

**Thermo-Calc
Software**

TC-Prisma Online Training Course

Day 2 - October 22, 2025



Day 2: TC-PRISMA (Precipitation Module)

09:00	Yesterday's home assignment
09:10	Theoretical Background: Growth Models
09:40	Examples: Ni alloys
10:30	Q & A
10:45	Examples: Steel
11:20	Example: Steel, Para-equilibrium models
11:50	Q & A

Home assignment: Al - Zr Alloy

System	
Database package	ALDEMO + MALDEMO (or TCAL9+MOBAL8)
Elements	Al, Zr, Si, (Fe is optional if TCAL9 used)
Matrix phase	FCC_A1
Precipitate phase	Al ₃ Sc (= AL3X in TCAL9)
Conditions	
Composition	Al - 0.23 Zr – 0.05 Si (- 0.1 Fe) (wt.%)
Temperature	650 K
Simulation time	500 hours
Nucleation properties	Nucleation Site Type: Bulk
Data Parameters	
Interfacial Energy	Calculated
Molar Volume (Matrix):	Fcc_A1: from database
Molar Volume (Precipitate):	Al ₃ Sc: from database

Home assignment: Al - Zr Alloy



The aim of this assignment is not only to simulate the precipitation treatment of this alloy, but to compare its simulated hardness (HV) with experimental data from the paper.

Following the paper, it is not the most stable Al-Zr phase that forms during the first several hundred hours of heat treatment. The most stable phase is called AL3ZR_D023 in TCAL and will probably become stable after even longer time at high T. Instead another phase of the same chemistry, AL3X, precipitates first (this phase is called AL3SC in ALDEMO).

The calculation of hardness in TC-Prisma is based on the same principle as the calculation of Yield strength. The selection and setting of parameters is done in the Plot renderer, see next slide.

Use the experimental file "Souza_data.exp" for comparison with your result.

Precipitation hardening in dilute Al-Zr alloys

Pedro Henrique Lamarão Souza^{a,*}, Carlos Augusto Silva de Oliveira^a,
José Maria do Vale Quaresma^b

J MATER RES TECHNOL. 2018; **7**(1): 66–72.

Home assignment: Al - Zr Alloy

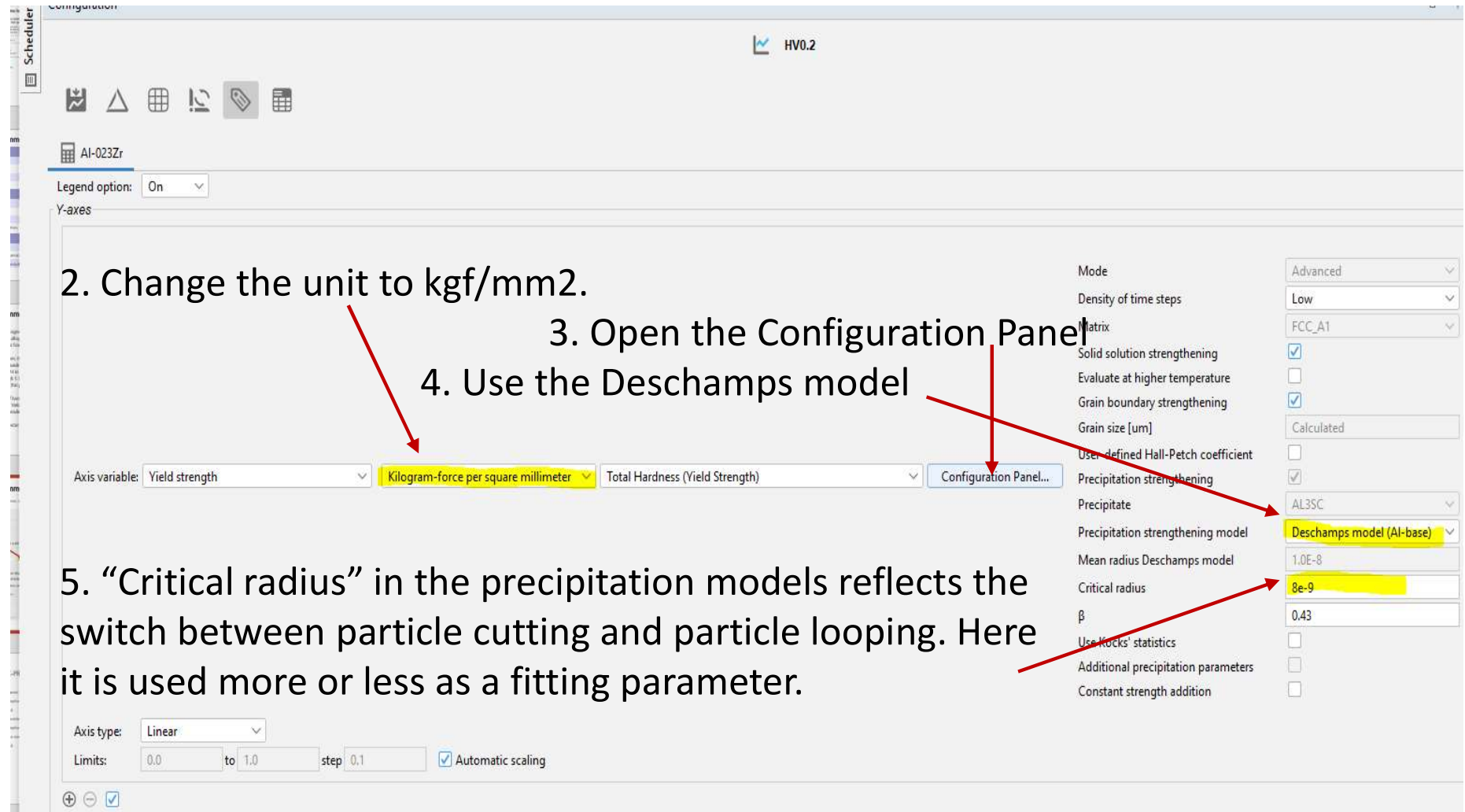
1. In the Plot renderer, select Yield Strength as Y-axis variable.

2. Change the unit to kgf/mm².

3. Open the Configuration Panel

4. Use the Deschamps model

5. “Critical radius” in the precipitation models reflects the switch between particle cutting and particle looping. Here it is used more or less as a fitting parameter.



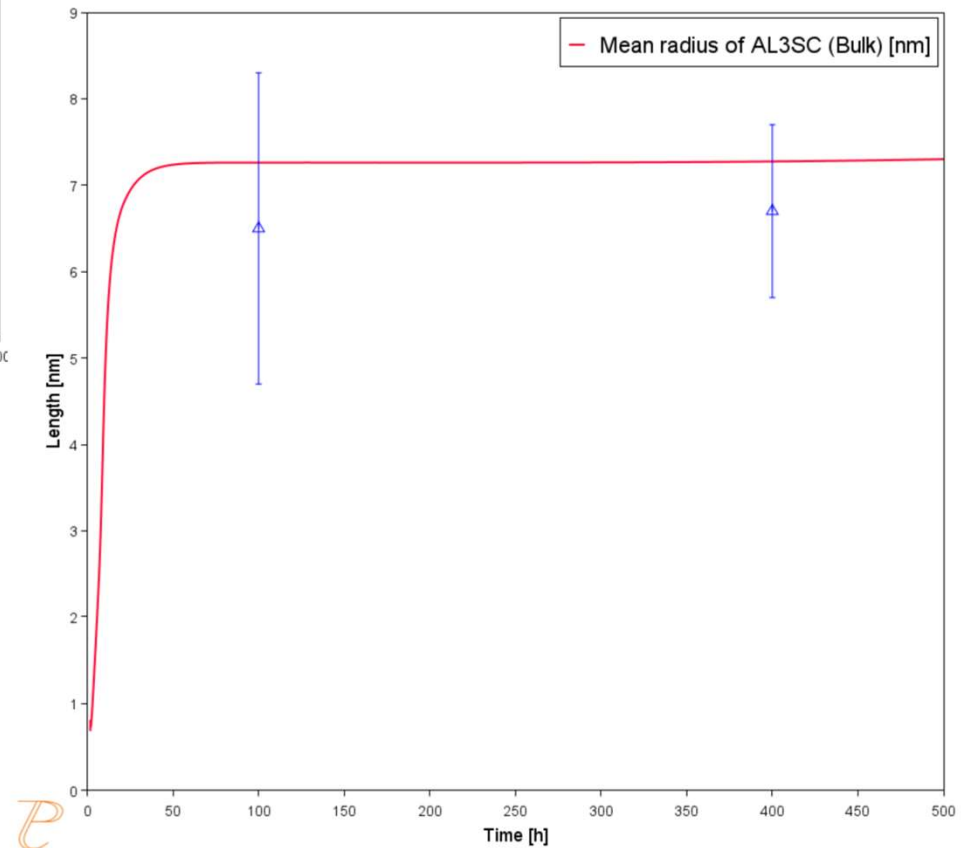
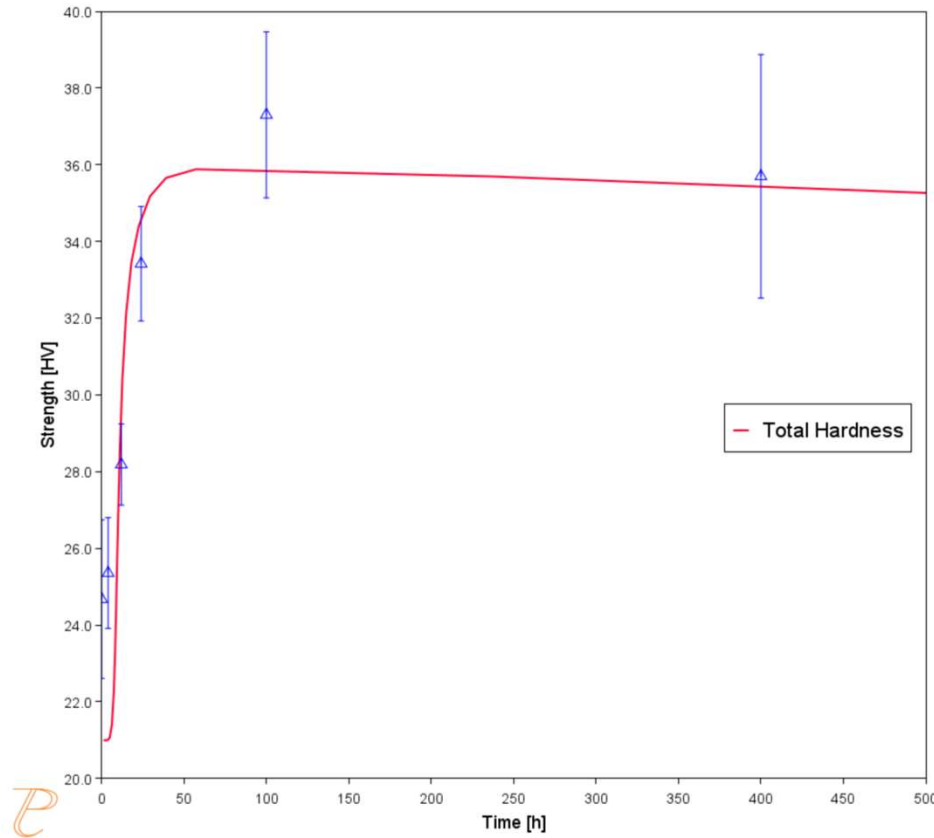
The screenshot shows the Thermo-Calc software interface. The main window displays the configuration for a plot of Yield strength (Y-axis variable) versus Total Hardness (Yield Strength). The unit is set to Kilogram-force per square millimeter. The Configuration Panel is open, showing various settings for the plot. The following settings are highlighted with red arrows:

- Axis variable: Yield strength
- Unit: Kilogram-force per square millimeter
- Configuration Panel...
- Mode: Advanced
- Density of time steps: Low
- Matrix: FCC_A1
- Solid solution strengthening: ☒
- Evaluate at higher temperature: ☐
- Grain boundary strengthening: ☒
- Grain size [um]: Calculated
- User defined Hall-Petch coefficient: ☐
- Precipitation strengthening: ☒
- Precipitate: AL3SC
- Precipitation strengthening model: Deschamps model (Al-base)
- Mean radius Deschamps model: 1.0E-8
- Critical radius: 8e-9
- β : 0.43
- Use Kocks' statistics: ☐
- Additional precipitation parameters: ☐
- Constant strength addition: ☐

At the bottom of the plot area, the following settings are visible:

- Axis type: Linear
- Limits: 0.0 to 1.0 step 0.1
- Automatic scaling: ☒

HA: Al - Zr Alloy - Results

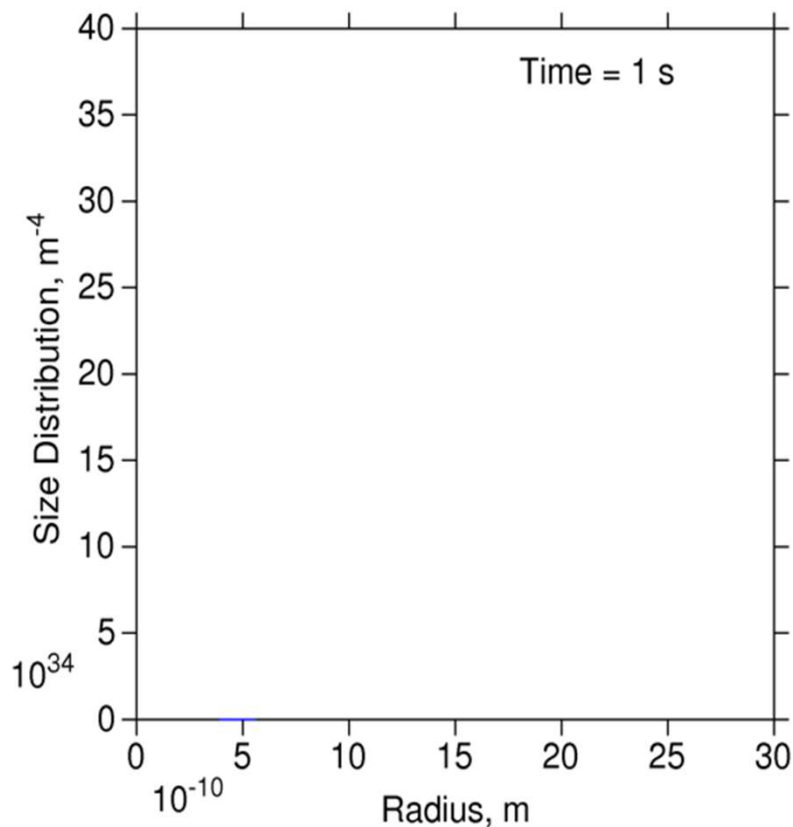


Theory: Growth

Models and Model Parameters

LS (Langer-Schwartz) and KWN (Kampmann and Wagner Numerical) Approach

$$J = \int_{r^*}^{\infty} j(r) dr$$



Continuity equation

$$\frac{\partial f(r,t)}{\partial t} = -\frac{\partial}{\partial r} [\nu(r)f(r,t)] + j(r,t)$$

$$C_0^\alpha = C^\alpha + (C^\beta - C^\alpha) \int_0^\infty \frac{4\pi}{3} f(r,t) r^3 dr$$

Mass balance

Models: Growth Rate

Available models

❑ Binary

$$v = \frac{c^\alpha - c^{\alpha/\beta}(r)}{c^\beta - c^{\alpha/\beta}(r)} \frac{D}{r}$$

H.B. Aaron, D. Fainstain, G. R. Kotler, J. Appl. Phys., 41(1970)4404

❑ Multi-Component Coarsening

$$v = \frac{2\sigma V_m^\alpha}{(\Delta C^{\alpha\beta})[M]^{-1}[\Delta C^{\alpha\beta})} \frac{1}{r} \left(\frac{1}{r^*} - \frac{1}{r} \right)$$

