIME-RECORD FOR TIME    26873.203
DELETING TIME-RECORD FOR TIME    26916.152
DELETING TIME-RECORD FOR TIME    26959.102
DELETING TIME-RECORD FOR TIME    27002.052
DELETING TIME-RECORD FOR TIME    27045.001
DELETING TIME-RECORD FOR TIME    27087.951
DELETING TIME-RECORD FOR TIME    27130.901
DELETING TIME-RECORD FOR TIME    27184.588
DELETING TIME-RECORD FOR TIME    27227.537
DELETING TIME-RECORD FOR TIME    27270.487
DELETING TIME-RECORD FOR TIME    27313.437
DELETING TIME-RECORD FOR TIME    27356.386
DELETING TIME-RECORD FOR TIME    27399.336
DELETING TIME-RECORD FOR TIME    27442.286
DELETING TIME-RECORD FOR TIME    27485.235
DELETING TIME-RECORD FOR TIME    27528.185
DELETING TIME-RECORD FOR TIME    27571.135
DELETING TIME-RECORD FOR TIME    27614.084
DELETING TIME-RECORD FOR TIME    27667.772
DELETING TIME-RECORD FOR TIME    27721.459
DELETING TIME-RECORD FOR TIME    27764.408
DELETING TIME-RECORD FOR TIME    27807.358
DELETING TIME-RECORD FOR TIME    27850.308
DELETING TIME-RECORD FOR TIME    27893.257
DELETING TIME-RECORD FOR TIME    27957.682
DELETING TIME-RECORD FOR TIME    28000.632
DELETING TIME-RECORD FOR TIME    28043.581
DELETING TIME-RECORD FOR TIME    28086.531
DELETING TIME-RECORD FOR TIME    28129.481
DELETING TIME-RECORD FOR TIME    28172.430
DELETING TIME-RECORD FOR TIME    28215.380
DELETING TIME-RECORD FOR TIME    28258.330
DELETING TIME-RECORD FOR TIME    28301.279
DELETING TIME-RECORD FOR TIME    28344.229
DELETING TIME-RECORD FOR TIME    28387.179
DELETING TIME-RECORD FOR TIME    28430.128
DELETING TIME-RECORD FOR TIME    28473.078
DELETING TIME-RECORD FOR TIME    28516.028
DELETING TIME-RECORD FOR TIME    28558.977
DELETING TIME-RECORD FOR TIME    28601.927
DELETING TIME-RECORD FOR TIME    28644.877
DELETING TIME-RECORD FOR TIME    28687.826
DELETING TIME-RECORD FOR TIME    28730.776
DELETING TIME-RECORD FOR TIME    28773.726
DELETING TIME-RECORD FOR TIME    28848.888
DELETING TIME-RECORD FOR TIME    28891.837
DELETING TIME-RECORD FOR TIME    28945.524
DELETING TIME-RECORD FOR TIME    28988.474
DELETING TIME-RECORD FOR TIME    29052.899
DELETING TIME-RECORD FOR TIME    29095.848
DELETING TIME-RECORD FOR TIME    29138.798
DELETING TIME-RECORD FOR TIME    29181.748
DELETING TIME-RECORD FOR TIME    29224.697
DELETING TIME-RECORD FOR TIME    29267.647
DELETING TIME-RECORD FOR TIME    29310.597
DELETING TIME-RECORD FOR TIME    29353.546
DELETING TIME-RECORD FOR TIME    29396.496
DELETING TIME-RECORD FOR TIME    29439.446
DELETING TIME-RECORD FOR TIME    29482.395
DELETING TIME-RECORD FOR TIME    29536.082
DELETING TIME-RECORD FOR TIME    29579.032
DELETING TIME-RECORD FOR TIME    29621.982
DELETING TIME-RECORD FOR TIME    29664.931
DELETING TIME-RECORD FOR TIME    29707.881
DELETING TIME-RECORD FOR TIME    29750.831
DELETING TIME-RECORD FOR TIME    29793.780
DELETING TIME-RECORD FOR TIME    29836.730
DELETING TIME-RECORD FOR TIME    29879.680
DELETING TIME-RECORD FOR TIME    29922.629
DELETING TIME-RECORD FOR TIME    29965.579
DELETING TIME-RECORD FOR TIME    30008.529
DELETING TIME-RECORD FOR TIME    30051.478
DELETING TIME-RECORD FOR TIME    30094.428
DELETING TIME-RECORD FOR TIME    30137.378
DELETING TIME-RECORD FOR TIME    30180.327
DELETING TIME-RECORD FOR TIME    30223.277
DELETING TIME-RECORD FOR TIME    30266.227
DELETING TIME-RECORD FOR TIME    30309.176

```

DELETING TIME-RECORD FOR TIME 30352.126
DELETING TIME-RECORD FOR TIME 30395.076
DELETING TIME-RECORD FOR TIME 30438.025
DELETING TIME-RECORD FOR TIME 30491.713
DELETING TIME-RECORD FOR TIME 30534.662
DELETING TIME-RECORD FOR TIME 30599.087
DELETING TIME-RECORD FOR TIME 30642.036
DELETING TIME-RECORD FOR TIME 30706.461
DELETING TIME-RECORD FOR TIME 30749.411
DELETING TIME-RECORD FOR TIME 30813.835
DELETING TIME-RECORD FOR TIME 30856.785
DELETING TIME-RECORD FOR TIME 30899.734
DELETING TIME-RECORD FOR TIME 30942.684
DELETING TIME-RECORD FOR TIME 30985.634
DELETING TIME-RECORD FOR TIME 31028.583
DELETING TIME-RECORD FOR TIME 31082.271
DELETING TIME-RECORD FOR TIME 31125.220
DELETING TIME-RECORD FOR TIME 31178.907
DELETING TIME-RECORD FOR TIME 31221.857
DELETING TIME-RECORD FOR TIME 31264.807
DELETING TIME-RECORD FOR TIME 31307.756
DELETING TIME-RECORD FOR TIME 31350.706
DELETING TIME-RECORD FOR TIME 31393.656
DELETING TIME-RECORD FOR TIME 31436.605
DELETING TIME-RECORD FOR TIME 31479.555
DELETING TIME-RECORD FOR TIME 31522.505
DELETING TIME-RECORD FOR TIME 31565.454
DELETING TIME-RECORD FOR TIME 31608.404
DELETING TIME-RECORD FOR TIME 31651.354
DELETING TIME-RECORD FOR TIME 31715.778
DELETING TIME-RECORD FOR TIME 31758.728
DELETING TIME-RECORD FOR TIME 31801.678
DELETING TIME-RECORD FOR TIME 31844.627
DELETING TIME-RECORD FOR TIME 31887.577
DELETING TIME-RECORD FOR TIME 31930.527
DELETING TIME-RECORD FOR TIME 31973.476
DELETING TIME-RECORD FOR TIME 32016.426
DELETING TIME-RECORD FOR TIME 32059.376
DELETING TIME-RECORD FOR TIME 32102.325
DELETING TIME-RECORD FOR TIME 32145.275
DELETING TIME-RECORD FOR TIME 32188.225
DELETING TIME-RECORD FOR TIME 32231.174
DELETING TIME-RECORD FOR TIME 32274.124
DELETING TIME-RECORD FOR TIME 32317.074

KEEPING TIME-RECORD FOR TIME 32360.023
AND FOR TIME 32400.000
WORKSPACE RECLAIMED

INTERPOLATION SCHEME USED THIS FRACTION OF
THE ALLOCATED MEMORY: 0.736891040822413
EFFICIENCY FACTOR: 34.6331929724956
MEMORY FRACTION USAGE PER BRANCH:
1.000000000000000
0.538497038689332
0.591018590064088

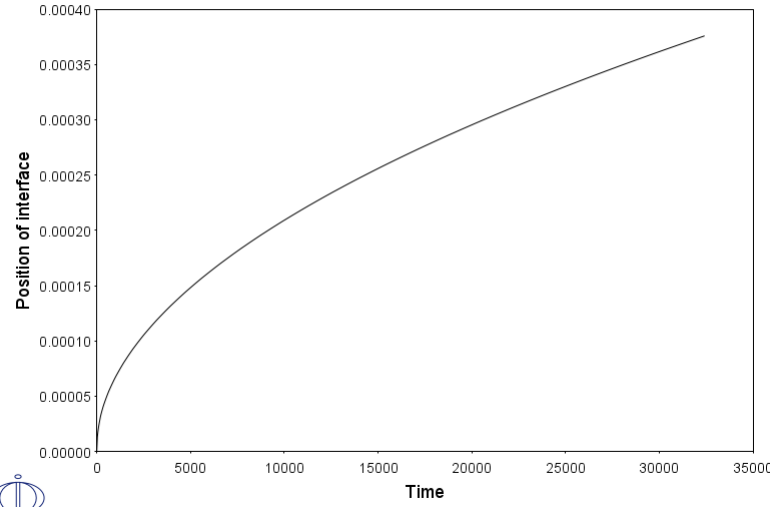
DEALLOCATING

TIMESTEP AT 32400.0000 SELECTED

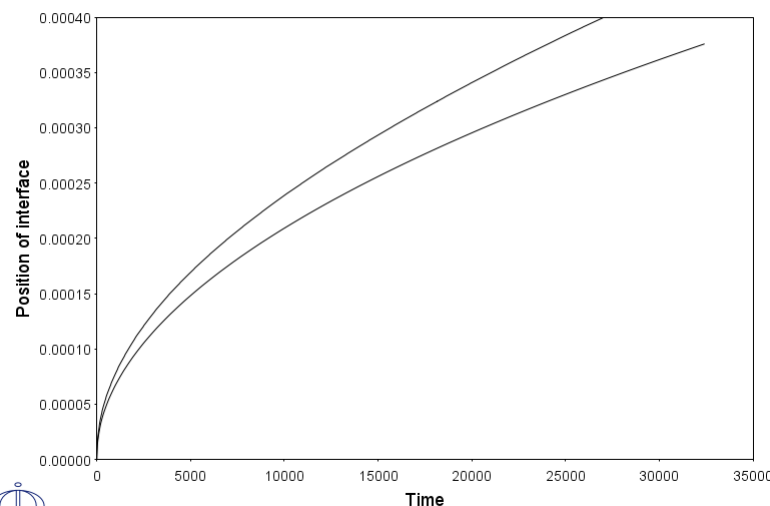
DIC>
DIC> set-inter
--OK---
DIC>

exb7-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exb7\plot.DCM"
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 3.24000E+04
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
*** ENTERING C1NB1_S AS A DIFFUSION NONE PHASE
DIC> read exb7
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ PLOT THE INTERFACE POSITION AS A FUNCTION OF TIME
POST-1: @@
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-d-a y po-o-in aus upp
POST-1:
POST-1: plot
2021.05.13.11.41.39
UPPER INTERFACE OF REGION "AUS#1"
CELL #1
```



```
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: @@
POST-1: @@ APPEND DATA FROM A CORRESPONDING SIMULATION
POST-1: @@ WITHOUT NIOBIUM
POST-1: @@
POST-1:
POST-1: app y fec.exp 0
DATASET NUMBER(s): /-1/: 1
POST-1:
POST-1: plot
2021.05.13.11.41.55
UPPER INTERFACE OF REGION "AUS#1"
CELL #1
```



POST-1:

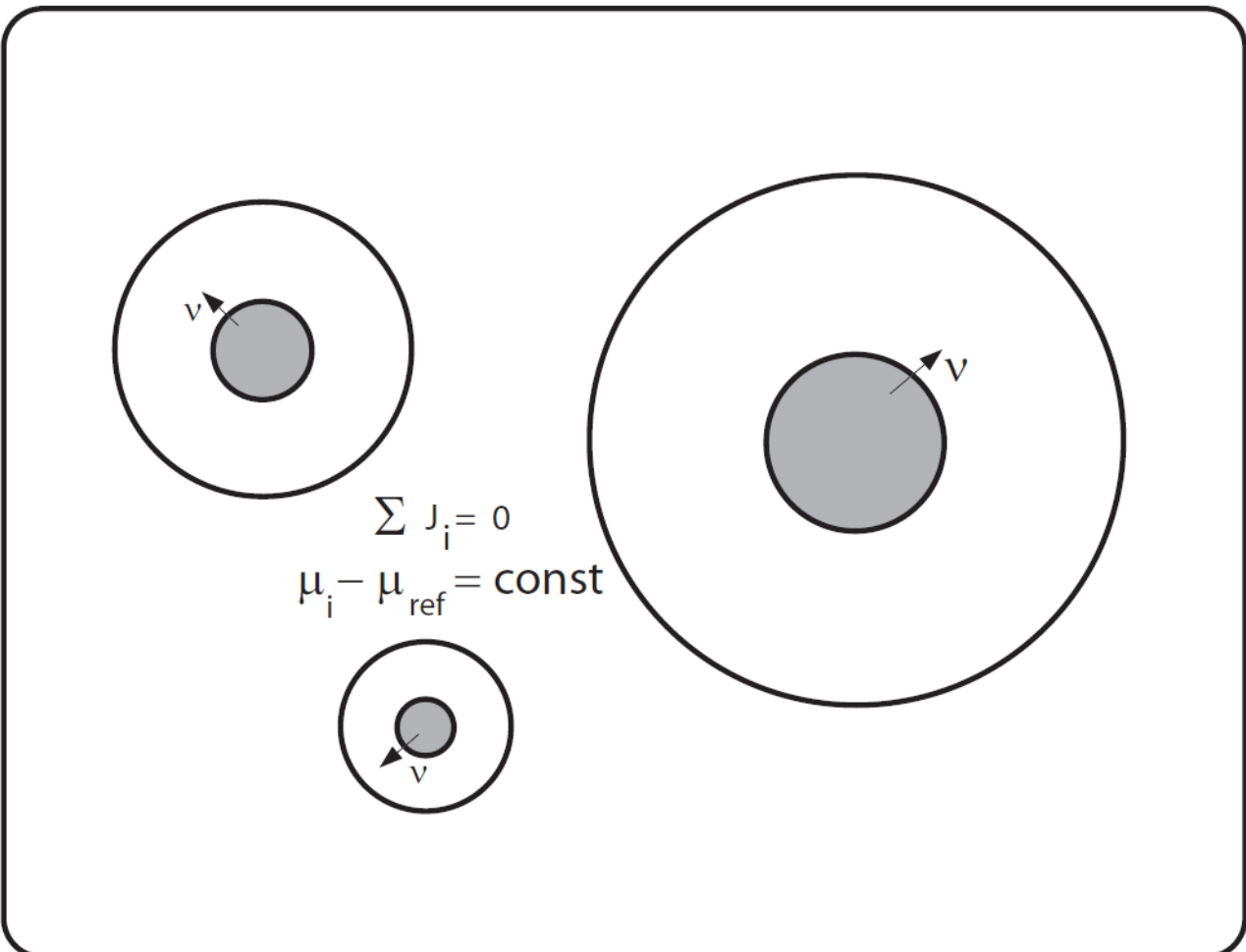
POST-1: set-inter

--OK---

POST-1:



Cell Calculations



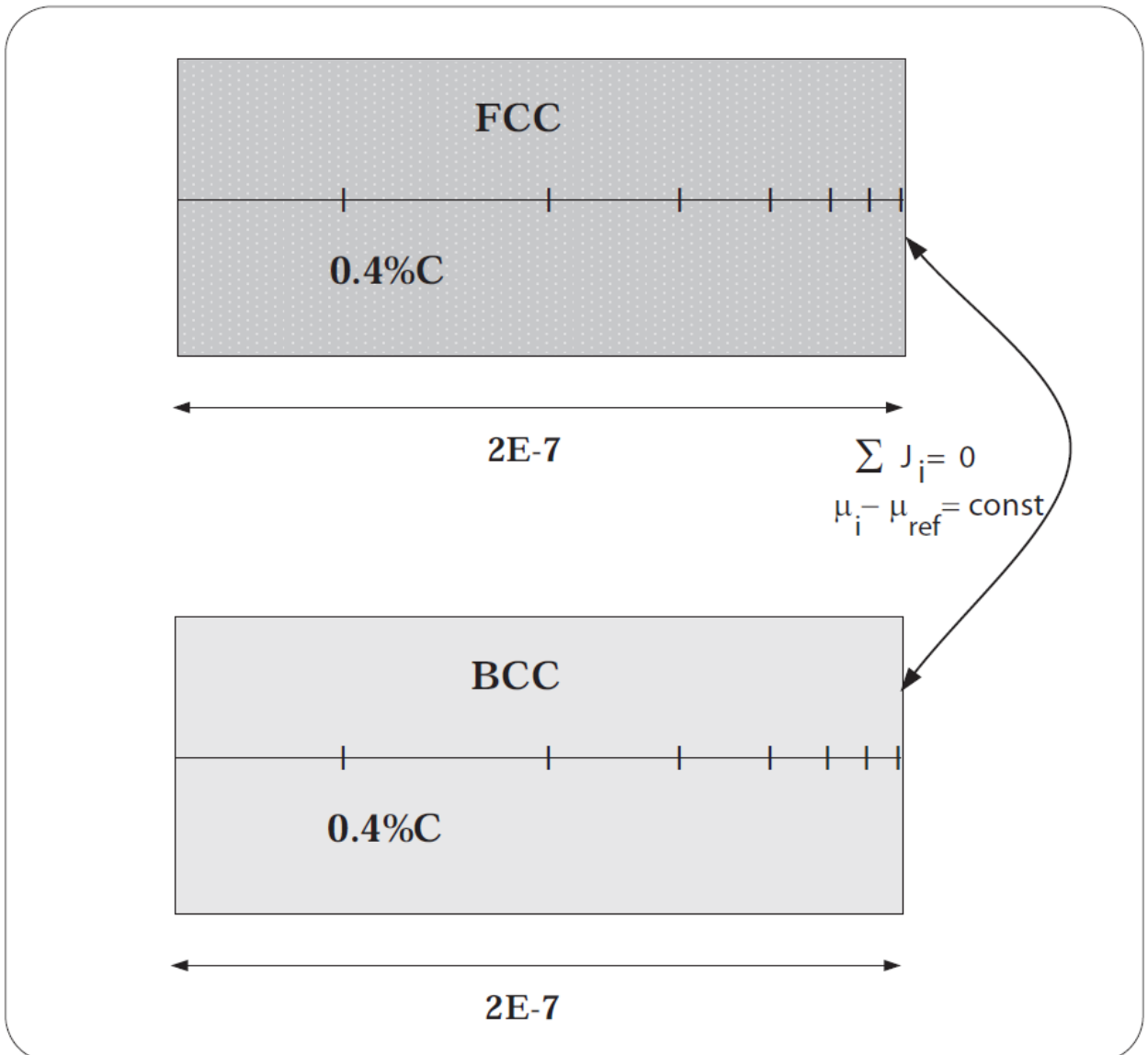


Example exc1

'Carbon cannon' in α/γ Fe-C system: Two-cell calculation

This example simulates what happens to a ferrite plate that has inherited the carbon content of its parent austenite. The ferrite plate formed is embedded in an austenite matrix. This setup corresponds to a proposed mechanism for formation of Widmannstätten ferrite or for the ferrite phase of the bainite structure. It is assumed that the phase boundary between ferrite and austenite is immobile, this is achieved in the simulation by putting the ferrite and the austenite in two different cells. See also M. Hillert, L. Höglund and J. Ågren: Acta Metall. Mater. 41 (1993), pp.1951-1957.

$T = 673\text{K}$



excl-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\excl\setup.DCM"

SYS: i>_@
NO SUCH COMMAND, USE HELP
SYS: @@ Cell calculation.
SYS: @@ Carbon cannon in ferrite/austenite: Fe-C system, 2-cell calculation
SYS: @@ This example simulates what happens to a ferrite plate that has
SYS: @@ inherited the carbon content of its parent austenite. The ferrite
SYS: @@ plate formed is embedded in an austenite matrix. This setup
SYS: @@ corresponds to a proposed mechanism for formation of Widmannst tten
SYS: @@ ferrite or for the ferrite phase of the bainite structure. It is
SYS: @@ assumed that the phase boundary between ferrite and austenite is
SYS: @@ immobile, this is achieved in the simulation by putting the ferrite
SYS: @@ and the austenite in two different cells. See also M. Hillert,
SYS: @@ L. H glund and J.  gren: Acta Metall. Mater. 41 (1993), pp.1951-1957.
SYS: -----
NO SUCH COMMAND, USE HELP

SYS:
SYS: @@ excl_setup.DCM
SYS:
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:
TDE_TCFE11: @@
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: @@
TDE_TCFE11: sw fedemo
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-sys fe c
FE C DEFINED
TDB_FEDEMO: rej ph * all
GAS:G LIQUID:L BCC_A2
LAVES_PHASE_C14 CRCC_A12 CEMENTITE
CUB_A13 DIAMOND_FCC_A4 FCC_A1
GRAPHITE HCP_A3 KSI_CARBIDE
M23C6 M5C2 M7C3
REJECTED
TDB_FEDEMO: res ph fcc,bcc
FCC_A1 BCC_A2 RESTORED
TDB_FEDEMO: get
11:43:13,672 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES
ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'P. Franke, estimated parameter within SGTE, 2007; Fe-C, Ni-C, Mo-C, C-Mn'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'

-OK-

TDB_FEDEMO: @@
TDE_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE DATA
TDE_FEDEMO: @@
TDB_FEDEMO: app mfdemo
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys fe c
FE C DEFINED
APP: rej ph * all
BCC_A2 FCC_A1 REJECTED
APP: res ph fcc,bcc
FCC_A1 BCC_A2 RESTORED
APP: get
ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Z. Metallkunde 85(1994)498-501; C and N diffusion in bcc Cr
-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'

```

-OK-
APP:
APP: @@ ENTER THE DICTRA MONITOR
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 673; * N
DIC>
DIC> @@
DIC> @@ IN THE FIRST CELL
DIC> @@
DIC> @@ ENTER REGION aus CONTAINING AUSTENITE
DIC> @@ ENTER A GEOMETRICAL GRID INTO THAT REGION
DIC> @@ ENTER THE INITIAL COMPOSITION INTO THE AUSTENITE
DIC> @@
DIC> enter-region aus
DIC> enter-grid aus 0.2e-6 AUTO
DIC> enter-phase act aus matrix fcc_a1#1
DIC>
DIC> enter-composition
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1/: fcc_a1#1
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: c
TYPE /LINEAR/: lin 0.4 0.4
DIC>
DIC> @@
DIC> @@ IN THE SECOND CELL
DIC> @@
DIC> create-new-cell
CELL DISTRIBUTION FACTOR /1/: 1
CREATING NEW CELL, NUMBER: 2
CELL 2 SELECTED
DIC-2>
DIC-2> @@
DIC-2> @@ ENTER REGION fer CONTAINING FERRITE
DIC-2> @@ ENTER A GEOMETRICAL GRID INTO THAT REGION
DIC-2> @@ ENTER THE INITIAL COMPOSITION INTO THE FERRITE
DIC-2> @@
DIC-2> enter-region fer
DIC-2>
DIC-2>
DIC-2>
DIC-2> enter-grid fer 0.2e-6 AUTO
DIC-2> enter-phase act fer matrix bcc_a2#1
DIC-2>
DIC-2> enter-composition
REGION NAME : /FER/: fer
PHASE NAME: /BCC_A2/: bcc_a2#1
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: c
TYPE /LINEAR/: lin 0.4 0.4
DIC-2>
DIC-2> @@
DIC-2> @@ SET THE SIMULATION TIME AND OTHER SIMULATION PARAMETERS
DIC-2> @@
DIC-2> set-simulation-time
END TIME FOR INTEGRATION /.1/: 0.5
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /.05/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC-2>
DIC-2>
DIC-2>
DIC-2> @@
DIC-2> @@ USE IMPLICIT (1) TIME INTEGRATION
DIC-2> @@
DIC-2> set-simulation-cond
NS01A PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA Timesteps IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/:
VARY POTENTIALS OR ACTIVITIES OR LNAC : /POTENTIAL/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/:
DEGREE OF IMPURITY WHEN INTEGRATING PDES (AUTO, 0 -> 0.5 -> 1): /AUTO/: 1.0
MAX TIMESTEP CHANGE PER Timestep : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/:
DIC-2> @@
DIC-2> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC-2> @@
DIC-2> save excl Y
DIC-2>
DIC-2> set-inter
--OK--
DIC-2>

```

excl-run

DIC-2>About
NO SUCH COMMAND, USE HELP
DIC-2>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\excl\run.DCM"
DIC-2>
DIC-2>
DIC-2> @@ excl_run.DCM
DIC-2>
DIC-2> @@
DIC-2> @@ READ THE WORKSPACE AND START THE SIMULATION
DIC-2> @@
DIC-2> go d-m
TIME STEP AT TIME 0.00000E+00
DIC-2> read excl
OK
DIC> sim
Region: AUS
single geometric dense at 0.20000E-06
0.89105 93
Region: FER
single geometric dense at 0.20000E-06
0.96940 72
U-FRACTION IN SYSTEM: C = .018673311178274 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
U-FRACTION IN SYSTEM: C = .018673311178274 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
3.02236653202599 3.02233686631279 1.73539217981896 0.149154115934069 3.651349914660366E-
003 1.241094884564162E-005 8.565304874058873E-010 1.957154250803553E-016 TIME = 0.1000000E-06 DT = 0.1000000E-
06 SUM OF SQUARES = 0.19571543E-15
U-FRACTION IN SYSTEM: C = .0186733111782041 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 0 seconds
1.814486158120001E-002 1.814519617402008E-002 1.136805123873663E-003 1.313921959983013E-004 2.632580953357226E-
006 8.905377903305629E-009 7.068475150799260E-013 1.930361806076383E-019 TIME = 0.3000000E-06 DT = 0.2000000E-
06 SUM OF SQUARES = 0.19303618E-18
U-FRACTION IN SYSTEM: C = .018673311178248 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 1 seconds
1.558906225565544E-002 1.558854133087159E-002 4.100498874343879E-004 7.060889806107446E-006 2.331165470476944E-
009 1.390124489563625E-014 2.721728775170181E-023 TIME = 0.7000000E-06 DT = 0.4000000E-
06 SUM OF SQUARES = 0.27217288E-22
U-FRACTION IN SYSTEM: C = .0186733111782479 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 1 seconds
1.821516411651870E-003 1.821361287293460E-003 6.815349783573381E-006 2.173574098267615E-008 2.249148547273783E-
013 7.485201286575788E-021 TIME = 0.1500000E-05 DT = 0.8000000E-06 SUM OF SQUARES = 0.74852013E-20
U-FRACTION IN SYSTEM: C = .0186733111782513 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 0 seconds
1.148030822150194E-003 1.147942845570995E-003 5.356292071700177E-006 2.090542623118400E-008 3.251711425242003E-
013 1.994571424996989E-020 TIME = 0.3100000E-05 DT = 0.1600000E-05 SUM OF SQUARES = 0.19945714E-19
U-FRACTION IN SYSTEM: C = .0186733111782626 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 1 seconds
4.719380441682447E-002 4.718971708926128E-002 1.715083336223400E-004 5.314202566247585E-007 5.019723862260593E-
012 1.537019069482865E-019 TIME = 0.6300000E-05 DT = 0.3200000E-05 SUM OF SQUARES = 0.15370191E-18
U-FRACTION IN SYSTEM: C = .0186733111782689 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 0 seconds
2.234625320733774E-002 2.234424967091886E-002 7.572089234366166E-005 2.203921096476018E-007 1.896529491363427E-
012 4.786375080913457E-020 TIME = 0.1270000E-04 DT = 0.6400000E-05 SUM OF SQUARES = 0.47863751E-19
U-FRACTION IN SYSTEM: C = .0186733111782759 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 1 seconds
1.086668823010137E-002 1.086569692023525E-002 3.555404233756766E-005 1.001885057921724E-007 8.076952345751924E-
013 1.848609630463012E-020 TIME = 0.2550000E-04 DT = 0.1280000E-04 SUM OF SQUARES = 0.18486096E-19
U-FRACTION IN SYSTEM: C = .0186733111782846 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 0 seconds
5.366713503686531E-003 5.366220092923794E-003 1.728153076345393E-005 4.798302038763194E-008 3.754841343847700E-
013 8.215615126752323E-021 TIME = 0.5110000E-04 DT = 0.2560000E-04 SUM OF SQUARES = 0.82156151E-20

output ignored...

... output resumed

6.039853793880095E-005 6.047186596748560E-005 4.848043613502960E-010 9.862891723403028E-015 1.467305206463634E-
024 TIME = 0.19893903 DT = 0.25600018E-01 SUM OF SQUARES = 0.14673052E-23
U-FRACTION IN SYSTEM: C = .0186733111740062 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 1 seconds
7.273325839371019E-005 7.280144598526400E-005 1.236135683692217E-009 3.257722575993787E-014 1.031073170069598E-
023 TIME = 0.22719739 DT = 0.28258357E-01 SUM OF SQUARES = 0.10310732E-22
U-FRACTION IN SYSTEM: C = .0186733111740106 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 1 seconds
9.378548650229065E-005 9.385005622307225E-005 2.935664553618166E-009 1.320416476491613E-013 1.676183956401913E-
022 TIME = 0.25952755 DT = 0.32330166E-01 SUM OF SQUARES = 0.16761840E-21
U-FRACTION IN SYSTEM: C = .0186733111740301 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 0 seconds
1.362089908213268E-004 1.362721964684466E-004 9.568726641728680E-009 8.452436749051035E-013 4.738384692808313E-
021 TIME = 0.29855549 DT = 0.39027939E-01 SUM OF SQUARES = 0.47383847E-20
U-FRACTION IN SYSTEM: C = .0186733111741636 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 1 seconds
1.869123526775527E-004 1.86970575317960E-004 2.773275155988044E-008 5.142445211417035E-012 1.275528427824585E-
019 TIME = 0.34855549 DT = 0.5000000E-01 SUM OF SQUARES = 0.12755284E-18
U-FRACTION IN SYSTEM: C = .018673311175056 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 0 seconds
1.969093575072288E-005 1.970752138708888E-005 3.235768579997403E-010 8.585338062027034E-015 3.766017451083959E-
024 TIME = 0.39855549 DT = 0.5000000E-01 SUM OF SQUARES = 0.37660175E-23
U-FRACTION IN SYSTEM: C = .0186733111750612 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 1 seconds
1.788524242369600E-006 1.784034764465464E-006 5.697860461543272E-012 6.94935993599005E-
018 TIME = 0.44855549 DT = 0.5000000E-01 SUM OF SQUARES = 0.69493599E-17
U-FRACTION IN SYSTEM: C = .0186733111684701 FE = 1
TOTAL SIZE OF SYSTEM: 4E-07 [m]
CPU time used in timestep 0 seconds

2.634178844977439E-005 2.632601716217299E-005 1.094912108301709E-009 4.599606206421634E-014 2.948843827665331E-
 020 TIME = 0.49855549 DT = 0.50000000E-01 SUM OF SQUARES = 0.29488438E-19
 U-FRACTION IN SYSTEM: C = .0186733111680409 FE = 1
 TOTAL SIZE OF SYSTEM: 4E-07 [m]
 CPU time used in timestep 1 seconds
 2.919561855910987E-002 2.919279873727918E-002 4.211812980878393E-005 5.331008256885776E-008 1.007569798349511E-
 013 1.360161492072539E-022 TIME = 0.50000000E-01 DT = 0.14445092E-02 SUM OF SQUARES = 0.13601615E-21
 U-FRACTION IN SYSTEM: C = .0186733111680417 FE = 1
 TOTAL SIZE OF SYSTEM: 4E-07 [m]
 MUST SAVE WORKSPACE ON FILE
 WORKSPACE SAVED ON FILE
 RECLAIMING WORKSPACE
 DELETING TIME-RECORD FOR TIME 0.0000000
 DELETING TIME-RECORD FOR TIME 0.10000000E-06
 DELETING TIME-RECORD FOR TIME 0.30000000E-06
 DELETING TIME-RECORD FOR TIME 0.70000000E-06
 DELETING TIME-RECORD FOR TIME 0.15000000E-05
 DELETING TIME-RECORD FOR TIME 0.31000000E-05
 DELETING TIME-RECORD FOR TIME 0.63000000E-05
 DELETING TIME-RECORD FOR TIME 0.12700000E-04
 DELETING TIME-RECORD FOR TIME 0.25500000E-04
 DELETING TIME-RECORD FOR TIME 0.51100000E-04
 DELETING TIME-RECORD FOR TIME 0.10230000E-03
 DELETING TIME-RECORD FOR TIME 0.20470000E-03
 DELETING TIME-RECORD FOR TIME 0.40950000E-03
 DELETING TIME-RECORD FOR TIME 0.81910000E-03
 DELETING TIME-RECORD FOR TIME 0.16383000E-02
 DELETING TIME-RECORD FOR TIME 0.32767000E-02
 DELETING TIME-RECORD FOR TIME 0.65535000E-02
 DELETING TIME-RECORD FOR TIME 0.13107100E-01
 DELETING TIME-RECORD FOR TIME 0.26214300E-01
 DELETING TIME-RECORD FOR TIME 0.45264232E-01
 DELETING TIME-RECORD FOR TIME 0.64872172E-01
 DELETING TIME-RECORD FOR TIME 0.84957979E-01
 DELETING TIME-RECORD FOR TIME 0.10565621
 DELETING TIME-RECORD FOR TIME 0.12712011
 DELETING TIME-RECORD FOR TIME 0.14957043
 DELETING TIME-RECORD FOR TIME 0.17333901
 DELETING TIME-RECORD FOR TIME 0.19893903
 DELETING TIME-RECORD FOR TIME 0.22719739
 DELETING TIME-RECORD FOR TIME 0.25952755
 DELETING TIME-RECORD FOR TIME 0.29855549
 DELETING TIME-RECORD FOR TIME 0.34855549
 DELETING TIME-RECORD FOR TIME 0.39855549
 DELETING TIME-RECORD FOR TIME 0.44855549

 KEEPING TIME-RECORD FOR TIME 0.49855549
 AND FOR TIME 0.50000000
 WORKSPACE RECLAIMED

 TIMESTEP AT 0.500000000 SELECTED

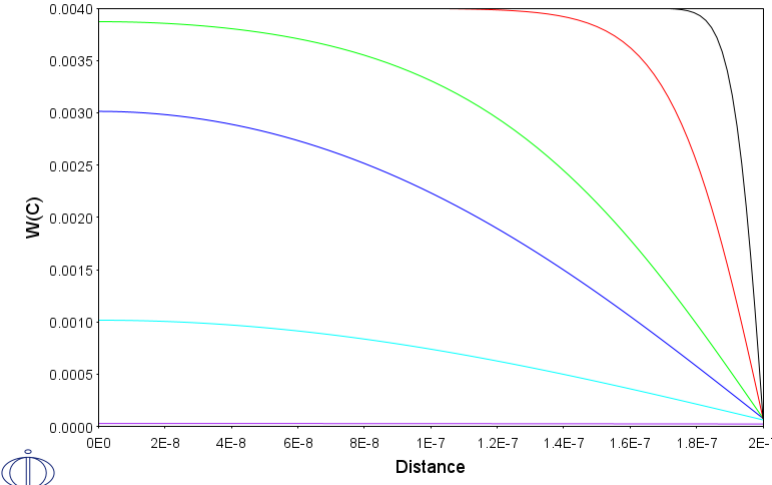
DIC>
DIC>
DIC>
DIC> set-inter
--OK--
DIC>

exc1-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exc1\plot.DCM"
DIC>
DIC>
DIC> @@ exc1_plot.DCM
DIC>
DIC> @@
DIC> FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE c1
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 5.00000E-01
DIC> read exc1
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ FIRST PLOT CARBON CONCENTRATION PROFILES IN FERRITE (CELL-2)
POST-1: @@ THEN SET THE DISTANCE AS X-AXIS (NOT THAT DISTANCE IS SET INDEPENDENT
POST-1: @@ VARIABLE AUTOMATICALLY) AND W-FRACTION CARBON AS Y-AXIS
POST-1: @@ REMEMBER THAT THE PLOT CONDITION ALSO MUST BE SET.
POST-1: @@
POST-1: select-cell
Number /NEXT/: 2
CELL 2 SELECTED
POST-2:
POST-2: @@
POST-2: @@ NOTICE THAT THE PROMPT INCLUDES THE CURRENT CELL NUMBER
POST-2: @@
POST-2: s-d-a x dist glo
INFO: Distance is set as independent variable
POST-2: s-d-a y w(c)
POST-2: s-p-c time .0001 .001 .01 .03 .1 .5
POST-2:
POST-2: @@
POST-2: @@ SET THE TITLE ON THE PLOTS
POST-2: @@
POST-2: set-title Figure c1.1
POST-2: plot
```

Figure c1.1

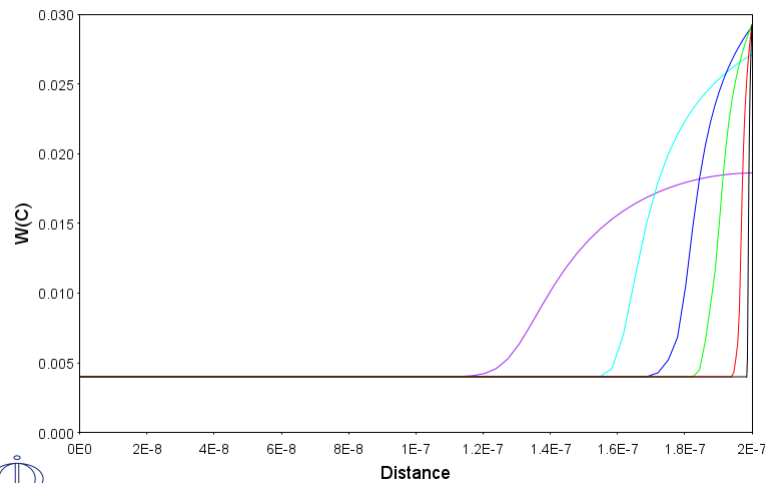
2021.05.13.11.45.38
Time=1E-04,0.001,0.01,0.03,0.1,0.5
CELL#2



```
POST-2:
POST-2:
POST-2:
POST-2:@?<_hit_return_to_continue_>
POST-2:
POST-2: @@
POST-2: @@ DO THE SAME THING FOR THE AUSTENITE (CELL-1)
POST-2: @@
POST-2: select-cell
Number /NEXT/: 1
CELL 1 SELECTED
POST-1: set-title Figure c1.2
POST-1: plot
```

Figure c1.2

2021.05.13.11.45.39
 Time = 1E-04, 001., 01., 03., 1., 5
 CELL #1

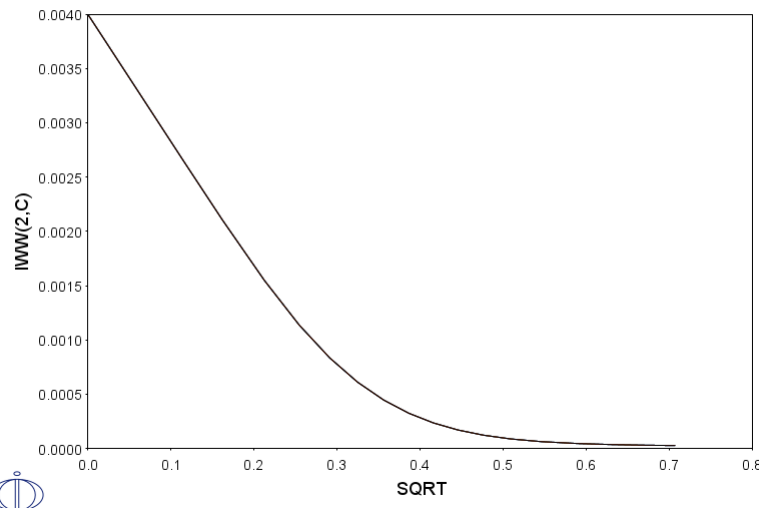


```

POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: @@
POST-1: @@ PLOT THE AVERAGE WEIGHT FRACTION OF CARBON IN FERRITE VS. SQUARE ROOT
POST-1: @@ OF TIME. START BY DEFINING A "SQUARE-ROOT-OF-TIME" FUNCTION.
POST-1: @@
POST-1: sel-cell 2
CELL 2 SELECTED
POST-2: enter func sqrt=sqrt(time);
POST-2: s-d-a x sqrt
POST-2: s-d-a y iww(2,c)
POST-2: s-i-v time
POST-2: set-title Figure c1.3
POST-2: plot
  
```

Figure c1.3

2021.05.13.11.45.40
 CELL #2

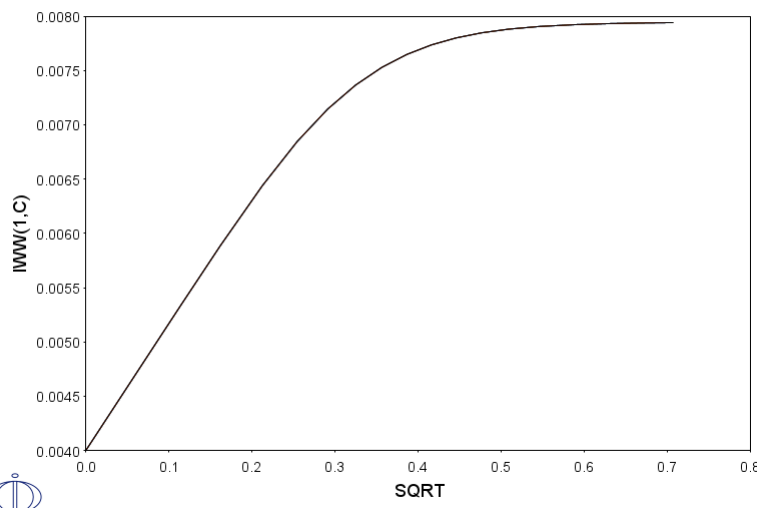


```

POST-2:
POST-2:
POST-2:
POST-2:@?<_hit_return_to_continue_>
POST-2:
POST-2: @@
POST-2: @@ DO THE SAME THING FOR THE AUSTENITE
POST-2: @@
POST-2: sel-cell 1
CELL 1 SELECTED
POST-1: s-d-a y iww(1,c)
POST-1: set-title Figure c1.4
POST-1: plot
  
```

Figure c1.4

2021.05.13.11.45.42
CELL #1



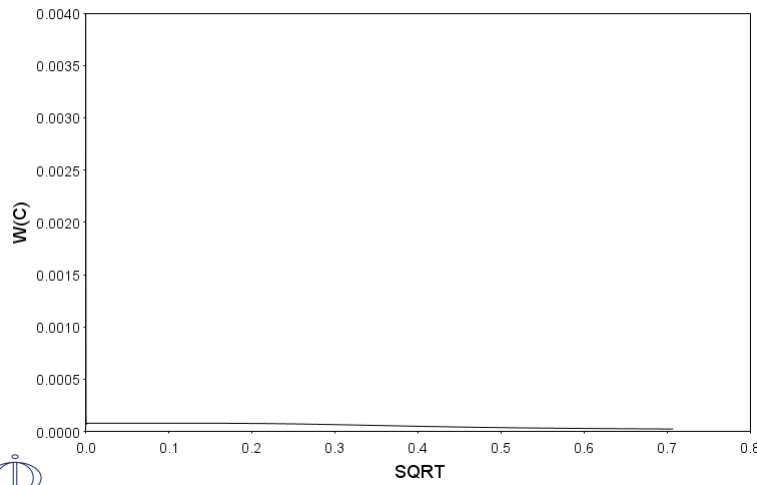
```

POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: @@
POST-1: @@ PLOT HOW THE CONCENTRATION IN FERRITE AT THE FERRITE/AUSTENITE BOUNDARY
POST-1: @@ V.S SQUARE ROOT OF TIME. THE FERRITE/AUSTENITE BOUNDARY IS REPRESENTED
POST-1: @@ BY THE CELL BOUNDARY I.E. THE "LAST" INTERFACE.
POST-1: @@
POST-1: sel-cell 2
CELL 2 SELECTED
POST-2: s-d-a y w(c)
POST-2: s-p-c interface last
POST-2: set-title Figure c1.5
POST-2: plot

```

Figure c1.5

2021.05.13.11.45.43
UPPER INTERFACE OF REGION "LAST"
CELL #2



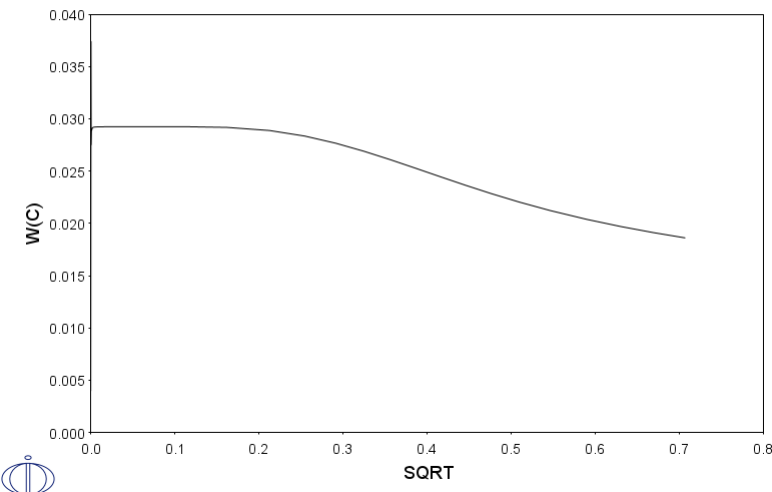
```

POST-2:
POST-2:
POST-2:
POST-2:@?<_hit_return_to_continue_>
POST-2:
POST-2: @@
POST-2: @@ DO THE SAME THING FOR THE AUSTENITE
POST-2: @@
POST-2: sel-cell 1
CELL 1 SELECTED
POST-1: set-title Figure c1.6
POST-1: plot

```

Figure c1.6

2021.05.13.11.45.44
UPPER INTERFACE OF REGION "LAST"
CELL #1



POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: set-inter
---OK---
POST-1:

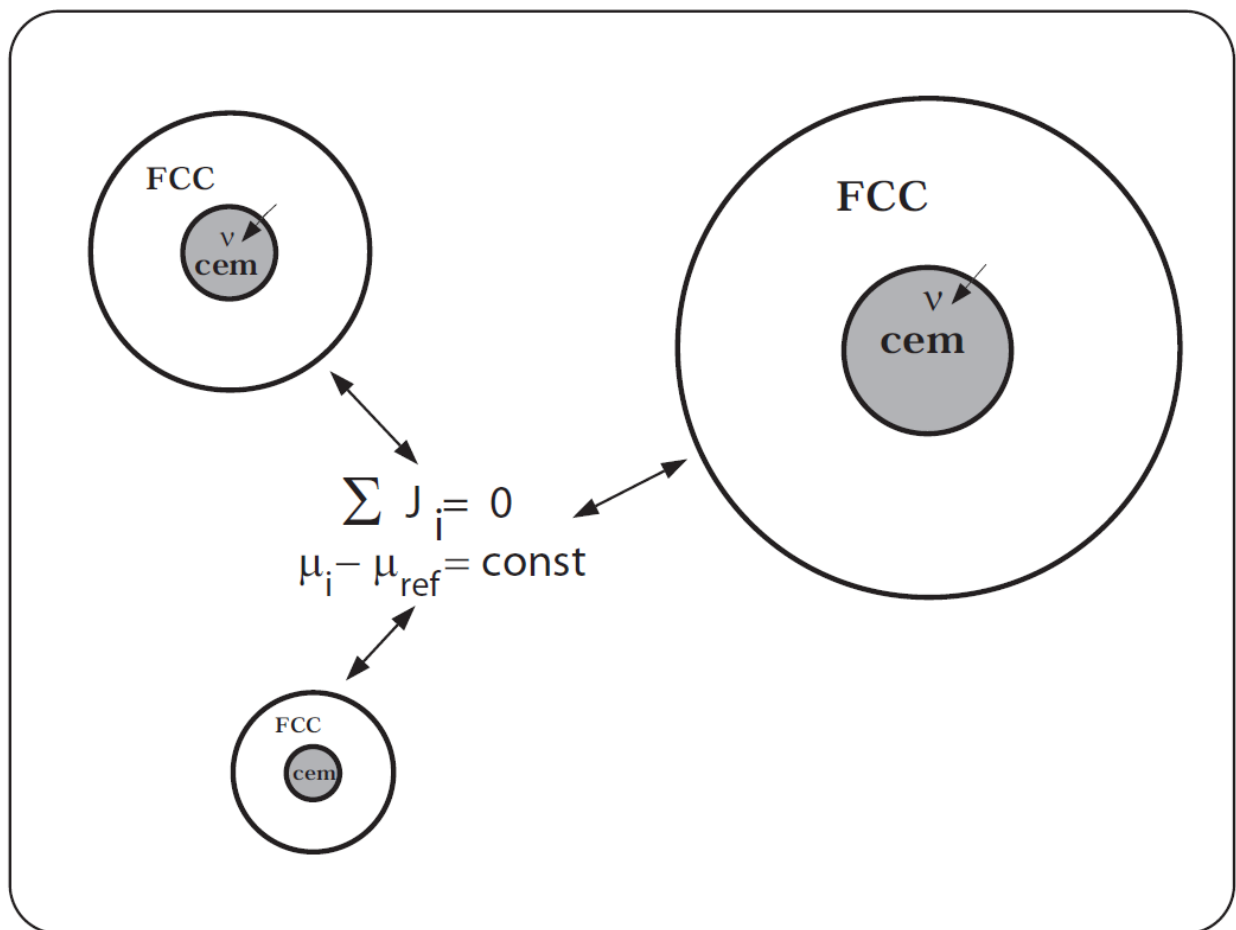


Example exc2

Cementite dissolution in an Fe-Cr-C alloy: Three particle sizes and three different cells

This example calculates the dissolution of cementite particles in an austenite matrix. This is the same as exc1 except that there are three particle sizes. Altogether six particles are considered using three different cells. This is to be able to represent some size distribution among the cementite particles. See also Z.-K. Liu, L. Höglund, B. Jönsson and J. Ågren: Metall.Trans.A, v. 22A (1991), pp. 1745-1752.

$$T = 118\text{K}$$



exc2-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exc2\setup.DCM"
SYS: i>_@
NO SUCH COMMAND, USE HELP
SYS: @@ Cell calculation.
SYS: @@ Cementite dissolution in an Fe-Cr-C alloy: Three particle sizes and
SYS: @@ three different cells
SYS: @@ This example calculates the dissolution of cementite particles
SYS: @@ in an austenite matrix. This example is the same as exc1 but
SYS: @@ instead there are three particle sizes. A total of six
SYS: @@ particles are considered using three different cells. This is to
SYS: @@ represent some size distribution among the cementite particles.
SYS: @@ See also Z.-K. Liu, L. Håglund, B. Jansson and J. Ågren:
SYS: @@ Metall.Trans.A, v. 22A (1991), pp. 1745-1752.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exc2_setup.DCM
SYS:
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: @@
TDB_TCFE11: switch fedemo
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-sys fe cr c
FE CR C
DEFINED
TDB_FEDEMO: rej ph * all
GAS:G LIQUID:L BCC_A2
LAVES_PHASE_C14 CBCC_A12 CEMENTITE
CHI_A12 CUB_A13 DIAMOND_FCC_A4
FCC_A1 GRAPHITE HCP_A3
KSI_CARBIDE M23C6 M3C2
M5C2 M7C3 SIGMA
REJECTED
TDB_FEDEMO: res ph fcc cementite
FCC_A1 CEMENTITE RESTORED
TDB_FEDEMO: get
11:47:01,675 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES
ELEMENTS
SPECIES
PHASES
Creating a new composition set FCC_A1#2
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'P. Villars, L.D. Calvert. Pearson's handbook of crystallographic data for intermetallic phases, Metals park, Ohio 1985: American society for metals'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'B. Hallstedt, D. Djurovic, J. von Appen, R. Dronskowsk, A. Dick, F. Koermann, T. Hickel, J. Neugebauer, CALPHAD, 34, 129 -33(2010); Fe-C'
'A.V. Khvan, B. Hallstedt, C. Broeckmann, CALPHAD, 46, 24 -33(2014); Cr-Fe -C'
'J. Bratberg, Z. Metallkd., 96 (2005) 335-344; Fe-Cr-Mo-C'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'P. Gustafsson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar volumes'
'R. Naraghi, Thermo-Calc Software AB, Sweden, 2016; FCC Fe-Cr-C and C-Cr-Ni'
'J-O. Andersson and B. Sundman, CALPHAD, 11 (1987) 83-92; TRITA 0270 (1986); CR-FE'
-OK-
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app mobfe4
Current database: Steels/Fe-Alloys Mobility v4.0

VA DEFINED
B2_BCC REJECTED
APP: def-sys fe cr c
FE CR C
DEFINED
APP: rej ph * all
BCC_A2 CEMENTITE FCC_A1
FE4N_LP1 HCP_A3 LIQUID:L
REJECTED
APP: res ph fcc cementite

```

FCC_A1          CEMENTITE RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Z. Metallkunde 85(1994)502-509; C diffusion in fcc Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'This parameter has been estimated'
-OK-
APP:
APP: @@ ENTER THE DICTRA MONITOR
APP: @@ go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob t 0 1183; * n
DIC>
DIC> @-----
DIC> @@ CELL NUMBER ONE
DIC> @@
DIC>
DIC> @@ ENTER REGIONS carb AND aus
DIC> @@
DIC> enter-region carb
DIC> enter-region aus
ATTACH TO REGION NAMED /CARB/:
ATTACHED TO THE RIGHT OF CARB /YES/:
DIC> @@
DIC> @@ ENTER GEOMTRICAL GRIDS INTO THE REGIONS
DIC> @@
DIC>
DIC> @@
DIC> @@ THE SIZE OF THE CEMENTITE PARTICLES ARE KNOWN AS WE ASSUME
DIC> @@ IT HAS BEEN MEASURED.
DIC> @@
DIC> enter-grid
REGION NAME : /CARB/: carb
WIDTH OF REGION /1/: 0.700000e-6
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ THE SIZE OF THE FCC REGION CAN BE CALCULATED FROM A MASS BALANCE
DIC> @@ AFTER ESTIMATING THE INITIAL COMPOSITIONS IN THE TWO PHASES.
DIC> @@
DIC> enter-grid
REGION NAME : /AUS/: aus
WIDTH OF REGION /1/: 7.1832993E-7
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER PHASES INTO REGIONS
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /CARB/: carb
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: cementite
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUS/: aus
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC> @@
DIC> @@ ENTER INITIAL VALUES FOR THE COMPOSITIONS IN THE PHASES
DIC> @@
DIC> enter-composition carb cementite w-f
PROFILE FOR /CR/: cr lin 0.12423326 0.12423326
DIC>
DIC> enter-composition aus fcc#1 fe w-f
PROFILE FOR /C/: cr lin 4.6615447E-3 4.6615447E-3
PROFILE FOR /CR/: c lin 1.5135207E-4 1.5135207E-4
DIC>
DIC> @@
DIC> @@ CELL NUMBER TWO
DIC> @@
DIC> create-new-cell
CELL DISTRIBUTION FACTOR /1/: 2
CREATING NEW CELL, NUMBER: 2
CELL 2 SELECTED
DIC-2>
DIC-2> @@
DIC-2> @@ ENTER REGIONS carb AND aus
DIC-2> @@
DIC-2> enter-region carb
DIC-2> enter-region aus
ATTACH TO REGION NAMED /CARB/:
ATTACHED TO THE RIGHT OF CARB /YES/:
DIC-2> @@
DIC-2> @@ ENTER GEOMTRICAL GRIDS INTO THE REGIONS
DIC-2> @@
DIC-2> enter-grid carb 0.300000e-6 AUTO
DIC-2> enter-grid aus 3.0785568E-7 AUTO
DIC-2>
DIC-2> @@
DIC-2> @@ ENTER PHASES INTO THE REGIONS

```

```

DIC-2> @@
DIC-2> enter-phase act carb matrix cementite
DIC-2> enter-phase act aus matrix fcc#1
DIC-2>
DIC-2> @@
DIC-2> @@ ENTER INITIAL VALUES FOR THE COMPOSITIONS IN THE PHASES
DIC-2> @@
DIC-2> enter-composition carb cementite w-f
PROFILE FOR /CR/: cr lin 0.12423326 0.12423326
DIC-2>
DIC-2> enter-composition aus fcc#1 fe w-f
PROFILE FOR /C/: cr lin 4.6615447E-3 4.6615447E-3
PROFILE FOR /CR/: c lin 1.5135207E-4 1.5135207E-4
DIC-2>
DIC-2> @@
-----  

DIC-2> @@ CELL NUMBER THREE
DIC-2> @@
-----  

DIC-2> create-new-cell
CELL DISTRIBUTION FACTOR /1/: 3
CREATING NEW CELL, NUMBER: 3
CELL 3 SELECTED
DIC-3>
DIC-3> @@
DIC-3> @@ ENTER REGIONS carb AND aus
DIC-3> @@
DIC-3> enter-region carb
DIC-3> enter-region aus
ATTACH TO REGION NAMED /CARB/:
ATTACHED TO THE RIGHT OF CARB /YES/:
DIC-3> @@
DIC-3> @@ ENTER GEOMTRICAL GRIDS INTO THE REGIONS
DIC-3> @@
DIC-3> enter-grid carb 0.525500e-6 AUTO
DIC-3> enter-grid aus 5.3926054E-7 AUTO
DIC-3>
DIC-3> @@
DIC-3> @@ ENTER PHASES INTO REGIONS
DIC-3> @@
DIC-3> enter-phase act carb matrix cementite
DIC-3> enter-phase act aus matrix fcc#1
DIC-3>
DIC-3> @@
DIC-3> @@ ENTER INITIAL VALUES FOR THE COMPOSITIONS IN THE PHASES
DIC-3> @@
DIC-3> enter-composition carb cementite w-f
PROFILE FOR /CR/: cr lin 0.12423326 0.12423326
DIC-3>
DIC-3> enter-composition aus fcc#1 fe w-f
PROFILE FOR /C/: cr lin 4.6615447E-3 4.6615447E-3
PROFILE FOR /CR/: c lin 1.5135207E-4 1.5135207E-4
DIC-3>
DIC-3> @@
-----  

DIC-3> @@ GLOBAL CONDITIONS
DIC-3> @@
-----  

DIC-3> @@
DIC-3> @@
DIC-3> @@ SET TO A SPHERICAL GEOMETRY
DIC-3> @@
DIC-3> enter-geo 2
DIC-3>
DIC-3> @s-n-l
's-n-l' is not recognized as an internal or external command,
operable program or batch file.
DIC-3> @
DIC-3> @1E-3
'1E-3' is not recognized as an internal or external command,
operable program or batch file.
DIC-3>
DIC-3>
DIC-3>
DIC-3>
DIC-3>
DIC-3> @@
DIC-3> @@ SET THE SIMULATION TIME
DIC-3> @@
DIC-3> set-simulation-time
END TIME FOR INTEGRATION /.1/: 10000
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /1000/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC-3>
DIC-3> @@
DIC-3> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC-3> @@
DIC-3> save exc2 Y
DIC-3>
DIC-3>
DIC-3> set-inter
--OK--
DIC-3>

```

exc2-run

DIC-3>About
NO SUCH COMMAND, USE HELP
DIC-3>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exc2\run.DCM"
DIC-3>
DIC-3>
DIC-3> @@ exc2_run.DCM
DIC-3>
DIC-3> @@
DIC-3> @@ READ THE SET UP FROM FILE AND START THE SIMULATION
DIC-3> @@
DIC-3>
DIC-3> go d-m
TIME STEP AT TIME 0.00000E+00
DIC-3> read exc2
OK
DIC> sim yes
Region: CARB
single geometric dense at 0.70000E-06
0.84611 96
Region: AUS
double geometric
dense at outer boundaries, coarse at 0.35916E-06
lower part 1.0486 22
upper part 0.95367 22
Region: CARB
single geometric dense at 0.30000E-06
0.86023 95
Region: AUS
double geometric
dense at outer boundaries, coarse at 0.15393E-06
lower part 1.0067 22
upper part 0.99334 22
Region: CARB
single geometric dense at 0.52550E-06
0.85084 96
Region: AUS
double geometric
dense at outer boundaries, coarse at 0.26963E-06
lower part 1.0222 22
upper part 0.97828 22
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
Trying old scheme 4
GENERATING STARTING VALUES FOR CELL # 1 INTERFACE # 2
DETERMINING INITIAL EQUILIBRIUM VALUES
CALCULATING STARTING VALUES: 9 EQUILIBRIUM CALCULATIONS DONE 6 OUT OF 9
89434.0516915
Trying old scheme 4
GENERATING STARTING VALUES FOR CELL # 2 INTERFACE # 2
DETERMINING INITIAL EQUILIBRIUM VALUES
CALCULATING STARTING VALUES: 9 EQUILIBRIUM CALCULATIONS DONE 6 OUT OF 9
89434.0516915
Trying old scheme 4
GENERATING STARTING VALUES FOR CELL # 3 INTERFACE # 2
DETERMINING INITIAL EQUILIBRIUM VALUES
CALCULATING STARTING VALUES: 9 EQUILIBRIUM CALCULATIONS DONE 6 OUT OF 9
89434.0516915
U-FRACTION IN SYSTEM: C = .0406910187346776 CR = .0214382352304608
FE = .978561764900046
TOTAL SIZE OF SYSTEM: 2.90023192349E-17 [m^3]
U-FRACTION IN SYSTEM: C = .0406910187346776 CR = .0214382352304608
FE = .978561764900046
TOTAL SIZE OF SYSTEM: 2.90023192349E-17 [m^3]
0.549881621022649 0.549917064641369 0.549938978099245 0.549921828506974 0.549881463442408 0.549881396770029
002 3.584789101548021E-004 4.707896221377434E-005 1.368538550753866E-006 9.616968061557840E-
006 1.052143983519847E-006 9.010518128506797E-007 8.267746918785806E-007 8.321573841423081E-
007 8.024773010768645E-007 8.036126148622570E-007 7.984608479736118E-007 8.050009080302152E-
007 8.000667876566477E-007 7.984620140534939E-007 7.984782243546979E-007 8.003049929195575E-
007 8.048504050393717E-007 8.013658009019990E-007 7.869048260277564E-007 7.845298608542452E-
007 7.828052493059758E-007 7.79510507725587E-007 7.765420050156961E-007 7.772966895091672E-
007 7.719168100416537E-007 7.729228850419728E-007 7.669031199707825E-007 7.600329286365391E-
007 7.604611933770322E-007 7.600341337033347E-007 7.600503467509922E-007 7.525562577031810E-
007 7.561970194002578E-007 7.540768031241613E-007 7.401555176787141E-007 7.236762721075363E-
007 7.065875829999436E-007 6.785553055152433E-007 6.48867199900529E-007 6.49503027254320E-
007 5.991860190087056E-007 6.000961677486331E-007 5.458302367113732E-007 4.547576398519841E-
007 4.549588570952311E-007 4.547588535329702E-007 4.547750660658187E-007 3.591727447626518E-007

output ignored...

... output resumed

CPU time used in timestep 5 seconds
5.621388737732521E-003 5.621388877210812E-003 5.621388821356869E-003 5.621348236853987E-003 5.621327819904863E-
003 5.621133707426804E-003 5.620509889262913E-003 7.742660461463399E-005 1.344307524693995E-
007 4.461596923762502E-011 3.571139507682395E-015 1.103138996234142E-
018 TIME = 7809.9533 DT = 1000.0000 SUM OF SQUARES = 0.11031390E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.11246535E-10 AND -0.11246535E-10
POSITION OF INTERFACE CARB / AUS IS 0.41766263E-06
CELL # 3 VELOCITY AT INTERFACE # 2 IS -0.12055158E-10 AND -0.12055158E-10
POSITION OF INTERFACE CARB / AUS IS 0.24050830E-06
U-FRACTION IN SYSTEM: C = .0407685766901636 CR = .0215929329214049
FE = .978407067209102
TOTAL SIZE OF SYSTEM: 2.90023192349E-17 [m^3]
2 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUS

CPU time used in timestep 4 seconds
7.934792337300709E-003 7.934792424859177E-003 7.934792358769652E-003 7.934749878634881E-003 7.934730557860125E-
003 7.934481903096044E-003 7.933720034973661E-003 1.064665425785131E-004 1.753609738866775E-
007 7.573759727881984E-011 1.006687963004298E-014 2.678574071231658E-
018 TIME = 8809.9533 DT = 1000.0000 SUM OF SQUARES = 0.26785741E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.10003883E-10 AND -0.10003883E-10
POSITION OF INTERFACE CARB / AUS IS 0.40765875E-06
CELL # 3 VELOCITY AT INTERFACE # 2 IS -0.11461547E-10 AND -0.11461547E-10
POSITION OF INTERFACE CARB / AUS IS 0.22904676E-06
U-FRACTION IN SYSTEM: C = .0407686263795191 CR = .021592733030047
FE = .97840726710046
TOTAL SIZE OF SYSTEM: 2.90023192349E-17 [m^3]
CPU time used in timestep 5 seconds
1.006183996174871E-002 1.006184002234698E-002 1.006183995420535E-002 1.006180104690770E-002 1.006178556514470E-
002 1.006148495142356E-002 1.006060282846608E-002 1.098035078318554E-004 2.176782535198634E-
007 9.073739073247344E-011 1.570179997998957E-014 3.489793724572546E-

