

TCS High Entropy Alloys Database (TCHEA7)

Technical Information

Available Starting with Thermo-Calc Version 2024a



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About the TCS High Entropy Alloys Database (TCHEA)

TCS High Entropy Alloys Database (TCHEA) is a thermodynamic database for high entropy alloys (HEA) [2004Yeh; 2006Yeh]. HEAs are a new class of materials consisting of more than one principal element in a multi-component system. They are also known as multi-principal element alloys (MPEAs) [2013Wan; 2015Sen] or complex concentrated alloys (CCAs). The coupling of our CALPHAD based computational tools and databases allows a high fidelity calculation of thermodynamic properties and phase equilibria in multi-component HEAs, thus shedding light on the formation mechanism and thermodynamic and kinetic stability of HEAs, providing an efficient way to design HEAs for desired materials properties based on the prediction of microstructures through process optimization. In addition to thermodynamic data, it has properties data available for molar volume, viscosity, surface tension, electrical resistivity, and thermal conductivity.



[TCHEA: TCS High Entropy Alloys Database Revision History](#). The current version of the database is TCHEA7. See the link for any subversion release details.



The database is compatible with the TCS High Entropy Alloy Mobility Database (MOBHEA). The current version is MOBHEA3.

The CALPHAD Method

The Thermo-Calc databases are developed with the CALPHAD approach based on various types of experimental data and theoretical values (e.g. those from first-principles calculations). It is based on the critical evaluation of all the binary systems and many ternary systems. A hybrid approach of experiments, first-principal calculations and CALPHAD modeling has been used to obtain reliable thermodynamic descriptions of the BCC, FCC and HCP solutions. That enables predictions to be made for multicomponent alloy systems, especially for HEAs.

The extrapolation to higher-order systems helps to understand the phase equilibria in HEAs, so as to predict the phase formation, phase fractions and phase compositions or to calculate the driving force of forming a phase. All necessary molar volume data and thermal expansion data are assessed or estimated for most of the phases.



Learn more on our website about the [CALPHAD Method](#) and how it is applied to the Thermo-Calc databases.

Use Case Examples

There are examples available to both demonstrate the *validation* of the database and to showcase the types of *calculations* that can be used for different materials or application area.

Some case examples of how the TCHEA7 database can be used include:

- Calculate various phase diagrams and property diagrams in the assessed systems and higher-order systems.
- Predict solidification behavior of HEAs with a Scheil-Gulliver calculation.
- Predict a wide variety of equilibrium properties such as phase amount and constitution as a function of composition or temperature, homogeneity range, or thermo-stability of HEAs.
- Use the Add-on Diffusion Module (DICTRA) and combine it with TCHEA and compatible kinetic databases to simulate typical diffusion-controlled phase transformations in HEAs under arbitrary heat treatment conditions.
- Use the Precipitation Module (TC-PRISMA) and combine it with TCHEA and compatible kinetic databases, to simulate the concurrent nucleation, growth and coarsening of precipitates.

Combining Databases

It is possible to combine several databases to make calculations using Thermo-Calc. For more information related to a specific type of problem, contact one of our support specialists at info@thermocalc.com. The experts are available to make recommendations on the most suitable database to use for your needs.

References

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- [2017Mir] D.B. Miracle, O.N. Senkov, A critical review of high entropy alloys and related concepts, *Acta Mater.* 122 (2017) 448–511.
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Suggested References to Cite this Database

[2017Mao] H. Mao, H.-L. Chen, Q. Chen, TCHEA1: A Thermodynamic Database Not Limited for “High Entropy” Alloys. *J. Phase Equilibria Diffus.* 38 (2017) 353–368.

[2018Chen] H.-L. Chen, H. Mao, Q. Chen, Database development and Calphad calculations for high entropy alloys: Challenges, strategies, and tips. *Mater. Chem. Phys.* 210 (2018) 279-290.

TCS High Entropy Alloys Database (TCHEA) Resources

Information about the database is available on our website and in the Thermo-Calc software online Help.

- **Website:** On our website the information is both searchable and the database specific PDFs are available to download.
- **Online Help:** Technical database information is included with the Thermo-Calc software online Help. When in Thermo-Calc, press F1 to search for the same information as is contained in the PDF documents described. Depending on the database, there are additional examples available on the website.

Database Specific Documentation

- The *TCS High Entropy Alloys Database (TCHEA) Technical Information* PDF document contains version specific information such as the binary and ternary assessed systems, phases and models. It also includes details about the properties data (e.g. viscosity, surface tension, etc.), a list of the included elements, and summaries of the database revision history by version.
- The *TCS High Entropy Alloys Database (TCHEA) Examples Collection* PDF document contains a series of validation examples using experimental data, and a set of calculation examples showing some of the ways the database can be used.



Go to the [High Entropy Alloys Databases](#) page on our website where you can access an examples collection and the technical information plus learn more about the compatible kinetic database. Also explore further [applications of Thermo-Calc to high entropy alloys](#) including links to resources such as examples, publications, and more.



Learn more on our website about the [CALPHAD Method](#) and how it is applied to the Thermo-Calc databases.

TCHEA7 Elements, Systems, Phases, and Properties Data

Included Elements

The database has been developed in a 26 element framework:

Al	B	C	Co	Cr	Cu	Fe	Hf	Ir
Mn	Mo	N	Nb	Ni	Re	Rh	Ru	Si
Sn	Ta	Ti	V	W	Y	Zn	Zr	

Assessed Systems and Phases

The most recent version of the database contains:

- 319 binary systems, with almost all of these assessed to the full range of composition and temperature. These can be calculated with the BINARY module in Thermo-Calc Console Mode.
- 526 ternaries are assessed, and 212 of them to the full range of composition and temperature. These can be calculated with the TERNARY module in Thermo-Calc Console Mode.
- 540 solution and intermetallic phases, where nearly all stable phases in all assessed binary systems and most ternary systems are modeled.

About the Included Phases

The ordered B2 and L1₂ phases, together with BCC_A2 and FCC_A1, respectively, are modeled with the so-called partitioning model, which describes an ordered phase and its disordered counterpart using a single Gibbs energy curve. This type of description is of particular importance to be able to predict second order transformations between a disordered phase and its ordered structures.

Also note that there may be several possible composition sets for the phases named FCC_L12 and BCC_B2 designated by #1, #2, and so on (e.g. FCC_L12#1 and FCC_L12#2), due to the co-existence of disordered and ordered structures or the presence of miscibility gap. The #n suffix (where n is an integer) is generated dynamically by Thermo-Calc when using global minimization and therefore the identification of the phases should be determined from their site occupations. It can be found by LIST_EQUILIBRIUM with the VXNS option in the Console Mode or showing the site fraction in moles of the constituent elements in the Graphical Mode. When the site occupancies of the first and second sublattices are equal the phase is disordered.



In Console Mode, you can list phases and constituents in the Database (TDB) module and the Gibbs (GES) module. For some phases, supplementary information is included in the definitions. To show the information, it is recommended in the Database (TDB) module to use the command `LIST_SYSTEM` with the option `Constituents`.

Properties Data



You can find information on our website about the [properties that can be calculated](#) with Thermo-Calc and the Add-on Modules. Additional resources are added on a regular basis so keep checking back or [subscribe to our newsletter](#).

For more information about the various thermophysical models, and when in Thermo-Calc, press F1 to search the online help. The details are found under a *General Reference* section.

Below is a summary of the available parameters and variables for the databases when working in Thermo-Calc. There are differences when you are working in Console Mode versus Graphical Mode as well as if you use the TC-Python SDK. More details are described in the online help.

<i>Property</i>	<i>Model Parameters</i>	<i>Variables to Show or Plot in Console Mode and TC-Python</i>
Electrical conductivity	ELQ**	ELCD for a system <code>ELCD (PHI)</code> for phase PHI
Electrical resistivity	ELRS, ESPD	ELRS for a system <code>ELRS (PHI)</code> for a phase PHI
Thermal conductivity	THCD	THCD for a system <code>THCD (PHI)</code> for phase PHI
Thermal resistivity		THRS for a system <code>THRS (PHI)</code> for phase PHI
Thermal diffusivity		THDF for a system <code>THDF (PHI)</code> for phase PHI
Surface tension	SIGM, XI*	<code>SURF (LIQUID)</code> <code>SURF (ION)</code> **
Dynamic viscosity	VISC	<code>DVIS (LIQUID)</code> <code>DVIS (ION)</code> **
Kinematic viscosity		<code>KVIS (LIQUID)</code> <code>KVIS (ION)</code> **

<i>Property</i>	<i>Model Parameters</i>	<i>Variables to Show or Plot in Console Mode and TC-Python</i>
Molar volume	V0, VA	VM for a system $V_{M(PHI)}$ for phase PHI

* XI is not used in the TCOX database (all versions). As of 2023b it is also not used starting with the following versions of these databases: TCFE13, TCNI12.1, TCTI5.1, TCNOBL3, TCPMAG2, and TCCU6. As of 2024a, TCMG7, TCAL9, and TCHEA7.

** I_{ON} is used in the TCS Metal Oxide Solutions Database (TCOX)

TCHEA7 Systems

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TCHEA7 Critically Assessed Ternary Systems

These are the assessed ternary systems in the full range of composition and temperature.

<i>Critically Assessed Ternary Systems</i>				
Al-B-Ti	Al-C-Co	Al-C-Fe	Al-Co-Ni	Al-Co-Ti
Al-Co-W	Al-Co-Zr	Al-Cr-Ni	Al-Cr-Sn	Al-Cr-Ti
Al-Cr-V	Al-C-Ti	Al-Cu-Fe	Al-Cu-Mn	Al-Cu-Ni
Al-Cu-Si	Al-Cu-Sn	Al-Cu-Zn	Al-Fe-Mn	Al-Fe-N
Al-Fe-Si	Al-Fe-Ti	Al-Mn-Ni	Al-Mn-Si	Al-Mn-Ti
Al-Mn-Zn	Al-Mo-Ni	Al-Mo-Ti	Al-Nb-Ni	Al-Nb-Ti
Al-Ni-Ru	Al-Ni-Si	Al-Ni-Ta	Al-Ni-Ti	Al-Ni-W
Al-Ni-Zn	Al-N-Ti	Al-Ru-Ti	Al-Si-Sn	Al-Si-Ti
Al-Si-Zn	Al-Sn-Ti	Al-Sn-Zn	Al-Ta-Ti	Al-Ti-V
Al-Ti-W	Al-Ti-Y	Al-Ti-Zr	Al-V-Zr	C-Co-Cr
C-Co-Fe	C-Co-Mo	C-Co-Nb	C-Co-Ni	C-Co-Ta
C-Co-Ti	C-Co-W	C-Cr-Fe	C-Cr-Hf	C-Cr-Si
C-Cr-Ti	C-Cr-V	C-Cr-Zr	C-Cu-Fe	C-Fe-Mn
C-Fe-Mo	C-Fe-N	C-Fe-Ni	C-Fe-Si	C-Fe-Ti
C-Fe-V	C-Fe-W	C-Hf-Mo	C-Hf-Ni	C-Mn-Si
C-Mo-Ni	C-Mo-Ta	C-Mo-Ti	C-Mo-V	C-Mo-W
C-Mo-Zr	C-Nb-Ni	C-Nb-Re	C-Nb-Ti	C-Nb-W
C-Ni-Ta	C-Ni-Ti	C-Ni-W	C-Ni-Zr	Co-Cr-Cu
Co-Cr-Fe	Co-Cr-Mn	Co-Cr-Ni	Co-Cr-Ti	Co-Cr-W
Co-Cu-Fe	Co-Cu-Mn	Co-Cu-Nb	Co-Cu-Ni	Co-Fe-Mn

Critically Assessed Ternary Systems

Co-Fe-Mo	Co-Fe-N	Co-Fe-Ni	Co-Fe-W	Co-Mn-Ni
Co-Mo-Ti	Co-Ni-Si	Co-Ni-V	Co-Ni-W	Co-Sn-Ti
Co-Ta-Ti	Co-W-Zr	Cr-Cu-Fe	Cr-Cu-Mo	Cr-Cu-Nb
Cr-Cu-Si	Cr-Cu-Sn	Cr-Cu-Zr	Cr-Fe-Mn	Cr-Fe-Mo
Cr-Fe-N	Cr-Fe-Ni	Cr-Fe-Si	Cr-Fe-V	Cr-Mn-N
Cr-Mn-Ni	Cr-Mn-Ti	Cr-Mo-Nb	Cr-Mo-Ni	Cr-Mo-Ti
Cr-Nb-Ni	Cr-Nb-V	Cr-Nb-W	Cr-Ni-Re	Cr-Ni-Si
Cr-Ni-Ta	Cr-Ni-Ti	Cr-Ni-W	Cr-Ni-Zr	Cr-N-Ni
Cr-Si-Ti	Cr-Ti-Zr	C-Ta-W	C-Ti-W	Cu-Fe-Mn
Cu-Fe-Mo	Cu-Fe-N	Cu-Fe-Nb	Cu-Fe-Ni	Cu-Fe-Si
Cu-Fe-Sn	Cu-Fe-Ti	Cu-Fe-V	Cu-Mn-Ni	Cu-Mn-Si
Cu-Mn-Sn	Cu-Mn-Zn	Cu-Mo-Ni	Cu-Ni-Ti	Cu-Ni-Zn
Cu-Ti-Zr	Fe-Mn-N	Fe-Mn-Si	Fe-Mo-Ni	Fe-Nb-Ni
Fe-Ni-Ru	Fe-Ni-Si	Fe-Ni-Ti	Fe-Ni-W	Fe-N-Nb
Fe-N-Ni	Fe-N-Ti	Fe-N-V	Fe-Ti-V	Hf-Nb-Si
Hf-Ni-Ti	Ir-Rh-Ru	Mn-Ni-Si	Mn-Si-Zn	Mo-Nb-Ti
Mo-Nb-V	Mo-Ni-Ta	Mo-N-Ni	Mo-Ta-Ti	Mo-Ti-V
Mo-Ti-W	Mo-Ti-Zr	Nb-Ni-Ti	Nb-Ni-W	Nb-Re-V
Nb-Sn-Ti	Nb-Ta-Ti	Nb-Ti-V	Nb-Ti-W	Nb-Ti-Zr
Nb-V-Zr	Ni-Si-Ti	Ni-Ta-Ti	Ni-Ta-W	Ni-Ti-W
Ni-Ti-Zr	N-Ni-Ti	Re-Ta-W	Re-V-W	Si-Ti-W
Ta-Ti-V	Ta-Ti-W	Ta-Ti-Zr	Ti-V-W	Ti-V-Zr

Critically Assessed Ternary Systems

Ti-W-Zr	V-W-Zr			
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TCHEA7 Tentatively Assessed Ternary Systems

