

TCS Steel and Fe-alloys Database (TCFE14)

Technical Information

Available Starting with Thermo-Calc Version 2025a



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About the TCS Steel and Fe-alloys Database (TCFE)

TCS Steel and Fe-alloys Database (TCFE) is a thermodynamic and properties database with applications to a variety of steels and Fe-based alloys, such as stainless steels, high-speed steels, tool steels, high-strength low alloy (HSLA) steels, cast irons, corrosion-resistant high strength steels, low-density steels, and also cemented carbides. The databases can be used for alloy compositions, but also their interaction with atmospheres containing oxygen, nitrogen, carbon, hydrogen, or mixtures thereof.

TCS Steel and Fe-alloys Database (TCFE) is developed to be used with our entire suite of products: Thermo-Calc, the Add-on Modules, and all available SDKs. In addition, it is integral to the Steel Model Library, which includes martensite, pearlite, and bainite models designed to help experts working in the steel industry to quickly and easily set up calculations using the Property Model Calculator.

 [TCFE: TCS Steels/Fe-Alloys Database Revision History](#). The current version of the database is TCFE14. See the link for any subversion release details.

 The database is compatible with the TCS Steels/Fe-Alloys Mobility Database (MOBFE). The current version is MOBFE8.1.

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 binary, ternary, and for some databases, important higher order systems. This enables predictions to be made for multicomponent systems and alloys of industrial importance. Among these, the thermodynamic database is of fundamental importance.

 For more learning resources about CALPHAD and our databases, visit the video tutorials on our [website](#) or our [YouTube playlist](#).

 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 areas such as process metallurgy, heat treatment, and more depending on the database. Sometimes an example is both a validation and a calculation example.

Some examples of how this database can be used include the following:

- Calculate various phase diagrams in the assessed systems as well as to extrapolate higher-order systems
- Simulate the solidification process.
- Predict the relative stability of matrix phases (austenite and ferrite).
- Predict the precipitation of secondary phases such as sulfides, borides, oxides, phosphides, carbides, nitrides, carbonitrides, and also intermetallic phases such as the sigma and laves phases.
- Predict the density, molar volume, and thermal expansion coefficients.
- Use thermophysical property data to predict the viscosity of metallic liquids, the surface tension of metallic liquids, electrical resistivity, and thermal conductivity.
- Use elastic property data to predict Young's modulus.
- Predict the lath and plate martensite start temperature.
- Describe the thermodynamics of bainite and pearlite formation.
- Predict oxide scale formation on various steels.
- Integrate with a compatible atomic mobility database and use it to simulate diffusion controlled phase transformations with the Add-on Diffusion Module (DICTRA) and multi-particle precipitation kinetics with the Add-on Precipitation Module (TC-PRISMA).

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.

TCS Steel and Fe-alloys Database (TCFE) 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 Steel and Fe-alloys Database (TCFE) Technical Information* PDF document contains version specific information such as the binary, ternary, and higher-order assessed systems, phases and models. It also includes details about the properties data (e.g. viscosity, surface tension, etc.), the elastic properties (elastic moduli and constants), a list of the included elements, and summaries of the database revision history by version.
- The *TCS Steel and Fe-alloys Database (TCFE) Validation and Calculation 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 [Steels and Fe-Alloys Databases](#) page on our website where you can access a Validation and Calculation Examples Collection and the Technical Information plus learn more about the compatible kinetic database. Also explore further [applications of Thermo-Calc to iron and steels](#) including links to resources such as publications, webinars, videos, and more.



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

TCFE14 Elements, Systems, Phases, and Properties

Included Elements

There are 30 elements included in the most recent version of the database.

Included Elements									
Ar	Al	B	C	Ca	Ce	Co	Cr	Cu	Fe
H	Mg	Mn	Mo	N	Nb	Ni	O	P	Ru
S	Si	Sn	Ta	Ti	V	W	Y	Zn	Zr

Assessed Systems



Also see information about [TCFE14 Composition Limits](#)

The database is developed on the basis of complete assessments of binary, ternary, and some higher order systems. However, many intermediate compounds that usually do not occur in steels/Fe-alloys are ignored in the database. Therefore, the database may not be suitable to calculate complete binary and ternary systems, but only rather in the Fe-rich corner.



Sometimes, for some special steels/Fe-alloys, you may prefer to append some other stoichiometric or solution phases (usually intermediate compound phases that have been ignored in the *TCFE Database*) from another compatible database (e.g. *SSOL: SGTE Solutions Database* and/or *SSUB: SGTE Substances Database*). But you must be careful about appropriately appending such data in the combination.

The most recent version of the database contains the following:

- 371 assessed binary systems
- 335 assessed ternary systems
- 84 assessed quaternary systems

Included Phases



[Common Phases for Steel and Fe-alloys](#) and [TCFE14 Models for the Included Phases](#)

The TCS Steel and Fe-alloys Database (TCFE) includes 465 phases. It contains an extensive GAS mixture phase for the main purpose of considering oxygen/nitrogen-gas controls in steel-making processes, and different gas atmospheres under e.g. heat treatments.

When it is really necessary to consider a more comprehensive ionic liquid phase for calculations of e.g. formations of complex oxides on steel surfaces or investigating various steelmaking metallurgical processes, an `IONIC_LIQ` solution phase can be appended from the TCS Metal Oxide Solutions Database (TCOX).

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

A variety of properties data are included with the TCS Steel and Fe-alloys Database (TCFE).



The properties data was gradually added to the database versions as follows. Molar volume with thermal expansion coefficients available starting with TCFE4, the viscosity of metallic liquids starting with TCFE10, surface tension of liquids starting with TCFE11, and electrical resistivity and thermal conductivity starting with TCFE12.



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, thermomechanical, and properties 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 an SDK such as TC-Python or TC-Toolbox for MATLAB®.

<i>Property (and Graphical Mode Variable Name)</i>	<i>Model Parameters</i>	<i>Variables to Show or Plot in Console Mode or the SDKs (TC-Python, or TC-Toolbox for MATLAB®)***</i>
Molar volume	V0, VA	VM for a system VM(PHI) for phase PHI
Electrical conductivity	ELQ**	ELCD for a system ELCD(PHI) for phase PHI
Electrical resistivity	ELRS, ESPD	ELRS for a system ELRS(PHI) for a phase PHI
Thermal conductivity	THCD	THCD for a system THCD(PHI) for phase PHI
Thermal resistivity		THRS for a system THRS(PHI) for phase PHI
Thermal diffusivity		THDF for a system THDF(PHI) for phase PHI
Surface tension	SIGM, XI*	SURF(LIQUID) SURF(ION)**
Dynamic viscosity	VISC	DVIS(LIQUID) DVIS(ION)**
Kinematic viscosity		KVIS(LIQUID) KVIS(ION)**

* 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. As of 2024b, TCSLD5.

** ION is used in the TCS Metal Oxide Solutions Database (TCOX)

*** The examples listed for the SDKs are using Console Mode syntax. The quantities can also be accessed in both `ThermodynamicQuantity` and `ScheilQuantity` classes. See the various model descriptions or the SDK help for details.

Elastic Properties Data



Starting with TCFE14, the elastic properties (elastic moduli and elastic constants) are added to the database.

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

GRAPHICAL MODE

In the **Plot Renderer** in Graphical Mode, elastic constants and moduli can be selected from the drop-down list of axis variables.

The independent elastic constants are selected on the **Plot Renderer** as an axis variable **Elastic constant** and then choose an option (**C11**, **C12**, **C13**, **C33**, or **C44**) from the drop-down list.

The elastic moduli, **Bulk modulus**, **Shear modulus**, and **Young's modulus**, are directly available from the **Axis variable** list.

All can be tabulated and plotted using the quantity names, with options for a specific phase or all phases.

CONSOLE MODE

The quantities corresponding to the individual elastic constants and elastic moduli (derived from the elastic constants) can be calculated in Console Mode for individual phases or all phases. The results can be shown in the POLY module with the command SHOW_VALUE or shown as a plot in the POST module with the command PLOT_DIAGRAM using:

- `Cij(<phase name>)` or `Cij(*)`
- **Bulk modulus:** `BULKMOD(<phase name>)` or `BULKMOD(*)`
- **Shear modulus:** `SHEARMOD(<phase name>)` or `SHEARMOD(*)`
- **Young's modulus:** `YOUNGMOD(<phase name>)` or `YOUNGMOD(*)`

TC-PYTHON AND TC-TOOLBOX FOR MATLAB®

For the Software Development Kits (SDKs), i.e. TC-Python and TC-Toolbox, the quantities of elastic constants, bulk modulus, shear modulus, and Young's modulus can be retrieved for individual phases or all phases via `get_value_of()` or `get_values_of()` from any equilibrium calculation types using:

- `Cij(<phase name>)` or `Cij(ALL_PHASES/*)`
- `ThermodynamicQuantity.bulk_modulus(<phase name>)` or `ThermodynamicQuantity.bulk_modulus(ALL_PHASES/*)`
- `ThermodynamicQuantity.shear_modulus(<phase name>)` or `ThermodynamicQuantity.shear_modulus(ALL_PHASES/*)`
- `ThermodynamicQuantity.youngs_modulus(<phase name>)` or `ThermodynamicQuantity.youngs_modulus(ALL_PHASES/*)`



See the relevant SDK documentation for details.

TCFE14 Composition Limits

The TCS Steel and Fe-alloys Database (TCFE) is applicable for various types of steels/Fe-alloys with an Fe-minimum of 50 wt.%.



The critical calculations must always be verified by equilibrium experimental data; it is the user's responsibility to verify the calculations.



A sensible calculation cannot be expected if all alloying elements are at their highest limits. Some combinations of elements at high values do not give reasonable results. However, some alloying elements can exceed their limits considerably and the calculations still give good results.



If you discover any significant deviations and want to help us improve future versions of the database, please contact Thermo-Calc Software [at one of our offices](#) or send us an email at info@thermocalc.com.

For alloying elements, the recommended composition limits are given in the table below.



Ar is only considered in the gas phase and no modeling of solubility in the solid solution phases or liquid is taken into account.