J.E. Morral, G.R. Purdy, Scripta Metall. Mater., 30(1994)905-908

❑ PrecipiCalc Model

$$v = \left(1 + r \sqrt{4\pi N_v r_a} \right) \frac{\Delta \bar{c}_i G_{ij}^\alpha \Delta c_j^\infty + \bar{c}_\delta^\beta (\bar{\mu}_\delta^\alpha - \bar{\mu}_\delta^\beta) - 2V_m^\beta \sigma / r}{r \Delta \bar{c}_i G_{ij}^\alpha D_{jk}^{-1} B_k + 1 / M}$$

H.J. Jou et al. Superalloy 2004, p.877-886

Available models

- Similarity-Supersaturation

$$v = \frac{S_1 \sqrt{D_1}}{2\sqrt{t}} = \frac{S_2 \sqrt{D_2}}{2\sqrt{t}} = \dots \quad S_i = f(\Omega_i)$$

T.Sourmail, Ph. D Thesis, Univ. Cambridge, 2002

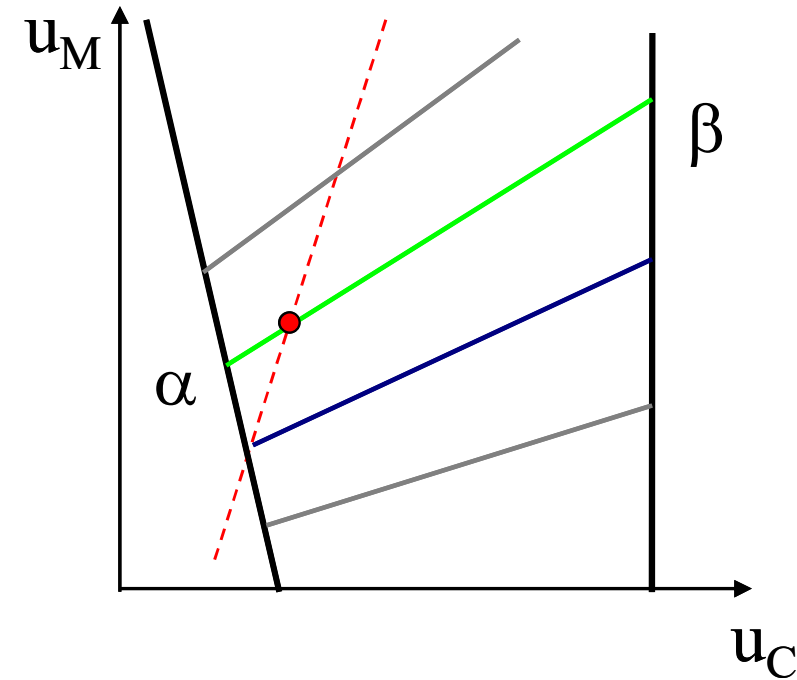
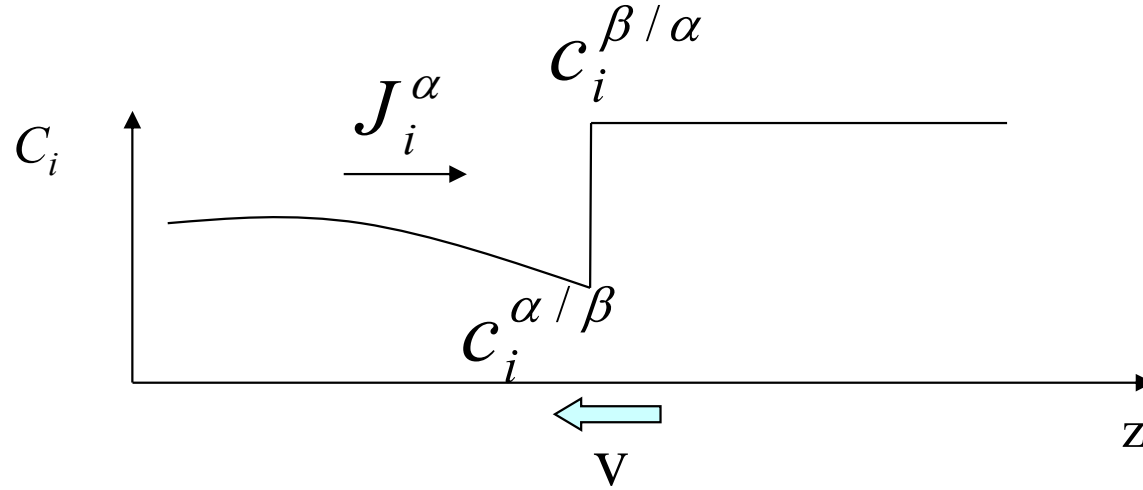
- Thermodynamic Extremum Principle

$$\dot{\rho}_k = \frac{F - (2\gamma_k/\rho_k)}{RT\rho_k} \left[\sum_{i=1}^n \frac{(c_{ki} - c_{0i})^2}{c_{0i}D_{0i}} \right]^{-1}$$

Svoboda J, Fischer FD, Fratzl P, Kozeschnik E. Mater Sci Eng A, 385(2004)166

Models: Growth Rate

- Local equilibrium at interface
- Flux balance equation



$$\mu_i^{\alpha/\beta} = \mu_i^{\beta/\alpha}$$

$$v(c_i^{\beta/\alpha} - c_i^{\alpha/\beta}) = -J_i^\alpha = \sum_j D_{ij}^\alpha \frac{\partial c_j^\alpha}{\partial z}$$

Models: Growth Rate

Advanced – Analytical Flux-balance Approximation

$$\mu_i^{\alpha/\beta} = \mu_i^{\beta/\alpha} + \frac{2\sigma V_m^\beta}{r}$$

Cross diffusion

$$v(c_i^{\beta/\alpha} - c_i^{\alpha/\beta}) = c_i^{\alpha/\beta} M_i (\mu_i^\alpha - \mu_i^{\alpha/\beta}) / \xi_i r$$

High supersaturation

General model – new in TC 2019a

$$v = \frac{K}{r} \left(\Delta G_m - \frac{2\sigma V_m}{r} \right)$$

Models: Growth Rate

General & Simplified –
different expressions for K

$$v = \frac{K}{r} \left(\Delta G_m - \frac{2\sigma V_m}{r} \right)$$

Simplified

Similar atomic
mobilities

$$K_{sphere}^{simplified} = \left[\sum_i \frac{\left(X_i^{\beta/\alpha}(r) - X_i^{\alpha/\beta}(r) \right)^2 \xi_i}{X_i^{\alpha/\beta}(r) M_i} \right]^{-1}$$

Number fixed
Coarsening

General

Very different
atomic
mobilities

$$K_{sphere}^{Morrall-Purdy} = \left[\left(\Delta X^{\alpha\beta} \right) [M]^{-1} \left(\Delta X^{\alpha\beta} \right) \right]^{-1}$$

Volume fixed
Coarsening

General

Very different
atomic
mobilities

$$K_{sphere}^{general} = \left[\left(\Delta X^{\alpha\beta} \right) [\ddot{G}] [\bar{D}]^{-1} \left(\Delta X^{\alpha\beta} \right) \right]^{-1}$$

Volume fixed
Growth

Para-equilibrium and Non-partitioning local equilibrium

$$v = \frac{K^{PE/NPLE}}{r} \left(\Delta G_m^{PE/NPLE} - \frac{2\sigma V_m}{r} \right)$$

$$K_{sphere}^{PE/NPLE} = \left[\sum_i \frac{\left(u_C^{\beta/\alpha}(r) - u_C^{\alpha/\beta}(r) \right)^2 \xi_C}{u_C^{\alpha/\beta}(r) M_C} \right]^{-1}$$

$$\beta^* = \frac{4\pi r^{*2} K}{a^4}$$

Models: NPLE and Para-eq.

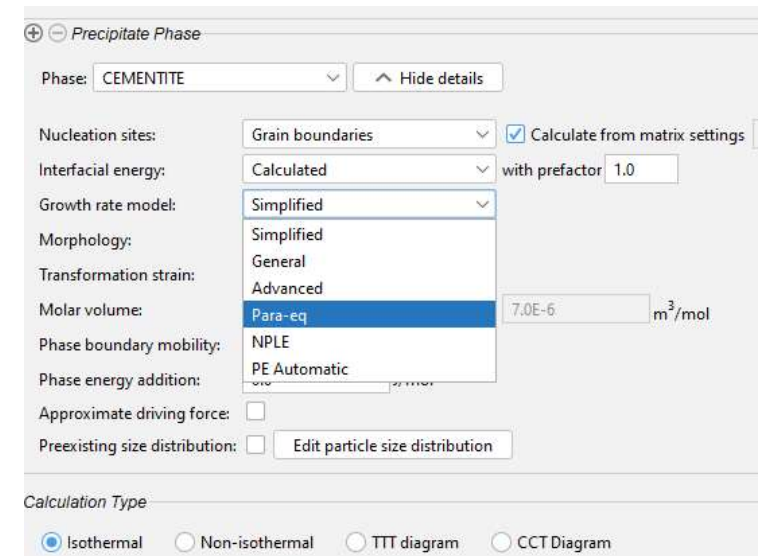


Models to handle NPLE (non partitioning local equilibrium) and para-equilibrium were introduced several years ago.

These models deal with fast diffusion processes and are additions to the Simplified growth rate model.

These models need only to consider the movement of the fast diffusing specie, usually an interstitial such as C or N.

The new (2023) **PE Automatic** model enables a smooth transition from Paraequilibrium growth rate model to Simplified growth rate model. The rate of transition process is dependent on the relative differences in diffusion between C and substitutional elements, as well as the differences in driving force between PE and Ortho-Equilibrium (i.e. Local Eq.).



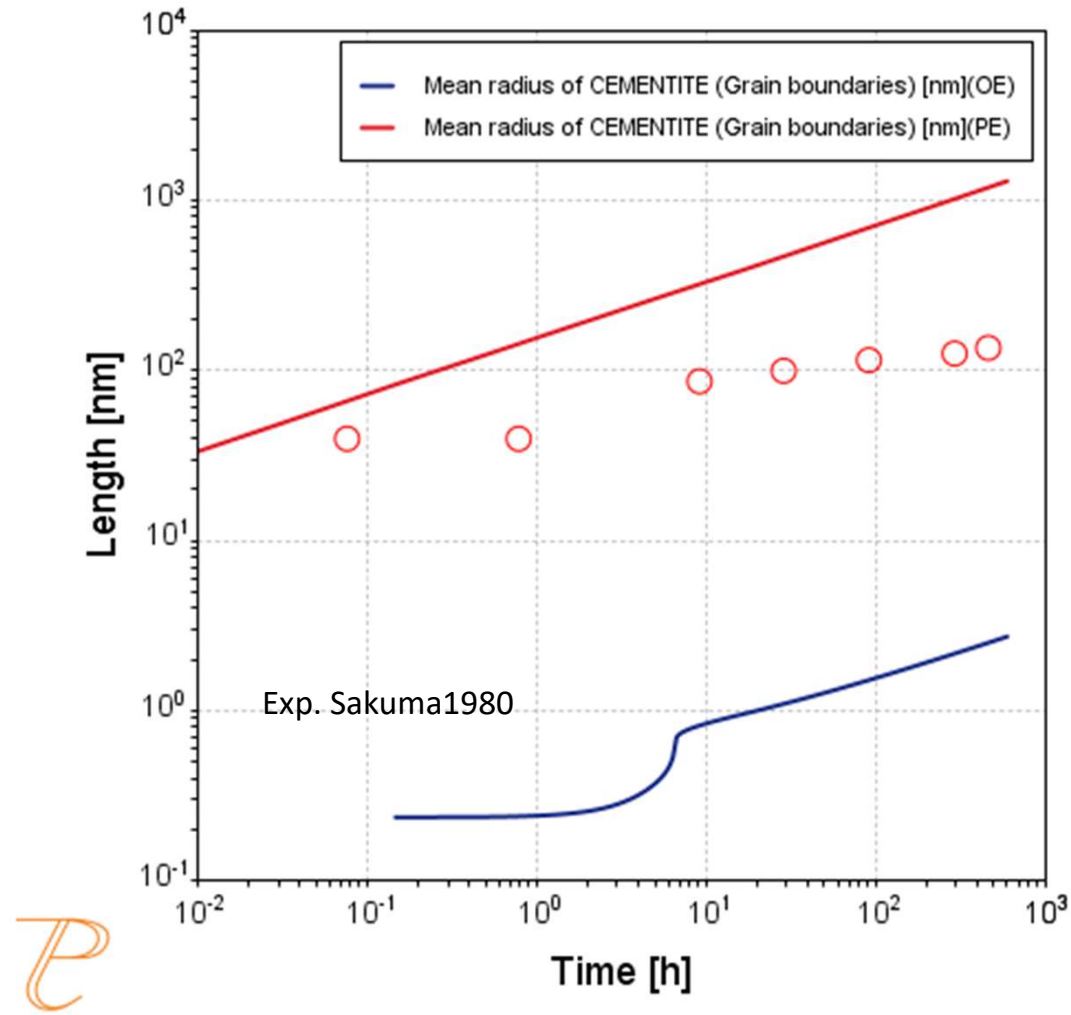
The non-zero volume correction

$$\nu' = \nu \left(1 + r \sqrt{4\pi N_V} \langle r \rangle \right)$$

Chen MK, Voorhees PW, Modeling and simulation in materials science and engineering 1993;1:591-612.

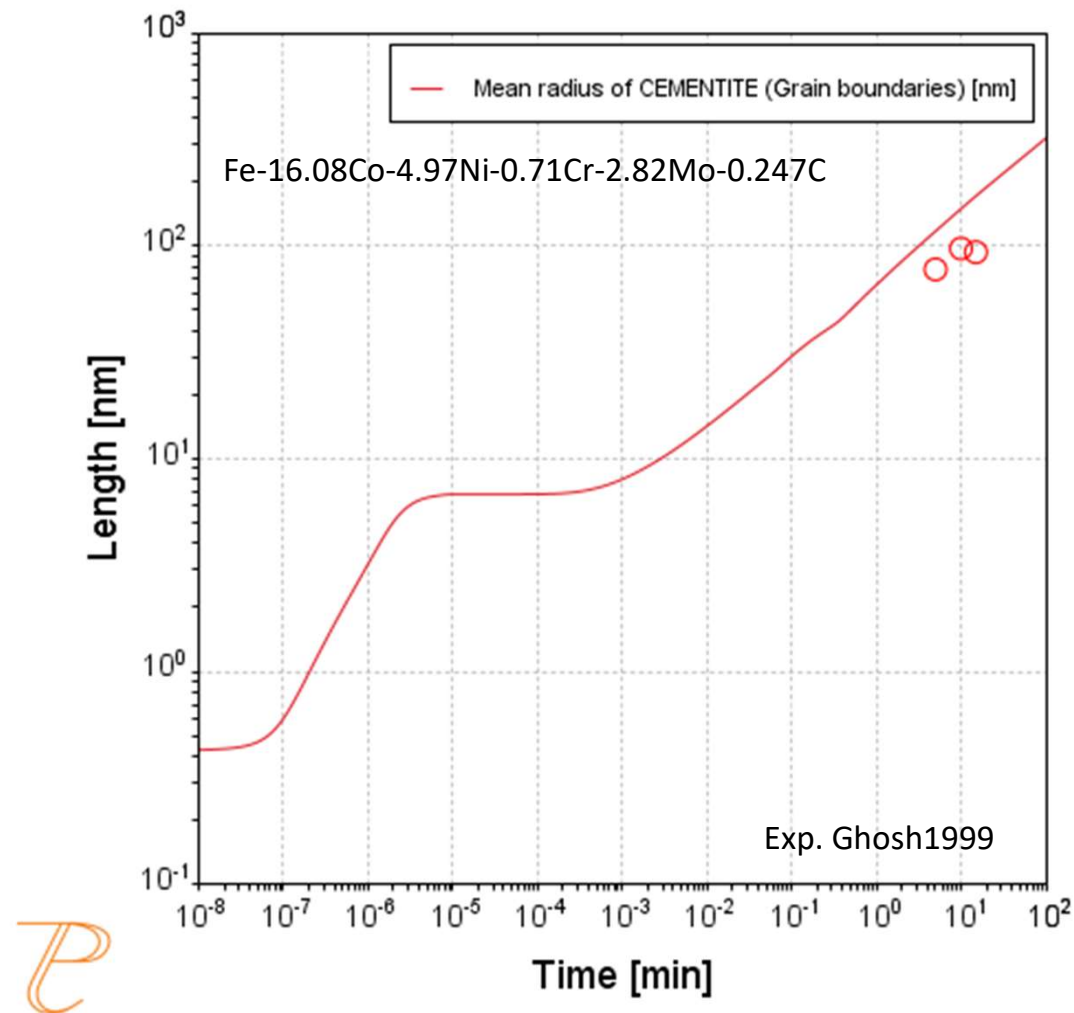
Ortho-eq vs Para-eq

Fe-0.26C-0.11Cr at T= 773 K



Ortho-eq vs Para-eq

Ultra-high-strength steel at $T = 783 \text{ K}$



P

Precipitate Shapes

➤ Interfacial Energy Anisotropy*

$$\frac{\sigma_l}{\sigma_r} = \frac{l}{r} = \alpha$$

➤ Elastic Strain Energy

- Elastically Isotropic or Cubic Systems
- First Approximation: Elastically Homogenous
- Eshelby's Theory**