<i>Tentatively Assessed Ternary Systems</i>				
Al-C-Cr	Al-C-Ni	Al-Co-Cr	Al-Co-Hf	Al-Co-Mo
Al-Co-Nb	Al-Co-Ru	Al-Co-Si	Al-Co-Ta	Al-Cr-Fe
Al-Cr-Mo	Al-Cr-Nb	Al-Cr-Re	Al-Cr-Ru	Al-Cr-Si
Al-Cr-Ta	Al-Cr-W	Al-Cr-Zr	Al-C-Si	Al-Fe-Hf
Al-Fe-Mo	Al-Fe-Nb	Al-Fe-Ni	Al-Fe-Re	Al-Fe-Ta
Al-Fe-W	Al-Fe-Zr	Al-Hf-Ni	Al-Hf-Ru	Al-Hf-Si
Al-Hf-Ti	Al-Mo-Nb	Al-Mo-Re	Al-Mo-Ru	Al-Mo-Si
Al-Mo-Ta	Al-Mo-W	Al-Mo-Zr	Al-Nb-Re	Al-Nb-Ru
Al-Nb-Si	Al-Nb-Ta	Al-Nb-W	Al-Ni-Re	Al-Ni-V
Al-Ni-Zr	Al-Re-Ru	Al-Re-Ta	Al-Re-Ti	Al-Re-W
Al-Ru-Ta	Al-Ru-W	Al-Ru-Zr	Al-Si-Zr	Al-Ta-W
C-Co-Re	C-Co-V	C-Cr-Mn	C-Cr-Mo	C-Cr-N
C-Cr-Nb	C-Cr-Ni	C-Cr-Re	C-Cr-Ta	C-Cr-W
C-Fe-Nb	C-Fe-Re	C-Fe-Ta	C-Hf-Nb	C-Hf-Ta
C-Hf-Ti	C-Hf-V	C-Hf-W	C-Mn-V	C-Mo-N
C-Mo-Re	C-Mo-Si	C-Nb-Ta	C-Nb-V	C-Nb-Zr
C-Ni-V	C-N-Nb	C-N-Ti	Co-Cr-Hf	Co-Cr-Mo
Co-Cr-Nb	Co-Cr-Re	Co-Cr-Ru	Co-Cr-Si	Co-Cr-Ta
Co-Cr-V	Co-Cu-Ti	Co-Fe-Hf	Co-Fe-Nb	Co-Fe-Ta
Co-Fe-Ti	Co-Fe-V	Co-Fe-Zr	Co-Hf-Ni	Co-Hf-Si
Co-Hf-Ti	Co-Mo-Nb	Co-Mo-Ni	Co-Mo-Re	Co-Mo-Ru

Tentatively Assessed Ternary Systems

Co-Mo-Ta	Co-Mo-V	Co-Mo-W	Co-Nb-Ni	Co-Nb-Si
Co-Nb-Ta	Co-Nb-Ti	Co-Nb-W	Co-Ni-Re	Co-Ni-Ru
Co-Ni-Ta	Co-Ni-Ti	Co-Ni-Zr	Co-Re-Ta	Co-Re-Ti
Co-Re-V	Co-Re-W	Co-Ru-Ta	Co-Ru-W	Co-Si-Ta
Co-Si-Ti	Co-Si-W	Co-Si-Zr	Co-Ta-W	Co-Ti-W
Co-Ti-Zr	Cr-Cu-Ni	C-Re-Ta	C-Re-V	C-Re-W
Cr-Fe-Hf	Cr-Fe-Nb	Cr-Fe-Re	Cr-Fe-Ta	Cr-Fe-Ti
Cr-Fe-W	Cr-Fe-Zr	Cr-Hf-Mo	Cr-Hf-Nb	Cr-Hf-Ni
Cr-Hf-Re	Cr-Hf-Si	Cr-Hf-Ta	Cr-Hf-W	Cr-Mo-N
Cr-Mo-Re	Cr-Mo-Ru	Cr-Mo-Si	Cr-Mo-Ta	Cr-Mo-W
Cr-Mo-Zr	Cr-Nb-Re	Cr-Nb-Si	Cr-Nb-Ta	Cr-Nb-Ti
Cr-Nb-Zr	Cr-Ni-Ru	Cr-Ni-V	Cr-N-Nb	Cr-N-V
Cr-Re-Ru	Cr-Re-Ta	Cr-Re-Ti	Cr-Re-V	Cr-Re-W
Cr-Re-Zr	Cr-Ru-Ta	Cr-Ru-Ti	Cr-Ru-W	Cr-Si-Ta
Cr-Si-W	Cr-Si-Zr	Cr-Ta-Ti	Cr-Ta-W	Cr-Ta-Zr
Cr-Ti-V	Cr-Ti-W	Cr-W-Zr	C-Si-Ti	C-Ta-Ti
C-Ta-V	C-Ta-Zr	C-Ti-V	C-Ti-Zr	Cu-Ni-Si
C-V-W	C-V-Zr	C-W-Zr	Fe-Hf-Mo	Fe-Hf-Nb
Fe-Hf-Ni	Fe-Hf-Re	Fe-Hf-Si	Fe-Hf-Ta	Fe-Hf-Ti
Fe-Hf-W	Fe-Hf-Zr	Fe-Mn-Ni	Fe-Mo-N	Fe-Mo-Nb
Fe-Mo-Re	Fe-Mo-Si	Fe-Mo-Ta	Fe-Mo-Ti	Fe-Mo-W
Fe-Mo-Zr	Fe-Nb-Re	Fe-Nb-Si	Fe-Nb-Ta	Fe-Nb-Ti

Tentatively Assessed Ternary Systems

Fe-Nb-W	Fe-Nb-Zr	Fe-Ni-Ta	Fe-Ni-V	Fe-Ni-Zr
Fe-Re-Ta	Fe-Re-Ti	Fe-Re-W	Fe-Re-Zr	Fe-Si-Ta
Fe-Si-Ti	Fe-Si-W	Fe-Si-Zr	Fe-Ta-Ti	Fe-Ta-W
Fe-Ta-Zr	Fe-Ti-W	Fe-W-Zr	Hf-Mo-Ni	Hf-Mo-Re
Hf-Mo-Si	Hf-Nb-Ni	Hf-Nb-Re	Hf-Ni-Re	Hf-Ni-Ru
Hf-Ni-Si	Hf-Ni-Ta	Hf-Ni-W	Hf-Re-Ta	Hf-Re-W
Hf-Ru-Ti	Hf-Ru-Zr	Hf-Si-Ta	Hf-Si-Ti	Hf-Si-W
Mn-Ni-V	Mo-Nb-Ni	Mo-Nb-Re	Mo-Nb-Ta	Mo-Ni-Re
Mo-Ni-Ru	Mo-Ni-Si	Mo-Ni-Ti	Mo-Ni-W	Mo-Ni-Zr
Mo-N-V	Mo-Re-Ru	Mo-Re-Ta	Mo-Re-Ti	Mo-Re-V
Mo-Re-W	Mo-Re-Zr	Mo-Ru-Si	Mo-Ru-Ta	Mo-Ru-W
Mo-Si-Zr	Nb-Ni-Re	Nb-Ni-Si	Nb-Ni-Ta	Nb-Ni-V
Nb-Ni-Zr	Nb-Re-Ta	Nb-Re-Ti	Nb-Re-W	Nb-Re-Zr
Nb-Ru-Si	Nb-Si-Ti	Ni-Re-Ta	Ni-Re-Ti	Ni-Re-W
Ni-Re-Zr	Ni-Ru-Ta	Ni-Ru-Ti	Ni-Ru-W	Ni-Ru-Zr
Ni-Si-Ta	Ni-Si-V	Ni-Si-W	Ni-Si-Zr	Ni-Ta-Zr
Ni-W-Zr	N-Nb-Ti	N-Ti-V	Re-Ru-Ta	Re-Ru-Ti
Re-Ru-W	Re-Ta-Ti	Re-Ta-V	Re-Ta-Zr	Re-Ti-W
Re-W-Zr	Ru-Si-Ti	Ru-Ta-Ti	Ru-Ta-W	Ru-Ti-Zr
Si-Ta-Zr	Si-Ti-Zr	Si-W-Zr	Ta-W-Zr	

TCHEA7 Phases

In this section:

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TCHEA7 Models for the Included Phases

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
LIQUID	Liquid					1	(AL, AL1N1, B, C, CO, CR, CU, FE, HF, HF1N1, IR, MN, MO, N, NB, NI, RE, RH, RU, SI, SN, TA, TI, V, W, Y, ZN, ZR)1.0
FCC_A1	Face-Centered Cubic (Cu, A1, fcc)	A1	cF4	(225, Fm-3m)	FCC_A1 will be combined to FCC_L12 if defined.	2	(AL, CO, CR, CU, FE, HF, IR, MN, MO, NB, NI, RE, RH, RU, SI, SN, TA, TI, V, W, Y, ZN, ZR)1.0(B, C, N, VA)1.0
FCC_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)	This phase has some contribution from FCC_A1.	3	(AL, CO, CR, CU, FE, HF, IR, MN, MO, NB, NI, RE, RH, RU, SI, SN, TA, TI, V, W, Y, ZN, ZR)0.75(AL, CO, CR, CU, FE, HF, IR, MN, MO, NB, NI, RE, RH, RU, SI, SN, TA, TI, V, W, Y, ZN, ZR)0.25(B, C, N, VA)1.0
BCC_A2	Body-Centered Cubic (W, A2, bcc)	A2	cI2	(229, Im-3m)	BCC_A2 will be combined to BCC_B2 if defined.	2	(AL, CO, CR, CU, FE, HF, IR, MN, MO, NB, NI, RE, RH, RU, SI, SN, TA, TI, V, VA, W, Y, ZN, ZR)1.0(B, C, N, VA)3.0
BCC_B2	CsCl (B2)	B2	cP2	(221, Pm-3m)	This phase has some contribution from BCC_A2.	3	(AL, CO, CR, CU, FE, HF, IR, MN, MO, NB, NI, RE, RH, RU, SI, SN, TA, TI, V, VA, W, Y, ZN, ZR)0.5(AL, CO, CR, CU, FE, HF, IR, MN, MO, NB, NI, RE, RH, RU, SI, SN, TA, TI, V, VA, W, Y, ZN, ZR)0.5(B, C, N, VA)3.0
HCP_A3	Hexagonal Close Packed (Mg, A3, hcp)	A3	hP2	(194, P6_3/mmc)	Disordered HCP_A3 solution phase.	2	(AL, CO, CR, CU, FE, HF, IR, MN, MO, NB, NI, RE, RH, RU, SI, SN, TA, TI, V, W, Y, ZN, ZR)1.0(B, C, N, VA)0.5
CBCC_A12	alpha-Mn (A12)	A12	cI58	(217, I-43m)		2	(AL, CO, CR, CU, FE, IR, MN, MO, NB, NI, RE, RU, SI, SN, TA, TI, V, W, Y, ZN, ZR)1.0(B, C, VA)1.0
CUB_A13	beta-Mn (A13)	A13	cP20	(213, P4_132)		2	(AL, CO, CR, CU, FE, HF, IR, MN, MO, NB, NI, RE, RU, SI, SN, TA, TI, V, W, Y, ZN, ZR)1.0(B, C, VA)1.0
DIS_FCC_A1	Face-Centered Cubic (Cu, A1, fcc)	A1	cF4	(225, Fm-3m)	A copy of the FCC_A1 phase just for the use in kinetic simulation.	2	(AL, CO, CR, CU, FE, HF, IR, MN, MO, NB, NI, RE, RH, RU, SI, SN, TA, TI, V, W, Y, ZN, ZR)1.0(B, C, N, VA)1.0
DIAMOND_A4	Diamond (A4)	A4	cF8	(227, Fd-3m)	Pure C, Si or solid solution phases based on them.	1	(AL, B, C, SI, SN)1.0
BETA_RHOMBO_B	beta-B (R-105)		hR105	(166, R-3m)		2	(B)93.0(B, C, CU, SI)12.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
GRAPHITE_A9	Hexagonal Graphite (A9)	A9	hP4	(194, P6 ₃ /mmc)		1	(B, C)1.0
BCT_A5	beta-Sn (A5)	A5	tI4	(141, I4 ₁ /amd)	Disordered BCT solution phase.	1	(AL, CU, NI, SN, ZN)1.0
M23B6_TAU_D84	Th6Mn23 (D8a)	D8a	cF116	(225, Fm-3m)	ternary boride.	4	(CO, HF, NI, RE)20.0(B)6.0(B, VA)6.0(AL, CR, HF, MO, RE, TA, TI, V, W, ZR)3.0
M3B_D011	Cementite (Fe3C, D011)	D011	oP16	(62, Pnma)	also NI3B_D011.	2	(CO, CR, FE, MO, NI)3.0(B)1.0
RE3B_E1A	Re3B		oS16	(63, Cmcm)		2	(CR, MO, RE, TA, W)3.0(B)1.0
M7B3_D102	Fe3Th7 (D102)	D102	hP20	(186, P6 ₃ mc)	also RH7B3, RE7B3_D102.	3	(CO, CR, MO, NB, RE, RH, RU, TA, W)7.0(B)3.0(B, VA)3.0
M2B_C16	Khatyrkite (Al2Cu, C16)	C16	tI12	(140, I4/mcm)	aka M2B_TETR.	2	(AL, CO, CR, FE, MN, MO, NB, NI, RE, TA, W)2.0(B)1.0
CR2B_CB	Mg2Cu (Cb)	Cb	oF48	(70, Fddd)	aka CR2B_ORTH.	2	(CR, FE, MO, RE)0.66666667(B)0.33333333
MN2B_D1F	Mg2Cu (Cb)	Cb	oF48	(70, Fddd)		2	(MN)0.6707(B)0.3293
CR5B3_D8I	Cr5B3 (D8I)	D8I	tI32	(140, I4/mcm)	also Mo5B3.	2	(CR, MO)0.625(B)0.375
M3B2_D5A	Si2U3 (D5a)	D5a	tP10	(127, P4/mbm)	also NB3B2, TA3B2, V3B2.	2	(FE, HF, MO, NB, TA, V)3.0(B)2.0
M3B2_T	Si2U3 (D5a)	D5a	tP10	(127, P4/mbm)	ternary boride Mo2FeB2 or Mo2CrB2.	3	(CR, FE, MO, NI, W)0.4(CR, FE, NI)0.2(B)0.4
IR4B3	Ir4B3		oF28	(42, Fmm2)		2	(IR)4.0(B)3.0
NI4B3	m-Ni4B3		mS28	(15, C2/c)		2	(NI)0.57142857(B)0.42857143
IR5B4	Ir5B4		tI36	(88, I4 ₁ /a)		2	(IR)5.0(B)4.0
MB_B27	FeB (B27)	B27	oP8	(62, Pnma)	also CoB, HfB, MnB, TiB.	2	(B)1.0(CO, CR, FE, HF, MN, MO, RE, TI, Y)1.0
MB_B33	CrB (B33)	B33	oS8	(63, Cmcm)	also NbB, NiB, TaB, VB.	2	(CR, FE, HF, MO, NB, NI, TA, TI, V)1.0(B)1.0