Element	Max	Element	Max
Al	10	Ni	20
B	Trace	O	Trace
C	7	P	Trace
Ca	Trace	Ru	15
Ce	Trace	S	Trace

<i>Element</i>	<i>Max</i>	<i>Element</i>	<i>Max</i>
Co	20	Si	5
Cr	30	Sn	5
Cu	5	Ta	10
H	Trace	Ti	3
Mg	Trace	V	15
Mn	30	W	15
Mo	10	Y	*
N	5	Zn	**
Nb	5	Zr	10

* Y systems are included mainly for the purpose of oxide dispersion strengthened (ODS) steels with many assessed oxygen containing binary and ternary systems within the Al-Cr-Cu-Fe-Mn-Ni-O-Si-Y-Zr frame of elements.

** The element Zn has been further treated with the focus on the Zn corner of Al-Cr-Fe-Zn system for galvanization process, but several other binaries and ternaries are also included.

TCFE14 Systems

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TCFE14 Assessed Binary Systems

371 binary systems are assessed.

TCFE14 Assessed Ternary Systems

335 ternary systems are assessed.

Assessed Ternary Systems							
Al-Ca-Fe	Al-Ca-O	Al-Ca-Si	Al-C-Fe	Al-C-Mn	Al-Co-Fe	Al-Co-Ni	Al-Co-Zr
Al-Cr-Fe	Al-Cr-Nb	Al-Cr-Ni	Al-Cr-O	Al-Cr-Zn	Al-C-Sn	Al-Cu-Fe	Al-Cu-Mn
Al-Cu-Ni	Al-Fe-H	Al-Fe-Mg	Al-Fe-Mn	Al-Fe-N	Al-Fe-Nb	Al-Fe-Ni	Al-Fe-O
Al-Fe-P	Al-Fe-S	Al-Fe-Si	Al-Fe-Ti	Al-Fe-Zn	Al-Fe-Zr	Al-H-Ti	Al-Mg-O
Al-Mg-Si	Al-Mg-Zn	Al-Mn-Ni	Al-Mn-O	Al-Nb-Ni	Al-Nb-Ti	Al-Ni-O	Al-Ni-Ti
Al-Ni-Zr	Al-O-Si	Al-O-Ti	Al-O-Y	Al-Ti-V	Al-Y-Zn	B-C-Fe	B-Co-Fe
B-Cr-Fe	B-Cr-Mn	B-Cr-Mo	B-Cr-Ni	B-Fe-Mn	B-Fe-Mo	B-Fe-Nb	B-Fe-Ni
B-Fe-Si	B-Fe-Ti	B-Fe-V	B-Fe-W	B-Fe-Zr	B-Mo-Ni	B-Mo-Ti	B-Ni-Si
B-Ni-Ti	B-Ni-Zr	B-Ti-Zr	Ca-Cr-O	Ca-Cr-S	Ca-Fe-O	Ca-Fe-S	Ca-Mg-O
Ca-Mg-S	Ca-Mn-O	Ca-Mn-S	Ca-Ni-O	Ca-O-Si	C-Ca-Fe	C-Co-Cr	C-Co-Fe
C-Co-Nb	C-Co-Ni	C-Co-Ti	C-Co-W	C-Co-Zn	C-Cr-Fe	C-Cr-Mn	C-Cr-Mo
C-Cr-N	C-Cr-Nb	C-Cr-Ni	C-Cr-Si	C-Cr-Ta	C-Cr-Ti	C-Cr-V	C-Cr-W
C-Cr-Zr	C-Cu-Fe	Ce-Fe-S	Ce-O-S	C-Fe-H	C-Fe-Mg	C-Fe-Mn	C-Fe-Mo
C-Fe-N	C-Fe-Nb	C-Fe-Ni	C-Fe-O	C-Fe-P	C-Fe-S	C-Fe-Si	C-Fe-Sn
C-Fe-Ti	C-Fe-V	C-Fe-W	C-Mg-Si	C-Mn-Si	C-Mn-V	C-Mo-N	C-Mo-Nb
C-Mo-Ta	C-Mo-Ti	C-Mo-V	C-Mo-W	C-Mo-Zr	C-Nb-Ti	C-Nb-V	C-Nb-W
C-Ni-Si	C-Ni-W	C-N-Nb	C-N-Ti	C-N-Zr	Co-Cr-Fe	Co-Cr-Ni	Co-Cr-W
Co-Cu-Fe	Co-Fe-Mo	Co-Fe-N	Co-Fe-Nb	Co-Fe-P	Co-Fe-S	Co-Fe-Si	Co-Fe-W
Co-Nb-Si	Co-Ni-W	Co-P-W	Co-Si-Ti	Co-Si-W	Co-Ti-Zr	Co-W-Zr	Cr-Cu-Fe
Cr-Cu-Mo	Cr-Cu-Ni	Cr-Cu-W	Cr-Fe-H	Cr-Fe-Mn	Cr-Fe-Mo	Cr-Fe-N	Cr-Fe-Nb

Assessed Ternary Systems							
Cr-Fe-Ni	Cr-Fe-O	Cr-Fe-P	Cr-Fe-Ru	Cr-Fe-S	Cr-Fe-Si	Cr-Fe-Sn	Cr-Fe-Ti
Cr-Fe-V	Cr-Fe-W	Cr-Fe-Zn	Cr-Fe-Zr	Cr-Mg-O	Cr-Mn-Mo	Cr-Mn-N	Cr-Mn-Ni
Cr-Mn-O	Cr-Mn-S	Cr-Mn-Si	Cr-Mo-N	Cr-Mo-Ni	Cr-Mo-Si	Cr-Nb-Ni	Cr-Nb-Si
Cr-Ni-O	Cr-Ni-Ru	Cr-Ni-S	Cr-Ni-Si	Cr-Ni-Ti	Cr-Ni-V	Cr-Ni-W	Cr-Ni-Zr
Cr-N-Nb	Cr-N-Ni	Cr-N-Si	Cr-N-Ti	Cr-N-V	Cr-N-W	Cr-O-Ti	Cr-O-Y
Cr-Si-Ti	Cr-Si-W	C-Si-Ti	C-Ta-W	C-Ti-V	C-Ti-W	C-Ti-Zr	Cu-Fe-Mn
Cu-Fe-Mo	Cu-Fe-N	Cu-Fe-Ni	Cu-Fe-P	Cu-Fe-S	Cu-Fe-Si	Cu-Fe-Sn	Cu-Fe-W
Cu-Fe-Zn	Cu-Fe-Zr	Cu-Mn-Ni	Cu-Mn-S	Cu-O-Sn	Cu-O-Y	C-V-W	C-V-Zr
C-W-Zr	Fe-H-Mo	Fe-H-Ni	Fe-H-Si	Fe-H-V	Fe-H-W	Fe-Mg-Ni	Fe-Mg-O
Fe-Mg-S	Fe-Mg-Si	Fe-Mn-Mo	Fe-Mn-N	Fe-Mn-Nb	Fe-Mn-Ni	Fe-Mn-O	Fe-Mn-P
Fe-Mn-S	Fe-Mn-Si	Fe-Mn-Sn	Fe-Mn-Ti	Fe-Mn-V	Fe-Mn-Zn	Fe-Mo-N	Fe-Mo-Ni
Fe-Mo-P	Fe-Mo-S	Fe-Mo-Si	Fe-Mo-V	Fe-Mo-W	Fe-Nb-Ni	Fe-Nb-O	Fe-Nb-P
Fe-Nb-S	Fe-Nb-Si	Fe-Nb-Sn	Fe-Nb-Ti	Fe-Nb-V	Fe-Nb-Zn	Fe-Nb-Zr	Fe-Ni-O
Fe-Ni-P	Fe-Ni-S	Fe-Ni-Si	Fe-Ni-Sn	Fe-Ni-Ti	Fe-Ni-V	Fe-Ni-W	Fe-Ni-Zn
Fe-N-Nb	Fe-N-Ni	Fe-N-Ti	Fe-N-V	Fe-N-W	Fe-O-S	Fe-O-Si	Fe-O-Sn
Fe-O-Y	Fe-P-Si	Fe-P-Ti	Fe-P-V	Fe-P-W	Fe-Si-Sn	Fe-Si-Ti	Fe-Si-W
Fe-Si-Zn	Fe-Si-Zr	Fe-Sn-W	Fe-Sn-Zn	Fe-Sn-Zr	Fe-S-Sn	Fe-S-Ta	Fe-S-Ti
Fe-S-V	Fe-S-W	Fe-S-Zn	Fe-S-Zr	Fe-Ti-V	Fe-Ti-Zr	H-Ni-Ti	H-Ti-Zr
Mg-Mn-O	Mg-Mn-S	Mg-Ni-O	Mg-O-Si	Mg-Si-Zn	Mn-Mo-Ni	Mn-Mo-Si	Mn-Ni-O
Mn-Ni-Si	Mn-O-S	Mn-O-Si	Mn-O-Y	Mn-Si-Zn	Mo-Ni-Si	Mo-Ni-Ti	Mo-N-Ni
Mo-N-V	Nb-Ni-Ti	Ni-O-Si	Ni-O-Ti	Ni-O-Y	Ni-P-W	Ni-Si-Ti	Ni-Si-W
Ni-Si-Zr	N-Nb-Ti	N-Nb-V	N-Ti-V	N-V-W	O-Si-Y	O-Y-Zr	

TCFE14 Assessed Quaternary Systems

84 quaternary systems are assessed.

Assessed Quaternary Systems					
Al-Ca-Mg-O	Al-Ca-O-Si	Al-C-Fe-Mn	Al-Cr-Fe-Ni	Al-Cr-Fe-O	Al-Cr-Fe-Zn
Al-Cr-Mg-O	Al-Cr-Mn-O	Al-Cr-Ni-O	Al-Cr-O-Y	Al-Fe-Mg-O	Al-Fe-Mn-O
Al-Fe-Ni-O	Al-Fe-O-Y	Al-Mg-Mn-O	Al-Mg-Ni-O	Al-Mg-O-Si	Al-Mn-Ni-O
Al-Mn-O-S	Al-O-Si-Y	B-Cr-Fe-Mo	Ca-Fe-Mg-S	Ca-Fe-Mn-S	Ca-Fe-O-Si
Ca-Mg-O-Si	C-Co-Cr-W	C-Co-Fe-Mo	C-Co-Fe-Ni	C-Co-Fe-W	C-Co-Nb-W
C-Co-Ni-W	C-Co-V-W	C-Cr-Fe-Mn	C-Cr-Fe-Mo	C-Cr-Fe-N	C-Cr-Fe-Ni
C-Cr-Fe-V	C-Cr-Fe-W	C-Cr-Mn-V	C-Cr-Mn-W	C-Cr-Mo-V	C-Cr-Mo-W
C-Cr-V-W	C-Fe-Mn-H	C-Fe-Mn-Nb	C-Fe-Mn-Si	C-Fe-Mo-Nb	C-Fe-Mo-Si
C-Fe-Mo-V	C-Fe-Mo-W	C-Fe-Nb-W	C-Fe-Ni-W	C-Fe-N-Ni	C-Fe-Si-W
C-Fe-V-W	C-Mo-N-Ni	C-Mo-Ti-V	C-Mo-V-W	C-N-Nb-Ti	C-N-Nb-V
C-N-Ti-V	Cr-Fe-H-Ni	Cr-Fe-Mg-O	Cr-Fe-Mn-N	Cr-Fe-Mn-O	Cr-Fe-Mo-N

TCFE14 Phases

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Common Phases for Steel and Fe-alloys



TCFE14 Models for the Included Phases

The following lists common phase names and the corresponding Thermo-Calc database phase names for some key steel and Fe-alloys.