```

018      TIME =  9809.9533    DT =  1000.0000    SUM OF SQUARES =  0.34897937E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.90228135E-11 AND -0.90228135E-11
POSITION OF INTERFACE CARB / AUS IS 0.39863594E-06
CELL # 3 VELOCITY AT INTERFACE # 2 IS -0.11230210E-10 AND -0.11230210E-10
POSITION OF INTERFACE CARB / AUS IS 0.21781655E-06
U-FRACTION IN SYSTEM: C = .0407686584004964 CR = .021592589420798
FE = .978407410709709
TOTAL SIZE OF SYSTEM: 2.90023192349E-17 [m^3]
CPU time used in timestep        4 seconds
1.781790411865416E-003      1.781790525604021E-003      1.781790643567297E-003      1.781728372351846E-003      1.781703673622629E-
003      1.781714254977339E-003      1.781674965403099E-003      4.687254999591863E-004      1.276511488825608E-
005      1.299830221207340E-008      2.034514363601619E-012      1.355975732407151E-
016      TIME = 10000.000      DT = 190.04670      SUM OF SQUARES = 0.13559757E-15
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.74898493E-11 AND -0.74898493E-11
POSITION OF INTERFACE CARB / AUS IS 0.39721251E-06
CELL # 3 VELOCITY AT INTERFACE # 2 IS -0.86365802E-11 AND -0.86365802E-11
POSITION OF INTERFACE CARB / AUS IS 0.21617519E-06
U-FRACTION IN SYSTEM: C = .0407686787456533 CR = .0215925564027491
FE = .978407443727758
TOTAL SIZE OF SYSTEM: 2.90023192349E-17 [m^3]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME      3161.2791
DELETING TIME-RECORD FOR TIME      3467.7760
DELETING TIME-RECORD FOR TIME      3467.7760
DELETING TIME-RECORD FOR TIME      3467.7761
DELETING TIME-RECORD FOR TIME      3467.7761
DELETING TIME-RECORD FOR TIME      3467.7762
DELETING TIME-RECORD FOR TIME      3467.7763
DELETING TIME-RECORD FOR TIME      3467.7767
DELETING TIME-RECORD FOR TIME      3467.7773
DELETING TIME-RECORD FOR TIME      3467.7786
DELETING TIME-RECORD FOR TIME      3467.7811
DELETING TIME-RECORD FOR TIME      3467.7863
DELETING TIME-RECORD FOR TIME      3467.7965
DELETING TIME-RECORD FOR TIME      3467.8170
DELETING TIME-RECORD FOR TIME      3467.8579
DELETING TIME-RECORD FOR TIME      3467.9399
DELETING TIME-RECORD FOR TIME      3468.1037
DELETING TIME-RECORD FOR TIME      3468.4314
DELETING TIME-RECORD FOR TIME      3469.0867
DELETING TIME-RECORD FOR TIME      3470.3975
DELETING TIME-RECORD FOR TIME      3473.0189
DELETING TIME-RECORD FOR TIME      3478.2618
DELETING TIME-RECORD FOR TIME      3488.7475
DELETING TIME-RECORD FOR TIME      3509.7191
DELETING TIME-RECORD FOR TIME      3551.6621
DELETING TIME-RECORD FOR TIME      3635.5482
DELETING TIME-RECORD FOR TIME      3803.3203
DELETING TIME-RECORD FOR TIME      4138.8647
DELETING TIME-RECORD FOR TIME      4809.9533
DELETING TIME-RECORD FOR TIME      5809.9533
DELETING TIME-RECORD FOR TIME      6809.9533
DELETING TIME-RECORD FOR TIME      7809.9533
DELETING TIME-RECORD FOR TIME      8809.9533

KEEPING TIME-RECORD FOR TIME      9809.9533
AND FOR TIME                      10000.000
WORKSPACE RECLAIMED

Timestep at 10000.0000      Selected

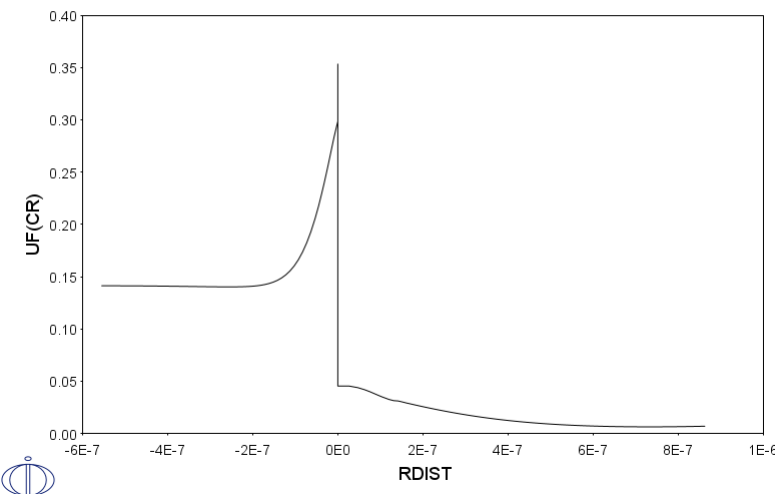
DIC>
DIC>
DIC>
DIC>
DIC>
DIC> set-inter
--OK--
DIC>
```

exc2-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exc2\plot.DCM"
DIC>
DIC>
DIC> @@ exc2_plot.DCM
DIC>
DIC> @@
DIC> FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE c2
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 1.00000E+04
DIC> read exc2
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ PLOT THE CHROMIUM CONCENTRATION PROFILES IN THE SAME WAY AS IN exb2
POST-1: @@ BUT NOW FOR EACH PARTICLE. LET US LOOK AT THE PROFILES AFTER 1000s.
POST-1: @@
POST-1: @@
POST-1: @@ FIRST CELL
POST-1: @@
POST-1: enter-symb
Function or table /FUNCTION/: func
NAME: rdist
FUNCTION: gd-poi(carb,u);
POST-1:
POST-1: s-d-a x rdist
POST-1:
POST-1: s-d-a y uf(cr)
POST-1:
POST-1: s-i-v
VARIABLE /TIME/: dist
DISTANCE : /GLOBAL/: glo
POST-1:
POST-1: s-p-c time 1000
POST-1:
POST-1: @@
POST-1: @@ SET THE TITLE ON THE PLOT
POST-1: @@
POST-1: set-title Figure C2.1
POST-1: plot
```

Figure C2.1

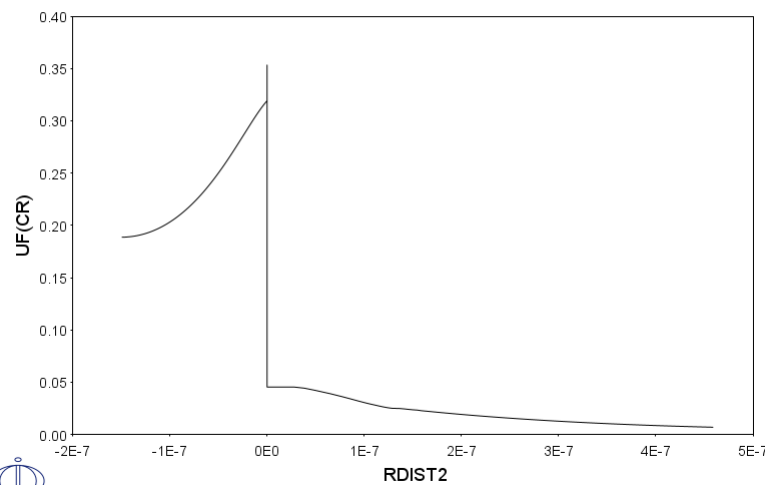
2021.05.13.12.29.42
Time = 1000
CELL #1



```
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: @@
POST-1: @@ SELECT CELL 2
POST-1: @@
POST-1: select-cell
Number /NEXT/: 2
CELL 2 SELECTED
POST-2:
POST-2: enter-symb
Function or table /FUNCTION/: func
NAME: rdist2
FUNCTION: gd-poi(carb,u);
POST-2:
POST-2: s-d-a x rdist2
POST-2:
POST-2: set-title Figure C2.2
POST-2: plot
```

Figure C2.2

2021.05.13.12.29.46
Time = 1000
CELL #2



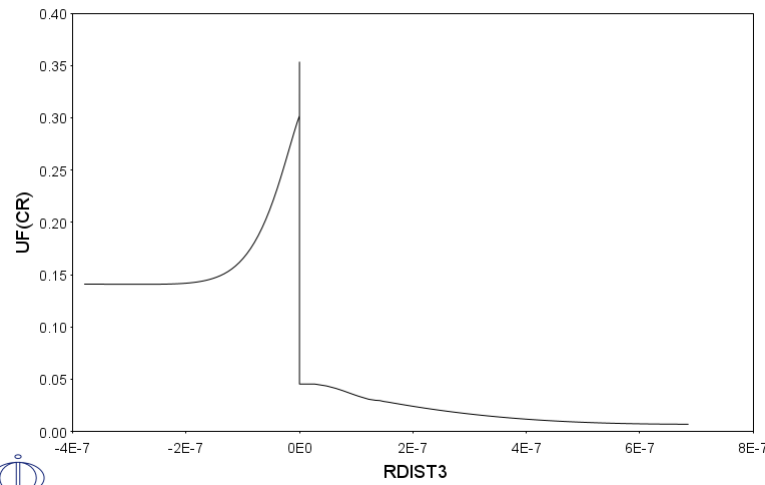
```

POST-2:
POST-2:
POST-2:
POST-2:@?<_hit_return_to_continue_>
POST-2:
POST-2: @@
POST-2: @@ SELECT CELL 3
POST-2: @@
POST-2: select-cell 3
CELL 3 SELECTED
POST-3:
POST-3: enter-symb
Function or table /FUNCTION/: func
NAME: rdist3
FUNCTION: gd-poi(carb,u);
POST-3:
POST-3: s-d-a x rdist3
POST-3:
POST-3: set-title Figure C2.3
POST-3: plot

```

Figure C2.3

2021.05.13.12.29.50
Time = 1000
CELL #3



```

POST-3:
POST-3:
POST-3:
POST-3:@?<_hit_return_to_continue_>
POST-3:
POST-3:
POST-3: @@
POST-3: @@ ALSO PLOT HOW THE DIAMETER OF THE CEMENTITE PARTICLE VARIES
POST-3: @@ WITH TIME IN THE THREE CELLS
POST-3: @@
POST-3:
POST-3: @@
POST-3: @@ SELECT THE FIRST CELL
POST-3: @@
POST-3: sel-cell 1
CELL 1 SELECTED
POST-1:
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-s-s x n .01 10000
POST-1: set-axis-type x log
POST-1:
POST-1: enter func diam1=2*poi(carb,u);
POST-1: s-d-a y diam1
POST-1: s-s-s y n 0 1.5e-6
POST-1:

```

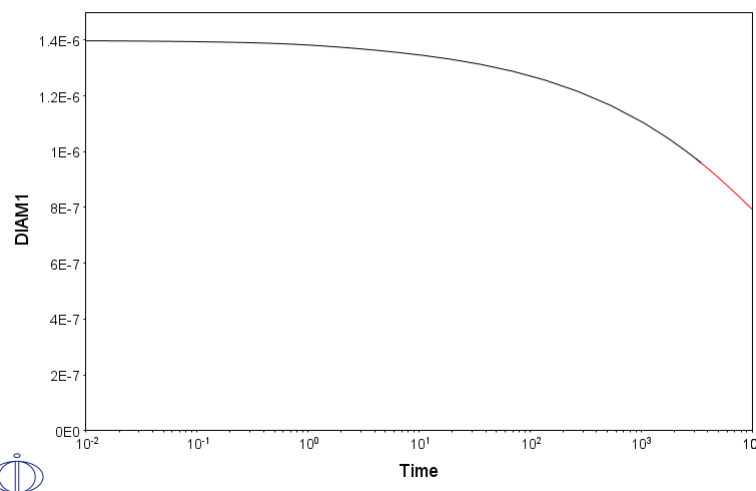
```

POST-1: s-p-c interf carb upp
POST-1:
POST-1: app n
POST-1: set-title Figure C2.4
POST-1: plot

```

Figure C2.4

2021.05.13.12.29.54
UPPER INTERFACE OF REGION "CARB#1"
CELL #1



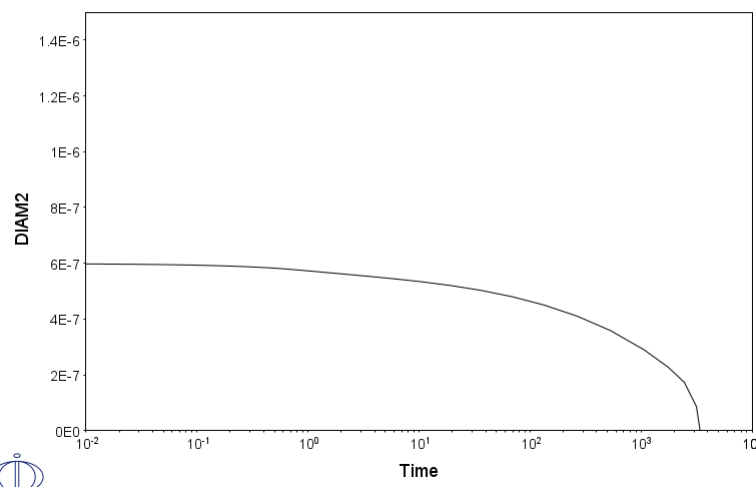
```

POST-1:
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1:
POST-1: @@
POST-1: @@ SELECT CELL 2
POST-1: @@
POST-1: sel-cell 2
CELL 2 SELECTED
POST-2:
POST-2: enter func diam2=2*poi(carb,u);
POST-2: s-d-a y diam2
POST-2: s-s-s y n 0 1.5e-6
POST-2:
POST-2: s-p-c interf carb upp
POST-2:
POST-2: set-title Figure C2.5
POST-2: plot

```

Figure C2.5

2021.05.13.12.29.54
UPPER INTERFACE OF REGION "CARB#2"
CELL #2



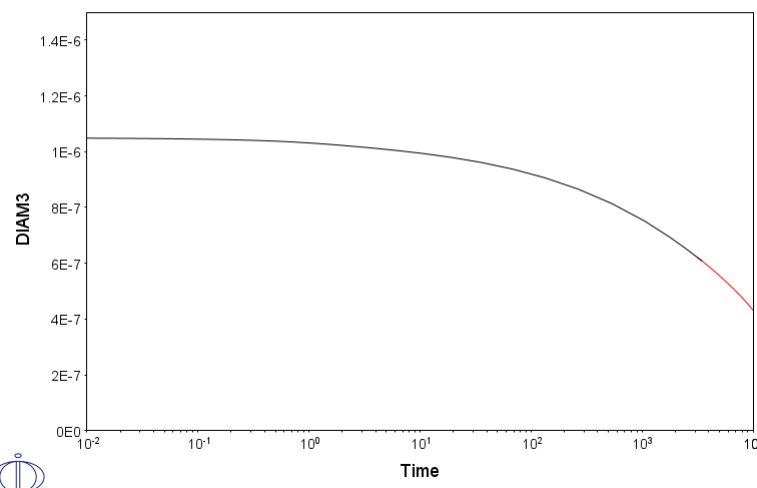
```

POST-2:
POST-2:
POST-2:
POST-2:
POST-2:@?<_hit_return_to_continue_>
POST-2:
POST-2: @@
POST-2: @@ SELECT CELL 3
POST-2: @@
POST-2: sel-cell 3
CELL 3 SELECTED
POST-3:
POST-3: enter func diam3=2*poi(carb,u);
POST-3: s-d-a y diam3
POST-3: s-s-s y n 0 1.5e-6
POST-3:
POST-3: s-p-c interf carb upp
POST-3:
POST-3: set-title Figure C2.6
POST-3: plot

```

Figure C2.6

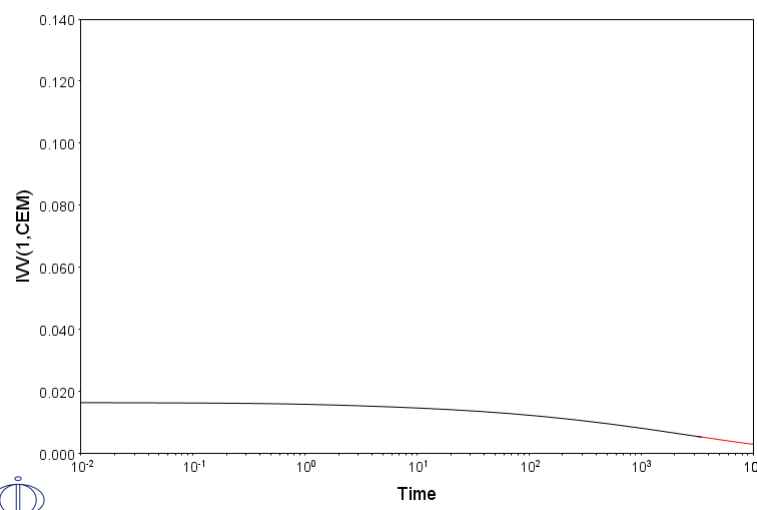
2021.05.13.12.30.01
UPPER INTERFACE OF REGION "CARB#3"
CELL #3



```
POST-3:  
POST-3:  
POST-3:  
POST-3:  
POST-3:@?<_hit_return_to_continue_>  
POST-3:  
POST-3: @@  
POST-3: @@ NOW PLOT THE VOLUME FRACTION OF CEMENTITE IN THE THREE CELLS  
POST-3: @@  
POST-3: s-d-a x time  
INFO: Time is set as independent variable  
POST-3: s-s-s x n .01 10000  
POST-3: set-axis-type x log  
POST-3:  
POST-3: @@  
POST-3: @@ CELL 1  
POST-3: @@  
POST-3: s-d-a y ivv(1,cem)  
POST-3: s-s-s y n 0 0.14  
POST-3:  
POST-3: s-p-c integral  
POST-3:  
POST-3: set-title Figure C2.7  
POST-3: plot
```

Figure C2.7

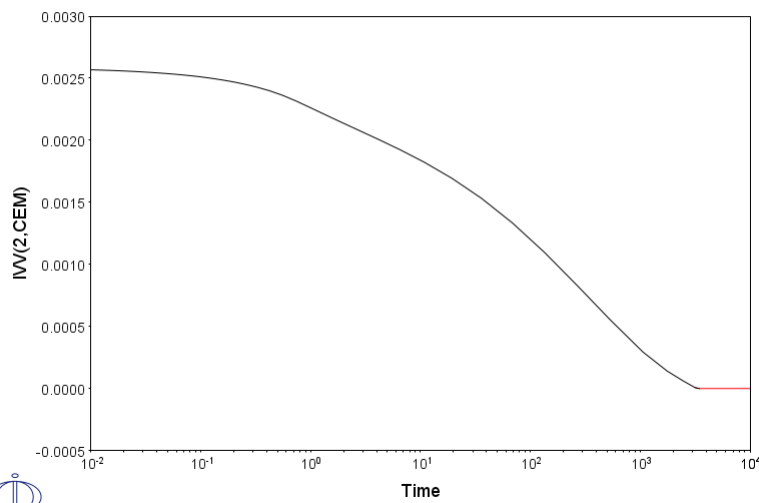
2021.05.13.12.30.06
CELL #3



```
POST-3:  
POST-3:  
POST-3:  
POST-3:  
POST-3:@?<_hit_return_to_continue_>  
POST-3:  
POST-3: @@  
POST-3: @@ CELL 2  
POST-3: @@  
POST-3: s-d-a y ivv(2,cem)  
POST-3:  
POST-3: set-title Figure C2.8  
POST-3: plot
```

Figure C2.8

2021.05.13.12.30.13
CELL #3



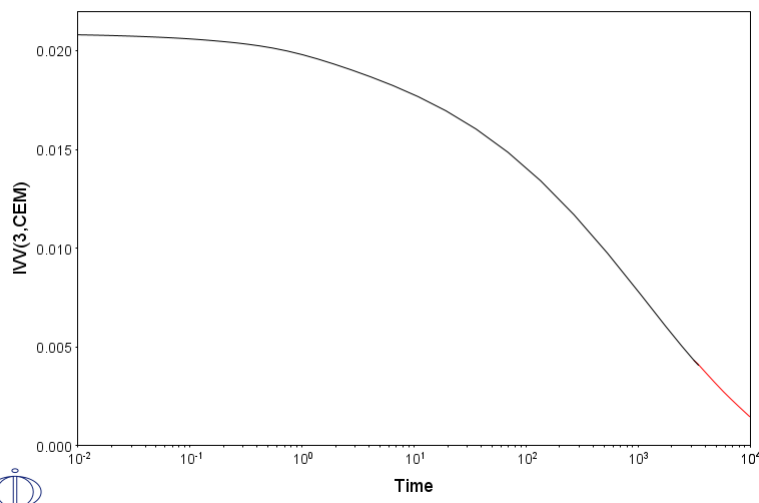
```

POST-3:
POST-3:
POST-3:
POST-3:
POST-3:@?<_hit_return_to_continue_>
POST-3:
POST-3: @@
POST-3: @@ CELL 3
POST-3: @@
POST-3: s-d-a y ivv(3,cem)
POST-3:
POST-3: set-title Figure C2.9
POST-3: plot

```

Figure C2.9

2021.05.13.12.30.21
CELL #3



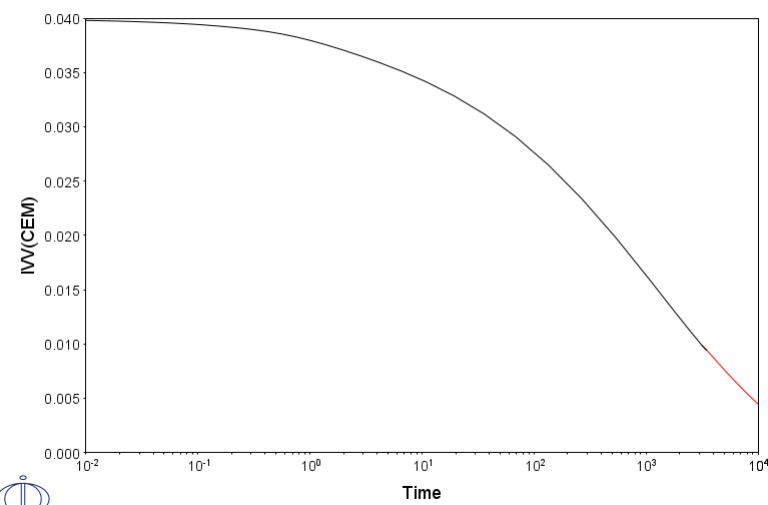
```

POST-3:
POST-3:
POST-3:
POST-3:
POST-3:@?<_hit_return_to_continue_>
POST-3:
POST-3: @@
POST-3: @@ FINALLY, PLOT HOW THE TOTAL VOLUME FRACTION OF CEMENTITE
POST-3: @@ VARIES WITH TIME.
POST-3: @@
POST-3: s-d-a y ivv(cem)
POST-3:
POST-3: set-title Figure C2.10
POST-3: plot

```

Figure C2.10

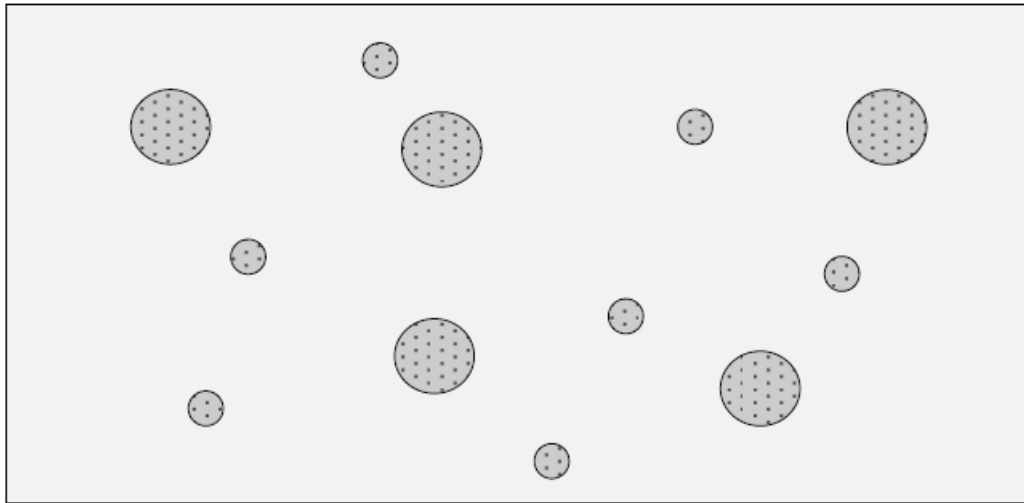
2021.05.13.12.30.29
CELL #3



POST-3:
POST-3:
POST-3:
POST-3:
POST-3:@?<_hit_return_to_continue_>
POST-3:
POST-3: set-inter
--OK---
POST-3:



Diffusion in Dispersed Systems



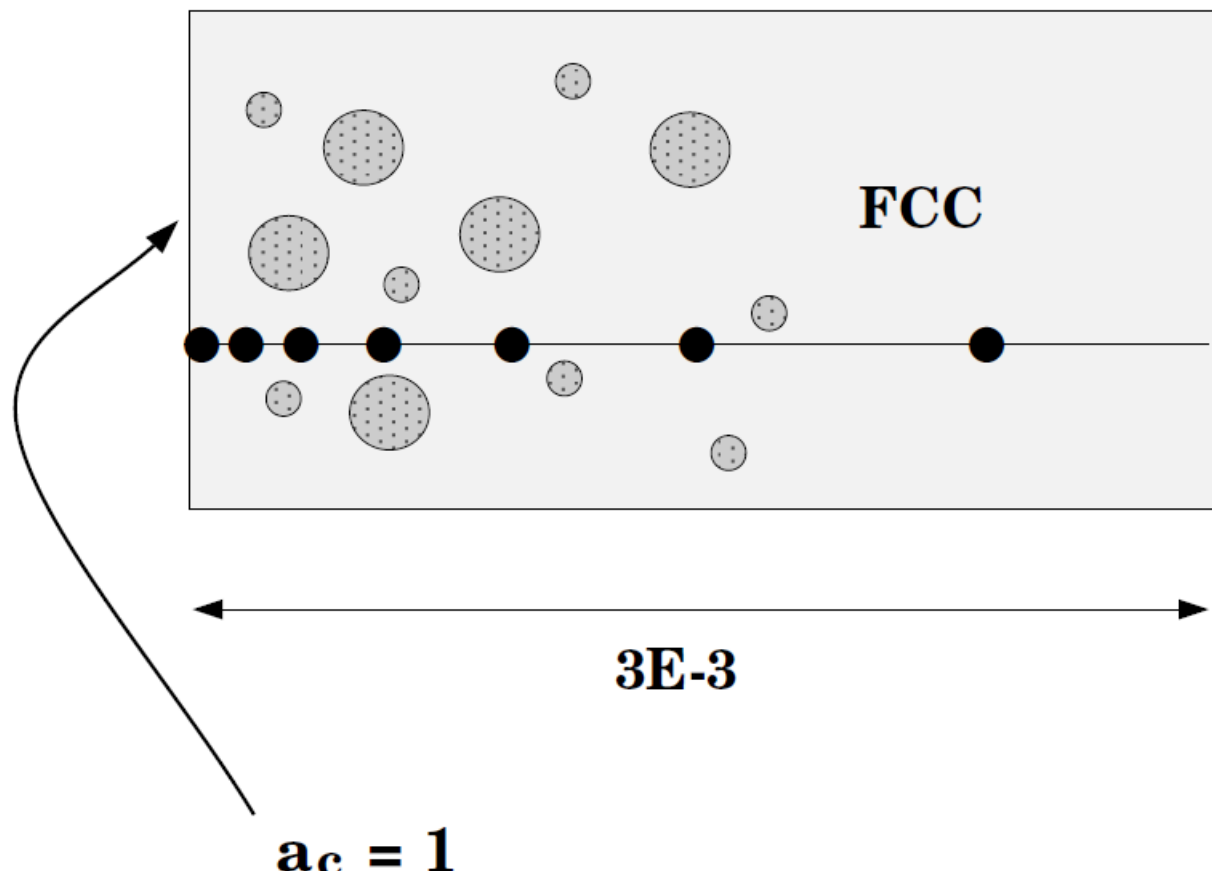


Example exd1a

Carburization of a Ni-25% Cr alloy: Dispersed system model

This example is about carburization of a Ni-25Cr alloy. In this case the M₃C₂ and M₇C₃ carbides are entered as spheroid phases in a FCC matrix. In this example the DISPERSED SYSTEM MODEL is used. This case is from A. Engström, L. Höglund and J. Ågren: Metall.Trans.A v. 25A (1994), pp. 1127-1134.

$$T = 1123 \text{ K}$$



exdla-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

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Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exdla\setup.DCM"
SYS: @@
SYS: @@ Diffusion in dispersed systems.
SYS: @@ Carburization of Ni-25%Cr alloy: Dispersed system model
SYS: @@ This example is about carburization of a Ni-25Cr alloy.
SYS: @@ In this case the M3C2 and M7C3 carbides are entered as
SYS: @@ spheroid phases in a FCC matrix. This simulation can be run
SYS: @@ with either the DISPERSED SYSTEM MODEL or the HOMOGENIZATION MODEL.
SYS: @@ In this example the DISPERSED SYSTEM MODEL is used, which requires
SYS: @@ that the default HOMOGENIZATION MODEL is disabled.
SYS: @@ With the DISPERSED SYSTEM MODEL the command
SYS: @@ ENTER_LABYRINTH_FUNCTION is used to take into account the
SYS: @@ impeding effect of dispersed phases on long-range diffusion.
SYS: @@ For the HOMOGENIZATION MODEL the command
SYS: @@ ENTER_HOMOGENIZATION_FUNCTION should be used.
SYS: @@ This case is from A. Engström, L. Häglund and J. Ågren:
SYS: @@ Metall.Trans.A v. 25A (1994), pp. 1127-1134.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exd1_setup.DCM
SYS:
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ USE THE SSOL DATABASE FOR THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw fedemo
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-sys ni cr c
NI CR C
 DEFINED
TDB_FEDEMO: rej ph * all
GAS:G LIQUID:L BCC_A2
LAVES_PHASE_C14 CBCC_A12 CEMENTITE
CHI_A12 CUB_A13 DIAMOND_FCC_A4
FCC_A1 GRAPHITE HCP_A3
KSI_CARBIDE M23C6 M3C2
M7C3 SIGMA REJECTED
TDB_FEDEMO: res ph fcc,m7c3,m3c2,grap
FCC_A1 M7C3 M3C2
GRAPHITE RESTORED
TDB_FEDEMO: get
12:31:49,473 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES
ELEMENTS
SPECIES
PHASES
Creating a new composition set FCC_A1#2
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'J. Bratberg, Z. Metallkd., 96 (2005) 335-344; Fe-Cr-Mo-C'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'A. Gabriel, C. Chatillon, and I. Ansara, published in High Temp. Sci.
(Parameters listed in CALPHAD, 11 (1987) 203-218); C-Ni'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
'R. Naraghi, Thermo-Calc Software AB, Sweden, 2016; FCC Fe-Cr-C and C-Cr-Ni'
'NPL, unpublished work (1989); C-Cr-Ni'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); CR -NI'
'B. Uhrenius, Int. J. Refract. Met. Hard Mater. 12 (1994) 121 -127; Molar
volumes'
'P. Villars, L.D. Calvert. Pearson's handbook of crystallographic data for
intermetallic phases, Metals park, Ohio 1985: American society for
metals'
'A.F. Guillermet and G. Grimvall, J. Phys. Chem. Solids, 1992, 53, 105-125;
Molar volumes'
-OK-
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app mfdemo
Current database: Fe-Alloys Mobility demo database v2.0

APP: DEFINED
APP: def-sys ni c cr
NI CR
 DEFINED

```

APP: rej ph * all
      FCC A2           FCC A1  REJECTED
APP: res ph fcc,m7c3,m3c2,grap
*** ERROR M7C3 INPUT IGNORED
*** ERROR M3C2 INPUT IGNORED
*** ERROR GRAP INPUT IGNORED
FCC_A1 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
Creating a new composition set FCC_A1#3
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: Z. Metallkunde 85(1994)502-509; C diffusion in fcc Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Ni diffusion fcc Cr-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'

-OK-
APP:
APP: @@ 
APP: @@ ENTER THE DICTRA MONITOR
APP: @@ 
APP: go d-m
NO TIME STEP DEFINED
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
*** ENTERING M3C2 AS A DIFFUSION NONE PHASE
*** ENTERING M7C3 AS A DIFFUSION NONE PHASE
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 1123; * N
DIC>
DIC> @@
DIC> @@ SET THE REFERENCE STATE FOR CARBON
DIC> @@
DIC> set-reference-state
Component: C
Reference state: grap
Temperature /*/: *
Pressure /100000/: 101325
DIC>
DIC> @@
DIC> @@ ENTER THE REGION aus
DIC> @@
DIC> enter-region aus
DIC>
DIC> @@
DIC> @@ ENTER A GEOMETRICAL GRID INTO THE REGION
DIC> @@
DIC> enter-grid aus 3e-3 geo 100 1.02
DIC>
DIC> @@
DIC> @@ ENTER A MATRIX PHASE IN THE REGION
DIC> @@
DIC> enter-phase act aus matrix fcc_a1#1
DIC>
DIC> @@
DIC> @@ ENTER THE START COMPOSITION FOR THE MATRIX PHASE
DIC> @@
DIC> enter-composition
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1#1/: fcc#1
DEPENDENT COMPONENT ? /NI/: ni
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: cr
TYPE /LINEAR/: lin 25 25
PROFILE FOR /CR/: c
TYPE /LINEAR/: lin 1e-4 1e-4
DIC>
DIC> @@
DIC> @@ ENTER SPHEROIDAL PHASES IN THE REGION
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUS/: aus
PHASE TYPE /MATRIX/: sph
PHASE NAME: /NONE/: m7c3
INFO: EQUILIBRIUM COMPOSITION AND FRACTION OF SPHEROID PHASES USED AS DEFAULT
DIC>
DIC> @@
DIC> @@ ENTER A STOICHIOMETRIC SPHEROIDAL PHASE IN THE REGION
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUS/: aus
PHASE TYPE /MATRIX/: sph
PHASE NAME: /NONE/: m3c2
INFO: EQUILIBRIUM COMPOSITION AND FRACTION OF SPHEROID PHASES USED AS DEFAULT
DIC>
DIC> @@
DIC> @@ ENTER A START COMPOSITION FOR THE SPHEROIDAL PHASES
DIC> @@
DIC> enter-composition
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1#1/: m7c3
USE EQUILIBRIUM VALUE /Y/: Y
DIC> enter-composition
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1#1/: m3c2
USE EQUILIBRIUM VALUE /Y/: Y
DIC>
DIC> @@
DIC> @@ SET THE BOUNDARY CONDITION
DIC> @@

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DIC> set-cond
GLOBAL OR BOUNDARY CONDITION /GLOBAL/: boundary
BOUNDARY /LOWER/: lower
CONDITION TYPE /CLOSED_SYSTEM/: mixed
Dependent substitutional element:NI
Dependent interstitial element:VA
TYPE OF CONDITION FOR COMPONENT C /ZERO_FLUX/: activity
LOW TIME LIMIT /0/: 0
ACR(C) (TIME)= 1;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
TYPE OF CONDITION FOR COMPONENT CR /ZERO_FLUX/: zero-flux
DIC>
DIC> @@
DIC> @@ ENTER THE LABYRINTH FACTOR
DIC> @@
DIC> enter-lab
REGION NAME : aus
f(T,P,VOLFR,X)= volfr**2;
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME. REMEMBER TO BE CAREFUL WITH THE
DIC> @@ Timestep WHEN THERE ARE SPHEROIDAL PHASES PRESENT. IN THIS CASE
DIC> @@ THE Timestep IS NOT ALLOWED TO BE LARGER THAN 1800s.
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 3600000
AUTOMATIC Timestep CONTROL /YES/: YES
MAX Timestep DURING INTEGRATION /360000/: 1800
INITIAL Timestep : /1E-07/:
SMALLEST ACCEPTABLE Timestep : /1E-07/:
DIC>
DIC> @@
DIC> @@ IN ORDER SO SAVE SOME SPACE ON THE DISK THE RESULT IS STORED
DIC> @@ SELECTIVELY. OTHERWISE THE STORE-RESULT-FILE FROM THIS EXAMPLE
DIC> @@ WOULD BE VERY LARGE.
DIC> @@
DIC> set-simulation-condition
NS01A PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA Timesteps IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/:
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/: 99
DEGREE OF IMPLICITY WHEN INTEGRATING PDES (AUTO, 0 -> 0.5 -> 1): /AUTO/:
MAX Timestep CHANGE PER Timestep : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/:
DIC>
DIC> @@ BY DEFAULT THE "HOMOGENIZATION MODEL" IS USED WHEN MULTIPLE PHASES
DIC> @@ ARE ENTERED IN A SINGLE REGION. FOR THIS EXAMPLE THE HOMOGENIZATION
DIC> @@ MODEL IS DISABLED.
DIC> ho n
HOMOGENIZATION DISABLED
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exd1 y
DIC>
DIC> set-inter
---OK---
DIC>

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exdla-run

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DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exdla\run.DCM"
DIC>
DIC>
DIC> @@ exd1_run.DCM
DIC>
DIC> @@
DIC> @@ READ THE SETUP FILE AND START THE SIMULATION
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
*** ENTERING M3C2 AS A DIFFUSION NONE PHASE
*** ENTERING M7C3 AS A DIFFUSION NONE PHASE
DIC> read exd1
OK
DIC> sim
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: C = 4.73399450059566E-06 CR = .273386452547573
NI = .726613547452427
TOTAL SIZE OF SYSTEM: .003 [m]
WARNING:M7C3 HAS NO VOLUME FRACTION, CREATING ONE
WARNING:M3C2 HAS NO VOLUME FRACTION, CREATING ONE
U-FRACTION IN SYSTEM: C = 4.73399450059566E-06 CR = .273386452547573
NI = .726613547452427
TOTAL SIZE OF SYSTEM: .003 [m]
TIME = 0.10000000E-06 DT = 0.10000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 8.32233829951615E-05 CR = .273386452547572
NI = .726613547452428
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.10010000E-03 DT = 0.10000000E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 1.14491977930612E-04 CR = .273386452547544
NI = .726613547452456
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 0.40010010 DT = 0.40000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 1.3439753194935E-04 CR = .273386452541228
NI = .726613547458772
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 1 seconds
TIME = 61.226617 DT = 60.826517 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 1.7323419086632E-04 CR = .273386451404806
NI = .726613548595194
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 136.73132 DT = 75.504702 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 2.15195361346137E-04 CR = .273386450378639
NI = .726613549621361
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 246.77986 DT = 110.04854 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 2.65132773423409E-04 CR = .273386449367934
NI = .726613550632066
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 417.73078 DT = 170.95092 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 3.26703519714746E-04 CR = .273386448427441
NI = .72661355157256
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 700.87762 DT = 283.14684 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 4.03852183677253E-04 CR = .273386447458974
NI = .726613552541025
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 1 seconds
TIME = 1196.7621 DT = 495.88443 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 5.03958315340991E-04 CR = .273386446322885
NI = .726613553677115
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 2136.4660 DT = 939.70399 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 6.43012086810231E-04 CR = .273386444988437
NI = .726613555011564
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3936.4660 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = 8.3851974916308E-04 CR = .273386443511134
NI = .726613556488866
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 5736.4660 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0010173697033215 CR = .273386442681224
NI = .726613557318776
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 7536.4660 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00118296902880158 CR = .273386442066142
NI = .726613557933858
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 1 seconds
TIME = 9336.4660 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00133879540957736 CR = .273386441011293
NI = .726613558988707
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 11136.466 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00148538430057559 CR = .273386439279029
NI = .726613560720971
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 12936.466 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00162329996141555 CR = .273386436825667
NI = .726613563174333
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 14736.466 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00175666371078121 CR = .273386434544898
NI = .726613565455102
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
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TIME = 16509.560 DT = 1773.0939 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00188419698676039 CR = .273386432317815
NI = .726613567682185
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 1 seconds
TIME = 18202.529 DT = 1692.9692 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00200374370728573 CR = .273386430131715
NI = .726613569868284
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 19871.496 DT = 1668.9668 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00212043403493914 CR = .273386427985689
NI = .726613572014311
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 21585.287 DT = 1713.7913 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00223651041851037 CR = .273386426227075
NI = .726613573772925
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 23385.287 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00235302551823152 CR = .273386424970864

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output ignored...

... output resumed

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CPU time used in timestep 1 seconds
TIME = 3563985.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0314537983526267 CR = .273386325210631
NI = .72661367478937
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3565785.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0314617329030149 CR = .273386325190173
NI = .726613674809827
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3567585.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0314696650642582 CR = .273386325169499
NI = .726613674830501
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3569385.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0314775948480468 CR = .273386325148607
NI = .726613674851393
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3571185.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0314855222664731 CR = .273386325127498
NI = .726613674872502
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3572985.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0314934473320116 CR = .273386325106171
NI = .726613674893829
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3574785.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315013700574985 CR = .273386325084626
NI = .726613674915374
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 1 seconds
TIME = 3576585.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315092904561121 CR = .273386325062863
NI = .726613674937137
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3578385.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315172085413539 CR = .273386325040882
NI = .726613674959118
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3580185.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315251243270301 CR = .273386325018683
NI = .726613674981317
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3581985.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315330378272334 CR = .273386324996265
NI = .726613675003735
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3583785.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315409490563253 CR = .273386324973628
NI = .726613675026371
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3585585.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315488580289196 CR = .273386324950773
NI = .726613675049227
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3587385.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315567647598653 CR = .273386324927699
NI = .726613675072302
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 1 seconds
TIME = 3589185.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .031564669264231 CR = .273386324904405
NI = .726613675095595
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3590985.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315725715572893 CR = .273386324880892
NI = .726613675119108
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3592785.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315804716545019 CR = .273386324857159
NI = .726613675142841
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds

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TIME = 3594585.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0315883695715048 CR = .273386324833206
NI = .726613675166794
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3596385.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .031596265324095 CR = .273386324809034
NI = .726613675190966
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3598185.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0316041589282161 CR = .273386324784641
NI = .726613675215359
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3599985.3 DT = 1800.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0316120503999463 CR = .273386324760028
NI = .726613675239973
TOTAL SIZE OF SYSTEM: .003 [m]
CPU time used in timestep 0 seconds
TIME = 3600000.0 DT = 14.712794 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0316121157454989 CR = .273386324759816
NI = .726613675240184
TOTAL SIZE OF SYSTEM: .003 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.1000000E-06
DELETING TIME-RECORD FOR TIME 0.1001000E-03
DELETING TIME-RECORD FOR TIME 163785.29
DELETING TIME-RECORD FOR TIME 341985.29
DELETING TIME-RECORD FOR TIME 520185.29
DELETING TIME-RECORD FOR TIME 698385.29
DELETING TIME-RECORD FOR TIME 876585.29
DELETING TIME-RECORD FOR TIME 1054785.3
DELETING TIME-RECORD FOR TIME 1232985.3
DELETING TIME-RECORD FOR TIME 1411185.3
DELETING TIME-RECORD FOR TIME 1589385.3
DELETING TIME-RECORD FOR TIME 1767585.3
DELETING TIME-RECORD FOR TIME 1945785.3
DELETING TIME-RECORD FOR TIME 2123985.3
DELETING TIME-RECORD FOR TIME 2302185.3
DELETING TIME-RECORD FOR TIME 2480385.3
DELETING TIME-RECORD FOR TIME 2658585.3
DELETING TIME-RECORD FOR TIME 2836785.3
DELETING TIME-RECORD FOR TIME 3014985.3
DELETING TIME-RECORD FOR TIME 3193185.3
DELETING TIME-RECORD FOR TIME 3371385.3
DELETING TIME-RECORD FOR TIME 3549585.3
DELETING TIME-RECORD FOR TIME 3596385.3
DELETING TIME-RECORD FOR TIME 3598185.3

KEEPING TIME-RECORD FOR TIME 3599985.3
AND FOR TIME 3600000.0
WORKSPACE RECLAIMED
```

```

TIMESTEP AT 3600000.00 SELECTED

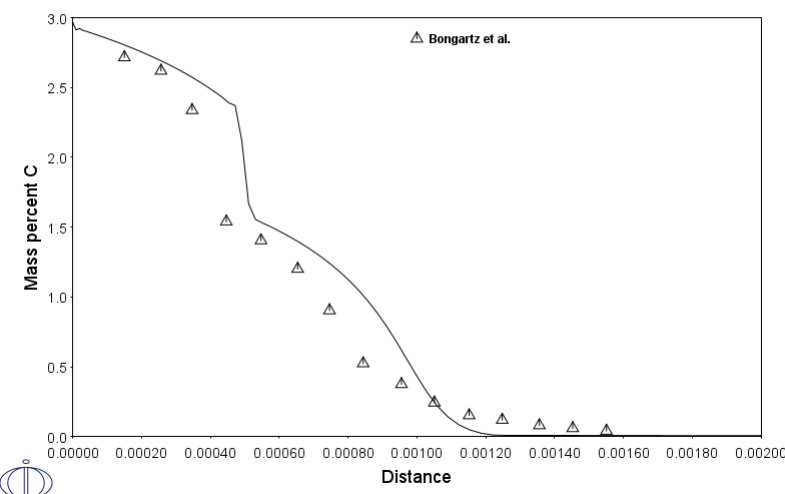
DIC>
DIC> set-inter
--OK---
DIC>
```

exd1a-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd1a\plot.DCM"
DIC>
DIC>
DIC> @@ exd1_plot.DCM
DIC>
DIC> @@
DIC> @@ FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE exd1
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 3.60000E+06
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
*** ENTERING M3C2 AS A DIFFUSION NONE PHASE
*** ENTERING M7C3 AS A DIFFUSION NONE PHASE
DIC> read exd1
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ LOT THE TOTAL CARBON CONCENTRATION PROFILE
POST-1: @@
POST-1: s-d-a y w-p c
POST-1: s-d-a x distance global
INFO: Distance is set as independent variable
POST-1: s-s-s x n 0 2e-3
POST-1: s-p-c time 3600000
POST-1:
POST-1: app y exd1.exp
PROLOGUE NUMBER: /0/: 1
DATASET NUMBER(s): /-1/: 1
POST-1:
POST-1: @@
POST-1: @@ SET THE TITLE ON THE PLOT
POST-1: @@
POST-1: set-tit d1.1
POST-1:
POST-1: plot
```

d1.1

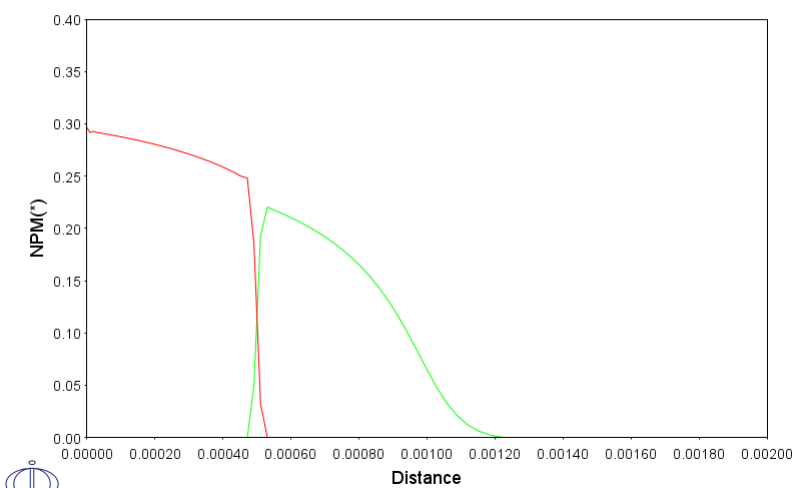
2021.05.13.12.39.27
Time = 3600000
CELL #1



```
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: @@
POST-1: @@ NOW PLOT THE AMOUNT OF CARBIDES FORMED
POST-1: @@
POST-1: s-d-a y npm(*)
POST-1: s-s-s y n 0 0.4
POST-1: app n
POST-1:
POST-1: set-tit d1.2
POST-1: plot
```

d1.2

2021.05.13.12.39.28
Time = 3600000
CELL #1



```
POST-1:  
POST-1:  
POST-1:  
POST-1:@?<_hit_return_to_continue_>  
POST-1:  
POST-1: set-inter  
---OK---  
POST-1:
```

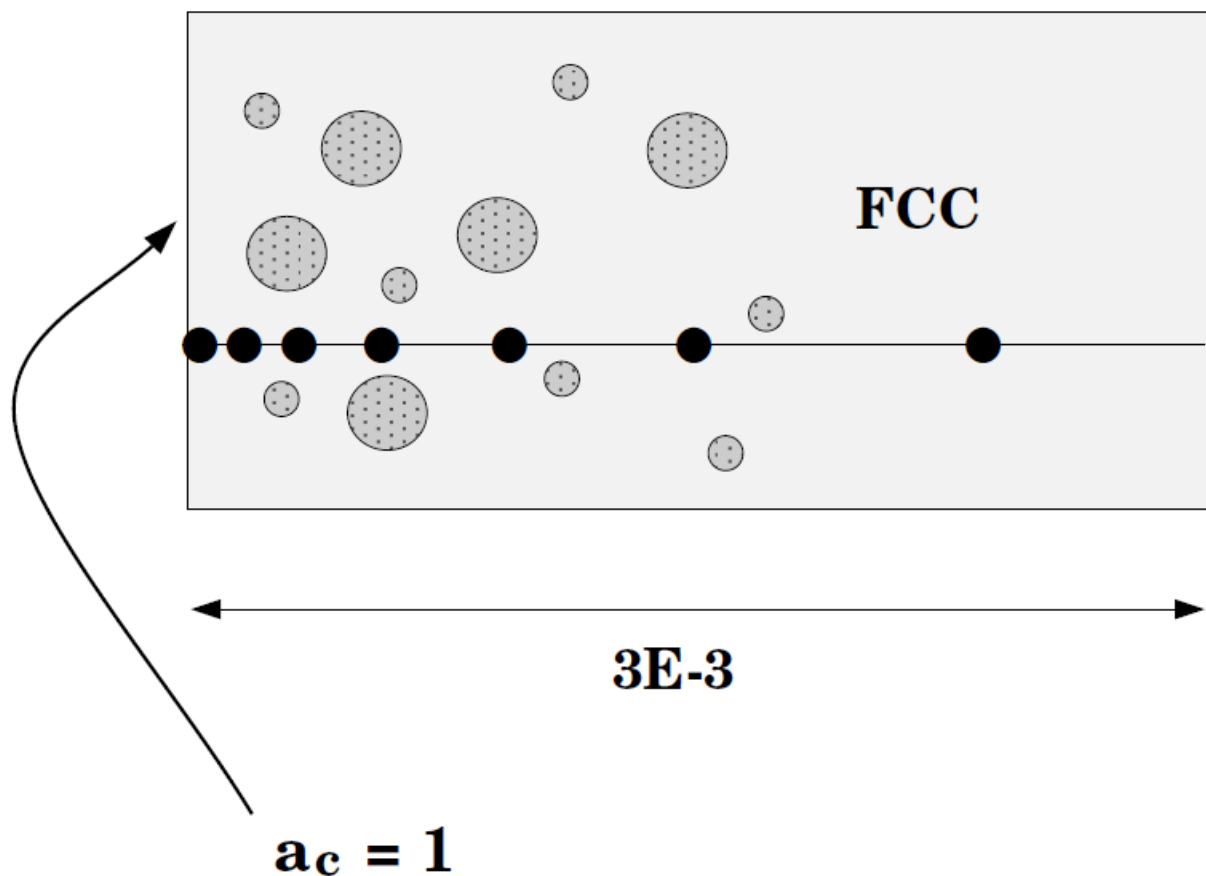


Example exd1b

Carburization of a Ni-25% Cr alloy: Homogenization model

This example is about carburization of a Ni-25Cr alloy. In this case the M₃C₂ and M₇C₃ carbides are entered as spheroid phases in a FCC matrix. It is similar to exd1a except the default HOMOGENIZATION MODEL is used and then ENTER_HOMOGENIZATION_FUNCTION should be used instead of ENTER_LABYRINTH_FUNCTION. This case is from A. Engström, L. Höglund and J. Ågren: Metall. Trans. A, v.25A (1994), pp. 1127-1134.

$$T = 1123 \text{ K}$$



exd1b-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd1b\setup.DCM"

SYS: i>@
NO SUCH COMMAND, USE HELP
SYS: @@ Diffusion in dispersed systems.
SYS: @@ Carburization of Ni-25%Cr alloy: Homogenization model
SYS: @@ This example is about carburization of a Ni-25Cr alloy.
SYS: @@ In this case the M3C2 and M7C3 carbides are entered as
SYS: @@ spheroid phases in a FCC matrix. This case is from
SYS: @@ A. Engström, L. Häglund and J. Ångren: Metall.Trans. A,
SYS: @@ v.25A (1994), pp. 1127-1134.
SYS: @@ This simulation can be run with the DISPERSED SYSTEM MODEL or
SYS: @@ HOMOGENIZATION MODEL. The default HOMOGENIZATION MODEL is used
SYS: @@ and then ENTER_HOMOGENIZATION_FUNCTION should be used instead of
SYS: @@ ENTER_LABYRINTH_FUNCTION.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exd1b_setup.DCM
SYS:
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASE
SYS: @@
SYS: @@
SYS: @@ This example modifies the database interactively, which is not
SYS: @@ yet supported by GES6. Therefore, we enforce the use of GES5.
SYS: set-ges-version 5
SYS:
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ USE THE SSOL DATABASE FOR THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw fedemo
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-sys ni cr c
NI CR C
 DEFINED
TDB_FEDEMO: rej ph * all
GAS:G LIQUID:L BCC_A2
LAVES_PHASE_C14 CBCC_A12 CEMENTITE
CHI_A12 CUB_A13 DIAMOND_FCC_A4
FCC_A1 GRAPHITE HCP_A3
KSI_CARBIDE M23C6 M3C2
M7C3 SIGMA REJECTED
TDB_FEDEMO: res ph fcc,m7c3,m3c2,grap
FCC_A1 M7C3 M3C2
GRAPHITE RESTORED
TDB_FEDEMO: get
REINITIATING GES
ELEMENTS
SPECIES
PHASES
Creating a new composition set FCC_A1#2
PARAMETERS ...
FUNCTIONS ...

List of references for assessed data

'J. Bratberg, Z. Metallkd., 96 (2005) 335-344; Fe-Cr-Mo-C'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'A. Gabriel, C. Chatillon, and I. Ansara, published in High Temp. Sci.
(Parameters listed in CALPHAD, 11 (1987) 203-218); C-Ni'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
'R. Naraghi, Thermo-Calc Software AB, Sweden, 2016; FCC Fe-Cr-C and C-Cr-Ni'
'NPL, unpublished work (1989); C-Cr-Ni'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); CR -Ni'
'B. Uhrenius, Int. J. Refract. Met. Hard Mater. 12 (1994) 121 -127; Molar
volumes'
'P. Villars, L.D. Calvert. Pearson's handbook of crystallographic data for
intermetallic phases, Metals park, Ohio 1985: American society for
metals'
'A.F. Guillermet and G. Grimvall, J. Phys. Chem. Solids, 1992, 53, 105-125;
Molar volumes'
-OK-
TDB_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app mfdemo
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys ni c cr
NI CR
 NI

```

DEFINED
APP: rej ph * all
BCC_A2           FCC_A1 REJECTED
APP: res ph fcc,m7c3,m3c2,grap
*** ERROR M7C3 INPUT IGNORED
*** ERROR M3C2 INPUT IGNORED
*** ERROR GRAP INPUT IGNORED
FCC_A1 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
Creating a new composition set FCC_A1#3
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: Z. Metallkunde 85(1994)502-509; C diffusion in fcc Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Ni diffusion fcc Cr-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'

-OK-
APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR
APP: @@
APP: go d-m
NO TIME STEP DEFINED
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
*** ENTERING M3C2 AS A DIFFUSION NONE PHASE
*** ENTERING M7C3 AS A DIFFUSION NONE PHASE
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 1123; * N
DIC>
DIC> @@
DIC> @@ SET THE REFERENCE STATE FOR CARBON
DIC> @@
DIC> set-reference-state
Component: C
Reference state: grap
Temperature /*/: *
Pressure /100000/: 101325
DIC>
DIC> @@
DIC> @@ ENTER THE REGION aus
DIC> @@
DIC> enter-region aus
DIC>
DIC> @@
DIC> @@ ENTER A GEOMETRICAL GRID INTO THE REGION
DIC> @@
DIC> enter-grid aus 3e-3 geo 100 1.02
DIC>
DIC> @@
DIC> @@ ENTER A MATRIX PHASE IN THE REGION
DIC> @@
DIC> enter-phase act aus matrix fcc_a1#1
DIC>
DIC> @@
DIC> @@ ENTER THE START COMPOSITION FOR THE MATRIX PHASE
DIC> @@
DIC> enter-composition
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1#1/: fcc#1
DEPENDENT COMPONENT ? /NI/: ni
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: cr
TYPE /LINEAR/: lin 25 25
PROFILE FOR /CR/: c
TYPE /LINEAR/: lin 1e-4 1e-4
DIC>
DIC> @@
DIC> @@ ENTER SPHEROIDAL PHASES IN THE REGION
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUS/: aus
PHASE TYPE /MATRIX/: sph
PHASE NAME: /NONE/: m7c3
INFO: EQUILIBRIUM COMPOSITION AND FRACTION OF SPHEROID PHASES USED AS DEFAULT
DIC>
DIC> @@
DIC> @@ ENTER A STOICHIOMETRIC SPHEROIDAL PHASE IN THE REGION
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /AUS/: aus
PHASE TYPE /MATRIX/: sph
PHASE NAME: /NONE/: m3c2
INFO: EQUILIBRIUM COMPOSITION AND FRACTION OF SPHEROID PHASES USED AS DEFAULT
DIC>
DIC>
DIC> @@
DIC> @@ ENTER A START COMPOSITION FOR THE SPHEROIDAL PHASES
DIC> @@
DIC> enter-composition
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1#1/: m7c3
USE EQUILIBRIUM VALUE /Y/: Y
DIC> enter-composition
REGION NAME : /AUS/: aus
PHASE NAME: /FCC_A1#1/: m3c2
USE EQUILIBRIUM VALUE /Y/: Y
DIC>
DIC> @@
DIC> @@ SET THE BOUNDARY CONDITION

```

```
DIC> @@
DIC> set-cond
GLOBAL OR BOUNDARY CONDITION /GLOBAL/: boundary
BOUNDARY /LOWER/: lower
CONDITION TYPE /CLOSED_SYSTEM/: mixed
Dependent substitutional element:NI
Dependent interstitial element:VA
TYPE OF CONDITION FOR COMPONENT C /ZERO_FLUX/: activity
LOW TIME LIMIT /0/: 0
ACR(C)(TIME)= 1;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
TYPE OF CONDITION FOR COMPONENT CR /ZERO_FLUX/: zero-flux
DIC>
DIC> @@
DIC> @@ SELECT THE HOMOGENIZATION FUNCTION
DIC> @@
DIC> enter-homo
ENTER HOMOGENIZATION FUNCTION # /5/: 8
SELECTED FUNCTION IS LABYRINTH FACTOR f**2 WITH PRESCRIBED MATRIX PHASE
PHASE NAME: fcc#1
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME. REMEMBER TO BE CAREFUL WITH THE
DIC> @@ Timestep WHEN SPHEROIDAL PHASES ARE PRESENT. IN THIS CASE
DIC> @@ THE Timestep IS NOT ALLOWED TO BE LARGER THAN 1800s.
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 3600000
AUTOMATIC Timestep CONTROL /YES/: YES
MAX Timestep DURING INTEGRATION /360000/:
INITIAL Timestep : /1E-07/:
SMALLEST ACCEPTABLE Timestep : /1E-07/:
DIC>
DIC> @@
DIC> @@ TO SAVE SOME SPACE ON THE DISK THE RESULTS ARE STORED SELECTIVELY,
DIC> @@ OTHERWISE THE STORE-RESULT-FILE FROM THIS EXAMPLE WOULD BE
DIC> @@ VERY LARGE.
DIC> @@
DIC> set-simulation-condition
NS01A PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA Timesteps IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/:
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/: 99
DEGREE OF IMPLICITY WHEN INTEGRATING PDEs (AUTO, 0 -> 0.5 -> 1): /AUTO/:
MAX Timestep CHANGE PER Timestep : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/:
DIC>
DIC> @@
DIC> @@ SAVE THE SETUP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exd1 y
DIC>
DIC> set-inter
--OK--
DIC>
```

exd1b-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd1b\run.DCM"
DIC>
DIC>
DIC> @@ exd1_run.DCM
DIC>
DIC> @@
DIC> @@ READ THE SETUP FILE AND START THE SIMULATION
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
*** ENTERING M3C2 AS A DIFFUSION NONE PHASE
*** ENTERING M7C3 AS A DIFFUSION NONE PHASE
DIC> read exd1
OK
DIC> sim
DEGREE OF IMPLICITY SET TO EULER BACKWARD
STARTING SIMULATION USING HOMOGENIZATION MODEL
-----
INFO: PHASE WITH LIMITED SOLUBILITY OF ELEMENT(S) EXIST
A FALLBACK PHASE ZZDICTRA_GHOST WILL BE DEFINED
ALONG WITH THE FOLLOWING PARAMETERS:
G(ZZDICTRA_GHOST,C;0)-H298(GRAPHITE,C;0)
G(ZZDICTRA_GHOST,CR;0)-H298(BCC_A2,CR;0)
G(ZZDICTRA_GHOST,NI;0)-H298(FCC_A1,NI;0)
L(ZZDICTRA_GHOST,C,CR;0)
L(ZZDICTRA_GHOST,C,NI;0)
L(ZZDICTRA_GHOST,CR,NI;0)
WARNING:M7C3 HAS NO VOLUME FRACTION, CREATING ONE
WARNING:M3C2 HAS NO VOLUME FRACTION, CREATING ONE
Starting time-step t0= 0.000000 dt= 0.10000000E-06
Starting time-step t0= 0.10000000E-06 dt= 0.20000000E-06
Starting time-step t0= 0.30000000E-06 dt= 0.40000000E-06
Starting time-step t0= 0.70000000E-06 dt= 0.80000000E-06
Starting time-step t0= 0.15000000E-05 dt= 0.16000000E-05
Starting time-step t0= 0.31000000E-05 dt= 0.32000000E-05
Starting time-step t0= 0.63000000E-05 dt= 0.64000000E-05
Starting time-step t0= 0.12700000E-04 dt= 0.12800000E-04
Starting time-step t0= 0.25500000E-04 dt= 0.25600000E-04
Starting time-step t0= 0.51100000E-04 dt= 0.51200000E-04
Starting time-step t0= 0.10230000E-03 dt= 0.10240000E-03
Starting time-step t0= 0.20470000E-03 dt= 0.20480000E-03
Starting time-step t0= 0.40950000E-03 dt= 0.40960000E-03
Starting time-step t0= 0.81910000E-03 dt= 0.81920000E-03
Starting time-step t0= 0.16383000E-02 dt= 0.16384000E-02
Starting time-step t0= 0.32767000E-02 dt= 0.32768000E-02
Starting time-step t0= 0.65535000E-02 dt= 0.65536000E-02
Starting time-step t0= 0.13107100E-01 dt= 0.13107200E-01
Starting time-step t0= 0.26214300E-01 dt= 0.26214400E-01
Starting time-step t0= 0.52428700E-01 dt= 0.52428800E-01
Starting time-step t0= 0.10485750 dt= 0.52428800E-01
Starting time-step t0= 0.15728630 dt= 0.52428800E-01
Starting time-step t0= 0.20971510 dt= 0.52428800E-01
Starting time-step t0= 0.26214390 dt= 0.52428800E-01
Starting time-step t0= 0.31457270 dt= 0.52428800E-01
Starting time-step t0= 0.36700150 dt= 0.10485760
Starting time-step t0= 0.47185910 dt= 0.20971520
Starting time-step t0= 0.68157430 dt= 0.41943040
Starting time-step t0= 1.1010047 dt= 0.41943040
Starting time-step t0= 1.5204351 dt= 0.83886080
Starting time-step t0= 2.3592959 dt= 0.83886080
Starting time-step t0= 3.1981567 dt= 0.83886080
Starting time-step t0= 4.0370175 dt= 0.83886080
Starting time-step t0= 4.8758783 dt= 0.83886080
Starting time-step t0= 5.7147391 dt= 0.83886080
Starting time-step t0= 6.5535999 dt= 0.83886080
Starting time-step t0= 7.3924607 dt= 0.83886080
Starting time-step t0= 8.2313215 dt= 0.83886080
Starting time-step t0= 9.0701823 dt= 0.83886080
Starting time-step t0= 9.9090431 dt= 0.83886080
Starting time-step t0= 10.747904 dt= 0.83886080
Starting time-step t0= 11.586765 dt= 0.83886080
Starting time-step t0= 12.425625 dt= 0.83886080
Starting time-step t0= 13.264486 dt= 0.83886080
Starting time-step t0= 14.103347 dt= 0.83886080
Starting time-step t0= 14.942208 dt= 0.83886080
Starting time-step t0= 15.781069 dt= 0.83886080
Starting time-step t0= 16.619929 dt= 0.83886080
Starting time-step t0= 17.458790 dt= 0.83886080
Starting time-step t0= 18.297651 dt= 0.83886080
Starting time-step t0= 19.136512 dt= 0.83886080
Starting time-step t0= 19.975373 dt= 0.83886080
Starting time-step t0= 20.814233 dt= 0.83886080
Starting time-step t0= 21.653094 dt= 0.83886080
Starting time-step t0= 22.491955 dt= 0.83886080
Starting time-step t0= 23.330816 dt= 0.83886080
Starting time-step t0= 24.169677 dt= 0.83886080
Starting time-step t0= 25.008537 dt= 0.83886080
Starting time-step t0= 25.847398 dt= 0.83886080
Starting time-step t0= 26.686259 dt= 0.83886080
Starting time-step t0= 27.525120 dt= 0.83886080
Starting time-step t0= 28.363981 dt= 0.83886080
Starting time-step t0= 29.202841 dt= 0.83886080
Starting time-step t0= 30.041702 dt= 0.83886080
Starting time-step t0= 30.880563 dt= 0.83886080
Starting time-step t0= 31.719424 dt= 0.83886080
Starting time-step t0= 32.558285 dt= 0.83886080
Starting time-step t0= 33.397145 dt= 0.83886080
Starting time-step t0= 34.236006 dt= 0.83886080
Starting time-step t0= 35.074867 dt= 0.83886080
Starting time-step t0= 35.913728 dt= 0.83886080
Starting time-step t0= 36.752589 dt= 0.83886080
Starting time-step t0= 37.591449 dt= 0.83886080
Starting time-step t0= 38.430310 dt= 1.6777216
Starting time-step t0= 40.108032 dt= 1.6777216
Starting time-step t0= 41.785753 dt= 1.6777216
Starting time-step t0= 43.463475 dt= 1.6777216
Starting time-step t0= 45.141197 dt= 1.6777216
Starting time-step t0= 46.818918 dt= 1.6777216
Starting time-step t0= 48.496640 dt= 1.6777216
```

```

Starting time-step t0= 50.174361 dt= 1.6777216
Starting time-step t0= 51.852083 dt= 1.6777216
Starting time-step t0= 53.529805 dt= 1.6777216
Starting time-step t0= 55.207526 dt= 1.6777216
Starting time-step t0= 56.885248 dt= 1.6777216
Starting time-step t0= 58.562969 dt= 1.6777216
Starting time-step t0= 60.240691 dt= 1.6777216
Starting time-step t0= 61.918413 dt= 1.6777216
Starting time-step t0= 63.596134 dt= 1.6777216
Starting time-step t0= 65.273856 dt= 1.6777216
Starting time-step t0= 66.951577 dt= 1.6777216
Starting time-step t0= 68.629299 dt= 1.6777216
Starting time-step t0= 70.307021 dt= 1.6777216
Starting time-step t0= 71.984742 dt= 1.6777216
Starting time-step t0= 73.662464 dt= 1.6777216
Starting time-step t0= 75.340185 dt= 1.6777216
Starting time-step t0= 77.017907 dt= 3.3554432
Starting time-step t0= 80.373350 dt= 3.3554432
Starting time-step t0= 83.728793 dt= 3.3554432
Starting time-step t0= 87.084237 dt= 3.3554432
Starting time-step t0= 90.439680 dt= 3.3554432
Starting time-step t0= 93.795123 dt= 3.3554432

```

output ignored...