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MOB_BG	MoB (Bg)	Bg	tI16	(141, I4_1/amd)		2	(CR, FE, MO)1.0(B)1.0
WB_ALPHA_BG	MoB (Bg)	Bg	tI16	(141, I4_1/amd)		2	(B, C, VA)1.0(W)1.0
RU1B1	Unknown Structure		cl*			2	(RU)1.0(B)1.0
WB_BETA_B33	CrB (B33)	B33	oS8	(63, Cmcm)		2	(B, C, VA)1.0(W)1.0
RHB_B81	NiAs (B81)	B81	hP4	(194, P6_3/mmc)		2	(RH)1.0(B)1.1
M5B6	V5B6		oS22	(65, Cmmm)	also Nb5B6.	2	(NB, V)5.0(B)6.0
IR4B5	Ir4B5		mS18	(8, Cm)		2	(IR)4.0(B)5.0
M3B4_D7B	Ta3B4 (D7b)	D7b	oI14	(71, Immm)	also ternary X1Y2B4 boride.	2	(B)4.0(AL, CR, HF, MN, NB, TA, TI, V)3.0
RU2B3	Ru2B3		hP10	(194, P6_3/mmc)		2	(RU)2.0(B)3.0
V2B3	V2B3		oS20	(63, Cmcm)		2	(V)0.4(B)0.6
MB2_C32	Hexagonal omega (C32)	C32	hP3	(191, P6/mmm)		2	(B)2.0(AL, CR, HF, MN, MO, NB, RU, TA, TI, V, Y, ZR)1.0
REB2	ReB2		hP6	(194, P6_3/mmc)		3	(RE)1.0(B)2.0(B, VA)2.0
MO2B5_D8i	Mo2B5 (D8i)	D8i	hR7	(166, R-3m)		2	(MO)0.32(B)0.68
W2B5_D8i	Mo2B5 (D8i)	D8i	hR7	(166, R-3m)	nonstoichiometric W2B5.	2	(B, C, VA)5.0(W)2.0
CRB4	CrB4		oI10	(71, Immm)		2	(CR)0.2(B)0.8
MNB4	MnB4		mS10	(12, C2/m)		2	(MN)0.2(B)0.8

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MOB4	MoB4		hP16	(194, P6 ₃ /mmc)		2	(MO)0.2(B)0.8
YB4_D1E	ThB4 (D1e)	D1e	tP20	(127, P4/mbm)		2	(Y)0.2(B)0.8
W2B9	W2B9		hP22	(147, P-3)		2	(B)9.0(W)2.0
YB6_D21	CaB6 (D21)	D21	cP7	(221, Pm-3m)		2	(Y)1.0(B)6.0
ZRB12_D2F	UB12 (D2f)	D2f	cF52	(225, Fm-3m)	also YB12.	2	(B)12.0(Y, ZR)1.0
ALB12_ALPHA	alpha-ALB12		tP216	(92, P4 ₁₂ -12)		2	(AL, TI)1.0(B)12.0
YB66	YB66		cF1936	(226, Fm-3c)		2	(Y)1.0(B)66.0
M12C	Fe6W6C		cF104	(227, Fd-3m)	also Mo6Ni6C	3	(CO, NI)6.0(MO, W)6.0(C)1.0
M6C_E93	Fe3W3C (E93)	E93	cF112	(227, Fd-3m)		4	(CO, FE, NI)2.0(MO, NB, TA, W)2.0(CO, CR, FE, MO, NB, NI, TA, V)2.0(C)1.0
ALM3C_E21	Cubic Perovskite (CaTiO ₃ , E21)	E21	cP5	(221, Pm-3m)	also AlCo3C, AlFe3C.	3	(AL)1.0(CO, FE)3.0(C)1.0
TI3ALC_E21	Cubic Perovskite (CaTiO ₃ , E21)	E21	cP5	(221, Pm-3m)	i.e. Ti3AlC1-x.	3	(TI)3.0(AL)1.0(C, VA)1.0
W3MC	W10Co3C3.4		hP34	(194, P6 ₃ /mmc)	also W3CoC, W3NiC.	3	(W)3.0(CO, NI)1.0(C)1.0
M23C6_D84	Cr23C6 (D84)	D84	cF116	(225, Fm-3m)		3	(CO, CR, FE, MN, NI, RE, V)20.0(CO, CR, FE, MN, MO, NI, RE, V)3.0(C)6.0
ALCR2C	AlCr2		hP8	(194, P6 ₃ /mmc)	MAX_PHASE.	3	(AL)1.0(C)1.0(CR)2.0
CEMENTITE_D011	Cementite (Fe3C, D011)	D011	oP16	(62, Pnma)		2	(CO, CR, FE, MN, MO, NI, V, W)3.0(C, N)1.0

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Ti2ALC	AlCr2		hP8	(194, P6 ₃ /mmc)	i.e. Ti2AlC1-x.	3	(Ti)2.0(Al)1.0(C, VA)1.0
M5C2	Mn5C2 (Fe5C2 Hagg carbide)		mS28	(15, C2/c)		2	(Fe, Mn)5.0(C)2.0
M7C3_D101	C3Cr7 (D101)	D101	oP40	(62, Pnma)		2	(Co, Cr, Fe, Mn, Mo, Ni, Re, V, W)7.0(C)3.0
Ti3ALC2	Ti3SiC2		hP12	(194, P6 ₃ /mmc)		3	(Ti)3.0(Al, Si)1.0(C, VA)2.0
M3C2_D510	Tungbaite (Cr3C2, D510)	D510	oP20	(62, Pnma)		2	(Co, Cr, Mo, V, W)3.0(C)2.0
Al4C3_D71	Al4C3 (D71)	D71	hR7	(166, R-3m)		2	(Al, Si)4.0(C)3.0
MC_ETA	CMo		hP12	(194, P6 ₃ /mmc)		2	(Mo, V, W)1.0(C, VA)1.0
MC_SHP	Tungsten Carbide (Bh)	Bh	hP2	(187, P-6m2)	also MoC_LT, Strukturbericht Bh	2	(Mo, W)1.0(C, N)1.0
YC_GAMMA_B1	Rock Salt (NaCl, B1)	B1	cF8	(225, Fm-3m)		2	(Y)1.0(C, C2, VA)1.0
Y15C19_BETA	Unknown Structure				the high temperature beta phase	2	(C)19.0(Y)15.0
Y15C19_ALPHA	alpha-Y15C19		oP18	(55, Pbam)	aka Y15C19_R.	2	(C)19.0(Y)15.0
Y2C3_BETA	Unknown Structure				HT phase	3	(Y)2.0(C)2.0(C, VA)1.0
Y2C3_ALPHA	Sc3C4		tP70	(128, P4/mnc)	aka Y2C3_R.	3	(Y)2.0(C)2.0(C, VA)1.0
YC2_C11A	CaC2-I (C11a)	C11a	tI6	(139, I4/mmm)		1	(C2Y1)1.0
PI_A13	beta-Mn (A13)	A13	cP20	(213, P4 ₁₃₂)	i.e. Cr-(Fe,Ni)-N nitride.	3	(Cr)12.8(Fe, Ni)7.2(N)4.0
FE4N_LP1	gama-Fe4N (L'10)	L'10	cP5	(221, Pm-3m)	Only stable in (Co-Cr-Fe)-N when gas suspended.	2	(Co, Cr, Fe, Mn, Ni)4.0(C, N)1.0

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ALTi3N_E21	Cubic Perovskite (CaTiO ₃ , E21)	E21	cP5	(221, Pm-3m)		3	(AL)1.0(Ti)3.0(N)1.0
ALTi2N	AICr2		hP8	(194, P6_3/mmc)		3	(AL)1.0(Ti)2.0(N)1.0
AL2Ti3N2	(Al2Ti3N2)		hP22	(186, P6_3mc)		3	(AL)2.0(Ti)3.0(N)2.0
FEcN_CHI	Mn5C2 (Fe5C2 Hagg carbide)		mS28	(15, C2/c)	Only stable in C-Fe-N when gas suspended.	2	(FE)2.2(C, N)1.0
Ti4N3	Sc2Te3		hR8	(166, R-3m)	aka Xi-TiN0.58 HT	2	(Ti)0.685(N)0.315
Ti2N_C4	Rutile (TiO ₂ , C4)	C4	tP6	(136, P4_2/mnm)		2	(Ti)2.0(N)1.0
Z_PHASE	CrNbN		tP6	(129, P4/nmm)		3	(CR, FE)1.0(MO, NB, V)1.0(N, VA)1.0
HF3N2	TiS-9R		hR6	(166, R-3m)		2	(HF)3.0(N)2.0
MN3N2	Mn3N2		tI10	(139, I4/mmm)	aka MN6N4.	2	(MN)6.0(N)4.0
Ti3N2	TiS-9R		hR6	(166, R-3m)		2	(Ti)0.71(N)0.29
HF4N3	Sc2Te3		hR8	(166, R-3m)		2	(HF)4.0(N)3.0
MN6N5	CoO		tI4	(139, I4/mmm)		2	(MN)6.0(N)5.0
ALN_B4	Wurtzite (ZnS, B4)	B4	hP4	(186, P6_3mc)		2	(AL)1.0(N)1.0
TAN_EPS	TaN-eps		hP6	(189, P-62m)		2	(TA)1.0(N)1.0
HFN_B1	Rock Salt (NaCl, B1)	B1	cF8	(225, Fm-3m)		2	(HF)1.0(N, VA)1.0
MN6Si	Fe7W6 (D85) mu-phase	D85	hR13	(166, R-3m)		2	(AL, MN)17.0(Si, ZN)3.0

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MN9SI2	Mn9Si2		oI186	(71, Immm)		2	(MN)33.0(SI)7.0
CR3NI5SI2	AlAu4		cP20	(198, P2_13)		4	(CR)3.0(NI)5.0(SI)2.0(C, VA)1.0
CR2NI2SI	NiTi2		cF96	(227, Fd-3m)	i.e. the Tau1 phase Cr5Ni5Si3.	3	(CR)5.0(NI)5.0(SI)3.0
CR3SI_A15	Cr3Si (A15)	A15	cP8	(223, Pm-3n)	also Cr3X, Nb3X, Ti3X, V3X.	3	(CR, FE, IR, MO, NB, NI, RE, RH, SI, SN, TA, TI, V, ZR)3.0(AL, CO, CR, IR, NB, NI, RH, RU, SI, SN, TA, TI, V, ZR)1.0(C, VA)3.0
CO3SI_D019	Ni3Sn (D019)	D019	hP8	(194, P6_3/mmc)		2	(CO)3.0(SI)1.0
IR3SI1	SiU3 (D0c)	D0c	tI16	(140, I4/mcm)		2	(IR)3.0(SI)1.0
M3SI1	Ti3P		tP32	(86, P4_2/n)	also Nb3Si, Ta3Si, Ti3Si, Zr3Si.	2	(HF, NB, TA, TI, ZR)3.0(SI)1.0
MN3SI_D03	BiF3 (D03)	D03	cF16	(225, Fm-3m)		2	(FE, MN)3.0(AL, SI)1.0
NI3SI_ORTHO_D011	Cementite (Fe3C, D011)	D011	oP16	(62, Pnma)		2	(NI)3.0(SI)1.0
NI3SI_MONOCL	Ge9Pd25		hP34	(147, P-3)		2	(NI)3.0(SI)1.0
NI5SI2	Ni31Si12		hP42	(150, P321)		2	(CO, CR, CU, FE, NI)5.0(SI)2.0
CO2SI1	Co2Si (C37)	C37	oP12	(62, Pnma)	also Ni2Si (delta)	2	(CO, CR, CU, FE, IR, NI, RH, TI)2.0(SI)1.0
CRNBSI	ZrNiAl		hP9	(189, P-62m)		3	(CR)1.0(NB)1.0(SI)1.0
FE2SI	AlNi2		hP6	(164, P-3m1)		2	(FE)0.666667(SI)0.333333
NI2SI_THETA	AlNi2		hP6	(164, P-3m1)	aka M2SI_TETA.	3	(CU, NI)1.0(NI, VA)1.0(AL, SI)1.0
RE2SI	Re2Si		mP24	(14, P2_1/c)		2	(RE)2.0(SI)1.0
RU2SI_C37	Co2Si (C37)	C37	oP12	(62, Pnma)		2	(RU)2.0(SI)1.0
M5SI3_D88	Mavlyanovite (Mn5Si3, D88)	D88	hP16	(193, P6_3/mmc)	also M5Sn3, M5Si3C.	4	(CR, CU, FE, HF, MN, MO, NB, NI, SI, TI, W, Y, ZR)2.0(AL, CR, SI, SN, TI)3.0(CR, CU, FE, HF, MN, MO, NB, NI, TI, Y, ZR)3.0(C,

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							SN, VA)1.0
RH5Si3	Rh5Ge3		oP16	(55, Pbam)		2	(RH)5.0(SI)3.0
TA5Si3_D8L	Cr5B3 (D8l)	D8l	tI32	(140, I4/mcm)	also alpha-Nb5Si3.	2	(HF, NB, TA)5.0(AL, SI)3.0
W5Si3_D8M	W5Si3 (D8m)	D8m	tI32	(140, I4/mcm)	also Cr5Si3, Mo5Si3, Nb5Si3, V5Si3.	3	(CR, FE, MO, NB, TI, V, W)4.0(CR, FE, MO, NB, SI, TI, V, W)1.0 (AL, SI, SN)3.0
RH20Si13	Rh20Si13		hP34	(176, P6_3/m)		2	(RH)20.0(SI)13.0
IR3Si2_B82	Ni2In (B82)	B82	hP6	(194, P6_3/mmc)		2	(IR)3.0(SI)2.0
M3Si2_D5A	Si2U3 (D5a)	D5a	tP10	(127, P4/mbm)	also Hf3Si2, Zr3Si2.	2	(HF, NB, ZR)3.0(SI)2.0
NI3Si2	Ni3Si2		oP80	(36, Cmc2_1)		2	(NI)3.0(SI)2.0
M11Si8	Cr11Ge8		oP76	(62, Pnma)	also Cr11Si8, Nb11Si8.	2	(CR, NB)11.0(SI)8.0
M4Si3	Ru4Si3		oP28	(62, Pnma)	also Cr4Si3, Nb4Si3.	2	(CR, NI, RU)4.0(SI)3.0
ZR5Si4_TP36	Si4Zr5		tP36	(92, P4_12_12)	also Hf5Si4, Ti5Si4.	2	(HF, NB, TI, ZR)5.0(AL, SI)4.0
Y5Si4	Gd5Si4		oP36	(62, Pnma)		2	(Y)5.0(SI)4.0
M6Si5	Si5V6		oI44	(72, Ibam)	also Cr6Si5, Nb6Si5.	2	(CR, NB, TI, V)6.0(SI)5.0
MSI_B20	FeSi (B20)	B20	cP8	(198, P2_13)	also CoSi, CrSi, MnSi, ReSi.	2	(CO, CR, FE, MN, NI, RE)1.0(AL, SI)1.0
IRS_B27	FeB (B27)	B27	oP8	(62, Pnma)	the HT phase.	2	(IR)1.0(SI)1.0
MSI_B27	FeB (B27)	B27	oP8	(62, Pnma)	also TiSi, HfSi, YSi, ZrSi(alpha).	2	(HF, NB, TI, Y, ZR)1.0(AL, SI)1.0
NISI_B31	MnP (B31)	B31	oP8	(62, Pnma)		2	(NI)1.0(SI)1.0