Name in the Database	Common Name and Description
FCC_A1	Austenite (γ) as well as cubic carbonitrides (MeX) such as MoC, TiC, WC and ZrC
BCC_A2	Ferrite (α)
HCP_A3	Hcp metals as well as Me ₂ X carbonitrides (ε) such as Cr ₂ N, Mo ₂ C, Ta ₂ C, V ₂ C, and W ₂ C
CEMENTITE_D011	Cementite (θ), Fe ₃ C, Mn ₃ C, Ni ₃ B, and Co ₃ B
C14_LAVES	This phase includes all MgZn ₂ -type phases such as Fe ₂ Mo, Fe ₂ Nb, Fe ₂ Ta, Fe ₂ Ti, and Fe ₂ W
CHI_A12	(χ), an intermetallic compound containing primarily Fe, Cr, and Mo
GRAPHITE	Graphite
M6C_E93	M6C carbides such as W ₃ Fe ₃ C
M7C3_D101	M7C3 carbides such as Mn ₇ C ₃ and Cr ₇ C ₃
M23C6_D84	M23C6 carbides such as Cr ₂₃ C ₆ and Mn ₂₃ C ₆
SIGMA_D8B	Sigma phase (σ), a chromium/molybdenum-rich intermetallic compound which is hard, brittle and non-magnetic.
MU_D85	μ phase, such as W ₆ Fe ₇
MS_B1	Rocksalt (B1) structure sulfides such as Alabandite (MnS) and Niningerite (MgS)
PI_A13	a Cr-Ni nitride (π)
M5C2	Mn ₅ C ₂ , also metastable Fe ₅ C ₂ (χ)
M3C2_D510	Cr ₃ C ₂ (β)
MC_ETA	MoC _{1-x} (η)
MC_SHP	MoC, WC

Name in the Database	Common Name and Description
R_PHASE	a high temperature Fe-Mo phase
CORUNDUM	M ₂ O ₃ oxides such as Al ₂ O ₃ (Corundum), Cr ₂ O ₃ (Eskolaite), and Fe ₂ O ₃ (Hematite)
HALITE	MO oxides such as FeO (Wustite), MnO, NiO
SPINEL	M ₃ O ₄ oxides such as Fe ₃ O ₄ (Magnetite) and Cr ₃ O ₄
M2P_C22	Phosphides such as Fe ₂ P, Cr ₂ P, Ni ₂ P
M3P_D0E	Phosphide such as Fe ₃ P, Cr ₃ P, Ni ₃ P

TCFE14 Models for the Included Phases

The crystal structure information for all included phases in the database is listed here. Similar information is also available when working in the Console Mode by using the DATABASE_RETRIEVAL module. You can display the information using the LIST_SYSTEM CONSTITUENTS and after the system is defined.



The DICTRA_FCC_A1 phases are always rejected by default. It can be restored in the TDB Module (in Console Mode) if it is necessary for your system.

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
GAS	Gas					1	(AL, AL1B1O2, AL1B3H12, AL1C1, AL1C2, AL1CU1, AL1CU1S1, AL1CU1S2, AL1H1, AL1H1O1_ALOH, AL1H1O1_HALO, AL1H1O2, AL1H2, AL1H2O2, AL1H3, AL1H3O3, AL1N1, AL1O1, AL1O2, AL1P1, AL1P2, AL1S1, AL1S2, AL2, AL2C2, AL2C6H18, AL2O1, AL2O2, AL2O3, AL2S1, AL2S2, AR, B, B10H14, B1C1, B1C1H3O1, B1C2, B1C2H7O2, B1C3H9, B1C3H9O3, B1C6H15, B1H1, B1H1O1_BOH, B1H1O1_HBO, B1H1O2, B1H1S1, B1H2, B1H2O1, B1H2O2, B1H3, B1H3O1, B1H3O2, B1H3O3, B1H6N1, B1N1, B1O1, B1O2, B1S1, B1S2, B2, B2C1, B2H4O4, B2H6, B2O1, B2O2, B2O3, B2S1, B2S2, B2S3, B3H3O3, B3H3O6, B3H6N3, B4S6, B5H9, C, C1H1, C1H1N1_HCN, C1H1N1_HNC, C1H1N1O1, C1H1N1S1, C1H1O1, C1H1O2, C1H1P1, C1H2, C1H2N4, C1H2O1, C1H2O2_CIS, C1H2O2_DIOXIRANE, C1H2O2_TRANS, C1H3, C1H3O1_CH2OH, C1H3O1_CH3O, C1H3P1, C1H4, C1H4N2O1, C1H4O1, C1H4S1, C1H5N1, C1H5O1P1, C1H5O3P1, C1H5P1, C1H5P1S1, C1H6N1P1_N, C1H6N1P1_P, C1H6P2, C1N1, C1N1O1, C1N1O1_NCO, C1N2_NCN, C1N2_NCN, C1O1, C1O1S1, C1O2, C1P1, C1P1S1, C1P1S1I2, C1P2, C1S1, C1S2, C1S1I, C1S2I, C1S3I, C1S4I, C2, C2H1, C2H1N1, C2H2, C2H2O1, C2H3, C2H4, C2H4O1_ACETALDEHYDE, C2H4O1_OXIRANE, C2H4O2_ACETICACID, C2H4O2_DIOXETANE, C2H4O3_123TRIOXOLANE, C2H4O3_124TRIOXOLANE, C2H5, C2H6, C2H6O1_1, C2H6O1_2, C2H6O1S1, C2H6O2, C2H7O1P1, C2H7O3P1, C2H7P1_1, C2H7P1_2, C2H7P1S1, C2H8N1P1_N, C2H8N1P1_P, C2H8S1I1)

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
							C2N1_CCN, C2N1_CNC, C2N2, C2O1, C2P1, C2P2, C2Si1, C2Si2, C2Si3, C3, C3H1, C3H1N1, C3H4_1, C3H4_2, C3H6_1, C3H6_2, C3H6O1_1, C3H6O1_2, C3H8, C3N1, C3O2, C4, C4H1, C4H10_1, C4H10_2, C4H12S1, C4H2_1, C4H2_2, C4H4_1, C4H4_2, C4H6_1, C4H6_2, C4H6_3, C4H6_4, C4H6_5, C4H8_1, C4H8_2, C4H8_3, C4H8_4, C4H8_5, C4H8_6, C4N1, C4N2, C4Ni1O4, C5, C5Fe1O5, C5H1N1, C5N1, C6O, C6H6, C6H6O1, C6MO1O6, C6N1, C6N2, C9N1, CA, CA1H1, CA1H1O1, CA1H2O2, CA1O1, CA1S1, CA2, CE, CE1O1, CE1S1, CO, CO1H1, CO1H1O1, CO1H2O2, CO1O1, CO1S1, CO2, CR, CR1H1, CR1H1O1, CR1H1O2, CR1H1O3, CR1H2O2, CR1H2O3, CR1H2O4, CR1H3O3, CR1H3O4, CR1H4O4, CR1H4O5, CR1N1, CR1O1, CR1O2, CR1O3, CR1S1, CR1S2, CR2, CR2O1, CR2O2, CR2O3, CU, CU1H1, CU1H1O1, CU1O1, CU1S1, CU2, CU2S1, FE, FE1H1, FE1H1O1, FE1H1O2, FE1H2O2, FE1O1, FE1O2, FE1S1, FE2, H, H1MG1, H1MG1O1, H1MN1, H1MN1O1, H1MO1O1, H1MO1O2, H1MO3, H1N1, H1N1O1, H1N1O2_CIS, H1N1O2_TRANS, H1N1O3, H1N3, H1N1I, H1N1O1, H1O1, H1O1P1, H1O1S1_HSO, H1O1S1_SOH, H1O1W1, H1O1Z1N1, H1O2, H1O2W1, H1P1, H1S1, H1S1I, H1Z1N1, H1Z1R1, H2, H2MG1O2, H2M01O2, H2M01O3, H2M01O4, H2N1, H2N2_1_1N2H2, H2N2_CIS, H2N2_TRANS, H2N2O2, H2N1O2, H2O1, H2O1S1_H2O, H2O1S1_HSOH, H2O2, H2O2W1, H2O2Z1N1, H2O3S1, H2O3W1, H2O4S1, H2O4W1, H2P1, H2S1, H2S2, H2S1I, H3N1, H3N1O1, H3P1, H3S1I, H4N2, H4O4S1, H4S1I, H4SN1, H6S12, MG, MG1N1, MG1O1, MG1S1, MG2, MN, MN1O1, MN1O2, MN1S1, MO, MO1N1, MO1O1, MO1O2, MO1O3, MO1S1, MO1S2, MO2, MO2O6, MO3O9, MO4O12, MO5O15, N, N1NB1, N1O1, N1O2, N1O3, N1P1, N1S1, N1S12, N1T1I, N1V1, N1ZR1, N2, N2O1, N2O2, N2O3, N2O4, N2O5, N3, NB, NB1O1, NB1O2, NB1S1, NI, NI1O1, NI1S1, NI2, O, O1O4P, O1O4V, O12W4, O15W5, O1P1, O1RU1, O1S1, O1S2, O1S1I, O1S1N1, O1TA1, O1T1I, O1V1, O1W1, O1Y1, O1Y2, O1Z1N1, O1Z1R1, O2, O2P1, O2RU1, O2S1, O2Si1, O2Si2, O2SN1, O2TA1, O2T1, O2V1, O2W1, O2Y1, O2Y2, O2Z1R1, O3, O3P2, O3RU1, O3S1, O3W1, O4P2, O4RU1, O5P2, O6P3, O6P4, O6W2, O7P4, O8P4, O8W3, O9P4, O9W3, P, P1S1, P1S1I, P1S12, P2, P2S1I, P3, P4, P4S3, RU, S, S1S1, S1S1N1, S1TA1, S1T1, S1V1, S1W1, S1Y1, S1Z1N1, S1Z1R1, S2, S2S1I, S2SN1, S2SN2, S2T1, S2W1, S2ZR1, S3, S4, S5, S6, S7, S8, SI, SI2, SI3, SN, SN2, TA, TI, TI2, V, W, Y, ZN, ZR, ZR2)
LIQUID	Liquid				1		(AL, AL2/3S, ALN, ALO3/2, B, B1N1, C, CA, CA2SN, CAC2, CAS, CE, CEO2, CEO3/2, CES, CO, COS, CR, CRO3/2, CRS, CU, CU2O, CU2S, CUO, FE, FEO, FEO3/2, FES, H, MG, MG2SN1, MGS, MIN, MNO, MNO3/2, MNS, MO, MO1/2S, N, NB, NBO, NBO2, NBS, NI, NIO, NIS, P, RU, S, S1Z1R1/2, SI, SI1/2S, S1O2, SN, SNO, SNO2, SNS, SZN, TA, TI, TIO, TIO2, TIO3/2, TIS, V, VS, W, Y,