➤ Particle Shape

- Determined by Minimization of Combined Interfacial Energy and Elastic Energy
- User-Defined, Fixed Value

Needle (Prolate Spheroid)

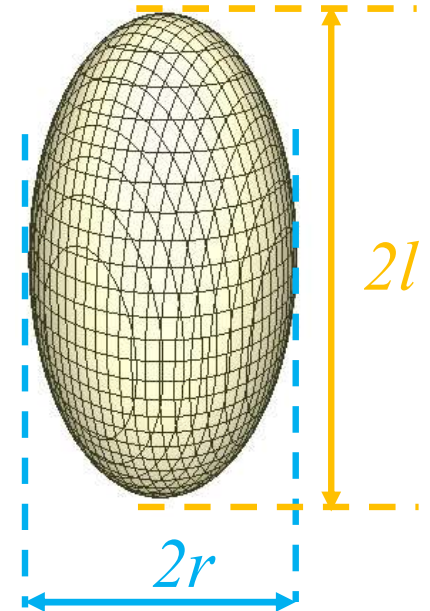
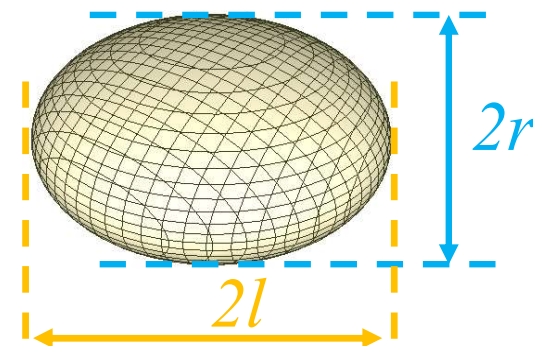


Plate (Oblate Spheroid)



* C.A. Johnson, *Surf. Sci.* 3(1965)429

** J.D. Eshelby, *Pro. Roy. Soc. A*, 241(1957)376

Effect on Growth Rate

K. Wu, Q. Chen, P. Mason, J Phase Eq. Diffus. 39(2018)571-583.

Needle***

(Prolate Spheroid)

$$\frac{dR}{dt} = K_{\sigma} \cdot K_{\text{shp}} \cdot \left(\frac{dR}{dt} \right)_{\text{sph}}$$

R : Radius of Equivalent Sphere

➤ Interfacial energy anisotropy*

- Generalized Gibbs-Thomson Effect

$$\mu(R) - \mu(\infty) = K_{\sigma} \frac{2\sigma_{\text{ch}}^{\text{sp}} V_m}{R}$$

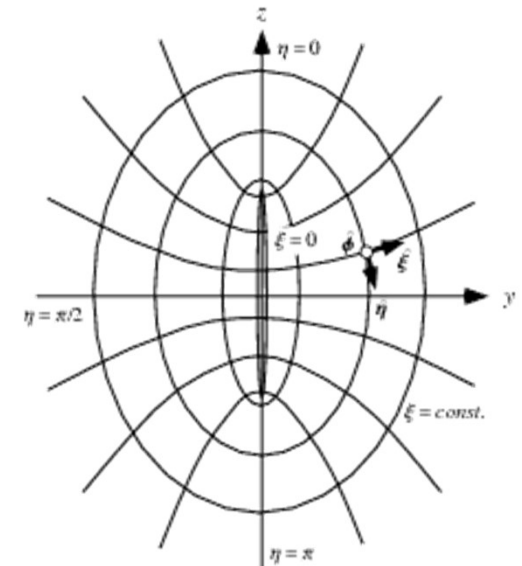
➤ Shape Effect

- Assumption of Shape Conserving Concentration Field**

* C.A. Johnson, *Surf. Sci.* 3(1965)429

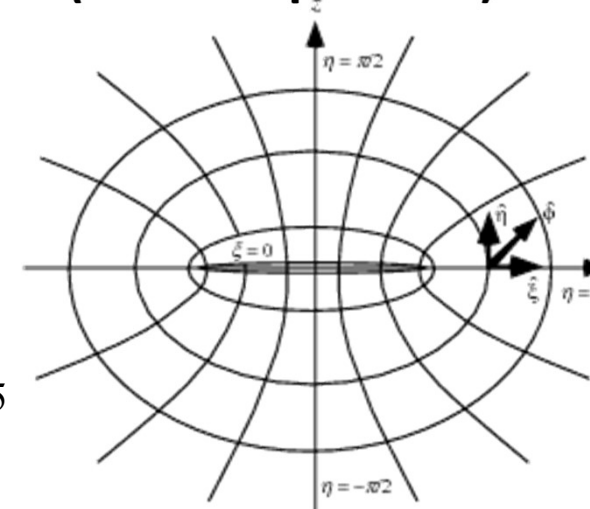
** F.S. Ham, *Quart. Appl. Math.*, 17(1959)137; *J Phys. Chem. Solids*, 6(1958)335

*** [http://http://mathworld.wolfram.com](http://mathworld.wolfram.com)



Plate***

(Oblate Spheroid)



Effect on Growth Rate

K. Wu, Q. Chen, P. Mason, J Phase Eq. Diffus. 39(2018)571-583.

$$\frac{dR}{dt} = K_{\sigma} \cdot K_{shp} \cdot \left(\frac{dR}{dt} \right)_{sph}$$

R : Radius of Equivalent Sphere

Aspect ratio

$$\alpha = \frac{l}{r} \geq 1$$

Eccentricity

$$e = \sqrt{1 - \frac{1}{\alpha^2}}$$

Needle

$$K_{\sigma} = \sqrt[3]{\alpha}$$

$$K_{shp} = \frac{2\sqrt[3]{\alpha^2}e}{\ln(1+e) - \ln(1-e)}$$

α	$K_{\sigma}K_{shp}$
3.0	1.6
10.0	3.3
15.0	4.4
20.0	5.4

Plate

$$K_{\sigma} = \sqrt[3]{\alpha^2}$$

$$K_{shp} = \frac{e\sqrt[3]{\alpha}}{\arccos(0) - \arccos(e)}$$

α	$K_{\sigma}K_{shp}$
10.0	6.8
15.0	10.0
20.0	13.1

Examples

Ni alloys



Available online at www.sciencedirect.com



Acta Materialia 56 (2008) 448–463



www.elsevier.com/locate/actamat

Effects of a tungsten addition on the morphological evolution, spatial correlations and temporal evolution of a model Ni–Al–Cr superalloy

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^c Department of Materials Science and Engineering, Massachusetts Institute of Technology, 77 Massachusetts Avenue, Cambridge, MA 02139, USA

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Received 8 July 2007; received in revised form 26 September 2007; accepted 28 September 2007

Available online 26 November 2007

Abstract

The effect of adding 2 at.% W to a model Ni–Al–Cr superalloy on the morphological evolution, spatial correlations and temporal evolution of $\gamma'(L1_2)$ -precipitates at 1073 K is studied with scanning electron microscopy and atomic force microscopy. Adding W yields a larger microhardness, earlier onset of spheroidal-to-cuboidal precipitate morphological transition, larger volume fraction (from ~20% to 30%), reduction in coarsening kinetics by one-third and a larger number density (N_v) of smaller mean radii ($\langle R \rangle$) precipitates. The kinetics of $\langle R \rangle$ and interfacial area per unit volume obey $t^{1/3}$ and $t^{-1/3}$ relationships, respectively, which is consistent with coarsening driven by interfacial energy reduction. The N_v power-law dependencies deviate, however, from model predictions, indicating that a stationary state is not achieved. Quantitative analyses with precipitate size distributions, pair correlation functions and edge-to-edge inter-precipitate distance distributions give insight into two-dimensional microstructural evolution, including the elastically driven transition from a uniform γ' -distribution to one-dimensional $\langle 001 \rangle$ -strings to eventually clustered packs of γ' -precipitates in the less densely packed Ni–Al–Cr alloy.

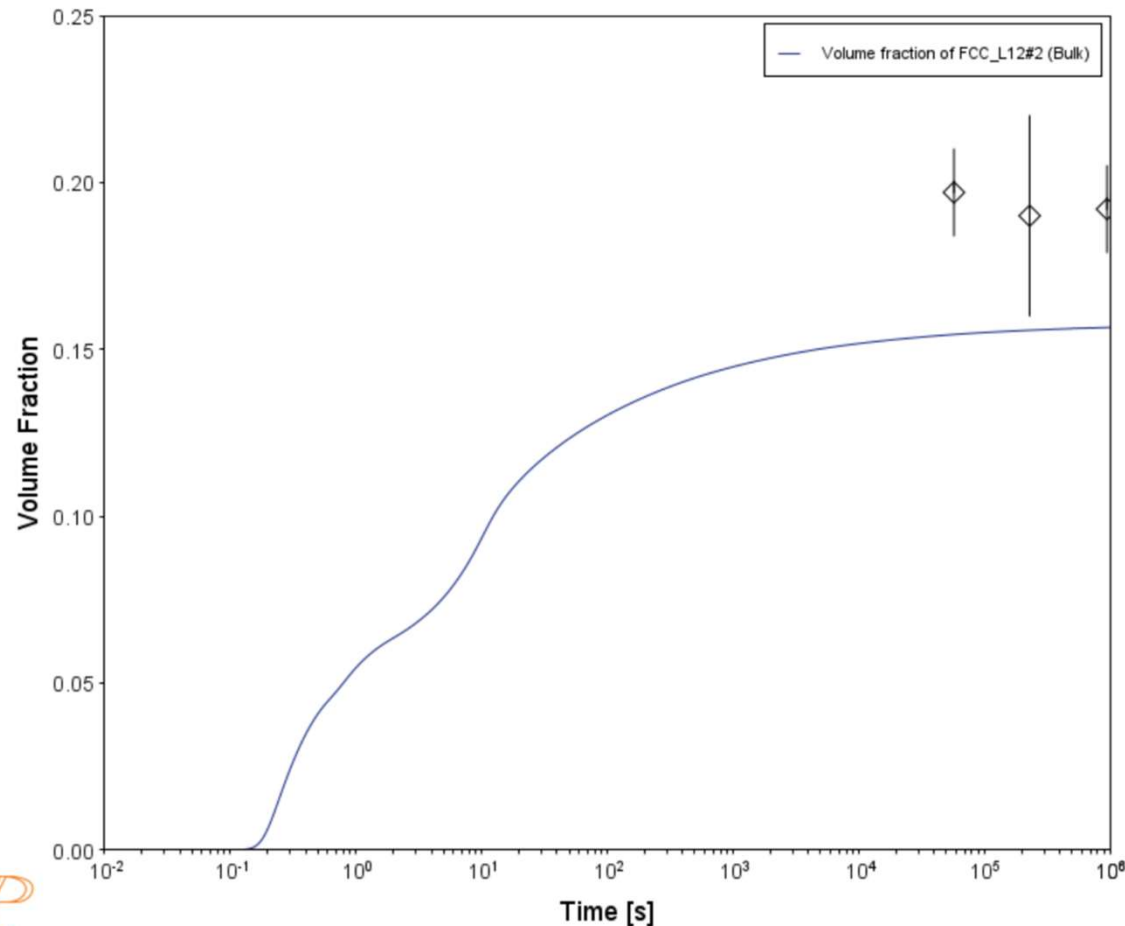
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Ni-alloy Example 1

System	
Database package	TCNI13 + MOBNi6
Elements	Ni, Al, Cr
Matrix phase	DIS_FCC_A1
Precipitate phases	FCC_L12#2
Conditions	
Composition	Ni – 9.8 Al – 8.3 Cr (at.%)
Temperature	800 °C
Simulation time	1E6 s
Nucleation properties	Nucleation Site Type: Bulk
Data Parameters – Interfacial Energies	
Interfacial Energy	Calculated
Molar Volume (Matrix phase):	DIS_FCC_A1: from database
Molar Volume (Precipitate phase):	FCC_L12#2: from database

Ni-alloy Example 1

Strange result for Volume fraction compared with experimental data from Sudbrack when the setup on previous page is used:



Results

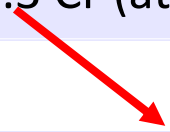
Volume fraction	Number density	Mean radius	Table renderer 1	
System				
Moles	1.00000			
Mass	55.02701	[g]		
Temperature	1073.15000	[K]		
Total Gibbs Energy	-66916.16735	[J]		
Enthalpy	-51528.57953	[J]		
Volume	6.97148E-6	[m³]		
Component	Mole Fraction	Mass Fraction	Activity	Potential
Al	0.09800	0.04805	8.82663E-10	-1.86021E5
Cr	0.08300	0.07843	0.00136	-58918.48387
Ni	0.81900	0.87352	0.00250	-53474.75936
Stable Phases				
	Moles	Mass	Volume Fraction	
DIS_FCC_A1#1	0.84221	46.67188	0.84210	Composition
Composition				
Component	Mole Fraction	Mass Fraction		
Ni	0.82880	0.87777		
Cr	0.08612	0.08081		
Al	0.08508	0.04142		
	Moles	Mass	Volume Fraction	
FCC_L12#2	0.15779	8.35514	0.15790	Composition
Composition				
Component	Mole Fraction	Mass Fraction		
Ni	0.76668	0.84977		
Al	0.16698	0.08509		
Cr	0.06634	0.06514		

In addition, the fit for mean radius as function of time, is not very good.