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RHSI_B31	MnP (B31)	B31	oP8	(62, Pnma)		2	(RH)1.0(SI)1.0
RHSI_B20	FeSi (B20)	B20	cP8	(198, P2_13)		2	(RH)1.0(SI)1.0
RUSI	FeSi (B20)	B20	cP8	(198, P2_13)	united HT_B2/LT_B20 phase.	2	(RU)1.0(SI)1.0
IR4Si5	Ru4Si5		mP18	(11, P2_1/m)		2	(IR)4.0(SI)5.0
RH4Si5	Ru4Si5		mP18	(11, P2_1/m)		2	(RH)4.0(SI)5.0
IR3Si4	Ru4Si3		oP28	(62, Pnma)		2	(IR)3.0(SI)4.0
RH3Si4	Rh3Si4		oP28	(62, Pnma)		2	(RH)3.0(SI)4.0
RU2Si3	Ge3Ru2		oP40	(60, Pbcn)	united HT/LT phase.	2	(RU)2.0(SI)3.0
IR3Si5	Ir3Si5		mP64	(14, P2_1/c)		2	(IR)3.0(SI)5.0
Y3Si5_HT_CC	alpha-ThSi2 (Cc)	Cc	tI12	(141, I4_1/amd)		2	(Y)3.0(SI)5.0
Y3Si5_LT_C32	Hexagonal omega (C32)	C32	hP3	(191, P6/mmm)		2	(Y)3.0(SI)5.0
MN11Si19	Mn11Si19		tP120	(118, P-4n2)		2	(MN)11.0(AL, SI)19.0
CRSi2_C40	CrSi2 (C40)	C40	hP9	(180, P6_222)	also NbSi2, TaSi2, VSi2.	2	(CR, CU, HF, MO, NB, SI, TA, TI, V, W)1.0(AL, CR, CU, SI)2.0
FESI2_HT	FeSi2-h		tP3	(123, P4/mmm)		2	(FE)0.3(SI)0.7
FESI2_LT	FeSi2-l		oS48	(64, Cmce)		2	(FE)0.333333(SI)0.666667
MSI2_C1	Fluorite (CaF2, C1)	C1	cF12	(225, Fm-3m)	also NiSi2, CoSi2.	2	(CO, CU, MN, NI)1.0(AL, CU, SI)2.0
MOSi2_C11B	MoSi2 (C11b)	C11b	tI6	(139, I4/mmm)	also AlCr2, Ti2Zn, ZnZr2, CUZR2_C11B.	2	(AL, CO, CU, FE, MO, NI, RH, W, ZN)1.0(AL, CR, HF, SI, TI, ZR)2.0

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RESi2_C11B	Re4Si7		mS44	(8, Cm)		2	(RE)0.357(SI)0.643
TiSi2_C54	TiSi2 (C54)	C54	oF24	(70, Fddd)	also MoSi2, RuAl2, ZrSn2.	2	(MO, NB, RU, TI, ZR)1.0(AL, SI, SN)2.0
YSi2_HT_CC	alpha-ThSi2 (Cc)	Cc	tI12	(141, I4_1/amd)		2	(Y)1.0(SI)2.0
YSi2_LT_C32	Hexagonal omega (C32)	C32	hP3	(191, P6/mmm)		2	(Y)1.0(SI)2.0
ZRSi2_C49	ZrSi2 (C49)	C49	oS12	(63, Cmcn)	also HfSi2.	2	(HF, NB, ZR)1.0(SI)2.0
IRSi3_LT	Unknown Structure		mS*			2	(IR)1.0(SI)3.0
IRSi3_HT	Unknown Structure		oS*			2	(IR)1.0(SI)3.0
ALFESI_ALPHA_TAU5	Fe23Al81Si15		hP246	(194, P6_3/mmc)		4	(AL)0.6612(Fe)0.19(SI)0.0496(AL, SI)0.0992
ALFESI_BETA_TAU6	Fe2Al9Si2		mS52	(15, C2/c)		3	(AL)14.0(Fe)3.0(SI)3.0
ALFESI_GAMMA_TAU2	Unknown Structure		mS*			3	(AL)3.0(Fe)1.0(SI)1.0
ALFESI_DELTA_TAU4	FeAl3Si2		oP24	(60, Pbcn)		3	(AL)0.55(Fe)0.15(SI)0.3
ALFESI_TAU1	Unknown Structure					3	(AL)2.0(Fe)2.0(SI)1.0
ALFESI_TAU3	Fe(Al0.67Si0.33)3		oS128	(67, Cmme)		3	(AL)2.0(Fe)1.0(SI)1.0
AL2MN2SI3	(Al2Mn2Si3)		hP21	(174, P-6)	Tau1	3	(AL)2.0(MN)2.0(SI)3.0
AL5MN6SI7	CrSi2 (C40)	C40	hP9	(180, P6_222)	Tau2	3	(AL)5.0(MN)6.0(SI)7.0
AL1MN1SI1	TiSi2 (C54)	C54	oF24	(70, Fddd)	Tau3	3	(AL)1.0(MN)1.0(SI)1.0
AL3MNSI2	(Al3MnSi2)		tP48	(85, P4/n)	Tau4	3	(AL)3.0(MN)1.0(SI)2.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
AL3MN4Si2	Unknown Structure				ternary tau5 or D phase	3	(AL)3.0(MN)4.0(Si)2.0
ALMNSI_TAU6	Unknown Structure					2	(AL, MN)4.0(Si)1.0
ALMNSI_TAU8	Al9Mn3Si (E9c)	E9c	hP26	(194, P6_3/mmc)		5	(MN, VA)6.0(MN, VA)2.0(AL)12.0(AL, Si)6.0(AL, Si)2.0
AL2MNSi3	Ga5Pd		tI24	(140, I4/mcm)	Tau10	3	(AL)2.0(MN)1.0(Si)3.0
MN15Ni45Si40	Unknown Structure				ternary T1 or N phase	3	(MN)0.15(Ni)0.45(Si)0.4
MN15Ni50Si35	Unknown Structure				ternary T2 or PHI phase	3	(MN)0.15(Ni)0.5(Si)0.35
MN6Ni16Si7	Th6Mn23 (D8a)	D8a	cF116	(225, Fm-3m)	ternary T3 or G phase	3	(MN)0.206897(Ni)0.551724(Si)0.241379
MN1Ni1Si1_C37	MnCuP		oP12	(62, Pnma)	ternary T4 or E phase	3	(MN)1.0(Ni)1.0(Si)1.0
MNNiSi_T5	MgZn2 Hexagonal Laves (C14)	C14	hP12	(194, P6_3/mmc)	ternary T5 or tao_1 phase	2	(MN)1.0(Ni, Si)2.0
MNNiSi_T6	Cu2Mg Cubic Laves (C15)	C15	cF24	(227, Fd-3m)	ternary T6 or tao_2 phase	2	(MN)1.0(Ni, Si)2.0
MN3Ni2Si	Mn3Ni2Si		cF96	(227, Fd-3m)	ternary T7 or Omega	3	(MN)3.0(Ni)2.0(Si)1.0
MN2NiSi	Unknown Structure				ternary T8 or S phase	2	(MN, Ni)3.0(Si)1.0
MN6NiSi3	R-(Co,Cr,Mo)		hR53	(148, R-3)	ternary T9 or R' phase	3	(MN)0.61(Ni)0.12(Si)0.27
MN66Ni4Si30	Unknown Structure				ternary T10 or U phase	3	(MN)0.66(Ni)0.04(Si)0.3
MN52Ni29Si19	Unknown Structure				ternary phase T11 or W phase	3	(MN)0.52(Ni)0.29(Si)0.19
B4C_D1G	B13C2 B4C (D1g)	D1g	hR15	(166, R-3m)		2	(B11C, B12)1.0(B2, C2B, CB2)1.0
BN_B4	Wurtzite (ZnS, B4)	B4	hP4	(186, P6_3mc)		2	(B)1.0(N)1.0
SIB3_D1G	B13C2 B4C (D1g)	D1g	hR15	(166, R-3m)		3	(B)6.0(Si)2.0(B, Si)6.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
SIB6	SiB6		oP280	(58, Pnm)		3	(B)210.0(Si)23.0(B, Si)48.0
BNSI_RHOMBO	alpha-B (hR12)		hR12	(166, R-3m)	aka BNSI, ALPHA_RHOMBO_B.	3	(B)61.0(Si)1.0(B, Si)8.0
SIC_B3	Zincblende (ZnS, B3)	B3	cF8	(216, F-43m)		2	(Si)1.0(C)1.0
SI3N4	Nierite (alpha-Si3N4)		hP28	(159, P31c)		2	(Si)3.0(N)4.0
AL4SiC4_E94	Al5C3N (E94)	E94	hP18	(186, P6_3mc)		3	(Al)4.0(Si)1.0(C)4.0
AL8SiC7	Unknown Structure		hP16			3	(Al)8.0(Si)1.0(C)7.0
FE8Si2C	Mn8Si2C		aP*	(1, P1)		3	(Fe)8.0(Si)2.0(C)1.0
TI3SiC2	Ti3SiC2		hP12	(194, P6_3/mmc)		3	(Ti)3.0(Si)1.0(C)2.0
DIS_MU	Fe7W6 (D85) mu-phase	D85	hR13	(166, R-3m)	Part of the description for the MU_PHASE	1	(AL, CO, CR, CU, FE, MN, MO, NB, NI, RE, TA, TI, W)1.0
MU_PHASE	Fe7W6 (D85) mu-phase	D85	hR13	(166, R-3m)	DIS_MU contribution added onto it.	4	(AL, CO, CR, CU, FE, MN, MO, NB, NI, RE, TA, TI, W)1.0(AL, CO, CR, CU, FE, MN, MO, NB, NI, RE, TA, TI, W)2.0(AL, CO, CR, CU, FE, MN, MO, NB, NI, RE, TA, TI, W)6.0(AL, CO, CR, CU, FE, MN, MO, NB, NI, RE, TA, TI, W)4.0
DIS_SIG	sigma-CrFe (D8b)	D8b	tP30	(136, P4_2/mnm)	Part of the description for the SIGMA phase.	1	(AL, CO, CR, FE, IR, MN, MO, NB, NI, RE, RH, RU, SI, TA, TI, V, W)1.0
SIGMA	sigma-CrFe (D8b)	D8b	tP30	(136, P4_2/mnm)	DIS_SIG contribution added onto it.	3	(AL, CO, CR, FE, IR, MN, MO, NB, NI, RE, RH, RU, SI, TA, TI, V, W)10.0(AL, CO, CR, FE, IR, MN, MO, NB, NI, RE, RH, RU, SI, TA, TI, V, W)4.0(AL, CO, CR, FE, IR, MN, MO, NB, NI, RE, RH, RU, SI, TA, TI, V, W)16.0
CHI_A12	alpha-Mn (A12)	A12	cI58	(217, I-43m)	also M5Re24, Mo2Re8, Ta3Re7, WRe3.	3	(CR, FE, NI, RE)24.0(AL, CR, HF, MO, NB, TA, TI, W, ZR)10.0(CR, FE, MO, NB, NI, RE, TA, W)24.0
G_PHASE_D8A	Th6Mn23 (D8a)	D8a	cF116	(225, Fm-3m)		3	(AL, CO, FE, MN, NI, TI)16.0(HF, NB, TI, Y, ZR)6.0(CO, FE, MN, NI, SI)7.0
GAMMA_D83	gamma-brass (Cu9Al4,	D83	cP52	(215, P-43m)	also Cu5Zn8, Ni5Zn8, Al5Cu8	3	(AL, NI, SI, ZN)4.0(AL, CU, NI, SI, ZN)1.0(CU, FE, MN, NI,

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	D83)				(rt).		ZN)8.0
GAMMA_HT_D82	gamma-brass (Cu5Zn8, D82)	D82	cI52	(217, I-43m)	aka GAMMA_H.	3	(AL, ZN)4.0(AL, CU, ZN)1.0(CU, FE, MN, NI)8.0
HEUSLER_L21	Heusler (L21)	L21	cF16	(225, Fm-3m)	aka H_L21.	3	(AL, CR, NI, SN, TI)0.5(AL, HF, NB, NI, TA, TI, ZR)0.5(CO, FE, NI, RU, VA)1.0
O_PHASE	NaHg		oS16	(63, Cmcmm)	aka Ti2NbAl, the O phase.	3	(NB, TA, TI)0.5(AL, NB, TA, TI)0.25(NB, TA, TI)0.25
P_PHASE	Cr9Mo21Ni20		oP56	(62, Pnma)		3	(CR, FE, NI, RE)24.0(CR, FE, MO, NI, RE)20.0(MO)12.0
R_PHASE	R-(Co,Cr,Mo)		hR53	(148, R-3)		3	(CO, CR, FE, NI, RE)27.0(MO, W)14.0(CO, CR, FE, MO, NI, RE, W)12.0
B82_OMEGA	Ni2In (B82)	B82	hP6	(194, P6_3/mmc)		3	(AL, SN)1.0(CO, NB, SN, TA, TI)1.0(TI)1.0
MNNI_L10	CuAu (L10)	L10	tP2	(123, P4/mmm)		2	(CO, CR, CU, FE, MN, NI)0.5(CO, CR, CU, FE, MN, NI)0.5
IRMN_L10	CuAu (L10)	L10	tP2	(123, P4/mmm)	also IrTi.	2	(IR, MN, TI)0.5(IR, MN, TI)0.5
ALTI_L10	CuAu (L10)	L10	tP2	(123, P4/mmm)	Solid solution of ordered L10.	2	(AL, CO, CR, MN, MO, NB, SN, TA, TI, V, W, ZR)1.0(AL, CO, CR, MN, MO, NB, SN, TA, TI, V, W, ZR)1.0
IRNB_L10	CuAu (L10)	L10	tP2	(123, P4/mmm)		2	(IR, NB)0.5(IR, NB)0.5
IRW_EPSILON_HCP	Hexagonal Close Packed (Mg, A3, hcp)	A3	hP2	(194, P6_3/mmc)	HT epsilon solution phase with HCP structure.	2	(IR, W)1.0(IR, W)1.0
IRW_LT_B19	beta'-AuCd (B19)	B19	oP4	(51, Pmma)		2	(IR)1.0(IR, W)1.0
IRV_L10	CuAu (L10)	L10	tP2	(123, P4/mmm)	aka Ir1.04V0.96 or IrV1-x.	2	(IR)0.5(IR, V)0.5
NBRH_L10	CuAu (L10)	L10	tP2	(123, P4/mmm)	aka Rh1.04V0.96.	2	(NB)0.96(RH)1.04