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
							Y2/3O, ZN, ZR, ZR1/2O)1
BCC_A2	Body-Centered Cubic (W, A2, bcc)	A2	cl2	(229, Im-3m)		2	(AL, CA, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, S, SI, SN, TA, Ti, V, W, Y, ZN, ZR)1(B, C, H, N, O, VA)3
A2_BCC	Body-Centered Cubic (W, A2, bcc)	A2	cl2	(229, Im-3m)		2	(AL, CA, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, S, SI, SN, TA, Ti, V, W, Y, ZN, ZR)1(B, C, H, N, O, VA)3
BCC_B2	CsCl (B2)	B2	cP2	(221, Pm-3m)		3	(AL, CA, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, S, SI, SN, TA, Ti, V, W, Y, ZN, ZR)0.5(AL, CA, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, S, SI, SN, TA, Ti, V, W, Y, ZN, ZR)0.5(B, C, H, N, O, VA)3
FCC_A1	Face-Centred Cubic (Cu, A1, fcc)	A1	cF4	(225, Fm-3m)		2	(AL, CA, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, S, SI, SN, TA, Ti, V, W, Y, ZN, ZR)1(B, C, H, N, O, VA)1
A1_FCC	Face-Centred Cubic (Cu, A1, fcc)	A1	cF4	(225, Fm-3m)		2	(AL, CA, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, S, SI, SN, TA, Ti, V, W, Y, ZN, ZR)1(B, C, H, N, O, VA)1
DICTRA_FCC_A1	Face-Centred Cubic (Cu, A1, fcc)	A1	cF4	(225, Fm-3m)		2	(AL, CA, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, S, SI, SN, TA, Ti, V, W, Y, ZN, ZR)1(B, C, H, N, O, VA)1
FCC_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)		3	(AL, CA, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, S, SI, SN, TA, Ti, V, W, Y, ZN, ZR)0.75(AL, CA, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, S, SI, SN, TA, Ti, V, W, Y, ZN, ZR)0.25(B, C, H, N, O, VA)1
HCP_A3	Hexagonal Close Packed (Mg, A3, hcp)	A3	hP2	(194, P6_3/mmc)		2	(AL, CA, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, S, SI, SN, TA, Ti, V, W, Y, ZN, ZR)1(B, C, H, N, O, VA)0.5
BCT_A5	beta-Sn (A5)	A5	tI4	(141, I4_1/amd)		1	(AL, CA, CU, S, SN, Ti, ZN)1
CBCC_A12	alpha-Mn (A12)	A12	cl58	(217, I-43m)		2	(AL, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, SI, SN, TA, Ti, V, ZN, ZR)1(B, C, N, VA)1
CUB_A13	beta-Mn (A13)	A13	cP20	(213, P4_132)		2	(AL, CE, CO, CR, CU, FE, MG, MN, MO, NB, NI, P, RU, SI, SN, TA, Ti, V, Y, ZN, ZR)1(B, C, N, VA)1
DIAMOND_A4	Diamond (A4)	A4	cF8	(227, Fd-3m)		1	(AL, B, C, MN, O, P, SI, SN, ZN)1

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
BETA_RHOMBO_B	beta-B (R-105)		hR105	(166, R-3m)		2	(B)93(B, C, Si)12
ORTHORHOMBIC_S	alpha-S (A16)	A16	oF128	(70, Fddd)		1	(S)1
RED_P	Unknown Structure					1	(P)1
WHITE_P	Unknown Structure					1	(P)1
GRAPHITE_A9	Hexagonal Graphite (A9)	A9	hP4	(194, P6_3/mmc)		1	(B, C)1
CEMENTITE_D011	Cementite (Fe3C, D011)	D011	oP16	(62, Pnma)		2	(Al, Co, Cr, Fe, Mn, Mo, Nb, Ni, Si, V, W)3(B, C, N)1
M23C6_D84	Cr23C6 (D84)	D84	cF116	(225, Fm-3m)		3	(Co, Cr, Fe, Mn, Ni, V)20(Co, Cr, Fe, Mn, Mo, Ni, V, W)3(B, C)6
M7C3_D101	C3Cr7 (D101)	D101	oP40	(62, Pnma)		2	(Al, Co, Cr, Fe, Mn, Mo, Nb, Ni, Si, Ti, V, W)7(B, C)3
M6C_E93	Fe3W3C (E93)	E93	cF112	(227, Fd-3m)		4	(Co, Fe, Ni)2(Mo, Nb, W)2(Co, Cr, Fe, Mo, Nb, Ni, Si, V, W)2(C)1
M5C2	Mn5C2 (Fe5C2 Hagg carbide)		mS28	(15, C2/c)		2	(Fe, Mn, Nb, V)5(C, N)2
M3C2_D510	Tongbaite (Cr3C2, D510)	D510	oP20	(62, Pnma)		2	(Co, Cr, Mo, Ti, V, W)3(C)2
MC_ETA	CMo		hP12	(194, P6_3/mmc)		2	(Mo, Ti, V, W)1(C, Va)1
MC_SHP	Tungsten Carbide (Bh)	Bh	hP2	(187, P-6m2)		2	(Mo, W)1(C, N)1
KSI_CARBIDE	Mo6Fe11C5		mS44	(12, C2/m)		2	(Cr, Fe, Mo, W)3(C)1
Z_PHASE	CrNbN		tP6	(129, P4/nmm)		3	(Cr, Fe)1(Mo, Nb, V)1(N, Va)1
FE4N_LP1	gama-Fe4N (L'10)	L'10	cP5	(221, Pm-3m)		2	(Co, Cr, Fe, Mn, Ni)4(C, N, Va)1
PI_A13	beta-Mn (A13)	A13	cP20	(213, P4_132)		3	(Cr)12.8(Fe, Ni, Si)7.2(N)4

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
SIGMA_D8B	sigma-CrFe (D8b)	D8b	tP30	(136, P4_2/mnm)		3	(AL, CO, CR, FE, MN, NI, RU, TA, V)10(CR, MO, NB, TA, TI, V, W)4(AL, CO, CR, FE, MN, MO, NB, NI, RU, SI, TA, TI, V, W)16
MU_D85	Fe7W6 (D85) mu-phase	D85	hR13	(166, R-3m)		3	(AL, CO, CR, FE, MN, NB, NI, TA)7(MO, NB, TA, TI, V, W)2(CO, CR, FE, MO, NB, NI, TA, TI, W)4
P_PHASE	Cr9Mo21Ni20		oP56	(62, Pnma)		3	(CR, FE, NI)24(CR, FE, MO, NI)20(MO)12
R_PHASE	R-(Co,Cr,Mo)		hR53	(148, R-3)		3	(CO, CR, FE, MN, NI)27(MO, V, W)14(CO, CR, FE, MN, MO, NI, W)12
CHI_A12	alpha-Mn (A12)	A12	cI58	(217, I-43m)		3	(CR, FE, NI)24(CR, MO, TI, W, ZR)10(CR, FE, MO, NI, W)24
C14_LAVES	MgZn2 Hexagonal Laves (C14)	C14	hP12	(194, P6_3/mmc)		2	(AL, CO, CR, CU, FE, MG, MN, MO, NB, NI, SI, TA, TI, V, W, Y, ZN, ZR)2(AL, CO, CR, CU, FE, MG, MN, MO, NB, NI, SI, TA, TI, V, W, Y, ZN, ZR)1
C15_LAVES	Cu2Mg Cubic Laves (C15)	C15	cF24	(227, Fd-3m)		2	(AL, CA, CE, CO, CR, CU, FE, MG, NB, NI, SI, SN, Y, ZN, ZR)2(AL, CA, CE, CO, CR, CU, FE, MG, NB, NI, SI, SN, Y, ZN, ZR)1
C36_LAVES	MgNi2 Hexagonal Laves (C36)	C36	hP24	(194, P6_3/mmc)		2	(FE, MG, NI, ZN)2(MG, NB, NI)1
M3SI_D03	BiF3 (D03)	D03	cF16	(225, Fm-3m)		2	(FE, MN)3(SI)1
MN9Si2	Mn9Si2		oI186	(71, Immm)		2	(MN)0.81818(SI)0.18182
MN11Si19	Mn11Si19		tP120	(118, P-4n2)		2	(MN)0.367(SI)0.633
MN6Si	Fe7W6 (D85) mu-phase	D85	hR13	(166, R-3m)		2	(MN)0.85714(SI)0.14286
G_PHASE	Th6Mn23 (D8a)	D8a	cF116	(225, Fm-3m)		3	(AL, CO, CU, FE, NI, TI)16(MN, NB, TI, Y, ZR)6(CO, CU, FE, NI, SI)7
ETA_M5SiN	Fe3W3C (E93)	E93	cF112	(227, Fd-3m)		4	(CR, MO)3(FE, NI)2(SI)1(N)1
CR3Si_A15	Cr3Si (A15)	A15	cP8	(223, Pm-3n)		2	(CO, CR, FE, MO, NB, SI, V)3(AL, CO, CR, NB, SI, V)1
FESi2_H	FeSi2-h		tP3	(123,		2	(FE)0.3(MG, SI)0.7