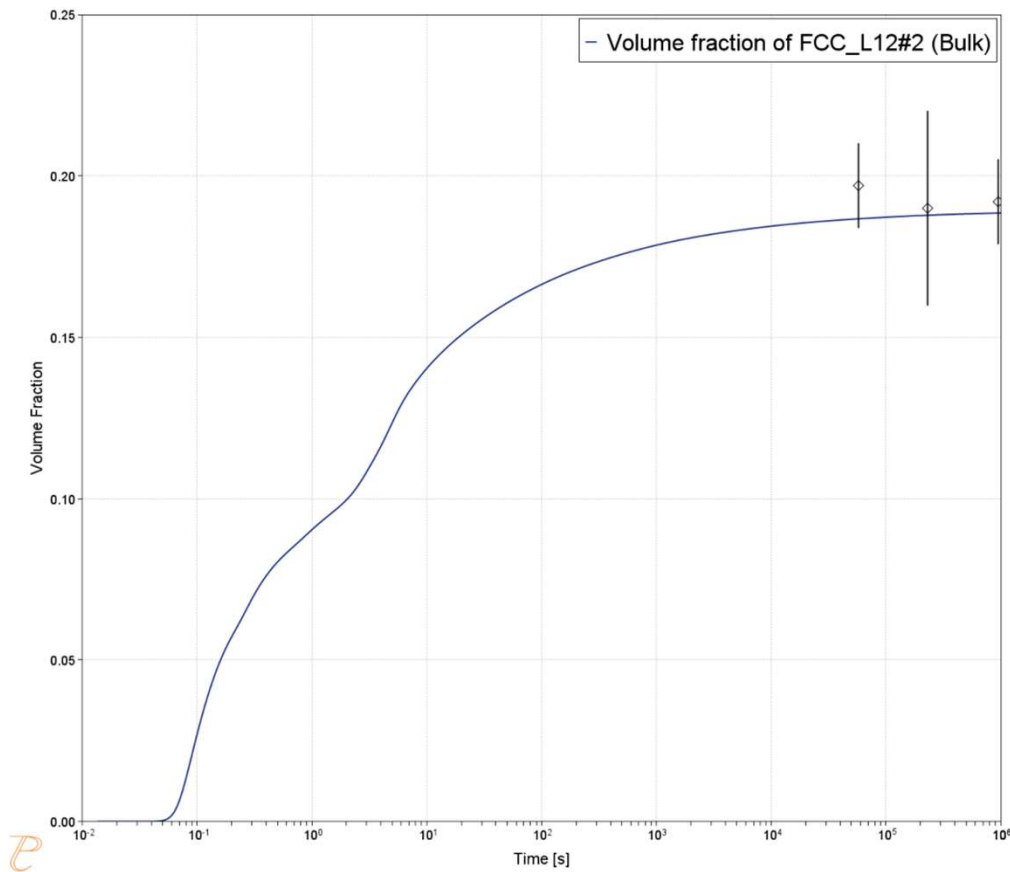
Ni-alloy Example 1

System	
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Elements	Ni, Al, Cr
Matrix phase	DIS_FCC_A1
Precipitate phases	FCC_L12#2
Conditions	
Composition	Ni – 9.8 Al – 8.3 Cr (at.%)
Temperature	800 °C
Simulation time	1E6 s
Nucleation properties	Nucleation Site Type: Bulk
Data Parameters – Interfacial Energies	
Interfacial Energy	Calculated
Molar Volume (Matrix phase):	DIS_FCC_A1: from database
Molar Volume (Precipitate phase):	FCC_L12#2: from database

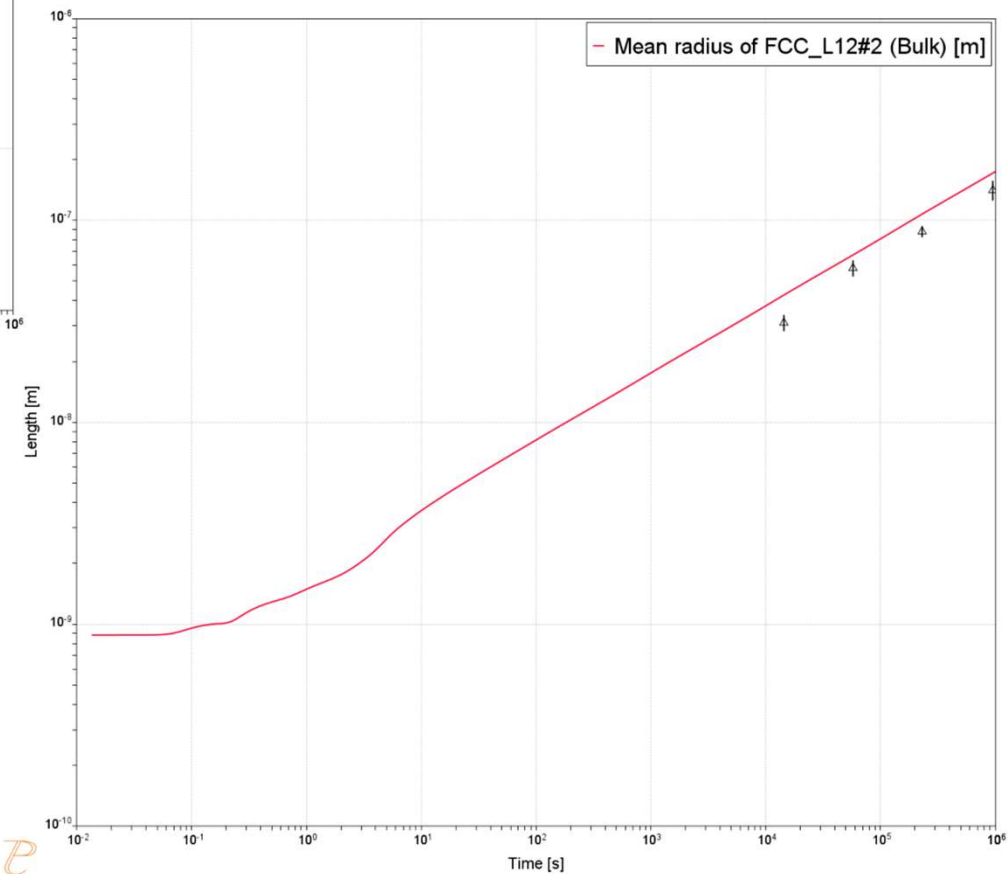
8.9 Cr



Ni-alloy Example 1



Using Ni – 9.8 Al – 8.9 Cr (at-%)



Influence of composition on monomodal versus multimodal γ' precipitation in Ni–Al–Cr alloys

T. Rojhirunsakool · S. Meher · J. Y. Hwang ·
S. Nag · J. Tiley · R. Banerjee

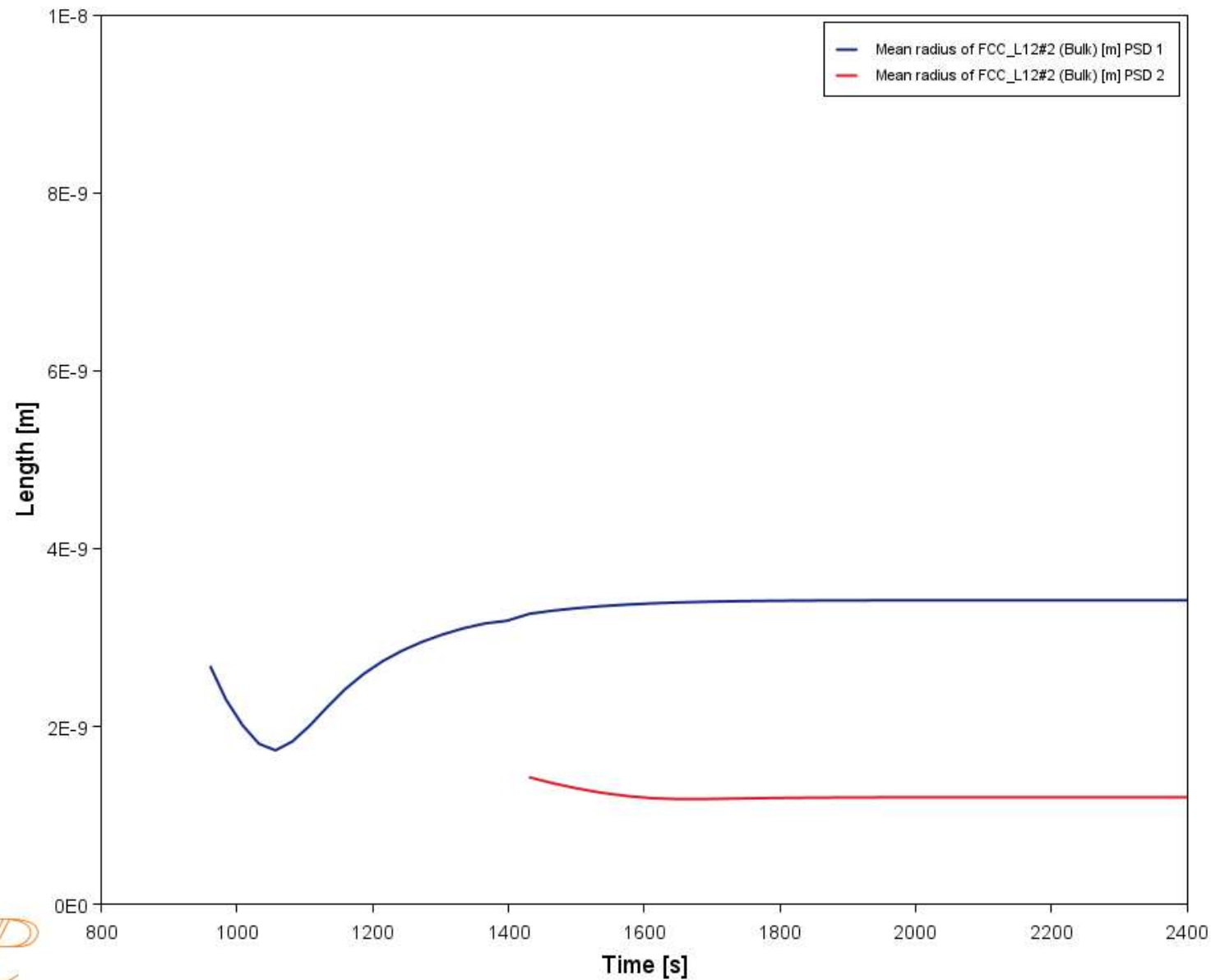
Abstract This study investigates the influence of alloy composition on γ' precipitation in Ni–8Al–8Cr and Ni–10Al–10Cr at.% during continuous cooling from a supersolvus temperature. When subjected to the same cooling rate, Ni–8Al–8Cr develops a monomodal population, whereas Ni–10Al–10Cr develops a multimodal (primarily bimodal) population of γ' precipitates. The bimodal γ' precipitate size distribution in Ni–10Al–10Cr alloy can be attributed to two successive nucleation bursts during continuous cooling while the monomodal γ' size distribution in Ni–8Al–8Cr results from a single nucleation burst followed by a longer time—wider temperature window for nucleation resulting in a larger number density of precipitates. Three-dimensional atom

Ni-alloy Example 2

System	
Database package	NIDEMO + MNIDEMO
Elements	Ni, Al, Cr
Matrix phase	DIS_FCC_A1
Precipitate phases	FCC_L12#2
Conditions	
Composition	Ni – 10 (8) Al – 10 (8) Cr (at.%)
Temperature	940 - 380 °C
Simulation time	2400 s
Nucleation properties	Nucleation Site Type: Bulk
Data Parameters – Interfacial Energies	
Interfacial Energy	Bulk: 0.023 J/m ²
Molar Volume (Matrix phase):	DIS_FCC_A1: from database
Molar Volume (Precipitate phase):	FCC_L12#2: from database
Mobility Adjustment Factor	1 (i.e. no change)