... output resumed

```

DELETING TIME-RECORD FOR TIME 2328955.9
DELETING TIME-RECORD FOR TIME 2335827.9
DELETING TIME-RECORD FOR TIME 2342699.8
DELETING TIME-RECORD FOR TIME 2349571.8
DELETING TIME-RECORD FOR TIME 2356443.7
DELETING TIME-RECORD FOR TIME 2363315.7
DELETING TIME-RECORD FOR TIME 2370187.6
DELETING TIME-RECORD FOR TIME 2377059.6
DELETING TIME-RECORD FOR TIME 2383931.5
DELETING TIME-RECORD FOR TIME 2390803.5
DELETING TIME-RECORD FOR TIME 2397675.4
DELETING TIME-RECORD FOR TIME 2404547.3
DELETING TIME-RECORD FOR TIME 2411419.3
DELETING TIME-RECORD FOR TIME 2418291.2
DELETING TIME-RECORD FOR TIME 2425163.2
DELETING TIME-RECORD FOR TIME 2432035.1
DELETING TIME-RECORD FOR TIME 2438907.1
DELETING TIME-RECORD FOR TIME 2445779.0
DELETING TIME-RECORD FOR TIME 2452651.0
DELETING TIME-RECORD FOR TIME 2459522.9
DELETING TIME-RECORD FOR TIME 2466394.9
DELETING TIME-RECORD FOR TIME 2473266.8
DELETING TIME-RECORD FOR TIME 2480138.8
DELETING TIME-RECORD FOR TIME 2487010.7
DELETING TIME-RECORD FOR TIME 2493882.7
DELETING TIME-RECORD FOR TIME 2500754.6
DELETING TIME-RECORD FOR TIME 2507626.6
DELETING TIME-RECORD FOR TIME 2514498.5
DELETING TIME-RECORD FOR TIME 2521370.5
DELETING TIME-RECORD FOR TIME 2528242.4
DELETING TIME-RECORD FOR TIME 2535114.4
DELETING TIME-RECORD FOR TIME 2541986.3
DELETING TIME-RECORD FOR TIME 2548858.3
DELETING TIME-RECORD FOR TIME 2555730.2
DELETING TIME-RECORD FOR TIME 2562602.1
DELETING TIME-RECORD FOR TIME 2569474.1
DELETING TIME-RECORD FOR TIME 2576346.0
DELETING TIME-RECORD FOR TIME 2583218.0
DELETING TIME-RECORD FOR TIME 2593525.9
DELETING TIME-RECORD FOR TIME 2600397.9
DELETING TIME-RECORD FOR TIME 2607269.8
DELETING TIME-RECORD FOR TIME 2614141.8
DELETING TIME-RECORD FOR TIME 2621013.7
DELETING TIME-RECORD FOR TIME 2627885.6
DELETING TIME-RECORD FOR TIME 2634757.6
DELETING TIME-RECORD FOR TIME 2641629.5
DELETING TIME-RECORD FOR TIME 2648501.5
DELETING TIME-RECORD FOR TIME 2655373.4
DELETING TIME-RECORD FOR TIME 2662245.4
DELETING TIME-RECORD FOR TIME 2669117.3
DELETING TIME-RECORD FOR TIME 2675989.3
DELETING TIME-RECORD FOR TIME 2682861.2
DELETING TIME-RECORD FOR TIME 2689733.2
DELETING TIME-RECORD FOR TIME 2696605.1
DELETING TIME-RECORD FOR TIME 2703477.1
DELETING TIME-RECORD FOR TIME 2710349.0
DELETING TIME-RECORD FOR TIME 2717221.0
DELETING TIME-RECORD FOR TIME 2724092.9
DELETING TIME-RECORD FOR TIME 2730964.9
DELETING TIME-RECORD FOR TIME 2737836.8
DELETING TIME-RECORD FOR TIME 2744708.8
DELETING TIME-RECORD FOR TIME 2758452.7
DELETING TIME-RECORD FOR TIME 2772196.6
DELETING TIME-RECORD FOR TIME 2785940.4
DELETING TIME-RECORD FOR TIME 2799684.3
DELETING TIME-RECORD FOR TIME 2813428.2
DELETING TIME-RECORD FOR TIME 2827172.1
DELETING TIME-RECORD FOR TIME 2840916.0
DELETING TIME-RECORD FOR TIME 2854659.9
DELETING TIME-RECORD FOR TIME 2868403.8
DELETING TIME-RECORD FOR TIME 2875275.8
DELETING TIME-RECORD FOR TIME 2882147.7
DELETING TIME-RECORD FOR TIME 2895891.6
DELETING TIME-RECORD FOR TIME 2909635.5
DELETING TIME-RECORD FOR TIME 2923379.4
DELETING TIME-RECORD FOR TIME 2937123.3
DELETING TIME-RECORD FOR TIME 2950867.2
DELETING TIME-RECORD FOR TIME 2964611.1
DELETING TIME-RECORD FOR TIME 2978355.0
DELETING TIME-RECORD FOR TIME 2992098.9
DELETING TIME-RECORD FOR TIME 3005842.8
DELETING TIME-RECORD FOR TIME 3019586.7
DELETING TIME-RECORD FOR TIME 3033330.6
DELETING TIME-RECORD FOR TIME 3047074.5
DELETING TIME-RECORD FOR TIME 3060818.4
DELETING TIME-RECORD FOR TIME 3074562.2

```

DELETING TIME-RECORD FOR TIME 3088306.1
DELETING TIME-RECORD FOR TIME 3102050.0
DELETING TIME-RECORD FOR TIME 3115793.9
DELETING TIME-RECORD FOR TIME 3129537.8
DELETING TIME-RECORD FOR TIME 3143281.7
DELETING TIME-RECORD FOR TIME 3157025.6
DELETING TIME-RECORD FOR TIME 3170769.5
DELETING TIME-RECORD FOR TIME 3184513.4
DELETING TIME-RECORD FOR TIME 3198257.3
DELETING TIME-RECORD FOR TIME 3212001.2
DELETING TIME-RECORD FOR TIME 3225745.1
DELETING TIME-RECORD FOR TIME 3232617.0
DELETING TIME-RECORD FOR TIME 3239489.0
DELETING TIME-RECORD FOR TIME 3246360.9
DELETING TIME-RECORD FOR TIME 3253232.9
DELETING TIME-RECORD FOR TIME 3260104.8
DELETING TIME-RECORD FOR TIME 3266976.8
DELETING TIME-RECORD FOR TIME 3273848.7
DELETING TIME-RECORD FOR TIME 3280720.7
DELETING TIME-RECORD FOR TIME 3287592.6
DELETING TIME-RECORD FOR TIME 3294464.6
DELETING TIME-RECORD FOR TIME 3301336.5
DELETING TIME-RECORD FOR TIME 3308208.5
DELETING TIME-RECORD FOR TIME 3315080.4
DELETING TIME-RECORD FOR TIME 3321952.4
DELETING TIME-RECORD FOR TIME 3328824.3
DELETING TIME-RECORD FOR TIME 3335696.3
DELETING TIME-RECORD FOR TIME 3349440.2
DELETING TIME-RECORD FOR TIME 3363184.1
DELETING TIME-RECORD FOR TIME 3376927.9
DELETING TIME-RECORD FOR TIME 3390671.8
DELETING TIME-RECORD FOR TIME 3404415.7
DELETING TIME-RECORD FOR TIME 3418159.6
DELETING TIME-RECORD FOR TIME 3431903.5
DELETING TIME-RECORD FOR TIME 3445647.4
DELETING TIME-RECORD FOR TIME 3459391.3
DELETING TIME-RECORD FOR TIME 3473135.2
DELETING TIME-RECORD FOR TIME 3486879.1
DELETING TIME-RECORD FOR TIME 3500623.0
DELETING TIME-RECORD FOR TIME 3514366.9
DELETING TIME-RECORD FOR TIME 3528110.8
DELETING TIME-RECORD FOR TIME 3541854.7
DELETING TIME-RECORD FOR TIME 3555598.6
DELETING TIME-RECORD FOR TIME 3562470.5
DELETING TIME-RECORD FOR TIME 3572778.5

KEEPING TIME-RECORD FOR TIME 3586522.3
AND FOR TIME 3600000.0
WORKSPACE RECLAIMED

INTERPOLATION SCHEME USED THIS FRACTION OF
THE ALLOCATED MEMORY: 0.745280446047089
EFFICIENCY FACTOR: 65.4571748263559

DEALLOCATING

TIMESTEP AT 3600000.00 SELECTED

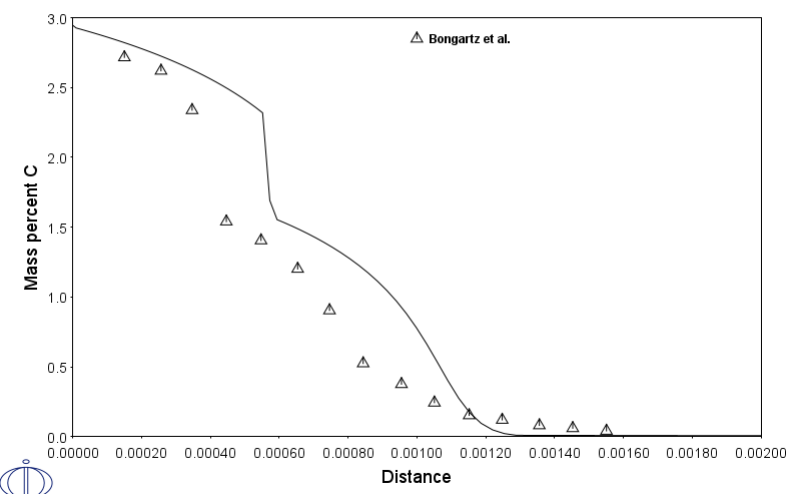
DIC>
DIC> set-inter
--OK---
DIC>

exd1b-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd1b\plot.DCM"
DIC>
DIC>
DIC> @@ exd1_plot.DCM
DIC>
DIC> @@
DIC> @@ FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE exd1b
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 3.60000E+06
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
*** ENTERING M3C2 AS A DIFFUSION NONE PHASE
*** ENTERING M7C3 AS A DIFFUSION NONE PHASE
DIC> read exd1
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ PLOT THE TOTAL CARBON CONCENTRATION PROFILE
POST-1: @@
POST-1: s-d-a y w-p c
POST-1: s-d-a x distance global
INFO: Distance is set as independent variable
POST-1: s-s-s x n 0 2e-3
POST-1: s-p-c time 3600000
POST-1:
POST-1: app y exd1.exp
PROLOGUE NUMBER: /0/: 1
DATASET NUMBER(s): /-1/: 1
POST-1:
POST-1: @@
POST-1: @@ SET A TITLE ON THE PLOT
POST-1: @@
POST-1: set-tit d1.1
POST-1:
POST-1: plot
```

d1.1

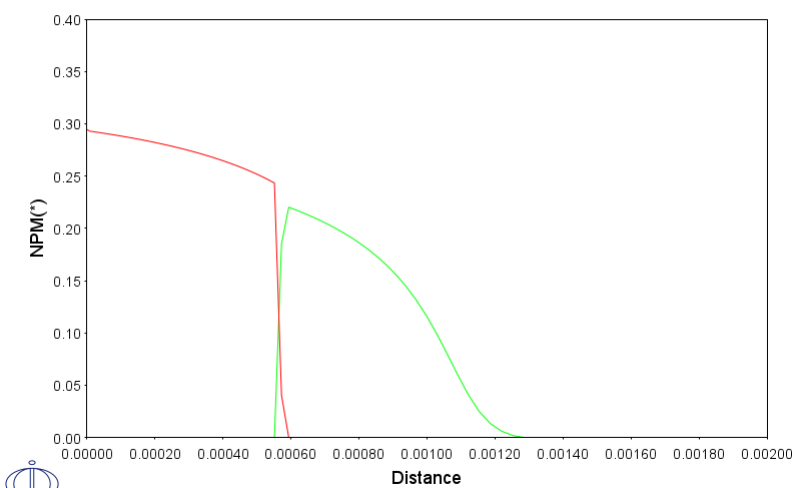
2021.05.13.12.44.01
Time = 3600000
CELL #1



```
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: @@
POST-1: @@ PLOT THE AMOUNT OF CARBIDES FORMED
POST-1: @@
POST-1: s-d-a y npm(*)
POST-1: s-s-s y n 0 0.4
POST-1: app n
POST-1:
POST-1: set-tit d1.2
POST-1: plot
```

d1.2

2021.05.13.12.44.06
Time = 3600000
CELL #1



POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1: set-inter
---OK---
POST-1:



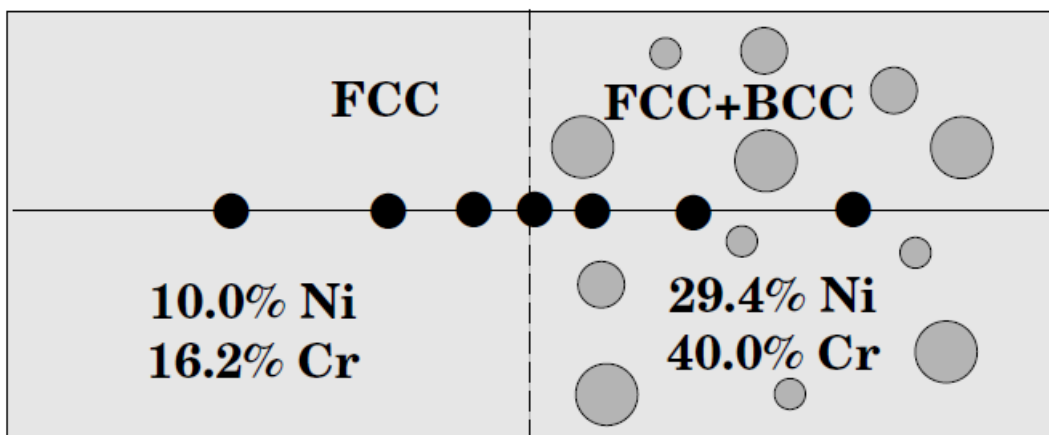
Example exd2a

Diffusion couple of Fe-Ni-Cr alloys: Step-profile

This example calculates the interdiffusion in a diffusion couple between a two-phase (FCC+BCC) and a single-phase (FCC) Fe-Ni-Cr alloy. Initially it uses a step profile. This simulation can be run with either the DISPERSED SYSTEM MODEL or the HOMOGENIZATION MODEL. In this example the DISPERSED SYSTEM MODEL is used.

This case is from A. Engström: Scand. J. Met., vol. 24, 1995, pp.12-20.

T = 1473 K



4E-3

exd2a-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd2a\setup.DCM"
SYS: @@
SYS: @@ Diffusion in dispersed systems.
SYS: @@ Diffusion couple of Fe-Cr-Ni alloys: Dispersed system model
SYS: @@ This example calculates the interdiffusion in a diffusion
SYS: @@ couple between a two-phase (FCC+BCC) and a single-phase (FCC)
SYS: @@ Fe-Ni-Cr alloy. This case is from A. Engström: Scand. J. Met., v. 24,
SYS: @@ 1995, pp.12-20. This simulation can be run with either the DISPERSED
SYS: @@ SYSTEM MODEL or the HOMOGENIZATION MODEL.
SYS: @@ In this example the DISPERSED SYSTEM MODEL is used, which requires
SYS: @@ that the default HOMOGENIZATION MODEL is disabled.
SYS: @@ With the DISPERSED SYSTEM MODEL the command
SYS: @@ ENTER_LABYRINTH_FUNCTION is used to take into account the
SYS: @@ impeding effect of dispersed phases on long-range diffusion.
SYS: @@ For the HOMOGENIZATION MODEL the command
SYS: @@ ENTER_HOMOGENIZATION_FUNCTION should be used.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exd2_setup.DCM
SYS:
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA          /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: sw fedemo
Current database: Iron Demo Database v4.0

VA          /- DEFINED
TDB_FEDEMO: def-sys fe ni cr
FE           NI           CR
DEFINED
TDB_FEDEMO: rej ph * all
LIQUID:L      BCC_A2      LAVES_PHASE_C14
CBCC_A12     CHI_A12     CUB_A13
FCC_A1       HCP_A3      SIGMA
REJECTED
TDB_FEDEMO: res ph fcc,bcc
FCC_A1       BCC_A2      RESTORED
TDB_FEDEMO: get
12:45:23,196 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
    volumes'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'J-O. Andersson and B. Sundman, CALPHAD, 11 (1987) 83-92; TRITA 0270
    (1986); CR-FE'
'B.J. Lee, CALPHAD, 17 (1993) 251-268; revision of Fe-Cr and Fe -Ni liquid'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); CR -NI'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
-OK-
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app mfdemo
Current database: Fe-Alloys Mobility demo database v2.0

VA  DEFINED
APP: def-sys fe ni cr
FE           NI           CR
DEFINED
APP: rej ph * all
BCC_A2      FCC_A1      REJECTED
APP: res ph fcc,bcc
FCC_A1       BCC_A2      RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Ni diffusion fcc Cr-Ni'
'B. Jonsson: Z. Metallkunde 86(1995)686-692; Cr, Fe and Ni diffusion fcc
    Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
```

```

-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
'B. Jonsson: ISIJ International, 35(1995)1415-1421; Cr, Fe and Ni
diffusion bcc Cr-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'

-OK-
APP:
APP: @@ 
APP: @@ ENTER THE DICTRA MONITOR
APP: @@ 
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@ 
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@ 
DIC> set-cond glob T 0 1473; * N
DIC>
DIC> @@ 
DIC> @@ ENTER THE REGION fer
DIC> @@ 
DIC> enter-region fer
DIC>
DIC>
DIC> @@ 
DIC> @@ ENTER A DOUBLE GEOMETRICAL GRID INTO THE REGION
DIC> @@ THIS GIVES A SHORT DISTANCE BETWEEN THE GRIDPOINTS
DIC> @@ IN THE MIDDLE OF THE REGION WHERE THE INITIAL INTERFACE IS
DIC> @@ 
DIC> enter-grid fer
WIDTH OF REGION /1/: 4e-3
TYPE /LINEAR/: double
NUMBER OF POINTS /50/: 200
VALUE OF R IN THE GEOMETRICAL SERIE FOR LOWER PART OF REGION: 0.97
VALUE OF R IN THE GEOMETRICAL SERIE FOR UPPER PART OF REGION: 1.03093
DIC>
DIC> @@ 
DIC> @@ ENTER A MATRIX PHASE IN THE REGION
DIC> @@ 
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act fer matrix fcc
DIC>
DIC> @@ 
DIC> @@ ENTER THE START COMPOSITION FOR THE MATRIX PHASE FROM FILES
DIC> @@ 
DIC> enter-composition
REGION NAME : /FER/: fer
PHASE NAME: /FCC_A1/: fcc
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /CR/: cr
TYPE /LINEAR/: read d2cr.dat
PROFILE FOR /NI/: ni
TYPE /LINEAR/: read d2ni.dat
DIC>
DIC> @@ 
DIC> @@ ENTER FERRITE AS THE SPHEROIDAL PHASE IN THE REGION.
DIC> @@ SINCE THE FRACTION OF FERRITE IS SMALL, AND THESE APPEAR
DIC> @@ AS ISOLATED PARTICLES, FERRITE IS ENTERED AS A SPHEROIDAL PHASE
DIC> @@ 
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /FER/: fer
PHASE TYPE /MATRIX/: sph
PHASE NAME: /NONE/: bcc
INFO: EQUILIBRIUM COMPOSITION AND FRACTION OF SPHEROID PHASES USED AS DEFAULT
DIC>
DIC> @@ 
DIC> @@ ENTER A COMPOSITION FOR THE SPHEROIDAL PHASE
DIC> @@ USE THE EQUILIBRIUM VALUE
DIC> @@ 
DIC> enter-composition
REGION NAME : /FER/: fer
PHASE NAME: /FCC_A1/: bcc
USE EQUILIBRIUM VALUE /Y/: y
DIC>
DIC>
DIC> @@ 
DIC> @@ ENTER A LABYRINTH FACTOR
DIC> @@ IN THIS CASE THE LOW DIFFUSIVITY PHASE IS THE MATRIX AND THE
DIC> @@ "EFFECTIVE" DIFFUSIVITY IN THE AUSTENITE+FERRITE TWO-PHASE
DIC> @@ REGION IS EXPECTED TO BE HIGHER THAN THE DIFFUSIVITY IN THE
DIC> @@ AUSTENITE.
DIC> enter-lab
REGION NAME : fer
f(T,P,VOLFR,X)= 1+3*(1-volfr)/volfr;
DIC>
DIC> @@ 
DIC> @@ SET THE SIMULATION TIME AND OTHER SIMULATION PARAMETERS
DIC> @@ 
DIC> set-simulation-time
END TIME FOR INTEGRATION /1/: 720000
AUTOMATIC TIMESTEP CONTROL /YES/: YES
MAX TIMESTEP DURING INTEGRATION /72000/: 5000
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC> set-simulation-condition
NS01A PRINT CONTROL : /0/:
FLUX CORRECTION FACTOR : /1/:
NUMBER OF DELTA Timesteps IN CALLING MULDIF: /2/:
CHECK INTERFACE POSITION /AUTO/:
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/:
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/:
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/: 99
DEGREE OF IMPLICITY WHEN INTEGRATING PDES (AUTO, 0 -> 0.5 -> 1): /AUTO/:
MAX TIMESTEP CHANGE PER TIMESTEP : /2/:
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/:
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/:

```

CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/:
DIC> @@ BY DEFAULT THE "HOMOGENIZATION MODEL" IS USED WHEN MULTIPLE PHASES
DIC> @@ ARE ENTERED IN A SINGLE REGION. THE HOMOGENIZATION MODEL NEEDS TO BE
DIC> @@ DISABLED FOR THIS EXAMPLE.
DIC> ho n
HOMOGENIZATION DISABLED
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC>
DIC> save exd2 y
DIC>
DIC> set-inter
--OK--
DIC>

exd2a-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd2a\run.DCM"
DIC>
DIC>
DIC> @@ exd2_run.DCM
DIC>
DIC> @@
DIC> FILE FOR RUNNING THE SIMULATION OF EXAMPLE D2
DIC> @@
DIC>
DIC> @@
DIC> @@ READ THE SETUP FROM FILE AND START THE SIMULATION
DIC> @@
DIC>
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exd2
OK
DIC> sim
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: CR = .297842647391913 FE = .517227320517284
NI = .184930032090802
TOTAL SIZE OF SYSTEM: .004 [m]
WARNING:BCC_A2 HAS NO VOLUME FRACTION, CREATING ONE
U-FRACTION IN SYSTEM: CR = .297842647391914 FE = .517227320517284
NI = .184930032090802
TOTAL SIZE OF SYSTEM: .004 [m]
TIME = 0.10000000E-06 DT = 0.10000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297842647356821 FE = .517227320600573
NI = .184930032042606
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 0.10010000E-03 DT = 0.10000000E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297842648005431 FE = .517227319061159
NI = .18493003293341
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 0.40010010 DT = 0.40000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297842676377595 FE = .517227251840507
NI = .184930071781899
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 281.72200 DT = 281.32190 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297847760270075 FE = .517216317308819
NI = .184935922421106
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 844.36579 DT = 562.64380 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297849615184743 FE = .517212946633452
NI = .184937438181806
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 1 seconds
TIME = 1969.6534 DT = 1125.2876 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297850550731974 FE = .517211426818697
NI = .184938022449329
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 4220.2286 DT = 2250.5752 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297851119000641 FE = .517210569323811
NI = .184938311675549
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 8721.3789 DT = 4501.1504 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297851494954336 FE = .517210020163941
NI = .184938484881723
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 13721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .29785173211369 FE = .517209740277053
NI = .184938527609257
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 18721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .29785188345351 FE = .517209584045532
NI = .184938532500958
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 1 seconds
TIME = 23721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297851977657557 FE = .517209483265972
NI = .184938539076472
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 28721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852029985828 FE = .517209412426188
NI = .184938557587983
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 33721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852049766013 FE = .517209361691075
NI = .184938588542912
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 38721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852045015471 FE = .517209326359908
NI = .184938628624622
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 43721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852052300694 FE = .517209299378172
NI = .184938648321134
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 48721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852064900644 FE = .517209276184261
NI = .184938658915095
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep 0 seconds
TIME = 53721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852069935262 FE = .517209260646723
NI = .184938669418015
TOTAL SIZE OF SYSTEM: .004 [m]
```

```

CPU time used in timestep          1 seconds
TIME = 58721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852077632328 FE = .517209244116551
NI = .184938678251121
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 63721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852087307341 FE = .517209229697541
NI = .184938682995117
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 68721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852092166835 FE = .517209219702968
NI = .184938688130197
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 73721.379 DT = 5000.0000 SUM OF SQUARES = 0.0000000

```

output ignored...

```

... output resumed

U-FRACTION IN SYSTEM: CR = .297852035243634 FE = .517209097316277
NI = .1849386744009
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 608721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852040343917 FE = .51720909582834
NI = .18493863827743
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 613721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852046756561 FE = .517209094403527
NI = .184938858839912
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 618721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852053277545 FE = .51720909326319
NI = .184938853459266
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 623721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852059288762 FE = .517209092514195
NI = .184938848197043
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 628721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852064436494 FE = .517209092217949
NI = .184938843345557
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 633721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852068500944 FE = .51720909241553
NI = .184938839083526
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          1 seconds
TIME = 638721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852071392547 FE = .51720909308393
NI = .184938835523523
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 643721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852073681692 FE = .517209091807033
NI = .184938834511275
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 648721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .29785207451878 FE = .517209091190446
NI = .184938834290774
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 653721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852073880058 FE = .517209091158173
NI = .184938834961769
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 658721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852071737293 FE = .51720909167323
NI = .184938836589476
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 663721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852068061707 FE = .517209092719061
NI = .184938839219232
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          1 seconds
TIME = 668721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852062825047 FE = .517209094290037
NI = .184938842884917
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 673721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852056000143 FE = .51720909638732
NI = .184938847612538
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 678721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852047560719 FE = .517209099016375
NI = .184938853422906
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 683721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .29785203748116 FE = .517209102185644
NI = .18493860333196
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 688721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852031720919 FE = .517209104528
NI = .184938863751081
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 693721.38 DT = 5000.0000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .29785203510703 FE = .517209102778461

```

```

NI = .184938862114509
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 698721.38      DT = 5000.0000      SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852041265681 FE = .517209100862436
NI = .184938857871883
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          1 seconds
TIME = 703721.38      DT = 5000.0000      SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852048239014 FE = .517209099115
NI = .184938852645985
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 708721.38      DT = 5000.0000      SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .297852055131709 FE = .517209097666817
NI = .184938847201474
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 713721.38      DT = 5000.0000      SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .29785206147074 FE = .51720909658047
NI = .18493884194879
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 718721.38      DT = 5000.0000      SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .29785206697996 FE = .517209095891388
NI = .184938837128653
TOTAL SIZE OF SYSTEM: .004 [m]
CPU time used in timestep          0 seconds
TIME = 720000.00      DT = 1278.6211      SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: CR = .29785207176732 FE = .517209095429353
NI = .184938832803327
TOTAL SIZE OF SYSTEM: .004 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME    0.0000000
DELETING TIME-RECORD FOR TIME    0.10000000E-06
DELETING TIME-RECORD FOR TIME    0.10010000E-03
DELETING TIME-RECORD FOR TIME    468721.38
DELETING TIME-RECORD FOR TIME    708721.38
DELETING TIME-RECORD FOR TIME    713721.38

KEEPING TIME-RECORD FOR TIME    718721.38
AND FOR TIME                   720000.00
WORKSPACE RECLAIMED

TIMESTEP AT      720000.000      SELECTED
```

```

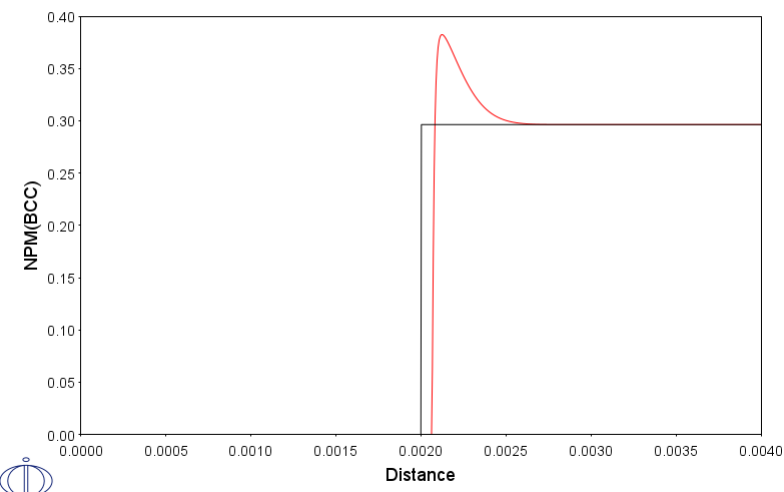
DIC>
DIC>
DIC>
DIC>
DIC> set-inter
--OK--
DIC>
```

exd2a-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd2a\plot.DCM"
DIC>
DIC>
DIC> @@ exd2_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE D2
DIC> @@
DIC> @@ ENTER THE DICTRA MODULE AND SPECIFY THE STORE-RESULT FILE
DIC> @@
DIC>
DIC>
DIC> go d-m
TIME STEP AT TIME 7.20000E+05
DIC> read exd2
OK
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1:
POST-1: @@
POST-1: @@ LET US SEE HOW THE FRACTION OF FERRITE HAS CHANGED
POST-1: @@ AS A RESULT OF THE DIFFUSION
POST-1: @@
POST-1: s-d-a y npm(bcc)
POST-1: s-d-a x distance global
INFO: Distance is set as independent variable
POST-1: s-p-c time 0 720000
POST-1: set-tit Figure D2.1
POST-1: plot
```

Figure D2.1

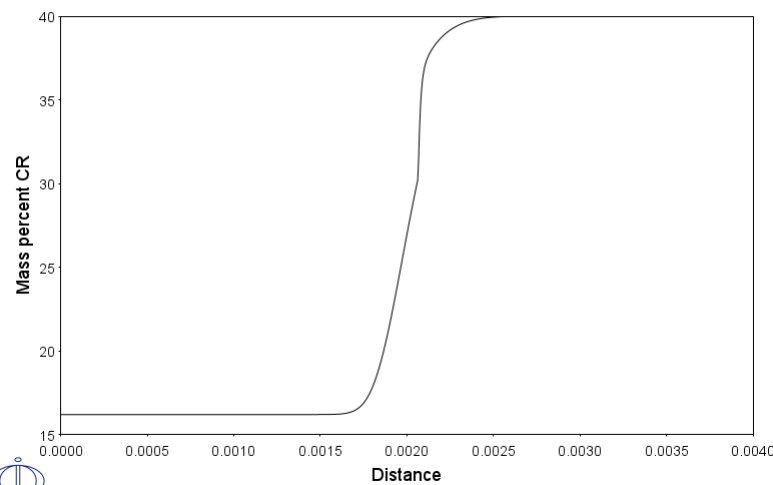
2021.05.13.12.47.50
Time = 0.720000
CELL #1



```
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: @@
POST-1: @@ NOW PLOT HOW THE AVERAGE Cr-CONCENTRATION VARIES WITH DISTANCE
POST-1: @@
POST-1: s-d-a y w-p cr
POST-1: s-p-c time last
POST-1: set-tit Figure D2.2
POST-1: plot
```

Figure D2.2

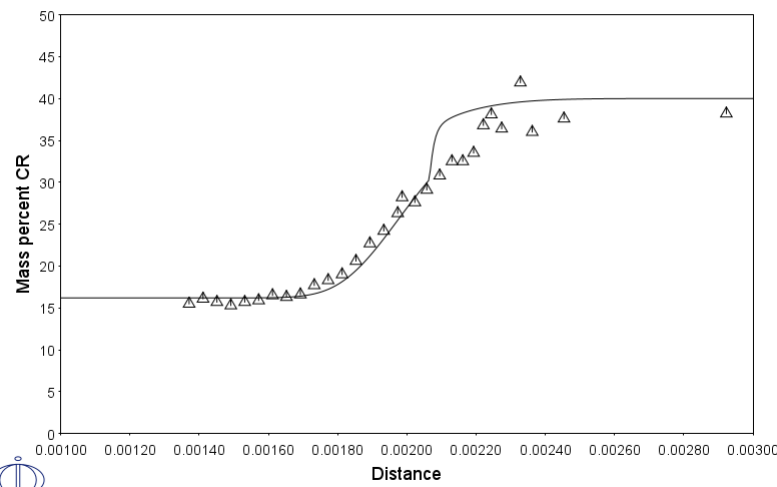
2021.05.13.12.47.51
Time = 720000
CELL #1



POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: @@
POST-1: @@ NOW SELECT A BETTER SCALING AND APPEND THE EXPERIMENTAL DATA
POST-1: @@
POST-1:
POST-1: app y exd2.exp
PROLOGUE NUMBER: /0/: 1
DATASET NUMBER(s): /-1/: 4
POST-1:
POST-1: s-s-s y n 0 50
POST-1: s-s-s x n 10e-4 30e-4
POST-1:
POST-1: set-tit Figure D2.3
POST-1: plot

Figure D2.3

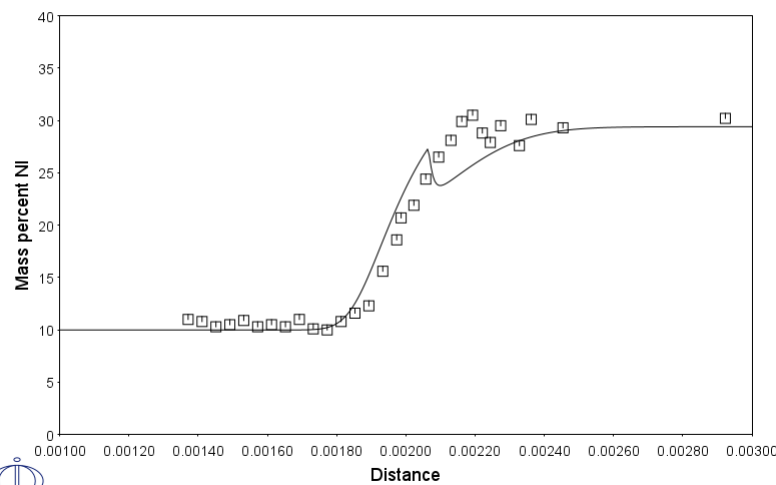
2021.05.13.12.47.52
Time = 720000
CELL #1



POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: @@
POST-1: @@ NOW WE DO THE SAME FOR NICKEL
POST-1: @@
POST-1:
POST-1: s-d-a y w-p ni
POST-1:
POST-1: app y exd2.exp
PROLOGUE NUMBER: /1/: 1
DATASET NUMBER(s): /-1/: 5
POST-1:
POST-1: s-s-s x n 10e-4 30e-4
POST-1: s-s-s y n 0 40
POST-1:
POST-1: set-tit Figure D2.4
POST-1: plot

Figure D2.4

2021.05.13.12.47.53
Time = 720000
CELL #1



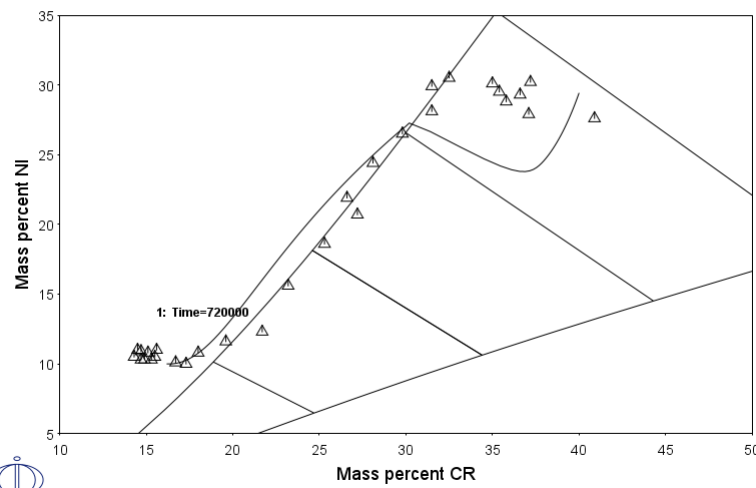
```

POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: @@
POST-1: @@ LET US PLOT THE DIFFUSION PATH FOR THE COUPLE.
POST-1: @@ WE APPEND THE TERNARY PHASE-DIAGRAM CALCULATED IN THERMO-CALC
POST-1: @@ AND THE EXPERIMENTAL DATA
POST-1: @@
POST-1: s-d-a x w-p cr
POST-1: s-d-a y w-p ni
POST-1: s-i-v dis gl
POST-1: s-p-c time last
POST-1:
POST-1: app y exd2.exp
PROLOGUE NUMBER: /1/: 1
DATASET NUMBER(s): /-1/: 6 7 8
POST-1:
POST-1: s-s-s x n 10 50
POST-1: s-s-s y n 5 35
POST-1:
POST-1: set-tit Figure D2.5
POST-1: plot

```

Figure D2.5

2021.05.13.12.47.54
Time = 720000
CELL #1



```

POST-1:
POST-1:
POST-1:
POST-1:
POST-1: set-interactive
--OK--
POST-1:

```

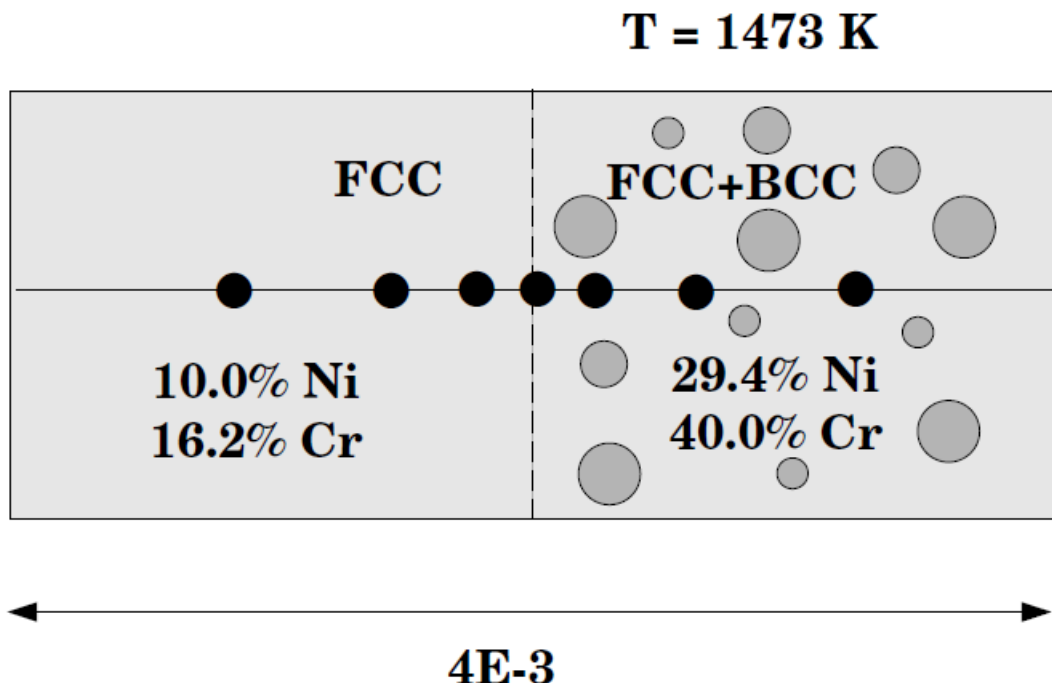


Example exd2b

Diffusion couple of Fe-Ni-Cr alloys: Homogenization model

This example calculates the interdiffusion in a diffusion couple between a two-phase (FCC+BCC) and a single-phase (FCC) Fe-Ni-Cr alloy. Initially it has a step profile. It is similar to exd2a except the default HOMOGENIZATION MODEL is used and then ENTER_HOMOGENIZATION_FUNCTION is used instead of ENTER_LABYRINTH_FUNCTION.

This case is from A. Engström: Scand. J. Met., vol. 24, 1995, pp.12-20.



exd2b-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd2b\setup.DCM"

SYS: i>_@
NO SUCH COMMAND, USE HELP
SYS: @@ Diffusion in dispersed systems.
SYS: @@ Diffusion couple of Fe-Cr-Ni alloys: Homogenization model
SYS: @@ This example calculates the interdiffusion in a diffusion
SYS: @@ couple between a two-phase (FCC+BCC) and a single-phase (FCC)
SYS: @@ Fe-Ni-Cr alloy. This case is from A. Engström: Scand. J. Met.,
SYS: @@ v. 24, 1995, pp.12-20. This simulation can be run with either
SYS: @@ the DISPERSED SYSTEM MODEL or the HOMOGENIZATION MODEL.
SYS: @@ Here the default HOMOGENIZATION MODEL is used and then
SYS: @@ ENTER_HOMOGENIZATION_FUNCTION should be used instead of
SYS: @@ ENTER_LABYRINTH_FUNCTION.

SYS: -----
NO SUCH COMMAND, USE HELP

SYS:
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: sw fedemo
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-sys fe ni cr
FE NI CR
DEFINED
TDB_FEDEMO: rej ph * all
LIQUID:L BCC_A2 LAVES_PHASE_C14
CBCC_A12 CHI_A12 CUB_A13
FCC_A1 HCP_A3 SIGMA
REJECTED
TDB_FEDEMO: res ph fcc,bcc
FCC_A1 BCC_A2 RESTORED
TDB_FEDEMO: get
12:49:11,623 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES
ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'J-O. Andersson and B. Sundman, CALPHAD, 11 (1987) 83-92; TRITA 0270
(1986); CR-FE'
'B.J. Lee, CALPHAD, 17 (1993) 251-268; revision of Fe-Cr and Fe -Ni liquid'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); CR -NI'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
-OK-

TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app mfedemo
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys fe ni cr
FE NI CR
DEFINED
APP: rej ph * all
BCC_A2 FCC_A1 REJECTED
APP: res ph fcc,bcc
FCC_A1 BCC_A2 RESTORED
APP: get
ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Ni diffusion fcc Cr-Ni'
'B. Jonsson: Z. Metallkunde 86(1995)686-692; Cr, Fe and Ni diffusion fcc
Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
'B. Jonsson: ISIJ International, 35(1995)1415-1421; Cr, Fe and Ni
diffusion bcc Cr-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion'

```

    in bcc Fe'
-OK-
APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 1473; * N
DIC>
DIC> @@
DIC> @@ ENTER THE REGION fer
DIC> @@
DIC> enter-region fer
DIC>
DIC>
DIC> @@
DIC> @@ ENTER A DOUBLE-GEOMETRICAL GRID INTO THE REGION
DIC> @@ THIS GIVES A SHORT DISTANCE BETWEEN THE GRIDPOINTS
DIC> @@ IN THE MIDDLE OF THE REGION WHERE THE INITIAL INTERFACE IS
DIC> @@
DIC> enter-grid fer
WIDTH OF REGION /1/: 4e-3
TYPE /LINEAR/: double
NUMBER OF POINTS /50/: 200
VALUE OF R IN THE GEOMETRICAL SERIE FOR LOWER PART OF REGION: 0.97
VALUE OF R IN THE GEOMETRICAL SERIE FOR UPPER PART OF REGION: 1.03093
DIC>
DIC> @@
DIC> @@ ENTER A MATRIX PHASE IN THE REGION
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act fer matrix fcc
DIC>
DIC> @@
DIC> @@ ENTER THE START COMPOSITION FOR THE MATRIX PHASE FROM FILES
DIC> @@
DIC> enter-composition
REGION NAME : /FER/: fer
PHASE NAME: /FCC_A1/: fcc
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /CR/: cr
TYPE /LINEAR/: read d2cr.dat
PROFILE FOR /NI/: ni
TYPE /LINEAR/: read d2ni.dat
DIC>
DIC> @@
DIC> @@ ENTER FERRITE AS THE SPHEROIDAL PHASE IN THE REGION
DIC> @@ SINCE THE FRACTION OF FERRITE IS SMALL, AND THESE APPEAR
DIC> @@ AS ISOLATED PARTICLES, FERRITE IS ENTERED AS A SPHEROIDAL PHASE
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /FER/: fer
PHASE TYPE /MATRIX/: sph
PHASE NAME: /NONE/: bcc
INFO: EQUILIBRIUM COMPOSITION AND FRACTION OF SPHEROID PHASES USED AS DEFAULT
DIC>
DIC> @@
DIC> @@ ENTER THE COMPOSITION FOR THE SPHEROIDAL PHASE
DIC> @@ USE THE EQUILIBRIUM VALUE
DIC> @@
DIC> enter-composition
REGION NAME : /FER/: fer
PHASE NAME: /FCC_A1/: bcc
USE EQUILIBRIUM VALUE /Y/: y
DIC>
DIC> @@ SELECT A HOMOGENIZATION FUNCTION
DIC> @@ IN THIS CASE THE LOWER HASHIN-SHTRIKMAN BOUND
DIC> en-ho 1
SELECTED FUNCTION IS HASHIN-SHTRIKMAN BOUND: GENERAL LOWER
DIC>
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME AND OTHER SIMULATION PARAMETERS
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 720000
AUTOMATIC TIMESTEP CONTROL /YES/: YES
MAX TIMESTEP DURING INTEGRATION /72000/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC> set-simulation-condition
NS01A PRINT CONTROL : /0/: 0
FLUX CORRECTION FACTOR : /1/: 1
NUMBER OF DELTA Timesteps IN CALLING MULDIF: /2/: 2
CHECK INTERFACE POSITION /AUTO/: n
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/: act
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/: y
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/: 99
DEGREE OF IMPLICITY WHEN INTEGRATING PDEs (AUTO, 0 -> 0.5 -> 1): /AUTO/: 1
MAX TIMESTEP CHANGE PER TIMESTEP : /2/: 2
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/: n
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/: y
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/: n
DIC>
DIC>
DIC> @@
DIC> @@ SAVE THE SETUP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC>
DIC> save exd2 y
DIC>
DIC> set-inter

```

--OK--

DIC>

exd2b-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd2b\run.DCM"
DIC>
DIC>
DIC> @@ exd2_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING THE SIMULATION OF EXAMPLE D2
DIC> @@
DIC>
DIC>
DIC> @@
DIC> @@ READ THE SETUP FROM FILE AND START THE SIMULATION
DIC> @@
DIC>
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exd2
OK
DIC> sim
STARTING SIMULATION USING HOMOGENIZATION MODEL
-----
WARNING:BCC_A2 HAS NO VOLUME FRACTION, CREATING ONE
Starting time-step t0= 0.0000000 dt= 0.10000000E-06
Starting time-step t0= 0.10000000E-06 dt= 0.20000000E-06
Starting time-step t0= 0.30000000E-06 dt= 0.40000000E-06
Starting time-step t0= 0.70000000E-06 dt= 0.80000000E-06
Starting time-step t0= 0.15000000E-05 dt= 0.16000000E-05
Starting time-step t0= 0.31000000E-05 dt= 0.32000000E-05
Starting time-step t0= 0.63000000E-05 dt= 0.64000000E-05
Starting time-step t0= 0.12700000E-04 dt= 0.12800000E-04
Starting time-step t0= 0.25500000E-04 dt= 0.25600000E-04
Starting time-step t0= 0.51100000E-04 dt= 0.51200000E-04
Starting time-step t0= 0.10230000E-03 dt= 0.10240000E-03
Starting time-step t0= 0.20470000E-03 dt= 0.20480000E-03
Starting time-step t0= 0.40950000E-03 dt= 0.40960000E-03
Starting time-step t0= 0.81910000E-03 dt= 0.81920000E-03
Starting time-step t0= 0.16383000E-02 dt= 0.16384000E-02
Starting time-step t0= 0.32767000E-02 dt= 0.32768000E-02
Starting time-step t0= 0.65535000E-02 dt= 0.65536000E-02
Starting time-step t0= 0.13107100E-01 dt= 0.13107200E-01
Starting time-step t0= 0.26214300E-01 dt= 0.26214400E-01
Starting time-step t0= 0.52428700E-01 dt= 0.52428800E-01
Starting time-step t0= 0.10485750 dt= 0.10485760
Starting time-step t0= 0.20971510 dt= 0.20971520
Starting time-step t0= 0.41943030 dt= 0.41943040
Starting time-step t0= 0.83886070 dt= 0.83886080
Starting time-step t0= 1.6777215 dt= 1.6777216
Starting time-step t0= 3.3554431 dt= 3.3554432
Starting time-step t0= 6.7108863 dt= 6.7108864
Starting time-step t0= 13.421773 dt= 13.421773
Starting time-step t0= 26.843545 dt= 26.843546
Starting time-step t0= 53.687091 dt= 26.843546
Starting time-step t0= 80.530637 dt= 26.843546
Starting time-step t0= 107.37418 dt= 26.843546
Starting time-step t0= 134.21773 dt= 26.843546
Starting time-step t0= 161.06127 dt= 26.843546
Starting time-step t0= 187.90482 dt= 26.843546
Starting time-step t0= 214.74836 dt= 53.687091
Starting time-step t0= 268.43546 dt= 53.687091
Starting time-step t0= 322.12255 dt= 53.687091
Starting time-step t0= 375.80964 dt= 53.687091
Starting time-step t0= 429.49673 dt= 53.687091
Starting time-step t0= 483.18382 dt= 53.687091
Starting time-step t0= 536.87091 dt= 53.687091
Starting time-step t0= 590.55800 dt= 107.37418
Starting time-step t0= 697.93219 dt= 107.37418
Starting time-step t0= 805.30637 dt= 107.37418
Starting time-step t0= 912.68055 dt= 107.37418
Starting time-step t0= 1020.0547 dt= 107.37418
Starting time-step t0= 1127.4289 dt= 214.74836
Starting time-step t0= 1342.1773 dt= 214.74836
Starting time-step t0= 1556.9256 dt= 214.74836
Starting time-step t0= 1771.6740 dt= 214.74836
Starting time-step t0= 1986.4224 dt= 214.74836
Starting time-step t0= 2201.1707 dt= 214.74836
Starting time-step t0= 2415.9191 dt= 214.74836
Starting time-step t0= 2630.6675 dt= 429.49673
Starting time-step t0= 3060.1642 dt= 429.49673
Starting time-step t0= 3489.6609 dt= 429.49673
Starting time-step t0= 3919.1577 dt= 429.49673
Starting time-step t0= 4348.6544 dt= 429.49673
Starting time-step t0= 4778.1511 dt= 429.49673
Starting time-step t0= 5207.6478 dt= 429.49673
Starting time-step t0= 5637.1446 dt= 858.99346
Starting time-step t0= 6496.1380 dt= 858.99346
Starting time-step t0= 7355.1315 dt= 858.99346
Starting time-step t0= 8214.1250 dt= 858.99346
Starting time-step t0= 9073.1184 dt= 858.99346
Starting time-step t0= 9932.1119 dt= 858.99346
Starting time-step t0= 10791.105 dt= 858.99346
Starting time-step t0= 11650.099 dt= 1717.9869
Starting time-step t0= 13368.086 dt= 1717.9869
Starting time-step t0= 15086.073 dt= 1717.9869
Starting time-step t0= 16804.060 dt= 1717.9869
Starting time-step t0= 18522.046 dt= 1717.9869
Starting time-step t0= 20240.033 dt= 1717.9869
Starting time-step t0= 21958.020 dt= 3435.9738
Starting time-step t0= 25393.994 dt= 3435.9738
Starting time-step t0= 28829.968 dt= 3435.9738
Starting time-step t0= 30547.955 dt= 3435.9738
Starting time-step t0= 33983.929 dt= 3435.9738
Starting time-step t0= 37419.903 dt= 3435.9738
Starting time-step t0= 40855.876 dt= 3435.9738
Starting time-step t0= 44291.850 dt= 6871.9477
Starting time-step t0= 51163.798 dt= 6871.9477
Starting time-step t0= 58035.746 dt= 6871.9477
Starting time-step t0= 64907.693 dt= 6871.9477
Starting time-step t0= 71779.641 dt= 6871.9477
Starting time-step t0= 78651.589 dt= 6871.9477
Starting time-step t0= 85523.536 dt= 6871.9477
Starting time-step t0= 88959.510 dt= 6871.9477
```

```

Starting time-step t0= 95831.458 dt= 13743.895
Starting time-step t0= 109575.35 dt= 13743.895
Starting time-step t0= 116447.30 dt= 13743.895
Starting time-step t0= 130191.20 dt= 13743.895
Starting time-step t0= 143935.09 dt= 13743.895
Starting time-step t0= 147371.07 dt= 6871.9477
Starting time-step t0= 150807.04 dt= 3435.9738
Starting time-step t0= 154243.01 dt= 6871.9477
Starting time-step t0= 161114.96 dt= 13743.895
Starting time-step t0= 174858.86 dt= 13743.895
Starting time-step t0= 188602.75 dt= 27487.791
Starting time-step t0= 216090.54 dt= 27487.791
Starting time-step t0= 229834.44 dt= 13743.895
Starting time-step t0= 243578.33 dt= 27487.791
Starting time-step t0= 271066.12 dt= 27487.791
Starting time-step t0= 277938.07 dt= 13743.895
Starting time-step t0= 291681.97 dt= 27487.791
Starting time-step t0= 319169.76 dt= 27487.791
Starting time-step t0= 346657.55 dt= 27487.791
Starting time-step t0= 374145.34 dt= 27487.791
Starting time-step t0= 387889.23 dt= 27487.791
Starting time-step t0= 394761.18 dt= 13743.895
Starting time-step t0= 408505.08 dt= 13743.895
Starting time-step t0= 422248.97 dt= 27487.791
Starting time-step t0= 449736.76 dt= 27487.791
Starting time-step t0= 463480.66 dt= 27487.791
Starting time-step t0= 470352.61 dt= 6871.9477
Starting time-step t0= 477224.55 dt= 13743.895
Starting time-step t0= 490968.45 dt= 27487.791
Starting time-step t0= 518456.24 dt= 27487.791
Starting time-step t0= 532200.14 dt= 27487.791
Starting time-step t0= 545944.03 dt= 27487.791
Starting time-step t0= 573431.82 dt= 27487.791
Starting time-step t0= 600919.61 dt= 54975.581
Starting time-step t0= 628407.40 dt= 27487.791
Starting time-step t0= 642151.30 dt= 13743.895
Starting time-step t0= 655895.19 dt= 27487.791
Starting time-step t0= 683382.98 dt= 36617.016

```

MUST SAVE WORKSPACE ON FILE

WORKSPACE SAVED ON FILE

RECLAIMING WORKSPACE