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
				P4/mmm)			
FESI2_L	FeSi2-I		oS48	(64, Cmce)		2	(FE)0.333333(SI)0.666667
MSI_B20	FeSi (B20)	B20	cP8	(198, P2_13)		2	(CO, CR, FE, MN, NI, RU)0.5(AL, MG, SI)0.5
M5Si3_D88	Mavlyanovite (Mn5Si3, D88)	D88	hP16	(193, P6_3/mcm)		3	(CR, FE, MN, NI, TI, Y)5(SI)3(C, VA)1
MG2NI_CA	Mg2Ni (Ca)	Ca	hP18	(180, P6_222)		2	(MG)2(NI)1
MG2Si_C1	Fluorite (CaF2, C1)	C1	cF12	(225, Fm-3m)		2	(MG)2(SI, SN)1
Ni3Nb_D0a	beta-TiCu3 (D0a)	D0a	oP8	(59, Pmmn)		2	(FE, NB, NI)3(FE, MO, NB, NI, TI)1
NiT2	NiTi2		cF96	(227, Fd-3m)		2	(FE, NI)0.3333(TI)0.6667
Ni3Ti_D024	Ni3Ti (D024)	D024	hP16	(194, P6_3/mmc)		2	(FE, NI, TI)3(MO, NB, NI, TI)1
NIY_B27	FeB (B27)	B27	oP8	(62, Pnma)		2	(NI)1(Y)1
FE17Y2	Fe17Lu2		hP80	(194, P6_3/mmc)		2	(CO, FE, NI, ZN)17(Y)2
NI2Y1	Ni2Tm		cF192	(216, F-43m)		2	(NI)2(Y)1
NI2Y3	Ni2Y3		tP80	(92, P4_12_12)		2	(NI)2(Y)3
NI3Y	Ni3Pu		hR12	(166, R-3m)		2	(CO, FE, NI)3(Y)1
NI4Y	Unknown Structure		hR*			2	(NI)4(Y)1
M7Y2	Co7Gd2		hR18	(166, R-3m)		2	(CO, NI)7(CA, Y)2
NI5Y_D2D	CaCu5 (D2d)	D2d	hP6	(191, P6/mmm)		2	(NI)5(Y)1

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
ALY_B33	CrB (B33)	B33	oS8	(63, Cmcm)		2	(AL)1(Y)1
AL2Y3	Zr3Al2		tP20	(136, P4_2/mnm)		2	(AL, ZN)2(Y, ZR)3
ALY2_C37	Co2Si (C37)	C37	oP12	(62, Pnma)		2	(AL)1(Y)2
AL3Y_HT	BaPb3		hR12	(166, R-3m)		2	(AL)0.75(Y)0.25
CO5Y_D2D	CaCu5 (D2d)	D2d	hP6	(191, P6/mmm)		3	(CO2, Y)1(CO)4(CO, VA)1
CO3Y2	Unknown Structure		cP*			2	(CO)3(Y)2
CO7Y6	Unknown Structure					2	(CO)7(Y)6
COY_B33	CrB (B33)	B33	oS8	(63, Cmcm)		2	(CO)1(Y)1
CO3Y4	Co3Ho4		hP22	(176, P6_3/m)		2	(CO)3(Y)4
CO5Y8	Co5Y8		mP52	(14, P2_1/c)		2	(CO)5(Y)8
SI2Y_C32	Hexagonal omega (C32)	C32	hP3	(191, P6/mmm)		2	(Y)1(SI)2
SI2Y_CC	alpha-ThSi2 (Cc)	Cc	tI12	(141, I4_1/amd)		2	(CE, Y)1(SI)2
SI5Y3_CC	alpha-ThSi2 (Cc)	Cc	tI12	(141, I4_1/amd)		2	(Y)3(SI)5
SI5Y3_C32	Hexagonal omega (C32)	C32	hP3	(191, P6/mmm)		2	(Y)3(SI)5
SI4Y5	Gd5Si4		oP36	(62, Pnma)		2	(Y)5(SI, SN)4
SIY_B33	CrB (B33)	B33	oS8	(63, Cmcm)		2	(Y)1(SI)1
CU2Y_H	Unknown Structure		hP*			2	(CU)2(Y)1

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
CU6Y	Cu7Tb		hP8	(191, P6/mmm)		2	(CU2, Y)1(CU)5
CU4Y	Cu5Y1.25		mP16	(11, P2_1/m)		2	(CU)4(Y)1
CU7Y2	Ag51Gd14		hP68	(175, P6/m)		2	(CU)7(Y)2
AL3NB_D022	Al3Ti (D022)	D022	tI8	(139, I4/mmm)		2	(AL)3(NB)1
CO3MO_D019	Ni3Sn (D019)	D019	hP8	(194, P6_3/mmc)		2	(CO)3(MO)1
CO3V	Al3Pu		hP24	(194, P6_3/mmc)		2	(CO, V)3(CO, V)1
MNTA	Unknown Structure					2	(MN)1(TA)1
MOSI2_C11B	MoSi2 (C11b)	C11b	tI6	(139, I4/mmm)		2	(MO)1(SI)2
MO5Si3_D8M	W5Si3 (D8m)	D8m	tI32	(140, I4/mcm)		2	(MO)5(SI)3
GAMMA2_ALFEZN	Unknown Structure					2	(AL, FE, ZN)0.255(ZN)0.745
FE2SITI_L21	Heusler (L21)	L21	cF16	(225, Fm-3m)		3	(FE)0.5(AL, FE, NB, SI, SN, TI)0.25(AL, FE, NB, SI, SN, TI)0.25
FESITI	TiFeSi		oI36	(46, Ima2)		3	(FE)1(SI)1(TI)1
FESI2TI	CrSi2Zr		oP48	(55, Pbam)		3	(FE)0.25(SI)0.5(TI)0.25
FE4Si3TI	Pd40Sn31Y13		hP168	(191, P6/mmm)		3	(FE)0.5(SI)0.375(TI)0.125
AL8Mn5_D810	Cr5Al8 (D810)	D810	hR26	(160, R3m)		3	(AL)12(MN)5(AL, FE, MN)9
NB5Si3_D8L	Cr5B3 (D8l)	D8l	tI32	(140, I4/mcm)		2	(CR, NB)5(SI)3
NB5Si3_D8M	W5Si3 (D8m)	D8m	tI32	(140, I4/mcm)		3	(NB)4(CR, NB, SI)1(SI)3

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
MSI2_C40	CrSi2 (C40)	C40	hP9	(180, P6_222)		2	(CR, NB)1(SI)2
CRNBSI	ZrNiAl		hP9	(189, P-62m)		3	(CR)1(NB)1(SI)1
M11Si8	Cr11Ge8		oP76	(62, Pnma)		2	(CR, NB)11(SI)8
M6Si5	Si5V6		oI44	(72, Ibam)		2	(CR, NB)6(SI)5
FENBSI2	CrSi2Zr		oP48	(55, Pbam)		3	(FE)1(NB)1(SI)2
FE4NB4Si7	Co4Ge7Zr4		tI60	(139, I4/mmm)		3	(FE)4(NB, ZR)4(SI)7
FENBSI_C37	MnCuP		oP12	(62, Pnma)		3	(FE)1(NB, ZR)1(SI)1
FE3Nb4Si5	Fe3Nb4Si5		oP72	(31, Pmn2_1)		3	(FE)3(NB)4(SI)5
FENB2Si2	FeNb2Si2		tP198	(132, P4_2/mcm)		3	(FE)1(NB)2(SI)2
KAPPA_E21	Cubic Perovskite (CaTiO3, E21)	E21	cP5	(221, Pm-3m)		3	(AL)1(FE, MN)3(C, VA)1
AL4C3_D71	Al4C3 (D71)	D71	hR7	(166, R-3m)		2	(AL, SI)4(C)3
B4C_D1G	B13C2 B4C (D1g)	D1g	hR15	(166, R-3m)		2	(B11C, B12)1(B2, C2B, CB2)1
M12C	Fe6W6C		cF104	(227, Fd-3m)		3	(CO)6(W)6(C)1
FE8Si2C	Mn8Si2C		aP*	(1, P1)		3	(FE, MN)8(SI)2(C)1
MGC2	MgC2		tP6	(136, P4_2/mnm)		2	(MG)1(C)2
SIC_B3	Zincblende (ZnS, B3)	B3	cF8	(216, F-43m)		2	(SI)1(C)1
MN5SiC	Mn5SiC		oS56	(36, Cmc2_1)		3	(MN)0.714(SI)0.143(C)0.143

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
Y15C19_ALPHA	alpha-Y15C19		oP18	(55, Pbam)		2	(C)19(Y)15
Y15C19_BETA	Unknown Structure					2	(C)19(Y)15
Y2C3_ALPHA	Sc3C4		tP70	(128, P4/mnc)		3	(Y)2(C)2(C, VA)1
Y2C3_BETA	Unknown Structure					3	(Y)2(C)2(C, VA)1
YC_B1	Rock Salt (NaCl, B1)	B1	cF8	(225, Fm-3m)		2	(Y)1(C, C2, VA)1
MY3_D011	Cementite (Fe3C, D011)	D011	oP16	(62, Pnma)		2	(CO, NI)1(Y)3
K_PHASE	Cubic Perovskite (CaTiO ₃ , E21)	E21	cP5	(221, Pm-3m)		3	(C, CO, ZN)51(ZN)32(C)17
CA3ZN_E1A	Re3B		oS16	(63, Cmcm)		2	(CA)3(ZN)1
CA5ZN3_D8L	Cr5B3 (D8I)	D8I	tl32	(140, I4/mcm)		2	(CA)5(SN, ZN)3
CAZN_B33	CrB (B33)	B33	oS8	(63, Cmcm)		2	(SN, ZN)1(CA)1
CAZN2	KHg2		ol12	(74, Imma)		2	(CU, ZN)2(CA, CE, Y)1
CAZN3	CaZn3		hP32	(194, P6_3/mmc)		2	(CA)1(ZN)3
CAZN11	BaCd11		tl48	(141, I4_1/amd)		2	(CA, CE)1(ZN)11
CAZN13_D23	NaZn13 (D23)	D23	cF112	(226, Fm-3c)		2	(CA)1(ZN)13
COZN7	CoZn7		hP60	(171, P6_2)		2	(CO)1(ZN)7
CO2ZN15	Co2Zn15		mS28	(12, C2/m)		2	(CO)2(ZN)15
COZN_A13	beta-Mn (A13)	A13	cP20	(213, P4_132)		1	(CO, ZN)1