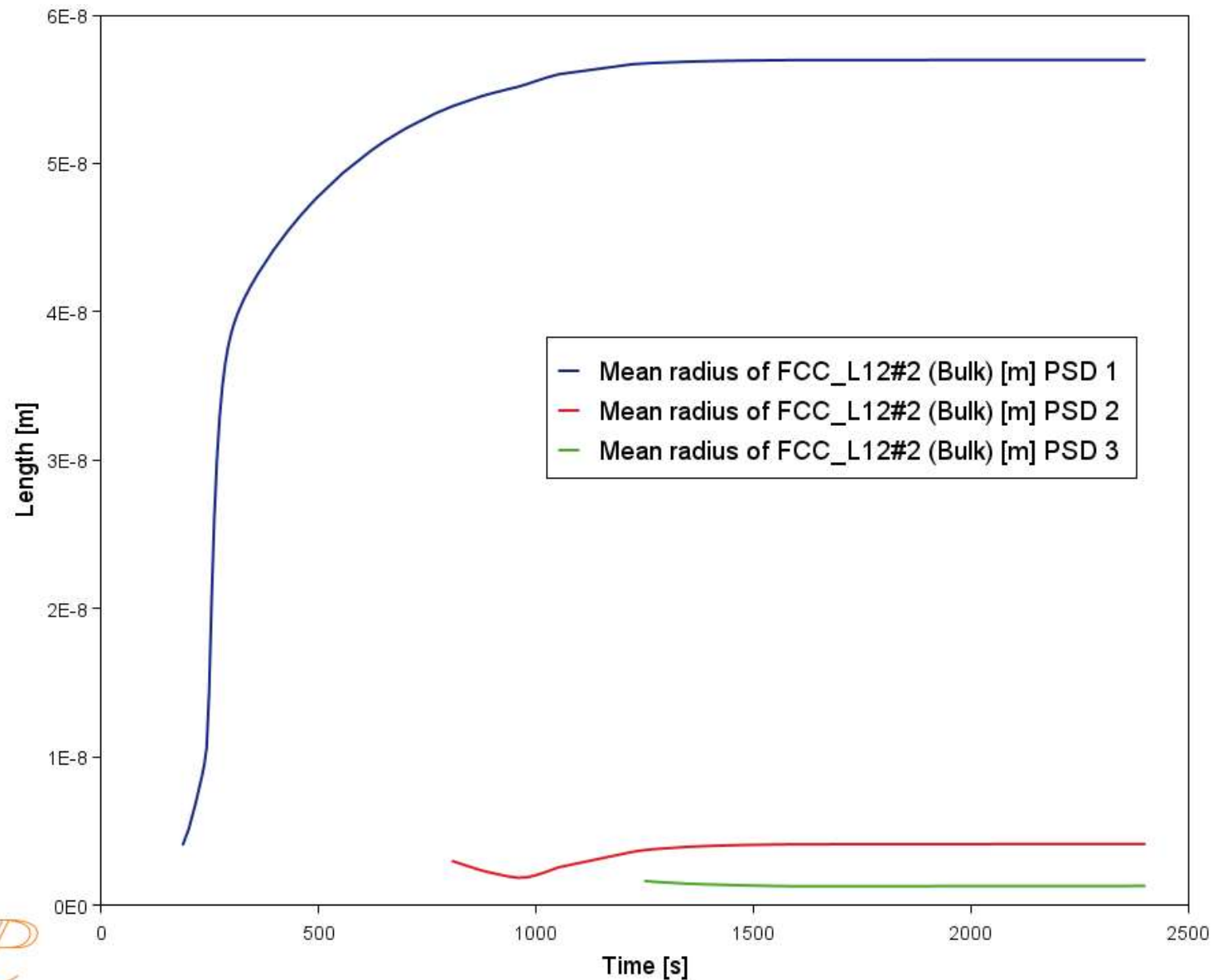
Ni-alloy Example 2

Ni – 8 Al – 8 Cr



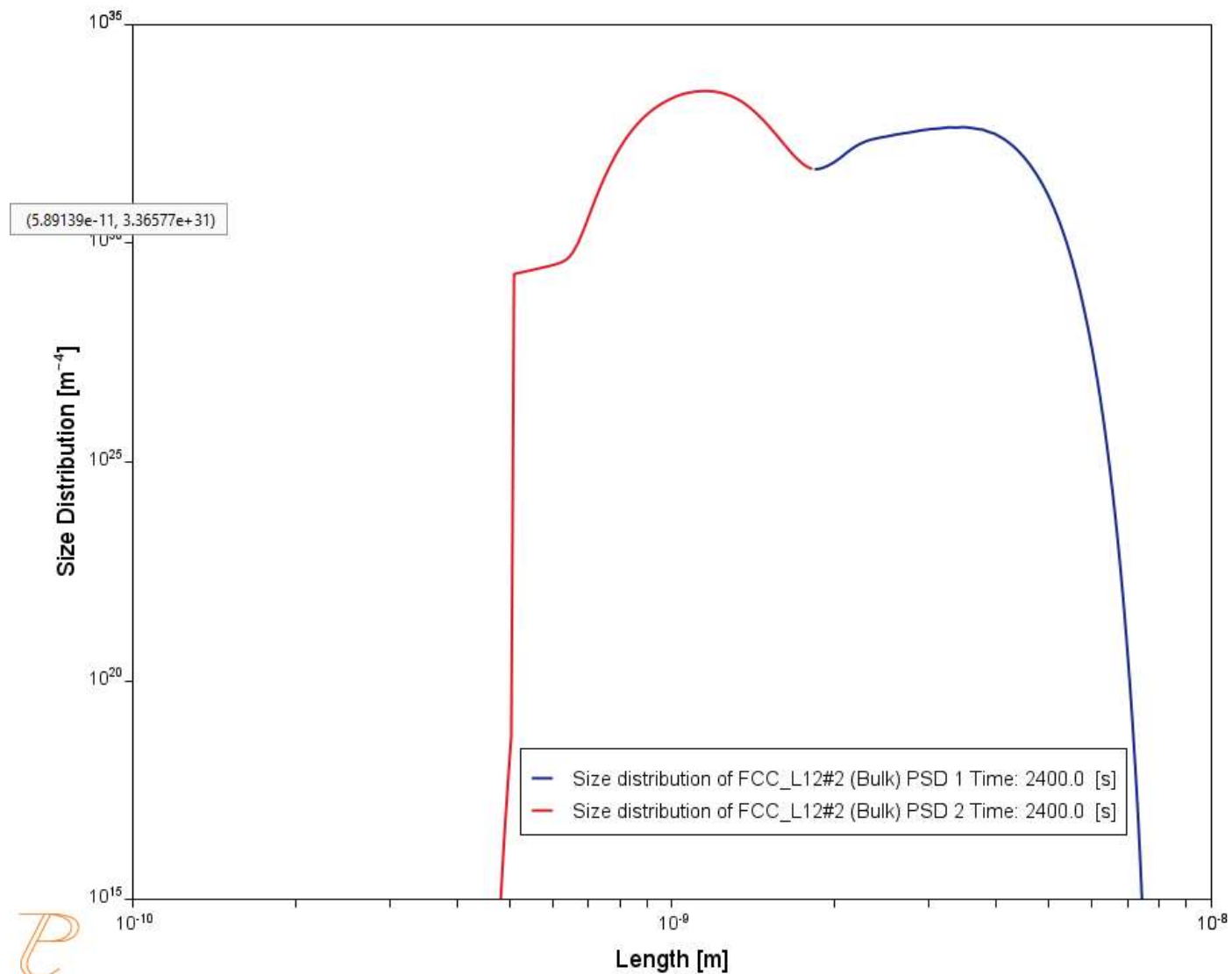
Ni-alloy Example 2

Ni – 10 Al – 10 Cr



Ni-alloy Example 2

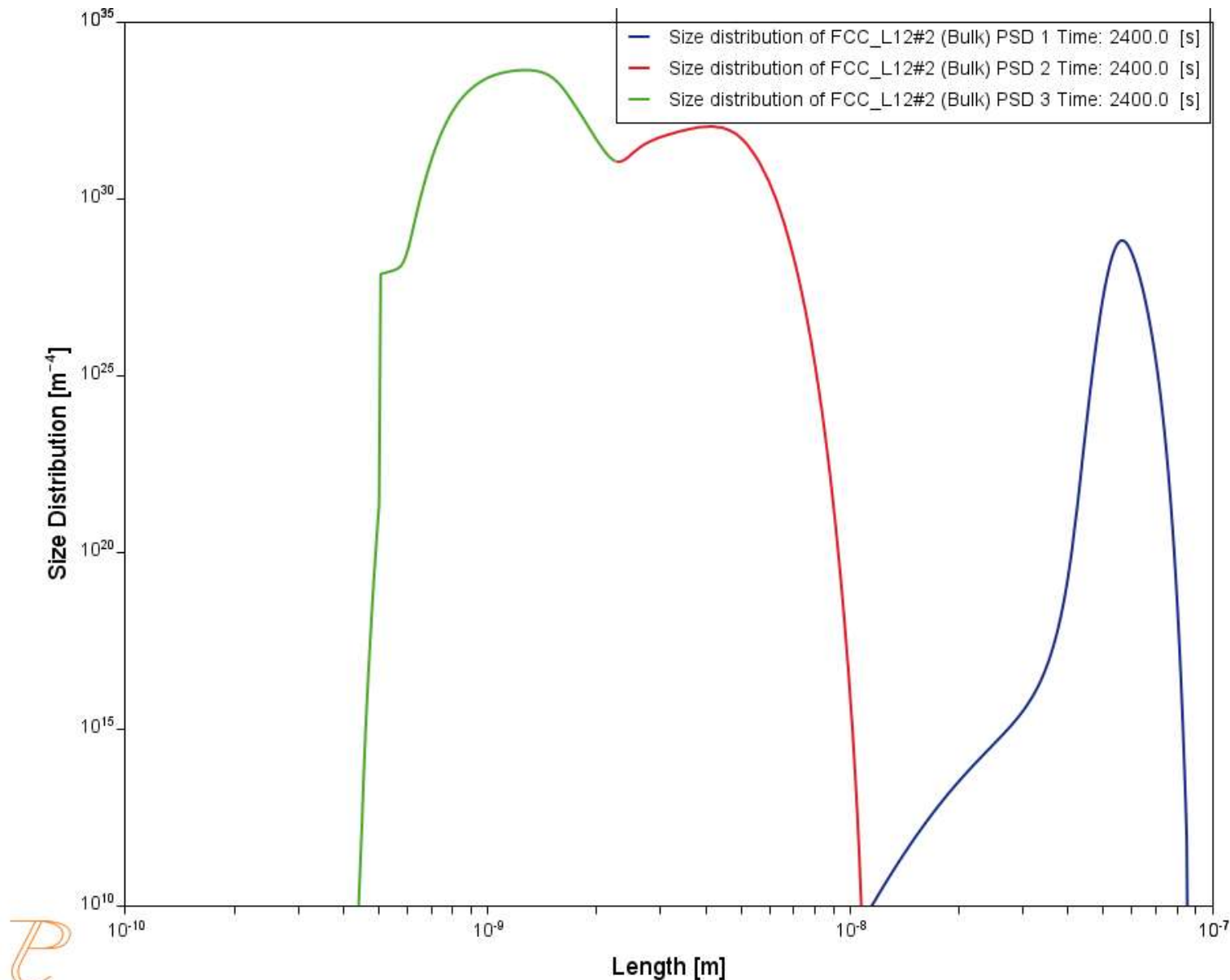
Ni – 8 Al – 8 Cr



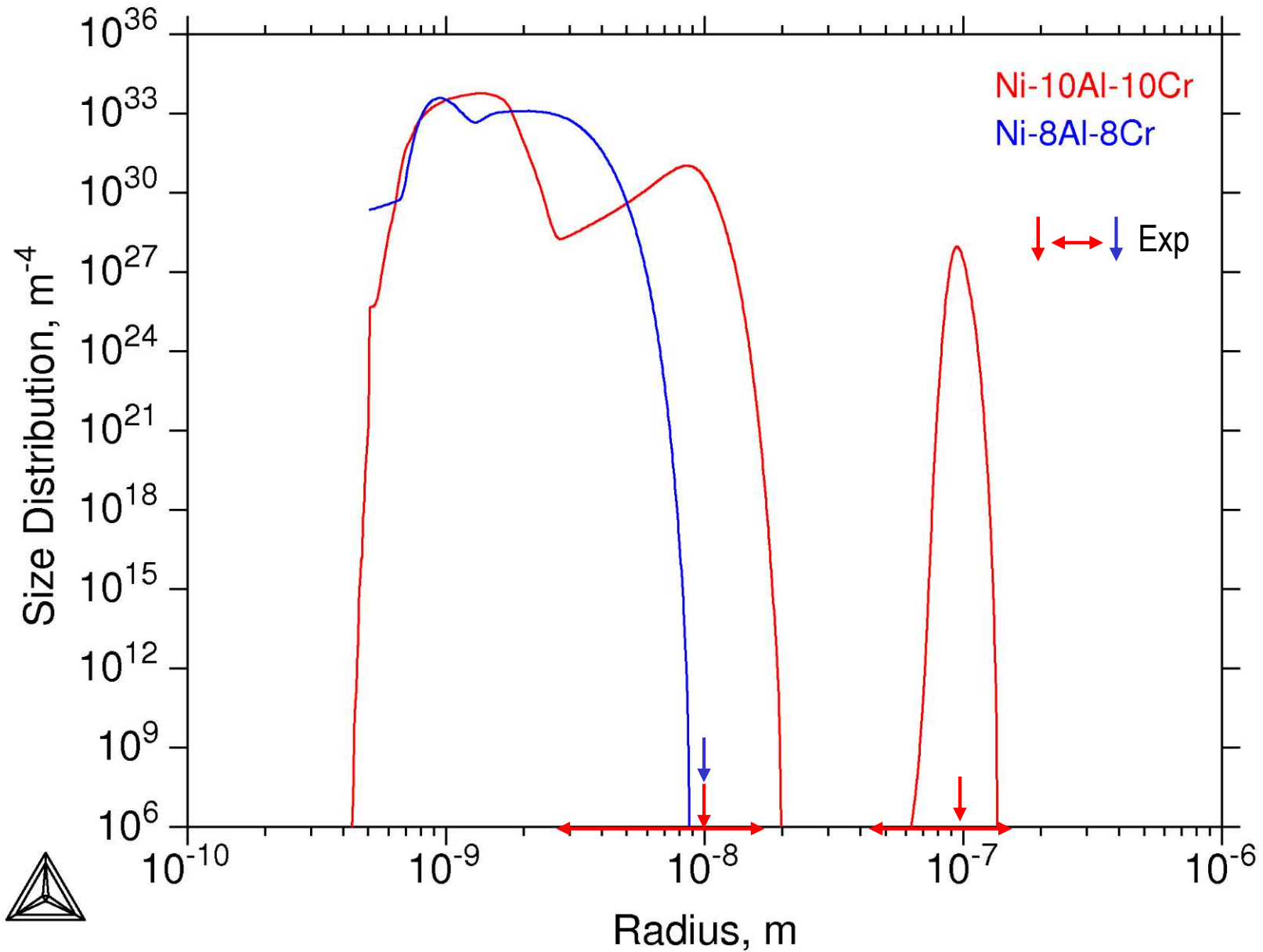
P

Ni-alloy Example 2

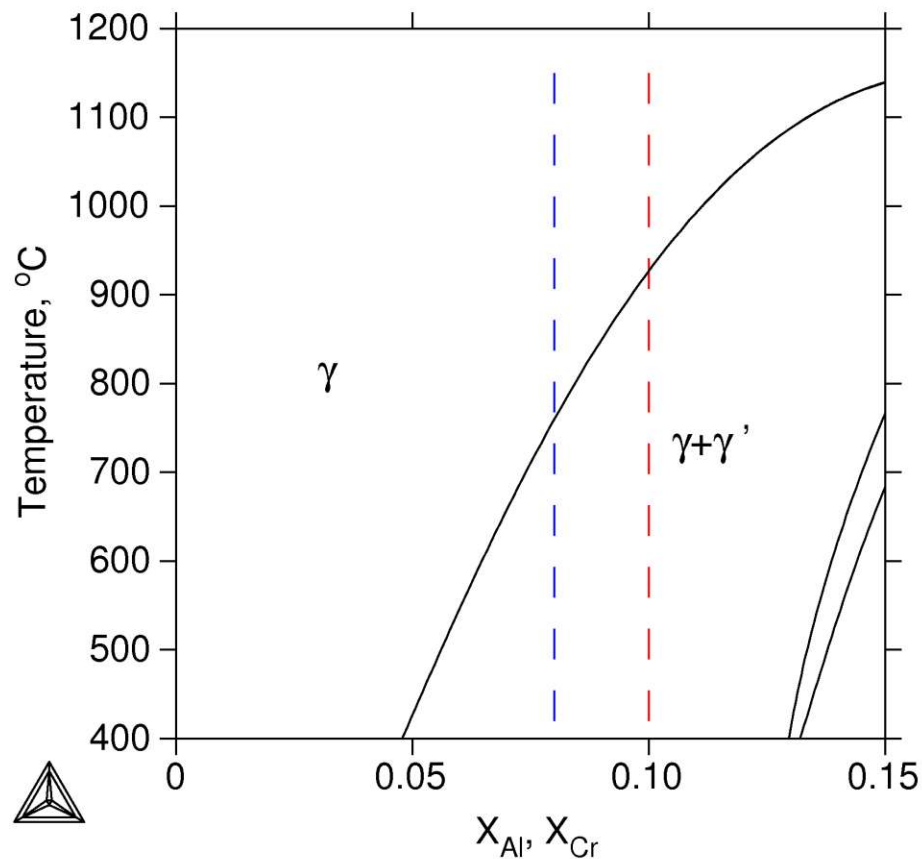
Ni – 10 Al – 10 Cr



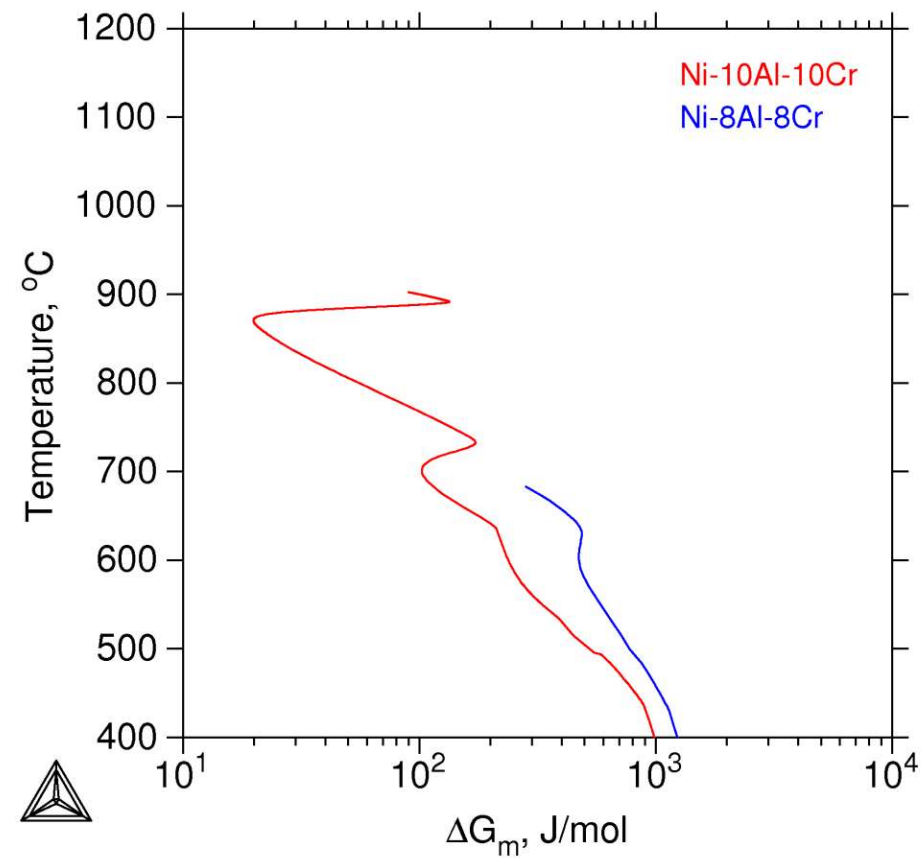
Ni-10Al-10Cr and Ni-8Al-8Cr



Ni-10Al-10Cr and Ni-8Al-8Cr

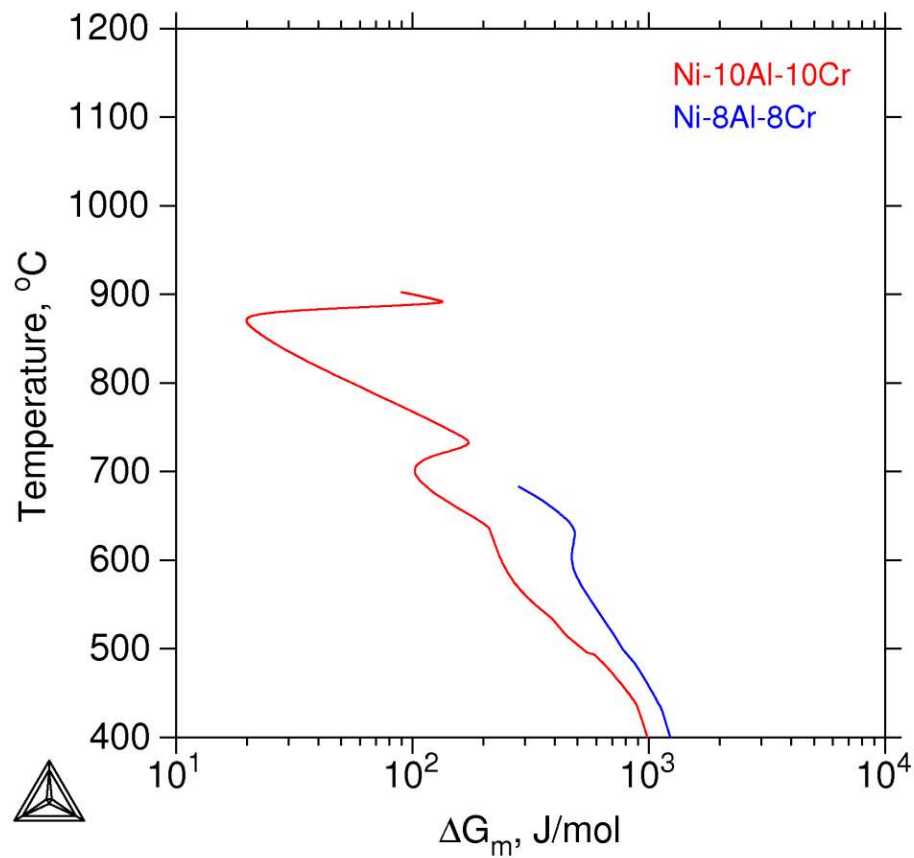


Vertical phase diagram
section in Ni-xAl-xCr

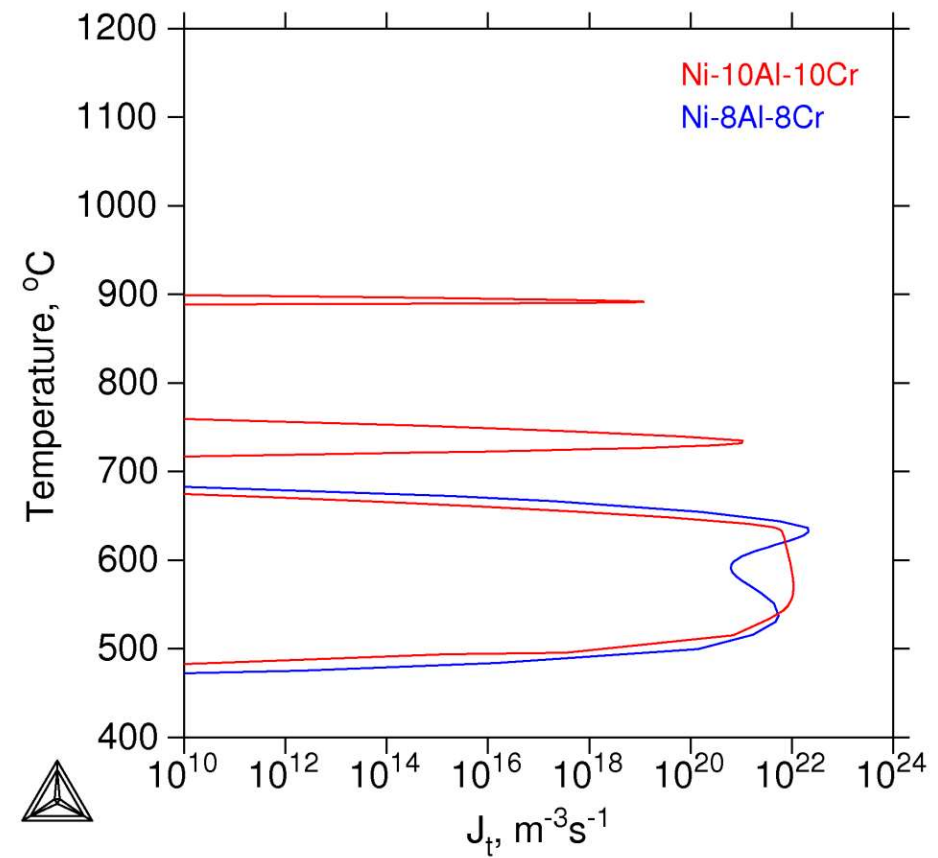


Thermodynamic driving force

Ni-10Al-10Cr and Ni-8Al-8Cr



Thermodynamic driving force

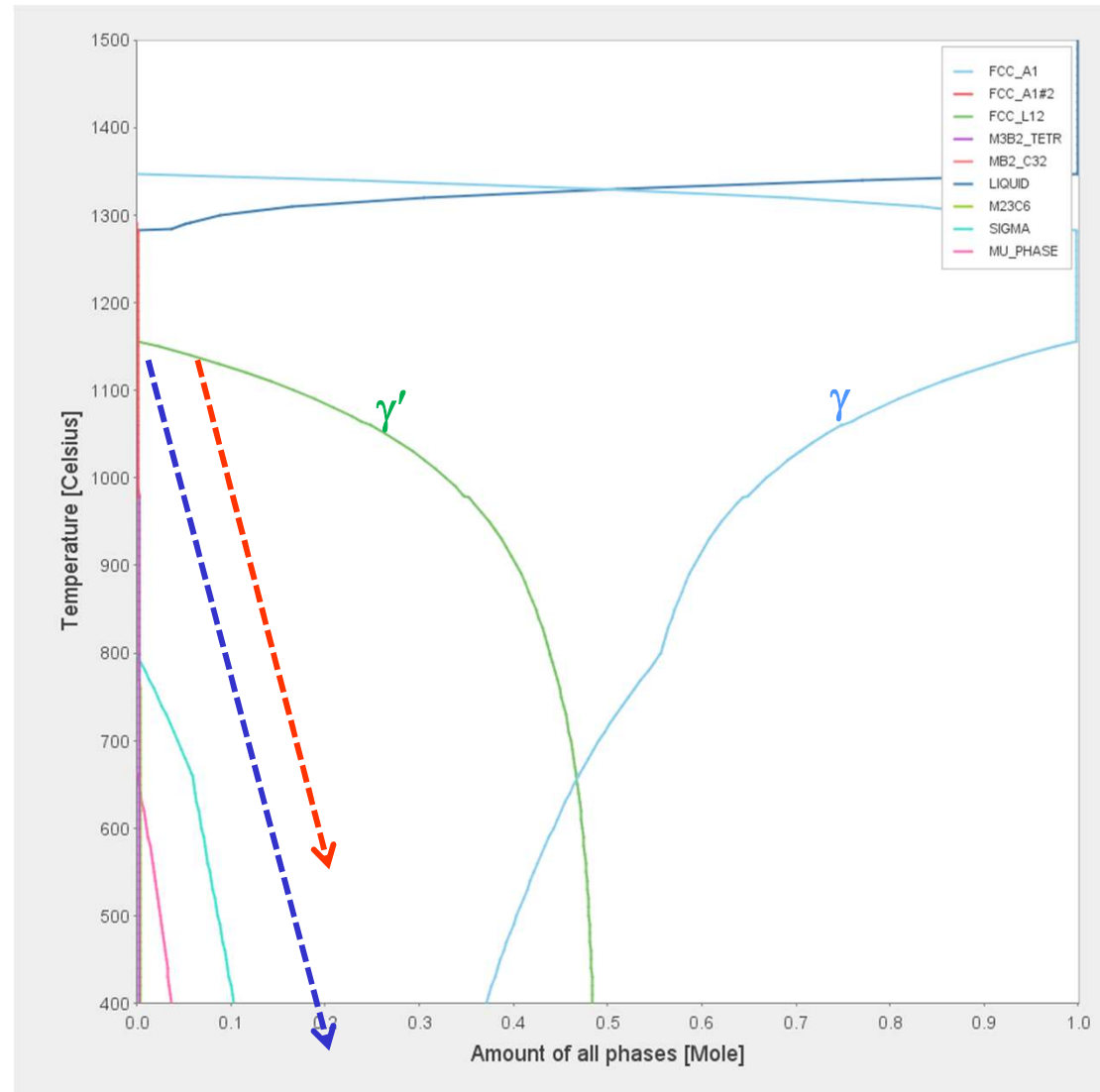


Nucleation rate

U720Li

Precipitation Kinetics during Continuous Cooling

wt.%	1*	2**
Al	2.53	2.46
B	0.014	
C	0.014	0.025
Co	14.43	14.75
Cr	15.92	16.35
Fe	0.09	0.06
Mo	2.96	3.02
Ti	4.96	4.99
W	1.26	1.3
Zr		0.035
Ni	Bal	Bal



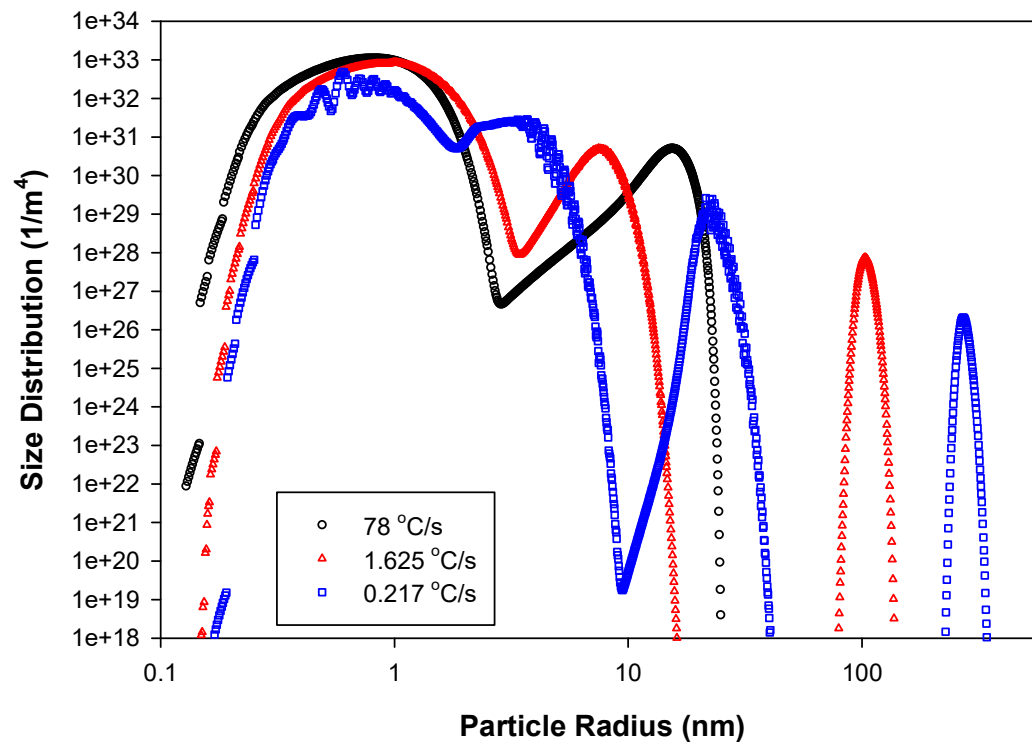
➤ Databases: TTNi8+MOBNI1

* Radis et al., *Superalloys* 2008

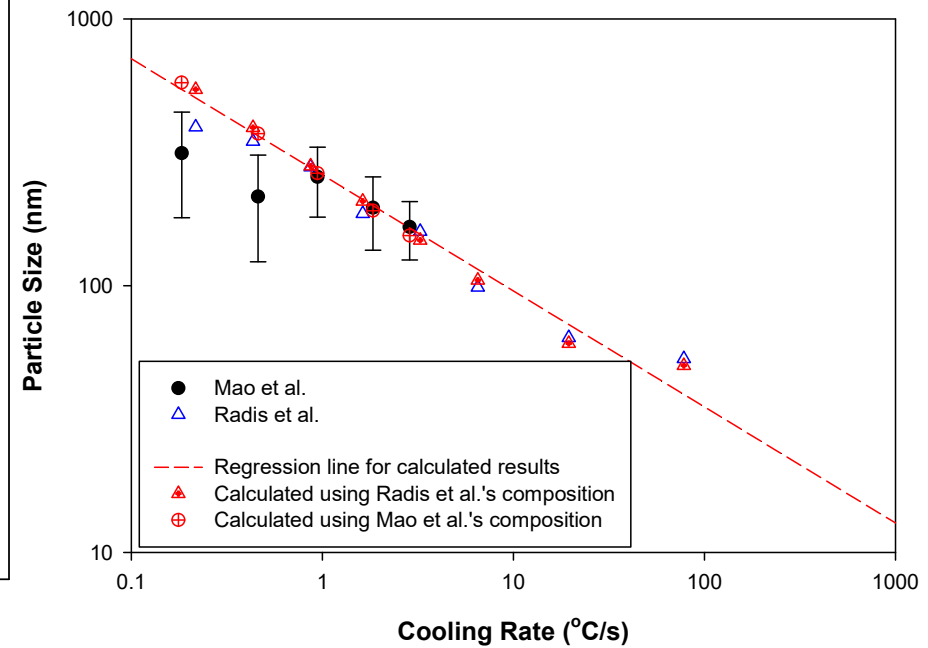
** Mao et al., *Metall. Mater. Trans. A*, 32A(10) 2441(2001)

U720Li : Cooling Rate Effect

Size Distribution



Mean Particle Size



Q & A

Examples

Steel



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Simulation of the kinetics of precipitation reactions in ferritic steels

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Abstract

Computer simulations of diffusion-controlled phase transformations in model alloys of Fe–Cr–C, Fe–Cr–W–C, Fe–Cr–Si–C, and Fe–Cr–Co–V–C are presented. The compositions considered are typical for ferritic steels. The simulations are performed using the software DICTRA and the thermodynamic calculations of phase equilibria are performed using Thermo-Calc. The thermodynamic driving forces and the kinetics of diffusion-controlled precipitation reactions of $M_{23}C_6$, M_7C_3 , cementite and Laves-phase (Fe, Cr)₂W are discussed. The simultaneous growth of stable and metastable phases is treated in a multi-cell approach. The results show remarkable effects on the growth kinetics due to the competition during simultaneous growth.