```

DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 805.30637
DELETING TIME-RECORD FOR TIME 1556.9256
DELETING TIME-RECORD FOR TIME 2415.9191
DELETING TIME-RECORD FOR TIME 3489.6609
DELETING TIME-RECORD FOR TIME 4348.6544
DELETING TIME-RECORD FOR TIME 5207.6478
DELETING TIME-RECORD FOR TIME 6496.1380
DELETING TIME-RECORD FOR TIME 7355.1315
DELETING TIME-RECORD FOR TIME 8214.1250
DELETING TIME-RECORD FOR TIME 9073.1184
DELETING TIME-RECORD FOR TIME 9932.1119
DELETING TIME-RECORD FOR TIME 10791.105
DELETING TIME-RECORD FOR TIME 11650.099
DELETING TIME-RECORD FOR TIME 13368.086
DELETING TIME-RECORD FOR TIME 15086.073
DELETING TIME-RECORD FOR TIME 16804.060
DELETING TIME-RECORD FOR TIME 18522.046
DELETING TIME-RECORD FOR TIME 20240.033
DELETING TIME-RECORD FOR TIME 21958.020
DELETING TIME-RECORD FOR TIME 25393.994
DELETING TIME-RECORD FOR TIME 28829.968
DELETING TIME-RECORD FOR TIME 30547.955
DELETING TIME-RECORD FOR TIME 33983.929
DELETING TIME-RECORD FOR TIME 37419.903
DELETING TIME-RECORD FOR TIME 40855.876
DELETING TIME-RECORD FOR TIME 44291.850
DELETING TIME-RECORD FOR TIME 51163.798
DELETING TIME-RECORD FOR TIME 58035.746
DELETING TIME-RECORD FOR TIME 64907.693
DELETING TIME-RECORD FOR TIME 71779.641
DELETING TIME-RECORD FOR TIME 78651.589
DELETING TIME-RECORD FOR TIME 85523.536
DELETING TIME-RECORD FOR TIME 88959.510
DELETING TIME-RECORD FOR TIME 95831.458
DELETING TIME-RECORD FOR TIME 109575.35
DELETING TIME-RECORD FOR TIME 116447.30
DELETING TIME-RECORD FOR TIME 130191.20
DELETING TIME-RECORD FOR TIME 143935.09
DELETING TIME-RECORD FOR TIME 147371.07
DELETING TIME-RECORD FOR TIME 150807.04
DELETING TIME-RECORD FOR TIME 154243.01
DELETING TIME-RECORD FOR TIME 161114.96
DELETING TIME-RECORD FOR TIME 174858.86
DELETING TIME-RECORD FOR TIME 188602.75
DELETING TIME-RECORD FOR TIME 216090.54
DELETING TIME-RECORD FOR TIME 229834.44
DELETING TIME-RECORD FOR TIME 243578.33
DELETING TIME-RECORD FOR TIME 271066.12
DELETING TIME-RECORD FOR TIME 277938.07
DELETING TIME-RECORD FOR TIME 291681.97
DELETING TIME-RECORD FOR TIME 319169.76
DELETING TIME-RECORD FOR TIME 346657.55
DELETING TIME-RECORD FOR TIME 374145.34
DELETING TIME-RECORD FOR TIME 387889.23
DELETING TIME-RECORD FOR TIME 394761.18
DELETING TIME-RECORD FOR TIME 408505.08
DELETING TIME-RECORD FOR TIME 422248.97
DELETING TIME-RECORD FOR TIME 449736.76
DELETING TIME-RECORD FOR TIME 463480.66
DELETING TIME-RECORD FOR TIME 470352.61
DELETING TIME-RECORD FOR TIME 477224.55
DELETING TIME-RECORD FOR TIME 490968.45
DELETING TIME-RECORD FOR TIME 518456.24
DELETING TIME-RECORD FOR TIME 532200.14
DELETING TIME-RECORD FOR TIME 545944.03
DELETING TIME-RECORD FOR TIME 573431.82
DELETING TIME-RECORD FOR TIME 600919.61
DELETING TIME-RECORD FOR TIME 628407.40
DELETING TIME-RECORD FOR TIME 642151.30
DELETING TIME-RECORD FOR TIME 655895.19

```

```

KEEPING TIME-RECORD FOR TIME 683382.98
AND FOR TIME 720000.00

```

WORKSPACE RECLAIMED

INTERPOLATION SCHEME USED THIS FRACTION OF
THE ALLOCATED MEMORY: 0.174895104174184
EFFICIENCY FACTOR: 36.7070439139405

DEALLOCATING

TIMESTEP AT 720000.000 SELECTED

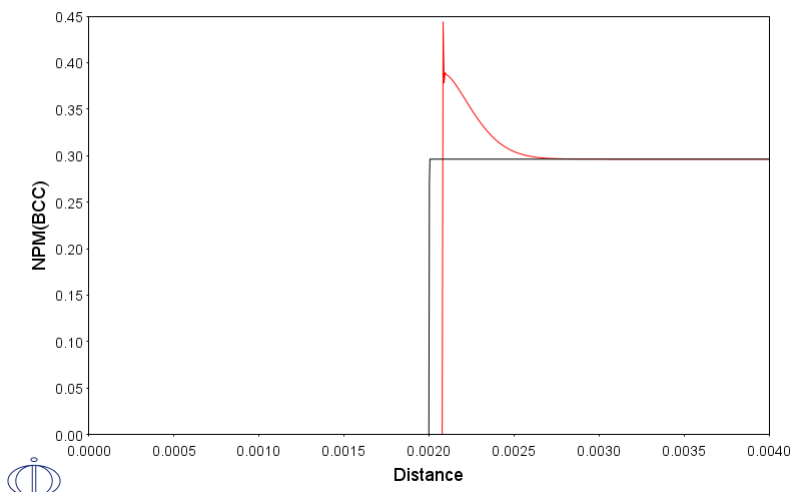
DIC>
DIC>
DIC>
DIC>
DIC> set-inter
--OK--
DIC>

exd2b-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd2b\plot.DCM"
DIC>
DIC>
DIC> @@ exd2_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE D2
DIC> @@
DIC> @@ ENTER THE DICTRA MODULE AND SPECIFY THE STORE-RESULT FILE
DIC> @@
DIC>
DIC>
DIC> go d-m
TIME STEP AT TIME 7.20000E+05
DIC> read exd2
OK
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1:
POST-1: @@
POST-1: @@ LET US FIRST SEE HOW THE FRACTION OF FERRITE HAS CHANGED
POST-1: @@ AS A RESULT OF THE DIFFUSION
POST-1: @@
POST-1: s-d-a y npm(bcc)
POST-1: s-d-a x distance global
INFO: Distance is set as independent variable
POST-1: s-p-c time 0 720000
POST-1: set-tit Figure D2.1
POST-1: plot
```

Figure D2.1

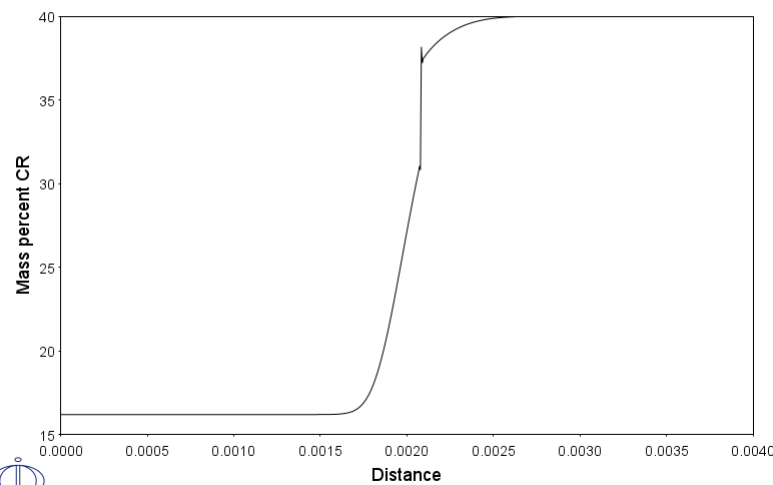
2021.05.13.12.51.20
Time = 0.720000
CELL #1



```
POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: @@
POST-1: @@ LET US NOW PLOT HOW THE AVERAGE CR-CONCENTRATION VARIES WITH DISTANCE
POST-1: @@
POST-1: s-d-a y w-p cr
POST-1: s-p-c time last
POST-1: set-tit Figure D2.2
POST-1: plot
```

Figure D2.2

2021.05.13.12.51.22
Time = 720000
CELL #1



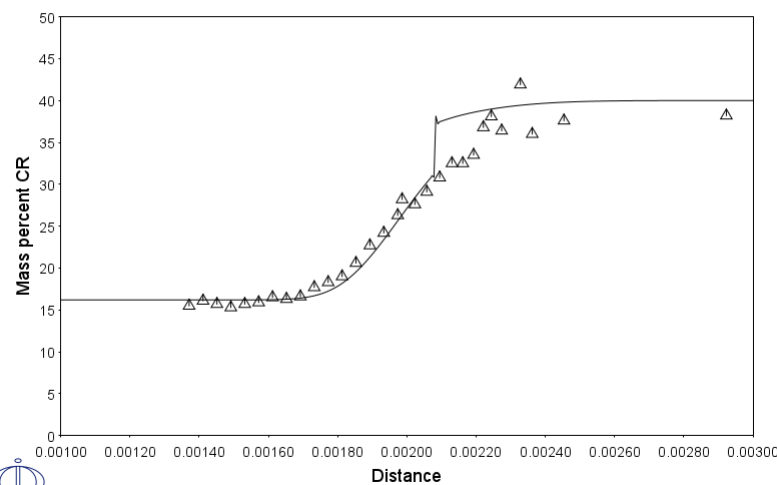
```

POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: @@
POST-1: @@ WE SELECT A BETTER SCALING AND APPEND EXPERIMENTAL DATA
POST-1: @@
POST-1: app y exd2.exp
PROLOGUE NUMBER: /0/: 1
DATASET NUMBER(s): /-1/: 4
POST-1:
POST-1: s-s-s y n 0 50
POST-1: s-s-s x n 10e-4 30e-4
POST-1:
POST-1: set-tit Figure D2.3
POST-1: plot

```

Figure D2.3

2021.05.13.12.51.25
Time = 720000
CELL #1



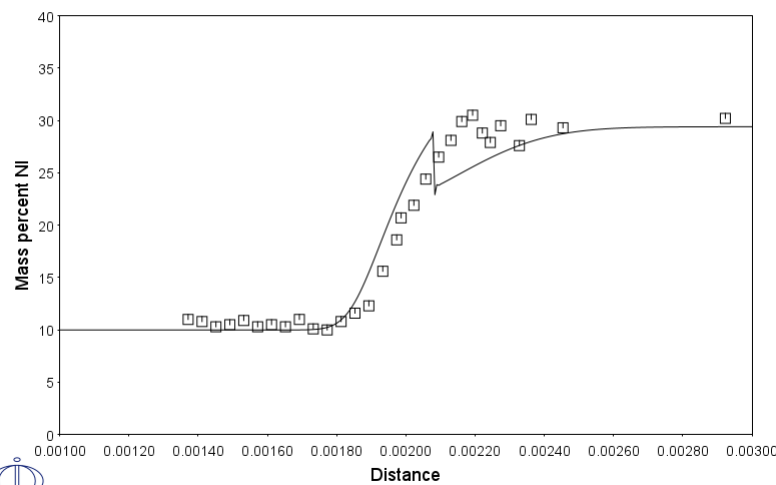
```

POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: @@
POST-1: @@ NOW WE DO THE SAME FOR NICKEL
POST-1: @@
POST-1:
POST-1: s-d-a y w-p ni
POST-1:
POST-1: app y exd2.exp
PROLOGUE NUMBER: /1/: 1
DATASET NUMBER(s): /-1/: 5
POST-1:
POST-1: s-s-s x n 10e-4 30e-4
POST-1: s-s-s y n 0 40
POST-1:
POST-1: set-tit Figure D2.4
POST-1: plot

```

Figure D2.4

2021.05.13.12.51.27
Time = 720000
CELL #1



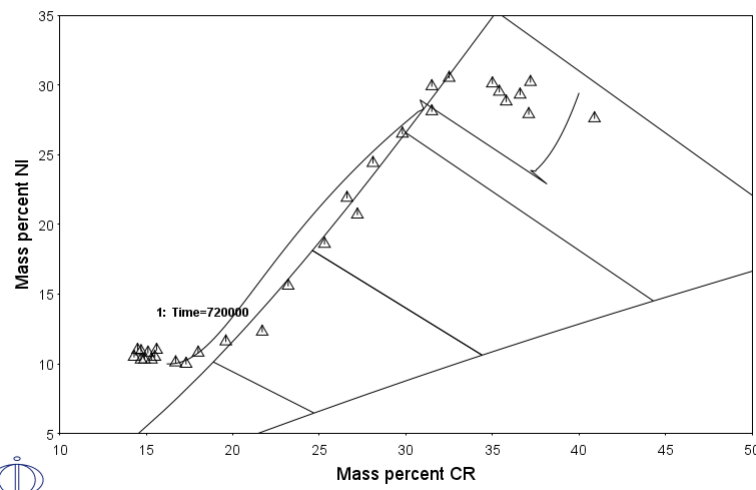
```

POST-1:
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: @@
POST-1: @@ LET US PLOT THE DIFFUSION PATH FOR THE COUPLE
POST-1: @@ WE APPEND THE TERNARY PHASE-DIAGRAM CALCULATED IN THERMO-CALC
POST-1: @@ AND THE EXPERIMENTAL DATA
POST-1: @@
POST-1: s-d-a x w-p cr
POST-1: s-d-a y w-p ni
POST-1: s-i-v dis gl
POST-1: s-p-c time last
POST-1:
POST-1: app y exd2.exp
PROLOGUE NUMBER: /1/: 1
DATASET NUMBER(s): /-1/: 6 7 8
POST-1:
POST-1: s-s-s x n 10 50
POST-1: s-s-s y n 5 35
POST-1:
POST-1: set-tit Figure D2.5
POST-1: plot

```

Figure D2.5

2021.05.13.12.51.29
Time = 720000
CELL #1



```

POST-1:
POST-1:
POST-1:
POST-1:
POST-1: set-interactive
--OK--
POST-1:

```

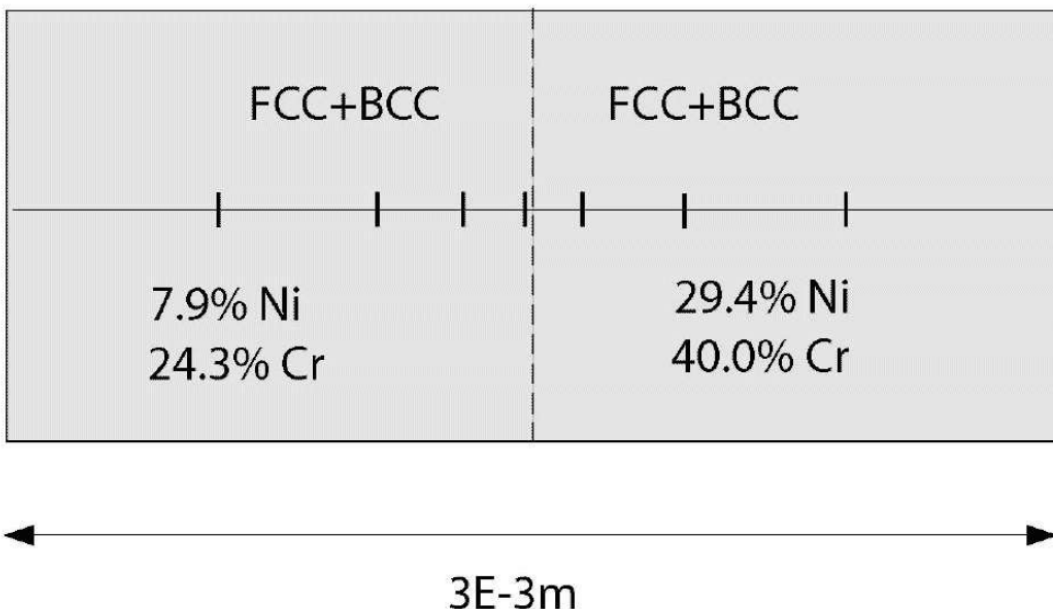


Example exd3

Diffusion couple of Fe-Ni-Cr alloys: Homogenization Model

This example shows the use of the homogenization model. It is taken from H. Larsson and A. Engström, Acta. Mater. v.54 (2006), pp. 2431-2439. Experimental data from A. Engström, Scand J Metall, v.243 (1995), p.12. The homogenization model can be used for multiphase simulations like the dispersed system model, but unlike the dispersed system model there is no need to have a single continuous matrix phase and, furthermore, there is no need to limit the size of time-steps. The set-up is performed in the same manner as for the dispersed system model, which means that a certain phase is entered as the matrix phase and the other phases are entered as spheroidal, but the choice of matrix phase will not affect the simulation.

$T = 1373 \text{ K}$



exd3-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd3\setup.DCM.test"

SYS: i>@
NO SUCH COMMAND, USE HELP
SYS: @@ Diffusion in dispersed systems.
SYS: @@ Diffusion couple of Fe-Cr-Ni alloys: Homogenization model
SYS: @@ This example uses the homogenization model. It is taken from
SYS: @@ H. Larsson and A. Engström, Acta. Mater. v.54 (2006), pp. 2431-2439.
SYS: @@ Experimental data from A. Engström, Scand J Metall, v.243 (1995), p.12.
SYS: @@ The homogenization model can be used for multiphase simulations
SYS: @@ like the dispersed system model, but unlike the dispersed system model
SYS: @@ there is no need to have a single continuous matrix phase and, furthermore,
SYS: @@ there is no need to limit the size of time-steps.
SYS: @@ The set-up is performed in the same manner as for the dispersed system
SYS: @@ model, which means that a certain phase is entered as the matrix phase
SYS: @@ and the other phases are entered as spheroidal, but the choice of matrix
SYS: @@ phase will not affect the simulation.
SYS: -----
NO SUCH COMMAND, USE HELP

SYS:
SYS: @@ exd3_setup.DCM
SYS:
SYS: go da

THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED

DICTRA_FCC_A1 REJECTED
TDB_TCFE11: sw fedemo

Current database: Iron Demo Database v4.0

VA /- DEFINED

TDB_FEDEMO: def-sys fe cr ni
FE CR NI
DEFINED

TDB_FEDEMO: rej-ph *
LIQUID:L BCC_A2 LAVES_PHASE_C14
CBCC_A12 CHI_A12 CUB_A13
FCC_A1 HCP_A3 SIGMA
REJECTED

TDB_FEDEMO: rest-ph bcc,fcc
BCC_A2 FCC_A1 RESTORED

TDB_FEDEMO: get

12:52:46,283 INFO *** Invoking Gibbs Energy System v6 ***

REINITIATING GES

ELEMENTS

SPECIES

PHASES

PARAMETERS ...

FUNCTIONS

List of references for assessed data

'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'J-O. Andersson and B. Sundman, CALPHAD, 11 (1987) 83-92; TRITA 0270
(1986); CR-FE'
'B.J. Lee, CALPHAD, 17 (1993) 251-268; revision of Fe-Cr and Fe -Ni liquid'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); CR -NI'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'

-OK-

TDB_FEDEMO:

TDB_FEDEMO: app mfdemo

Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys fe cr ni
FE CR NI
DEFINED

APP: rej-ph *
BCC_A2 FCC_A1 REJECTED
APP: rest-ph bcc,fcc
BCC_A2 FCC_A1 RESTORED

APP: get

ELEMENTS

SPECIES

PHASES

PARAMETERS ...

FUNCTIONS

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Fe diffusion fcc Cr-Fe'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Cr and Ni diffusion fcc Cr-Ni'
'B. Jonsson: Z. Metallkunde 86(1995)686-692; Cr, Fe and Ni diffusion fcc
Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
'B. Jonsson: ISIJ International, 35(1995)1415-1421; Cr, Fe and Ni
diffusion bcc Cr-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion

```

    in bcc Fe'
-OK-
APP:
APP: go -m
NO TIME STEP DEFINED
DIC>
DIC> set-cond glob T 0 1373.15; * N
DIC>
DIC> ent-geo 0
DIC>
DIC> ent-reg
REGION NAME : fecrni
DIC>
DIC> ent-grid
REGION NAME : /FECRNI/: fecrni
WIDTH OF REGION /1/: 3e-3
TYPE /LINEAR/: double
NUMBER OF POINTS /50/: 60
VALUE OF R IN THE GEOMETRICAL SERIE FOR LOWER PART OF REGION: 0.85
VALUE OF R IN THE GEOMETRICAL SERIE FOR UPPER PART OF REGION: 1.15
DIC>
DIC> ent-ph
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /FECRNI/: fecrni
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC> ent-ph
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /FECRNI/: fecrni
PHASE TYPE /MATRIX/: sph
PHASE NAME: /NONE/: bcc
INFO: EQUILIBRIUM COMPOSITION AND FRACTION OF SPHEROID PHASES USED AS DEFAULT
DIC>
DIC> ent-comp
REGION NAME : /FECRNI/: fecrni
PHASE NAME: /FCC_A1/: fcc#1
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: m-f
PROFILE FOR /CR/: cr
TYPE /LINEAR/: read cr.dat
PROFILE FOR /NI/: ni
TYPE /LINEAR/: read ni.dat
DIC>
DIC> ent-comp
REGION NAME : /FECRNI/: fecrni
PHASE NAME: /FCC_A1/: bcc
USE EQUILIBRIUM VALUE /Y/: y
DIC>
DIC> se-si-ti
END TIME FOR INTEGRATION /.1/: 3.6e5
AUTOMATIC TIMESTEP CONTROL /YES/: yes
MAX TIMESTEP DURING INTEGRATION /36000/: 3.6e4
INITIAL TIMESTEP : /1E-07/: 1
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/: 1e-7
DIC>
DIC> @@ SIMULATIONS ARE FASTER WHEN THE RESULTS ARE NOT SAVED
DIC> @@ FOR EVERY TIME STEP
DIC> s-s-c
NS01A PRINT CONTROL : /0/: 0
FLUX CORRECTION FACTOR : /1/: 1
NUMBER OF DELTA TIMESTEPS IN CALLING MULDIF: /2/: 2
CHECK INTERFACE POSITION /AUTO/: n
VARY POTENTIALS OR ACTIVITIES OR LNAC : /ACTIVITIES/: act
ALLOW AUTOMATIC SWITCHING OF VARYING ELEMENT : /YES/: y
SAVE WORKSPACE ON FILE (YES,NO,0-999) /YES/: 99
DEGREE OF IMPLICITY WHEN INTEGRATING PDES (AUTO, 0 -> 0.5 -> 1): /AUTO/: .5
MAX TIMESTEP CHANGE PER TIMESTEP : /2/: 2
USE FORCED STARTING VALUES IN EQUILIBRIUM CALCULATION /AUTO/: n
ALWAYS CALCULATE STIFFNES MATRIX IN MULDIF /YES/: y
CALCULATE RESIDUAL FOR DEPENDENT COMPONENT /NO/:
DIC>
DIC> @@ There are a several options available for the homogenization
DIC> @@ model. There is also an interpolation scheme that may reduce
DIC> @@ simulation times. However, for this example, the default settings
DIC> @@ are kept and the interpolation scheme is turned off.
DIC> @@
DIC> homogen yes yes
INFO: HOMOGENIZATION MODEL ENABLED
DIC>
DIC> @@ There are a large number of homogenization functions
DIC> @@ available. These determine how the average kinetics
DIC> @@ of the multiphase mixture is evaluated. For this example,
DIC> @@ the General lower Hashin-Shtrikman bound is a good choice.
DIC> en-ho
ENTER HOMOGENIZATION FUNCTION # /5/: 1
SELECTED FUNCTION IS HASHIN-SHTRIKMAN BOUND: GENERAL LOWER
DIC>
DIC>
DIC>
DIC> save exd3 Y
DIC>
DIC>
DIC> set-inter
--OK--
DIC>
```

exd3-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd3\run.DCM.test"
DIC>
DIC>
DIC> @@ exd3_run.DCM
DIC>
DIC> @@
DIC> @@ READ THE SETUP FROM FILE AND START THE SIMULATION
DIC> @@
DIC>
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exd3
OK
DIC>
DIC> sim
STARTING SIMULATION USING HOMOGENIZATION MODEL
-----
WARNING:BCC_A2 HAS NO VOLUME FRACTION, CREATING ONE
Starting time-step t0= 0.0000000 dt= 1.0000000
Starting time-step t0= 1.0000000 dt= 2.0000000
Starting time-step t0= 3.0000000 dt= 4.0000000
Starting time-step t0= 7.0000000 dt= 8.0000000
Starting time-step t0= 15.0000000 dt= 16.0000000
Starting time-step t0= 31.0000000 dt= 32.0000000
Starting time-step t0= 63.0000000 dt= 64.0000000
Starting time-step t0= 127.0000000 dt= 128.0000000
Starting time-step t0= 255.0000000 dt= 256.0000000
Starting time-step t0= 511.0000000 dt= 256.0000000
Starting time-step t0= 767.0000000 dt= 256.0000000
Starting time-step t0= 1023.0000000 dt= 512.0000000
Starting time-step t0= 1535.0000000 dt= 512.0000000
Starting time-step t0= 2047.0000000 dt= 1024.0000000
Starting time-step t0= 3071.0000000 dt= 1024.0000000
Starting time-step t0= 4095.0000000 dt= 2048.0000000
Starting time-step t0= 6143.0000000 dt= 2048.0000000
Starting time-step t0= 8191.0000000 dt= 2048.0000000
Starting time-step t0= 10239.0000000 dt= 2048.0000000
Starting time-step t0= 12287.0000000 dt= 4096.0000000
Starting time-step t0= 16383.0000000 dt= 8192.0000000
Starting time-step t0= 24575.0000000 dt= 8192.0000000
Starting time-step t0= 32767.0000000 dt= 8192.0000000
Starting time-step t0= 40959.0000000 dt= 8192.0000000
Starting time-step t0= 49151.0000000 dt= 16384.0000000
Starting time-step t0= 65535.0000000 dt= 16384.0000000
Starting time-step t0= 81919.0000000 dt= 16384.0000000
Starting time-step t0= 98303.0000000 dt= 16384.0000000
Starting time-step t0= 114687.0000000 dt= 16384.0000000
Starting time-step t0= 131071.0000000 dt= 16384.0000000
Starting time-step t0= 147455.0000000 dt= 16384.0000000
Starting time-step t0= 163839.0000000 dt= 16384.0000000
Starting time-step t0= 180223.0000000 dt= 16384.0000000
Starting time-step t0= 196607.0000000 dt= 16384.0000000
Starting time-step t0= 212991.0000000 dt= 16384.0000000
Starting time-step t0= 229375.0000000 dt= 16384.0000000
Starting time-step t0= 245759.0000000 dt= 16384.0000000
Starting time-step t0= 262143.0000000 dt= 16384.0000000
Starting time-step t0= 278527.0000000 dt= 16384.0000000
Starting time-step t0= 294911.0000000 dt= 16384.0000000
Starting time-step t0= 311295.0000000 dt= 16384.0000000
Starting time-step t0= 327679.0000000 dt= 16384.0000000
Starting time-step t0= 344063.0000000 dt= 15937.0000000
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 1.0000000
DELETING TIME-RECORD FOR TIME 511.0000000
DELETING TIME-RECORD FOR TIME 1023.0000000
DELETING TIME-RECORD FOR TIME 1535.0000000
DELETING TIME-RECORD FOR TIME 2047.0000000
DELETING TIME-RECORD FOR TIME 3071.0000000
DELETING TIME-RECORD FOR TIME 4095.0000000
DELETING TIME-RECORD FOR TIME 6143.0000000
DELETING TIME-RECORD FOR TIME 8191.0000000
DELETING TIME-RECORD FOR TIME 10239.0000000
DELETING TIME-RECORD FOR TIME 12287.0000000
DELETING TIME-RECORD FOR TIME 16383.0000000
DELETING TIME-RECORD FOR TIME 24575.0000000
DELETING TIME-RECORD FOR TIME 32767.0000000
DELETING TIME-RECORD FOR TIME 40959.0000000
DELETING TIME-RECORD FOR TIME 49151.0000000
DELETING TIME-RECORD FOR TIME 65535.0000000
DELETING TIME-RECORD FOR TIME 81919.0000000
DELETING TIME-RECORD FOR TIME 98303.0000000
DELETING TIME-RECORD FOR TIME 114687.0000000
DELETING TIME-RECORD FOR TIME 131071.0000000
DELETING TIME-RECORD FOR TIME 147455.0000000
DELETING TIME-RECORD FOR TIME 163839.0000000
DELETING TIME-RECORD FOR TIME 180223.0000000
DELETING TIME-RECORD FOR TIME 196607.0000000
DELETING TIME-RECORD FOR TIME 212991.0000000
DELETING TIME-RECORD FOR TIME 229375.0000000
DELETING TIME-RECORD FOR TIME 245759.0000000
DELETING TIME-RECORD FOR TIME 262143.0000000
DELETING TIME-RECORD FOR TIME 278527.0000000
DELETING TIME-RECORD FOR TIME 294911.0000000
DELETING TIME-RECORD FOR TIME 311295.0000000
DELETING TIME-RECORD FOR TIME 327679.0000000
KEEPING TIME-RECORD FOR TIME 344063.0000000
AND FOR TIME 360000.0000000
WORKSPACE RECLAIMED
-----
INTERPOLATION SCHEME USED THIS FRACTION OF
THE ALLOCATED MEMORY: 0.146664786348893
EFFICIENCY FACTOR: 7.09082167832168
-----
DEALLOCATING
-----
```

TIMESTEP AT 360000.000 SELECTED

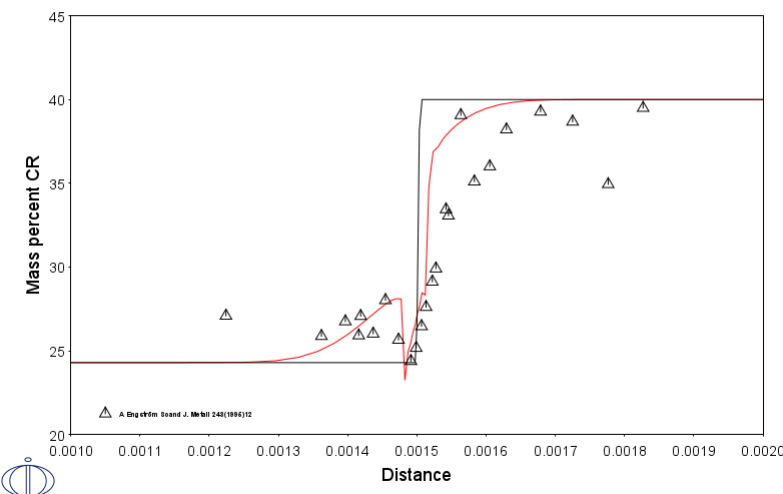
DIC>
DIC>
DIC> set-inter
---OK---
DIC>

exd3-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exd3\plot.DCM.test"
DIC>
DIC>
DIC> @@ exd3_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE d3
DIC> @@
DIC> @@ ENTER THE DICTRA MODULE AND SPECIFY THE STORE-RESULT FILE
DIC> @@
DIC>
DIC>
DIC> go d-m
TIME STEP AT TIME 3.60000E+05
DIC> read exd3
OK
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1:
POST-1: @@
POST-1: @@ First study the composition profiles of Cr and Ni
POST-1: @@
POST-1: s-d-a x distance global
INFO: Distance is set as independent variable
POST-1: s-d-a y w-p Cr
POST-1: s-p-c time 0 360000
POST-1: set-tit Figure D3.1
POST-1:
POST-1: app yes k5k7cr.exp 0; 1
POST-1:
POST-1: s-s-s x n 1e-3 2e-3
POST-1:
POST-1: s-s-s y n 20 45
POST-1:
POST-1: SET_EXP_FILE_FORMAT 3
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Figure D3.1

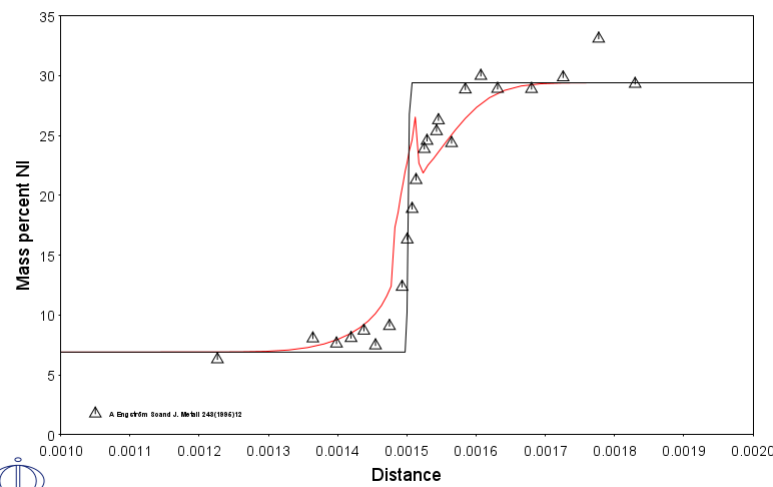
2021.05.13.12.54.49
Time = 0.360000
CELL #1



```
POST-1:
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: s-d-a y w-p Ni
POST-1: set-tit Figure D3.2
POST-1:
POST-1: app yes k5k7ni.exp 0; 1
POST-1:
POST-1: s-s-s y n 0 35
POST-1:
POST-1: SET_EXP_FILE_FORMAT 3
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Figure D3.2

2021.05.13.12.54.49
Time = 0.360000
CELL #1



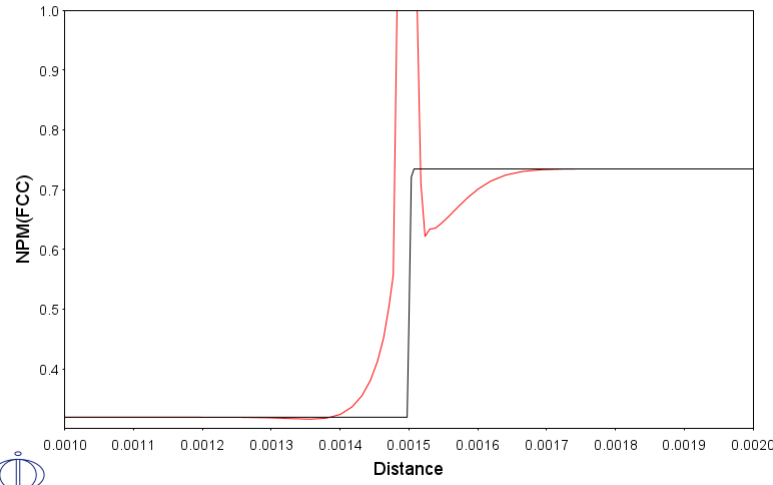
```

POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: @@
POST-1: @@ Then study the amount of FCC and BCC
POST-1: @@
POST-1: app no
POST-1: s-d-a y npm(fcc)
POST-1: set-tit Figure D3.3
POST-1:
POST-1: SET_EXP_FILE_FORMAT 3
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

```

Figure D3.3

2021.05.13.12.54.51
Time = 0.360000
CELL #1



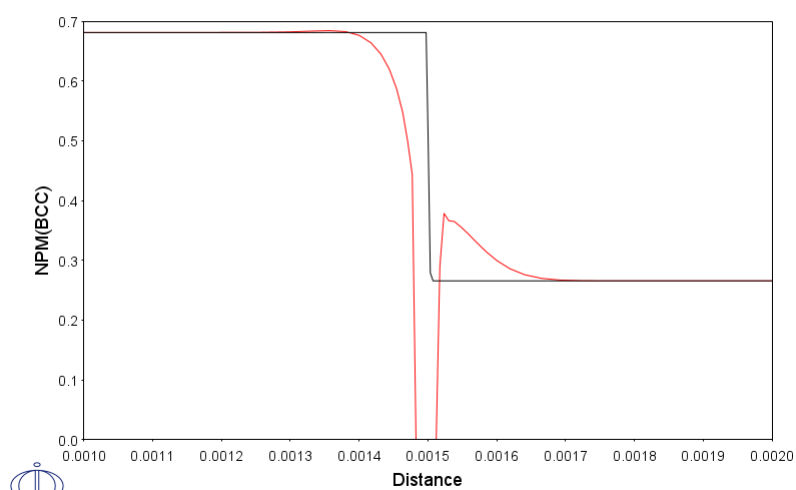
```

POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1: s-d-a y npm(bcc)
POST-1: set-tit Figure D3.4
POST-1:
POST-1: SET_EXP_FILE_FORMAT 3
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot

```

Figure D3.4

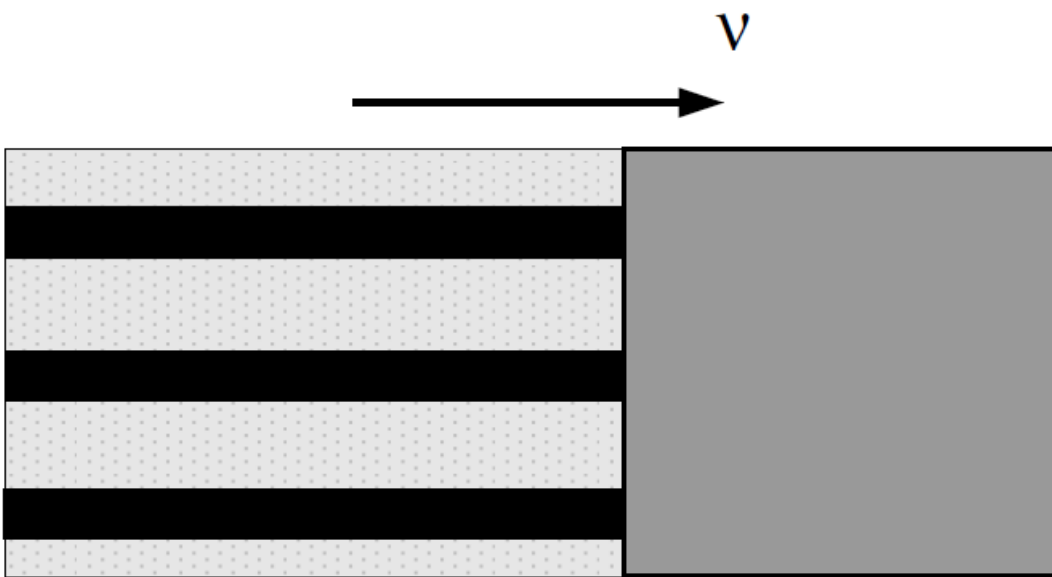
2021.05.13.12.54.52
Time = 0.360000
CELL #1



POST-1:
POST-1:
POST-1: set-interactive
--OK---
POST-1:



Cooperative Growth



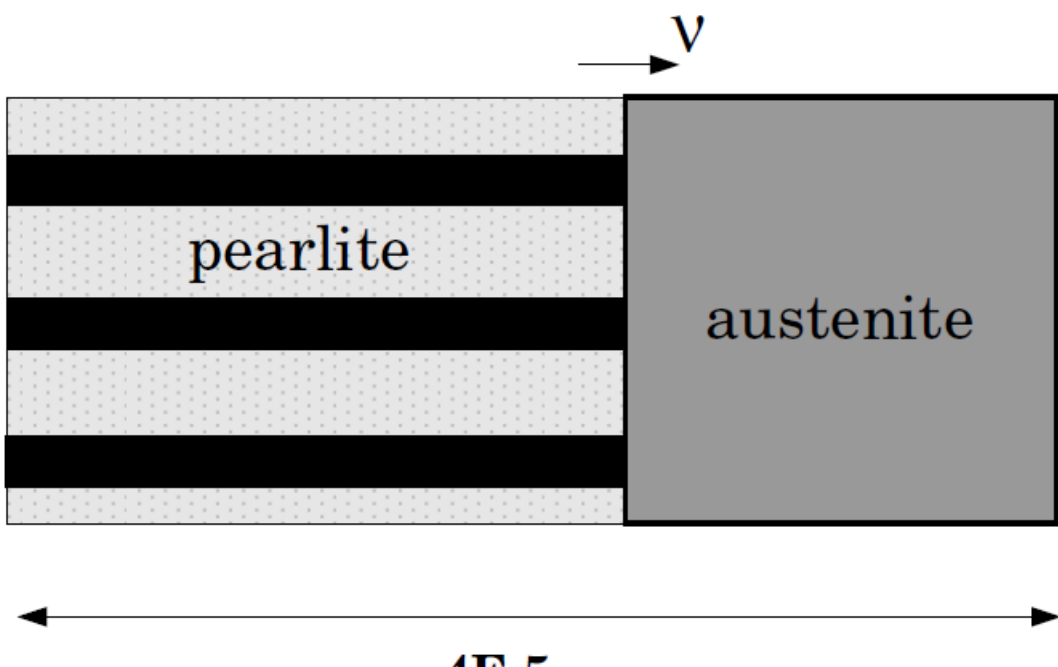


Example exe1

Growth of pearlite in an Fe-Mn-C alloy

This is an example of pearlite growth in an Fe-0.50wt%C - 0.91wt%Mn steel.

$$T = 900 - \text{Time} * 10$$




```

FE4N_LP1           HCP_A3          LIQUID:L
REJECTED
APP: rest-ph bcc,fcc,cem
BCC_A2           FCC_A1          CEMENTITE
RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'Y. Liu, L. Zhang, et al. CALPHAD 33(2009)614-23; Fe-Mn-C (fcc)'
'B. Jonsson: Z. Metallkunde 85(1994)498-501; C and N diffusion in bcc Cr
-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'
'S. Deng, et al., CALPHAD, 56 (2017) 230-240.'
'Assessed from data presented in Landolt-Bornstein, Vol. 26, ed. H.
Mehrer, Springer (1990); Impurity diff of Mn in bcc Fe.'

-OK-
APP:
APP: @@
APP: @@ ALL THE THERMODYNAMIC AND KINETIC DATA HAVE BEEN RETRIEVED.
APP: @@ GO TO THE DICTRA MONITOR TO SET UP YOUR PROBLEM.
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ SET THE CONDITION FOR TEMPERATURE
DIC> @@
DIC> set-cond glob t 0 900-time*10; * n
DIC>
DIC> @@
DIC> @@ ENTER A REGION
DIC> @@
DIC> enter-reg pearlite
DIC>
DIC> @@
DIC> @@ ENTER A SMALL INITIAL SIZE OF THE GRID IN THE 'PEARLITE' REGION
DIC> @@
DIC> enter-grid pearlite 5e-10 lin 5
DIC>
DIC> @@
DIC> @@ ENTER INTO THE 'PEARLITE' REGION THE PHASES 'BCC' AND 'CEM' AND SPECIFY
DIC> @@ THAT ARE PRESENT IN THE FORM OF A 'LAMELLAR AGGREGATE. SET THE STATUS
DIC> @@ TO 'ACTIVE'. SEVERAL PROMPTS FOLLOW ABOUT THE VALUES OF THE PARAMETERS
DIC> @@ IN THE PEARLITE GROWTH MODEL, FOR EXAMPLE AS SURFACE TENSION, OPTIMUM
DIC> @@ GROWTH RATE FACTOR, AND BOUNDARY DIFFUSION COEFFICIENTS.
DIC> @@
DIC> @@ CARBON(C) IS TREATED IN A SPECIAL WAY. IF 'AUTOMATIC' IS ENTERED
DIC> @@ THE DIFFUSION OF C IS CALCULATED ACCORDING TO AN EQUATION FOR
DIC> @@ MIXED BOUNDARY AND VOLUME DIFFUSION. YOU CAN CHOOSE BETWEEN
DIC> @@ MANUAL OR AUTOMATIC START VALUES FOR ALL VARIABLES EXCEPT THE GROWTH
DIC> @@ RATE. IN THIS EXAMPLE WE WILL TRY 1E-6
DIC> @@
DIC> @@ FOR MORE INFORMATION ABOUT THE PEARLITE GROWTH MODEL SEE
DIC> @@ B. JA-NSSON: TRITA-MAC-0478, 1992 (ROYAL INSTITUTE OF TECHNOLOGY)
DIC> @@ STOCKHOLM, SWEDEN, 1992.
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /PEARLITE/: pearlite
PHASE TYPE /MATRIX/: lam
    Eutectoid reaction is "GAMMA" ==> "ALPHA" + "BETA"
Enter name of "ALPHA" phase /BCC_A2/: bcc_a2
Enter name of "BETA" phase /CEMENTITE/: cementite
Enter name of "GAMMA" phase /FCC_A1/: fcc_al
Enter "ALPHA"/"BETA" surface tension
LOW TIME LIMIT /0/: 0
Surface tension(T,P,TIME)= 1;
HIGH TIME LIMIT /*/: 1000
ANY MORE RANGES /N/: N
Enter "ALPHA"/"GAMMA" surface tension
LOW TIME LIMIT /0/: 0
Surface tension(T,P,TIME)= 1;
HIGH TIME LIMIT /*/: 1000
ANY MORE RANGES /N/: N
Optimum growth condition factor /2/: 2
Name of dependent element /FE/: fe
INPUT OF DIFFUSION DATA
Growth model (VOLUME/BOUNDARY/KIRKALDY) for element C /BOUNDARY/: boundary
DF(C) = /value/AUTOMATIC/MIXED/: auto
Growth model (VOLUME/BOUNDARY/KIRKALDY) for element MN /BOUNDARY/: boundary
DF(MN) = /value/MIXED/: 5.4e-14
DQ(MN): 155000
Automatic start values for the S0 determination /Y/: Y
Growth rate V: 1E-6
Automatic start values on other variables /Y/: Y
DIC>
DIC> @@
DIC> @@ INITIATE THE COMPOSITION RECORDS FOR THE 'PEARLITE'
DIC> @@
DIC> enter-composition
REGION NAME : /PEARLITE/: pearlite
DIC>
DIC> @@
DIC> @@ NOW CONTINUE BY DEFINING A MATRIX PHASE INTO WHICH THE PEARLITE
DIC> @@ WILL GROW. START BY ENTERING A REGION NAME, 'AUSTENITE'

```

```

DIC> @@
DIC> enter-region austenite
ATTACH TO REGION NAMED /PEARLITE/:
ATTACHED TO THE RIGHT OF PEARLITE /YES/:
DIC> @@
DIC> @@ SPECIFY WHAT PHASE 'FCC' WILL BE PRESENT IN THE 'AUSTENITE' REGION
DIC> @@ AND WHAT TYPE OF PHASE 'MATRIX' IT IS AND ITS INITIAL STATE 'ACTIVE'
DIC> @@
DIC> enter-phase act austenite matrix fcc
DIC>
DIC> @@
DIC> @@ WE ALSO NEED TO HAVE A SPATIAL GRID IN THE 'AUSTENITE' REGION.
DIC> @@ CHOSE SIZE '4E-5' GRIDTYPE 'GEOMETRICAL', '30' GRIDPOINTS AND '1.5'
DIC> @@ AS VALUES FOR THE GEOMETRICAL FACTOR OF THE GRID.
DIC> @@
DIC> enter-grid austenite 4e-5 geo 30 1.5
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL CONCENTRATION PROFILES IN THE 'FCC' PHASE OF THE
DIC> @@ 'AUSTENITE' REGION. CONCENTRATIONS MUST BE GIVEN IN Y-FRACTIONS.
DIC> @@
DIC> enter-composition
REGION NAME : /AUSTENITE/: austenite
PHASE NAME: /FCC_A1/: fcc
DEPENDENT COMPONENT ? /MN/: fe
COMPOSITION TYPE /MOLE_FRACTION/: site-fraction
PROFILE FOR MN
TYPE /LINEAR/: lin 9.29232973E-3 9.29232973E-3
PROFILE FOR C
TYPE /LINEAR/: lin 2.3384332E-2 2.3384332E-2
DIC>
DIC> @@
DIC> @@ THE MATRIX PHASE IS NOW COMPLETE.
DIC> @@
DIC>
DIC> @@
DIC> @@ SPECIFY A SPHERICAL '2' GEOMETRY
DIC> @@
DIC> enter-geo 2
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 5
AUTOMATIC TIMESTEP CONTROL /YES/: YES
MAX TIMESTEP DURING INTEGRATION /.5/: 0.1
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exel Y
DIC>
DIC> set-inter
--OK---
DIC>

```

exe1-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exe1\run.DCM.test"
DIC>
DIC>
DIC> @@ exe1_run.DCM
DIC>
DIC> @@
DIC> FILE FOR RUNNING EXAMPLE e1
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR
DIC> @@
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC>
DIC> @@
DIC> @@ READ SETUP FROM FILE
DIC> @@
DIC> read exe1
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> simulate
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
Trying old scheme 4
U-FRACTION IN SYSTEM: C = .0233843320030518 FE = .990707670399293
MN = .0092923297312127
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
U-FRACTION IN SYSTEM: C = .0233843320030518 FE = .990707670399293
MN = .0092923297312127
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
17 GRIDPOINT(S) ADDED TO CELL #1 REGION: AUSTENITE
TIME = 0.10000000E-06 DT = 0.10000000E-06 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.77954867E-05 AND 0.77954867E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.50077955E-09
U-FRACTION IN SYSTEM: C = .0233843320030518 FE = .990707670399294
MN = .0092923297312127
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
10 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

CPU time used in timestep 3 seconds
4 GRIDPOINT(S) ADDED TO CELL #1 REGION: AUSTENITE
TIME = 0.10010000E-03 DT = 0.10000000E-03 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.77957274E-05 AND 0.77957274E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.12803523E-08
U-FRACTION IN SYSTEM: C = .0233843320030518 FE = .990707670399294
MN = .0092923297312127
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
10 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE

CPU time used in timestep 1 seconds
1 GRIDPOINT(S) REMOVED FROM CELL# 1 REGION # 1
TIME = 0.13027597E-02 DT = 0.12026597E-02 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.77986225E-05 AND 0.77986225E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.10659441E-07
U-FRACTION IN SYSTEM: C = .0233843320030518 FE = .990707670399295
MN = .0092923297312127
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
CPU time used in timestep 0 seconds
2 GRIDPOINT(S) REMOVED FROM CELL# 1 REGION # 1
TIME = 0.37080790E-02 DT = 0.24053193E-02 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.78044138E-05 AND 0.78044138E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.29431548E-07
U-FRACTION IN SYSTEM: C = .0233843320030517 FE = .990707670399294
MN = .0092923297312127
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
CPU time used in timestep 0 seconds
1 GRIDPOINT(S) REMOVED FROM CELL# 1 REGION # 1
TIME = 0.85187177E-02 DT = 0.48106387E-02 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.78160011E-05 AND 0.78160011E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.67031506E-07
U-FRACTION IN SYSTEM: C = .0233843320030517 FE = .990707670399294
MN = .0092923297312127
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
CPU time used in timestep 0 seconds
1 GRIDPOINT(S) REMOVED FROM CELL# 1 REGION # 1
TIME = 0.18139995E-01 DT = 0.96212773E-02 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.78391945E-05 AND 0.78391945E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.14245457E-06
U-FRACTION IN SYSTEM: C = .0233843320030517 FE = .990707670399294
MN = .00929232973121269
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
CPU time used in timestep 0 seconds
1 GRIDPOINT(S) REMOVED FROM CELL# 1 REGION # 1
TIME = 0.37382550E-01 DT = 0.19242555E-01 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.78856562E-05 AND 0.78856562E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.29419474E-06
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399294
MN = .00929232973121268
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
CPU time used in timestep 0 seconds
1 GRIDPOINT(S) REMOVED FROM CELL# 1 REGION # 1
TIME = 0.75867659E-01 DT = 0.38485109E-01 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.79788762E-05 AND 0.79788762E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.60126266E-06
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399294
MN = .00929232973121268
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
CPU time used in timestep 0 seconds
1 GRIDPOINT(S) REMOVED FROM CELL# 1 REGION # 1
TIME = 0.15283788 DT = 0.76970219E-01 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.81664857E-05 AND 0.81664857E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.12298389E-05
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399294
MN = .00929232973121267
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
CPU time used in timestep 0 seconds
TIME = 0.25283788 DT = 0.10000000 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.84125012E-05 AND 0.84125012E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.20710890E-05
U-FRACTION IN SYSTEM: C = .0233843320030515 FE = .990707670399294
```

```

        MN = .00929232973121267
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
CPU time used in timestep          1 seconds
1 GRIDPOINT(S) REMOVED FROM CELL# 1 REGION # 1
TIME = 0.35283788 DT = 0.10000000 SUM OF SQUARES = 0.0000000
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.86610019E-05 AND 0.86610019E-05
POSITION OF INTERFACE PEARLITE / AUSTENITE IS 0.29371892E-05
U-FRACTION IN SYSTEM: C = .0233843320030515 FE = .990707670399295
MN = .00929232973121272
TOTAL SIZE OF SYSTEM: 2.68092626329E-13 [m^3]
CPU time used in timestep          0 seconds

output ignored...

... output resumed

WORKSPACE RECLAIMED
INFO: CELL 1 REGION AUSTENITE DELETED
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
TIME = 3.2743323 DT = 0.10000000E-04 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 3.2843323 DT = 0.10000000E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 3.3843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 3.4843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 3.5843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 3.6843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 3.7843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 3.8843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 3.9843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 4.0843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 4.1843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 4.2843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 4.3843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 4.4843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 4.5843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 4.6843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 4.7843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep          0 seconds
TIME = 4.8843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
```

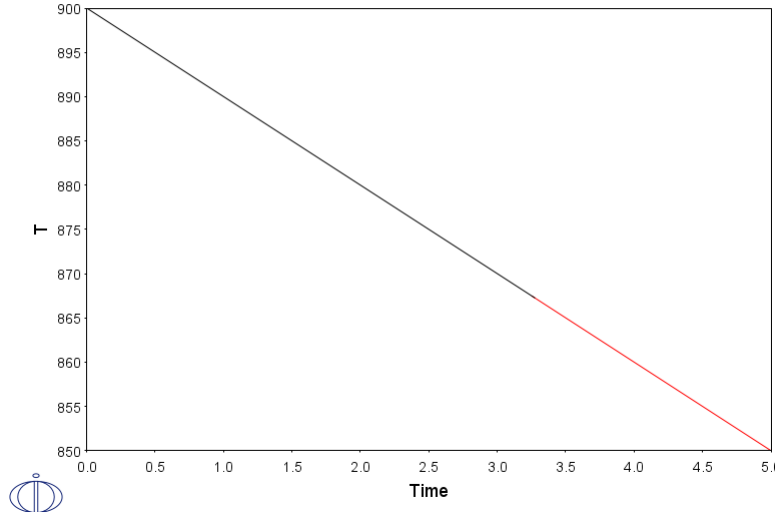
CPU time used in timestep 0 seconds
TIME = 4.9843323 DT = 0.10000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
CPU time used in timestep 0 seconds
TIME = 5.0000000 DT = 0.15667747E-01 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0233843320030516 FE = .990707670399324
MN = .00929232973118252
TOTAL SIZE OF SYSTEM: 2.68072664408E-13 [m^3]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 3.2528379
DELETING TIME-RECORD FOR TIME 3.2743223
DELETING TIME-RECORD FOR TIME 3.2743323
DELETING TIME-RECORD FOR TIME 3.2843323
DELETING TIME-RECORD FOR TIME 3.3843323
DELETING TIME-RECORD FOR TIME 3.4843323
DELETING TIME-RECORD FOR TIME 3.5843323
DELETING TIME-RECORD FOR TIME 3.6843323
DELETING TIME-RECORD FOR TIME 3.7843323
DELETING TIME-RECORD FOR TIME 3.8843323
DELETING TIME-RECORD FOR TIME 3.9843323
DELETING TIME-RECORD FOR TIME 4.0843323
DELETING TIME-RECORD FOR TIME 4.1843323
DELETING TIME-RECORD FOR TIME 4.2843323
DELETING TIME-RECORD FOR TIME 4.3843323
DELETING TIME-RECORD FOR TIME 4.4843323
DELETING TIME-RECORD FOR TIME 4.5843323
DELETING TIME-RECORD FOR TIME 4.6843323
DELETING TIME-RECORD FOR TIME 4.7843323
DELETING TIME-RECORD FOR TIME 4.8843323
KEEPING TIME-RECORD FOR TIME 4.9843323
AND FOR TIME 5.0000000
WORKSPACE RECLAIMED

TIMESTEP AT 5.0000000 SELECTED

DIC>
DIC> @@
DIC> @@ THE SIMULATION IS FINISHED
DIC> @@
DIC>
DIC> set-inter
--OK---
DIC>

exe1-plot

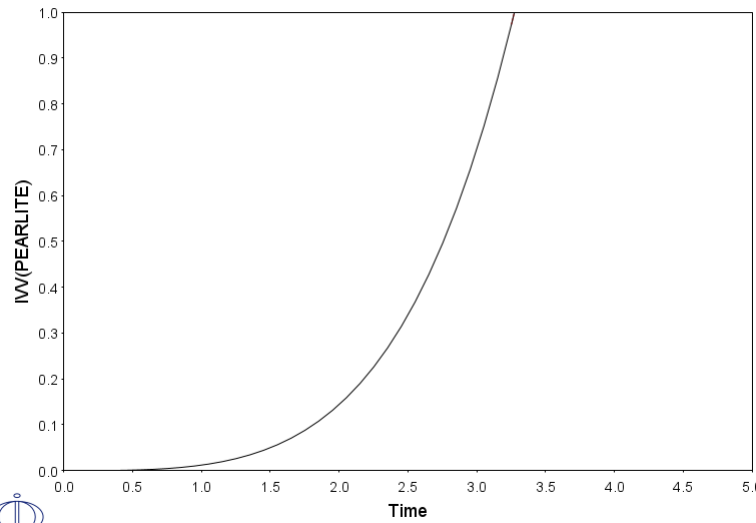
```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exe1\plot.DCM.test"
DIC>
DIC>
DIC> @@ exe1_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE e1
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 5.00000E+00
DIC> read exe1
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ PLOT THE TEMPERATURE AS A FUNCTION OF TIME
POST-1: @@
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-d-a y t
POST-1: s-p-c interface first
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 3
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13 12:58:10
"FIRST" INTERFACE OF SYSTEM
CELL #1
```



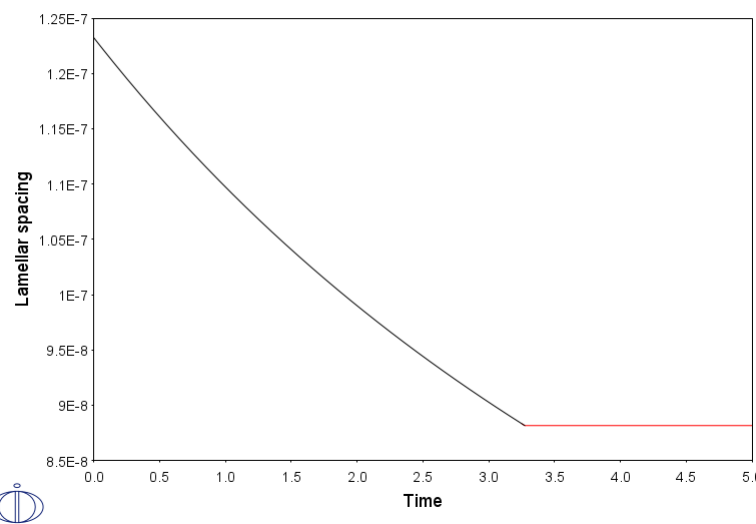
```
POST-1:
POST-1:
POST-1:
POST-1:Hit RETURN to continue
POST-1:
POST-1: @@
POST-1: @@ NOW PLOT THE FRACTION OF PEARLITE VS. TIME
POST-1: @@
POST-1: s-d-a y ivv(pearlite)
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 3
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

2021.05.13.12.58.11

CELL #1

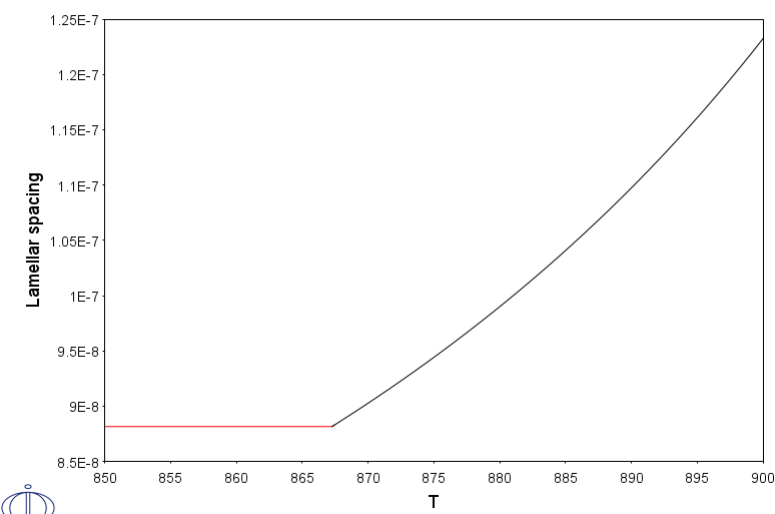


```
POST-1:  
POST-1:  
POST-1:  
POST-1:Hit RETURN to continue  
POST-1:  
POST-1: @@  
POST-1: @@ PLOT THE LAMELLAR SPACING AS A FUNCTION OF TIME  
POST-1: @@  
POST-1: s-d-a  
AXIS (X, Y OR Z) : y  
VARIABLE : lamellar-sp  
IN REGION: /*/: pearlite  
POST-1:  
POST-1: s-p-c  
CONDITION /INTEGRAL/: interface  
INTERFACE : pearlite  
UPPER OR LOWER INTERFACE OF REGION PEARLITE#1 /LOWER/: upper  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 3  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ... OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot  
2021.05.13.12.58.13  
UPPER INTERFACE OF REGION "PEARLITE#1"  
CELL #1
```

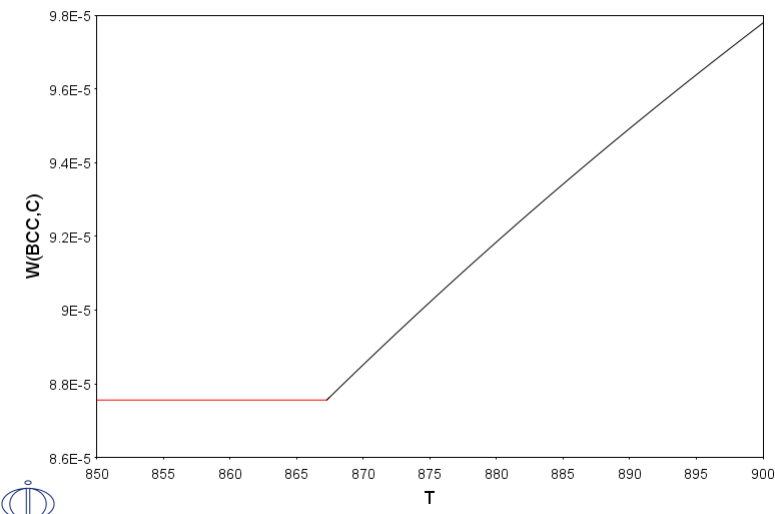


```
POST-1:  
POST-1:  
POST-1:  
POST-1:Hit RETURN to continue  
POST-1:  
POST-1: @@  
POST-1: @@ LET'S LOOK AT THE LAMELLAR SPACING VS. TEMPERATURE INSTEAD  
POST-1: @@  
POST-1: s-d-a x t  
POST-1:  
POST-1: s-p-c interface pearlite upper  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 3  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ... OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

2021.05.13.12.58.14
UPPER INTERFACE OF REGION "PEARLITE#1"
CELL #1

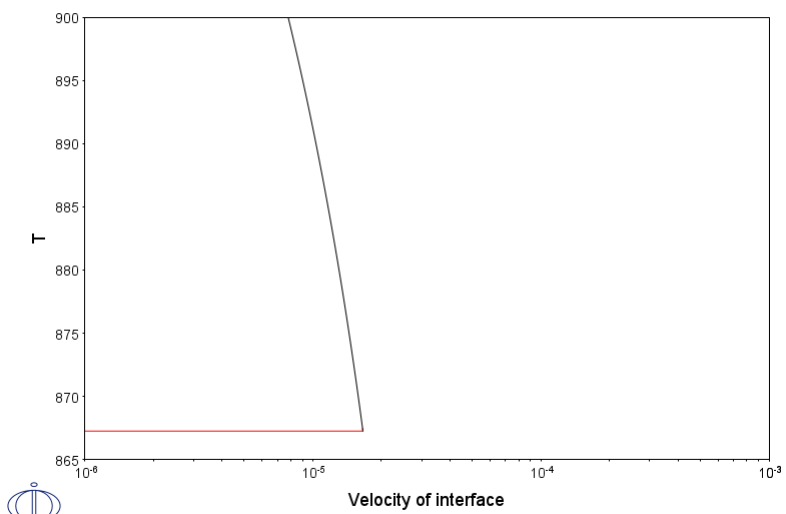


```
POST-1:  
POST-1:  
POST-1:  
POST-1:Hit RETURN to continue  
POST-1:  
POST-1: @@  
POST-1: @@ AND THE C COMPOSITION IN THE FERRITE VS. TEMP  
POST-1: @@  
POST-1: s-d-a y w(bcc,c)  
POST-1:  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 3  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ... OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot  
2021.05.13.12.58.15  
UPPER INTERFACE OF REGION "PEARLITE#1"  
CELL #1
```



```
POST-1:  
POST-1:  
POST-1:  
POST-1:Hit RETURN to continue  
POST-1:  
POST-1: @@  
POST-1: @@ FINALLY, LET'S LOOK AT THE VELOCITY OF THE INTERFACE VS. TEMP  
POST-1: @@  
POST-1: s-d-a y t  
POST-1: s-d-a x velocity  
INTERFACE : pearlite  
UPPER OR LOWER INTERFACE OF REGION PEARLITE#1 /LOWER/: upper  
POST-1: set-ax-ty x log  
POST-1: s-s-s x n 1e-6 1e-4  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 3  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
ORKING ... OST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

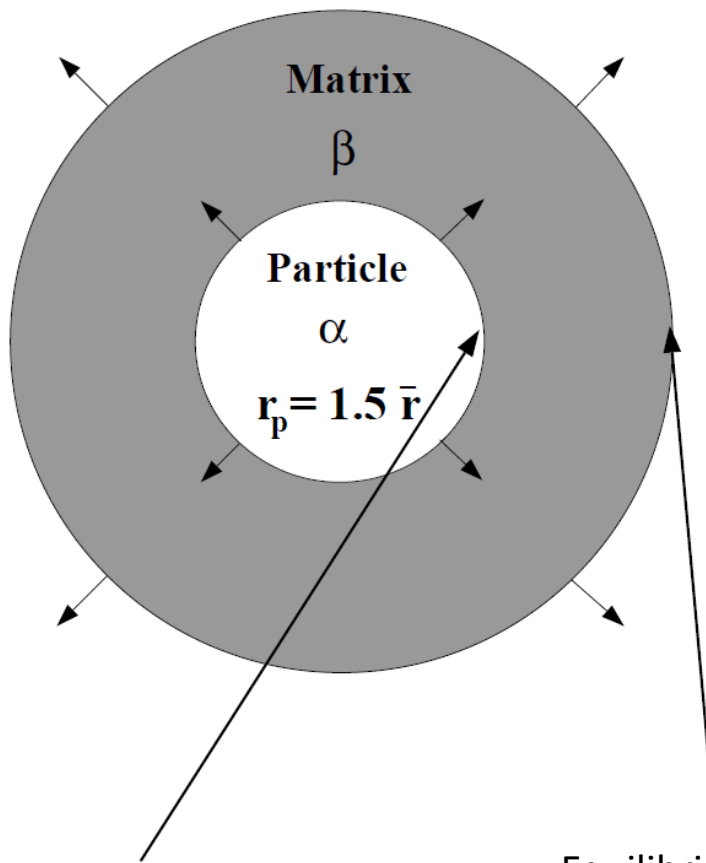
2021.05.13.12.58.16
UPPER INTERFACE OF REGION "PEARLITE#1"
CELL #1



POST-1:
POST-1:
POST-1:
POST-1: Hit RETURN to continue
POST-1:
POST-1:
POST-1:
POST-1: set-inter
--OK--
POST-1:



Coarsening



Moving phase interface
with α and β in local
equilibrium.