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
CRZN17	Unknown Structure		hP*			2	(AL, CR)1(FE, ZN)17
FE3ZN7_D82	gamma-brass (Cu5Zn8, D82)	D82	cI52	(217, I-43m)		4	(CO, CU, FE, MN, NI, ZN)0.154(CU, FE, MN, ZN)0.154(AL, CO, CU, FE, MN, NI, SI, ZN)0.231(ZN)0.461
FEZN4	Fe11Zn40		cF408	(216, F-43m)		3	(FE)0.137(AL, FE, NI, SI, SN, ZN)0.118(MN, ZN)0.745
FEZN10	FeZn10		hP632	(194, P6 ₃ /mmc)		4	(CR, FE)0.058(AL, FE, MN, NI, SI, SN, ZN)0.18(ZN)0.525 (ZN)0.237
FEZN13	CoZn13		mS28	(12, C2/m)		3	(CO, CR, FE, MN, NI, VA)0.072(AL, SI, VA, ZN)0.072(AL, ZN)0.856
FENIZN_T	Zn89(Fe0.5Ni0.5)13.8		cF432	(216, F-43m)		2	(FE, NI, ZN)1(ZN)5
CUZN_EPSILON	Hexagonal Close Packed (Mg, A3, hcp)	A3	hP2	(194, P6 ₃ /mmc)		2	(CU, MN, ZN)1(VA)0.5
MG2ZN3	Mg4Zn7		mS110	(12, C2/m)		2	(MG)2(AL, ZN)3
MG12ZN13	Zr21Re25		hR92	(167, R-3c)		2	(MG)12(AL, ZN)13
MG51ZN20	Mg51Zn20		oI158	(71, Immm)		2	(MG)51(ZN)20
MG2ZN11_D8C	Mg2Zn11 (D8c)	D8c	cP39	(200, Pm-3)		2	(MG)2(AL, SI, ZN)11
MNZN9	Unknown Structure		h**			2	(MN)0.1(ZN)0.9
MOZN7	Ca7Ge		cF32	(225, Fm-3m)		2	(MO)1(ZN)7
MOZN22	Zn93(Zn0.43Mo0.57)Mo4		cF420	(216, F-43m)		2	(MO)1(ZN)22
NIZN8	Ni3Zn22		mS50	(12, C2/m)		2	(FE, NI)1(ZN)8
NIZN_L2A	delta-CuTi (L2a)	L2a	tP2	(123, P4/mmm)		2	(NI, ZN)1(NI, ZN)1
ZN3P2	Zn3P2 (D59)	D59	tP40	(137, P4 ₂ /nmc)		2	(ZN)3(P)2

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
ZNP2	ZnAs2		mP24	(14, P2_1/c)		2	(ZN)1(P)2
ZNS_B3	Zincblende (ZnS, B3)	B3	cF8	(216, F-43m)		2	(FE, ZN)1(S)1
ZNS_B4	Wurtzite (ZnS, B4)	B4	hP4	(186, P6_3mc)		2	(FE, ZN)1(S)1
Ti2ZN	CuZr2		tI6	(139, I4/mmm)		2	(TI, ZR)2(ZN)1
TiZN3_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)		2	(NB, TI, V)1(ZN)3
TiZN5	Unknown Structure					2	(TI)1(ZN)5
TiZN10	Ti3Zn22		tP100	(135, P4_2/mbc)		2	(TI)1(ZN)10
TiZN15	TiZn16		oS68	(63, Cmcm)		2	(NB, TI)1(ZN)15
V4ZN5	V4Zn5		tI18	(139, I4/mmm)		2	(V)4(ZN)5
A_YZN2	Unknown Structure					2	(Y)0.3333(ZN)0.6667
B_YZN2	Unknown Structure					2	(Y)0.3333(ZN)0.6667
YZN3	Zn3Y		oP16	(62, Pnma)		2	(Y)0.25(ZN)0.75
Y3ZN11	Al11La3		oI28	(71, Immm)		2	(CE, Y)0.2143(ZN)0.7857
Y13ZN58	Y13Zn58		hP146	(194, P6_3/mmc)		2	(Y)0.1831(ZN)0.8169
YZN5	ErZn5		hP36	(194, P6_3/mmc)		2	(Y)0.1667(ZN)0.8333
YZN12_D2B	Mn12Th (D2b)	D2b	tI26	(139, I4/mmm)		2	(Y)0.0769(ZN)0.9231
YZN_B2	CsCl (B2)	B2	cP2	(221, Pm-3m)		2	(CE, TI, Y, ZR)0.5(ZN)0.5

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
ZN22ZR	Zn22Zr		cF184	(227, Fd-3m)		2	(ZN)0.9565(ZR)0.0435
ZN39ZR5	Zn39Zr5		mS88	(12, C2/m)		2	(ZN)0.8864(ZR)0.1136
ZN3ZR_D023	Al3Zr (D023)	D023	tI16	(139, I4/mmm)		2	(ZN)0.75(ZR)0.25
ZN3ZR_HT	Unknown Structure		c**			2	(ZN)0.75(ZR)0.25
ALMGZN_T2	Mg46Zn37Al17		cP640	(205, Pa-3)		3	(AL)0.15(MG)0.43(ZN)0.42
ALMGZN_Q	Quasicrystal					3	(AL)0.15(MG)0.44(ZN)0.41
ALMG_BETA	Al45Mg28		cF1832	(227, Fd-3m)		2	(AL, ZN)140(MG)89
ALMG_EPS	Al30Mg23		hR53	(148, R-3)		2	(AL, ZN)30(MG)23
AL12MG17_A12	alpha-Mn (A12)	A12	cI58	(217, I-43m)		3	(MG)10(AL, MG, ZN)24(AL, MG, ZN)24
ALMGZN_PHI	Mg21(Al,Zn)17		oP152	(57, Pbcm)		2	(MG)21(AL, ZN)17
ALMGZN_T1	Bergman [Mg32(Al,Zn)49, D8e]	D8e	cI162	(204, Im-3)		4	(MG)26(AL, MG)6(AL, MG, ZN)48(AL)1
AL2FE	Al2Fe		aP18	(1, P1)		3	(AL)2(FE, MN)1(VA, ZN)0.035
AL5FE4	gamma-brass (Cu5Zn8, D82)	D82	cI52	(217, I-43m)		1	(AL, FE)1
AL5FE2	Al2.8Fe		oS24	(63, Cmcm)		3	(AL)5(FE, MN)2(VA, ZN)3
AL13FE4	Al13Fe4		mS102	(12, C2/m)		3	(AL)0.6275(FE, MN, ZN)0.235(AL, VA, ZN)0.1375
AL7CR	Al45V7		mS104	(12, C2/m)		3	(AL)6(CR)1(AL, ZN)1
AL2CR3	MoSi2 (C11b)	C11b	tI6	(139, I4/mmm)		3	(AL)0.4(CR)0.5(CR, ZN)0.1

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
ALCRZN_TAU4	Unknown Structure			(166, R-3m)		3	(AL, ZN)0.48(CR)0.1207(ZN)0.3993
ALTI3_D019	Ni3Sn (D019)	D019	hP8	(194, P6_3/mmc)	Ti3Al	3	(AL, Ti)3(AL, Ti)1(VA)2
ALTI_L10	CuAu (L10)	L10	tP2	(123, P4/mmm)	TiAl	3	(AL, Ti)1(AL, Ti)1(VA)2
AL3Ti_D022	Al3Ti (D022)	D022	tl8	(139, I4/mmm)	also Al3V	2	(AL, Ti)3(AL, Ti)1
AL5Ti2	Al5Ti2		tP28	(123, P4/mmm)		2	(AL, Ti)5(AL, Ti)2
ALN_B4	Wurtzite (ZnS, B4)	B4	hP4	(186, P6_3mc)		2	(AL)1(N)1
BN_BK	BN (Bk)	Bk	hP4	(194, P6_3/mmc)		2	(B)1(N)1
BN_B3	Zincblende (ZnS, B3)	B3	cF8	(216, F-43m)		2	(B)1(N)1
Si3N4	Nierite (alpha-Si3N4)		hP28	(159, P31c)		2	(Si)3(N)4
Ti2N_C4	Rutile (TiO2, C4)	C4	tP6	(136, P4_2/mnm)		2	(Ti)2(C, N)1
MN3N2	Mn3N2		tl10	(139, I4/mmm)		2	(MN)3(N)2
MN6N5	CoO		tl4	(139, I4/mmm)		2	(MN)6(N)5
TAN_EPS	TaN-eps		hP6	(189, P-62m)		2	(TA)1(N)1
MB_B27	FeB (B27)	B27	oP8	(62, Pnma)		2	(B)1(CO, CR, FE, MN, MO, NI, TI, V, ZR)1
MB_B33	CrB (B33)	B33	oS8	(63, Cmcm)		2	(CR, FE, MN, MO, NB, NI, TA, TI, V)1(B)1
MB_BG	MoB (Bg)	Bg	tl16	(141, I4_-)		2	(CR, FE, MO)0.5(B)0.5