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Keywords: Ferritic steels; Phase transformation kinetics; Thermodynamics; Kinetics

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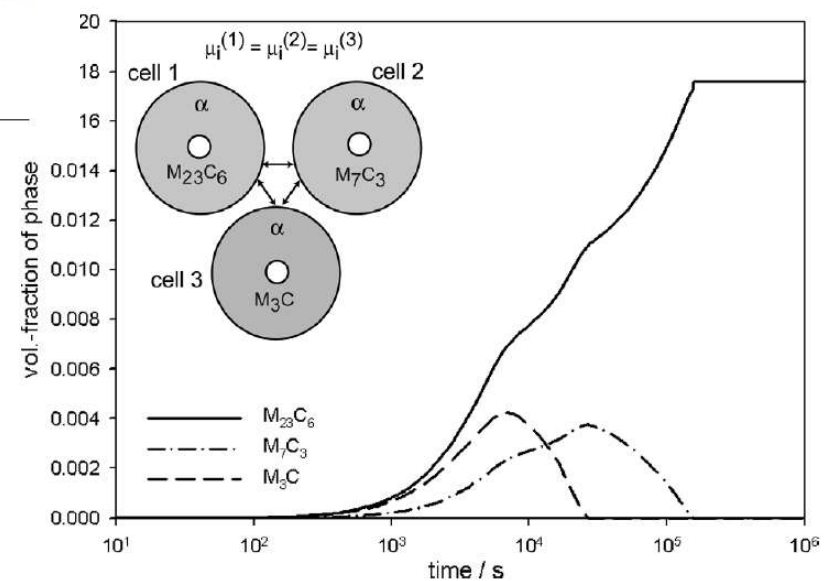


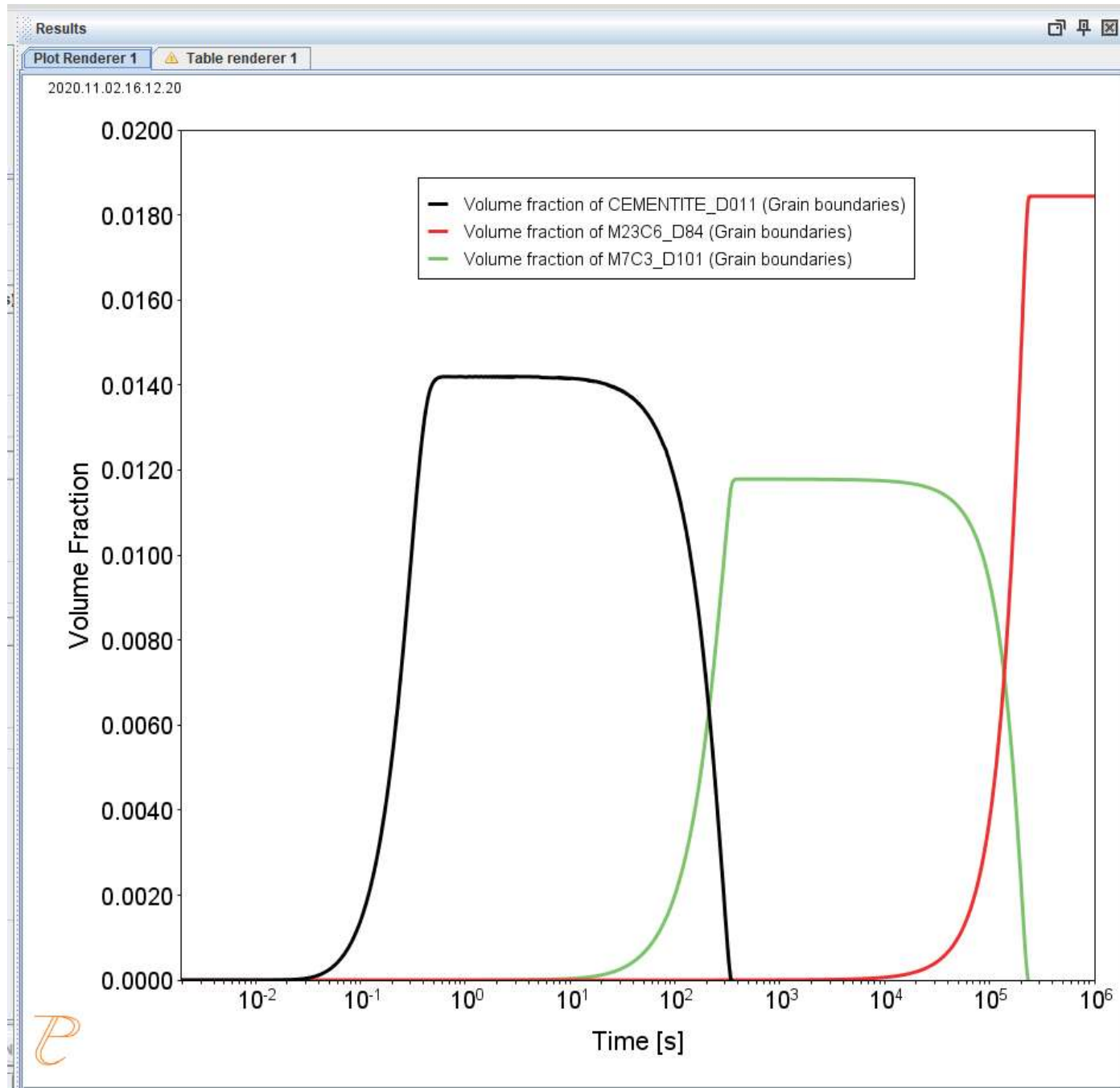
Fig. 6. Three-cell simulation of competitive growth of stable $M_{23}C_6$ and of metastable M_7C_3 and M_3C in Fe–12Cr–0.1C at 1053 K were 5 μ m and the nucleus sizes were 1 nm in each cell.

Steel Example 1

System	
Database package	TCFE14 + MOBFE8
Elements	Fe, Cr, C
Matrix phase	Bcc_A2
Precipitate phases	M ₂₃ C ₆ , M ₇ C ₃ , Cementite
Conditions	
Composition	Fe – 12 Cr – 0.1 C (wt.%)
Temperature	1053 K
Simulation time	1e6 s
Nucleation properties	Nucleation Site Type: Grain Boundaries
Data Parameters–Interfacial Energies	
Cementite	0.167 J/m ²
M ₂₃ C ₆	0.252 J/m ²
M ₇ C ₃	0.282 J/m ²

Including already existing size distributions – see Example 2

Steel Example 1



Steel Example 2

Same as Ex 1 but starting from already existing particle distributions of all three carbides, with small average radii and small volume fractions.

LSW
Normal
Log normal
Weibull

Precipitate Phase

Phase:

Nucleation sites: ☒ Calculate f

Interfacial energy:

Growth rate model:

Morphology:

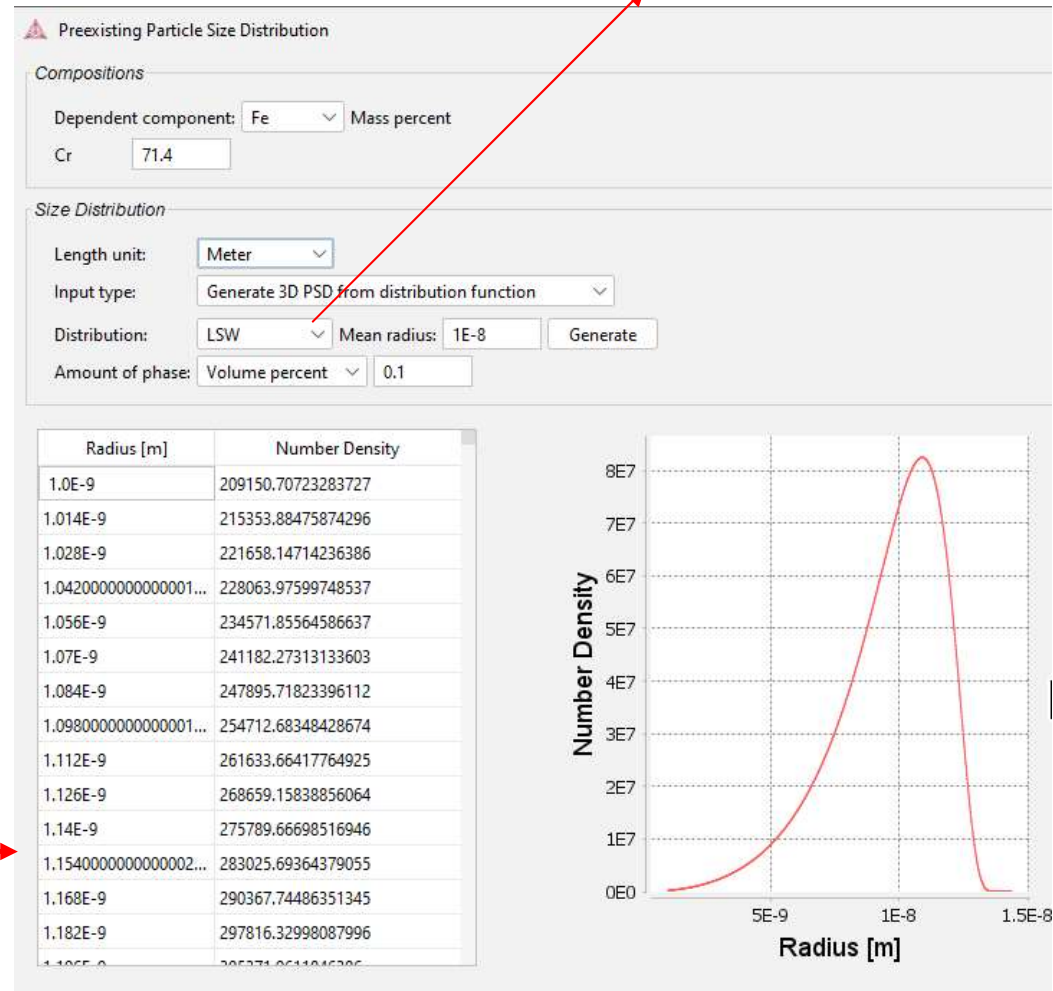
Transformation strain:

Molar volume:

Phase boundary mobility:

Approximate driving force: ☐

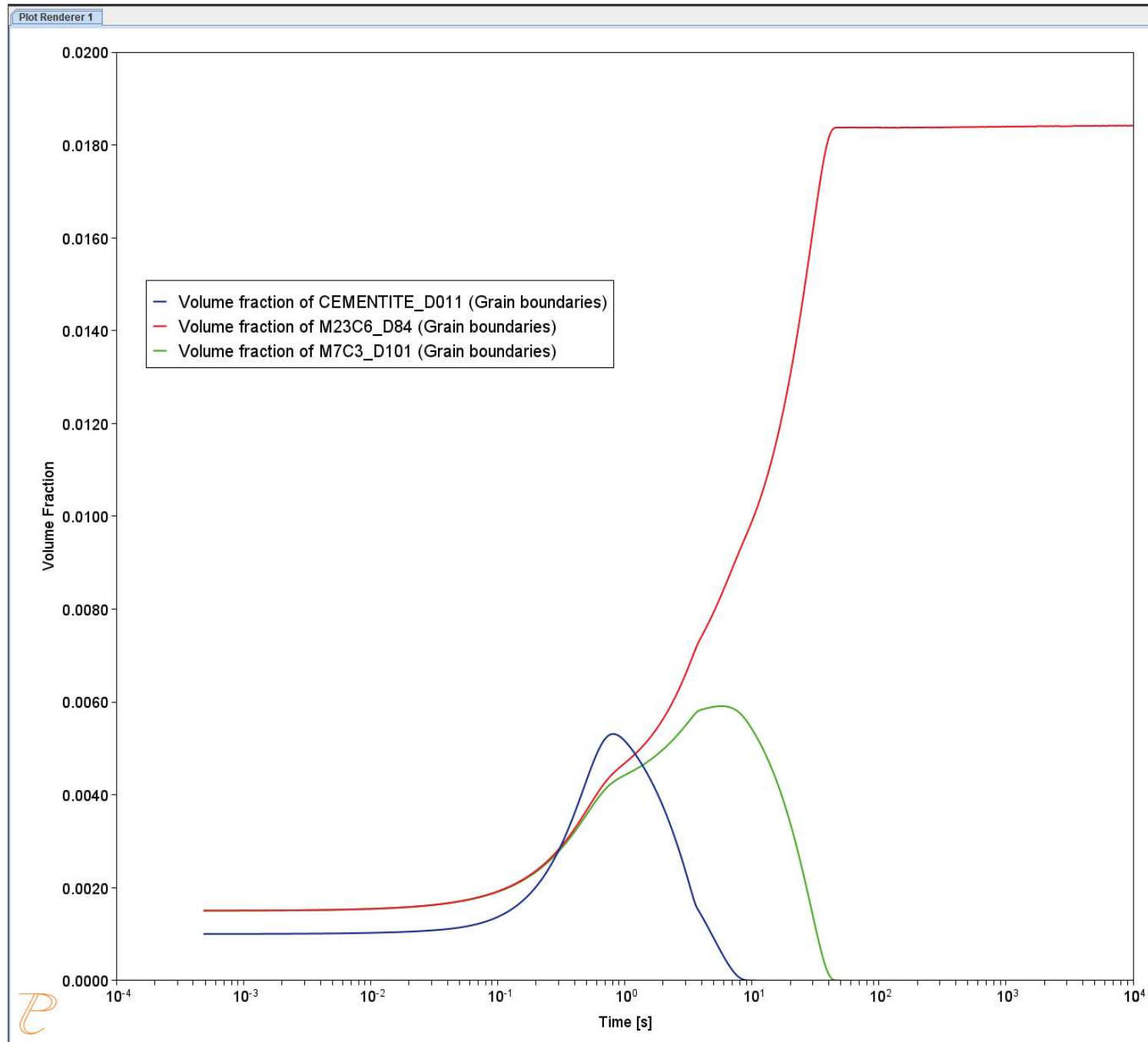
Preexisting size distribution: ☒



Steel Example 2

Initial Particle Size Distribution	
Phase	CEMENTITE
Initial composition	Cr 71.4 wt.%
Distribution	LSW with mean radius 1.0e-8m
Amount	0.001 (volume fraction)
Initial Particle Size Distribution	
Phase	M23C6
Initial composition	Cr 69.3 wt.%
Distribution	LSW with mean radius 1.0e-8m
Amount	0.0015 (volume fraction)
Initial Particle Size Distribution	
Phase	M7C3
Initial composition	Cr 82.9 wt.%
Distribution	LSW with mean radius 1.0e-8m
Amount	0.0015 (volume fraction)

Steel Example 2



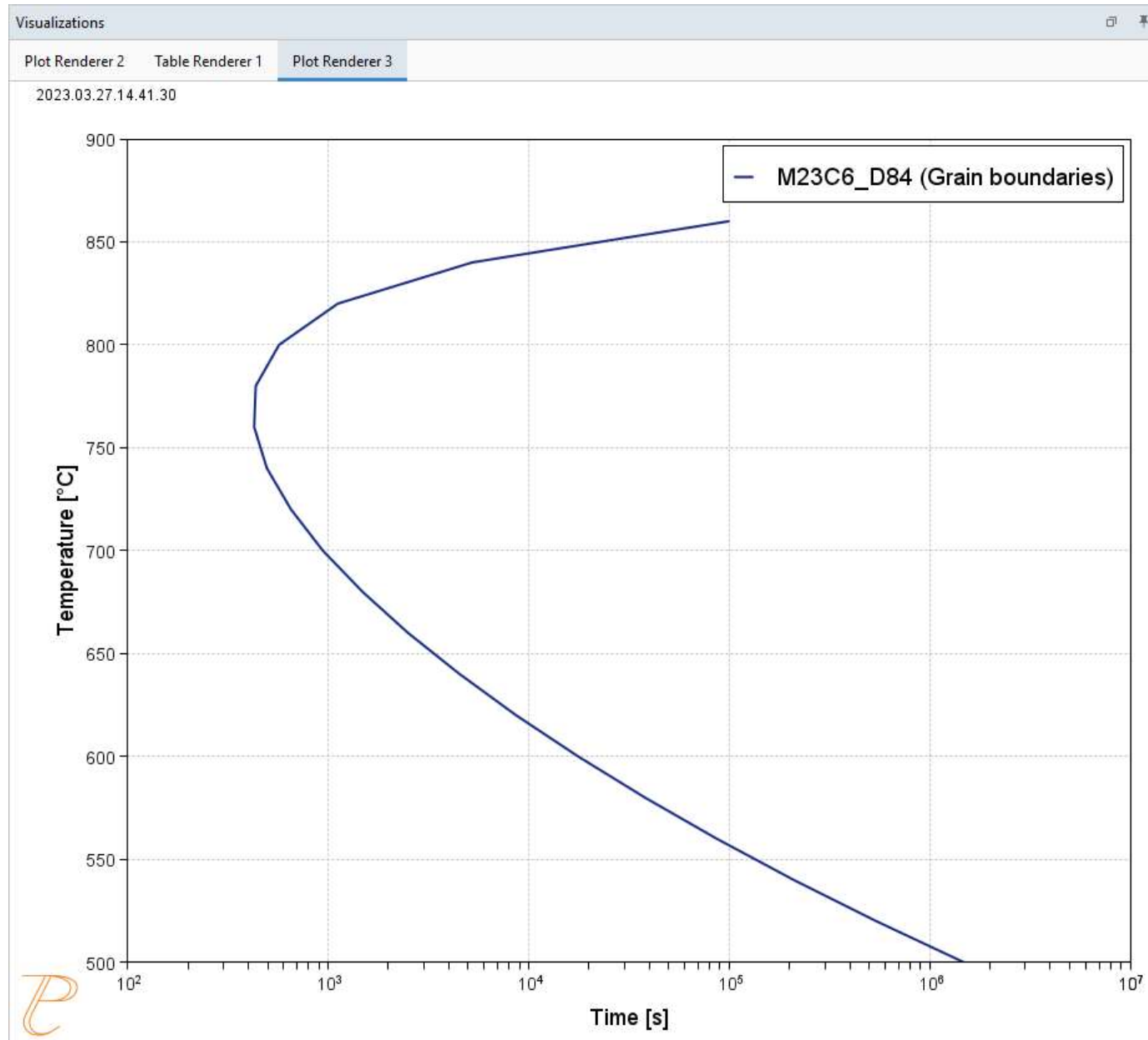
Steel Example 3 – TTP

System	
Database package	TCFE14 + MOBFE8
Elements	Fe,C,Cr,Mn,Ni,Si
Matrix phase	Fcc_A1
Precipitate phase	M ₂₃ C ₆
Conditions – TTT diagram	
Composition	Fe-0.068C-20.89Cr-1.61Mn-10.28Ni-0.49Si (wt.%)
Temperature	500 °C, 860 °C, 20 °C
Simulation time	1E7 s
Nucleation properties	Nucleation Site Type: Grain Boundary, Grain size 100 μm
Data Parameters	
Interfacial Energy	Grain Boundary: Calculated