$\frac{2\sigma V_m}{r}$ Interfacial energy
contribution for α phase

Equilibrium as defined by
the average composition
in the system.

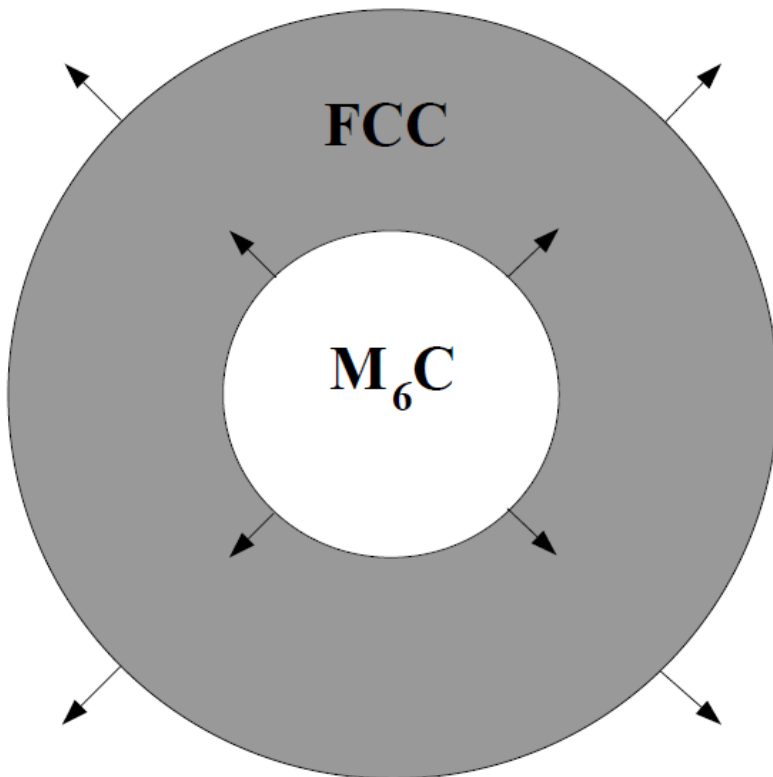
$\frac{2\sigma V_m}{r_p}$ Interfacial energy
contribution for α phase



Example exf1

Coarsening of an M_6C precipitate in an Fe-Mo-C alloy

This example calculates the Ostwald-ripening of a spherical M_6C carbide in an austenite matrix.



$$T = 1173\text{K}$$

$$r_p = 0.228 \mu\text{m}$$

exf1-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exf1\setup.DCM.test"
SYS: @@
SYS: @@ Coarsening problem.
SYS: @@ Coarsening of M6C precipitate in an Fe-Mo-C alloy
SYS: @@ This example calculates the Ostwald-ripening of a spherical
SYS: @@ M6C carbide in an austenite matrix.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASES
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA          /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: switch TCFE9
Current database: Steels/Fe-Alloys v9.3

VA          /- DEFINED
L12_FCC      B2_BCC           DICTRA_FCC_A1
REJECTED
TDB_TCFE9: def-sys fe mo c
FE           MO               C
DEFINED
TDB_TCFE9: rej ph * all
GAS:G        LIQUID:L          BCC_A2
FCC_A1       HCP_A3           CBCC_A12
CUB_A13      DIAMOND_FCC_A4  GRAPHITE
CEMENTITE    M23C6            M7C3
M6C          M5C2             M3C2
MC_ETA       MC_SHP          KSI_CARBIDE
Z_PHASE      FE4N_LP1         FECN_CHI
SIGMA        MU_PHASE         P_PHASE
R_PHASE      CHI_A12         LAVES_PHASE_C14
AL5FE4      M2O3C:I          REJECTED
TDB_TCFE9: res ph fcc m6c
FCC_A1       M6C RESTORED
TDB_TCFE9: get
12:59:34,755 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'A. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.-G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
    volumes'
'X.-G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'B. Hallstedt, unpublished work (2016); C-Fe-Mn Epsilon martensite.'
'J-O. Andersson, CALPHAD, 12 (1988) 1-8; TRITA 0317 (1986); C-MO'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'A. Fernandez Guillermet, CALPHAD, 6 (1982) 127-140; (sigma phase revised
    1986); TRITA-MAC 200 (1982); FE-MO'
'J-O. Andersson, CALPHAD, 12 (1988) 9-23; TRITA 0321 (1986); C-FE-MO'
'S. Nagakura, Transactions of the Iron and Steel Institute of Japan, 8
    (1968) 265-294; Molar volumes'
'I.K. Kupalova, V.I. Pavlova, High Speed Steels: Physical Properties,
    Prop. Data Updat. 2 (1988) 67-78; Molar volumes'
-OK-
TDB_TCFE9: @@
TDB_TCFE9: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_TCFE9: @@
TDB_TCFE9: app
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
TCAL7 = Al-Alloys v7.1
TCAL6 = Al-Alloys v6.0
```

TCAL5 = Al-Alloys v5.1
TCAL4 = Al-Alloys v4.0
TCAL3 = Al-Alloys v3.0
TCAL2 = Al-Alloys v2.1
TCAL1 = Al-Alloys v1.2
TCMG6 = Mg-Alloys v6.1
TCMG5 = Mg-Alloys v5.1
TCMG4 = Mg-Alloys v4.0
TCMG3 = Mg-Alloys v3.0
TCMG2 = Mg-Alloys v2.0
TCMG1 = Mg-Alloys v1.1
TCTI3 = Ti-Alloys v3.0
TCTI2 = Ti-Alloys v2.2
TCTI1 = Ti-Alloys v1.0
TCCU4 = Cu-Alloys v4.0
TCCU3 = Cu-Alloys v3.1
TCCU2 = Cu-Alloys v2.0
TCCU1 = Cu-Alloys v1.0
TCCC1 = Cemented carbide v1.0
TCHEA5 = High Entropy Alloy v5.0
TCHEA4 = High Entropy Alloy v4.2
TCHEA3 = High Entropy Alloy v3.1
TCHEA2 = High Entropy Alloy v2.1.1
TCHEA1 = High Entropy Alloy v1.0
SSOL7 = SGTE Alloy Solutions Database v7.0
SSOL6 = SGTE Alloy Solutions Database v6.0
SSOL5 = SGTE Alloy Solutions Database v5.0
SSOL4 = SGTE Alloy Solutions Database v4.9g
SSOL2 = SGTE Alloy Solutions Database v2.1
SSUB6 = SGTE Substances Database v6.0
SSUB5 = SGTE Substances Database v5.2
SSUB4 = SGTE Substances Database v4.1
SSUB3 = SGTE Substances Database v3.3
SSUB2 = SGTE Substances Database v2.2
SNOB3 = SGTE Noble Metal Alloys Database v3.1
SNOB2 = SGTE Noble Metal Alloys Database v2.1
SNOB1 = SGTE Noble Metal Alloys Database v1.2
STBC2 = SGTE Thermal Barrier Coating TDB v2.2
STBC1 = SGTE Thermal Barrier Coating TDB v1.1
SALT1 = SGTE Molten Salts Database v1.3
SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
SEMC2 = TC Semi-Conductors v2.1
SLAG4 = Fe-containing Slag v4.1
SLAG3 = Fe-containing Slag v3.2
SLAG2 = Fe-containing Slag v2.2
SLAG1 = Fe-containing Slag v1.2
TCOX11 = Metal Oxide Solutions v11.0
TCOX10 = Metal Oxide Solutions v10.1
TCOX9 = Metal Oxide Solutions v9.0
TCOX8 = Metal Oxide Solutions v8.0
TCOX7 = Metal Oxide Solutions v7.0
TCOX6 = Metal Oxide Solutions v6.0
TCOX5 = Metal Oxide Solutions v5.1
TCOX4 = Metal Oxide Solutions v4.1
ION3 = Ionic Solutions v3.0
ION2 = Ionic Solutions v2.6
ION1 = Ionic Solutions v1.5
NOX2 = NPL Oxide Solutions Database v2.1
TCNOBL1 = Noble Metals Alloys v1.0
TCNOBL2 = Noble Metals Alloys v2.0
TCSLD4 = Solder Alloys v4.0
TCSLD3 = Solder Alloys v3.3
TCSLD2 = Solder Alloys v2.0
TCSLD1 = Solder Alloys v1.1
TCSI1 = Ultrapure Silicon v1.2
TCPMP2 = Materials Processing v2.5
TCES1 = Combustion/Sintering v1.1
TCSC1 = Super Conductor v1.0
TCFC1 = SOFC Database v1.0
TCNF2 = Nuclear Fuels v2.1b
NUMT2 = Nuclear Materials v2.1
NUOX4 = Nuclear Oxides v4.2
NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
NUCL19 = IRSN NUCLEA-19
NUCL15 = IRSN NUCLEA-15_4
NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
MEPH19 = IRSN Mephista-19
MEPH15 = IRSN Mephista-15_1
MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
TCAQ3 = Aqueous Solution v3.0
TCAQ2 = Aqueous Solution v2.7
AQ52 = TGG Aqueous Solution Database v2.6
GCE2 = TGG Geochemical/Environmental TDB v2.3
FEDEMO = Iron Demo Database v4.0
ALDEMO = Aluminum Demo Database v4.0
NIDEMO = Nickel Demo Database v2.0
CUDEMO = Copper Demo Database v1.0
SLDEMO = Solder Demo Database v1.0
OXDEMO = Oxide Demo Database v3.0
SUBDEMO = Substance Demo Database v1.0
PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
PG35 = PG35 Binary Semi-Conductors TDB v1.3
PURE5 = SGTE Unary (Pure Elements) TDB v5.2
MOB2 = Alloys Mobility v2.7
MOB1 = Alloys Mobility v1.3
MOBFE1 = Steels/Fe-Alloys Mobility v1.1
MOBFE2 = Steels/Fe-Alloys Mobility v2.0
MOBFE3 = Steels/Fe-Alloys Mobility v3.0
MOBFE4 = Steels/Fe-Alloys Mobility v4.0
MOBFE5 = Steels/Fe-Alloys Mobility v5.0
MOBFE6 = Steels/Fe-Alloys Mobility v6.0
MOBN15 = Ni-Alloys Mobility v5.1
MOBN14 = Ni-Alloys Mobility v4.1
MOBN13 = Ni-Alloys Mobility v3.2
MOBN12 = Ni-Alloys Mobility v2.4
MOBN11 = Ni-Alloys Mobility v1.10
MOBAL6 = Al-Alloys Mobility v6.0
MOBAL5 = Al-Alloys Mobility v5.0
MOBAL4 = Al-Alloys Mobility v4.0
MOBAL3 = Al-Alloys Mobility v3.0
MOBAL2 = Al-Alloys Mobility v2.0
MOBAL1 = Al-Alloys Mobility v1.0
MOBCU1 = Cu-Alloys Mobility v1.0
MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0

```

MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBSI1 = Si-Alloys Mobility v1.0
MOBSDL1 = Solder-Alloys Mobility v1.1
MOBTI4 = Ti-Alloys Mobility v4.0
MOBTI3 = Ti-Alloys Mobility v3.1
MOBTI2 = Ti-Alloys Mobility v2.0
MOBTI1 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

```

```

DATABASE NAME /TCFE9/: mobfe4
Current database: Steels/Fe-Alloys Mobility v4.0

```

```

VA DEFINED
B2_BCC REJECTED
APP: def-sys fe mo c
FE MO C
DEFINED
APP: rej ph * all
BCC_A2 CEMENTITE FCC_A1
FE4N_LP1 HCP_A3 LIQUID:L
REJECTED
APP: res ph fcc m6c
*** ERROR M6C INPUT IGNORED
FCC_A1 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
FUNCTIONS .....

```

List of references for assessed data

```

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'Y. Liu et al., Calphad 34(2010)253-262; fcc Cu-Fe.'
'W. Zheng, et al., Metall Mater Trans, 48 (2017) 536-550; Fe-Mn-Mo (fcc
and bcc)'

```

-OK-

```

APP:
APP:
APP: @@ ENTER THE DICTRA MONITOR
APP: @@
APP: go d-m
NO TIME STEP DEFINED
*** ENTERING M6C AS A DIFFUSION NONE PHASE
DIC>

```

```

DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> s-cond glob t 0 1173; * N
DIC>

```

```

DIC> @@
DIC> @@ ENTER REGIONS part AND aus
DIC> @@
DIC> enter-region

```

```

REGION NAME : part
DIC> enter-region aus

```

```

ATTACH TO REGION NAMED /PART/:
ATTACHED TO THE RIGHT OF PART /YES/:

```

```

DIC> @@
DIC> @@ ENTER GEOMTRICAL GRIDS INTO THE REGIONS
DIC> @@
DIC>
DIC> @@
DIC> @@ THE INITIAL SIZE OF THE CARBIDE PARTICLE IS ASSUMED TO BE KNOWN
DIC> @@ (IN THIS CASE THE VALUE IS FROM NISHIZAWA ET. AL.). THE
DIC> @@ AVERAGE PARTICLE SIZE IS ASSUMED TO BE 0.152E-6 METERS. HOWEVER, THE
DIC> @@ CALCULATIONS ARE PERFORMED ON A MAXIMUM SIZE PARTICLE, WHICH IS ASSUMED
DIC> @@ TO BE 1.5 TIMES THE AVERAGE SIZE. THE SURROUNDING AUSTENITIC MATRIX
DIC> @@ SIZE IS CHOOSEN TO MAINTAIN THE AVERAGE COMPOSITION.
DIC> @@

```

```

DIC> enter-grid
REGION NAME : /PART/: part

```

```

WIDTH OF REGION /1/: 0.228E-6

```

```

TYPE /LINEAR/: AUTO

```

```

DIC>

```

```

DIC> enter-grid

```

```

REGION NAME : /AUS/: aus

```

```

WIDTH OF REGION /1/: 4.53147041E-7

```

```

TYPE /LINEAR/: AUTO

```

```

DIC>

```

```

DIC> @@

```

```

DIC> @@ ENTER PHASES INTO REGIONS

```

```

DIC> @@

```

```

DIC> enter-phase active part matrix m6c

```

```

DIC>

```

```

DIC> enter-phase active aus matrix fcc#1

```

```

DIC>

```

```

DIC> @@

```

```

DIC> @@ ENTER THE INITIAL COMPOSITIONS IN THE PHASES

```

```

DIC> @@

```

```

DIC> enter-composition

```

```

REGION NAME : /PART/: part

```

```

PHASE NAME: /M6C/: m6c

```

```

DEPENDENT COMPONENT ? /MO/: fe

```

```

COMPOSITION TYPE /MOLE_FRACTION/: w-f

```

```

PROFILE FOR /MO/: mo lin 6.20117E-01 6.20117E-01

```

```

DIC>

```

```

DIC> ent-composition

```

```
REGION NAME : /AUS/: aus
PHASE NAME: /FCC A1/: fcc#1
DEPENDENT COMPONENT ? /MO/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-f
PROFILE FOR /C/: mo lin 1.82099E-02 1.82099E-02
PROFILE FOR /MO/: c lin 2.83351E-03 2.83351E-03
DIC>
DIC>
DIC> @@
DIC> @@ SET A SPHERICAL GEOMETRY
DIC> @@
DIC> ent-geo 2
DIC>
DIC> @@
DIC> @@ ENTER THE SURFACE TENSION ENERGY CONTRIBUTION AS A FUNCTION OF
DIC> @@ THE INTERFACE POSITION (THE RADIUS OF THE PARTICLE).
DIC> @@ ALSO ENTER THE MOLAR VOLUME OF THE PHASE CORRECTED TO BE THE
DIC> @@ MOLAR VOLUME PER SUBSTITUTIONAL ATOM.
DIC> @@
DIC> @@ THE SURFACE TENSION IS 0.7, THE MOLAR VOLUME IS 0.71 AND THE
DIC> @@ TRANSFORMATION TO MOLAR VOLUME PER SUBSTITUTIONAL ATOM IS 7/6.
DIC> @@
DIC> set-surf 2*0.7*0.71*(7/6)/X;
ENTERED FUNCTION :2*.7*.71*7/6/X FOR CELL #1
DIC>
DIC>
DIC> @@
DIC> @@ ENABLE THE SIMPLIFIED MODEL FOR THE COARSENING (OSTWALD-RIPENING)
DIC> @@
DIC> coarse YES
DIC>
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME AND OTHER SIMULATION PARAMETERS
DIC> @@
DIC> set-simulation-time 1E6
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /100000/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exf1 Y
DIC>
DIC> set-inter
--OK--
DIC>
```

exf1-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exf1\run.DCM.test"
DIC> @@ exf1_run.DCM
DIC>
DIC> @@
DIC> @@ READ THE SETUP FROM FILE AND START THE SIMULATION
DIC> @@
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
*** ENTERING M6C AS A DIFFUSION NONE PHASE
DIC> read exf1
OK
DIC> sim
Region: PART
single geometric dense at 0.22800E-06
0.80000 64
Region: AUS
single geometric dense at 0.0000
1.00600 63
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
Trying old scheme 4
GENERATING STARTING VALUES FOR CELL # 1 INTERFACE # 2
DETERMINING INITIAL EQUILIBRIUM VALUES
CALCULATING STARTING VALUES: 9 EQUILIBRIUM CALCULATIONS DONE 6 OUT OF 9
U-FRACTION IN SYSTEM: C = .0190652843033664 FE = .970761291162784
MO = .0292387089677228
TOTAL SIZE OF SYSTEM: 1.32376603026E-18 [m^3]
U-FRACTION IN SYSTEM: C = .0190652843033664 FE = .970761291162784
MO = .0292387089677228
TOTAL SIZE OF SYSTEM: 1.32376603026E-18 [m^3]
0.610222970497625 0.610391048253360 0.610222911134902 2.564830792704480E-002 2.191933775609301E-
003 1.266865150888817E-004 1.096501654713504E-003 1.278319049399803E-004 4.432656450601522E-
007 1.566395317892607E-009 1.94733765313978E-009 3.659996125120900E-009 1.418022110198698E-
009 1.309139507131020E-009 1.103984934282495E-009 3.800021059935427E-009 7.446660066835133E-
010 2.344008708335402E-010 1.080152259465027E-013 1.867047546680927E-018 TIME = 0.1000000E-06 DT = 0.1000000E-
06 SUM OF SQUARES = 0.18670475E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.50349408E-05 AND -0.50349408E-05
POSITION OF INTERFACE PART / AUS IS 0.22799950E-06
U-FRACTION IN SYSTEM: C = .0190659317665348 FE = .97075948763499
MO = .0292405124955171
TOTAL SIZE OF SYSTEM: 1.32375726043E-18 [m^3]
6 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: PART

CPU time used in timestep 1 seconds
5.960103082661315E-005 5.961330244477064E-005 5.959458009987109E-005 3.934421751033465E-007 1.790908700287879E-
012 8.025303437845193E-020 SWITCHING ACTIVITIES FOR INTERFACE #2, CELL #1
FROM: C TO: MO
TIME = 0.254446964E-05 DT = 0.244446964E-05 SUM OF SQUARES = 0.80253034E-19
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.39293667E-08 AND -0.39293667E-08
POSITION OF INTERFACE PART / AUS IS 0.22799949E-06
U-FRACTION IN SYSTEM: C = .01906647907566638 FE = .970759484159471
MO = .029240515971036
TOTAL SIZE OF SYSTEM: 1.32375709311E-18 [m^3]
CPU time used in timestep 0 seconds
2.51515852968503E-008 2.515761992652319E-008 2.499573886069715E-008 2.005534172040529E-008 1.975331989271387E-
008 1.917061703973256E-008 1.798309031226448E-008 1.798198229048188E-008 1.576453640079517E-
008 1.173116059610038E-008 5.468961566157615E-009 5.475061910470316E-009 2.640249640042667E-
011 1.077675289092714E-019 TIME = 0.74340891E-05 DT = 0.48893927E-05 SUM OF SQUARES = 0.10776753E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.56129636E-09 AND -0.56129636E-09
POSITION OF INTERFACE PART / AUS IS 0.22799948E-06
U-FRACTION IN SYSTEM: C = .0190671017629171 FE = .970759483986865
MO = .0292405161436416
TOTAL SIZE OF SYSTEM: 1.32375704531E-18 [m^3]
CPU time used in timestep 0 seconds
4.333438087006518E-010 4.334345197139853E-010 4.419215189345052E-010 4.026611773982020E-010 4.018893335048845E-
010 4.003483966913515E-010 3.972574076251299E-010 3.977091072732849E-010 3.911287932296859E-
010 3.789974358433123E-010 3.553242202315046E-010 3.559796456900178E-010 3.102532494760969E-
010 2.292896947168586E-010 1.040111353482194E-010 1.045003421997081E-010 9.690622551426697E-
014 4.706856943694132E-024 TIME = 0.17212874E-04 DT = 0.97787854E-05 SUM OF SQUARES = 0.47068569E-23
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.11440105E-09 AND -0.11440105E-09
POSITION OF INTERFACE PART / AUS IS 0.22799948E-06
U-FRACTION IN SYSTEM: C = .0190679130516543 FE = .970759484521662
MO = .0292405156088451
TOTAL SIZE OF SYSTEM: 1.32375702582E-18 [m^3]
CPU time used in timestep 1 seconds

output ignored...

... output resumed

2.171297345832007E-011 9.341085264458381E-013 1.185624034686549E-013 1.155534463862785E-
022 TIME = 667998.25 DT = 100000.00 SUM OF SQUARES = 0.11555345E-21
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.22669488E-13 AND 0.22669488E-13
POSITION OF INTERFACE PART / AUS IS 0.24398484E-06
U-FRACTION IN SYSTEM: C = .0195561971882094 FE = .970692497669248
MO = .0293075024612589
TOTAL SIZE OF SYSTEM: 1.62216577705E-18 [m^3]
29 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: PART
29 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUS

CPU time used in timestep 0 seconds
8.463228722651036E-012 8.732923363774625E-012 1.106305642801926E-011 1.671261459244899E-012 8.083185529438911E-
013 2.031417575905656E-015 4.552642885278637E-
025 TIME = 767998.25 DT = 100000.00 SUM OF SQUARES = 0.45526429E-24
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.22252996E-13 AND 0.22252996E-13
POSITION OF INTERFACE PART / AUS IS 0.24621014E-06
U-FRACTION IN SYSTEM: C = .0195587075255261 FE = .970694293020125
MO = .0293057071103818
TOTAL SIZE OF SYSTEM: 1.66695743697E-18 [m^3]
16 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: PART
16 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUS

CPU time used in timestep 0 seconds
4.716189884869470E-012 4.737893973260528E-012 6.250071117694707E-012 2.547264430727638E-012 1.906488344119326E-
012 8.178985251311430E-013 4.505542982616667E-
022 TIME = 867998.25 DT = 100000.00 SUM OF SQUARES = 0.45055430E-21
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.21855116E-13 AND 0.21855116E-13
```

POSITION OF INTERFACE PART / AUS IS 0.24839565E-06
 U-FRACTION IN SYSTEM: C = .0195611435099182 FE = .970696048456854
 MO = .0293039516736537
 TOTAL SIZE OF SYSTEM: 1.71174343902E-18 [m^3]
 25 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: PART
 25 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUS

CPU time used in timestep	0 seconds	3.789984686980590E-012	4.665295481791764E-012	3.497964005114297E-012	3.136690411875925E-
012	2.396321665117972E-012	1.082065620523175E-012	1.796667887015045E-015	5.918392824720176E-	
025	TIME = 967998.25	DT = 100000.00	SUM OF SQUARES = 0.59183928E-24		
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.21474553E-13 AND 0.21474553E-13					
POSITION OF INTERFACE PART / AUS IS 0.25054311E-06					
U-FRACTION IN SYSTEM: C = .0195635100799567 FE = .970697758696434					
MO = .029302241434073					
TOTAL SIZE OF SYSTEM: 1.75652397336E-18 [m^3]					
13 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: PART					
13 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUS					

CPU time used in timestep	1 seconds	3.128551219381802E-010	3.018312578535717E-010	3.724165660663593E-011	1.354265960616163E-
011	1.037000287214743E-019	TIME = 1000000.0	DT = 32001.746	SUM OF SQUARES = 0.10370003E-18	
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.21247167E-13 AND 0.21247167E-13					
POSITION OF INTERFACE PART / AUS IS 0.25122306E-06					
U-FRACTION IN SYSTEM: C = .0195660575765549 FE = .970698831747598					
MO = .0293011683829095					
TOTAL SIZE OF SYSTEM: 1.77086385838E-18 [m^3]					
MUST SAVE WORKSPACE ON FILE					
WORKSPACE SAVED ON FILE					
RECLAIMING WORKSPACE					
DELETING TIME-RECORD FOR TIME 0.000000					
DELETING TIME-RECORD FOR TIME 0.10000000E-06					
DELETING TIME-RECORD FOR TIME 0.25446964E-05					
DELETING TIME-RECORD FOR TIME 0.74340891E-05					
DELETING TIME-RECORD FOR TIME 0.17212874E-04					
DELETING TIME-RECORD FOR TIME 0.36770445E-04					
DELETING TIME-RECORD FOR TIME 0.75885587E-04					
DELETING TIME-RECORD FOR TIME 0.15411587E-03					
DELETING TIME-RECORD FOR TIME 0.31057644E-03					
DELETING TIME-RECORD FOR TIME 0.62349757E-03					
DELETING TIME-RECORD FOR TIME 0.12493398E-02					
DELETING TIME-RECORD FOR TIME 0.25010244E-02					
DELETING TIME-RECORD FOR TIME 0.50043934E-02					
DELETING TIME-RECORD FOR TIME 0.10011132E-01					
DELETING TIME-RECORD FOR TIME 0.20024608E-01					
DELETING TIME-RECORD FOR TIME 0.40051560E-01					
DELETING TIME-RECORD FOR TIME 0.80105465E-01					
DELETING TIME-RECORD FOR TIME 0.16021328					
DELETING TIME-RECORD FOR TIME 0.32042890					
DELETING TIME-RECORD FOR TIME 0.64086014					
DELETING TIME-RECORD FOR TIME 1.2817226					
DELETING TIME-RECORD FOR TIME 2.5634476					
DELETING TIME-RECORD FOR TIME 5.1268975					
DELETING TIME-RECORD FOR TIME 10.253797					
DELETING TIME-RECORD FOR TIME 20.507597					
DELETING TIME-RECORD FOR TIME 41.015196					
DELETING TIME-RECORD FOR TIME 82.030395					
DELETING TIME-RECORD FOR TIME 164.06079					
DELETING TIME-RECORD FOR TIME 328.12159					
DELETING TIME-RECORD FOR TIME 656.24318					
DELETING TIME-RECORD FOR TIME 1312.4864					
DELETING TIME-RECORD FOR TIME 2624.9727					
DELETING TIME-RECORD FOR TIME 5249.9454					
DELETING TIME-RECORD FOR TIME 10499.891					
DELETING TIME-RECORD FOR TIME 20999.782					
DELETING TIME-RECORD FOR TIME 41999.564					
DELETING TIME-RECORD FOR TIME 83999.127					
DELETING TIME-RECORD FOR TIME 167998.25					
DELETING TIME-RECORD FOR TIME 267998.25					
DELETING TIME-RECORD FOR TIME 367998.25					
DELETING TIME-RECORD FOR TIME 467998.25					
DELETING TIME-RECORD FOR TIME 567998.25					
DELETING TIME-RECORD FOR TIME 667998.25					
DELETING TIME-RECORD FOR TIME 767998.25					
DELETING TIME-RECORD FOR TIME 867998.25					

KEEPING TIME-RECORD FOR TIME 967998.25
 AND FOR TIME 1000000.0
 WORKSPACE RECLAIMED

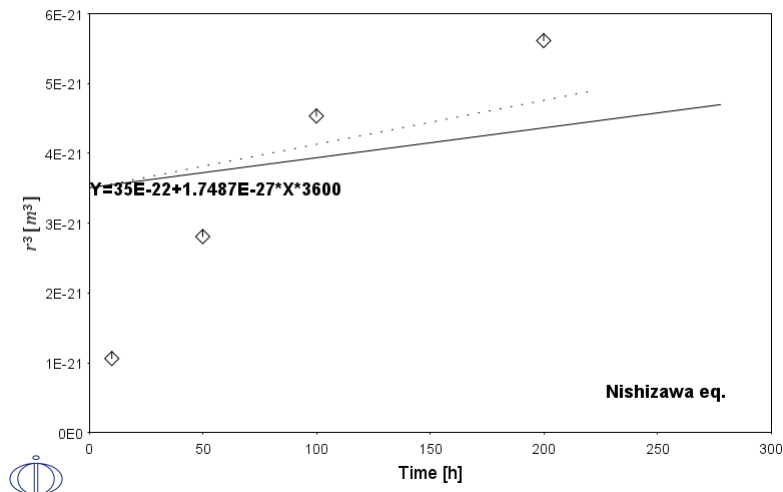
TIMESTEP AT	1000000.00	SELECTED
DIC>		
DIC>		
DIC> set-inter		
--OK--		
DIC>		

exf1-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exf1\plot.DCM.test"
DIC> @@ exf1_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE f1
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 1.00000E+06
*** ENTERING M6C AS A DIFFUSION NONE PHASE
DIC> read exf1
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ PLOT THE AVERAGE PARTICLE SIZE (CUBED) AS THIS ASSUMED TO
POST-1: @@ SCALE LINEARLY WITH TIME. THEN A FUNCTION IS ENTERED SO
POST-1: @@ THIS QUANTITY CAN BE ACCESSED. WE ALSO WANT TO PLOT THIS
POST-1: @@ QUANTITY VERSUS TIME (IN HOURS) SO A FUNCTION IS ENTERED.
POST-1: @@
POST-1: enter-symbol func rr3=(poi(part,u)/1.5)**3;
POST-1: enter-symbol func hours=time/3600;
POST-1: s-d-a x hours
POST-1: s-d-a y rr3
POST-1: @@
POST-1: @@ AS WE ARE PLOTTING FUNCTIONS ON BOTH AXES WE MUST EXPLICITLY
POST-1: @@ DEFINE THE INDEPENDENT VARIABLE AND THE PLOT CONDITION
POST-1: @@
POST-1: s-ind time
POST-1: s-p-c inter
INTERFACE : part upper
POST-1:
POST-1:
POST-1: set-axis-text-status x n
AXIS TEXT : Time [h]
POST-1:
POST-1: @@
POST-1: @@ WHEN THIS IS PLOTTED, THIS AXIS TEXT NOTATION WORKS WELL FOR
POST-1: @@ THE AVERAGE RADIUS CUBED. FOR MORE INFORMATION ABOUT HOW TO
POST-1: @@ ADJUST TEXT IN THE POST PROCESSOR USING THE DATAPLOT LANGUAGE,
POST-1: @@ SEARCH THE ONLINE HELP (FROM THE MAIN MENU -> HELP > ONLINE HELP)
POST-1:
POST-1:
POST-1: set-axis-text-status y n
AXIS TEXT : \ latex r^3\, [m^3]
POST-1:
POST-1:
POST-1: @@
POST-1: @@ COMPARE WITH EXPERIMENTAL DATA FROM NISHIZAWA ET AL.
POST-1: @@ TRANS. JPN. INST. MET. VOL. 22 1981 PP. 733-742.
POST-1: @@
POST-1: @@
POST-1: app y exf1
PROLOGUE NUMBER: /0/: 0
DATASET NUMBER(s): /-1/: 1
POST-1:
POST-1: s-s-s y n 0 6e-21
POST-1:
POST-1: @@
POST-1: @@ SET A TITLE ON THE PLOT
POST-1: @@
POST-1: set-title Figure f1.1
POST-1:
POST-1: @@ SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Figure f1.1

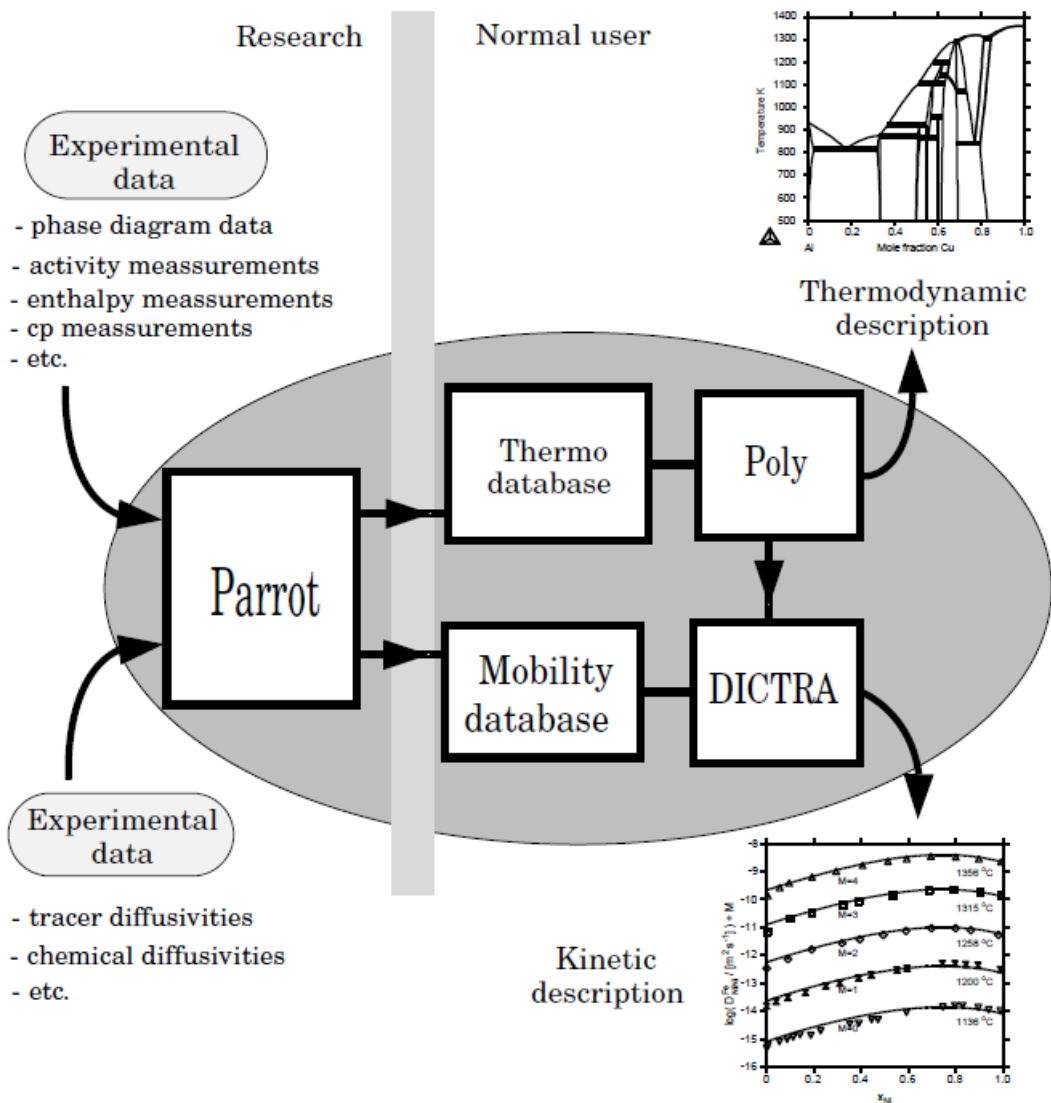
2021.05.13.13.01.49
UPPER INTERFACE OF REGION "PART#1"
CELL #1



```
POST-1:  
POST-1:@?<_hit_return_to_continue_>  
POST-1: @@  
POST-1: @@ THE DIFFERENCE BETWEEN THE CALCULATION AND THE EQUATION USED BY  
POST-1: @@ NISHIZAWA ET AL. IS MAINLY DUE TO DIFFERENT THERMODYNAMIC  
POST-1: @@ DESCRIPTIONS AND DIFFUSIVITIES.  
POST-1: @@  
POST-1:  
POST-1: set-inter  
--OK--  
POST-1:
```



Kinetic Data

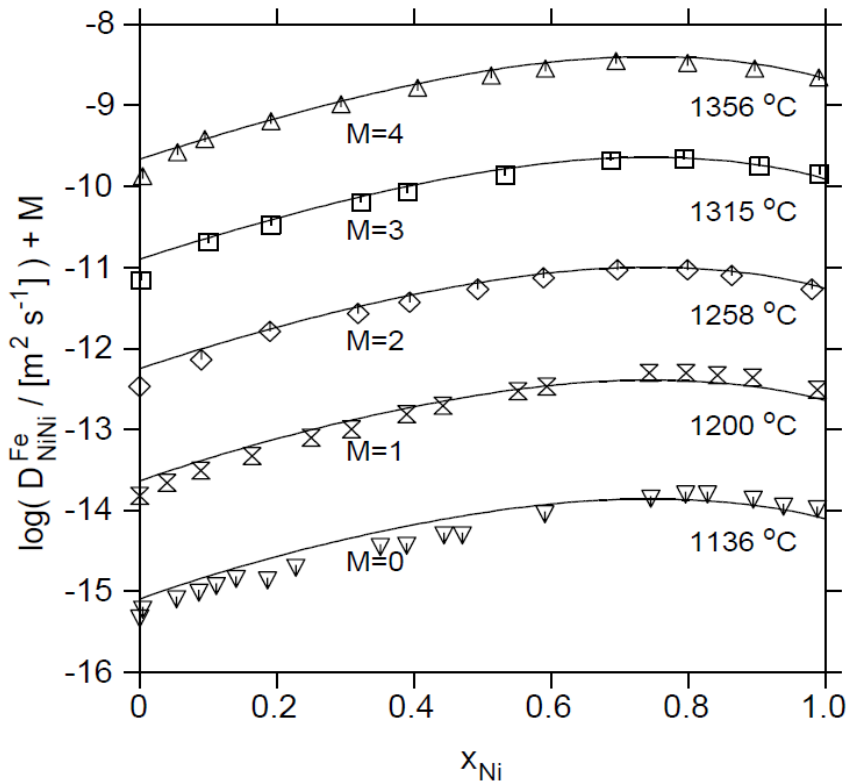




Example exg1

Checking diffusivities in an Fe-Ni alloy

This is an example file to check the mobilities and diffusivities in an Fe-Ni alloy.



exgl-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exgl\setup.DCM.test"
SYS: @@
SYS: @@ Kinetic data example.
SYS: @@ Checking mobilities and diffusivities in an Fe-Ni alloy
SYS: @@ This is an example file to check the mobilities and diffusivities
SYS: @@ in an Fe-Ni alloy.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exgl_setup.DCM
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA                                /- DEFINED
DICTRA_FCC_A1  REJECTED
TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ SELECT A DATABASE FOR THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw fedemo
Current database: Iron Demo Database v4.0

VA                                /- DEFINED
TDB_FEDEMO: @@
TDB_FEDEMO: @@ DEFINE THE SYSTEM TO WORK WITH
TDB_FEDEMO: @@
TDB_FEDEMO: def-sys fe ni
FE                               NI  DEFINED
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ EXCLUDE THE THERMODYNAMIC DATA FOR THE PHASES THAT ARE NOT NEEDED
TDB_FEDEMO: @@
TDB_FEDEMO: rej ph * all
LIQUID:L          BCC_A2           LAVES_PHASE_C14
CBCC_A12         CUB_A13          FCC_A1
HCP_A3  REJECTED
TDB_FEDEMO: res ph fcc
FCC_A1  RESTORED
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ RETRIEVE DATA FROM THE DATABASE FILE
TDB_FEDEMO: @@
TDB_FEDEMO: get
13:03:07,799 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
    volumes'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
-OK-
TDB_FEDEMO: @@
TDB_FEDEMO: @@ MOBILITY/DIFFUSIVITY DATA ARE STORED IN A SEPARATE DATABASE FILE.
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app
Use one of these databases

TCFE11 = Steels/Fe-Alloys v11.0
TCFE10 = Steels/Fe-Alloys v10.1
TCFE9 = Steels/Fe-Alloys v9.3
TCFE8 = Steels/Fe-Alloys v8.2
TCFE7 = Steels/Fe-Alloys v7.0
TCFE6 = Steels/Fe-Alloys v6.2
TCFE5 = Steels/Fe-Alloys v5.0
TCFE4 = Steels/Fe-Alloys v4.1
TCFE3 = Steels/Fe-Alloys v3.1
TCFE2 = Steels/Fe-Alloys v2.1
TCFE1 = Steels/Fe-Alloys v1.0
FEDAT = TCS/TT Steels Database v1.0
TCNI11 = Ni-Alloys v11.0
TCNI10 = Ni-Alloys v10.0
TCNI9 = Ni-Alloys v9.1
TCNI8 = Ni-Alloys v8.2
TCNI7 = Ni-Alloys v7.2
TCNI6 = Ni-Alloys v6.1
TCNI5 = Ni-Alloys v5.1
TCNI4 = Ni-Alloys v4.0
TCNI1 = Ni-Alloys v1.3
TCAL8 = Al-Alloys v8.0
```

TCAL7 = Al-Alloys v7.1
 TCAL6 = Al-Alloys v6.0
 TCAL5 = Al-Alloys v5.1
 TCAL4 = Al-Alloys v4.0
 TCAL3 = Al-Alloys v3.0
 TCAL2 = Al-Alloys v2.1
 TCAL1 = Al-Alloys v1.2
 TCMG6 = Mg-Alloys v6.1
 TCMG5 = Mg-Alloys v5.1
 TCMG4 = Mg-Alloys v4.0
 TCMG3 = Mg-Alloys v3.0
 TCMG2 = Mg-Alloys v2.0
 TCMG1 = Mg-Alloys v1.1
 TCTI3 = Ti-Alloys v3.0
 TCTI2 = Ti-Alloys v2.2
 TCTI1 = Ti-Alloys v1.0
 TCCU4 = Cu-Alloys v4.0
 TCCU3 = Cu-Alloys v3.1
 TCCU2 = Cu-Alloys v2.0
 TCCU1 = Cu-Alloys v1.0
 TCCC1 = Cemented carbide v1.0
 TCHEA5 = High Entropy Alloy v5.0
 TCHEA4 = High Entropy Alloy v4.2
 TCHEA3 = High Entropy Alloy v3.1
 TCHEA2 = High Entropy Alloy v2.1.1
 TCHEA1 = High Entropy Alloy v1.0
 SSOL7 = SGTE Alloy Solutions Database v7.0
 SSOL6 = SGTE Alloy Solutions Database v6.0
 SSOL5 = SGTE Alloy Solutions Database v5.0
 SSOL4 = SGTE Alloy Solutions Database v4.9g
 SSOL2 = SGTE Alloy Solutions Database v2.1
 SSUB6 = SGTE Substances Database v6.0
 SSUB5 = SGTE Substances Database v5.2
 SSUB4 = SGTE Substances Database v4.1
 SSUB3 = SGTE Substances Database v3.3
 SSUB2 = SGTE Substances Database v2.2
 SNOB3 = SGTE Noble Metal Alloys Database v3.1
 SNOB2 = SGTE Noble Metal Alloys Database v2.1
 SNOB1 = SGTE Noble Metal Alloys Database v1.2
 STBC2 = SGTE Thermal Barrier Coating TDB v2.2
 STBC1 = SGTE Thermal Barrier Coating TDB v1.1
 SALT1 = SGTE Molten Salts Database v1.3
 SNUX6 = SGTE In-Vessel Nuclear Oxide TDB v6.2
 SEMC2 = TC Semi-Conductors v2.1
 SLAG4 = Fe-containing Slag v4.1
 SLAG3 = Fe-containing Slag v3.2
 SLAG2 = Fe-containing Slag v2.2
 SLAG1 = Fe-containing Slag v1.2
 TCOX11 = Metal Oxide Solutions v11.0
 TCOX10 = Metal Oxide Solutions v10.1
 TCOX9 = Metal Oxide Solutions v9.0
 TCOX8 = Metal Oxide Solutions v8.0
 TCOX7 = Metal Oxide Solutions v7.0
 TCOX6 = Metal Oxide Solutions v6.0
 TCOX5 = Metal Oxide Solutions v5.1
 TCOX4 = Metal Oxide Solutions v4.1
 ION3 = Ionic Solutions v3.0
 ION2 = Ionic Solutions v2.6
 ION1 = Ionic Solutions v1.5
 NOX2 = NPL Oxide Solutions Database v2.1
 TCNOBL1 = Noble Metals Alloys v1.0
 TCNOBL2 = Noble Metals Alloys v2.0
 TCSLD4 = Solder Alloys v4.0
 TCSLD3 = Solder Alloys v3.3
 TCSLD2 = Solder Alloys v2.0
 TCSLD1 = Solder Alloys v1.1
 TCSI1 = Ultrapure Silicon v1.2
 TCMR2 = Materials Processing v2.5
 TCES1 = Combustion/Sintering v1.1
 TCSC1 = Super Conductor v1.0
 TCFCl = SOFC Database v1.0
 TCNF2 = Nuclear Fuels v2.1b
 NUMT2 = Nuclear Materials v2.1
 NUOX4 = Nuclear Oxides v4.2
 NUTO1 = U-Zr-Si Ternary Oxides TDB v1.1
 NUTA1 = Ag-Cd-In Ternary Alloys TDB v1.1
 NUCL19 = IRSN NUCLEA-19
 NUCL15 = IRSN NUCLEA-15_4
 NUCL10 = ThermoData NUCLEA Alloys-oxides TDB v10.2
 MEPH19 = IRSN Mephisto-19
 MEPH15 = IRSN Mephisto-15_1
 MEPH11 = ThermoData MEPHISTA Nuclear Fuels TDB v11.2
 TCAQ3 = Aqueous Solution v3.0
 TCAQ2 = Aqueous Solution v2.7
 AQS2 = TGG Aqueous Solution Database v2.6
 GCE2 = TGG Geochemical/Environmental TDB v2.3
 FEDEMO = Iron Demo Database v4.0
 ALDEMO = Aluminum Demo Database v4.0
 NI DEMO = Nickel Demo Database v2.0
 CUDEMO = Copper Demo Database v1.0
 SLDEMO = Solder Demo Database v1.0
 OXDEMO = Oxide Demo Database v3.0
 SUBDEMO = Substance Demo Database v1.0
 PAQ2 = Public Aqueous Soln (SIT) TDB v2.5
 PG35 = PG35 Binary Semi-Conductors TDB v1.3
 PURE5 = SGTE Unary (Pure Elements) TDB v5.2
 MOB2 = Alloys Mobility v2.7
 MOB1 = Alloys Mobility v1.3
 MOBFE1 = Steels/Fe-Alloys Mobility v1.1
 MOBFE2 = Steels/Fe-Alloys Mobility v2.0
 MOBFE3 = Steels/Fe-Alloys Mobility v3.0
 MOBFE4 = Steels/Fe-Alloys Mobility v4.0
 MOBFE5 = Steels/Fe-Alloys Mobility v5.0
 MOBFE6 = Steels/Fe-Alloys Mobility v6.0
 MOBN15 = Ni-Alloys Mobility v5.1
 MOBN14 = Ni-Alloys Mobility v4.1
 MOBN13 = Ni-Alloys Mobility v3.2
 MOBN12 = Ni-Alloys Mobility v2.4
 MOBN11 = Ni-Alloys Mobility v1.10
 MOBAL6 = Al-Alloys Mobility v6.0
 MOBAL5 = Al-Alloys Mobility v5.0
 MOBAL4 = Al-Alloys Mobility v4.0
 MOBAL3 = Al-Alloys Mobility v3.0
 MOBAL2 = Al-Alloys Mobility v2.0
 MOBAL1 = Al-Alloys Mobility v1.0
 MOBCU1 = Cu-Alloys Mobility v1.0

```

MOBCU2 = Cu-Alloys Mobility v2.0
MOBCU3 = Cu-Alloys Mobility v3.0
MOBCU4 = Cu-Alloys Mobility v4.0
MOBHEA1 = High Entropy Alloys Mobility v1.0
MOBHEA2 = High Entropy Alloys Mobility v2.0
MOBMG2 = Mg-Alloys Mobility v2.0
MOBMG1 = Mg-Alloys Mobility v1.0
MOBSI1 = Si-Alloys Mobility v1.0
MOBSDL1 = Solder-Alloys Mobility v1.1
MOBTI4 = Ti-Alloys Mobility v4.0
MOBTI3 = Ti-Alloys Mobility v3.1
MOBTI2 = Ti-Alloys Mobility v2.0
MOBTI1 = Ti-Alloys Mobility v1.0
MALDEMO = Al-Alloys Mobility demo database v2.0
MFEDEMO = Fe-Alloys Mobility demo database v2.0
MNIDEMO = Ni-Alloys Mobility demo database v1.0
MCUDEMO = Cu-Alloys Mobility demo database v1.0
USER = User defined Database

DATABASE NAME /FEDEMO/: mfedemo
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys fe ni
FE NI DEFINED
APP: rej ph * all FCC_A1 REJECTED
BCC_A2 FCC_A1 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
-OK-
APP:
APP: @@ ENTER THE DICTRA MONITOR WHERE THE SYSTEM IS SET UP
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@ CHECK THE DIFFUSIVITES
DIC> @@
DIC> check-diffusion-matrix
OUTPUT TO SCREEN OR FILE /SCREEN/:
PHASE NAME : fcc
DEPENDENT COMPONENT ? /NI/: fe
CONCENTRATION OF NI IN U-FRACTION /1/: 0.3
Pressure /100000/: 101325
Temperature /298.15/: 1409
OPTION ( dlphmx0ez or * ) /D/: dl

Dkj (reduced n=FE)
k / j NI
NI +3.83335E-15
LOkj = Uk*Mvak IF (kES) ELSE Uk*Yva*Mvak
k / j FE NI
FE +9.0901E-20
NI +9.0901E-20

Volume = 1.000000000000000E-005

DIC>
DIC>
DIC>@@<Hit_return_to_continue>
DIC>
DIC> @@
DIC> @@ USE STEPPING IN POLY-3 TO CALCULATE THE DIFFUSIVITIES VS. COMPOSITION
DIC> @@
DIC> go p-3
POLY: s-c t=1409,p=101325,n=1,x(ni)=0.3
POLY: c-e
Using global minimization procedure
Calculated 209 grid points in 2 s
Found the set of lowest grid points in 0 s
Calculated POLY solution 0 s, total time 2 s
POLY:

POLY: s-a-v
Axis number: /1/: 1
Condition /NONE/: x(ni)
Min value /0/: 0
Max value /1/: 1
Increment /.025/: 1e-3
POLY:
POLY: step
Option? /NORMAL/: normal
No initial equilibrium, using default
Step will start from axis value 0.300000
...OK

Phase Region from 0.300000 for:
FCC_A1
Global test at 3.08000E-01 .... OK
Global test at 3.18000E-01 .... OK
Global test at 3.28000E-01 .... OK
Global test at 3.38000E-01 .... OK
Global test at 3.48000E-01 .... OK
Global test at 3.58000E-01 .... OK
Global test at 3.68000E-01 .... OK
Global test at 3.78000E-01 .... OK
Global test at 3.88000E-01 .... OK
Global test at 3.98000E-01 .... OK

```

```
Global test at 4.08000E-01 .... OK
Global test at 4.18000E-01 .... OK
Global test at 4.28000E-01 .... OK
Global test at 4.38000E-01 .... OK
Global test at 4.48000E-01 .... OK
Global test at 4.58000E-01 .... OK
Global test at 4.68000E-01 .... OK
Global test at 4.78000E-01 .... OK
Global test at 4.88000E-01 .... OK
Global test at 4.98000E-01 .... OK
Global test at 5.08000E-01 .... OK
Global test at 5.18000E-01 .... OK
Global test at 5.28000E-01 .... OK
Global test at 5.38000E-01 .... OK
Global test at 5.48000E-01 .... OK
Global test at 5.58000E-01 .... OK
Global test at 5.68000E-01 .... OK
Global test at 5.78000E-01 .... OK
Global test at 5.88000E-01 .... OK
Global test at 5.98000E-01 .... OK
Global test at 6.08000E-01 .... OK
Global test at 6.18000E-01 .... OK
Global test at 6.28000E-01 .... OK
Global test at 6.38000E-01 .... OK
Global test at 6.48000E-01 .... OK
Global test at 6.58000E-01 .... OK
Global test at 6.68000E-01 .... OK
Global test at 6.78000E-01 .... OK
Global test at 6.88000E-01 .... OK
Global test at 6.98000E-01 .... OK
Global test at 7.08000E-01 .... OK
Global test at 7.18000E-01 .... OK
Global test at 7.28000E-01 .... OK
Global test at 7.38000E-01 .... OK
Global test at 7.48000E-01 .... OK
Global test at 7.58000E-01 .... OK
Global test at 7.68000E-01 .... OK
Global test at 7.78000E-01 .... OK
Global test at 7.88000E-01 .... OK
Global test at 7.98000E-01 .... OK
Global test at 8.08000E-01 .... OK
Global test at 8.18000E-01 .... OK
Global test at 8.28000E-01 .... OK
Global test at 8.38000E-01 .... OK
Global test at 8.48000E-01 .... OK
Global test at 8.58000E-01 .... OK
Global test at 8.68000E-01 .... OK
Global test at 8.78000E-01 .... OK
Global test at 8.88000E-01 .... OK
Global test at 8.98000E-01 .... OK
Global test at 9.08000E-01 .... OK
Global test at 9.18000E-01 .... OK
Global test at 9.28000E-01 .... OK
Global test at 9.38000E-01 .... OK
Global test at 9.48000E-01 .... OK
Global test at 9.58000E-01 .... OK
Global test at 9.68000E-01 .... OK
Global test at 9.78000E-01 .... OK
Global test at 9.88000E-01 .... OK
Global test at 9.98000E-01 .... OK
Terminating at 1.00000
Calculated 703 equilibria
```

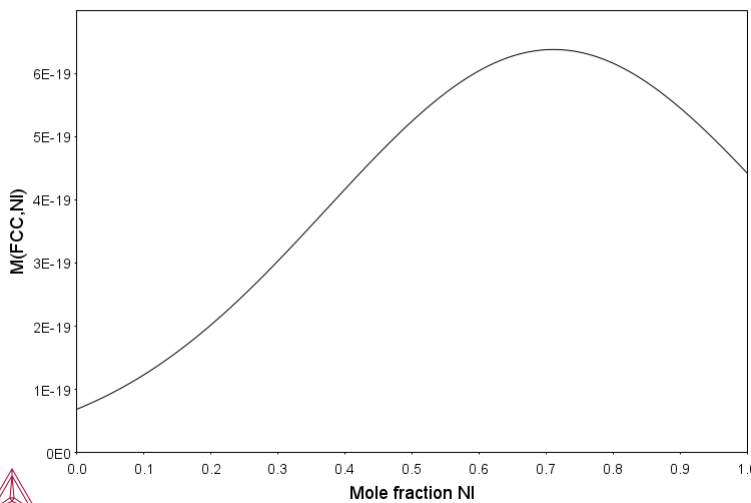
```
Phase Region from 0.300000 for:
FCC_A1
Global test at 2.92000E-01 .... OK
Global test at 2.82000E-01 .... OK
Global test at 2.72000E-01 .... OK
Global test at 2.62000E-01 .... OK
Global test at 2.52000E-01 .... OK
Global test at 2.42000E-01 .... OK
Global test at 2.32000E-01 .... OK
Global test at 2.22000E-01 .... OK
Global test at 2.12000E-01 .... OK
Global test at 2.02000E-01 .... OK
Global test at 1.92000E-01 .... OK
Global test at 1.82000E-01 .... OK
Global test at 1.72000E-01 .... OK
Global test at 1.62000E-01 .... OK
Global test at 1.52000E-01 .... OK
Global test at 1.42000E-01 .... OK
Global test at 1.32000E-01 .... OK
Global test at 1.22000E-01 .... OK
Global test at 1.12000E-01 .... OK
Global test at 1.02000E-01 .... OK
Global test at 9.20000E-02 .... OK
Global test at 8.20000E-02 .... OK
Global test at 7.20000E-02 .... OK
Global test at 6.20000E-02 .... OK
Global test at 5.20000E-02 .... OK
Global test at 4.20000E-02 .... OK
Global test at 3.20000E-02 .... OK
Global test at 2.20000E-02 .... OK
Global test at 1.20000E-02 .... OK
Global test at 2.00000E-03 .... OK
Terminating at 0.100000E-11
Calculated 303 equilibria
*** Buffer saved on file: C:\Users\vagrant\AppData\Local\Temp\RESULT_002.POLY3
```

```
POLY:
POLY: @@
POLY: @@ ENTER THE POST PROCESSOR AND PLOT THE RESULT
POLY: @@
POLY: post
    POLY-3 POSTPROCESSOR VERSION 3.2
POST:
POST:
POST:
POST: @@
POST: @@ PLOT THE MOBILITY OF Ni VS. X(Ni)
POST: @@
POST: s-d-a y m(fcc,nii)
POST: s-d-a x m-f ni
POST:
POST:
POST: SET_EXP_FILE_FORMAT 5
```

```

POST: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST: SET_EXP_FILE_FORMAT 10
POST:
POST: plot
2021.05.13.13.03.12
MFEDEMO: FE,Ni
T=1409,P=1.01325E5,N=1.