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RHV_L10	CuAu (L10)	L10	tP2	(123, P4/mmm)	aka Rh1.12V0.88.	2	(RH, V)1.12(RH, V)0.88
ALCU_EPS_B82	Ni2In (B82)	B82	hP6	(194, P6 ₃ /mmc)	epsilon2 HT.	2	(AL, CU, NI)1.0(CU, FE)1.0
ALCU_ETA	AlCu(r)		mS20	(12, C2/m)	united HT-eta1 & LT-eta2.	2	(AL, CU)1.0(CU, FE, NI, ZN)1.0
ALPHA_B19	beta'-AuCd (B19)	B19	oP4	(51, Pmma)	an ordered HCP.	2	(MO, NB, TI, V, ZR)1.0(MO, NB, TI, V, ZR)1.0
ALRE_B11	gamma-CuTi (B11)	B11	tP4	(129, P4/nmm)		2	(AL)1.0(RE)1.0
ALZR_B33	CrB (B33)	B33	oS8	(63, Cmcmm)	also AlHf, ALY_B33.	2	(AL)1.0(HF, Y, ZR)1.0
COSN_B35	CoSn (B35)	B35	hP6	(191, P6/mmm)	also FeSn, COSN_HP6.	2	(CO, FE, NI)0.5(SN)0.5
COY_B33	CrB (B33)	B33	oS8	(63, Cmcmm)	aka COY_BF.	2	(CO)1.0(Y)1.0
CUTI_B11	gamma-CuTi (B11)	B11	tP4	(129, P4/nmm)	aka B11.	2	(CO, CU, NI, TI)1.0(CU, NI, TA, TI)1.0
HF1IR1	Unknown Structure				united HT/LT phase	2	(HF)1.0(IR)1.0
HFMN	NiTi2		cF96	(227, Fd-3m)		2	(HF)0.5(MN)0.5
HFNI_ALPHA	CrB (B33)	B33	oS8	(63, Cmcmm)		2	(HF)0.5(NI)0.5
HFRE	Zr21Re25		hR92	(167, R-3c)	also Hf21Re25.	2	(HF)1.0(RE)1.0
HFRH_B2	CsCl (B2)	B2	cP2	(221, Pm-3m)		2	(HF, RH)1.0(RH)1.0
IRV_RT	alpha-IrV		oS8	(65, Cmmm)		2	(IR)1.0(V)1.0
IRZR_BETA_B2	CsCl (B2)	B2	cP2	(221, Pm-3m)	i.e. HT- or beta- phase.	2	(IR, ZR)1.0(IR, ZR)1.0
IRZR_ALPHA	Mn3As (D0d)	D0d	oS16	(63, Cmcmm)	LT phase.	2	(IR, ZR)1.0(ZR)1.0

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MNTA	Unknown Structure					2	(MN)1.0(TA)1.0
MNTI_LT	Zr ₂₁ Re ₂₅		hR92	(167, R-3c)	aka Ti ₂₁ Mn ₂₅ _RT.	2	(MN)1.0(TI)1.0
MOIR_HT	Hexagonal Close Packed (Mg, A3, hcp)	A3	hP2	(194, P6 ₃ /mmc)		2	(IR, MO)1.0(IR, MO)1.0
MOIR_LT_B19	beta'-AuCd (B19)	B19	oP4	(51, Pmma)		2	(MO)1.0(IR)1.0
NBRH_HT	Unknown Structure					2	(NB, RH)1.0(NB, RH)1.0
NB1ZN1	Unknown Structure					2	(NB)0.5(ZN)0.5
NIZN_TP2	CuAu (L10)	L10	tP2	(123, P4/mmm)	united HT/LT phase.	2	(AL, CU, NI, ZN)0.5(AL, NI, ZN)0.5
NIY_B27	FeB (B27)	B27	oP8	(62, Pnma)		2	(NI)1.0(Y)1.0
NIZR_B33	CrB (B33)	B33	oS8	(63, Cmcmm)		2	(NI)1.0(TI, ZR)1.0
RHSN_B20	FeSi (B20)	B20	cP8	(198, P2 ₁₃)		2	(RH)1.0(SN)1.0
RHZR_LT	IrZr		oS16	(63, Cmcmm)	i.e. alpha_RhZr	2	(RH)1.0(RH, ZR)1.0
RHV_RT	alpha-IrV		oS8	(65, Cmmm)		2	(RH)1.0(V)1.0
RHV_HT	Unknown Structure					2	(RH, V)1.0(V)1.0
RHZR_HT_B2	CsCl (B2)	B2	cP2	(221, Pm-3m)	i.e. beta_RhZr.	2	(RH)1.0(RH, ZR)1.0
ZNZR_B2	CsCl (B2)	B2	cP2	(221, Pm-3m)		2	(ZN)1.0(ZR)1.0
ALCU_PRIME	Al ₉ Cu ₁₁ (h)		oF88	(42, Fmm2)	aka THETA_PRIME.	2	(AL)2.0(CU)1.0
AL2FE	Al ₂ Fe		aP18	(1, P1)		2	(AL, CU)2.0(Fe, MN)1.0
ALRE2_C11B	MoSi ₂ (C11b)	C11b	tI6	(139, I4/mmm)		2	(AL)1.0(RE)2.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
ALRH2	Unknown Structure					2	(AL)1.0(RH)2.0
AL2TI_LT	Ga2Hf		tI24	(141, I4_1/amd)		2	(AL, NB, TI)2.0(AL, CO, NB, TA, TI, V, ZR)1.0
Al2W_C40	CrSi2 (C40)	C40	hP9	(180, P6_222)		2	(AL)2.0(W)1.0
ALY2_C37	Co2Si (C37)	C37	oP12	(62, Pnma)		2	(AL)1.0(Y)2.0
ALZR2_B82	Ni2In (B82)	B82	hP6	(194, P6_3/mmc)		2	(AL)1.0(TI, ZR)2.0
C14_LAVES	MgZn2 Hexagonal Laves (C14)	C14	hP12	(194, P6_3/mmc)	also CuZn2.	2	(AL, CO, CR, CU, FE, HF, MN, MO, NB, NI, RE, RU, SI, TA, TI, V, W, Y, ZN, ZR)2.0(AL, CO, CR, CU, FE, HF, MN, MO, NB, NI, RE, RU, SI, TA, TI, V, W, Y, ZN, ZR)1.0
C15_LAVES	Cu2Mg Cubic Laves (C15)	C15	cF24	(227, Fd-3m)		2	(AL, CO, CR, CU, FE, HF, IR, MN, MO, NB, NI, RE, RH, RU, SI, TA, TI, V, W, Y, ZN, ZR)2.0(AL, CO, CR, CU, FE, HF, MO, NB, NI, RE, RH, RU, SI, TA, TI, V, W, Y, ZR)1.0
C16_THETA	Khatyrkite (Al2Cu, C16)	C16	tI12	(140, I4/mcm)		2	(AL, HF, MN, MO, NB, SN, TA, TI, W, ZR)2.0(AL, CO, CR, CU, FE, IR, MN, NI, RH, SI)1.0
C36_LAVES	MgNi2 Hexagonal Laves (C36)	C36	hP24	(194, P6_3/mmc)		2	(AL, CO, CR, CU, FE, HF, MO, NB, NI, TA, TI, W, ZR)2.0(AL, CO, CR, CU, FE, HF, MO, NB, NI, TA, TI, W, ZR)1.0
CRNI2_OP6	MoPt2		oI6	(71, Immm)		2	(CR, MO, W)1.0(MO, NI, W)2.0
CU2TI1	Au2V		oS12	(63, Cmcn)		2	(CO, CU, NI)2.0(TI)1.0
CU2Y_HT	Unknown Structure		hP*			2	(CU)2.0(Y)1.0
CU2Y_LT	CeCu2		oI12	(74, Imma)		2	(CU)2.0(Y)1.0
HF2IR	NiTi2		cF96	(227, Fd-3m)		2	(HF)2.0(IR)1.0
HF2RH	NiTi2		cF96	(227, Fd-3m)		2	(HF)2.0(RH)1.0

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HFSN2_C40	CrSi ₂ (C40)	C40	hP9	(180, P6 ₂ 22)		2	(HF)1.0(SN)2.0
MNNI2	Unknown Structure					2	(MN, NI)1.0(NI)2.0
NBZN2_C36	MgNi ₂ Hexagonal Laves (C36)	C36	hP24	(194, P6 ₃ /mmc)		2	(NB)0.333(ZN)0.667
NI2V	MoPt ₂		oI6	(71, Immm)		2	(MO, NI)2.0(MO, NB, TA, V)1.0
NI2TA_C11B	MoSi ₂ (C11b)	C11b	tI6	(139, I4/mmm)		2	(CO, NI)2.0(TA, TI)1.0
NITI2	NiTi ₂		cF96	(227, Fd-3m)		2	(CO, CR, CU, FE, NI, RE, TI)1.0(AL, CR, CU, HF, NI, TA, TI, ZR)2.0
NI2Y1	Ni ₂ Tm		cF192	(216, F-43m)		2	(NI)2.0(Y)1.0
REZR2	Zr ₂₁ Re ₂₅		hR92	(167, R-3c)	aka Zr ₂₁ Re ₂₅ .	2	(NI, RE)1.0(ZR)2.0
RHSN2_RT	RhSn ₂		tI26	(139, I4/mmm)		2	(RH)0.33333(SN)0.66667
RH2SN	Co ₂ Si (C37)	C37	oP12	(62, Pnma)		2	(RH)2.0(SN)1.0
RH2TA	Co ₂ Si (C37)	C37	oP12	(62, Pnma)		2	(RH)2.0(TA)1.0
SNTI2_B82	Ni ₂ In (B82)	B82	hP6	(194, P6 ₃ /mmc)		2	(SN)1.0(TI)2.0
SN2Y_C49	ZrSi ₂ (C49)	C49	oS12	(63, Cmcm)		2	(SN)2.0(Y)1.0
RHZR2	NiTi ₂		cF96	(227, Fd-3m)		2	(RH)1.0(ZR)2.0
TASN2_CB	Mg ₂ Cu (Cb)	Cb	oF48	(70, Fddd)	also SN2TA_CB.	2	(TA)1.0(SN)2.0
VSN2_CB	Mg ₂ Cu (Cb)	Cb	oF48	(70, Fddd)		2	(SN)0.6(V)0.4
YZN2_LT	Unknown Structure					2	(Y)1.0(ZN)2.0

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YZN2_HT	Unknown Structure					2	(Y)1.0(ZN)2.0
AL3CO1	Os4Al13		mS34	(12, C2/m)	aka CoAl3_cub.	2	(AL)3.0(CO)1.0
AL3IR_D018	Na3As (D018)	D018	hP8	(194, P6_3/mmc)		2	(AL)0.75(IR)0.25
AL3NI_D011	Cementite (Fe3C, D011)	D011	oP16	(62, Pnma)		2	(AL)0.75(NI)0.25
AL3RH_LT	(Al3Rh)		oP*	(62, Pnma)		2	(AL)3.0(RH)1.0
ALTi3_D019	Ni3Sn (D019)	D019	hP8	(194, P6_3/mmc)	also Ni3Sn_LT, Mn3Sn, Ti3Sn.	2	(AL, CO, CR, CU, MN, MO, NB, NI, SN, TA, TI, V, W, ZR)3.0(AL, C, CR, MO, NB, NI, SI, SN, TA, TI, V, W)1.0
AL3TI_D022	Al3Ti (D022)	D022	tI8	(139, I4/mmm)	gamma double prime, AL3TI_D022, Al3M, Ni3V.	2	(AL, CO, CR, FE, MO, NB, NI, SI, TI, V)3.0(AL, CO, CR, MO, NB, NI, SI, TA, TI, V, ZR)1.0
AL3Y_HT	BaPb3		hR12	(166, R-3m)		2	(AL)0.75(Y)0.25
AL3Y_LT	Ni3Sn (D019)	D019	hP8	(194, P6_3/mmc)		2	(AL)0.75(Y)0.25
AL3ZR_D023	Al3Zr (D023)	D023	tI16	(139, I4/mmm)	also Al3Hf, Zn3Zr.	2	(AL, ZN)3.0(HF, TI, ZR)1.0
CO1SN3	Au3Zn		oS32	(64, Cmce)		2	(CO)0.25(SN)0.75
CO3V1	Al3Pu		hP24	(194, P6_3/mmc)		2	(CO, NI, V)3.0(CO, V)1.0
CO3Y1	Ni3Pu		hR12	(166, R-3m)		2	(CO)3.0(Y)1.0
CU3SN_LT	Cu3Sn		oS80	(63, Cmcm)	epsilon	2	(CU, SN)3.0(CU, SN)1.0
CUTi3_L60	CuTi3 (L60)	L60	tP4	(123, P4/mmm)		2	(CU, TI)1.0(TI)3.0
HFIR3_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)		2	(HF)1.0(IR)3.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
HFNI3_ALPHA	PdRh2Ta		hP40	(194, P6 ₃ /mmc)	the LT phase.	2	(HF)0.25(NI)0.75
HFNI3_BETA	BaPb3		hR12	(166, R-3m)	the HT phase.	2	(HF)0.25(NI)0.75
IRNB_ALPHA2	TaIr		oP12	(51, Pmma)		2	(IR)1.0(IR, NB)1.0
IR3W_D019	Ni3Sn (D019)	D019	hP8	(194, P6 ₃ /mmc)		2	(IR, W)3.0(IR, W)1.0
IRY3_D011	Cementite (Fe3C, D011)	D011	oP16	(62, Pnma)		2	(IR)1.0(Y)3.0
IR3Y1	Ni3Pu		hR12	(166, R-3m)		2	(IR)3.0(Y)1.0
IRZR3_D0E	alpha-V3S		tI32	(121, I-42m)		2	(IR)1.0(ZR)3.0
MY3_D011	Cementite (Fe3C, D011)	D011	oP16	(62, Pnma)	also CoY3, NiY3.	2	(CO, NI)1.0(Y)3.0
MZR3_E1A	MgCuAl2 (E1a)		oS16	(63, Cmcmm)	also CoZr3, FeZr3.	2	(CO, FE, NI)1.0(ZR)3.0
MN3TI1	Unknown Structure				HT phase	2	(MN)3.0(TI)1.0
MO1IR3	Ni3Sn (D019)	D019	hP8	(194, P6 ₃ /mmc)		2	(MO)1.0(IR, MO)3.0
MO3IR_A15	Cr3Si (A15)	A15	cP8	(223, Pm-3n)		2	(MO)3.06(IR)0.94
NBZN3_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)		2	(NB)0.25(ZN)0.75
NI3TA_D0A	beta-TiCu3 (D0a)	D0a	oP8	(59, Pmmn)	also delta, Ni3Mo, Ni3Nb.	2	(AL, CO, CR, FE, NB, NI)3.0(AL, FE, MO, NB, NI, TA, TI, V, W)1.0
NI3TI_D024	Ni3Ti (D024)	D024	hP16	(194, P6 ₃ /mmc)	also Eta, AlNi6Ta.	2	(AL, CO, CR, CU, FE, HF, NI, TA, TI, W, ZR)0.75(AL, CR, CU, HF, MO, NB, NI, SI, TA, TI, W, ZR)0.25
M3Y	Ni3Pu		hR12	(166, R-3m)	also Fe3Y, Ni3Y.	2	(FE, NI)3.0(Y)1.0
RHY3_D011	Cementite (Fe3C, D011)	D011	oP16	(62, Pnma)		2	(RH)1.0(Y)3.0