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
				1/amd)			
M2B_C16	Khatyrkite (Al2Cu, C16)	C16	tI12	(140, I4/mcm)		2	(CO, CR, FE, MN, MO, NI, V, W)2(B)1
M2B_CB	Mg2Cu (Cb)	Cb	oF48	(70, Fddd)		2	(CR, FE, MN, MO, NI)0.667(B)0.333
M3B2_D5A	Si2U3 (D5a)	D5a	tP10	(127, P4/mbm)		3	(CR, FE, MO, NI, W)0.4(CR, FE, NI)0.2(B)0.4
NB3B2_D5A	Si2U3 (D5a)	D5a	tP10	(127, P4/mbm)		2	(FE, NB, V)0.6(B)0.4
MB2_C32	Hexagonal omega (C32)	C32	hP3	(191, P6/mmm)		2	(B)2(AL, CR, MG, TI, Y, ZR)1
MOB2_C32	Hexagonal omega (C32)	C32	hP3	(191, P6/mmm)		2	(MO)0.38(B)0.62
B3SI_D1G	B13C2 B4C (D1g)	D1g	hR15	(166, R-3m)		3	(B)6(SI)2(B, SI)6
M2B3	V2B3		oS20	(63, Cmcm)		2	(NB, V)0.4(B)0.6
CR5B3_D8L	Cr5B3 (D8l)	D8l	tI32	(140, I4/mcm)		2	(CR, MO)0.625(B)0.375
YB4_D1E	ThB4 (D1e)	D1e	tP20	(127, P4/mbm)		2	(CE, Y)1(B)4
M3B4_D7B	Ta3B4 (D7b)	D7b	oI14	(71, Immm)		2	(CR, FE, NB, V)0.429(B)0.571
MO2B5_D8I	Mo2B5 (D8i)	D8i	hR7	(166, R-3m)		2	(MO)0.32(B)0.68
M1B6_D21	CaB6 (D21)	D21	cP7	(221, Pm-3m)		2	(CA, CE, Y)1(B)6
M5B6	V5B6		oS22	(65, Cmmm)		2	(NB, V)0.455(B)0.545
YB12_D2F	UB12 (D2f)	D2f	cF52	(225, Fm-3m)		2	(B)12(Y)1
YB66	YB66		cF1936	(226, Fm-3c)		2	(Y)1(B)66

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
FE1NB1B1_C22	Revised Fe2P (C22)	C22(II)	hP9	(189, P-62m)		3	(FE)0.333(NB)0.333(B)0.333
FE3NB3B4	Unknown Structure					3	(FE)0.3(NB)0.3(B)0.4
FEWB_C37	MnCuP		oP12	(62, Pnma)		3	(FE)1(W)1(B)1
FE5SiB2	Mo5SiB2		tI32	(140, I4/mcm)		3	(FE)5(SI)1(B)2
FE10Si2B3	Unknown Structure					3	(FE)2(SI)0.4(B)0.6
FE5Si2B	Nb5Sn2Si		tI32	(140, I4/mcm)		3	(FE)4.7(SI)2(B)1
MNP_B31	MnP (B31)	B31	oP8	(62, Pnma)		2	(CO, CR, FE, MN, NI, W)1(P, SI)1
CU3P_D021	Cu3P (D021)	D021	hP24	(165, P-3c1)		2	(CU, FE)3(P)1
M2P_C22	Revised Fe2P (C22)	C22(II)	hP9	(189, P-62m)		2	(AL, CO, CR, FE, MN, MO, NB, NI, TI, V, W)2(P, SI)1
M3P_D0E	Ni3P (D0e)	D0e	tI32	(82, I-4)		2	(AL, CO, CR, CU, FE, MN, MO, NI, TI)3(B, P)1
MO3P	alpha-V3S		tI32	(121, I-42m)		2	(MO)3(P)1
MOP_BH	Tungsten Carbide (Bh)	Bh	hP2	(187, P-6m2)		2	(MO)1(P)1
TI3P	Ti3P		tP32	(86, P4_2/n)		2	(NB, TI)3(P, SI)1
FEMP_C37	Co2Si (C37)	C37	oP12	(62, Pnma)		3	(CO, CR, FE, NB, NI, TI, V, W)1(CO, CR, FE, NB, NI, TI, V, W)1(P)1
FENB2P	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)		3	(FE)1(NB)2(P)1
FENB4P	Nb4CoSi		tP12	(124, P4/mcc)		3	(FE)1(NB)4(P, SI)1
NBP	NbAs		tI8	(109, I4_1md)		2	(NB)1(P)1
NB7P4	Nb7P4		mS44	(12, C2/m)		2	(NB)7(P)4

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
FESI4P4	FeSi4P4		aP9	(1, P1)		3	(FE)1(SI)4(P)4
ZINCBLENDE_B3	Zincblende (ZnS, B3)	B3	cF8	(216, F-43m)		2	(AL)1(P)1
SIP1	(SiP)		oS48	(36, Cmc2_1)		2	(P)1(SI)1
SIP2	Pyrite (FeS2, C2)	C2	cP12	(205, Pa-3)		2	(P)2(SI)1
WP2	MoP2		oS12	(36, Cmc2_1)		2	(W)1(P)2
CORUNDUM	Corundum (Al2O3, D51)	D51	hR10	(167, R-3c)		3	(AL+3, CR+2, CR+3, FE+2, FE+3, MN+3, SN+4, TI+3)2(CR+3, FE+3, NI+2, VA)1(O-2)3
HALITE	Rock Salt (NaCl, B1)	B1	cF8	(225, Fm-3m)		2	(AL+3, CA+2, CR+3, FE+2, FE+3, MG+2, MN+2, MN+3, NI+2, NI+3, SI+4, VA)1(O-2)1
SPINEL	Spinel (Al2MgO4, H11)	H11	cF56	(227, Fd-3m)		4	(AL+3, CR+2, CR+3, FE+2, FE+3, MG+2, MN+2, NI+2)1(AL+3, CR+3, FE+2, FE+3, MG+2, MN+2, MN+3, MN+4, NI+2, VA)2(CR+2, FE+2, MG+2, MN+2, VA)2(O-2)4
ALPHA_SPINEL	Haussmannite (Mn3O4)		tI28	(141, I4_1/amd)		4	(MG+2, MN+2, MN+3, NI+2)1(AL+3, CR+3, FE+3, MN+2, MN+3, VA)2(MN+2, VA)2(O-2)4
NIMNO3	Ilmenite (FeTiO3, E22)	E22	hR10	(148, R-3)		2	(MN+3, MN+4, NI+2)2(O-2)3
NI6MNO8	Rock Salt (NaCl, B1)	B1	cF8	(225, Fm-3m)		3	(MG+2, NI+2)6(MN+4)1(O-2)8
QUARTZ	alpha-Quartz (low Quartz)		hP9	(152, P3_121)		1	(SiO2)1
CRISTOBALITE	Ideal beta-Cristobalite (SiO2, C9)	C9	cF24	(227, Fd-3m)		1	(SiO2)1
TRIDYMITE	Monoclinic (Cc) Low Tridymite (SiO2)		mS144	(9, Cc)		1	(SiO2)1
RHODONITE	Rhodonite (MnSiO3-b)		aP50	(2, P-1)		3	(MN+2)1(SI+4)1(O-2)3
OLIVINE	Forsterite (Mg2SiO4, S12)	S12	oP28	(62, Pnma)		4	(CA+2, FE+2, MG+2, MN+2, NI+2)1(CA+2, FE+2, MG+2, MN+2, NI+2)1(SI+4)1(O-2)4

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
CAMNO3	CaTiO ₃ Pnma Perovskite		oP20	(62, Pnma)		3	(CA+2)1(MN+4)1(O-2)3
CAMN2O4	CaMn ₂ O ₄		oP28	(57, Pbcm)		3	(CA+2)1(MN+3)2(O-2)4
CA1CR2O ₄ _A	SrCr ₂ O ₄		oP28	(59, Pmmn)		1	(CA1CR2O ₄)1
C1A1	Al ₂ CaO ₄		mP84	(14, P2_1/c)		3	(CA+2)1(AL+3)2(O-2)4
AF	FeGaO ₃		oP40	(33, Pna2_1)		2	(AL ₂ O ₃)1(FE ₂ O ₃)1
CAV2O4	CaV ₂ O ₄		oP28	(62, Pnma)		3	(CA+2)1(CR+3, FE+3)2(O-2)4
C1A2	Al ₄ CaO ₇		mS48	(15, C2/c)		3	(CA+2)1(AL+3)4(O-2)7
CF2	Ca _{3.5} Fe ₁₄ O _{24.5}		mS172	(5, C2)		3	(CA+2)1(FE+3)4(O-2)7
C1A6	Magnetoplumbite (PbFe ₁₂ O ₁₉)		hP64	(194, P6_3/mmc)		3	(CA+2)1(AL+3)12(O-2)19
C2F	Ca ₂ Fe ₂ O ₅		oP36	(62, Pnma)		3	(CA+2)2(FE+3)2(O-2)5
C3A1	Ca ₃ Al ₂ O ₆		cP264	(205, Pa-3)		3	(CA+2)3(AL+3)2(O-2)6
C12A7	Mayenite (12CaO ₇ Al ₂ O ₃ , K74, C12A7)	K74	cl152	(220, I-43d)		4	(CA+2)6(AL+3)6(AL+3, FE+3)1(O-2)16.5
CWF	CaFe ₃ O ₅		oS36	(63, Cmcm)		4	(CA+2)1(FE+2)1(FE+3)2(O-2)5
CW3F	CaFe ₅ O ₇		oS52	(63, Cmcm)		4	(CA+2)1(FE+2)3(FE+3)2(O-2)7
C4WF4	Ca ₄ Fe ₉ O ₁₇		mS60	(5, C2)		4	(CA+2)4(FE+2)1(FE+3)8(O-2)17
C4WF8	Sr ₂ Fe ₂ O ₅		oI44	(74, Imma)		4	(CA+2)4(FE+2)1(FE+3)16(O-2)29
C3A2M1	3CaO ₂ Al ₂ O ₃ .MgO		oP72	(57, Pbcm)		3	(CAO)3(AL ₂ O ₃)2(MGO)1
C1A8M2	CaMg ₂ Al ₁₆ O ₂₇		hP94	(187, P-6m2)		3	(CAO)1(AL ₂ O ₃)8(MGO)2