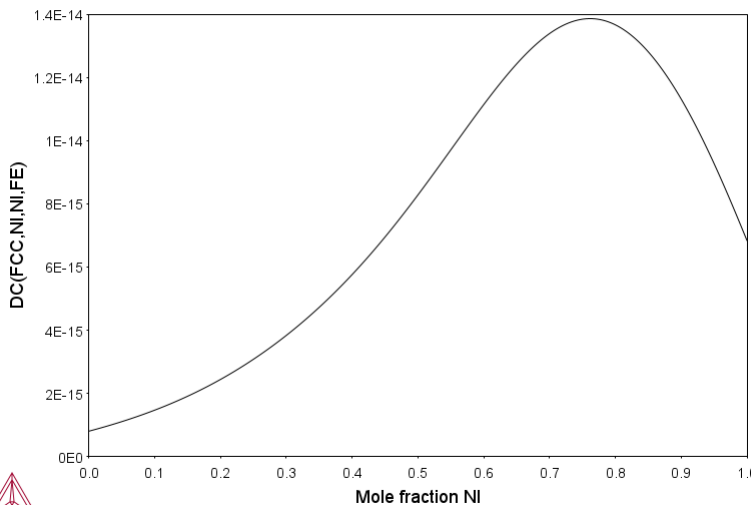
```



```

POST:
POST:
POST:
POST:@?<Hit_return_to_continue>
POST:
POST: @@
POST: @@ THEN PLOT THE DIFFUSIVITY OF Ni VS. X(Ni)
POST: @@
POST: s-d-a y dc(fcc,ni,ni,fe)
POST:
POST:
POST: SET_EXP_FILE_FORMAT 5
POST: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST: SET_EXP_FILE_FORMAT 10
POST:
POST: plot
2021.05.13.13.03.13
MFEDEMO: FE,Ni
T=1409,P=1.01325E5,N=1.

```

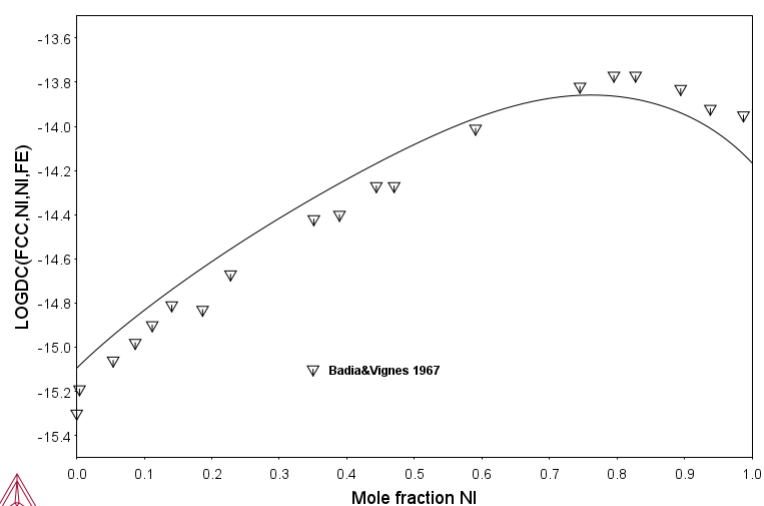


```

POST:
POST:
POST:@?<Hit_return_to_continue>
POST:
POST: @@
POST: @@ PLOT THE LOGARITHM OF DC AND APPEND THE EXPERIMENTAL DATA
POST: @@
POST: s-d-a y logdc(fcc,ni,ni,fe)
POST:
POST: app y feni.exp
PROLOGUE NUMBER: /0/: 1
DATASET NUMBER(s): /-1/: 1
POST:
POST: s-s-s y n -15.5 -13.5
POST:
POST:
POST: SET_EXP_FILE_FORMAT 5
POST: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST: SET_EXP_FILE_FORMAT 10
POST:
POST: plot

```

2021.05.13.03.14
MFEDEMO: FE, NI
T=1409, P=1.01325E5, N=1.



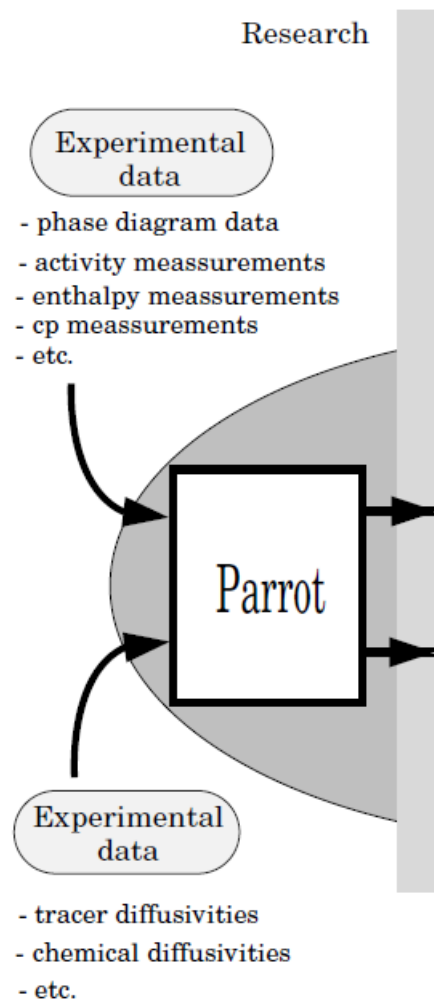
POST:
POST:
POST:@?<Hit_return_to_continue>
POST:
POST: set-inter
POST:



Example exg2

Optimization of mobilities in Ni-Al fcc alloys

A file for reading thermodynamic data and setting up the kinetic parameters which are needed for an optimization of the FCC phase in the binary Ni-Al system. See also A. Engström and J. Ågren: ("Assessment of Diffusional Mobilities in Face-Centered Cubic Ni-Cr-Al Alloys" in Z. METALLKUNDE, Feb. 1996).



exg2-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exg2\setup.DCM.test"
SYS: i>{@
NO SUCH COMMAND, USE HELP
SYS: @@ Kinetic data example.
SYS: @@ Optimization of mobilities in Ni-Al fcc alloys
SYS: @@ A file for reading thermodynamic data and setting up the kinetic
SYS: @@ parameters that are needed for an optimization of the FCC phase
SYS: @@ in the binary Ni-Al system.
SYS: @@ See also A. Engström and J. Ågren: ("Assessment of Diffusional
SYS: @@ Mobilities in Face-Centered Cubic Ni-Cr-Al Alloys" in
SYS: @@ Z. Metallkunde, Feb. 1996).
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exg2_setup.DCM
SYS:
SYS:
SYS: @@
SYS: @@ EXPLICITLY SELECTING GES VERSION 5 BECAUSE PARAMETER OPTIMIZATION
SYS: @@ IS NOT SUPPORTED IN GES VERSION 6
SYS: @@
SYS: set ges-version 5
SYS:
SYS: @@
SYS: @@ RETRIEVE THERMODYNAMIC DATA FROM A USER-DEFINED DATABASE
SYS: @@
SYS: go data
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: sw us tdata.TDB
Current database: User defined Database
This database does not support the DATABASE_INFORMATION command

VA   DEFINED
TDB_USER: def-sys al ni
AL          NI   DEFINED
TDB_USER: rej ph *
LIQUID      B2_BCC
FCC_A1      GAMMA_PRIME REJECTED
TDB_USER: rest ph fcc_a1
FCC_A1 RESTORED
TDB_USER: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....
-OK-
TDB_USER: @@
TDB_USER: @@ APPEND THE KINETIC DATA FROM THE MOBILITY DATABASE IN ORDER TO
TDB_USER: @@ HAVE SOME DUMMY PARAMETERS.
TDB_USER: @@
TDB_USER: app mob2
Current database: Alloys Mobility v2.7

VA   DEFINED
GAS:G REJECTED
APP: def-sys al ni
AL          NI   DEFINED
APP: rej ph *
BCC_A2      FCC_A1          M4N
HCP_A3      LIQUID:L REJECTED
APP: res ph fcc_a1
FCC_A1 RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data
'This parameter has not been assessed'
'A. Engstrom and J. Agren: Z. Metallkunde 87(1996)92-97;
'Al, Cr and Ni diffusion in fcc Al-Cr-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27;
'Ni self-diffusion'
-OK-
APP: @@
APP: @@ GO TO THE DICTRA MODULE AND DEFINE THE KINETIC PARAMETERS. THE
APP: @@ VARIABLES V1,V2,V3 AND V4 ARE TO BE OPTIMIZED. NOTE THAT IF
APP: @@ YOU ARE OPTIMIZING PARAMETERS FOR A PHASE WITH MAGNETIC
APP: @@ CONTRIBUTION I.E. USING BOTH MF- AND MQ-PARAMETERS, YOU
APP: @@ MIGHT HAVE TO ENTER THE PARROT MODULE AND GO BACK BEFORE
APP: @@ ENTERING PARAMETERS CONTAINING VARIABLES.
APP: @@
APP: go dic_par
PARROT VERSION 5.3d RUNNING ON PC/WINDOWS NT
Developed at the Division of Physical Metallurgy
```

```
PARROT:  
PARROT:  
PARROT: go d-m  
NO TIME STEP DEFINED  
DIC>  
DIC> @@ MOBILITY OF A1 IN A1  
DIC> ENTER-MOB-DATA  
PARAMETER: MQ(FCC_A1&AL,AL:VA) 298.15 -142000+R*T*LN(1.71E-4); 6000 N  
MQ(FCC_A1&AL#1,AL:VA;0)  
DIC>  
DIC> @@ MOBILITY OF A1 IN Ni  
DIC> ENTER-MOB-DATA  
PARAMETER: MQ(FCC_A1&AL,NI:VA) 298.15 -284000+R*T*LN(7.5E-4); 6000 N  
MQ(FCC_A1&AL#1,NI:VA;0)  
DIC>  
DIC> @@ MOBILITY OF A1 INTERACTION BETWEEN A1 AND Ni  
DIC> ENTER-MOB-DATA  
PARAMETER: MQ(FCC_A1&AL,AL,NI:VA;0) 298.15 V1+V2*T; 6000 N  
MQ(FCC_A1&AL#1,AL,NI:VA;0)  
DIC>  
DIC> @@ MOBILITY OF Ni IN A1  
DIC> ENTER-MOB-DATA  
PARAMETER: MQ(FCC_A1&NI,AL:VA) 298.15 -145900+R*T*LN(4.4E-4); 6000 N  
MQ(FCC_A1&NI#1,AL:VA;0)  
DIC>  
DIC> @@ MOBILITY OF Ni IN Ni  
DIC> ENTER-MOB-DATA  
PARAMETER: MQ(FCC_A1&NI,NI:VA) 298.15 -287000-69.8*T; 6000 N  
MQ(FCC_A1&NI#1,NI:VA;0)  
DIC>  
DIC> @@  
DIC> @@ GO TO PARROT AND SAVE THE SET UP TO FILE  
DIC> @@  
DIC> go dic_parrot  
PARROT VERSION 5.3d RUNNING ON PC/WINDOWS NT  
PARROT: create-new-store-file opt  
PARROT:  
PARROT: set-inter  
--OK--  
PARROT:
```

exg2-run

```
PARROT:About
NO SUCH COMMAND, USE HELP
PARROT:PARROT:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exg2\run.DCM.test"
PARROT: @@ exg2_run.DCM
PARROT:
PARROT: -----
PARROT: @@ FILE FOR DOING THE OPTIMIZATION IN PARROT
PARROT: -----
PARROT:
PARROT: @@
PARROT: @@ GO TO PARROT AND READ THE SETUP
PARROT: @@
PARROT: go dic_parrot
PARROT: VERSION 5.3d RUNNING ON PC/WINDOWS NT
PARROT: set-store-file opt
PARROT:
PARROT:
PARROT: @@
PARROT: @@ COMPILE THE EXPERIMENTAL DATA IN exp.DOP INTO STRUCTURED BINARY DATA.
PARROT: @@
PARROT: compile-experiments exp
OUTPUT TO SCREEN OR FILE /SCREEN/:
INITIATE STORE FILE: /Y/:

-----+
$ DOP-FILE CONTAINING EXPERIMENTAL INFORMATION USED DURING THE
$ OPTIMIZATION IN PARROT (COMPARE WITH POP-FILE USED WHEN EVALUATING
$ THERMODYNAMIC DATA). THE EXPERIMENTAL DATA HERE STEAM FROM A STUDY BY
$ YAMAMOTO ET AL. TRANS. JPN. INST. MET. VOL. 21, NO. 9 (1980), P. 601.
$ CONSULT THE THERMO-CALC USER'S GUIDE TO LEARN MORE ABOUT SYNTAXES
$ FOR OPTIMIZATION OF THERMODYNAMIC DATA.
$-----+


ENTER CONST P0=101325

TABLE_HEAD 10
CREATE_NEW 0010,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.01055
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.6:.1
CREATE_NEW 0011,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.02032
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.56:.1
CREATE_NEW 0012,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.02957
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.65:.1
CREATE_NEW 0013,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.03884
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.52:.1
CREATE_NEW 0014,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.03884
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.52:.1
CREATE_NEW 0015,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.04927
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.48:.1
CREATE_NEW 0016,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.06062
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.43:.1
CREATE_NEW 0017,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.07029
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.41:.1
CREATE_NEW 0018,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.08113
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.37:.1
CREATE_NEW 0019,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.09166
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.32:.1
CREATE_NEW 0020,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.0945
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.28:.1
CREATE_NEW 0021,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.1099
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.24:.1
CREATE_NEW 0022,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.1207
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.2:.1
CREATE_NEW 0023,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.129
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.18:.1
CREATE_NEW 0024,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.1392
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.16:.1
```

```

CREATE_NEW 0025,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.1503
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.1:.
CREATE_NEW 0026,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0
S-C X(AL)=.1589
EXPERIMENT LOGDC(FCC_A1,AL,AL,NI)=-12.08:.
CREATE_NEW 0027,1
C-S PH FCC=ENT 1
S-C T=1573,N=1,P=P0

```

output ignored...

... output resumed

V3	-1.32972463E+05	-1.32959167E+05	-1.32959167E+05	3.97449919E+00	3.97449919E+00
	3.97449919E+00	3.97449919E+00	3.97449919E+00	3.97449919E+00	3.97449919E+00
	3.97449919E+00	3.97449919E+00	3.97449919E+00	3.97449919E+00	3.97449919E+00
	3.97449919E+00	3.97449919E+00	3.97449919E+00	3.97449919E+00	3.97449919E+00
V4	7.81857805E+01	7.81857805E+01	7.81857805E+01	4.46532140E+00	4.46532140E+00
	4.46532140E+00	4.46532140E+00	4.46532140E+00	4.46532140E+00	4.46532140E+00
	4.46532140E+00	4.46532140E+00	4.46532140E+00	4.46532140E+00	4.46532140E+00
	4.46532140E+00	4.46532140E+00	4.46532140E+00	4.46532140E+00	4.46532140E+00

NUMBER OF OPTIMIZING VARIABLES : 4
ALL OTHER VARIABLES ARE FIX WITH THE VALUE ZERO
THE SUM OF SQUARES HAS CHANGED FROM 2.02977003E+01 TO 2.02976923E+01
DEGREES OF FREEDOM 113. REDUCED SUM OF SQUARES 1.79625595E-01

Sorry, LIST-DATA disabled for this database

===== BLOCK NUMBER 1

DEFINED CONSTANTS	P0=101325
10 LOGDC(F...,AL,NI)=-12.6	-12.56 0.10 4.3827E-02 0.4383
11 LOGDC(F...,AL,NI)=-12.56	-12.55 0.10 1.2341E-02 0.1234
12 LOGDC(F...,AL,NI)=-12.65	-12.53 0.10 0.1167 1.167
13 LOGDC(F...,AL,NI)=-12.52	-12.51 0.10 5.8843E-03 5.8843E-02
14 LOGDC(F...,AL,NI)=-12.52	-12.51 0.10 5.8843E-03 5.8843E-02
15 LOGDC(F...,AL,NI)=-12.48	-12.49 0.10 -8.4123E-03 -8.4123E-02
16 LOGDC(F...,AL,NI)=-12.43	-12.46 0.10 -2.6851E-02 -0.2685
17 LOGDC(F...,AL,NI)=-12.41	-12.43 0.10 -1.7821E-02 -0.1782
18 LOGDC(F...,AL,NI)=-12.37	-12.39 0.10 -2.3493E-02 -0.2349
19 LOGDC(F...,AL,NI)=-12.32	-12.36 0.10 -3.8626E-02 -0.3863
20 LOGDC(F...,AL,NI)=-12.28	-12.33 0.10 -5.1933E-02 -0.5193
21 LOGDC(F...,AL,NI)=-12.24	-12.29 0.10 -5.4919E-02 -0.5492
22 LOGDC(F...,AL,NI)=-12.2	-12.26 0.10 -5.5128E-02 -0.5513
23 LOGDC(F...,AL,NI)=-12.18	-12.22 0.10 -4.3400E-02 -0.4340
24 LOGDC(F...,AL,NI)=-12.16	-12.18 0.10 -2.2927E-02 -0.2293
25 LOGDC(F...,AL,NI)=-12.1	-12.14 0.10 -3.6888E-02 -0.3689
26 LOGDC(F...,AL,NI)=-12.08	-12.10 0.10 -1.9703E-02 -0.1970
27 LOGDC(F...,AL,NI)=-12.02	-12.04 0.10 -2.2272E-02 -0.2227
28 LOGDC(F...,AL,NI)=-11.98	-11.99 0.10 -9.0917E-03 -9.0917E-02
29 LOGDC(F...,AL,NI)=-11.94	-11.95 0.10 -9.1618E-03 -9.1618E-02
30 LOGDC(F...,AL,NI)=-13	-12.86 0.10 0.1360 1.360
31 LOGDC(F...,AL,NI)=-12.96	-12.85 0.10 0.1059 1.059
32 LOGDC(F...,AL,NI)=-12.92	-12.84 0.10 8.4799E-02 0.8480
33 LOGDC(F...,AL,NI)=-12.9	-12.82 0.10 8.2549E-02 0.8255
34 LOGDC(F...,AL,NI)=-12.77	-12.79 0.10 -1.5173E-02 -0.1517
35 LOGDC(F...,AL,NI)=-12.74	-12.75 0.10 -1.1192E-02 -0.1119
36 LOGDC(F...,AL,NI)=-12.82	-12.72 0.10 9.5223E-02 0.9522
37 LOGDC(F...,AL,NI)=-12.82	-12.72 0.10 9.5223E-02 0.9522
38 LOGDC(F...,AL,NI)=-12.69	-12.68 0.10 5.7856E-03 5.7856E-02
39 LOGDC(F...,AL,NI)=-12.65	-12.65 0.10 2.6229E-03 2.6229E-02
40 LOGDC(F...,AL,NI)=-12.64	-12.62 0.10 1.9222E-02 0.1922
41 LOGDC(F...,AL,NI)=-12.61	-12.58 0.10 2.8492E-02 0.2849
42 LOGDC(F...,AL,NI)=-12.55	-12.54 0.10 9.2091E-03 9.2091E-02
43 LOGDC(F...,AL,NI)=-12.53	-12.51 0.10 2.1034E-02 0.2103
44 LOGDC(F...,AL,NI)=-12.47	-12.47 0.10 2.0402E-03 2.0402E-02
45 LOGDC(F...,AL,NI)=-12.41	-12.42 0.10 -1.0621E-02 -0.1062
46 LOGDC(F...,AL,NI)=-12.38	-12.38 0.10 -1.6323E-04 -1.6323E-03
47 LOGDC(F...,AL,NI)=-12.36	-12.32 0.10 3.6256E-02 0.3626
48 LOGDC(F...,AL,NI)=-12.36	-12.32 0.10 3.6256E-02 0.3626
49 LOGDC(F...,AL,NI)=-12.3	-12.27 0.10 3.0494E-02 0.3049
50 LOGDC(F...,AL,NI)=-13.23	-13.19 0.10 3.8416E-02 0.3842
51 LOGDC(F...,AL,NI)=-13.23	-13.19 0.10 3.8416E-02 0.3842
52 LOGDC(F...,AL,NI)=-13.19	-13.18 0.10 1.3209E-02 0.1321
53 LOGDC(F...,AL,NI)=-13.15	-13.16 0.10 -5.5530E-03 -5.5530E-02
54 LOGDC(F...,AL,NI)=-13.12	-13.13 0.10 -1.4736E-02 -0.1474
55 LOGDC(F...,AL,NI)=-13.09	-13.10 0.10 -1.2913E-02 -0.1291
56 LOGDC(F...,AL,NI)=-13.06	-13.07 0.10 -6.9397E-03 -6.9397E-02
57 LOGDC(F...,AL,NI)=-13.02	-13.04 0.10 -1.5971E-02 -0.1597
58 LOGDC(F...,AL,NI)=-12.99	-13.00 0.10 -1.0672E-02 -0.1067
59 LOGDC(F...,AL,NI)=-12.96	-12.96 0.10 8.0069E-04 8.0069E-03
60 LOGDC(F...,AL,NI)=-12.91	-12.93 0.10 -2.0470E-02 -0.2047
61 LOGDC(F...,AL,NI)=-12.88	-12.89 0.10 -7.3947E-03 -7.3947E-02
62 LOGDC(F...,AL,NI)=-12.86	-12.85 0.10 1.4312E-02 0.1431
63 LOGDC(F...,AL,NI)=-12.86	-12.85 0.10 1.4312E-02 0.1431
64 LOGDC(F...,AL,NI)=-12.83	-12.81 0.10 1.7076E-02 0.1708
65 LOGDC(F...,AL,NI)=-12.8	-12.77 0.10 2.9558E-02 0.2956
66 LOGDC(F...,AL,NI)=-12.75	-12.72 0.10 2.5293E-02 0.2529
67 LOGDC(F...,AL,NI)=-12.71	-12.68 0.10 2.6861E-02 0.2686
68 LOGDC(F...,AL,NI)=-12.67	-12.63 0.10 4.4033E-02 0.4403
70 LOGDC(F...,AL,NI)=-13.5	-13.54 0.10 -4.2456E-02 -0.4246
71 LOGDC(F...,AL,NI)=-13.47	-13.53 0.10 -5.5160E-02 -0.5516
72 LOGDC(F...,AL,NI)=-13.45	-13.50 0.10 -5.0384E-02 -0.5038
73 LOGDC(F...,AL,NI)=-13.42	-13.48 0.10 -5.6139E-02 -0.5614
74 LOGDC(F...,AL,NI)=-13.39	-13.44 0.10 -5.3034E-02 -0.5303
75 LOGDC(F...,AL,NI)=-13.36	-13.40 0.10 -4.3865E-02 -0.4386
76 LOGDC(F...,AL,NI)=-13.34	-13.37 0.10 -3.0943E-02 -0.3094
77 LOGDC(F...,AL,NI)=-13.31	-13.33 0.10 -2.0539E-02 -0.2054
78 LOGDC(F...,AL,NI)=-13.24	-13.29 0.10 -5.0311E-02 -0.5031
79 LOGDC(F...,AL,NI)=-13.22	-13.26 0.10 -4.1318E-02 -0.4132
80 LOGDC(F...,AL,NI)=-13.19	-13.22 0.10 -2.6230E-02 -0.2623
81 LOGDC(F...,AL,NI)=-13.13	-13.17 0.10 -4.3098E-02 -0.4310
82 LOGDC(F...,AL,NI)=-13.12	-13.14 0.10 -1.9773E-02 -0.1977

83	LOGDC(F...,AL,NI)=-13.08	-13.09	0.10	-1.4457E-02	-0.1446
84	LOGDC(F...,AL,NI)=-13.04	-13.05	0.10	-9.4618E-03	-9.4618E-02
85	LOGDC(F...,AL,NI)=-13.03	-13.01	0.10	2.2404E-02	0.2240
90	LOGDC(F...,AL,NI)=-13.97	-13.92	0.10	5.1241E-02	0.5124
91	LOGDC(F...,AL,NI)=-13.92	-13.90	0.10	2.0497E-02	0.2050
92	LOGDC(F...,AL,NI)=-13.88	-13.87	0.10	9.3920E-03	9.3920E-02
93	LOGDC(F...,AL,NI)=-13.85	-13.84	0.10	5.2264E-03	5.2264E-02
94	LOGDC(F...,AL,NI)=-13.82	-13.81	0.10	1.1044E-02	0.1104
95	LOGDC(F...,AL,NI)=-13.78	-13.77	0.10	1.3352E-02	0.1335
96	LOGDC(F...,AL,NI)=-13.9	-13.74	0.10	0.1640	1.640
97	LOGDC(F...,AL,NI)=-13.85	-13.69	0.10	0.1576	1.576
98	LOGDC(F...,AL,NI)=-13.65	-13.65	0.10	1.5604E-03	1.5604E-02
99	LOGDC(F...,AL,NI)=-13.62	-13.62	0.10	4.9052E-03	4.9052E-02
100	LOGDC(F...,AL,NI)=-13.57	-13.57	0.10	-9.5589E-04	-9.5589E-03
101	LOGDC(F...,AL,NI)=-13.52	-13.53	0.10	-5.4819E-03	-5.4819E-02
102	LOGDC(F...,AL,NI)=-13.47	-13.49	0.10	-2.0262E-02	-0.2026
103	LOGDC(F...,AL,NI)=-13.45	-13.45	0.10	4.9628E-03	4.9628E-02
104	LOGDC(F...,AL,NI)=-13.4	-13.40	0.10	2.8073E-03	2.8073E-02
110	LOGDC(F...,AL,NI)=-14.32	-14.32	0.10	-3.9760E-03	-3.9760E-02
111	LOGDC(F...,AL,NI)=-14.32	-14.30	0.10	1.9307E-02	0.1931
112	LOGDC(F...,AL,NI)=-14.28	-14.27	0.10	1.0664E-02	0.1066
113	LOGDC(F...,AL,NI)=-14.25	-14.24	0.10	8.9412E-03	8.9412E-02
114	LOGDC(F...,AL,NI)=-14.22	-14.20	0.10	1.8329E-02	0.1833
115	LOGDC(F...,AL,NI)=-14.17	-14.16	0.10	1.2994E-02	0.1299
116	LOGDC(F...,AL,NI)=-14.15	-14.12	0.10	3.0489E-02	0.3049
117	LOGDC(F...,AL,NI)=-14.1	-14.07	0.10	2.5593E-02	0.2559
118	LOGDC(F...,AL,NI)=-14.03	-14.03	0.10	-1.4483E-03	-1.4483E-02
119	LOGDC(F...,AL,NI)=-14	-14.00	0.10	2.7074E-03	2.7074E-02
120	LOGDC(F...,AL,NI)=-13.95	-13.95	0.10	-1.9282E-03	-1.9282E-02
121	LOGDC(F...,AL,NI)=-13.9	-13.91	0.10	-5.2349E-03	-5.2349E-02
122	LOGDC(F...,AL,NI)=-13.85	-13.87	0.10	-1.6645E-02	-0.1665
130	LOGDC(F...,AL,NI)=-14.73	-14.76	0.10	-3.0493E-02	-0.3049
131	LOGDC(F...,AL,NI)=-14.71	-14.73	0.10	-2.3757E-02	-0.2376
132	LOGDC(F...,AL,NI)=-14.68	-14.70	0.10	-1.9446E-02	-0.1945
133	LOGDC(F...,AL,NI)=-14.66	-14.67	0.10	-8.5997E-03	-8.5997E-02
134	LOGDC(F...,AL,NI)=-14.61	-14.63	0.10	-1.5314E-02	-0.1531
135	LOGDC(F...,AL,NI)=-14.61	-14.63	0.10	-1.5314E-02	-0.1531
136	LOGDC(F...,AL,NI)=-14.58	-14.58	0.10	3.4408E-03	3.4408E-02
137	LOGDC(F...,AL,NI)=-14.54	-14.54	0.10	3.3897E-03	3.3897E-02
138	LOGDC(F...,AL,NI)=-14.5	-14.49	0.10	9.2113E-03	9.2113E-02
139	LOGDC(F...,AL,NI)=-14.46	-14.44	0.10	1.5390E-02	0.1539
140	LOGDC(F...,AL,NI)=-14.41	-14.41	0.10	1.6005E-03	1.6005E-02
141	LOGDC(F...,AL,NI)=-14.35	-14.36	0.10	-1.1351E-02	-0.1135
142	LOGDC(F...,AL,NI)=-14.27	-14.31	0.10	-4.1487E-02	-0.4149
143	LOGDC(F...,AL,NI)=-14.2	-14.27	0.10	-7.4774E-02	-0.7477

PARROT:
PARROT:
PARROT: set-inter
--OK--
PARROT:

exg2-plot

```
PARROT:About
NO SUCH COMMAND, USE HELP
PARROT:PARROT:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exg2\plot.DCM.test"
PARROT: @@ exg2_plot.DCM
PARROT:
PARROT: @-----
PARROT: @@ FILE FOR PLOTTING THE RESULT AFTER THE OPTIMIZATION. HERE
PARROT: @@ DIFFUSIVITIES CALCULATED FROM THE OPTIMIZED VARIABLES ARE
PARROT: @@ COMPARED WITH EXPERIMENTALLY MEASURED ONES.
PARROT: @-----
PARROT:
PARROT: @@
PARROT: @@ GO TO PARROT AND READ THE FILE CONTAINING THE RESULT FROM
PARROT: @@ THE OPTIMIZATION.
PARROT: @@
PARROT: go dic_parrot
PARROT VERSION 5.3d RUNNING ON PC/WINDOWS NT
PARROT: set-store-file opt
PARROT:
PARROT: @@
PARROT: @@ GO TO POLY3 AND STEP IN X(AL)
PARROT: @@
PARROT: go p-3
POLY: s-c n=1,p=101325,t=1573
POLY: s-c x(al)=.1
POLY: c-e,,
Using global minimization procedure
Calculated          209 grid points in           0 s
Found the set of lowest grid points in           0 s
Calculated POLY solution      0 s, total time   0 s
POLY: add,
POLY:

POLY: s-a-v 1 x(al) 0.0001 .20 0.001,,
POLY: step
Option? /NORMAL/:
Step will start from axis value  0.100000
...OK

Phase Region from  0.100000    for:
  FCC_A1
Global test at  1.08000E-01 .... OK
Global test at  1.18000E-01 .... OK
Global test at  1.28000E-01 .... OK
Global test at  1.38000E-01 .... OK
Global test at  1.48000E-01 .... OK
Global test at  1.58000E-01 .... OK
Global test at  1.68000E-01 .... OK
Global test at  1.78000E-01 .... OK
Global test at  1.88000E-01 .... OK
Global test at  1.98000E-01 .... OK
Terminating at  0.200000
Calculated  103 equilibria

Phase Region from  0.100000    for:
  FCC_A1
Global test at  9.20000E-02 .... OK
Global test at  8.20000E-02 .... OK
Global test at  7.20000E-02 .... OK
Global test at  6.20000E-02 .... OK
Global test at  5.20000E-02 .... OK
Global test at  4.20000E-02 .... OK
Global test at  3.20000E-02 .... OK
Global test at  2.20000E-02 .... OK
Global test at  1.20000E-02 .... OK
Global test at  2.00000E-03 .... OK
Terminating at  0.100000E-03
Calculated  103 equilibria
*** Buffer saved on file: C:\Users\vagrant\AppData\Local\Temp\RESULT_002.POLY3
POLY: @@
POLY: @@ REPEAT THE PROCEDURE FOR SOME OTHER TEMPERATURES
POLY: @@
POLY: s-c t=1523,x(al)=.1
POLY: c-e,,
Using global minimization procedure
Calculated          209 grid points in           0 s
Found the set of lowest grid points in           0 s
Calculated POLY solution      0 s, total time   0 s
POLY: s-a-v 1 x(al) 0.0001 .20 0.001,,
POLY: step
Option? /NORMAL/:
Step will start from axis value  0.100000
...OK

Phase Region from  0.100000    for:
  FCC_A1
Global test at  1.08000E-01 .... OK
Global test at  1.18000E-01 .... OK
Global test at  1.28000E-01 .... OK
Global test at  1.38000E-01 .... OK
Global test at  1.48000E-01 .... OK
Global test at  1.58000E-01 .... OK
Global test at  1.68000E-01 .... OK
Global test at  1.78000E-01 .... OK
Global test at  1.88000E-01 .... OK
Global test at  1.98000E-01 .... OK
Terminating at  0.200000
Calculated  103 equilibria

Phase Region from  0.100000    for:
  FCC_A1
Global test at  9.20000E-02 .... OK
Global test at  8.20000E-02 .... OK
Global test at  7.20000E-02 .... OK
Global test at  6.20000E-02 .... OK
Global test at  5.20000E-02 .... OK
Global test at  4.20000E-02 .... OK
Global test at  3.20000E-02 .... OK
Global test at  2.20000E-02 .... OK
Global test at  1.20000E-02 .... OK
Global test at  2.00000E-03 .... OK
Terminating at  0.100000E-03
Calculated  103 equilibria
```

```

Terminating at 0.100000E-03
Calculated 103 equilibria
*** Buffer saved on file: C:\Users\vagrant\AppData\Local\Temp\RESULT_002.POLY3
POLY: s-c t=1473,x(al)=.1
POLY: c-e,,,,,
Using global minimization procedure
Calculated 209 grid points in 0 s
Found the set of lowest grid points in 0 s
Calculated POLY solution 0 s, total time 0 s
POLY: s-a-v 1 x(al) 0.0001 .20 0.001,,,
POLY: step
Option? /NORMAL/:
Step will start from axis value 0.100000
...OK

Phase Region from 0.100000 for:
FCC_A1
Global test at 1.08000E-01 .... OK
Global test at 1.18000E-01 .... OK
Global test at 1.28000E-01 .... OK
Global test at 1.38000E-01 .... OK
Global test at 1.48000E-01 .... OK
Global test at 1.58000E-01 .... OK
Global test at 1.68000E-01 .... OK
Global test at 1.78000E-01 .... OK
Global test at 1.88000E-01 .... OK
Global test at 1.98000E-01 .... OK
Terminating at 0.200000
Calculated 103 equilibria

Phase Region from 0.100000 for:
FCC_A1
Global test at 9.20000E-02 .... OK
Global test at 8.20000E-02 .... OK
Global test at 7.20000E-02 .... OK
Global test at 6.20000E-02 .... OK
Global test at 5.20000E-02 .... OK
Global test at 4.20000E-02 .... OK
Global test at 3.20000E-02 .... OK
Global test at 2.20000E-02 .... OK
Global test at 1.20000E-02 .... OK
Global test at 2.00000E-03 .... OK
Terminating at 0.100000E-03
Calculated 103 equilibria
*** Buffer saved on file: C:\Users\vagrant\AppData\Local\Temp\RESULT_002.POLY3
POLY: s-c t=1423,x(al)=.1
POLY: c-e,,,,,
Using global minimization procedure
Calculated 209 grid points in 0 s
Found the set of lowest grid points in 0 s
Calculated POLY solution 0 s, total time 0 s
POLY: s-a-v 1 x(al) 0.0001 .20 0.001,,,
POLY: step
Option? /NORMAL/:
Step will start from axis value 0.100000
...OK

Phase Region from 0.100000 for:
FCC_A1
Global test at 1.08000E-01 .... OK
Global test at 1.18000E-01 .... OK
Global test at 1.28000E-01 .... OK
Global test at 1.38000E-01 .... OK
Global test at 1.48000E-01 .... OK
Global test at 1.58000E-01 .... OK
Global test at 1.68000E-01 .... OK
Global test at 1.78000E-01 .... OK
Global test at 1.88000E-01 .... OK
Global test at 1.98000E-01 .... OK
Terminating at 0.200000
Calculated 103 equilibria

Phase Region from 0.100000 for:
FCC_A1
Global test at 9.20000E-02 .... OK
Global test at 8.20000E-02 .... OK
Global test at 7.20000E-02 .... OK
Global test at 6.20000E-02 .... OK
Global test at 5.20000E-02 .... OK
Global test at 4.20000E-02 .... OK
Global test at 3.20000E-02 .... OK
Global test at 2.20000E-02 .... OK
Global test at 1.20000E-02 .... OK
Global test at 2.00000E-03 .... OK
Terminating at 0.100000E-03
Calculated 103 equilibria
*** Buffer saved on file: C:\Users\vagrant\AppData\Local\Temp\RESULT_002.POLY3
POLY: s-c t=1373,x(al)=.1
POLY: c-e,,,,,
Using global minimization procedure
Calculated 209 grid points in 0 s
Found the set of lowest grid points in 0 s
Calculated POLY solution 0 s, total time 0 s
POLY: s-a-v 1 x(al) 0.0001 .20 0.001,,,
POLY: step
Option? /NORMAL/:
Step will start from axis value 0.100000
...OK

Phase Region from 0.100000 for:
FCC_A1
Global test at 1.08000E-01 .... OK
Global test at 1.18000E-01 .... OK
Global test at 1.28000E-01 .... OK
Global test at 1.38000E-01 .... OK
Global test at 1.48000E-01 .... OK
Global test at 1.58000E-01 .... OK
Global test at 1.68000E-01 .... OK
Global test at 1.78000E-01 .... OK
Global test at 1.88000E-01 .... OK
Global test at 1.98000E-01 .... OK
Terminating at 0.200000
Calculated 103 equilibria

Phase Region from 0.100000 for:

```

```

FCC_A1
Global test at 9.20000E-02 .... OK
Global test at 8.20000E-02 .... OK
Global test at 7.20000E-02 .... OK
Global test at 6.20000E-02 .... OK
Global test at 5.20000E-02 .... OK
Global test at 4.20000E-02 .... OK
Global test at 3.20000E-02 .... OK
Global test at 2.20000E-02 .... OK
Global test at 1.20000E-02 .... OK
Global test at 2.00000E-03 .... OK
Terminating at 0.10000E-03
Calculated 103 equilibria
*** Buffer saved on file: C:\Users\vagrant\AppData\Local\Temp\RESULT_002.POLY3
POLY: s-c t=1323,x(al)=.1
POLY: c-e,,,
Using global minimization procedure
Calculated 209 grid points in 0 s
Found the set of lowest grid points in 0 s
Calculated POLY solution 0 s, total time 0 s
POLY: s-a-v 1 x(al) 0.0001 .20 0.001,,
POLY: step
Option? /NORMAL/:
Step will start from axis value 0.100000
...OK

Phase Region from 0.100000 for:
FCC_A1
Global test at 1.08000E-01 .... OK
Global test at 1.18000E-01 .... OK
Global test at 1.28000E-01 .... OK
Global test at 1.38000E-01 .... OK
Global test at 1.48000E-01 .... OK
Global test at 1.58000E-01 .... OK
Global test at 1.68000E-01 .... OK
Global test at 1.78000E-01 .... OK
Global test at 1.88000E-01 .... OK
Global test at 1.98000E-01 .... OK
Terminating at 0.200000
Calculated 103 equilibria

Phase Region from 0.100000 for:
FCC_A1
Global test at 9.20000E-02 .... OK
Global test at 8.20000E-02 .... OK
Global test at 7.20000E-02 .... OK
Global test at 6.20000E-02 .... OK
Global test at 5.20000E-02 .... OK
Global test at 4.20000E-02 .... OK
Global test at 3.20000E-02 .... OK
Global test at 2.20000E-02 .... OK
Global test at 1.20000E-02 .... OK
Global test at 2.00000E-03 .... OK
Terminating at 0.10000E-03
Calculated 103 equilibria
*** Buffer saved on file: C:\Users\vagrant\AppData\Local\Temp\RESULT_002.POLY3
POLY: s-c t=1273,x(al)=.1
POLY: c-e,,,
Using global minimization procedure
Calculated 209 grid points in 0 s
Found the set of lowest grid points in 0 s
Calculated POLY solution 0 s, total time 0 s
POLY: s-a-v 1 x(al) 0.0001 .20 0.001,,
POLY: step
Option? /NORMAL/:
Step will start from axis value 0.100000
...OK

Phase Region from 0.100000 for:
FCC_A1
Global test at 1.08000E-01 .... OK
Global test at 1.18000E-01 .... OK
Global test at 1.28000E-01 .... OK
Global test at 1.38000E-01 .... OK
Global test at 1.48000E-01 .... OK
Global test at 1.58000E-01 .... OK
Global test at 1.68000E-01 .... OK
Global test at 1.78000E-01 .... OK
Global test at 1.88000E-01 .... OK
Global test at 1.98000E-01 .... OK
Terminating at 0.200000
Calculated 103 equilibria

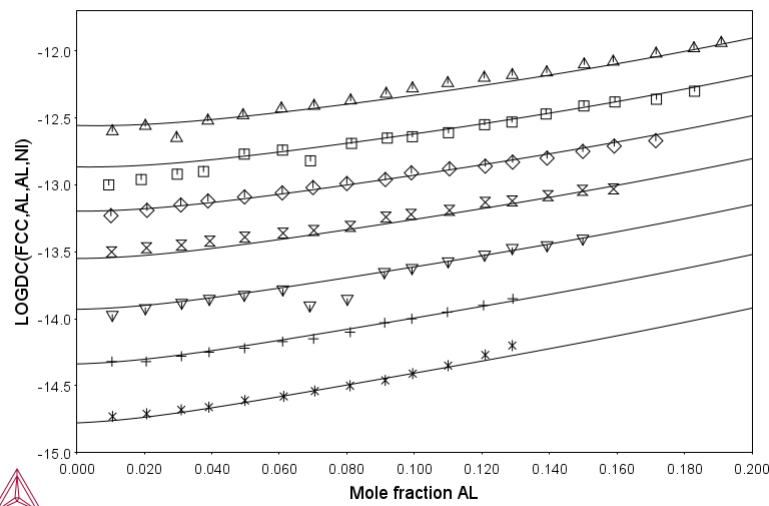
Phase Region from 0.100000 for:
FCC_A1
Global test at 9.20000E-02 .... OK
Global test at 8.20000E-02 .... OK
Global test at 7.20000E-02 .... OK
Global test at 6.20000E-02 .... OK
Global test at 5.20000E-02 .... OK
Global test at 4.20000E-02 .... OK
Global test at 3.20000E-02 .... OK
Global test at 2.20000E-02 .... OK
Global test at 1.20000E-02 .... OK
Global test at 2.00000E-03 .... OK
Terminating at 0.10000E-03
Calculated 103 equilibria
*** Buffer saved on file: C:\Users\vagrant\AppData\Local\Temp\RESULT_002.POLY3
POLY: @@
POLY: @@ ENTER THE POST MODULE, PLOT THE DIFFUSIVITY ON THE Y-AXIS
POLY: @@ AND MOLE-FRACTION Al ON THE X-AXIS.
POLY: @@
POLY: post
  POLY-3 POSTPROCESSOR VERSION 3.2
POST:
POST: s-d-a x m-f al
POST: s-d-a y logdc(fcc,al,al,ni)
POST:
POST: app y yama.exp
PROLOGUE NUMBER: /0/: 1
DATASET NUMBER(s): /-/: 1 2 3 4 5 6 7
POST:
POST:
POST: s-t-m-s y

```

```

COMMAND NOT SUPPORTED IN THIS PLOT DRIVER
POST: s-s-s y -15 -11.7
POST: s-t-m-s y
COMMAND NOT SUPPORTED IN THIS PLOT DRIVER
POST:
POST: SET_EXP_FILE_FORMAT 5
POST: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST: SET_EXP_FILE_FORMAT 10
POST:
POST: plot
2021.05.13.13.06.28
MOB2:AL_NI
T=1273, N=1., P=1.01325E5

```



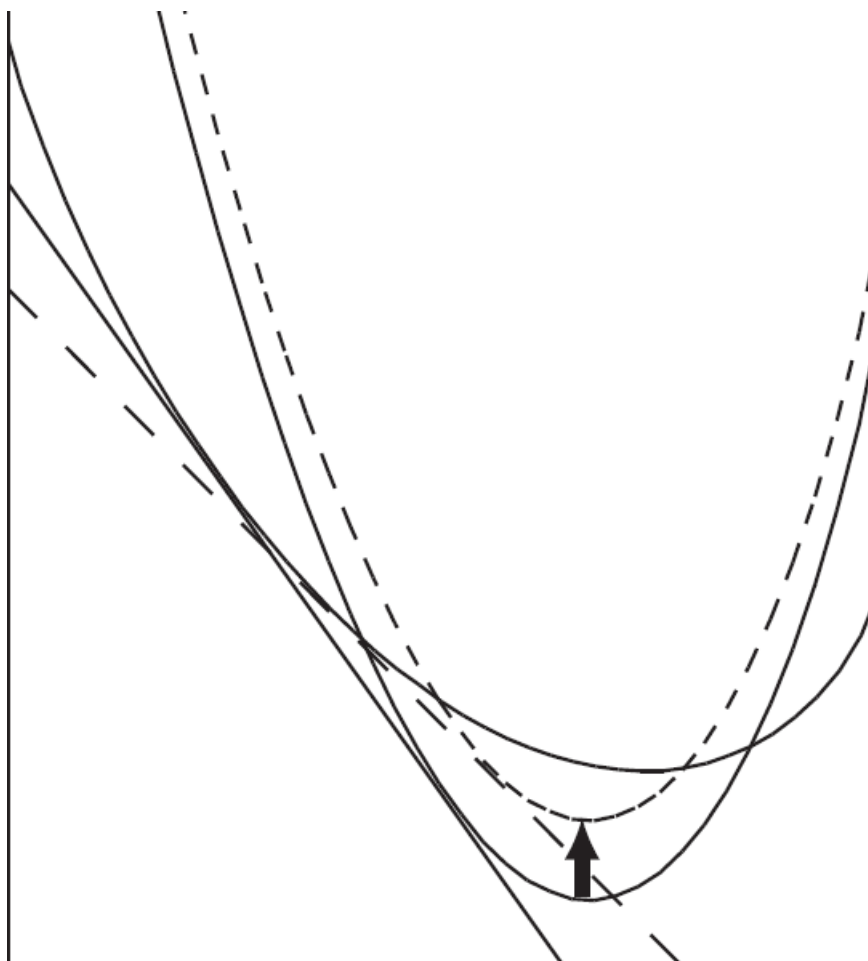
```

POST:
POST:@?<Hit_return_to_continue>
POST:
POST: set-inter
POST:

```



Deviation From Local Equilibrium

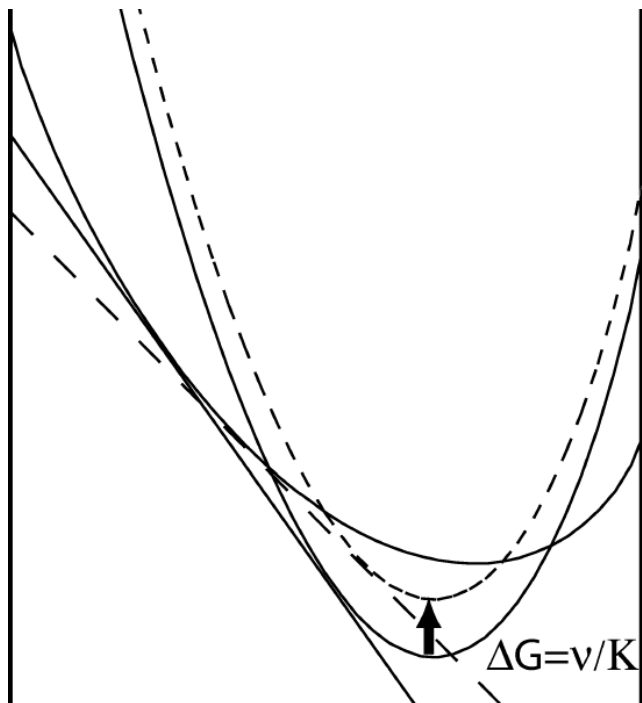




Example exh1

σ/γ diffusion couple with limited interface mobility

This example calculates the growth of ferrite (α) into austenite (γ) with a limited interface mobility. This is achieved by adding a Gibbs-energy contribution to the ferrite using the SET-SURFACE-ENERGY command.



exh1-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exh1\setup.DCM.test"

SYS: @@
SYS: @@ Deviation from local equilibrium.
SYS: @@ Ferrite/austenite diffusion couple with interface mobility
SYS: @@ This example calculates the growth of ferrite into austenite with
SYS: @@ a limited interface mobility. this is done by adding a Gibbs-energy
SYS: @@ contribution to the ferrite using the SET-SURFACE-ENERGY command.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: @@
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE THE DATA
TDB_TCFE11: @@
TDB_TCFE11: sw FEDEMO
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-sys fe c
FE C DEFINED
TDB_FEDEMO: rej ph * all
GAS:G LIQUID:L BCC_A2
LAVES_PHASE_C14 C8CC_A12 CEMENTITE
CUB_A13 DIAMOND_FCC_A4 FCC_A1
GRAPHITE HCP_A3 KSI_CARBIDE
M23C6 M5C2 M7C3
REJECTED
TDB_FEDEMO: res ph bcc fcc
BCC_A2 FCC_A1 RESTORED
TDB_FEDEMO: get
13:07:46,199 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES
ELEMENTS
SPECIES
PHASES
PARAMETERS
FUNCTIONS

List of references for assessed data

'P. Franke, estimated parameter within SGTE, 2007; Fe-C, Ni-C, Mo-C, C-Mn'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
-OK-
TDB_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app mfedemo
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys fe c
FE C DEFINED
APP: rej ph * all
BCC_A2 FCC_A1 REJECTED
APP: res ph bcc fcc
BCC_A2 FCC_A1 RESTORED
APP: get
ELEMENTS
SPECIES
PHASES
PARAMETERS
FUNCTIONS

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Z. Metallkunde 85(1994)498-501; C and N diffusion in bcc Cr
-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'
-OK-
APP: @@
APP: @@ ENTER THE DICTRA MONITOR
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@

```

DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 1000; * N
DIC> @@
DIC> @@
DIC> @@ START BY ENTERING THE REGIONS ferrite AND austenite WHERE
DIC> @@ THE BCC AND FCC PHASES ARE PUT, RESPECTIVELY. THE FERRITE REGION IS
DIC> @@ ASSUMED INITIALLY TO BE VERY THIN, 1E-9 METERS.
DIC> @@
DIC> enter-region
REGION NAME : ferrite
DIC> enter-region
REGION NAME : austenite
ATTACH TO REGION NAMED /FERRITE/:
ATTACHED TO THE RIGHT OF FERRITE /YES/:
DIC>
DIC> @@
DIC> @@ ENTER GRIDS INTO THE REGIONS
DIC> @@
DIC> enter-grid
REGION NAME : /FERRITE/: ferrite
WIDTH OF REGION /1/: 1e-9
TYPE /LINEAR/: AUTO
DIC>
DIC> enter-grid austenite
WIDTH OF REGION /1/: 0.999e-6
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER THE active PHASES INTO THE REGIONS
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /FERRITE/: ferrite
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: bcc
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /AUSTENITE/: austenite
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITION INTO BCC
DIC> @@
DIC> enter-composition
REGION NAME : /FERRITE/: ferrite
PHASE NAME: /BCC_A2/: bcc
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: C
TYPE /LINEAR/: linear
VALUE OF FIRST POINT : 0.019091893
VALUE OF LAST POINT : /1.9091893E-2/: 0.019091893
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITION INTO FCC
DIC> @@
DIC> enter-composition
REGION NAME : /AUSTENITE/: austenite
PHASE NAME: /FCC_A1/: fcc#1
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: C
TYPE /LINEAR/: linear
VALUE OF FIRST POINT : 0.0191
VALUE OF LAST POINT : /1.91E-2/: 0.0191
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exhl y
DIC>
DIC> set-inter
--OK--
DIC>

```

exh1-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exh1\run.DCM.test"
DIC>
DIC>
DIC> @@
DIC> @@ READ THE SETUP FROM FILE AND START THE SIMULATION
DIC> @@
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC>
DIC> read exh1
OK
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-sim-time
END TIME FOR INTEGRATION /1/: 2.5E-3
AUTOMATIC TIMESTEP CONTROL /YES/: YES
MAX TIMESTEP DURING INTEGRATION /2.5E-04/:
INITIAL TIMESTEP : /1E-07/: 1E-7
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/: 1E-7
DIC>
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim
Region: FERRITE
single geometric dense at 0.10000E-08
0.99998 24
Region: AUSTENITE
single geometric dense at 0.0000
1.0666 84
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
Trying old scheme 3
U-FRACTION IN SYSTEM: C = 8.8825328568563E-04 FE = 1
TOTAL SIZE OF SYSTEM: 1E-06 [m]
U-FRACTION IN SYSTEM: C = 8.8825328568563E-04 FE = 1
TOTAL SIZE OF SYSTEM: 1E-06 [m]
2.137709938543949E-004 2.137625593783627E-004 13.8289494082011 2.120998911035789E-004 2.112667987438772E-004
2.108508673179424E-004 2.106430552934355E-004 2.104353457279473E-004 2.101948159848891E-004
2.097141691265310E-004 2.087545259209826E-004 2.068418415541200E-004 2.030428809977812E-004
1.955505925945084E-004 1.80985466255661E-004 1.535545780378763E-004 1.054471342636204E-004
3.627422031933180E-005 13.8289493973143 3.615700659637290E-005 3.609847086651284E-005
3.606922078447431E-005 3.605460018916993E-005 3.603998255771458E-005 3.602477866736845E-005
3.599438050954132E-005 3.593362268572064E-005 3.581226100506191E-005 3.557015351193050E-005
3.508840199823097E-005 3.413475286112764E-005 3.226687014812406E-005 2.868876696680698E-005
2.216320958311595E-005 1.163469073143171E-005 7.48347184238138 1.161266681061649E-005
1.160166267452470E-005 1.159616256257289E-005 1.159341299561144E-005 1.159066375463646E-005
1.158790566549404E-005 1.158239047172325E-005 1.157136402252789E-005 1.154932687740117E-005
1.150531560030047E-005 1.141754509870998E-005 1.124301230593695E-005 1.089797956220244E-005
1.022404544182420E-005 1.490118163906617E-002 1.019021768213511E-005 1.017332482286429E-005
0.005 1.016066437488455E-005 1.016066437488455E-005 1.015644597727906E-005 1.015204956612225E-005
1.014325959902083E-005 1.012569108573464E-005 1.009059974271581E-005 1.002059979098280E-005
9.881330824735401E-006 4.762707176426177E-006 4.438313282537391E-003 2.058831396523542E-006
2.181365075882774E-007 6.215862814898044E-009 1.005966772581293E-011 2.807707694406453E-008
1.740878115116929E-015 2.916483858551415E-008 3.123161389050897E-019 TIME = 0.10000000E-06 DT = 0.10000000E-06
06 SUM OF SQUARES = 0.31231614E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS 4.9961063 AND 4.9961063
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.50061063E-06

output ignored...

... output resumed

TIME = 0.24974221E-02 DT = 0.18903327E-05 SUM OF SQUARES = 0.87230774E-22
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.83033508E-03 AND 0.83033508E-03
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.93676957E-06
U-FRACTION IN SYSTEM: C = 8.98900765800063E-04 FE = 1
TOTAL SIZE OF SYSTEM: 1E-06 [m]
CPU time used in timestep 1 seconds
7.855334004404191E-008 7.845460801976257E-008 2.895741356481490E-010 1.377394816594271E-012 3.299224416274729E-012
017 TIME = 0.24986191E-02 DT = 0.11969968E-05 SUM OF SQUARES = 0.32992244E-16
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.95215868E-03 AND 0.95215868E-03
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.93790930E-06
U-FRACTION IN SYSTEM: C = 8.98953471792101E-04 FE = 1
TOTAL SIZE OF SYSTEM: 1E-06 [m]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FERRITE

CPU time used in timestep 0 seconds
1.224785545856436E-007 1.223255985773290E-007 2.395878434457649E-010 6.518557879551556E-013 4.308840151991679E-013
018 TIME = 0.24993550E-02 DT = 0.73586306E-06 SUM OF SQUARES = 0.43088402E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.10897996E-02 AND 0.10897996E-02
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.93871125E-06
U-FRACTION IN SYSTEM: C = 8.98994461738538E-04 FE = 1
TOTAL SIZE OF SYSTEM: 1E-06 [m]
CPU time used in timestep 0 seconds
3.060408393253354E-007 3.057362302978318E-007 3.253001941359429E-010 5.404634182862258E-013 1.316152297969096E-013
018 TIME = 0.249997404E-02 DT = 0.38538384E-06 SUM OF SQUARES = 0.13161523E-17
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.12900929E-02 AND 0.12900929E-02
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.93920843E-06
U-FRACTION IN SYSTEM: C = 8.99052140693921E-04 FE = 1
TOTAL SIZE OF SYSTEM: 1E-06 [m]
1 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FERRITE

CPU time used in timestep 0 seconds
3.534169304619747E-007 3.530044302696894E-007 1.634713996639121E-010 8.223318251085297E-014 2.121719153613546E-014
020 TIME = 0.24999375E-02 DT = 0.19709686E-06 SUM OF SQUARES = 0.21217192E-19
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.14857381E-02 AND 0.14857381E-02
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.93950126E-06
U-FRACTION IN SYSTEM: C = 8.99110504327324E-04 FE = 1
TOTAL SIZE OF SYSTEM: 1E-06 [m]
CPU time used in timestep 0 seconds
1.382373138175631E-006 1.381370631987372E-006 1.794178772981907E-010 9.492227445077501E-015 3.810159827993530E-015
022 TIME = 0.25000000E-02 DT = 0.62514869E-07 SUM OF SQUARES = 0.38101598E-21
```

CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.18529423E-02 AND 0.18529423E-02
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.93961710E-06
U-FRACTION IN SYSTEM: C = 8.99171784523352E-04 FE = 1
TOTAL SIZE OF SYSTEM: 1E-06 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 0.30000000E-06
DELETING TIME-RECORD FOR TIME 0.70000000E-06
DELETING TIME-RECORD FOR TIME 0.15000000E-05
DELETING TIME-RECORD FOR TIME 0.28565341E-05
DELETING TIME-RECORD FOR TIME 0.52015356E-05
DELETING TIME-RECORD FOR TIME 0.98915386E-05
DELETING TIME-RECORD FOR TIME 0.19271545E-04
DELETING TIME-RECORD FOR TIME 0.33687181E-04
DELETING TIME-RECORD FOR TIME 0.53417477E-04
DELETING TIME-RECORD FOR TIME 0.78654361E-04
DELETING TIME-RECORD FOR TIME 0.10952899E-03
DELETING TIME-RECORD FOR TIME 0.14613094E-03
DELETING TIME-RECORD FOR TIME 0.18853082E-03
DELETING TIME-RECORD FOR TIME 0.23679896E-03
DELETING TIME-RECORD FOR TIME 0.29104871E-03
DELETING TIME-RECORD FOR TIME 0.35145426E-03
DELETING TIME-RECORD FOR TIME 0.41823583E-03
DELETING TIME-RECORD FOR TIME 0.48723339E-03
DELETING TIME-RECORD FOR TIME 0.55628740E-03
DELETING TIME-RECORD FOR TIME 0.62546249E-03
DELETING TIME-RECORD FOR TIME 0.69481220E-03
DELETING TIME-RECORD FOR TIME 0.76438013E-03
DELETING TIME-RECORD FOR TIME 0.83419603E-03
DELETING TIME-RECORD FOR TIME 0.90425851E-03
DELETING TIME-RECORD FOR TIME 0.97473728E-03
DELETING TIME-RECORD FOR TIME 0.10456577E-02
DELETING TIME-RECORD FOR TIME 0.11169997E-02
DELETING TIME-RECORD FOR TIME 0.11887868E-02
DELETING TIME-RECORD FOR TIME 0.12609692E-02
DELETING TIME-RECORD FOR TIME 0.13331752E-02
DELETING TIME-RECORD FOR TIME 0.14056137E-02
DELETING TIME-RECORD FOR TIME 0.14782827E-02
DELETING TIME-RECORD FOR TIME 0.15511176E-02
DELETING TIME-RECORD FOR TIME 0.16239995E-02
DELETING TIME-RECORD FOR TIME 0.16973710E-02
DELETING TIME-RECORD FOR TIME 0.17714237E-02
DELETING TIME-RECORD FOR TIME 0.18459454E-02
DELETING TIME-RECORD FOR TIME 0.19208360E-02
DELETING TIME-RECORD FOR TIME 0.19960899E-02
DELETING TIME-RECORD FOR TIME 0.20715733E-02
DELETING TIME-RECORD FOR TIME 0.21477959E-02
DELETING TIME-RECORD FOR TIME 0.22245451E-02
DELETING TIME-RECORD FOR TIME 0.23020845E-02
DELETING TIME-RECORD FOR TIME 0.23637496E-02
DELETING TIME-RECORD FOR TIME 0.24066139E-02
DELETING TIME-RECORD FOR TIME 0.24364991E-02
DELETING TIME-RECORD FOR TIME 0.24572378E-02
DELETING TIME-RECORD FOR TIME 0.24715984E-02
DELETING TIME-RECORD FOR TIME 0.24814555E-02
DELETING TIME-RECORD FOR TIME 0.24881136E-02
DELETING TIME-RECORD FOR TIME 0.24925865E-02
DELETING TIME-RECORD FOR TIME 0.24955318E-02
DELETING TIME-RECORD FOR TIME 0.24974221E-02
DELETING TIME-RECORD FOR TIME 0.24986191E-02
DELETING TIME-RECORD FOR TIME 0.24993550E-02
DELETING TIME-RECORD FOR TIME 0.24997404E-02

KEEPING TIME-RECORD FOR TIME 0.24999375E-02
AND FOR TIME 0.25000000E-02
WORKSPACE RECLAIMED

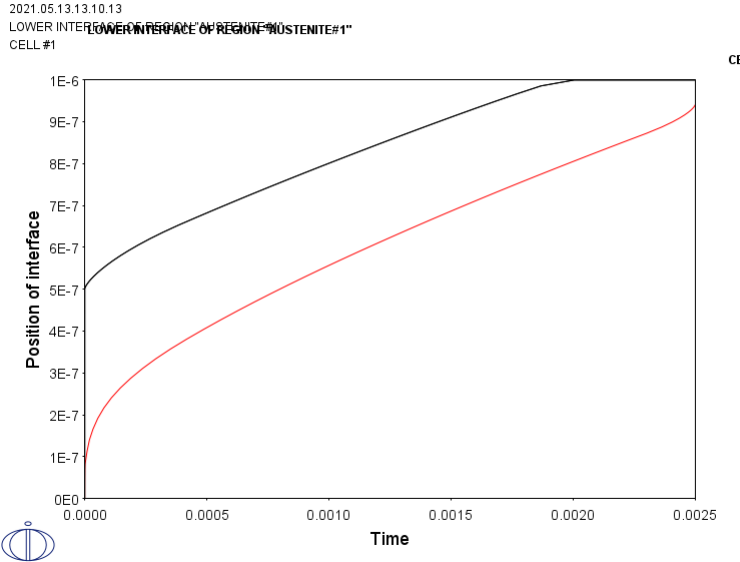
TIMESTEP AT 0.250000000E-02 SELECTED

DIC>
DIC>
DIC> set-inter
--OK--
DIC>

exh1-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exh1\plot.DCM.test"
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 2.50000E-03
DIC>
DIC> read exh1
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ SET THE DATA APPENDED FROM THE "EXP" FILE TO BE READ
POST-1: @@
POST-1: set-col for for red
COMMAND NOT SUPPORTED IN THIS PLOT DRIVER
POST-1:
POST-1: @@
POST-1: @@ COMPARE THE POSITION OF THE INTERFACE AS A FUNCTION OF TIME
POST-1: @@
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-d-a y posi aus low
POST-1:
POST-1: @@
POST-1: @@ APPEND THE SIMULATION (WITHOUT THE ENERGY CONTRIBUTION) FROM FILE
POST-1: @@
POST-1: app y noadd.exp 1; 1
POST-1:
POST-1: @@
POST-1: @@ SET A TITLE ON THE PLOT
POST-1: @@
POST-1: set-title Figure h1
POST-1:
POST-1: @@
POST-1: @@ PLOT THE RESULTS
POST-1: @@
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
ORKING ...
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Figure h1



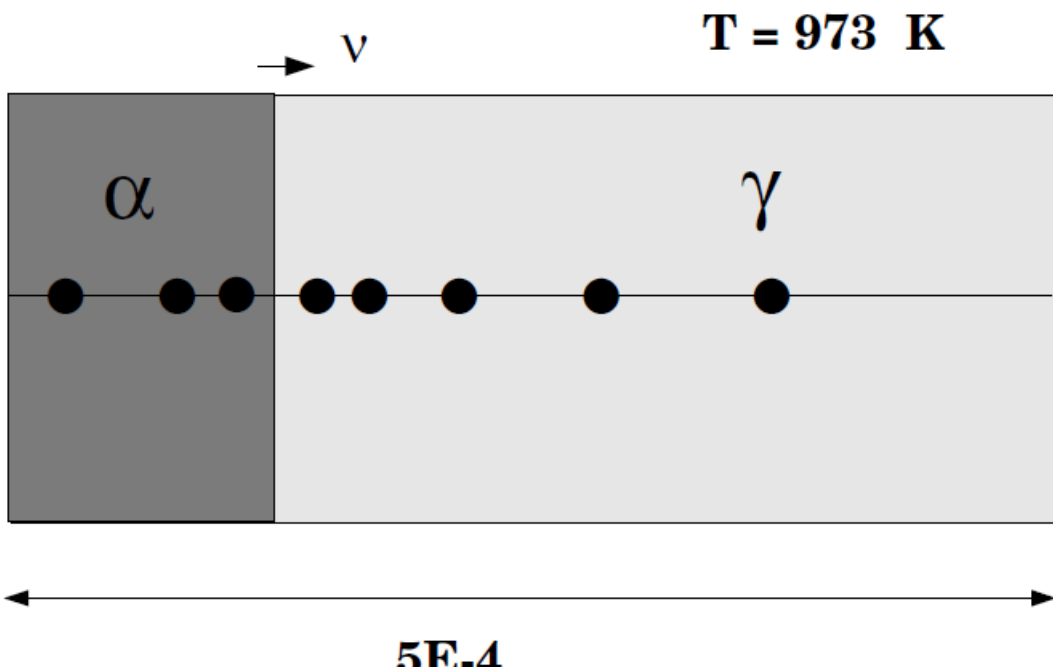
```
POST-1:
POST-1:
POST-1: set-inter
--OK---
POST-1:
```



Example exh2

σ/γ para-equilibrium in an Fe-Ni-C alloy

This example calculates the growth of ferrite (α) into austenite (γ) in an Fe-2.02%Ni-0.0885%C alloy using the para-equilibrium model. The results are compared with experimental information from Hutchinson, C. R., A. Fuchsmann, and Yves Brechet. "The diffusional formation of ferrite from austenite in Fe-C-Ni alloys." Met. Mat. Trans A 35.4 (2004): 1211-1221.



exh2-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exh2\setup.DCM.test"

SYS: @@
SYS: @@ Deviation from local equilibrium.
SYS: @@ Ferrite/austenite para-equilibrium in an Fe-Ni-C alloy
SYS: @@ This example calculates the growth of ferrite into austenite
SYS: @@ in an Fe-2.02%Ni-0.0885%C alloy using the para-equilibrium model.
SYS: @@ The results are compared with experimental information from
SYS: @@ Hutchinson, C. R., A. Fuchsmann, and Yves Brechet. "The diffusional
SYS: @@ formation of ferrite from austenite in Fe-C-Ni alloys." Metall.
SYS: @@ Mat. Trans. A 35.4 (2004): 1211-1221.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:

SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASE
SYS: @@
SYS: go da

THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: @@
TDB_TCFE11: @@ SELECT A DATABASE FOR THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw fedemo
Current database: Iron Demo Database v4.0

VA /- DEFINED
TDB_FEDEMO: def-sys fe ni c
FE NI C
DEFINED
TDB_FEDEMO: rej ph * all
GAS:G LIQUID:L BCC_A2
LAVES_PHASE_C14 CBC_A12 CEMENTITE
CUB_A13 DIAMOND_FCC_A4 FCC_A1
GRAPHITE HCP_A3 KSI CARBIDE
M23C6 MSC2 M7C3
REJECTED
TDB_FEDEMO: res ph bcc fcc
BCC_A2 FCC_A1 RESTORED
TDB_FEDEMO: get
13:11:30,360 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES
ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'P. Franke, estimated parameter within SGTE, 2007; Fe-C, Ni-C, Mo-C, C-Mn'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A. Gabriel, C. Chatillon, and I. Ansara, published in High Temp. Sci.
(Parameters listed in CALPHAD, 11 (1987) 203-218); C-NI'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
'A. Gabriel, P. Gustafson, and I. Ansara, CALPHAD, 11 (1987) 203 -218;
TRITA-MAC 285 (1986); C-FE-NI'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
-OK-

TDB_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app mfdemo
Current database: Fe-Alloys Mobility demo database v2.0

VA DEFINED
APP: def-sys fe ni c
FE NI C
DEFINED
APP: rej ph * all
BCC_A2 FCC_A1 REJECTED
APP: res ph bcc fcc
BCC_A2 FCC_A1 RESTORED
APP: get
ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Z. Metallkunde 85(1994)502-509; C diffusion in fcc Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'