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RH3Y1	CeNi3		hP24	(194, P6 ₃ /mmc)		2	(RH)3.0(Y)1.0
RH3ZR_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)		2	(RH, ZR)3.0(RH, ZR)1.0
RUY3_D011	Cementite (Fe3C, D011)	D011	oP16	(62, Pnma)		2	(RU)0.25(Y)0.75
SN3Y1	GdSn2.75		oS16	(38, Amm2)		2	(SN)3.0(Y)1.0
TA3SN_A15	Cr3Si (A15)	A15	cP8	(223, Pm-3n)		2	(TA)3.0(SN)1.0
V3SN_A15	Cr3Si (A15)	A15	cP8	(223, Pm-3n)		2	(SN)0.205(V)0.795
VZN3_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)		2	(V)1.0(ZN)3.0
Y1ZN3	Zn3Y		oP16	(62, Pnma)		2	(Y)1.0(ZN)3.0
ZN3ZR_HT	Unknown Structure		c**			2	(ZN)3.0(ZR)1.0
AL13CO4	Orthorhombic Co4Al13		oP102	(31, Pmn2 ₁)		2	(AL)13.0(CO)4.0
AL5CO2_D811	Co2Al5 (D811)	D811	hP28	(194, P6 ₃ /mmc)	also Al5Rh2.	2	(AL)5.0(CO, RH)2.0
AL9CO2	Co2Al9 (D8d)	D8d	mP22	(14, P2 ₁ /c)	also Al9Rh2	2	(AL)9.0(CO, RH)2.0
AL11CR2	Al5Cr		mS732	(15, C2/c)		3	(AL)10.0(AL)1.0(CR)2.0
AL13CR2	Al45V7		mS104	(12, C2/m)		2	(AL)13.0(CR)2.0
AL4CR	mu-Al4Mn		hP574	(194, P6 ₃ /mmc)		2	(AL)4.0(CR)1.0
AL8CR5_LT_D810	Cr5Al8 (D810)	D810	hR26	(160, R3m)		2	(AL)8.0(CR, V)5.0
AL9CR4_HT	Unknown Structure					2	(AL)9.0(CR)4.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
AL9CR4_LT	Unknown Structure					2	(AL)9.0(CR)4.0
ALCU_DELTA	Al5Cu8		hR52	(160, R3m)		2	(AL)2.0(CU, FE)3.0
ALCU_ZETA	Al9Cu11(h)		oF88	(42, Fmm2)	united HT-zeta1 and LT-zeta2.	2	(AL)9.0(CU, FE)11.0
AL23CUFE4_D2H	MnAl6 (D2h)	D2h	oS28	(63, Cmcmm)		3	(AL)23.0(CU)1.0(FE)4.0
AL62CU25FE13	Quasicrystal					3	(FE)0.125(AL, CU)0.255(AL)0.62
AL7CU2FE	FeCu2Al7 (E9a)	E9a	tP40	(128, P4/mnc)		3	(FE, NI)1.0(CU)2.0(AL)7.0
AL10CU10FE	(Al10Cu10Fe)		oF116	(42, Fmm2)		3	(FE)1.0(AL, CU)10.0(AL)10.0
AL7CU4NI	(Cu0.8Ni0.2)2.53Al3.5		hR14	(166, R-3m)		2	(AL)1.0(CU, FE, NI, VA)1.0
AL28CU4MN7	Unknown Structure				Tau1, Mn6+xCu4+yAl29-x-y.	3	(AL)28.0(MN)7.0(CU)4.0
AL11CU5MN3	Unknown Structure		oP380		Tau2	3	(AL)11.0(MN)3.0(CU)5.0
ALCU3MN2_C15	Cu2Mg Cubic Laves (C15)	C15	cF24	(227, Fd-3m)	Tau3.	3	(AL)1.0(MN)2.0(CU)3.0
AL5CU4ZN	Unknown Structure					4	(AL, CU)1.0(AL)4.0(CU)4.0(ZN)1.0
AL13FE4	Al13Fe4		mS102	(12, C2/m)	solid- solution of Al13Fe4 (aka Al3Fe) & Al13Ru4.	3	(AL, CU)0.6275(FE, MN, RU)0.235(AL, SI, VA)0.1375
AL5FE2	Al2.8Fe		oS24	(63, Cmcmm)		2	(AL, CU)5.0(FE, MN)2.0
AL5FE4_D82	gamma-brass (Cu5Zn8, D82)	D82	cI52	(217, I-43m)	also AL8FE5_D82	1	(AL, CU, FE)1.0
AL16FEMN3	mu-Al4Mn		hP574	(194, P6_3/mmc)	Tau1	2	(AL)4.0(FE, MN)1.0
AL13FE2MN2	Al13Fe4		mS102	(12, C2/m)	Tau2.	2	(FE, MN)4.0(AL)13.0
AL10FEMN2	Mn3Al10		hP26	(194, P6_	Tau3.	2	(FE, MN)3.0(AL)10.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
				3/mmc			
AL9IR2	Co2Al9 (D8d)	D8d	mP22	(14, P2_1/c)		2	(AL)0.818(IR)0.182
AL45Ir13	Al45Ir13		oP236	(62, Pnma)		2	(AL)0.776(IR)0.224
AL13IR4	Unknown Structure					2	(AL)0.765(IR)0.235
AL28IR9	Al28Ir9		hP236	(159, P31c)		2	(AL)0.757(IR)0.243
AL5IR2	Al2.75Ir		cP60	(195, P23)		2	(AL)0.73(IR)0.27
AL12MN_GPHASE	Al12W		cl26	(204, Im-3)		2	(AL)12.0(MN)1.0
AL4MN_LAMBDA	lambda-Al4Mn		hP586	(194, P6_3/mmc)	also AL461MN107	2	(AL)461.0(Fe, Mn)107.0
AL4MN_MU	mu-Al4Mn		hP574	(194, P6_3/mmc)		2	(AL)4.0(MN)1.0
AL11MN4_LT	Al11Mn4		aP15	(2, P-1)		2	(AL)11.0(Fe, Mn)4.0
AL11MN4_HT	Mn6(Mn0.5Al0.5)8Al25		oP156	(62, Pnma)		2	(AL, MN)29.0(MN)10.0
AL8MN5_D810	Cr5Al8 (D810)	D810	hR26	(160, R3m)		3	(AL, Ti, Zn)12.0(MN)5.0(AL, Cu, Mn, Si, Ti)9.0
AL6MN_D2H	MnAl6 (D2h)	D2h	oS28	(63, Cmcm)	also Al6Re, Al6Ru	2	(AL)6.0(Fe, Mn, Re, Ru)1.0
AL31MN6NI2	mu-Al4Mn		hP574	(194, P6_3/mmc)		3	(AL)31.0(MN)6.0(Ni)2.0
TI25MN9AL66_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)	also Ti25Mn8Al67 or Ti43Mn11Al66, Tau1.	2	(AL, MN, Ti)0.75(AL, MN, Ti)0.25
AL24MN5ZN	Unknown Structure				ternary Tau1 phase	3	(MN, Zn)5.0(Zn)1.0(AL)24.0
AL9MN2ZN	Unknown Structure				ternary Tau2 phase	3	(MN)2.0(Zn)1.0(AL)9.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
AL11MN3ZN2	Unknown Structure		oS152		ternary Tau3 phase	3	(MN)3.0(ZN)2.0(AL)11.0
AL63MO37	Unknown Structure				Mo3Al5_HT	2	(AL)63.0(MO)37.0
AL8MO3	Al8Mo3		mS22	(12, C2/m)		2	(AL)8.0(MO)3.0
ALMO_A2	Body-Centered Cubic (W, A2, bcc)	A2	cl2	(229, Im-3m)	improper modelling.	2	(AL, MO)1.0(AL, MO)1.0
AL3NI5	Ga3Pt5		oS16	(65, Cmmm)		2	(AL)0.375(NI)0.625
AL3NI2_D519	Al3Ni2 (D513)	D513	hP5	(164, P-3m1)	also Al3Ru2.	3	(AL, SI, SN, ZN)3.0(AL, CU, NI, RU)2.0(NI, RU, VA)1.0
AL13NI2TI5_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)	Tau1	3	(AL)13.0(NI)2.0(TI)5.0
ALNI2ZN	Unknown Structure					3	(AL)0.25(NI)0.5(ZN)0.25
AL13NI38ZN49	Unknown Structure					3	(AL)0.13(NI)0.38(ZN)0.49
AL11RE4	Al11Mn4		aP15	(2, P-1)		2	(AL)11.0(RE)4.0
AL4RE	Al4Re		aP71	(2, P-1)		2	(AL)4.0(RE)1.0
AL5RH2_HT	beta-Al5Rh2		cP54	(195, P23)		2	(AL)2.0(RH, VA)1.0
AL7RH3_HT	Unknown Structure		mP*			2	(AL)7.0(RH)3.0
AL15SI2M4_TAU9	Al15(Mn,Fe)3Si2		cl168	(204, Im-3)		3	(AL)14.0(Fe, Mn)4.0(AL, Si)5.0
ALSI3TI2	Zr3Al4Si5		tl24	(141, I4_1/amd)	aka Ti7Al5Si12, Tau1	3	(AL)0.166667(SI)0.5(TI)0.333333
AL3TI_LT	Al3Ti-LT		tl32	(139, I4/mmm)		2	(AL, TI)3.0(AL, TI, ZR)1.0
AL5TI2_HT	Al5Ti2		tP28	(123, P4/mmm)		2	(AL, TI)5.0(AL, NB, TA, TI, V, ZR)2.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
AL5Ti3	Al5Ti3		tP32	(127, P4/mbm)		2	(Al)5.0(TA, TI)3.0
AL21V2	Al10V		cF176	(227, Fd-3m)		2	(Al)21.0(V)2.0
AL45V7	Al45V7		mS104	(12, C2/m)	aka Al7V	2	(Al)45.0(V)7.0
AL23V4	Al23V4		hP54	(194, P6 ₃ /mmc)		2	(Al)23.0(V)4.0
AL8V5_D82	gamma-brass (Cu5Zn8, D82)	D82	cI52	(217, I-43m)		2	(Al, V)8.0(Al, CR, V, Zr)5.0
AL77W23	Unknown Structure					2	(Al)77.0(W)23.0
AL7W3	Unknown Structure					2	(Al)7.0(W)3.0
AL12W_GPHASE	Al12W		cI26	(204, Im-3)	also Al12Mo, Al12Re.	2	(Al)12.0(MO, RE, W)1.0
AL4W	Al4W		mS30	(8, Cm)	also Al4Mo.	2	(Al)4.0(MO, W)1.0
AL5W	Al5W		hP12	(182, P6 ₃ 22)	also Al5MO.	2	(Al)5.0(MO, W)1.0
AL2ZR3	Zr3Al2		tP20	(136, P4 ₂ /mnm)	also Al2Hf3, Al2Y3.	2	(Al)2.0(HF, TI, Y, ZR)3.0
AL3ZR2	Zr2Al3		oF40	(43, Fdd2)	also Al3Hf2.	2	(Al)3.0(HF, ZR)2.0
AL3ZR4	Al3Zr4		hP7	(191, P6/mmm)	also Al3Hf4.	2	(Al)3.0(HF, TI, ZR)4.0
AL3ZR5_D8M	W5Si3 (D8m)	D8m	tI32	(140, I4/mcm)		2	(Al)3.0(TI, ZR)5.0
AL4ZR5	Ti5Ga4		hP18	(193, P6 ₃ /mcm)		2	(Al)4.0(ZR)5.0
CO10CU57TI33	MoSi2 (C11b)	C11b	tI6	(139, I4/mmm)	Strukturbericht C11B.	3	(CO)0.1(CU)0.57(TI)0.33