Number of grid points: 15
Maximum number of grid points: 20
Minimum number of grid points: 10

Use the
General model

Steel Example 3

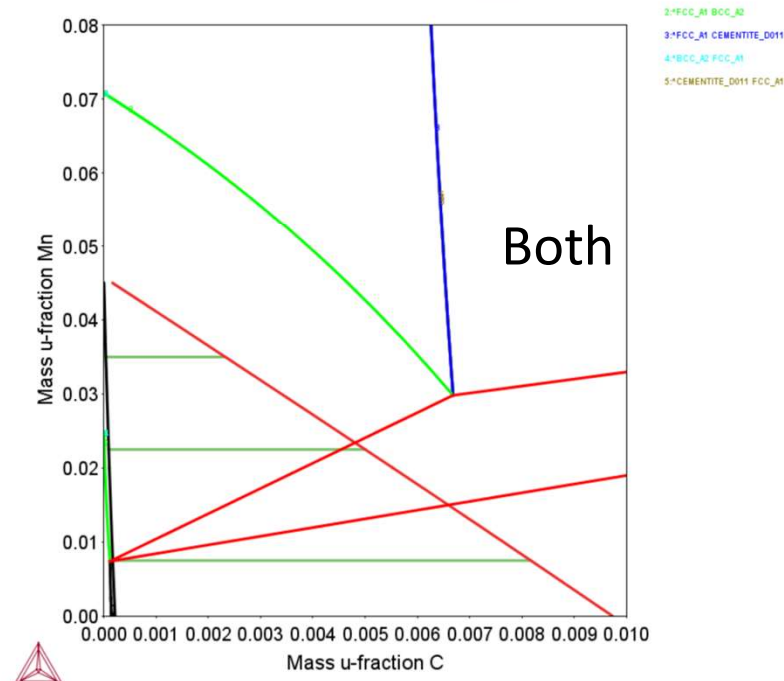
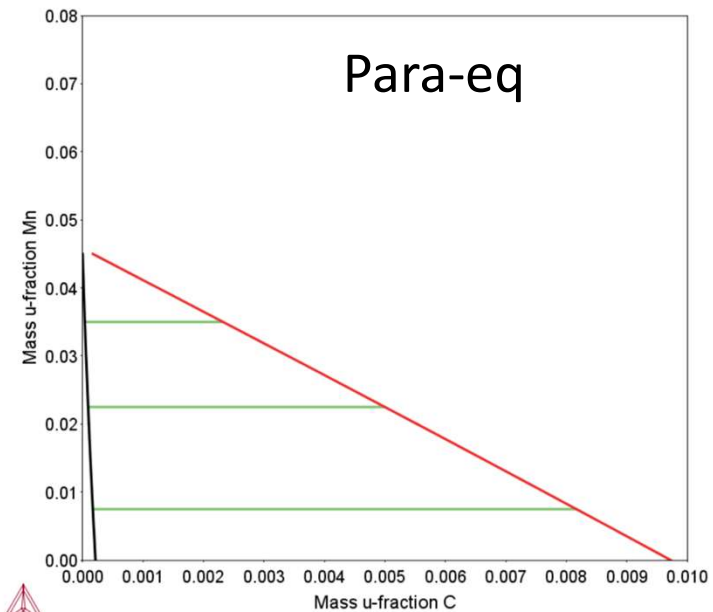
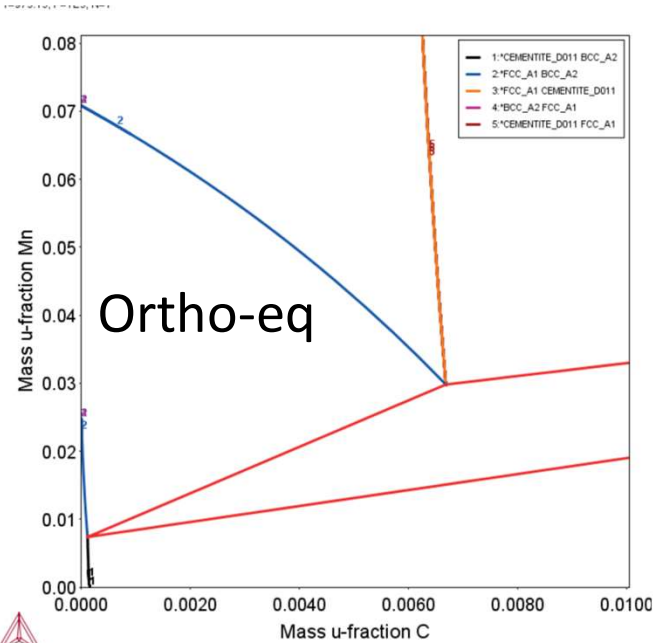


P

Example

Steel, Para-Equilibrium Models

Para-equilibrium



Bcc + Fcc equilibria
in Fe – Mn – C

Example: Para-equilibrium or not?

Precipitation of cementite during tempering of martensite in a Fe-Mn-C steel.

Consider three different phase interface conditions: the usual ortho-equilibrium condition, para-equilibrium condition, and a smooth transition from para-equilibrium to ortho-equilibrium condition using the recent **PE Automatic** growth rate model.

The simulation results can be compared with the experimental data from Miyamoto et al. [2007Miy].

Martensite will be represented by BCC_A2 which requires some rather unusual settings for mobility, grain size and grain shape.

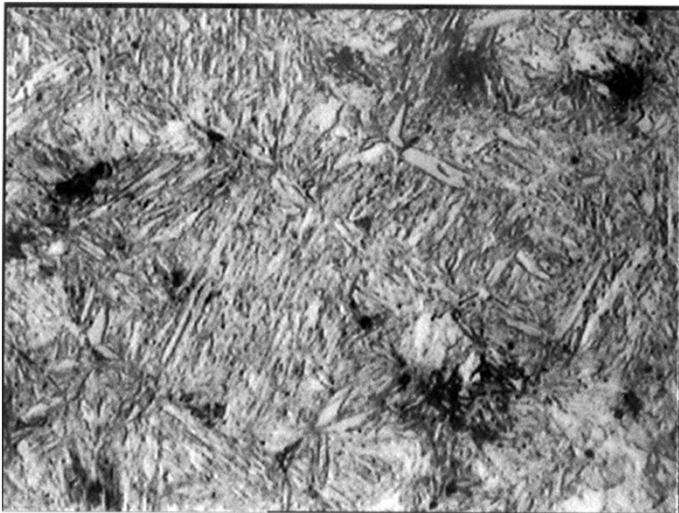


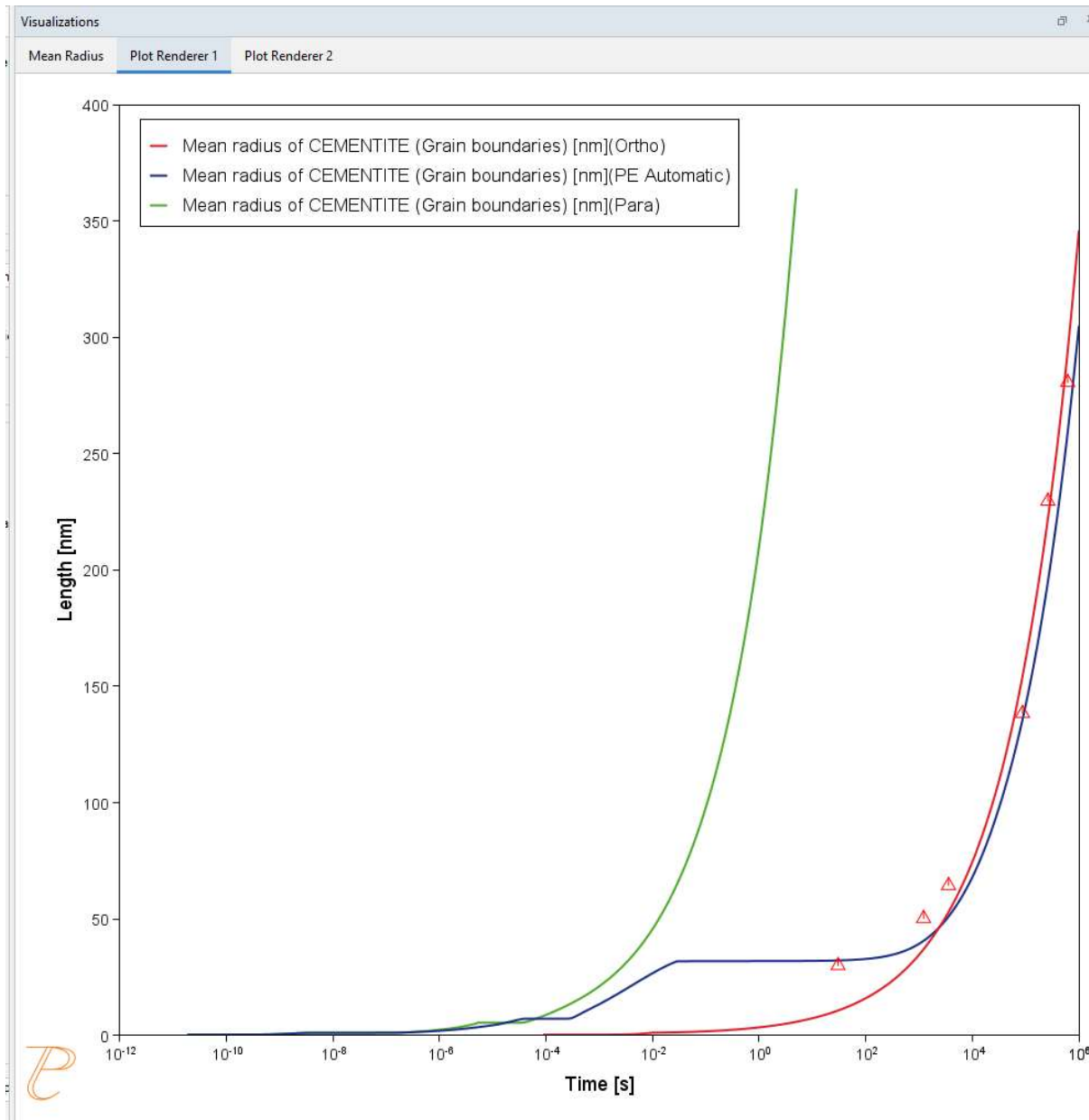
Image from Wikipedia

[2007Miy] G. Miyamoto, J. Oh, K. Hono, T. Furuhashi, T. Maki, Effect of partitioning of Mn and Si on the growth kinetics of cementite in tempered Fe-0.6 mass% C martensite. Acta Mater. 55, 5027–5038 (2007).

Example – Para-equilibrium or not?

System	
Database package	FEDEMO + MFEDEMO
Elements	Fe, Mn, C
Matrix phase	BCC_A2
Precipitate phase	Cementite
Conditions	
Composition	Fe- 0.61 C- 1.96 Mn (wt-%)
Temperature	650 °C
Simulation time	1E6 s (5 seconds for PE)
Growth rate models	Simplified / Para-eq / PE Automatic
Nucleation properties	Nucleation Site: Grain boundaries
Data Parameters	
Interfacial Energy	Calculated
Molar Volumes	Database
Grain size / Grain aspect ratio	1E-7 m / 100
Mobility Adjustment factor	0.008
Activation energy	- 70000 J/mol

Example – Para-equilibrium or not?



For those of you who will take part in the courses next week:

AM Online 28-29 Oct.

You already have the software and database installation you need. Just keep it.

You will be mailed new license credentials, and new details for Zoom meetings and material.

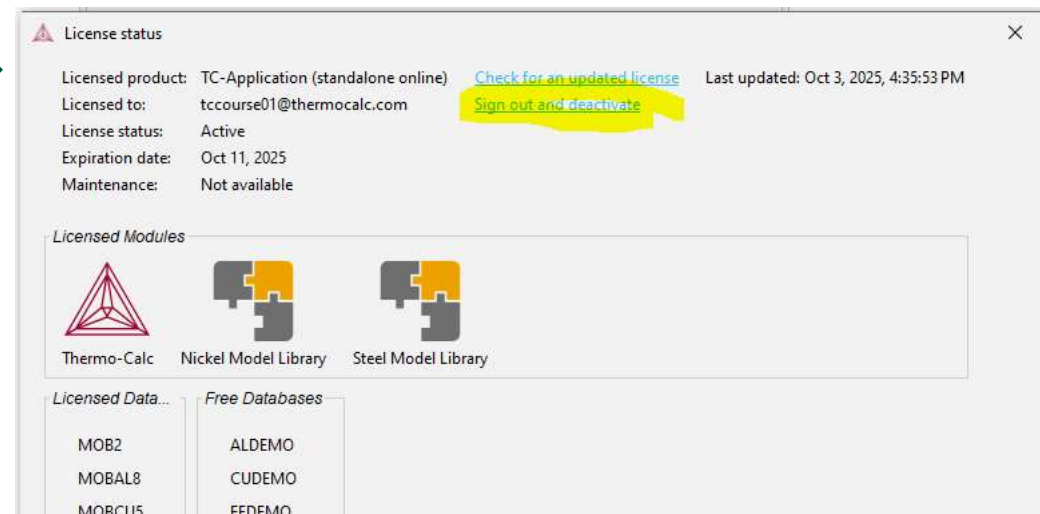
The current course license will be valid on Friday 24 Oct.,
but not on Saturday.

For those of you participating in

AM Online 28-29 Oct.

It will be easiest if you deactivate the current license at
the latest on Friday 24 Oct. You will be sent a new
license for TC + AM Module.

Help > Show License Info



We will send a **certificate of course completion** by email after also the AM Online course is finished next week.

Email ake@thermocalc.se if you think we have your name and affiliation incorrect.

Q & A

End of course.