```

'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
'B. Jonsson: Z. Metallkunde 85(1994)498-501; C and N diffusion in bcc Cr
-Fe-Ni'
'B. Jonsson: Z. Metallkunde 83(1992)349-355; Cr, Co, Fe and Ni diffusion
in bcc Fe'
'B. Jonsson: ISIJ International, 35(1995)1415-1421; Cr, Fe and Ni
diffusion bcc Cr-Fe-Ni'
-OK-
APP:
APP: @@
APP: @@ ENTER THE DICTRA MONITOR
APP: @@
APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 973; * N
DIC>
DIC> @@
DIC> @@ START BY ENTERING THE REGIONS ferrite AND austenite WHERE THE
DIC> @@ BCC AND FCC PHASES ARE PUT, RESPECTIVELY. THE FERRITE REGION IS
DIC> @@ ASSUMED INITIALLY TO BE VERY THIN, 1E-9 METERS.
DIC> @@
DIC> enter-region
REGION NAME : ferrite
DIC>
DIC> enter-region
REGION NAME : austenite
ATTACH TO REGION NAMED /FERRITE/:
ATTACHED TO THE RIGHT OF FERRITE /YES/:
DIC>
DIC> @@
DIC> @@ ENTER GRIDS INTO THE REGIONS
DIC> @@
DIC> enter-grid
REGION NAME : /FERRITE/: ferrite
WIDTH OF REGION /1/: 1e-9
TYPE /LINEAR/: AUTO
DIC>
DIC> enter-grid austenite
WIDTH OF REGION /1/: 50e-6
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER active PHASES INTO THE REGIONS
DIC> @@
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /FERRITE/: ferrite
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: bcc
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: active
REGION NAME : /AUSTENITE/: austenite
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: fcc#1
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITION INTO BCC
DIC> @@
DIC> enter-composition
REGION NAME : /FERRITE/: ferrite
PHASE NAME: /BCC_A2/: bcc
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: c
TYPE /LINEAR/: linear
VALUE OF FIRST POINT : 0.0885
VALUE OF LAST POINT : /8.85E-2/: 0.0885
PROFILE FOR /NI/: ni
TYPE /LINEAR/: linear
VALUE OF FIRST POINT : 2.02
VALUE OF LAST POINT : /2.02/: 2.02
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITION INTO FCC
DIC> @@
DIC> enter-composition
REGION NAME : /AUSTENITE/: austenite
PHASE NAME: /FCC_A1/: fcc#1
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: w-p
PROFILE FOR /C/: c
TYPE /LINEAR/: linear
VALUE OF FIRST POINT : 0.0885
VALUE OF LAST POINT : /8.85E-2/: 0.0885
PROFILE FOR /NI/: ni
TYPE /LINEAR/: linear
VALUE OF FIRST POINT : 2.02
VALUE OF LAST POINT : /2.02/: 2.02
DIC>
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exh2 y
DIC>
DIC> set-inter
--OK--
DIC>
```

exh2-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exh2\run.DCM.test"
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ FILE TO RUN EXAMPLE exh2
DIC> @@
DIC>
DIC> @@
DIC> @@ READ THE SETUP FROM FILE
DIC> @@
DIC>
DIC>
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC>
DIC> read exh2
OK
DIC>
DIC> @@
DIC> @@ SET THE SIMULATION TIME
DIC> @@
DIC> set-sim-time 50,,,
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ ENABLE THE PARA-EQUILIBRIUM MODEL
DIC> @@
DIC> para
ENABLE PARAEQ : /NO/: YES
U-FRACTION OF COMPONENT FE /AUTO/: AUTO
U-FRACTION OF COMPONENT NI /AUTO/: AUTO
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim
Region: FERRITE
single geometric dense at 0.10000E-08
0.99997 24
Region: AUSTENITE
single geometric dense at 0.0000
1.1602 98
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
Trying old scheme 4
GENERATING STARTING VALUES FOR CELL # 1 INTERFACE # 2
DETERMINING INITIAL EQUILIBRIUM VALUES
CALCULATING STARTING VALUES: 9 EQUILIBRIUM CALCULATIONS DONE 6 OUT OF 9
DETERMINED ACTIVITIES ACR(C) .161541295585
UNABLE TO OBTAIN GOOD STARTING VALUE USING THE OLD SCHEME
USE NEW SCHEME /YES/:
Trying new scheme
GENERATING STARTING VALUES FOR CELL # 1 INTERFACE # 2
DETERMINING INITIAL EQUILIBRIUM VALUES
CALCULATING STARTING VALUES: 18 EQUILIBRIUM CALCULATIONS DONE 1 OUT OF 18
U-FRACTION IN SYSTEM: C = .00412262676333 FE = .980742621143594
NI = .0192573788564064
TOTAL SIZE OF SYSTEM: 5.0001E-05 [m]
U-FRACTION IN SYSTEM: C = .00412262676333 FE = .980742621143594
NI = .0192573788564064
TOTAL SIZE OF SYSTEM: 5.0001E-05 [m]
2232.10931280802 2232.10931281135 2228.18551830736 2.32383611703743 1.947804402559963E-
003 8.416186789939047E-003 7.623216312568318E-006 1.892250135006316E-003 5.168572771943315E-
006 2.142557866573553E-006 3.440441220515192E-013 TIME = 0.10000000E-06 DT = 0.10000000E-
06 SUM OF SQUARES = 0.34404412E-12
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.20272106E-01 AND 0.20272106E-01
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.30272106E-08
U-FRACTION IN SYSTEM: C = .00412273631313628 FE = .980742621143594
NI = .0192573788564064
TOTAL SIZE OF SYSTEM: 5.0001E-05 [m]
11 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: AUSTENITE
CPU time used in timestep 1 seconds
2189.02035753682 2189.02035753763 2185.12779966492 2.22490243108414 1.816589661367364E-
003 7.875020729155016E-003 7.030647432450283E-007 1.892093511098311E-003
output ignored...
... output resumed
1.035830433581525E-006 4.070247910639300E-010 2.083959365948200E-003 3.057789734168416E-
019 TIME = 39.294723 DT = 4.0585241 SUM OF SQUARES = 0.30577897E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.19678971E-06 AND 0.19678971E-06
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.12958016E-04
U-FRACTION IN SYSTEM: C = .00412955175752938 FE = .980742621143593
NI = .0192573788564066
TOTAL SIZE OF SYSTEM: 5.0001E-05 [m]
18 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FERRITE
CPU time used in timestep 1 seconds
1039.07500724594 1039.07500724600 1036.24504247252 0.435758664141731 1.582365474451798E-
004 9.540009899475173E-007 3.683098567472132E-010 2.086873937912341E-003 2.668683642838332E-
019 TIME = 43.471055 DT = 4.1763314 SUM OF SQUARES = 0.26686836E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.18739190E-06 AND 0.18739190E-06
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.13740626E-04
U-FRACTION IN SYSTEM: C = .00413007211190805 FE = .980742621143594
NI = .0192573788564066
TOTAL SIZE OF SYSTEM: 5.0001E-05 [m]
10 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FERRITE
CPU time used in timestep 1 seconds
1018.79898889626 1018.79898889766 1015.99361196596 0.417671928605375 1.485016196423534E-
004 8.663587825737338E-007 3.298190609469457E-010 2.089742782700098E-003 2.668683406999758E-
019 TIME = 47.752852 DT = 4.2817972 SUM OF SQUARES = 0.26686834E-18
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.17907221E-06 AND 0.17907221E-06
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.14507377E-04
U-FRACTION IN SYSTEM: C = .00413054999711265 FE = .980742621143594
```

NI = .0192573788564067
 TOTAL SIZE OF SYSTEM: 5.000E-05 [m]
 13 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FERRITE

CPU time used in timestep	1 seconds			
998.914356505714	998.914356506641	996.133394484083	0.400344844967206	1.393711337747759E-
004 2.022471141752238E-004	8.988595106467371E-010	2.089218460490361E-003	1.682155864785915E-	
013 TIME = 49.893750 DT = 2.1408986	SUM OF SQUARES = 0.16821559E-12			
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.20276681E-06 AND 0.20276681E-06				
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.14941480E-04				
U-FRACTION IN SYSTEM: C = .00413164825082925 FE = .980742621143593				
NI = .0192573788564067				

TOTAL SIZE OF SYSTEM: 5.000E-05 [m]
 5 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FERRITE

CPU time used in timestep	1 seconds			
979.413690130442	979.413690129987	976.656970399060	0.383745048982190	1.308066951988766E-
004 5.518410352312049E-004	1.410377245535091E-008	2.084198669096521E-003	1.796303971242632E-	
009 1.193316244909533E-009	3.537387538513500E-010	8.509763447052110E-		
015 TIME = 50.000000 DT = 0.10624963	SUM OF SQUARES = 0.85097634E-14			
CELL # 1 VELOCITY AT INTERFACE # 2 IS 0.39315202E-06 AND 0.39315202E-06				
POSITION OF INTERFACE FERRITE / AUSTENITE IS 0.14983253E-04				
U-FRACTION IN SYSTEM: C = .00413198323104672 FE = .980742621143593				
NI = .0192573788564067				

TOTAL SIZE OF SYSTEM: 5.000E-05 [m]
 MUST SAVE WORKSPACE ON FILE
 WORKSPACE SAVED ON FILE
 RECLAIMING WORKSPACE

DELETING TIME-RECORD FOR TIME	0.0000000		
DELETING TIME-RECORD FOR TIME	0.10000000E-06		
DELETING TIME-RECORD FOR TIME	0.20000000E-06		
DELETING TIME-RECORD FOR TIME	0.40000000E-06		
DELETING TIME-RECORD FOR TIME	0.80000000E-06		
DELETING TIME-RECORD FOR TIME	0.16000000E-05		
DELETING TIME-RECORD FOR TIME	0.32000000E-05		
DELETING TIME-RECORD FOR TIME	0.64000000E-05		
DELETING TIME-RECORD FOR TIME	0.12800000E-04		
DELETING TIME-RECORD FOR TIME	0.25600000E-04		
DELETING TIME-RECORD FOR TIME	0.48352417E-04		
DELETING TIME-RECORD FOR TIME	0.79491429E-04		
DELETING TIME-RECORD FOR TIME	0.12664747E-03		
DELETING TIME-RECORD FOR TIME	0.21423327E-03		
DELETING TIME-RECORD FOR TIME	0.38940485E-03		
DELETING TIME-RECORD FOR TIME	0.73974803E-03		
DELETING TIME-RECORD FOR TIME	0.14404344E-02		
DELETING TIME-RECORD FOR TIME	0.27785297E-02		
DELETING TIME-RECORD FOR TIME	0.54207754E-02		
DELETING TIME-RECORD FOR TIME	0.10705267E-01		
DELETING TIME-RECORD FOR TIME	0.21274250E-01		
DELETING TIME-RECORD FOR TIME	0.42412216E-01		
DELETING TIME-RECORD FOR TIME	0.84688147E-01		
DELETING TIME-RECORD FOR TIME	0.16924001		
DELETING TIME-RECORD FOR TIME	0.33834374		
DELETING TIME-RECORD FOR TIME	0.67655119		
DELETING TIME-RECORD FOR TIME	1.3529661		
DELETING TIME-RECORD FOR TIME	2.7057959		
DELETING TIME-RECORD FOR TIME	5.2358266		
DELETING TIME-RECORD FOR TIME	7.8217627		
DELETING TIME-RECORD FOR TIME	10.658658		
DELETING TIME-RECORD FOR TIME	13.690783		
DELETING TIME-RECORD FOR TIME	16.905178		
DELETING TIME-RECORD FOR TIME	20.286292		
DELETING TIME-RECORD FOR TIME	23.819608		
DELETING TIME-RECORD FOR TIME	27.493178		
DELETING TIME-RECORD FOR TIME	31.300719		
DELETING TIME-RECORD FOR TIME	35.236199		
DELETING TIME-RECORD FOR TIME	39.294723		
DELETING TIME-RECORD FOR TIME	43.471055		
DELETING TIME-RECORD FOR TIME	47.752852		

KEEPING TIME-RECORD FOR TIME 49.893750
 AND FOR TIME 50.000000
 WORKSPACE RECLAIMED

TIMESTEP AT	50.000000	SELECTED	
-------------	-----------	----------	--

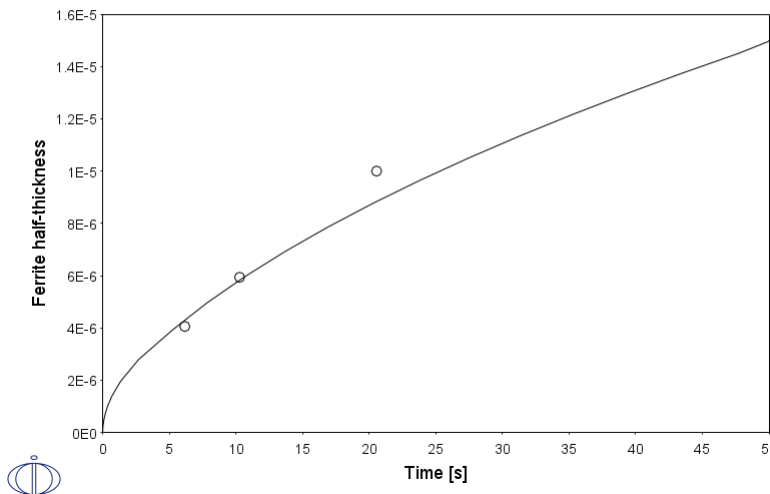
DIC>
DIC> set-inter
--OK--
DIC>

exh2-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exh2\plot.DCM.test"
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 5.00000E+01
DIC>
DIC> read exh2
OK
DIC>
DIC> @@
DIC> @@ GO TO THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ WE WANT TO PLOT THE POSITION OF THE INTERFACE AS A FUNCTION OF TIME
POST-1: @@ I.E. THE FERRITE HALF-THICKNESS
POST-1: @@
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: s-d-a y posi aus low
POST-1: @@
POST-1: @@ APPEND THE EXPERIMENTAL INFORMATION
POST-1: @@
POST-1: app y exh2.exp 1; 1
POST-1: @@
POST-1: @@ SET A TITLE ON THE PLOT
POST-1: @@
POST-1: set-title Figure h2
POST-1: @@
POST-1: @@ RENAME THE AXIS LABELS
POST-1: @@
POST-1: set-axis-text-status
AXIS (X, Y OR Z) : x
AUTOMATIC AXIS TEXT (Y OR N) /N/: NO
AXIS TEXT : Time [s]
POST-1: @@
POST-1: set-axis-text-status
AXIS (X, Y OR Z) : y
AUTOMATIC AXIS TEXT (Y OR N) /N/: NO
AXIS TEXT : Ferrite half-thickness
POST-1: @@
POST-1: @@
POST-1: @@ PLOT THE RESULTS
POST-1: @@
POST-1: @@ PLOT THE RESULTS
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

Figure h2

2021.05.13.13.14.13
LOWER INTERFACE OF REGION "AUSTENITE#1"
CELL #1



```
POST-1:
POST-1:
POST-1:
POST-1: set-inter
--OK--
POST-1:
```

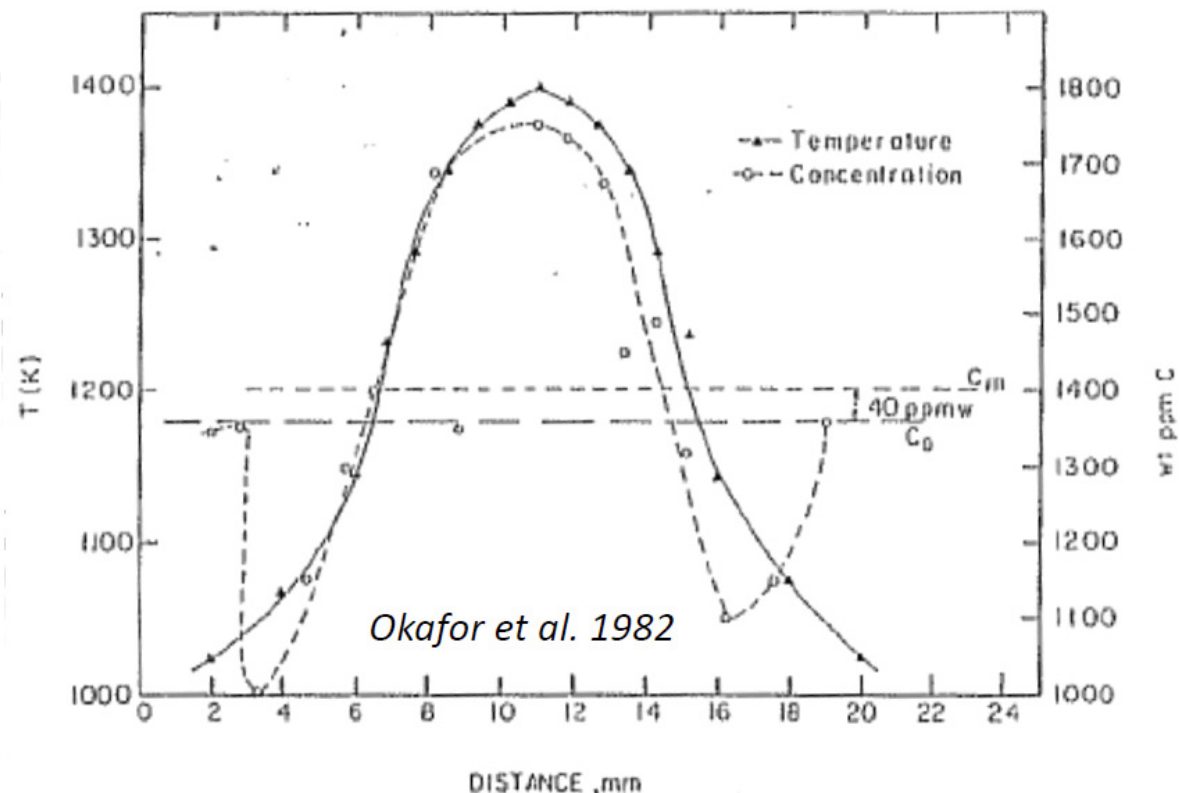


Example exh3

Diffusion induced by a temperature gradient (thermomigration)

This calculation shows how a temperature gradient induces diffusion.

$$J_C = -\frac{u_C}{V_s} y_{Va} M_{CVa} \left(\frac{\partial \mu_C}{\partial x} + \frac{Q_C^*}{T} \frac{\partial T}{\partial x} \right)$$



exh3-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exh3\setup.DCM.test"
SYS: @@
SYS: @@ Deviation from local equilibrium.
SYS: @@ Diffusion induced by a temperature gradient (thermomigration)
SYS: @@ This calculation shows how a temperature gradient induces
SYS: @@ diffusion.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: sw fedemo
Current database: Iron Demo Database v4.0

VA           /- DEFINED
TDB_FEDEMO: def-sys fe ni c
FE           NI           C
DEFINED
TDB_FEDEMO: rej ph * all
GAS:G          LIQUID:L          BCC_A2
LAVES_PHASE_C14 CBCC_A12         CEMENTITE
CUB_A13        DIAMOND_FCC_A4   FCC_A1
GRAPHITE       HCP_A3          KSI_CARBIDE
M23C6          M5C2            M7C3
REJECTED
TDB_FEDEMO: res ph fcc graph
FCC_A1          GRAPHITE RESTORED
TDB_FEDEMO: get
13:15:29,983 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'B.J. Lee, CALPHAD, 16 (1992) 121-149; C-Cr-Fe-Ni'
'A. Gabriel, C. Chatillon, and I. Ansara, published in High Temp. Sci.
(Parameters listed in CALPHAD, 11 (1987) 203-218); C-NI'
'A. Markstrom, Swerea KIMAB, Sweden; Molar volumes'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
'A. Gabriel, P. Gustafson, and I. Ansara, CALPHAD, 11 (1987) 203 -218;
TRITA-MAC 285 (1986); C-FE-NI'
'A.T. Dinsdale, T. Chart, MTDS NPL, Unpublished work (1986); FE-NI'
'B. Uhrenius, Int. J. Refract. Met. Hard Mater. 12 (1994) 121 -127; Molar
volumes'
-OK-
TDB_FEDEMO: app mfdemo
Current database: Fe-Alloys Mobility demo database v2.0

VA  DEFINED
APP: def-sys fe ni c
FE           NI           C
DEFINED
APP: rej ph * all
BCC_A2          FCC_A1  REJECTED
APP: res ph fcc
FCC_A1          RESTORED
APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'This parameter has not been assessed'
'J. Agren: Scripta Met. 20(1986)1507-1510; C diff in fcc C-Fe'
'B. Jonsson: Z. Metallkunde 85(1994)502-509; C diffusion in fcc Cr-Fe-Ni'
'B. Jonsson: Scand. J. Metall. 23(1994)201-208; Fe and Ni diffusion fcc Fe
-Ni'
'B. Jonsson: Scand. J. Metall. 24(1995)21-27; Ni self-diffusion'
-OK-
APP: go d-m
NO TIME STEP DEFINED
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC>
DIC> @@ ENTER A GAUSSIAN-SHAPED TEMPERATURE GRADIENT
DIC> set-cond glob T 0 1000+400*exp(-3.35074E4*(x-11e-3)**2); * N
DIC>
DIC> set-ref C grap,,,
DIC>
DIC> ent-reg aus,,,
DIC>
```

```
DIC> ent-grid aus 25e-3 auto
DIC>
DIC> ent-phd act aus matrix fcc#1
DIC>
DIC> ent-comp aus fcc#1 fe w-p
PROFILE FOR /C/: c lin 0.14 0.14
PROFILE FOR /NI/: ni lin 32.5 32.5
DIC>
DIC> s-s-time 5E7,.....
DIC>
DIC> @@ ENTER THE HEAT OF TRANSFER PARAMETER FOR CARBON
DIC> ent-heat-tra-p
HEAT TRANSFER PARAMETER FOR PHASE: fcc
ELEMENT: C
PARAMETER /0/: -42000
DIC>
DIC>
DIC> save exh3 y
DIC>
DIC> set-inter
--OK--
DIC>
```

exh3-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exh3\run.DCM.test"
DIC>
DIC> go d-m
TIME STEP AT TIME  0.00000E+00
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC>
DIC> read exh3
OK
DIC>
DIC> sim
Region: AUS
linear grid  75
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
U-FRACTION IN SYSTEM: C = .00662305741857946  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
U-FRACTION IN SYSTEM: C = .00662305741857946  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
TIME = 0.10000000E-06 DT = 0.10000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741857945  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 0.10010000E-03 DT = 0.10000000E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741857945  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 0.40010010 DT = 0.40000000  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741857945  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 1600.4001 DT = 1600.0000  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741857938  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 75402.175 DT = 73801.774  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741850602  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 223005.72 DT = 147603.55  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741820696  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 518212.82 DT = 295207.10  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741820656  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 1108627.0 DT = 590414.20  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741820616  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 2289455.4 DT = 1180828.4  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741818172  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 4651112.2 DT = 2361656.8  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0066230574181914  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 7580400.9 DT = 2929288.7  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741820453  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 10062302. DT = 2481900.8  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741821166  FE = .68534915460493
NI = .31465084539509
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 12331243. DT = 2268941.7  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741821537  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 14490802. DT = 2159558.6  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741821863  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 16584438. DT = 2093636.5  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .0066230574182206  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 18634851. DT = 2050412.1  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741791114  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 20655319. DT = 2020468.0  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741735322  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 22654176. DT = 1998857.1  SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: C = .00662305741687841  FE = .68534915460493
NI = .31465084539507
TOTAL SIZE OF SYSTEM: .025 [m]
CPU time used in timestep          0 seconds
TIME = 24636917. DT = 1982741.3  SUM OF SQUARES = 0.0000000
```

U-FRACTION IN SYSTEM: C = .00662305741647036 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 26607291. DT = 1970374.6 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741611502 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 28567937. DT = 1960645.2 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741580548 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 1 seconds
 TIME = 30520755. DT = 1952818.3 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741553341 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 32467152. DT = 1946397.4 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741529459 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 34408194. DT = 1941042.1 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741508377 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 1 seconds
 TIME = 36344708. DT = 1936513.6 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741489939 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 38277348. DT = 1932640.4 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741473561 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 40206647. DT = 1929298.2 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741459067 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 42133040. DT = 1926393.8 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741446232 FE = .68534915460493
 NI = .314650845395069
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 1 seconds
 TIME = 44056896. DT = 1923856.0 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741434744 FE = .68534915460493
 NI = .314650845395069
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 45978525. DT = 1921628.7 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741424537 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 47618324. DT = 1639799.2 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741417177 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 48913853. DT = 1295528.3 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741412362 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 1 seconds
 TIME = 49750529. DT = 836676.92 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741410038 FE = .68534915460493
 NI = .314650845395069
 TOTAL SIZE OF SYSTEM: .025 [m]
 CPU time used in timestep 0 seconds
 TIME = 50000000. DT = 249470.53 SUM OF SQUARES = 0.0000000
 U-FRACTION IN SYSTEM: C = .00662305741409993 FE = .68534915460493
 NI = .31465084539507
 TOTAL SIZE OF SYSTEM: .025 [m]
 MUST SAVE WORKSPACE ON FILE
 WORKSPACE SAVED ON FILE
 RECLAIMING WORKSPACE
 DELETING TIME-RECORD FOR TIME 0.0000000
 DELETING TIME-RECORD FOR TIME 0.10000000E-06
 DELETING TIME-RECORD FOR TIME 0.10010000E-03
 DELETING TIME-RECORD FOR TIME 0.40010010
 DELETING TIME-RECORD FOR TIME 1600.4001
 DELETING TIME-RECORD FOR TIME 75402.175
 DELETING TIME-RECORD FOR TIME 223005.72
 DELETING TIME-RECORD FOR TIME 518212.82
 DELETING TIME-RECORD FOR TIME 1108627.0
 DELETING TIME-RECORD FOR TIME 2289455.4
 DELETING TIME-RECORD FOR TIME 4651112.2
 DELETING TIME-RECORD FOR TIME 7580400.9
 DELETING TIME-RECORD FOR TIME 10062302.
 DELETING TIME-RECORD FOR TIME 12331243.
 DELETING TIME-RECORD FOR TIME 14490802.
 DELETING TIME-RECORD FOR TIME 16584438.
 DELETING TIME-RECORD FOR TIME 18634851.
 DELETING TIME-RECORD FOR TIME 20655319.
 DELETING TIME-RECORD FOR TIME 22654176.
 DELETING TIME-RECORD FOR TIME 24636917.
 DELETING TIME-RECORD FOR TIME 26607291.
 DELETING TIME-RECORD FOR TIME 28567937.
 DELETING TIME-RECORD FOR TIME 30520755.
 DELETING TIME-RECORD FOR TIME 32467152.
 DELETING TIME-RECORD FOR TIME 34408194.
 DELETING TIME-RECORD FOR TIME 36344708.
 DELETING TIME-RECORD FOR TIME 38277348.
 DELETING TIME-RECORD FOR TIME 40206647.
 DELETING TIME-RECORD FOR TIME 42133040.
 DELETING TIME-RECORD FOR TIME 44056896.
 DELETING TIME-RECORD FOR TIME 45978525.
 DELETING TIME-RECORD FOR TIME 47618324.
 DELETING TIME-RECORD FOR TIME 48913853.

KEEPING TIME-RECORD FOR TIME 49750529.

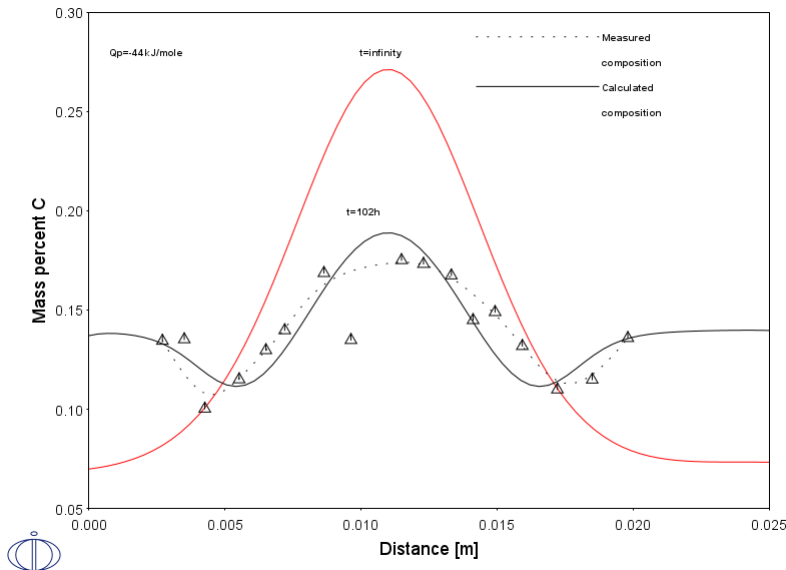
AND FOR TIME 50000000.
WORKSPACE RECLAIMED

TIMESTEP AT 50000000.0 SELECTED

DIC>
DIC> set-inter
--OK---
DIC>

exh3-plot

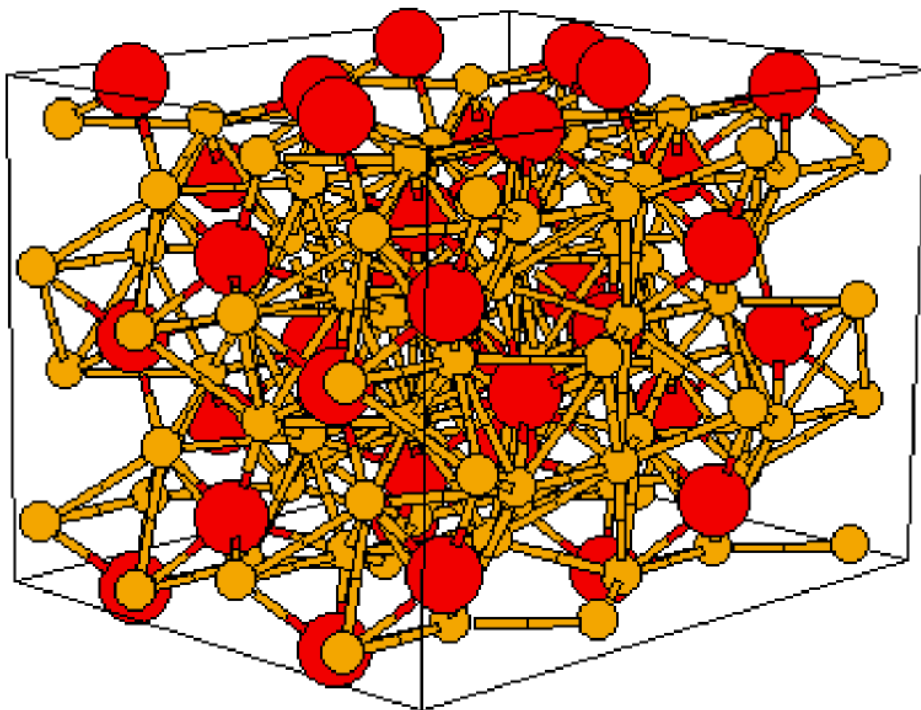
```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exh3\plot.DCM.test"
DIC> go d-m
TIME STEP AT TIME 5.00000E+07
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC>
DIC>
DIC> read exh3
OK
DIC>
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1: s-d-a x dist glob
INFO: Distance is set as independent variable
POST-1: s-d-a y w-p c
POST-1: s-p-c time 367200 SE7
POST-1: s-s-s y n 0.1 0.18
POST-1: s-s-s Y Y
POST-1: app y exh3.exp 0; 1 3;
POST-1: s-p-o n y y n y n n,,,...,
POST-1:
POST-1: set-ax-text-st x
AUTOMATIC AXIS TEXT (Y OR N) /N/: n
AXIS TEXT : Distance [m]
POST-1:
POST-1: set-ax-text-st y
AUTOMATIC AXIS TEXT (Y OR N) /N/: n
AXIS TEXT : Mass percent C
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```



```
POST-1:
POST-1:
POST-1:
POST-1:
POST-1: set-inter
--OK--
POST-1:
```



Diffusion in Complex Phases

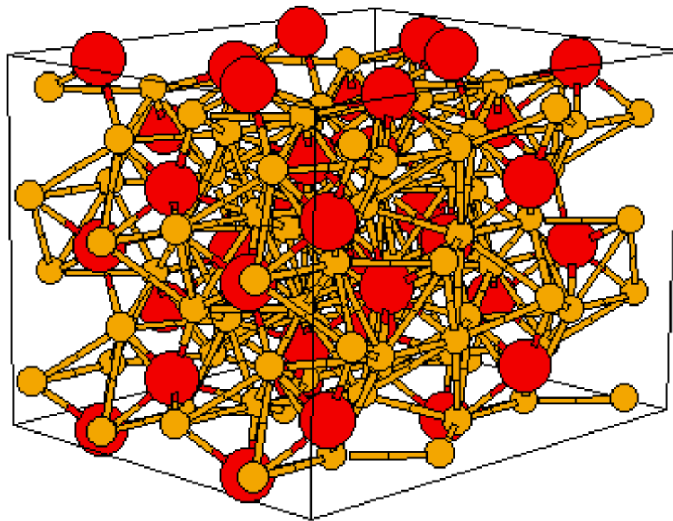




Example exi1

Diffusion in system with B2 ordering

Diffusion in including effects from chemical ordering. In this example folder, there is also a datafile AlFeNi-data.TDB, which contains both a thermodynamic and kinetic description for the ordered and disordered bcc.



exil-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exil\setup.DCM.test"
SYS: @@
SYS: @@ Diffusion in complex phases.
SYS: @@ Diffusion in a system with B2 ordering
SYS: @@ This example shows diffusion in a system with B2 ordering.
SYS: @@ The datafile AlFeNi-data.TDB contains both a thermodynamic
SYS: @@ and kinetic description for the ordered and disordered BCC.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exil_setup.DCM
SYS:
SYS:
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: sw user AlFeNi-data.TDB
Current database: User defined Database
This database does not support the DATABASE_INFORMATION command

VA DEFINED
13:18:54,470 INFO  USER_1151090780_12, number of lines read: 609
13:18:55,000 INFO  Parsing of USER_1151090780_12 completed in 607 ms
TDB_USER: def-sys fe al ni
FE           AL           NI
DEFINED
TDB_USER: rej ph *
B2_BCC          BCC_DIS          B2_ORD
REJECTED
TDB_USER: res ph bcc_dis b2_ord
BCC_DIS         B2_ORD  RESTORED
TDB_USER: get
13:18:55,232 INFO  *** Invoking Gibbs Energy System v6 ***
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
FUNCTIONS .....
INFO: Forcing option USE_POLY3 for complex phase B2_ORD
-OK-
TDB_USER: go -m
NO TIME STEP DEFINED
DIC> set-cond glob T 0 1277; * N
DIC>
DIC> enter-region beta
DIC>
DIC> enter-grid beta
WIDTH OF REGION /1/: 2e-3
TYPE /LINEAR/: double
NUMBER OF POINTS /50/: 50
VALUE OF R IN THE GEOMETRICAL SERIE FOR LOWER PART OF REGION: 0.85
VALUE OF R IN THE GEOMETRICAL SERIE FOR UPPER PART OF REGION: 1.1765
DIC>
DIC> enter-phase
ACTIVE OR INACTIVE PHASE /ACTIVE/: act
REGION NAME : /BETA/: beta
PHASE TYPE /MATRIX/: matrix
PHASE NAME: /NONE/: b2_ord
DIC>
DIC> enter-composition
REGION NAME : /BETA/: beta
PHASE NAME: /B2_ORD/: b2_ord
DEPENDENT COMPONENT ? /NI/: fe
COMPOSITION TYPE /MOLE_FRACTION/: mole-fraction
PROFILE FOR /AL/: ni
TYPE /LINEAR/: function
Function F(X)= 0.28-0.277*erf((x-1e-3)/3e-6);
PROFILE FOR /NI/: al
TYPE /LINEAR/: function
Function F(X)= 0.4295-0.0105*erf((x-1e-3)/3e-6);
13:18:55,975 INFO  Preparing system for use: USER_1151090780_12
13:18:57,074 INFO  Preparing phase for use: B2_ORD
13:18:57,724 INFO  Preparing phase for use: BCC_DIS
DIC>
DIC> set-simulation-time
END TIME FOR INTEGRATION /1/: 345600
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /34560/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC> s-a-s-v
AUTOMATIC STARTING VALUES FOR PHASE COMPOSITIONS /YES/: no
START VALUES FOR PHASES IN REGION BETA
PHASE: B2_ORD
MAJOR CONSTITUENTS IN PHASE B2_ORD: NI;AL
DIC>
DIC>
DIC>
DIC>
DIC>
DIC> save exil yes
```

DIC>
DIC> set-inter
--OK---
DIC>

exil-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exil\run.DCM.test"
DIC>
DIC>
DIC> @@ exil_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE i1
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exil
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
INFO: FORCED STARTING VALUES TURNED ON
U-FRACTION IN SYSTEM: AL = .429499345639258 FE = .290517917020317
NI = .279982737340425
TOTAL SIZE OF SYSTEM: .002 [m]
U-FRACTION IN SYSTEM: AL = .429499345639258 FE = .290517917020317
NI = .279982737340425
TOTAL SIZE OF SYSTEM: .002 [m]
6 GRIDPOINT(S) ADDED TO CELL #1 REGION: BETA
TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639258 FE = .290517917020317
NI = .279982737340425
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 0.1001000E-03 DT = 0.1000000E-03 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639258 FE = .290517917020317
NI = .279982737340425
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 0.40010010 DT = 0.40000000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639258 FE = .290517917020317
NI = .279982737340425
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 0 seconds
TIME = 513.08856 DT = 512.68846 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639248 FE = .290517917020327
NI = .279982737340426
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 1538.4655 DT = 1025.3769 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639233 FE = .290517917020342
NI = .279982737340425
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 3589.2193 DT = 2050.7538 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639231 FE = .290517917020345
NI = .279982737340424
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 7690.7270 DT = 4101.5077 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639227 FE = .290517917020346
NI = .279982737340427
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 15893.742 DT = 8203.0154 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639233 FE = .290517917020342
NI = .279982737340426
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 32299.773 DT = 16406.031 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .42949934563921 FE = .290517917020363
NI = .279982737340427
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 65111.835 DT = 32812.062 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639174 FE = .290517917020383
NI = .279982737340443
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 99671.835 DT = 34560.000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639088 FE = .290517917020497
NI = .279982737340415
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 134231.83 DT = 34560.000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639131 FE = .290517917020483
NI = .279982737340385
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 168791.83 DT = 34560.000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639167 FE = .290517917020446
NI = .279982737340387
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 2 seconds
TIME = 203351.83 DT = 34560.000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639157 FE = .290517917020456
NI = .279982737340387
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 237911.83 DT = 34560.000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639151 FE = .290517917020441
NI = .279982737340408
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 272471.83 DT = 34560.000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639152 FE = .290517917020425
NI = .279982737340423
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
```

```
TIME = 307031.83 DT = 34560.000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .42949934563916 FE = .290517917020412
NI = .279982737340428
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 341591.83 DT = 34560.000 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639172 FE = .290517917020402
NI = .279982737340425
TOTAL SIZE OF SYSTEM: .002 [m]
CPU time used in timestep 1 seconds
TIME = 345600.00 DT = 4008.1652 SUM OF SQUARES = 0.0000000
U-FRACTION IN SYSTEM: AL = .429499345639173 FE = .290517917020401
NI = .279982737340426
TOTAL SIZE OF SYSTEM: .002 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.1000000E-06
DELETING TIME-RECORD FOR TIME 0.10010000E-03
DELETING TIME-RECORD FOR TIME 0.40010010
DELETING TIME-RECORD FOR TIME 513.08856
DELETING TIME-RECORD FOR TIME 1538.4655
DELETING TIME-RECORD FOR TIME 3589.2193
DELETING TIME-RECORD FOR TIME 7690.7270
DELETING TIME-RECORD FOR TIME 15893.742
DELETING TIME-RECORD FOR TIME 32299.773
DELETING TIME-RECORD FOR TIME 65111.835
DELETING TIME-RECORD FOR TIME 99671.835
DELETING TIME-RECORD FOR TIME 134231.83
DELETING TIME-RECORD FOR TIME 168791.83
DELETING TIME-RECORD FOR TIME 203351.83
DELETING TIME-RECORD FOR TIME 237911.83
DELETING TIME-RECORD FOR TIME 272471.83
DELETING TIME-RECORD FOR TIME 307031.83

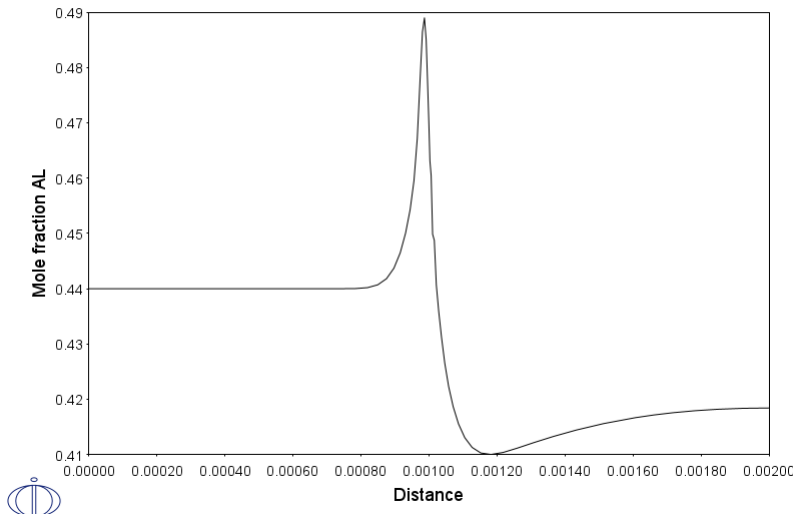
KEEPING TIME-RECORD FOR TIME 341591.83
AND FOR TIME 345600.00
WORKSPACE RECLAIMED
```

TIMESTEP AT 345600.000 SELECTED

DIC>
DIC> set-inter
--OK---
DIC>

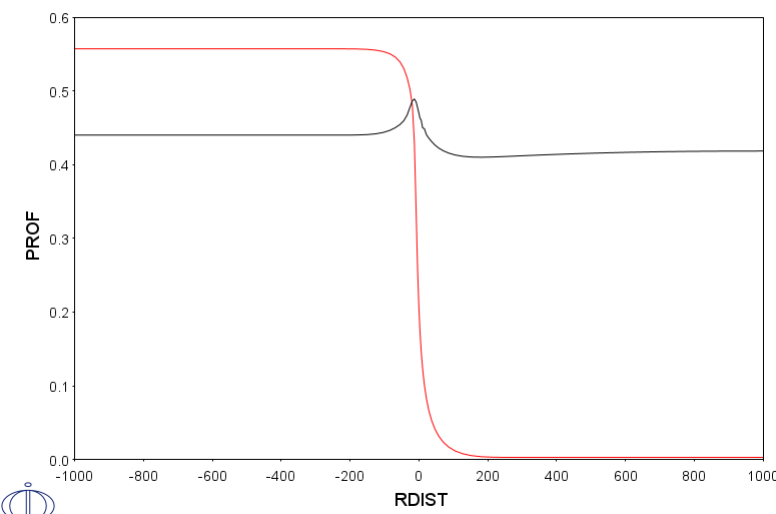
ex11-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\ex11\plot.DCM.test"
DIC>
DIC> @@ ex11_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE i1
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 3.45600E+05
DIC> read ex11
OK
DIC>
DIC> @@
DIC> @@ ENTER THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: s-d-a x dist glob
INFO: Distance is set as independent variable
POST-1:
POST-1: s-d-a y m-f al
POST-1:
POST-1: s-p-c time last
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.13.21.11
Time = 345600
CELL #1
```



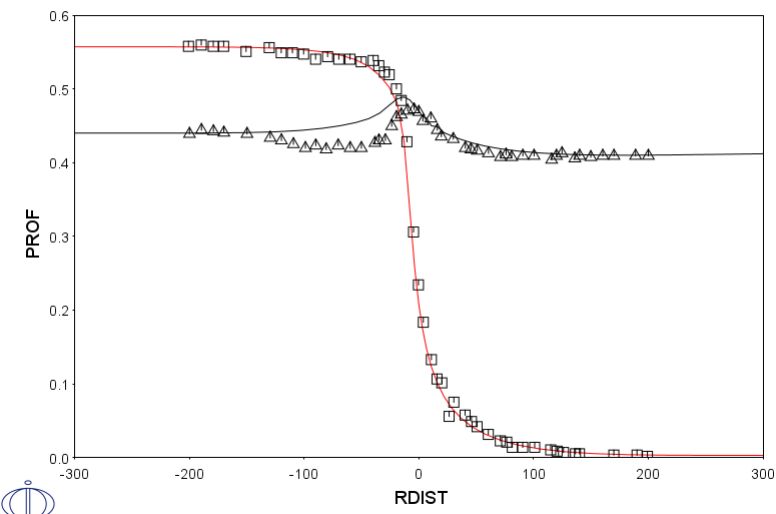
```
POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1:
POST-1: ent tab prof
Variable(s) x(al) x(ni)
POST-1:
POST-1: ent fun rdist
FUNCTION: 1e6*(gd-10e-4)
&
POST-1: s-d-a y prof
COLUMN NUMBER /*/: 1 2
POST-1:
POST-1: s-d-a x rdist
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

2021.05.13.21.12
Time = 345600
CELL #1



```
POST-1:  
POST-1:  
POST-1:@?<Hit_return_to_continue>  
POST-1:  
POST-1: app y exil.exp  
PROLOGUE NUMBER: /0/: 1  
DATASET NUMBER(s): /-1/: 1  
POST-1:  
POST-1: s-s-s x n -300 300  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
POST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

2021.05.13.21.12
Time = 345600
CELL #1



```
POST-1:  
POST-1:  
POST-1: set-inter  
--OK--  
POST-1:
```



Example exi2

Diffusion of carbon in cementite

This example demonstrates the use of the model for calculation of diffusion through a stoichiometric phase. The flux of a component in the stoichiometric phase is assumed to be proportional to the difference in chemical potential at each side of the stoichiometric phase multiplied with the mobility for the component in the phase. The mobility is assessed from experimental information and is basically the tracer diffusivity for the component. This calculation is compared with experimental data where a sample of pure iron has been exposed to a gas atmosphere with a certain carbon activity. The weight gain is then measured as a function of time. The experimental data is obtained from Ozturk B., Fearing V. L., Ruth A. Jr. and Simkovich G., Met. Trans A, vol 13A (1982), pp. 1871-1873.

$$J \sim \Delta\mu$$

exi2-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exi2\setup.DCM.test"
SYS: @@
SYS: @@ Diffusion in complex phases.
SYS: @@ Diffusion of carbon in cementite
SYS: @@ This example demonstrates the use of the model for calculation of
SYS: @@ diffusion through a stoichiometric phase. The flux of a component in
SYS: @@ the stoichiometric phase is assumed to be proportional to the
SYS: @@ difference in chemical potential at each side of the stoichiometric
SYS: @@ phase multiplied with the mobility for the component in the phase. The
SYS: @@ mobility is assessed from experimental information and is basically
SYS: @@ the tracer diffusivity for the component.
SYS: @@
SYS: @@ This calculation is compared with experimental data where a sample of
SYS: @@ pure iron has been exposed to a gas atmosphere with a certain carbon
SYS: @@ activity. The weight gain is then measured as a function of time.
SYS: @@ The experimental data is obtained from Ozturk B., Fearing V. L.,
SYS: @@ Ruth A. Jr. and Simkovich G., Met. Trans A, vol 13A (1982), pp. 1871-1873.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@
SYS: @@ RETRIEVE DATA FROM THE DATABASES
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA          /- DEFINED
DICTRA_FCC_A1  REJECTED
TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ USE A THERMODYNAMIC DATABASE TO RETRIEVE DATA
TDB_TCFE11: @@
TDB_TCFE11: switch FEDEMO
Current database: Iron Demo Database v4.0

VA          /- DEFINED
TDB_FEDEMO: def-sys fe c
FE          C  DEFINED
TDB_FEDEMO: rej ph * all
GAS:G          LIQUID:L
LAVES_PHASE_C14    CBCC_A12      CEMENTITE
CUB_A13        DIAMOND_FCC_A4   FCC_A1
GRAPHITE       HCP_A3         KSI_CARBIDE
M23C6          M5C2          M7C3
REJECTED

TDB_FEDEMO: res ph bcc fcc cementite grap
BCC_A2          FCC_A1      CEMENTITE
GRAPHITE RESTORED
TDB_FEDEMO: get
13:22:29,911 INFO *** Invoking Gibbs Energy System v6 ***
REINITIATING GES .....
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....

List of references for assessed data

'P. Franke, estimated parameter within SGTE, 2007; Fe-C, Ni-C, Mo-C, C-Mn'
'P. Gustafson, Scan. J. Metall., 14 (1985) 259-267; TRITA 0237 (1984); C-FE'
'X.G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'A.T. Dinsdale, SGTE Data for Pure Elements, CALPHAD, 15 (1991) 317-425'
'X.G. Lu, M. Selleby and B. Sundman, CALPHAD, 29, 2005, 68-89; Molar
volumes'
'B. Hallstedt, D. Djurovic, J. von Appen, R. Dronskowski, A. Dick, F.
Koermann, T. Hickel, J. Neugebauer, CALPHAD, 34, 129 -33(2010); Fe-C'
'B. Uhrenius, Int. J. Refract. Met. Hard Mater. 12 (1994) 121 -127; Molar
volumes'
-OK-
TDB_FEDEMO:
TDB_FEDEMO: @@
TDB_FEDEMO: @@ SWITCH TO A MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_FEDEMO: @@
TDB_FEDEMO: app user cembccstoik.TDB
Current database: User defined Database
This database does not support the DATABASE_INFORMATION command

VA  DEFINED
13:22:30,198 INFO USER_876738828_12, number of lines read: 51
13:22:30,353 INFO Parsing of USER_876738828_12 completed in 200 ms
TDB_APP: def-sys fe c
FE          C  DEFINED
TDB_APP: rej ph * all
BCC_A2          CEMENTITE  REJECTED
TDB_APP: res ph fcc bcc cementite
*** ERROR FCC INPUT IGNORED
BCC_A2          CEMENTITE  RESTORED
TDB_APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS .....
```

```

-OK-
TDB_APP: @@
TDE_APP: @@ ENTER THE DICTRA MONITOR
TDB_APP: @@
TDB_APP: go d-m
NO TIME STEP DEFINED
*** ENTERING FCC_A1 AS A DIFFUSION NONE PHASE
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC>
DIC> set-ref c grap,,,,,,,,,,,
13:22:31,300 INFO Preparing system for use: USER_876738828_12
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob t 0 723; * n
DIC>
DIC> @@
DIC> @@ ENTER THE REGIONS carb AND fer
DIC> @@
DIC> enter-region
REGION NAME : fer
DIC>
DIC> enter-region
REGION NAME : carb
ATTACH TO REGION NAMED /FER/:
ATTACHED TO THE RIGHT OF FER /YES/:
DIC> @@
DIC> @@ ENTER LINEAR GRIDS INTO THE REGIONS
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER A SIZE FOR THE FERRITE
DIC> @@
DIC> enter-grid
REGION NAME : /FER/: fer
WIDTH OF REGION /1/: 3.3E-6
TYPE /LINEAR/: AUTO
DIC>
DIC> @@
DIC> @@ ENTER A SIZE (VERY SMALL) FOR THE CEMENTITE LAYER
DIC> @@
DIC> @@
DIC> enter-grid
REGION NAME : /CARB/: carb
WIDTH OF REGION /1/: 1E-12
TYPE /LINEAR/: AUTO
DIC>
DIC>
DIC> @@
DIC> @@ ENTER PHASES INTO THE REGIONS
DIC> @@
DIC> enter-phase act carb matrix cementite
COMPOSITION RECORD FOR STOICHIOMETRIC PHASE CEMENTITE IN REGION CARB CREATED
DIC> enter-phase act fer matrix bcc#1
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITIONS IN THE PHASES
DIC> @@
DIC> enter-composition
REGION NAME : /FER/: carb
PHASE NAME: /CEMENTITE/: cementite
DIC>
DIC> enter-composition
REGION NAME : /FER/: fer
PHASE NAME: /BCC_A2/: bcc#1
COMPOSITION TYPE /MOLE_FRACTION/: weig-fraction
PROFILE FOR /C/: C lin 1E-5 1E-5
13:22:32,425 INFO Preparing phase for use: BCC_A2
DIC>
DIC> set-cond bound upp
CONDITION TYPE /CLOSED_SYSTEM/: mix
Dependent substitutional element:FE
Dependent interstitial element:VA
LOW TIME LIMIT /0/: 0
ACR(C) (TIME)= 9;
HIGH TIME LIMIT /*/: *
ANY MORE RANGES /N/: N
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ SIMULATE FOR 150 MINUTES
DIC> @@
DIC> set-simulation-time
END TIME FOR INTEGRATION /.1/: 9000
AUTOMATIC TIMESTEP CONTROL /YES/:
MAX TIMESTEP DURING INTEGRATION /900/:
INITIAL TIMESTEP : /1E-07/:
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exi2 Y
DIC>
DIC> set-inter
--OK---
DIC>
```

exi2-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exi2\run.DCM.test"
DIC>
DIC>
DIC> @@ exi2_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE i2
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
*** ENTERING FCC_A1 AS A DIFFUSION NONE PHASE
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC> read exi2
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim
Region: FER
single geometric dense at 0.33000E-05
0.91945 87
Region: CARB
double geometric
dense at outer boundaries, coarse at 0.50000E-12
lower part 1.2500 9
upper part 0.80000 9
DEGREE OF IMPLICITY SET TO TRAPEZOIDAL RULE
Trying old scheme 3
U-FRACTION IN SYSTEM: C = 4.6598005784384E-05 FE = 1
TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
U-FRACTION IN SYSTEM: C = 4.6598005784384E-05 FE = 1
TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
13:23:28,263 INFO Preparing phase for use: CEMENTITE
0.111094107740055 0.111116327660160 1.168933545268372E-022 TIME = 0.10000000E-06 DT = 0.10000000E-06
06 SUM OF SQUARES = 0.1169335E-21
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.55732855E-06 AND -0.55732855E-06
POSITION OF INTERFACE FER / CARB IS 0.32999999E-05
U-FRACTION IN SYSTEM: C = 4.65969237107415E-05 FE = 1
TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
2 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FER

CPU time used in timestep 0 seconds
1.985503622653896E-006 1.986016682715921E-006 8.569006692077880E-023 TIME = 0.25999666E-05 DT = 0.24999666E-05
05 SUM OF SQUARES = 0.85690067E-22
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.12341808E-06 AND -0.12341808E-06
POSITION OF INTERFACE FER / CARB IS 0.32999996E-05
U-FRACTION IN SYSTEM: C = 4.66223850218017E-05 FE = 1
TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
CPU time used in timestep 0 seconds
9.632817666939915E-007 9.640732404382838E-007 1.188386949892457E-025 TIME = 0.75998999E-05 DT = 0.49999333E-05
05 SUM OF SQUARES = 0.11883869E-24
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.91502734E-07 AND -0.91502734E-07
POSITION OF INTERFACE FER / CARB IS 0.32999992E-05
U-FRACTION IN SYSTEM: C = 4.66617913784151E-05 FE = 1
TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
CPU time used in timestep 1 seconds
5.520598281728859E-007 5.525040595679611E-007 1.288882881968142E-026 TIME = 0.17599767E-04 DT = 0.99998666E-04
05 SUM OF SQUARES = 0.12888829E-25
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.67380405E-07 AND -0.67380405E-07
POSITION OF INTERFACE FER / CARB IS 0.32999985E-05
U-FRACTION IN SYSTEM: C = 4.67208116371187E-05 FE = 1
TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
CPU time used in timestep 0 seconds
3.294904029785889E-007 3.297431202262135E-007 1.295444436331959E-026 TIME = 0.37599500E-04 DT = 0.19999733E-04
04 SUM OF SQUARES = 0.12954444E-25
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.48810906E-07 AND -0.48810906E-07
POSITION OF INTERFACE FER / CARB IS 0.32999975E-05
U-FRACTION IN SYSTEM: C = 4.68069809139129E-05 FE = 1
TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
CPU time used in timestep 0 seconds
1.842051604944532E-007 1.843420424822769E-007 2.883351411658548E-025 TIME = 0.77598966E-04 DT = 0.39999466E-04
04 SUM OF SQUARES = 0.28833514E-24
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.34957854E-07 AND -0.34957854E-07
POSITION OF INTERFACE FER / CARB IS 0.32999961E-05
U-FRACTION IN SYSTEM: C = 4.69308608646657E-05 FE = 1
TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
14 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARB

CPU time used in timestep 0 seconds
9.778279834954463E-008 9.785422397002213E-008 4.603903155444994E-026 TIME = 0.15759790E-03 DT = 0.79998933E-04
04 SUM OF SQUARES = 0.46039032E-25
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.24876846E-07 AND -0.24876846E-07
POSITION OF INTERFACE FER / CARB IS 0.32999941E-05
U-FRACTION IN SYSTEM: C = 4.7107484088289E-05 FE = 1
TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
49 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARB

CPU time used in timestep 0 seconds
5.038045607965860E-006 5.041694023499918E-006

output ignored...

... output resumed

TIME = 3142.1594 DT = 900.00000 SUM OF SQUARES = 0.30310202E-26
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.44854482E-11 AND -0.44854482E-11
POSITION OF INTERFACE FER / CARB IS 0.32722305E-05
U-FRACTION IN SYSTEM: C = .00283057618776225 FE = 1
TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
55 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARB

CPU time used in timestep 0 seconds
3.512901236418004E-008 3.518395504613353E-008 9.193038624674948E-
```

029 TIME = 4042.1594 DT = 900.00000 SUM OF SQUARES = 0.91930386E-28
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.38334138E-11 AND -0.38334138E-11
 POSITION OF INTERFACE FER / CARB IS 0.32687804E-05
 U-FRACTION IN SYSTEM: C = .00317904137490811 FE = 1
 TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
 9 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARB