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
CO7HF	(Co11Hf2)		oP*	(50, Pban)		2	(CO)7.0(HF)1.0
COSNTI_TAU2	Half-Heusler (C1b)	C1b	cF12	(216, F-43m)	aka TiCoSn.	3	(CO, NI)1.0(SN)1.0(TI)1.0
CO3Y2	Unknown Structure		cP*			2	(CO)3.0(Y)2.0
CO3Y4	Co3Ho4		hP22	(176, P6_3/m)		2	(CO)3.0(Y)4.0
CO5Y_D2D	CaCu5 (D2d)	D2d	hP6	(191, P6/mmm)		3	(CO2, Y)1.0(CO)4.0(CO, VA)1.0
CO5Y8	Co5Y8		mP52	(14, P2_1/c)		2	(CO)5.0(Y)8.0
CO7NB2	(Co7Nb2)		mS18	(12, C2/m)		2	(CO)7.0(NB)2.0
CO7TA2	BaPb3		hR12	(166, R-3m)		2	(CO)7.0(TA)2.0
CO7Y6	Unknown Structure					2	(CO)7.0(Y)6.0
CO17Y2	Ni17Th2		hP38	(194, P6_3/mmc)	united HT/LT phase.	3	(CO2, Y)1.0(CO2, Y)2.0(CO)15.0
COZN_LT_A13	beta-Mn (A13)	A13	cP20	(213, P4_132)		2	(CO, ZN)1.0(VA)1.0
COZN_HT	Unknown Structure					2	(CO, ZN)1.0(VA)1.0
COZN_GAMMA_D82	gamma-brass (Cu5Zn8, D82)	D82	cI52	(217, I-43m)	aka Zn11Co2, COZN4_D83.	2	(CO, ZN)1.0(VA)1.0
COZN_DELTA	Unknown Structure				Zn15Co2 HT phase	2	(CO)0.117647(ZN)0.882353
COZN_GAMMA1	Co2Zn15		mS28	(12, C2/m)	aka CoZn7.8	2	(CO)0.125(ZN)0.875
COZN_GAMMA2	CoZn13		mS28	(12, C2/m)	aka CoZn13	2	(CO)0.0714286(ZN)0.9285714
CO11ZR2	(Co11Hf2)		oP*	(50, Pban)		2	(CO)11.0(ZR)2.0
CRMN3_HT_SIGMA	sigma-CrFe (D8b)	D8b	tP30	(136, P4_		3	(MN)8.0(CR)4.0(CR, MN)18.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
				2/mnm			
CR3MN5	alpha-Mn (A12)	A12	cI58	(217, I-43m)		2	(CR)3.0(MN)5.0
CRZN13	Unknown Structure		m**			2	(CR)1.0(ZN)13.0
CRZN17	Unknown Structure		hP*			2	(CR)1.0(ZN)17.0
CU51HF14	Ag51Gd14		hP68	(175, P6/m)		2	(CU)51.0(HF)14.0
CU8HF3	Cu8HF3		oP44	(62, Pnma)		2	(CU)8.0(HF)3.0
CU10HF7	Ni10Zr7		oS68	(64, Cmce)		2	(CU)10.0(HF)7.0
CU5MN4SI	Unknown Structure					3	(CU)0.5(MN)0.37(SI)0.13
CU4MNSN_TAU1	MgCu4Sn		cF24	(216, F-43m)		3	(CU)0.6666(SN)0.1667(MN)0.1667
CUMNZN_TAU1	Cu2Mg Cubic Laves (C15)	C15	cF24	(227, Fd-3m)		3	(CU)0.334(MN)0.333(ZN)0.333
CU6NISI3	Unknown Structure					2	(CU, NI)0.732(SI)0.268
CU46NI25SI29	Unknown Structure					3	(CU)0.458(NI)0.25(SI)0.292
CU33SI7_DELTA	Unknown Structure				HT phase	2	(CU)0.825(SI)0.175
CU15SI4_EPS_D86	Cu15Si4 (D86)	D86	cI76	(220, I-43d)		2	(CU, MN)0.789474(AL, SI)0.210526
CU56SI11_GAMMA_A13	Mg3Ru2		cP20	(213, P4_132)		2	(CU, MN, NI, SI)0.835821(SI)0.164179
CU3SI_ETA	Cu3Si-h2		hR*	(162, P-31m)	United 3-allotropes: eta''-RT (oS* *), eta'-HT1(hR27 R-3), eta-HT2(hR* P-31m)	2	(CU, MN, NI)0.76(SI)0.24
CU6SN5_HT	Ni2In (B82)	B82	hP6	(194, P6_3/mmc)	also Co3Sn2, Mn(2-x)Sn, Ni3Sn2.	3	(CO, CU, MN, NI, VA)1.0(AL, CU, NI, SN)1.0(CO, CU, MN, NI, VA)1.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
CU6SN5_LT	Cu6Sn5		mS44	(15, C2/c)	eta-prime.	3	(CU)1.0(CU, SN)1.0(SN)1.0
CU10SN3	Cu10Sn3		hP26	(173, P6_3)	zeta.	2	(CU, NI)0.769(SN)0.231
CU41SN11	Cu41Sn11		cF416	(216, F-43m)	delta.	2	(CU, SN, ZN)41.0(CU, SN, ZN)11.0
CU3TI2	Cu3Ti2		tP10	(129, P4/nmm)		2	(CU, FE, NI)3.0(CO, TI)2.0
CU4TI1	Au4Zr		oP20	(62, Pnma)		2	(CU, TI)4.0(CU, TI)1.0
CU4TI3	Cu4Ti3		tI14	(139, I4/mmm)		2	(CO, CU, NI)4.0(TI)3.0
CU2TIZR	MgZn2 Hexagonal Laves (C14)	C14	hP12	(194, P6_3/mmc)		3	(CU)0.5(TI)0.25(ZR)0.25
CU7Y1	Cu7Tb		hP8	(191, P6/mmm)	aka Cu6Y or Cu5.4Y0.8	2	(CU2, Y)1.0(CU)5.0
CU4Y	Cu5Y1.25		mP16	(11, P2_1/m)		2	(CU)4.0(Y)1.0
CU7Y2	Ag51Gd14		hP68	(175, P6/m)		2	(CU)7.0(Y)2.0
CUMNZN_EPSILON_HCP	Hexagonal Close Packed (Mg, A3, hcp)	A3	hP2	(194, P6_3/mmc)		2	(CU, MN, ZN)1.0(VA)0.5
CU10ZR7	Ni10Zr7		oS68	(64, Cmce)		2	(CU)10.0(ZR)7.0
CU51ZR14	Ag51Gd14		hP68	(175, P6/m)		2	(CU)51.0(ZR)14.0
CU8ZR3	Cu8Hf3		oP44	(62, Pnma)		2	(CU)8.0(ZR)3.0
FENBZR_CFC2_C15	Cu2Mg Cubic Laves (C15)	C15	cF24	(227, Fd-3m)		3	(FE, NB, ZR)2.0(NB, ZR)1.0(NB, ZR)3.0
FE3SN2	Fe3Sn2		hR10	(166, R-3m)		2	(FE)3.0(SN)2.0
FE5SN3_B82	Ni2In (B82)	B82	hP6	(194, P6_3/mmc)		2	(FE)5.0(SN)3.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
FE3ZN7_GAMMA_D82	gamma-brass (Cu5Zn8, D82)	D82	cI52	(217, I-43m)		4	(FE, ZN)0.154(FE, ZN)0.154(FE, ZN)0.231(ZN)0.461
FEZN4_GAMMA1_D81	Fe11Zn40		cF408	(216, F-43m)		3	(FE)0.137(FE, ZN)0.118(ZN)0.745
FEZN10_DELTA	FeZn10		hP632	(194, P6_3/mmc)		4	(FE)0.058(FE, ZN)0.18(ZN)0.525(ZN)0.237
FEZN13_ZETA	CoZn13		mS28	(12, C2/m)		3	(FE, VA)0.072(ZN)0.856(VA, ZN)0.072
CU3SN_HT_GAMMA	BiF3 (D03)	D03	cF16	(225, Fm-3m)	Cu3Sn HT solution phase	1	(CU, MN, NI, SN, ZN)1.0
HF5IR3	Ir3Zr5		hP48	(178, P6_122)		2	(HF)5.0(IR)3.0
HF3NI7	HF3Ni7		aP20	(2, P-1)		2	(HF)0.3(NI)0.7
HF8NI21	Hf8Ni21		aP29	(2, P-1)	also ZR8NI21.	2	(HF, ZR)8.0(NI)21.0
HF3RH4	Unknown Structure					2	(HF)3.0(RH)4.0
HF3RH5	Rh5Ge3		oP16	(55, Pbam)		2	(HF)3.0(RH)5.0
HF5SN4	Ti5Ga4		hP18	(193, P6_3/mcm)		2	(HF)5.0(SN)4.0
IR2Y3	Y3Rh2		tI140	(140, I4/mcm)		2	(IR)2.0(Y)3.0
IR2Y5	Mn5C2 (Fe5C2 Hagg carbide)		mS28	(15, C2/c)		2	(IR)2.0(Y)5.0
IR3Y5	Mavlyanovite (Mn5Si3, D88)	D88	hP16	(193, P6_3/mcm)	united HT/LT phase.	2	(IR)3.0(Y)5.0
IR3ZR5	Ir3Zr5		hP48	(178, P6_122)		2	(IR)3.0(ZR)5.0
MN3SN2	Tongbaite (Cr3C2, D510)	D510	oP20	(62, Pnma)		2	(MN)3.0(SN)2.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
MNTI_HT	Unknown Structure		t**		aka TiMn Mn+ or Ti0.47Mn0.53	2	(MN)0.515(Ti)0.485
MN4TI	R-(Co,Cr,Mo)		hR53	(148, R-3)	aka Ti9Mn42	2	(CR, MN)0.815(Ti)0.185
MN12Y_D2B	Mn12Th (D2b)	D2b	tI26	(139, I4/mmm)		2	(MN)12.0(Y)1.0
MNZN9	Unknown Structure		h**			2	(MN)1.0(ZN)9.0
MO7IR3	sigma-CrFe (D8b)	D8b	tP30	(136, P4_2/mnm)		2	(MO)0.7(IR)0.3
MONI4_BETA_D1A	Ni4Mo (D1a)	D1a	tI10	(87, I4/m)	also WNi4.	2	(MO, W)1.0(NI)4.0
MONI_DELTA	MoNi		oP56	(19, P2_12_12_1)		3	(CO, CR, FE, NI, RE)24.0(CO, CR, FE, MO, NI, RE, W)20.0(CU, MO, W)12.0
MOZN7	Ca7Ge		cF32	(225, Fm-3m)		2	(MO)1.0(ZN)7.0
MOZN22	Zn93(Zn0.43Mo0.57)Mo4		cF420	(216, F-43m)		2	(MO)1.0(ZN)22.0
NB15NI56TI29_XA	Unknown Structure		o*100			3	(NB)0.15(NI)0.56(TI)0.29
NB8NI9TI3_XB	Unknown Structure					3	(NB)0.4(NI)0.45(TI)0.15
NB5NI75TI20_XC	Ni3Sn (D019)	D019	hP8	(194, P6_3/mmc)		3	(NB)0.05(NI)0.75(TI)0.2
NB13NI75TI12_XD	Unknown Structure					3	(NB)0.13(NI)0.75(TI)0.12
NB15NI80TI5_XE	Unknown Structure					3	(NB)0.15(NI)0.8(TI)0.05
NB3RU5	Rh5Ge3		oP16	(55, Pbam)	united Nb3Ru5_HT and NbRu3_LT phase	2	(NB, RU)0.375(RU)0.625
NBRH_ETA	Al3Pu		hP24	(194, P6_3/mmc)		2	(NB, RH)1.3(RH)2.7

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
NBRH_ZETA	Unknown Structure					2	(NB)3.0(NB, RH)5.0
NBRH_EPSILON	beta'-AuCd (B19)	B19	oP4	(51, Pmma)		2	(NB, RH)0.85(RH)1.15
NBRH_DELTA	TaIr		oP12	(51, Pmma)		2	(NB)0.9(RH)1.1
NBSN2_CB	Mg2Cu (Cb)	Cb	oF48	(70, Fddd)		2	(NB, SN, V)1.0(NB, SN)2.0
NBZN7	Unknown Structure					2	(NB)0.125(ZN)0.875
NBZN15	TiZn16		oS68	(63, Cmcm)		2	(NB)0.0625(ZN)0.9376
NB2ZN3_D85	Fe7W6 (D85) mu-phase	D85	hR13	(166, R-3m)		2	(NB)0.4(ZN)0.6
NI8M	Pt8Ti		tI18	(139, I4/mmm)	also Ni8Ta, Ni8Nb.	2	(NI)8.0(NB, TA)1.0
NI7ZR2	Ni7Zr2		mS36	(12, C2/m)	also Ni7HF2, Ni7Y2, CO7HF2 and CO7Y2.	2	(AL, CO, CR, NI)7.0(HF, Y, ZR)2.0
NI3SN4	delta-Ni3Sn4 (D7a)	D7a	mS14	(12, C2/m)		3	(CU, NI)0.25(NI, SN)0.25(SN)0.5
NI2Y3	Ni2Y3		tP80	(92, P4_12_12)		2	(NI)2.0(Y)3.0
NI4Y	Unknown Structure		hR*			2	(NI)4.0(Y)1.0
NI17Y2	Fe17Lu2		hP80	(194, P6_3/mmc)	also Fe17Y2.	2	(AL, FE, NI)1.0(Y)0.1176
NI2N8_DELTA	Ni3Zn22		mS50	(12, C2/m)		2	(NI)0.1111111(AL, ZN)0.8888889
NI11ZR9	Pt11Zr9		tI40	(87, I4/m)	also Ni11Hf9.	2	(NI)11.0(HF, ZR)9.0
NI10ZR7	Ni10Zr7		oS68	(64, Cmce)	also Ni10Hf7.	2	(NI)23.0(HF, ZR)17.0
NI5ZR_C15B	AuBe5 (C15b)	C15b	cF24	(216, F-43m)	also Ni5Y/Ni5Hf/Cu5Hf/Cu5Zr, aka ZRM5_C15B.	2	(AL, CU, NI)5.0(HF, Y, ZR)1.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
RE3NB_A12	alpha-Mn (A12)	A12	cI58	(217, I-43m)		3	(RE)24.0(NB, V, W)10.0(NB, RE, V, W)24.0
RHSN4	IrGe4		hP15	(152, P3_121)		2	(RH)1.0(SN)4.0
RH3SN2	Ni2In (B82)	B82	hP6	(194, P6_3/mmc)		3	(RH)0.125(RH)0.5(SN)0.375
RH12TA8	Rh12Ta8		oP12	(51, Pmma)		2	(RH, TA)1.2(TA)0.8
RH5TA4	Unknown Structure					2	(RH, TA)5.0(TA)4.0
RH5TI3	Rh5Ge3		oP16	(55, Pbam)		2	(RH)5.0(TI)3.0
RH5V3	Rh5V3		oS16	(63, Cmcm)		2	(RH)5.0(V)3.0
RH5Y_HT	CaCu5 (D2d)	D2d	hP6	(191, P6/mmm)		2	(RH)5.0(Y)1.0
RH2Y3	Y3Rh2		tI140	(140, I4/mcm)		2	(RH)2.0(Y)3.0
RH3Y5	Unknown Structure					2	(RH)3.0(Y)5.0
RH3Y7	Fe3Th7 (D102)	D102	hP20	(186, P6_3mc)		2	(RH)3.0(Y)7.0
RH4ZR3_LT	Pd4Pu3		hR14	(148, R-3)	alpha_Rh4Zr3.	2	(RH, ZR)4.0(ZR)3.0
RH4ZR3_HT	Unknown Structure				beta_Rh4Zr3	2	(RH, ZR)4.0(ZR)3.0
RH5ZR3	Pd5Pu3		oS32	(63, Cmcm)		2	(RH)5.0(RH, ZR)3.0
RU3SN7_D8F	Ir3Ge7 (D8f)	D8f	cI40	(229, Im-3m)		2	(RU)0.3(SN)0.7
RU2SN3	Ru2Sn3		tP20	(116, P-4c2)		2	(RU)0.4(SN)0.6
RU2Y3	Er3Ru2		hP10	(176, P6_3/m)		2	(RU)0.4(Y)0.6
RU25Y44	Ru25Y44		oP276	(52, Pnna)		2	(RU)0.362(Y)0.638

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
RU2Y5	Mn5C2 (Fe5C2 Hagg carbide)		mS28	(15, C2/c)		2	(RU)0.286(Y)0.714
SN3Ti2	Unknown Structure					2	(SN)3.0(Ti)2.0
SN5Ti6	Sn5Ti6-beta		hP22	(194, P6 ₃ /mmc)	also Sn5Nb6.	2	(Al, SN)5.0(Nb, Ti)6.0
SN5Y2	Shcherbinaite (V2O5) (Revised)		oP14	(59, Pmmn)		2	(SN)5.0(Y)2.0
SN10Y11	Ge10Ho11		tI84	(139, I4/mmm)		2	(SN)10.0(Y)11.0
SN4Y5	Gd5Si4		oP36	(62, Pnma)		2	(SN)4.0(Y)5.0
TA1Al1	Al38Ta48		mP86	(14, P2 ₁ /c)		2	(TA)0.51515(Al)0.48485
TAAL2_HT	Al69Ta39		cF444	(216, F-43m)		2	(TA)0.35(Al)0.65
TA41IR59	CuAu (L10)	L10	tP2	(123, P4/mmm)	aka Ta0.82Ir1.18_rt, gamma.	2	(TA)0.41(IR)0.59
TA43IR57	TaIr		oP12	(51, Pmma)	aka Ta0.86Ir1.14, delta.	2	(TA)0.43(IR, TA)0.57
T1CUNITI_C11B	MoSi2 (C11b)	C11b	tI6	(139, I4/mmm)	the Cu-Ni-Ti ternary phase Tau1.	2	(Cu, Ni)2.0(Ti)1.0
T2CUNITI	Cu3Ti2		tP10	(129, P4/nmm)	the Cu-Ni-Ti ternary phase Tau2.	3	(Cu)0.175(Ni)2.825(Ti)2.0
T4CUNITI	BaPb3		hR12	(166, R-3m)	the Cu-Ni-Ti ternary phase Tau4.	3	(Cu)0.05(Ni)0.7(Ti)0.25
T6CUNITI	Unknown Structure				the Cu-Ni-Ti ternary phase Tau6	3	(Cu)0.25(Ni)0.5(Ti)0.25
T1CUFETI_CU2TI	Au2V		oS12	(63, Cmcn)	ternary Ti0.33FexCu0.67-x, Tau1.	2	(Cu, Fe)2.0(Ti)1.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
T2CUFETI_CU3TI2	Cu3Ti2		tP10	(129, P4/nmm)	ternary Ti0.4FexCu0.6-x, Tau2.	2	(CU, FE)3.0(TI)2.0
T3CUFETI_CU4TI3	Cu4Ti3		tI14	(139, I4/mmm)	ternary Ti0.43FexCu0.57-x, Tau3.	2	(CU, FE)4.0(TI)3.0
T4CUFETI	Unknown Structure				ternary Ti0.37FexCu0.63-x, Tau4	2	(CU, FE)0.63(TI)0.37
T5CUFETI	Unknown Structure				ternary Ti0.45FexCu0.55-x, Tau5.	2	(CU, FE)0.55(TI)0.45
TIZN5	Unknown Structure					2	(TI)1.0(ZN)5.0
TIZN10	Ti3Zn22		tP100	(135, P4_2/mbc)		2	(TI)1.0(ZN)10.0
TIZN15	TiZn16		oS68	(63, Cmcm)		2	(TI)1.0(ZN)15.0
V4ZN5	V4Zn5		tI18	(139, I4/mmm)		2	(V)4.0(ZN)5.0
Y2ZN17	Ni17Th2		hP38	(194, P6_3/mmc)		2	(Y)2.0(ZN)17.0
Y13ZN58	Y13Zn58		hP146	(194, P6_3/mmc)		2	(Y)13.0(ZN)58.0
YZN5	ErZn5		hP36	(194, P6_3/mmc)	aka H_RZN5	2	(Y)1.0(ZN)5.0
ZN12Y_D2B	Mn12Th (D2b)	D2b	tI26	(139, I4/mmm)		2	(ZN)12.0(Y)1.0
ZN11Y3	Al11La3		oI28	(71, Immm)		2	(ZN)11.0(Y)3.0
ZN22ZR	Zn22Zr		cF184	(227, Fd-3m)		2	(ZN)22.0(ZR)1.0
ZN39ZR5	Zn39Zr5		mS88	(12, C2/m)		2	(ZN)39.0(ZR)5.0

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
ZN2ZR3	Zr3Al2		tP20	(136, P4_2/mnm)		2	(ZN)2.0(ZR)3.0
GAS	Gas					1	(AL, AL1C1, AL1C2, AL1CU1, AL1N1, AL2, AL2C2, B, B1C1, B1C2, B1N1, B2, B2C1, C, C1R1, C1N1, C1N2_CNN, C1N2_NCN, C1RH1, C1S1, C1S2, C1S3, C1S4, C2, C2N1_CCN, C2N1_CNC, C2N2, C2S1, C2S2, C2S3, C3, C3N1, C4, C4N1, C4N2, C5, C5N1, C6, C6N1, C6N2, C9N1, CO, CO2, CR, CR1N1, CR2, CU, CU2, FE, FE2, HF, IR, MN, MO, MO1N1, MO2, N, N1NB1, N1S1, N1S2, N1TI1, N1V1, N1ZR1, N2, N3, NB, NI, NI2, RE, RH, RU, SI, S2, S3, SN, SN2, TA, TI, TI2, V, W, Y, ZN, ZR, ZR2)1.0

TCHEA7 Properties Data

The properties data for molar volume, viscosity, surface tension, electrical resistivity, and thermal conductivity are included with the database. Molar volume is available starting with version 1 (TCHEA1), viscosity is available starting with version 4 (TCHEA4), and surface tension, electrical resistivity, and thermal conductivity are available starting with version 5 (TCHEA5).

Model Descriptions

For more information about the various thermophysical models, and when in Thermo-Calc, press F1 to search the online help. The details are found under a *General Reference* section.



You can find information on our website about the [properties that can be calculated](#) with Thermo-Calc and the Add-on Modules. Additional resources are added on a regular basis so keep checking back or [subscribe to our newsletter](#).

Examples



Go to the [High Entropy Alloys Databases](#) page on our website where you can access an examples collection and the technical information plus learn more about the compatible kinetic database. Also explore further [applications of Thermo-Calc to high entropy alloys](#) including links to resources such as examples, publications, and more.

TCHEA: TCS High Entropy Alloys Database Revision History

Current Database Version

Database name (acronym):	TCS High Entropy Alloys Database (TCHEA)
Database owner:	Thermo-Calc Software AB
Database version:	7.0
First release:	TCHEA1 was released with 2015b

Changes in the Most Recent Database Release

TCHEA6.1 to TCHEA7.0

Software release 2024a (December 2023/January 2024)

Binary Systems

- Newly assessed Mo-Sn, N-Y, Rh-Nb, Rh-Si, Rh-Ta, Rh-V, and Rh-Y.
- Reassessed Mn-W.

Ternary Systems

- Newly assessed Nb-V-Zr and V-W-Zr
- Revision of SIGMA phase stability: Al-Co-Cr, Al-Co-V, Al-Cr-Ni, Al-Cr-V, Al-Fe-Ni, Al-Fe-V, Al-Mo-Ni, Al-Mo-V, Al-Ni-V, Co-Cr-V, Co-Fe-V, Co-Mo-V, Cr-Fe-V, Fe-Mo-Ni, Fe-Mo-V, and Mo-Ni-V.
- Revision of BCC phase stability: Co-Cr-Ti, Co-Cu-Nb, Co-Cu-Ni, Co-Ni-Si, Co-Ni-V, Co-Ni-W, Cr-Cu-Si, Cr-Mo-Ni, Cr-Ni-Si, Cu-Fe-Ni, Cu-Mo-Ni, Fe-Ni-Ti, Fe-Ni-W, and Ni-Si-Ti.
- Revision of FCC phase stability: Al-Co-Zr, Al-Cr-Ti, Al-Fe-Ti, Al-Hf-Ti, Al-Mn-Ti, Al-Mo-Ti, Al-Nb-Ti, Al-Si-Ti, Al-Sn-Ti, Al-Ta-Ti, Al-Ti-Zr, Al-V-Zr, Co-Ta-Ti, Cr-Cu-Mo, Cr-Cu-Zr, Cr-Fe-Si, Cr-Fe-V, Cr-Mo-Nb, Cr-Ti-Zr, Fe-Ni-V, Fe-Ti-V, Mo-Nb-Ti, Mo-Ni-Ti, Mo-Re-Ta, Mo-Ta-Ti, Mo-Ti-W, Mo-Ti-Zr, Nb-Ti-W, Si-Ti-W, Ta-Ti-V, Ti-V-W, and Ti-V-Zr.
- Revision of HCP phase stability: Al-Co-Ni, Al-Co-W, Al-Co-Zr, Al-Cr-Ni, Al-Cu-Ni, Al-Fe-Mn, Al-Fe-Ni, Al-Nb-Ni, Al-Ni-Si, Al-Ni-Ti, Al-Ni-W, Al-Ni-Zn, Co-Cr-Ti, Co-Cu-Fe, Co-Cu-Nb, Co-Cu-Ni, Co-Ni-V, Co-Ni-W, Co-Ta-Ti, Cr-Cu-Mo, Cr-Cu-Nb, Cr-Cu-Si, Cr-Cu-Zr, Cr-Fe-Mo, Cr-Fe-Ni, Cr-Fe-Si, Cr-Fe-V, Cr-Mo-Nb,

Cr-Mo-Ni, Cr-Ni-Re, Cr-Ni-Si, Cr-Ni-W, Cr-Ti-Zr, Cu-Fe-Mn, Cu-Fe-Ni, Cu-Fe-Si, Cu-Fe-Sn, Cu-Fe-V, Cu-Mn-Ni, Cu-Mn-Si, Cu-Mn-Sn, Cu-Mo-Ni, Cu-Ni-Ti, Cu-Ni-Zn, Fe-Mo-Ni, Fe-Ni-Ti, Fe-Ni-V, Fe-Ni-W, Hf-Ni-Ti, Mo-Nb-Ti, Mo-Ni-Ti, Mo-Re-Ta, Mo-Ti-W, Mo-Ti-Zr, Ni-Si-Ti, Si-Ti-W, Ta-Ti-V, and Ti-V-W.

- Revision of LIQUID phase stability: Al-Cr-Ti, Mo-Nb-Ti, Mo-Ni-Ti, Mo-Re-Ru, Mo-Re-Ta, Mo-Ta-Ti, Ta-Ti-V, and Ti-V-W.

Surface Tension Re-assessed

The surface tension was re-assessed based on the Redlich-Kister-Muggianu (R-K-M) sub-regular solution model.

Previous Releases

TCHEA6.0 to TCHEA6.1

Software release 2023a (December 2022/January 2023)

Reassessed Binary, Ternary, and Quinary Systems

- Binary: Two reassessments, Cr-Mn, Ir-Nb.
- Ternary: Seventeen (re-)assessments: Al-Cr-V, Co-Cr-Fe, Co-Cr-Mn, Co-Cr-Ni, Co-Fe-Mn, Co-Fe-Ni, Co-Mn-Ni, Cr-Fe-Mn, Cr-Mn-Ni, Cr-Mo-Nb, Cr-Mo-Ti, Cr-Nb-V, Fe-Mn-Ni, Fe-Ni-W, Mo-Nb-V, Re-Nb-V, and Re-V-W.
- Quinary: Validation of the Co-Cr-Fe-Mn-Ni system in full temperature and composition space (7 isopleths).

TCHEA5.1 to TCHEA6.0

Software release version: 2022b (June 2022)

In this release of the database, there is one new ternary system assessment plus major changes to 18 critically assessed systems. In addition, 19 ternary systems also had some minor changes and a complete description of the gas phase is added.

- One new ternary assessment added: Al-V-Zr.
- Reassessed 18 ternary systems: Al-Co-Ti, Al-Cr-Ti, Al-Mn-Ti, Al-Mo-Ti, Al-Ni-Ti, Al-Sn-Ti, Al-Ta-Ti, Al-V-Zr, Co-Mo-Ti, Cr-Mn-Ti, Cr-Ti-Zr, Fe-Ti-V, Mo-Ti-V, Mo-Ti-Zr, Nb-Ti-W, Ta-Ti-V, Ti-V-W, and Ti-V-Zr.
- Minor modifications to 19 ternary systems: Al-B-Ti, Al-C-Ti, Al-Fe-Ti, Al-N-Ti, Al-Si-Ti, Co-Sn-Ti, Cr-Si-Ti, Cu-Ti-Zr, Mo-Nb-Ti, Mo-Ta-Ti, Mo-Ti-W, Nb-Sn-Ti, Nb-Ta-Ti, Nb-Ti-V, Nb-Ti-Zr, Si-Ti-W, Ta-Ti-W, Ta-Ti-Zr, and Ti-W-Zr.
- Added a complete description of the gas phase in the 26 element framework.

TCHEA5.0 to TCHEA5.1

Software release version: 2022a (December 2021/January 2022)

- Improved the BCC (A2/B2) phase descriptions in the framework of the Al-Cr-Nb-Ti-V-Zr system. Updated assessments of the Al-V, Mo-V, and Ti-V binaries. Updated assessments of the Al-Nb-Ti, Al-Ti-V, and Al-Ti-Zr ternaries.
- Improved the MC carbide description in the framework of the (Hf, Nb, Ta, Ti, V, Zr)C system. Added assessments of the C-Hf-Nb, C-Hf-V, C-Nb-Ta, C-Ta-V, C-Ta-Zr, C-Ti-V, and C-V-Zr ternaries. Updated assessment of the C-Hf-Ta, C-Hf-Ti, C-Nb-Ti, C-Nb-V, C-Nb-Zr, and C-Ti-Zr ternaries.
- Minor bug fixed for ternary-related calculations of the Al-Fe-W, Al-Hf-Nb, Al-Mo-Nb, and Al-Mo-Ni systems.

TCHEA4.2 to TCHEA5.0

Software release version: 2021b (June 2021)

- Addition of surface tension of liquid phase.
- Addition of electrical resistivity and thermal conductivity of liquid, BCC (A2, B2), FCC (A1, L12), and HCP solution phases.

TCHEA4.1 to TCHEA4.2

Software release version: 2021a (January 2021)

- Added the assessment of three binary systems- Ir-W, Mo-Rh, and Rh-W.

TCHEA4.0 to TCHEA4.1

Software release version: 2020b (June 2020)

- Crystal structure information is included for all phases.
- HCP_ZN is merged into HCP_A3. The epsilon phase with HCP structure in the Cu-Zn, Mn-Zn and Cu-Mn-Zn systems is separated from HCP_A3 and renamed as CUMNZN_EPSILON_HCP.
- Better estimation of excess energy for metastable solution phases and compound energy for metastable end-members in binary systems.

TCHEA3.1 to TCHEA4.0

Software release version: 2020a (January 2020)

- 49 ternaries added.
- 12 ternaries improved.
- 13 binaries added (mainly Ir-, or Rh-).
- 1 binary updated (Al-Ti).
- Includes the viscosity of metallic liquid data.

TCHEA3.0 to TCHEA3.1

Software release 2019a (December 2018).

- The Mn-Ni-Si ternary is critically assessed in full composition and temperature ranges.
- Some bug fixes e.g. avoid the fictitious HCP_ZN phase appears in Zn-free systems, adjust the phase stability of GAMMA_D03, CRSI2_C40, C15- & C36- laves phases in some systems.

TCHEA2.0 to TCHEA3.0

Software release version: 2018a (April 2018)

- Six new elements (B, Ir, Rh, Sn, Y and Zn) added.
- More than 100 new binary systems and >150 new phases added.

TCHEA2.0 to TCHEA2.1

Software release version: 2017b (October 2017)

- The database has improved stability description of the sigma phase.

TCHEA1.0 to TCHEA2.0

Software release version: 2017a (March 2017)

- Added 5 new elements (C, N, Re, Ru, Si) and 100 phases.
 - Assessed most of new binary systems that contain one or two of these 5 new elements in the 20-element framework of this database.
 - Assessed 200 additional ternary systems relevant to the 5 new elements.
 - The application of TCHEA2 extends from BCC and FCC HEAs to HCP HEA as well.
 - Revised some subsystems based on the validation against updated experimental information. This includes the phase stability of solid solutions such as BCC and FCC and intermetallic phases such as sigma and laves in some ternary and quaternary systems.
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