CPU time used in timestep 0 seconds
 1.337273691002185E-008 1.340171249711458E-008 1.442674507444166E-
 026 TIME = 4942.1594 DT = 900.00000 SUM OF SQUARES = 0.14426745E-25
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.34097971E-11 AND -0.34097971E-11
 POSITION OF INTERFACE FER / CARB IS 0.32657116E-05
 U-FRACTION IN SYSTEM: C = .00348899893656278 FE = 1
 TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
 9 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARB

CPU time used in timestep 0 seconds
 6.238597393560772E-009 6.256204153621230E-009 6.091408152458267E-
 030 TIME = 5842.1594 DT = 900.00000 SUM OF SQUARES = 0.60914082E-29
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.31046285E-11 AND -0.31046285E-11
 POSITION OF INTERFACE FER / CARB IS 0.32629174E-05
 U-FRACTION IN SYSTEM: C = .0037712160470436 FE = 1
 TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
 20 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARB

CPU time used in timestep 0 seconds
 3.280281446382724E-009 3.291907995858972E-009 1.340043917426806E-
 030 TIME = 6742.1594 DT = 900.00000 SUM OF SQUARES = 0.13400439E-29
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.28707015E-11 AND -0.28707015E-11
 POSITION OF INTERFACE FER / CARB IS 0.32603338E-05
 U-FRACTION IN SYSTEM: C = .00403216870914771 FE = 1
 TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
 6 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARB

CPU time used in timestep 0 seconds
 1.864873577148230E-007 1.872981046499102E-007 1.7666718431636158E-
 029 TIME = 7642.1594 DT = 900.00000 SUM OF SQUARES = 0.176667184E-28
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.26837249E-11 AND -0.26837249E-11
 POSITION OF INTERFACE FER / CARB IS 0.32579185E-05
 U-FRACTION IN SYSTEM: C = .00427612481768571 FE = 1
 TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
 7 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARB

CPU time used in timestep 0 seconds
 1.118941666112022E-007 1.124814013565127E-007 5.965760595733902E-
 030 TIME = 8542.1594 DT = 900.00000 SUM OF SQUARES = 0.59657606E-29
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.25296909E-11 AND -0.25296909E-11
 POSITION OF INTERFACE FER / CARB IS 0.32556417E-05
 U-FRACTION IN SYSTEM: C = .00450607892012243 FE = 1
 TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
 6 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: CARB

CPU time used in timestep 1 seconds
 6.975845441408309E-008 7.019562007454278E-008 8.604937630166184E-
 030 TIME = 9000.0000 DT = 457.84063 SUM OF SQUARES = 0.86049376E-29
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.23998555E-11 AND -0.23998555E-11
 POSITION OF INTERFACE FER / CARB IS 0.32545430E-05
 U-FRACTION IN SYSTEM: C = .00461705531635636 FE = 1
 TOTAL SIZE OF SYSTEM: 3.300001E-06 [m]
 MUST SAVE WORKSPACE ON FILE
 WORKSPACE SAVED ON FILE
 RECLAIMING WORKSPACE
 DELETING TIME-RECORD FOR TIME 0.0000000
 DELETING TIME-RECORD FOR TIME 0.10000000E-06
 DELETING TIME-RECORD FOR TIME 0.25999666E-05
 DELETING TIME-RECORD FOR TIME 0.75998999E-05
 DELETING TIME-RECORD FOR TIME 0.17599767E-04
 DELETING TIME-RECORD FOR TIME 0.37599500E-04
 DELETING TIME-RECORD FOR TIME 0.77598966E-04
 DELETING TIME-RECORD FOR TIME 0.15759790E-03
 DELETING TIME-RECORD FOR TIME 0.31759576E-03
 DELETING TIME-RECORD FOR TIME 0.63759149E-03
 DELETING TIME-RECORD FOR TIME 0.12775830E-02
 DELETING TIME-RECORD FOR TIME 0.25575659E-02
 DELETING TIME-RECORD FOR TIME 0.51175317E-02
 DELETING TIME-RECORD FOR TIME 0.10237463E-01
 DELETING TIME-RECORD FOR TIME 0.20477327E-01
 DELETING TIME-RECORD FOR TIME 0.40957054E-01
 DELETING TIME-RECORD FOR TIME 0.81916507E-01
 DELETING TIME-RECORD FOR TIME 0.16383541
 DELETING TIME-RECORD FOR TIME 0.32767323
 DELETING TIME-RECORD FOR TIME 0.65534886
 DELETING TIME-RECORD FOR TIME 1.3107001
 DELETING TIME-RECORD FOR TIME 2.6214026
 DELETING TIME-RECORD FOR TIME 5.2428076
 DELETING TIME-RECORD FOR TIME 10.485618
 DELETING TIME-RECORD FOR TIME 20.971238
 DELETING TIME-RECORD FOR TIME 41.942478
 DELETING TIME-RECORD FOR TIME 83.884958
 DELETING TIME-RECORD FOR TIME 167.76992
 DELETING TIME-RECORD FOR TIME 335.53984
 DELETING TIME-RECORD FOR TIME 671.07968
 DELETING TIME-RECORD FOR TIME 1342.1594
 DELETING TIME-RECORD FOR TIME 2242.1594
 DELETING TIME-RECORD FOR TIME 3142.1594
 DELETING TIME-RECORD FOR TIME 4042.1594
 DELETING TIME-RECORD FOR TIME 4942.1594
 DELETING TIME-RECORD FOR TIME 5842.1594
 DELETING TIME-RECORD FOR TIME 6742.1594
 DELETING TIME-RECORD FOR TIME 7642.1594

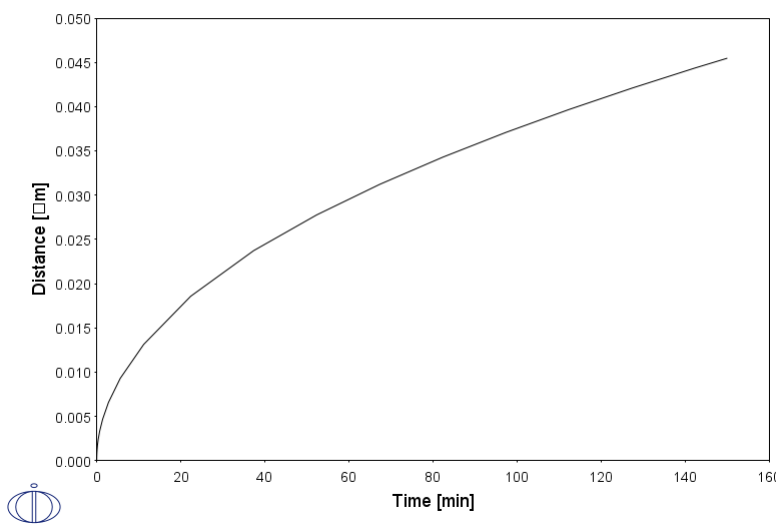
KEEPING TIME-RECORD FOR TIME 8542.1594
 AND FOR TIME 9000.0000
 WORKSPACE RECLAIMED

TIMESTEP AT 9000.00000 SELECTED

DIC>
DIC> set-inter
--OK--
DIC>

exi2-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exi2\plot.DCM.test"
DIC>
DIC> @@ exi2_plot.DCM
DIC>
DIC> @@
DIC> FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE i2
DIC> @@
DIC>
DIC> @@
DIC> GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 9.00000E+03
*** ENTERING FCC_A1 AS A DIFFUSION NONE PHASE
*** ENTERING GRAPHITE AS A DIFFUSION NONE PHASE
DIC>
DIC>
DIC> read exi2 Y
OK
DIC>
DIC>
DIC> @@
DIC> @@ PLOT THE SIZE OF THE CEMENTITE LAYER AS A FUNCTION OF TIME
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: ent-symb func csize
FUNCTION: 1e6*(poi(car,u)-poi(car,l));
POST-1:
POST-1: ent-symb func minutes
FUNCTION: time/60;
POST-1:
POST-1: s-d-a x minutes
POST-1: s-d-a y csize
POST-1:
POST-1: s-p-c inter first
POST-1:
POST-1: s-a-t-s x n Time [min]
POST-1: s-a-t-s y n Distance [ $\mu$ m]
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.13.24.30
"FIRST" INTERFACE OF SYSTEM
CELL #1
```

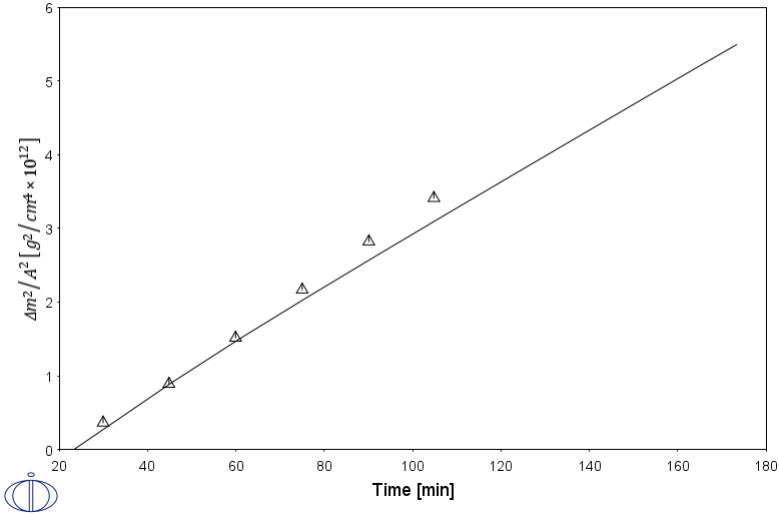


```
POST-1:
POST-1:@?<_hit_return_to_continue_>
POST-1:
POST-1:
POST-1: @@
POST-1: @@ ASSUME A CERTAIN TIME FOR NUCLEATION OF THE CEMENTITE LAYER
POST-1: @@
POST-1: ent-symb func cortim
FUNCTION: (time+1400)/60;
POST-1:
POST-1: @@
POST-1: @@ PLOT THE WEIGHT GAIN AS A FUNCTION OF TIME
POST-1: @@
POST-1: ent-symb func cwei
FUNCTION: 1e12*((poi(car,u)-poi(car,l)-1E-12)*12.01/2.33E-5*1e-4)**2;
POST-1:
POST-1:
POST-1: s-d-a x cortim
POST-1: s-d-a y cwei
POST-1:
POST-1: @@
POST-1: @@ COMPARE WITH EXPERIMENTAL DATA
POST-1: @@
POST-1: app y exi2.exp 0; 1
POST-1:
POST-1:
```

```

POST-1: s-a-t-s x n Time [min]
POST-1: s-a-t-s y n \Delta m^2/A^2, [g^2/cm^4\times 10^{12}]
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.13.24.30
"FIRST" INTERFACE OF SYSTEM
CELL #1

```



```

POST-1:
POST-1:
POST-1: set-inter
--OK---
POST-1:

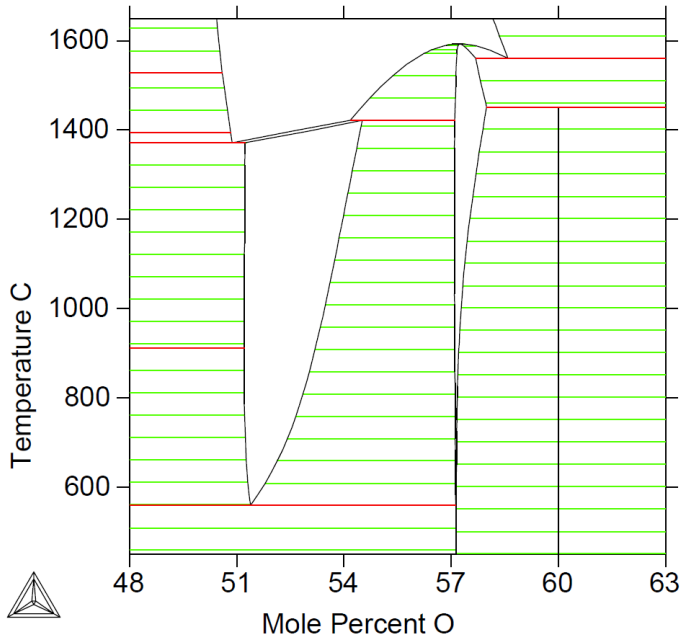
```



Example exi3a

Diffusion in iron oxide (FeO)

This example shows the oxidation of an iron sample and the consequent growth of an oxide layer using the grain boundary diffusion contribution model.



exi3a-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

```
SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exi3a\setup.DCM.test"
SYS: @@
SYS: @@ Diffusion in complex phases.
SYS: @@ Diffusion in iron oxide (FeO)
SYS: @@ This example shows the oxidation of an iron sample and the
SYS: @@ consequent growth of an oxide layer.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exi3_setup.DCM
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA           /- DEFINED
DICTRA_FCC_A1 REJECTED

TDB_TCFE11:
TDB_TCFE11: @@
TDB_TCFE11: @@ SELECT A USER DATABASE FOR READING THE THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw user FeO.TDB
Current database: User defined Database
This database does not support the DATABASE_INFORMATION command

VA           /- DEFINED
13:25:48,100 INFO  USER_467798867_12, number of lines read: 217
13:25:48,460 INFO  Parsing of USER_467798867_12 completed in 425 ms
TDB_USER: def-sys fe o
FE           O      DEFINED
TDB_USER: rej sp *
/-          VA           FE
O            FE+2         FE+3
FE+4        FE2O3        FEO
FEO3/2      O-2          O2
REJECTED

TDB_USER: res sp fe fe+2 fe+3 o o2 o-2 va
FE           FE+2         FE+3
O            O2           O-2
VA RESTORED
TDB_USER: rej ph * all
GAS:G        BCC_A2       SPINEL:I
REJECTED

TDB_USER: res ph bcc spinel gas
BCC_A2       SPINEL:I   GAS:G
RESTORED

TDB_USER:
TDB_USER: get
13:25:48,646 INFO  *** Invoking Gibbs Energy System v6 ***
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS ...
FUNCTIONS ...

List of references for assessed data

'A. Dinsdale, SGTE Data for Pure Elements, Calphad, 15 (1991), 317 -425'
'M. Kowalski and P.J. Spencer, Calphad, 19 (1995), 229-243; Cr-O, Fe-O and
'Ni-O'
'X.-G. Lu, Thermo-Calc Software AB, Sweden,2006; Molar volumes'
'X.-G. Lu, M. Selleby and B. Sundman, CALPHAD, Vol. 29, 2005, pp. 68-89;
'Molar volumes'
'B. Sundman, J. Phase Equil., 12 (1991), 127-140; Fe-O'
-OK-

*** WARNING: One or more elements have been rejected only as species but not as elements. This is not allowed with GES6, temporarily reverting

TDB_USER:
TDB_USER: @@
TDB_USER: @@ SWITCH TO A USER-DEFINED MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_USER: @@
TDB_USER: app user FeQmob.TDB
Current database: User defined Database
test database

VA           /-           O
DEFINED
13:25:48,760 INFO  USER_1724833957_12, number of lines read: 129
13:25:48,811 INFO  Parsing of USER_1724833957_12 completed in 52 ms
TDB_APP: def-sys fe o
FE DEFINED
TDB_APP: rej sp *
/-          VA           FE
O            FE+2         FE+3
FE2O3        FEO          FEO3/2
O-2          O2           O-2
REJECTED

TDB_APP: res sp fe fe+2 fe+3 o o2 o-2 va
FE           FE+2         FE+3
O            O2           O-2
VA RESTORED
TDB_APP: rej ph * all
SPINEL:I     GAS:G       BCC_A2
```

```

REJECTED
TDB_APP: res ph bcc spinel gas
BCC_A2 SPINEL:I GAS:G
RESTORED
TDB_APP:
TDB_APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
*** ERROR 2018 IN DESHPH: DIFFUSION DATA NOT POSSIBLE FOR GAS PHASE
*** ERROR 2018 IN DESHPH: DIFFUSION DATA NOT POSSIBLE FOR GAS PHASE
FUNCTIONS .....
-OK-
TDB_APP:
TDB_APP:
TDB_APP: @@
TDB_APP: @@ ENTER THE DICTRA MONITOR
TDB_APP: @@
TDB_APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 823; * N
DIC>
DIC>
DIC> @@
DIC> @@ SET THE REFERENCE STATE FOR O TO O2 (GAS)
DIC> @@
DIC> set-ref o gas,.....
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE REGIONS fer AND sp
DIC> @@
DIC> ent-reg fer
DIC> ent-reg sp,.....
DIC>
DIC>
DIC> @@
DIC> @@ ENTER PHASES INTO THE REGIONS
DIC> @@
DIC> ent-phase act fer matrix bcc#1
DIC> ent-phase act sp matrix spinel
DIC>
DIC>
DIC> @@
DIC> @@ ENTER GRIDS INTO THE REGIONS
DIC> @@
DIC>
DIC>
DIC> @@
DIC> @@ ENTER A SIZE FOR THE FERRITE
DIC> @@
DIC> @@
DIC> ent-grid fer 4.99999e-3 AUTO
DIC>
DIC> @@
DIC> @@ ENTER A THIN INITIAL SIZE FOR THE OXIDE
DIC> @@
DIC> @@
DIC> ent-grid sp 1.00e-10 AUTO
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITIONS IN BCC
DIC> @@
DIC> ent-comp fer bcc#1 m-f
PROFILE FOR /O/: o lin 1e-9 1e-9
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITIONS IN THE OXIDE
DIC> @@
DIC> ent-comp sp spinel m-f
this is a phase with charged species
with more than 2 sublattices
PROFILE FOR /FE/: FE lin 4.28771E-01 4.28549E-01
DIC>
DIC>
DIC> @@
DIC> @@ ENTER A BOUNDARY CONDITION "GAS" ON THE UPPER (RIGHT-MOST) INTERFACE
DIC> @@ OF THE OXIDE. THIS ALLOWS THE SYSTEM TO EXPAND AND THE OXIDE LAYER
DIC> @@ TO GROW EXTERNALLY. FOR THIS EXAMPLE AN OXYGEN ACTIVITY IS SPECIFIED
DIC> @@ THAT IS LOW ENOUGH NOT TO FORM CORUNDUM (FE2O3). WE ALSO SPECIFY
DIC> @@ THAT THERE IS NO FLUX OF Fe ACROSS THIS INTERFACE, I.E. NO Fe
DIC> @@ IS ALLOWED TO ENTER OR LEAVE THE SYSTEM.
DIC> @@
DIC> set-cond boundary upper gas
TYPE OF CONDITION FOR COMPONENT FE /ZERO_FLUX/: zero-flux
TYPE OF CONDITION FOR COMPONENT O /ZERO_FLUX/: act
LOW TIME LIMIT /0/: 0 4.5e-4; * N
DIC>
DIC>
DIC> @@
DIC> @@ ENTER START VALUES FOR THE INITIAL INTERFACE VELOCITIES
DIC> @@
DIC> s-a-s-v -1e-5 1e-5 yes
STARTING VALUES WILL BE TAKEN FROM PROFILES
DIC>
DIC>
DIC> @@
DIC> @@ SIMULATE FOR 24 HOURS
DIC> @@
DIC> s-s-time 86400,,
SMALLEST ACCEPTABLE Timestep : /1E-07/:
DIC> @@
DIC> @@ SPECIFY THAT POTENTIALS AND NOT ACTIVITIES ARE VARIED AT THE PHASE

```

```
DIC> @@ INTERFACE. ALSO USE A FULLY IMPLICIT SCHEME FOR TIME INTEGRATION.  
DIC> @@  
DIC> s-s-c 0 1 1 YES POT YES YES 1 2,.....  
RELEASING OLD STARTING VALUES  
DIC>  
DIC>  
DIC> @@  
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT  
DIC> @@  
DIC> save exi3.DIC Y  
DIC>  
DIC> set-inter  
--OK---  
DIC>
```

exi3a-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exi3a\run.DCM.test"
DIC>
DIC>
DIC> @@ exi3_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE i3
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exi3
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim y
Region: FER
single geometric dense at 0.50000E-02
0.87701      95
Region: SP
double geometric
dense at outer boundaries, coarse at 0.50000E-10
lower part 1.2500      9
upper part 0.80000      9
Trying old scheme
U-FRACTION IN SYSTEM: FE = 0.99999999500542 O = 2.10000395011622E-08
TOTAL SIZE OF SYSTEM: .0049999901 [m]
U-FRACTION IN SYSTEM: FE = 0.99999999500542 O = 2.10000395011622E-08
TOTAL SIZE OF SYSTEM: .0049999901 [m]
0.347387301620389 0.347386528700301 0.347386701355537 4.618089872590951E-002 6.656544255012604E-
023 TIME = 0.10000000E-06 DT = 0.10000000E-06 SUM OF SQUARES = 0.66565443E-22
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.39646054E-03 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49999900E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.66982674E-03 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.49999901E-02
U-FRACTION IN SYSTEM: FE = 0.99999998954903 O = 3.44025119032269E-08
TOTAL SIZE OF SYSTEM: .00499999012734 [m]
9 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FER

CPU time used in timestep 1 seconds
1.507552605981734E-005 1.507739146458280E-005 1.507822847278430E-005 2.226308397553746E-007 5.019252732530892E-
032 TIME = 0.30000000E-06 DT = 0.20000000E-06 SUM OF SQUARES = 0.50192527E-31
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.14849229E-03 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49999989E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.20393047E-03 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.49999901E-02
U-FRACTION IN SYSTEM: FE = 0.99999998713864 O = 4.25594164341045E-08
TOTAL SIZE OF SYSTEM: .00499999013842 [m]
CPU time used in timestep 0 seconds
1.398975101884493E-008 1.397926494986202E-008 1.395940586749686E-008 3.137487097463036E-011 3.385423730368188E-
036 TIME = 0.70000000E-06 DT = 0.40000000E-06 SUM OF SQUARES = 0.33854237E-35
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.14913262E-03 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49999989E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.21655229E-03 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.49999902E-02
U-FRACTION IN SYSTEM: FE = 0.99999998167656 O = 5.98849352760284E-08
TOTAL SIZE OF SYSTEM: .00499999016539 [m]
CPU time used in timestep 0 seconds
5.609742547736858E-007 5.611248463497880E-007 5.611264634683927E-007 2.444637656461759E-010 1.543482857347205E-
033 TIME = 0.15000000E-05 DT = 0.80000000E-06 SUM OF SQUARES = 0.15434829E-32
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.93393028E-04 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49999989E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.13339356E-03 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.49999902E-02
U-FRACTION IN SYSTEM: FE = 0.99999997507588 O = 8.12308039073763E-08
TOTAL SIZE OF SYSTEM: .00499999019739 [m]
23 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 1 seconds
8.108701238861243E-006 8.112502967034163E-006 8.112138099431511E-006 1.121067579181819E-008 2.140565808892579E-
032 TIME = 0.31000000E-05 DT = 0.16000000E-05 SUM OF SQUARES = 0.21405658E-31
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.70760985E-04 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49999897E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.10183290E-03 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.49999902E-02
U-FRACTION IN SYSTEM: FE = 0.99999996490465 O = 1.13822384430487E-07
TOTAL SIZE OF SYSTEM: .00499999024711 [m]
35 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 0 seconds
8.100481734460367E-006 8.103302413600414E-006 8.103162850520866E-006 7.195354229709166E-009 1.458658714794128E-
031 TIME = 0.63000000E-05 DT = 0.32000000E-05 SUM OF SQUARES = 0.14586587E-30
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.49011444E-04 AND 0.0000000

output ignored...

... output resumed

POSITION OF INTERFACE FER / SP IS 0.49497263E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.85406534E-09 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.50216007E-02
U-FRACTION IN SYSTEM: FE = .995601423066387 O = .0143132204530815
TOTAL SIZE OF SYSTEM: .005021600735 [m]
23 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 0 seconds
2.789556771575429E-007 2.79405758322531E-007 2.793655766630949E-007 3.800585454837353E-010 2.411174012406676E-
033 TIME = 56943.895 DT = 8640.0000 SUM OF SQUARES = 0.24111740E-32
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.54743813E-09 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49449965E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.78319309E-09 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.50236377E-02
```

U-FRACTION IN SYSTEM: FE = .995194725257641 O = .0156544064678654
TOTAL SIZE OF SYSTEM: .00502363765793 [m]
8 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 0 seconds
1.595105700155058E-007 1.598217866208553E-007 1.597953811124363E-007 2.020602965724977E-010 1.145025537248974E-
033 TIME = 65583.895 DT = 8640.0000 SUM OF SQUARES = 0.11450255E-32
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.50751127E-09 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49406116E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.72677808E-09 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.50255321E-02
U-FRACTION IN SYSTEM: FE = .994817249745987 O = .0168979971924892
TOTAL SIZE OF SYSTEM: .0050255321232 [m]
7 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 0 seconds
9.568367496895681E-008 9.590684461689169E-008 9.588874680435883E-008 1.147888044286007E-010 6.018531076210112E-
036 TIME = 74223.895 DT = 8640.0000 SUM OF SQUARES = 0.60185311E-35
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.47495053E-09 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49365080E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.68062878E-09 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.50273092E-02
U-FRACTION IN SYSTEM: FE = .99446375481309 O = .0180617615442488
TOTAL SIZE OF SYSTEM: .00502730918328 [m]
10 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 0 seconds
5.962030964014512E-008 5.978502104198708E-008 5.977212462770474E-008 6.869420013219069E-011 3.190573786775886E-
033 TIME = 82863.895 DT = 8640.0000 SUM OF SQUARES = 0.31905738E-32
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.44778107E-09 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49326392E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.64203085E-09 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.50289875E-02
U-FRACTION IN SYSTEM: FE = .994130359456913 O = .0191587681410927
TOTAL SIZE OF SYSTEM: .00502898750134 [m]
8 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 1 seconds
9.162180973662354E-009 9.108164932083602E-009 9.224421743951149E-009 1.011580317218201E-011 3.385423730368188E-
034 TIME = 86400.000 DT = 3536.1047 SUM OF SQUARES = 0.33854237E-33
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.44500201E-09 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49310656E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.61834793E-09 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.50296005E-02
U-FRACTION IN SYSTEM: FE = .994008758985554 O = .0195911680902242
TOTAL SIZE OF SYSTEM: .00502960047066 [m]
MUST SAVE WORKSPACE ON FILE
WORKSPACE SAVED ON FILE
RECLAIMING WORKSPACE
DELETING TIME-RECORD FOR TIME 0.0000000
DELETING TIME-RECORD FOR TIME 0.10000000E-06
DELETING TIME-RECORD FOR TIME 0.30000000E-06
DELETING TIME-RECORD FOR TIME 0.70000000E-06
DELETING TIME-RECORD FOR TIME 0.15000000E-05
DELETING TIME-RECORD FOR TIME 0.31000000E-05
DELETING TIME-RECORD FOR TIME 0.63000000E-05
DELETING TIME-RECORD FOR TIME 0.12700000E-04
DELETING TIME-RECORD FOR TIME 0.25500000E-04
DELETING TIME-RECORD FOR TIME 0.51100000E-04
DELETING TIME-RECORD FOR TIME 0.10230000E-03
DELETING TIME-RECORD FOR TIME 0.20470000E-03
DELETING TIME-RECORD FOR TIME 0.40950000E-03
DELETING TIME-RECORD FOR TIME 0.81910000E-03
DELETING TIME-RECORD FOR TIME 0.16383000E-02
DELETING TIME-RECORD FOR TIME 0.32767000E-02
DELETING TIME-RECORD FOR TIME 0.65535000E-02
DELETING TIME-RECORD FOR TIME 0.13107100E-01
DELETING TIME-RECORD FOR TIME 0.26214300E-01
DELETING TIME-RECORD FOR TIME 0.52428700E-01
DELETING TIME-RECORD FOR TIME 0.10485750
DELETING TIME-RECORD FOR TIME 0.20971510
DELETING TIME-RECORD FOR TIME 0.41943030
DELETING TIME-RECORD FOR TIME 0.83886070
DELETING TIME-RECORD FOR TIME 1.6777215
DELETING TIME-RECORD FOR TIME 3.3554431
DELETING TIME-RECORD FOR TIME 6.7108863
DELETING TIME-RECORD FOR TIME 13.421773
DELETING TIME-RECORD FOR TIME 26.843545
DELETING TIME-RECORD FOR TIME 53.687091
DELETING TIME-RECORD FOR TIME 107.37418
DELETING TIME-RECORD FOR TIME 214.74836
DELETING TIME-RECORD FOR TIME 429.49673
DELETING TIME-RECORD FOR TIME 858.99346
DELETING TIME-RECORD FOR TIME 1717.98669
DELETING TIME-RECORD FOR TIME 3435.9738
DELETING TIME-RECORD FOR TIME 6871.9477
DELETING TIME-RECORD FOR TIME 13743.895
DELETING TIME-RECORD FOR TIME 22383.895
DELETING TIME-RECORD FOR TIME 31023.895
DELETING TIME-RECORD FOR TIME 39663.895
DELETING TIME-RECORD FOR TIME 48303.895
DELETING TIME-RECORD FOR TIME 56943.895
DELETING TIME-RECORD FOR TIME 65583.895
DELETING TIME-RECORD FOR TIME 74223.895

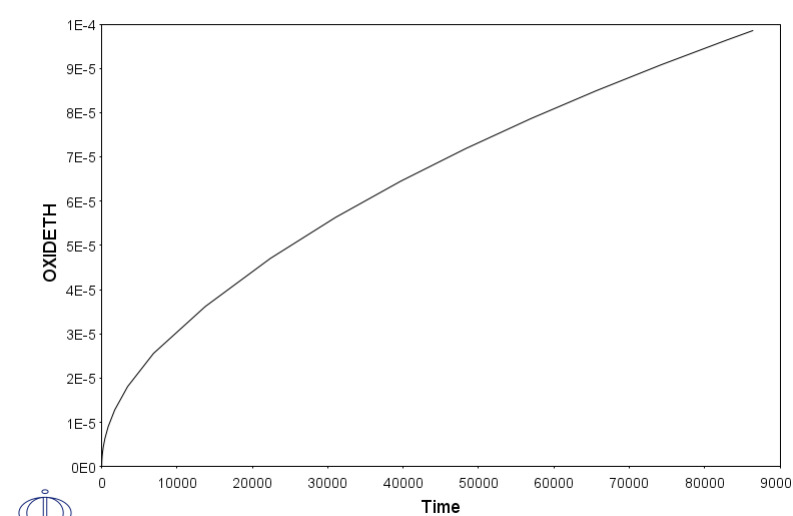
KEEPING TIME-RECORD FOR TIME 82863.895
AND FOR TIME 86400.000
WORKSPACE RECLAIMED

TIMESTEP AT 86400.0000 SELECTED

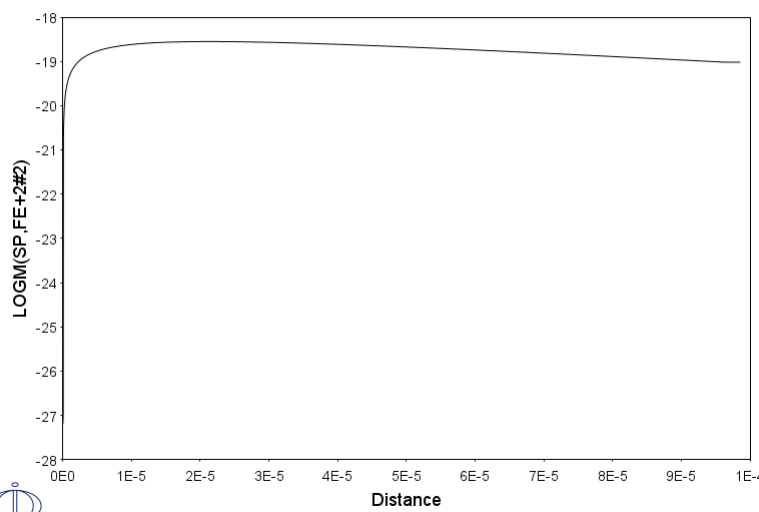
DIC>
DIC> set-inter
--OK--
DIC>

exi3a-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exi3a\plot.DCM.test"
DIC>
DIC> @@ exi3_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE i3
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 8.64000E+04
DIC> read exi3
OK
DIC>
DIC> @@
DIC> @@ ENTER THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ PLOT THE THICKNESS OF THE OXIDE LAYER GROWING AT THE SURFACE.
POST-1: @@ FOR THIS WE NEED TO ENTER A FUNCTION ACCORDING TO THE FOLLOWING.
POST-1: ent func oxideth
FUNCTION: poi(sp,upper)-poi(sp,lower)
&
POST-1: @@
POST-1: @@ PUT THIS FUNCTION ON THE Y-AXIS
POST-1: @@
POST-1: s-d-a y oxideth
POST-1: @@
POST-1: @@
POST-1: @@ AND PLOT THE OXIDE THICKNESS VERSUS TIME
POST-1: @@
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1: @@
POST-1: @@
POST-1: @@ SINCE WE ARE PLOTTING A FUNCTION, SPECIFY A PLOT CONDITION
POST-1: @@
POST-1: s-p-c interface sp upper
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.13.27.53
UPPER INTERFACE OF REGION "SP#1"
CELL #1

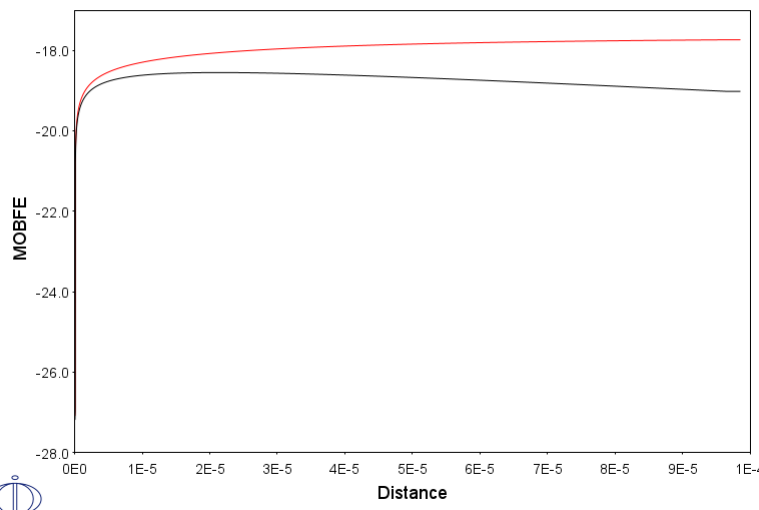

POST-1:
POST-1:@?<Hit_return_to_continue>
POST-1:
POST-1: @@
POST-1: @@ NOW PLOT THE MOBILITY IN A SPINEL FOR Fe+2 ON THE SECOND SUBLATTICE
POST-1: @@
POST-1: s-d-a y logm(sp,fe+2#2)
POST-1:
POST-1: @@
POST-1: @@ LIMIT THE PLOT TO THE SPINEL PHASE
POST-1: @@
POST-1: s-d-a x dis local sp
INFO: Distance is set as independent variable
POST-1:
POST-1: s-p-c time 86400
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 5
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
POST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
```

2021.05.13.13.27.54
Time = 86400
CELL #1



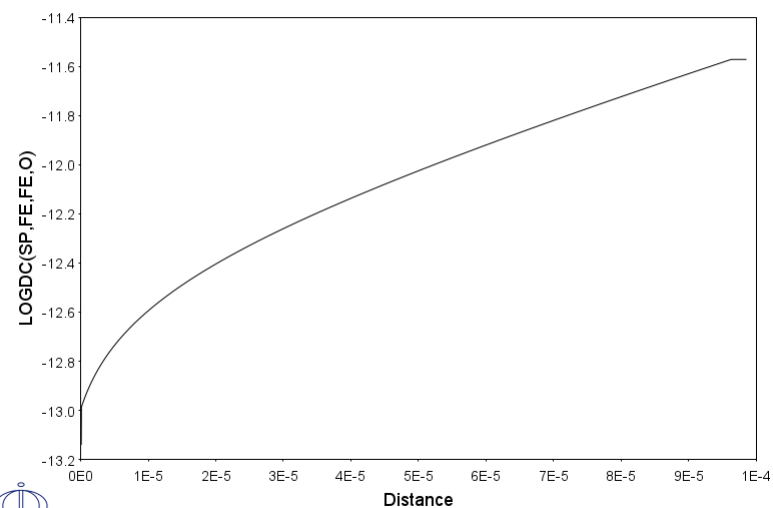
```
POST-1:  
POST-1:  
POST-1:@?<Hit_return_to_continue>  
POST-1:  
POST-1: @@  
POST-1: @@ COMPARE MOBILITIES IN A SPINEL FOR Fe+2 AND Fe+3 SPECIES PRESENT ON THE  
POST-1: @@ SECOND SUBLATTICE. FOR THIS WE NEED TO ENTER A TABLE.  
POST-1: @@  
POST-1: ent table mobfe  
Variable(s) logm(sp,fe+2#2) logm(sp,fe+3#2)  
POST-1:  
POST-1: s-d-a y mobfe  
COLUMN NUMBER /*:  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
POST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

2021.05.13.13.27.55
Time = 86400
CELL #1



```
POST-1:  
POST-1:  
POST-1:@?<Hit_return_to_continue>  
POST-1: @@  
POST-1: @@ NOW PLOT THE INTERDIFFUSION COEFFICIENT OF Fe IN A SPINEL  
POST-1: @@  
POST-1: s-d-a y logdc(sp,fe,fe,o)  
POST-1:  
POST-1:  
POST-1: SET_EXP_FILE_FORMAT 5  
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y  
POST-1: SET_EXP_FILE_FORMAT 10  
POST-1:  
POST-1: plot
```

2021.05.13.13.27.57
Time = 86400
CELL #1



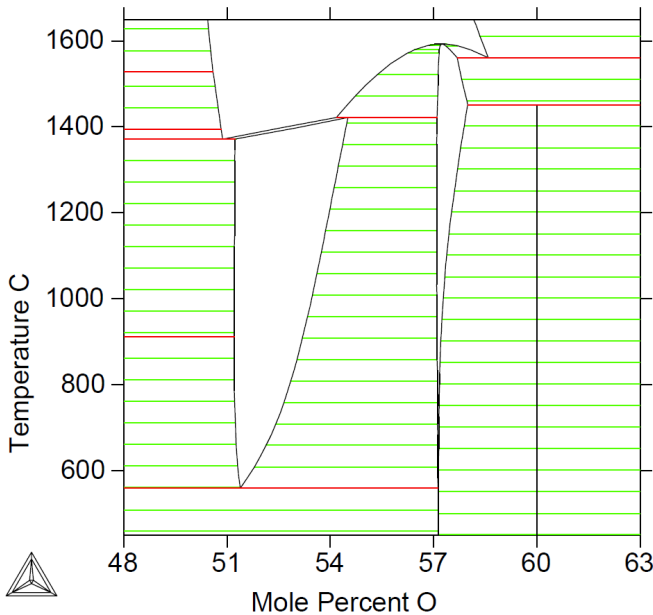
POST-1:
POST-1:
POST-1: set-inter
--OK---
POST-1:



Example i3b

Diffusion in iron oxide (FeO) with grain boundary contribution

Oxidation of iron sample and consequent growth of an oxide layer using the grain boundary diffusion contribution model.



exi3b-setup**SYS:About**

Thermo-Calc / DICTRA is software package for calculation of phase diagrams, simulation of phase transformation kinetics and much more.

Copyright Foundation for Computational Thermodynamics,
Stockholm, Sweden

Software (build 24339) running on WinNT 64-bit wordlength
Compiler: Intel(R) Visual Fortran Compiler Version 16.0.4.246 Build 20160811
License library version: 8.5.1.0017
Linked: Wed May 12 14:13:16 2021

SYS:SYS:MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exi3b\setup.DCM.test"

SYS: @@
SYS: @@ Diffusion in complex phases.
SYS: @@ Diffusion in iron oxide (FeO) with a grain boundary contribution
SYS: @@ This example shows the oxidation of an iron sample and consequent
SYS: @@ growth of an oxide layer using the grain boundary diffusion
SYS: @@ contribution model.
SYS: -----
NO SUCH COMMAND, USE HELP
SYS:
SYS: @@ exi3_setup.DCM
SYS:
SYS: @@
SYS: @@ START BY GOING TO THE DATABASE MODULE
SYS: @@
SYS: go da
THERMODYNAMIC DATABASE module
Database folder: C:\jenkins\workspace\dictra_generate_console_examples\data
Current database: Steels/Fe-Alloys v11.0

VA /- DEFINED
DICTRA_FCC_A1 REJECTED
TDB_TCFE11: @@
TDB_TCFE11: @@ SELECT A USER DATABASE TO READ THE THERMODYNAMIC DATA
TDB_TCFE11: @@
TDB_TCFE11: sw user FeO.TDB
Current database: User defined Database
This database does not support the DATABASE_INFORMATION command

VA /- DEFINED
13:29:14,721 INFO USER_2019713966_12, number of lines read: 217
13:29:15,093 INFO Parsing of USER_2019713966_12 completed in 439 ms
TDB_USER: def-sys fe o
FE O DEFINED
TDB_USER: rej sp *
/- VA FE
O FE+2 FE+3
FE+4 FE2O3 FEO
FE03/2 O-2 O2
REJECTED
TDB_USER: res sp fe fe+2 fe+3 o o2 o-2 va
FE FE+2 FE+3
O O2 O-2
VA RESTORED
TDB_USER: rej ph * all
GAS:G BCC_A2 SPINEL:I
REJECTED
TDB_USER: res ph bcc spinel gas
BCC_A2 SPINEL:I GAS:G
RESTORED
TDB_USER:
TDB_USER: get
13:29:15,298 INFO *** Invoking Gibbs Energy System v6 ***
ELEMENTS
SPECIES
PHASES
PARAMETERS ...
FUNCTIONS

List of references for assessed data

'A. Dinsdale, SGTE Data for Pure Elements, Calphad, 15 (1991), 317 -425'
'M. Kowalski and P.J. Spencer, Calphad, 19 (1995), 229-243; Cr-O, Fe-O and
Ni-O'
'X.-G. Lu, Thermo-Calc Software AB, Sweden, 2006; Molar volumes'
'X.-G. Lu, M. Selleby and B. Sundman, CALPHAD, Vol. 29, 2005, pp. 68-89;
Molar volumes'
'B. Sundman, J. Phase Equil., 12 (1991), 127-140; Fe-O'
-OK-

*** WARNING: One or more elements have been rejected only as species but not as elements. This is not allowed with GES6, temporarily reverting

TDB_USER: @@
TDB_USER: @@ SWITCH TO A USER-DEFINED MOBILITY DATABASE TO RETRIEVE MOBILITY DATA
TDB_USER: @@
TDB_USER: app user FeOmob.TDB
Current database: User defined Database
test database

VA /- O
DEFINED
13:29:15,385 INFO USER_1595751238_12, number of lines read: 128
13:29:15,433 INFO Parsing of USER_1595751238_12 completed in 51 ms
TDB_APP: def-sys fe o
FE DEFINED
TDB_APP: rej sp *
/- VA FE
O FE+2 FE+3
FE2O3 FEO FE03/2
O-2 O2 O2 REJECTED
TDB_APP: res sp fe fe+2 fe+3 o o2 o-2 va
FE FE+2 FE+3
O O2 O-2
VA RESTORED
TDB_APP: rej ph * all

```

SPINEL:I           GAS:G          BCC_A2
REJECTED
TDB_APP: res ph bcc spinel gas
BCC_A2           SPINEL:I        GAS:G
RESTORED
TDB_APP:
TDB_APP: get
ELEMENTS .....
SPECIES .....
PHASES .....
PARAMETERS .....
*** ERROR 2018 IN DESHPH: DIFFUSION DATA NOT POSSIBLE FOR GAS PHASE
*** ERROR 2018 IN DESHPH: DIFFUSION DATA NOT POSSIBLE FOR GAS PHASE
FUNCTIONS .....
-OK-
TDB_APP:
TDB_APP:
TDB_APP: @@ @
TDB_APP: @@ ENTER THE DICTRA MONITOR
TDB_APP: @@
TDB_APP: go d-m
NO TIME STEP DEFINED
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE GLOBAL CONDITION T
DIC> @@
DIC> set-cond glob T 0 823; * N
DIC>
DIC>
DIC> @@
DIC> @@ SET THE REFERENCE STATE FOR O TO O2 (GAS)
DIC> @@
DIC> set-ref o gas,,,,,,,
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE REGIONS fer AND sp
DIC> @@
DIC> ent-reg fer
DIC> ent-reg sp,,,,,,,
DIC>
DIC>
DIC> @@
DIC> @@ ENTER PHASES INTO THE REGIONS
DIC> @@
DIC> ent-phase act fer matrix bcc#1
DIC> ent-phase act sp matrix spinel
DIC>
DIC>
DIC> @@
DIC> @@ ENTER GRIDS INTO THE REGIONS
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER A SIZE FOR THE FERRITE
DIC> @@
DIC>
DIC> ent-grid fer 4.99999e-3 AUTO
DIC>
DIC> @@
DIC> @@ ENTER A THIN INITIAL SIZE FOR THE OXIDE
DIC> @@
DIC>
DIC> ent-grid sp 1.00e-10 AUTO
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITIONS IN BCC
DIC> @@
DIC> ent-comp fer bcc#1 m-f
PROFILE FOR /O/: o lin 1e-9 1e-9
DIC>
DIC>
DIC> @@
DIC> @@ ENTER THE INITIAL COMPOSITIONS IN THE OXIDE
DIC> @@
DIC> ent-comp sp spinel m-f
this is a phase with charged species
with more than 2 sublattices
PROFILE FOR /FE/: fe lin 4.28771E-01 4.28549E-01
DIC>
DIC>
DIC> @@
DIC> @@ ENTER A BOUNDARY CONDITION "GAS" ON THE UPPER (RIGHT-MOST) INTERFACE
DIC> @@ OF THE OXIDE. THIS ALLOWS THE SYSTEM TO EXPAND AND THE OXIDE LAYER
DIC> @@ TO GROW EXTERNALLY. FOR THIS EXAMPLE AN OXYGEN ACTIVITY IS SPECIFIED
DIC> @@ THAT IS LOW ENOUGH NOT TO FORM CORUNDUM (FE2O3). ALSO SPECIFY THAT
DIC> @@ THERE IS NO FLUX OF Fe ACROSS THIS INTERFACE, I.E. NO Fe IS
DIC> @@ ALLOWED TO ENTER OR LEAVE THE SYSTEM.
DIC> @@
DIC> set-cond boundary upper gas
TYPE OF CONDITION FOR COMPONENT FE /ZERO_FLUX/: zero-flux
TYPE OF CONDITION FOR COMPONENT O /ZERO_FLUX/: act
LOW TIME LIMIT /0/: 0 4.5e-4; * N
DIC>
DIC>
DIC> @@
DIC> @@ ENTER START VALUES FOR THE INITIAL INTERFACE VELOCITIES
DIC> @@
DIC> s-a-s-v -1e-5 1e-5 yes
STARTING VALUES WILL BE TAKEN FROM PROFILES
DIC>
DIC>
DIC> @@
DIC> @@ SIMULATE FOR 24 HOURS
DIC> @@
DIC> s-s-time 86400,,
SMALLEST ACCEPTABLE TIMESTEP : /1E-07/:
DIC>
DIC> @@

```

```
DIC> @@ SPECIFY THAT POTENTIALS AND NOT ACTIVITIES ARE VARIED AT THE PHASE
DIC> @@ INTERFACE. ALSO USE A FULLY IMPLICIT SCHEME FOR TIME INTEGRATION.
DIC> @@
DIC> s-s-c 0 1 1 YES POT YES YES 1 2.....,,
RELEASING OLD STARTING VALUES
DIC>
DIC>
DIC> @@ ENABLE THE GRAIN BOUNDARY DIFFUSION CONTRIBUTION MODEL
DIC> GB
REGION NAME : /SP/: SP
PHASE NAME: /SPINEL/: SPINEL
Enable model for grainboundary contribution to diffusion /YES/: YES
Grainboundary thickness /5E-10/: 5e-10
Grainsize(T,P,TIME)= 10.0e-6;
Bulkdiffusion activation energy multiplier /.5/: 0.333333
Enable model for dislocation contribution to diffusion /YES/: NO
DIC>
DIC>
DIC>
DIC> @@
DIC> @@ SAVE THE SET UP TO A NEW STORE FILE AND EXIT
DIC> @@
DIC> save exi3b.DIC Y
DIC>
DIC>
DIC>
DIC> set-inter
--OK---
DIC>
```

exi3b-run

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exi3b\run.DCM.test"
DIC>
DIC>
DIC> @@ exi3_run.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR RUNNING EXAMPLE i3
DIC> @@
DIC>
DIC> @@
DIC> @@ ENTER THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 0.00000E+00
DIC> read exi3b
OK
DIC>
DIC> @@
DIC> @@ START THE SIMULATION
DIC> @@
DIC> sim y
Region: FER
single geometric dense at 0.50000E-02
0.87701      95
Region: SP
double geometric
dense at outer boundaries, coarse at 0.50000E-10
lower part 1.2500      9
upper part 0.80000      9
Trying old scheme
U-FRACTION IN SYSTEM: FE = 0.99999999500542 O = 2.10000395011622E-08
TOTAL SIZE OF SYSTEM: .0049999901 [m]
U-FRACTION IN SYSTEM: FE = 0.99999999500542 O = 2.10000395011622E-08
TOTAL SIZE OF SYSTEM: .0049999901 [m]
6.141047161402763E-002   6.141016485504380E-002   6.141020906836839E-002   8.444643043596949E-003   1.457879775798498E-
024 TIME = 0.1000000E-06 DT = 0.1000000E-06 SUM OF SQUARES = 0.14578798E-23
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.16337999E-03 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49999900E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.29859730E-03 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.49999901E-02
U-FRACTION IN SYSTEM: FE = 0.99999999230859 O = 2.69779084025034E-08
TOTAL SIZE OF SYSTEM: .0049999011352 [m]
9 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: FER

CPU time used in timestep          0 seconds
8.109452882793873E-006   8.108765481852460E-006   8.108714358263970E-006   3.429203689632118E-007   3.460921124583648E-
030 TIME = 0.3000000E-06 DT = 0.2000000E-06 SUM OF SQUARES = 0.34609211E-29
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.37478240E-03 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49999989E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.57006118E-03 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.49999902E-02
U-FRACTION IN SYSTEM: FE = 0.99999998461782 O = 4.97800698647711E-08
TOTAL SIZE OF SYSTEM: .0049999015258 [m]
CPU time used in timestep          0 seconds
8.244220084356107E-006   8.245624820001861E-006   8.245834323740700E-006   5.265259157283041E-009   3.040545442780437E-
031 TIME = 0.7000000E-06 DT = 0.4000000E-06 SUM OF SQUARES = 0.30405454E-30
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.17607366E-03 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49999989E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.24106846E-03 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.49999902E-02
U-FRACTION IN SYSTEM: FE = 0.9999997890226 O = 6.90668859329386E-08
TOTAL SIZE OF SYSTEM: .0049999017858 [m]
CPU time used in timestep          1 seconds
1.134920301888734E-007   1.135894804039261E-007   1.135506444176019E-007   5.608707817274067E-010   5.589428618385944E-
033 TIME = 0.1500000E-05 DT = 0.8000000E-06 SUM OF SQUARES = 0.55894286E-32
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.14432292E-03 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49999989E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.20898651E-03 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.49999902E-02
U-FRACTION IN SYSTEM: FE = 0.99999996836327 O = 1.02507650482238E-07
TOTAL SIZE OF SYSTEM: .0049999023031 [m]
8 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep          0 seconds
4.495117154993389E-007   4.49645319495406E-007   4.496431998391493E-007   2.514741666006164E-010   2.513206922058051E-
034 TIME = 0.3100000E-05 DT = 0.1600000E-05 SUM OF SQUARES = 0.25132069E-33
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.94208025E-04 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49999986E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.13424763E-03 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.49999903E-02
U-FRACTION IN SYSTEM: FE = 0.9999999509026 O = 1.4547196415591E-07
TOTAL SIZE OF SYSTEM: .0049999029437 [m]
32 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep          0 seconds
1.153179467205265E-005   1.153635784462131E-005   1.153592952376297E-005   1.797713989031012E-008   1.284911245787787E-
030 TIME = 0.6300000E-05 DT = 0.3200000E-05 SUM OF SQUARES = 0.12849112E-29
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.67615626E-04 AND 0.0000000

output ignored...

... output resumed

POSITION OF INTERFACE FER / SP IS 0.49333469E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.11009831E-08 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.50286092E-02
U-FRACTION IN SYSTEM: FE = .99418084422836 O = .0189442387175316
TOTAL SIZE OF SYSTEM: .00502860924784 [m]
21 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep          1 seconds
5.273117098378293E-007   5.281046129101103E-007   5.280427494561641E-007   7.076339562288592E-010   1.841670509320294E-
032 TIME = 56943.895 DT = 8640.0000 SUM OF SQUARES = 0.18416705E-31
CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.70247282E-09 AND 0.0000000
POSITION OF INTERFACE FER / SP IS 0.49272776E-02
CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.10045406E-08 AND 0.0000000
POSITION OF INTERFACE SP / gas interface IS 0.50312191E-02
```

U-FRACTION IN SYSTEM: FE = .99366131709631 O = .0206594865694811
 TOTAL SIZE OF SYSTEM: .00503121911345 [m]
 9 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 0 seconds
 2.946689604425387E-007 2.952129684690918E-007 2.951641712357007E-007 3.823709736797433E-010 1.813082486708296E-

034 TIME = 65583.895 DT = 8640.0000 SUM OF SQUARES = 0.18130825E-33
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.64894663E-09 AND 0.0000000
 POSITION OF INTERFACE FER / SP IS 0.49216707E-02
 CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.92920435E-09 AND 0.0000000
 POSITION OF INTERFACE SP / gas interface IS 0.50336405E-02
 U-FRACTION IN SYSTEM: FE = .993180429172029 O = .0222444823738337
 TOTAL SIZE OF SYSTEM: .00503364054012 [m]
 8 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 1 seconds
 1.646046081816719E-007 1.649792107009017E-007 1.649481519432754E-007 1.988144432406818E-010 1.123208362097712E-

033 TIME = 74223.895 DT = 8640.0000 SUM OF SQUARES = 0.11232084E-32
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.60654812E-09 AND 0.0000000
 POSITION OF INTERFACE FER / SP IS 0.49164301E-02
 CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.86919604E-09 AND 0.0000000
 POSITION OF INTERFACE SP / gas interface IS 0.50359098E-02
 U-FRACTION IN SYSTEM: FE = .992730593159095 O = .0237257189756388
 TOTAL SIZE OF SYSTEM: .00503590981819 [m]
 11 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 0 seconds
 1.056445882444572E-007 1.059232720122668E-007 1.059039690796196E-007 1.220606026900290E-010 5.059327685939125E-

033 TIME = 82863.895 DT = 8640.0000 SUM OF SQUARES = 0.50593277E-32
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.57100511E-09 AND 0.0000000
 POSITION OF INTERFACE FER / SP IS 0.49114966E-02
 CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.81838278E-09 AND 0.0000000
 POSITION OF INTERFACE SP / gas interface IS 0.50380472E-02
 U-FRACTION IN SYSTEM: FE = .992307543343385 O = .0251191391110372
 TOTAL SIZE OF SYSTEM: .00503804716129 [m]
 7 GRIDPOINT(S) REMOVED FROM CELL #1 REGION: SP

CPU time used in timestep 1 seconds
 1.518627609602451E-008 1.512781289831391E-008 1.530717153876908E-008 1.085155039390110E-011 1.301507345230437E-

034 TIME = 86400.000 DT = 3536.1047 SUM OF SQUARES = 0.13015073E-33
 CELL # 1 VELOCITY AT INTERFACE # 2 IS -0.56483678E-09 AND 0.0000000
 POSITION OF INTERFACE FER / SP IS 0.49094993E-02
 CELL # 1 VELOCITY AT INTERFACE # 3 IS 0.78548555E-09 AND 0.0000000
 POSITION OF INTERFACE SP / gas interface IS 0.50388274E-02
 U-FRACTION IN SYSTEM: FE = .992153350362077 O = .0256664807147801
 TOTAL SIZE OF SYSTEM: .00503882739841 [m]

MUST SAVE WORKSPACE ON FILE

WORKSPACE SAVED ON FILE

RECLAIMING WORKSPACE

DELETING TIME-RECORD FOR TIME 0.0000000
 DELETING TIME-RECORD FOR TIME 0.10000000E-06
 DELETING TIME-RECORD FOR TIME 0.30000000E-06
 DELETING TIME-RECORD FOR TIME 0.70000000E-06
 DELETING TIME-RECORD FOR TIME 0.15000000E-05
 DELETING TIME-RECORD FOR TIME 0.31000000E-05
 DELETING TIME-RECORD FOR TIME 0.63000000E-05
 DELETING TIME-RECORD FOR TIME 0.12700000E-04
 DELETING TIME-RECORD FOR TIME 0.25500000E-04
 DELETING TIME-RECORD FOR TIME 0.51100000E-04
 DELETING TIME-RECORD FOR TIME 0.10230000E-03
 DELETING TIME-RECORD FOR TIME 0.20470000E-03
 DELETING TIME-RECORD FOR TIME 0.40950000E-03
 DELETING TIME-RECORD FOR TIME 0.81910000E-03
 DELETING TIME-RECORD FOR TIME 0.16383000E-02
 DELETING TIME-RECORD FOR TIME 0.32767000E-02
 DELETING TIME-RECORD FOR TIME 0.65535000E-02
 DELETING TIME-RECORD FOR TIME 0.13107100E-01
 DELETING TIME-RECORD FOR TIME 0.26214300E-01
 DELETING TIME-RECORD FOR TIME 0.52428700E-01
 DELETING TIME-RECORD FOR TIME 0.10485750
 DELETING TIME-RECORD FOR TIME 0.20971510
 DELETING TIME-RECORD FOR TIME 0.41943030
 DELETING TIME-RECORD FOR TIME 0.83886070
 DELETING TIME-RECORD FOR TIME 1.6777215
 DELETING TIME-RECORD FOR TIME 3.3554431
 DELETING TIME-RECORD FOR TIME 6.7108863
 DELETING TIME-RECORD FOR TIME 13.421773
 DELETING TIME-RECORD FOR TIME 26.843545
 DELETING TIME-RECORD FOR TIME 53.687091
 DELETING TIME-RECORD FOR TIME 107.37418
 DELETING TIME-RECORD FOR TIME 214.74836
 DELETING TIME-RECORD FOR TIME 429.49673
 DELETING TIME-RECORD FOR TIME 858.99346
 DELETING TIME-RECORD FOR TIME 1717.98669
 DELETING TIME-RECORD FOR TIME 3435.9738
 DELETING TIME-RECORD FOR TIME 6871.9477
 DELETING TIME-RECORD FOR TIME 13743.895
 DELETING TIME-RECORD FOR TIME 22383.895
 DELETING TIME-RECORD FOR TIME 31023.895
 DELETING TIME-RECORD FOR TIME 39663.895
 DELETING TIME-RECORD FOR TIME 48303.895
 DELETING TIME-RECORD FOR TIME 56943.895
 DELETING TIME-RECORD FOR TIME 65583.895
 DELETING TIME-RECORD FOR TIME 74223.895

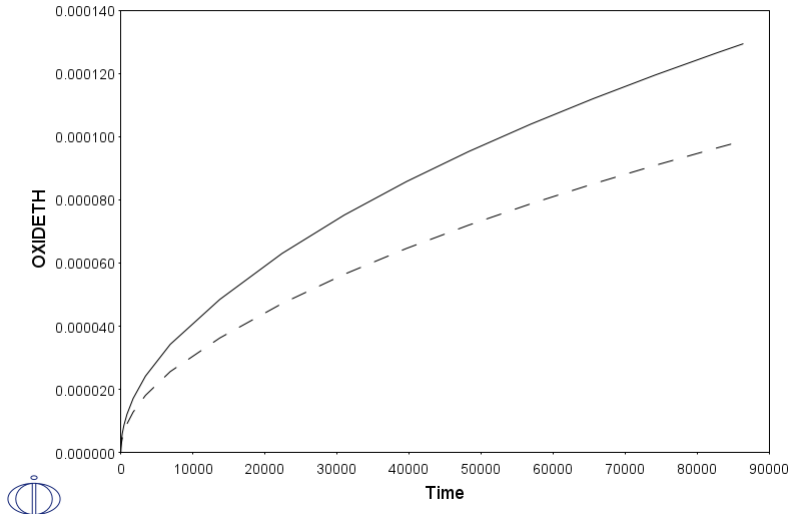
KEEPING TIME-RECORD FOR TIME 82863.895
 AND FOR TIME 86400.000
 WORKSPACE RECLAIMED

TIMESTEP AT 86400.0000 SELECTED

DIC>
DIC> set-inter
--OK--
DIC>

exi3b-plot

```
DIC>About
NO SUCH COMMAND, USE HELP
DIC>MACRO "c:\jenkins\workspace\dictra_generate_console_examples\examples\exi3b\plot.DCM.test"
DIC>
DIC> @@ exi3_plot.DCM
DIC>
DIC> @@
DIC> @@ FILE FOR GENERATING GRAPHICAL OUTPUT FOR EXAMPLE i3
DIC> @@
DIC>
DIC> @@
DIC> @@ GO TO THE DICTRA MONITOR AND READ THE STORE RESULT FILE
DIC> @@
DIC> go d-m
TIME STEP AT TIME 8.64000E+04
DIC> read exi3b
OK
DIC>
DIC> @@
DIC> @@ ENTER THE POST PROCESSOR
DIC> @@
DIC> post
POST PROCESSOR VERSION 1.7
Implemented by Bjorn Jonsson
POST-1:
POST-1: @@
POST-1: @@ PLOT THE THICKNESS OF THE OXIDE LAYER GROWING AT THE SURFACE.
POST-1: @@ FOR THIS WE NEED TO ENTER A FUNCTION ACCORDING TO THE FOLLOWING.
POST-1: ent func oxideth
FUNCTION: poi(sp,upper)-poi(sp,lower)
&
POST-1: @@
POST-1: @@ PUT THIS FUNCTION ON THE Y-AXIS
POST-1: @@
POST-1: s-d-a y oxideth
POST-1: s-d-a y
POST-1: @@
POST-1: @@ AND PLOT THE OXIDE THICKNESS VERSUS TIME
POST-1: @@
POST-1: s-d-a x time
INFO: Time is set as independent variable
POST-1:
POST-1: @@
POST-1: @@ SINCE WE ARE PLOTTING A FUNCTION, SPECIFY A PLOT CONDITION
POST-1: @@
POST-1: s-p-c interface sp upper
POST-1:
POST-1: app y exi3a.exp 0; 1;
POST-1:
POST-1:
POST-1: SET_EXP_FILE_FORMAT 4
POST-1: MAKE c:/jenkins/workspace/dictra_generate_console_examples/unite/distribution\macroResult.exp Y
OST-1: SET_EXP_FILE_FORMAT 10
POST-1:
POST-1: plot
2021.05.13.13.31.31
UPPER INTERFACE OF REGION "SP#1"
CELL #1
```



```
POST-1:
POST-1:
POST-1: set-inter
--OK---
POST-1:
```

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