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C2A14M2	Unknown Structure					3	(CAO)2(AL2O3)14(MGO)2
CORDIERITE	Na0.04 (Mg0.5Fe0.5)2Al4Si5O18		oS120	(66, Cccm)		1	(AL4MG2O18Si5)1
SAPPHIRINE	CaMg3Si3O10		aP68	(2, P-1)		1	(AL18MG7O40Si3)1
CA2SiO4_ALPHA	Ca2SiO4		hP24	(194, P6-3/mmc)		3	(CA+2, MG+2)2(Si+4)1(O-2)4
CA2SiO4_AP	K2CoCl4		oP84	(33, Pna2_1)		3	(CA+2, FE+2, MG+2)2(Si+4)1(O-2)4
LARNITE	Parawollastonite (CaSiO3, S33(II))	S33(II)	mP60	(14, P2_1/c)		3	(CA+2)2(Si+4)1(O-2)4
LOWCLINO_PYROXENE	Diopside [CaMg(SiO3)2, S41]	S41	mS40	(15, C2/c)		4	(CA+2, MG+2)1(MG+2)1(Si+4)2(O-2)6
CLINO_PYROXENE	Diopside [CaMg(SiO3)2, S41]	S41	mS40	(15, C2/c)		4	(CA+2, FE+2, MG+2)1(FE+2, MG+2)1(Si+4)2(O-2)6
ORTHO_PYROXENE	Enstatite (MgSiO3, S43)	S43	oP80	(61, Pbca)		4	(CA+2, MG+2)1(MG+2)1(Si+4)2(O-2)6
PROTO_PYROXENE	MgSiO3		oP40	(60, Pbcn)		3	(CA+2, MG+2)1(Si+4)1(O-2)3
WOLLASTONITE	Wollastonite (CaSiO3)		aP30	(2, P-1)		3	(CA+2, FE+2, MG+2)1(Si+4)1(O-2)3
PSEUDO_WOLLASTONITE	CaSiO3		mS120	(15, C2/c)		3	(CA+2)1(Si+4)1(O-2)3
ANDALUSITE	Andalusite (Al2SiO5, S02)	S02	oP32	(58, Pnnm)		4	(AL+3)1(AL+3)1(Si+4)1(O-2)5
SILLIMANITE	Sillimanite (Al2SiO5, S03)	S03	oP32	(62, Pnma)		4	(AL+3)1(AL+3)1(Si+4)1(O-2)5
MULLITE	Al(Al0.7Si0.3)2O4.8		oP24	(55, Pbam)		4	(AL+3)1(AL+3)1(AL+3, Si+4)1(O-2, VA)5
KYANITE	Kyanite (Al2SiO5, S01)	S01	aP32	(2, P-1)		4	(AL+3)1(AL+3)1(Si+4)1(O-2)5
HATRURITE	Ca3(SiO4)O-b		hR81	(160, R3m)		3	(CA+2)3(Si+4)1(O-2)5

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MELILITE	Akermanite (Ca ₂ MgSi ₂ O ₇ , S53)	S53	tP24	(113, P-42_1m)		5	(CA+2)2(AL+3, MG+2)1(AL+3, SI+4)1(SI+4)1(O-2)7
RANKINITE	3CaO·2SiO ₂		mP48	(14, P2_1/c)		3	(CA+2)3(SI+4)2(O-2)7
ANORTHITE	Ca(Al _{0.5} Si _{0.5}) ₄ O ₈		aP104	(2, P-1)		4	(CA+2)1(AL+3)2(SI+4)2(O-2)8
MERWINITE	Ca ₃ Mg(SiO ₄) ₂		mP56	(14, P2_1/c)		4	(CA+2)3(MG+2)1(SI+4)2(O-2)8
TIO_B1	Rock Salt (NaCl, B1)	B1	cF8	(225, Fm-3m)		3	(TI+2, TI+3, VA)1(TI, VA)1(O-2)1
TIO_ALPHA	alpha-TiO		mS20	(12, C2/m)		2	(TI+2)1(O-2)1
TI3O2	(Ti ₃ O ₂)		hP5	(191, P6/mmm)		3	(TI+2)2(TI)1(O-2)2
RUTILE_C4	Rutile (TiO ₂ , C4)	C4	tP6	(136, P4_2/mnm)		2	(MN+4, NB+4, SN+4, TI+4)1(O-2)2
NBO	NbO		cP6	(221, Pm-3m)		2	(NB+2)1(O-2)1
FE4NB2O9	Nb ₂ Mn ₄ O ₉		hP30	(165, P-3c1)		4	(FE+3)4(NB+2)1(NB+4)1(O-2)9
AL2TiO5	Pseudobrookite (Fe ₂ TiO ₅ , E41)	E41	oS32	(63, Cmcm)		3	(AL+3)2(TI+4)1(O-2)5
FLUORITE_C1	Fluorite (CaF ₂ , C1)	C1	cF12	(225, Fm-3m)		2	(CE+3, CE+4, MN+2, MN+3, NI+2, Y+3, ZR, ZR+4)2(O-2, VA)4
ZRO2_TETR	HgI ₂ (C13)	C13	tP6	(137, P4_2/nmc)		2	(MN+2, MN+3, NI+2, Y+3, ZR+4)2(O-2, VA)4
ZRO2_C43	Baddeleyite (ZrO ₂ , C43)	C43	mP12	(14, P2_1/c)		2	(Y+3, ZR+4)2(O-2, VA)4
S2ZR1	CdI ₂		hP3	(164, P-3m1)		2	(TA, ZR)1(S)2
M2O3C_D53	Bixbyite (Mn ₂ O ₃ , D53)	D53	cl80	(206, Ia-3)		3	(AL+3, CE+3, CE+4, CR+3, FE+3, MN+3, NI+2, Y, Y+3, ZR+4)2(O-2, VA)3(O-2, VA)1
M2O3H_D52	La ₂ O ₃ (D52)	D52	hP5	(164, P-3m1)		3	(MN+3, Y, Y+3, ZR+4)2(O-2, VA)3(O-2, VA)1

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
YAG	Garnet (Co3Al2Si3O12, S14)	S14	cl160	(230, Ia-3d)		3	(AL+3, CR+3, FE+3)5(Y+3)3(O-2)12
YAP	CaTiO3 Pnma Perovskite		oP20	(62, Pnma)		3	(AL+3, CR+3, FE+3)1(Y+3)1(O-2)3
YAM	Y4Al2O9		mP60	(14, P2_1/c)		4	(AL+3, SI+4)2(Y+3)4(O-2, VA)1(O-2)9
Y2S2D_Y2Si2O7	Possible delta-Y2Si2O7		oP44	(62, Pnma)		3	(Y+3)1(Y+3)1(Si2O7-6)1
Y2S2G_Y2Si2O7	Y2Si2O7-b		mP22	(14, P2_1/c)		3	(Y+3)1(Y+3)1(Si2O7-6)1
Y2S2B_Y2Si2O7	La4Ge3[GeO4]O10		aP44	(2, P-1)		3	(Y+3)1(Y+3)1(Si2O7-6)1
Y2S2A_Y2Si2O7	Thortveitite ([Sc,Y]2Si2O7, S21)	S21	mS22	(12, C2/m)		3	(Y+3)1(Y+3)1(Si2O7-6)1
Y2SiO5	Y2SiO5 (RE2SiO5 X2)		mS64	(15, C2/c)		4	(Y+3)1(Y+3)1(SiO4-4)1(O-2)1
ZR3Y4O12	UY6O12		hR19	(148, R-3)		3	(ZR+4)3(Y+3)4(O-2)12
MN2YO5	HoMn2O5		oP32	(55, Pbam)		4	(Y+3)1(MN+3)1(MN+4)1(O-2)5
MNYO3_HEX	LuMnO3		hP30	(185, P6_3cm)		3	(Y+3)1(MN+3)1(O-2)3
CUPRITE_C3	Cuprite (Cu2O, C3)	C3	cP6	(224, Pn-3m)		2	(CU+1)2(O-2)1
CUO_B26	Tenorite (CuO, B26)	B26	mS8	(15, C2/c)		2	(CU+2)1(O-2)1
YCUO2	Hexagonal Delafossite (CuAlO2)		hP8	(194, P6_3/mmc)		3	(Y+3)1(CU+1)1(O-2)2
Y2CU2O5	Cu2Ho2O5		oP36	(33, Pna2_1)		3	(Y+3)2(CU+2)2(O-2)5
YFE2O4	MnBi2Te4		hR7	(166, R-3m)		3	(FE+2, FE+3)2(Y+3)1(O-2)4
MS_B81	NiAs (B81)	B81	hP4	(194, P6_3/mmc)		2	(AL, CO, CR, CU, FE, MN, NB, NI, TI, V, VA, ZR)1(S)1

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MS2_C2	Pyrite (FeS2, C2)	C2	cP12	(205, Pa-3)		2	(CO, FE, MN, NI, RU, TI)1(S)2
CR1S1	CrS		mS8	(15, C2/c)	This is low-temp CrS.	2	(CR)1.03(S)1
CR2S3	Dolomite [MgCa(CO3)2, G11]	G11	hR10	(148, R-3)		2	(CR, FE)2(S)3
CR3S4	Brezinaite (Cr3S4)		mS14	(12, C2/m)		2	(CR, FE, NI)3(S)4
CR5S6	Cr5S6		hP22	(163, P-31c)		2	(CR)5(S)6
CR7S8	Cr7Se8		mS30	(12, C2/m)		2	(CR)7(S)8
CU2S_GAMMA	Cu2Se		cF44	(225, Fm-3m)		3	(CU, FE, VA)2(CU, VA)1(S)1
HEAZLEWOODITE_B1	Cu1.9S		cF12	(216, F-43m)	This is non-stoichiometric high temperature Ni3S2.	2	(FE, NI, VA)2(S)1
HEAZLEWOODITE_B2	Unknown Structure				This is non-stoichiometric high temperature Ni4S3.	2	(FE, NI, VA)2(S)1
MS_B1	Rock Salt (NaCl, B1)	B1	cF8	(225, Fm-3m)		2	(CA, CE, CO, CR, CU, FE, MG, MN, Y, ZR)1(S)1
NIS_LT	Millerite (NiS, B13)	B13	hR6	(160, R3m)	This is low temperature NiS.	2	(NI)1(S)1
NI3S2_LT	Hazelwoodite (Ni3S2, D5e)	D5e	hR5	(155, R32)		2	(NI)3(S)2
NI7S6	Unknown Structure		t**			2	(FE, NI)7(S)6
NI9S8	Ni9S8		oS68	(21, C222)		2	(FE, NI)9(S)8
FEMO3S4	FeMo3S4			(1, P1)		3	(FE)1(MO)3(S)4
TA3FES6	Ta3FeS6			(182, P6_3/mmc)		3	(FE)1(TA)3(S)6
TI4C2S2	AlCr2		hP8	(194, P6_3/mmc)		3	(TI)4(C)2(S)2

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
FESO4	Chalcocyanite (CuSO4)		oP24	(62, Pnma)		3	(FE)1(S)1(O)4
FE2S3O12	Unknown Structure					3	(FE)2(S)3(O)12
SIS2_C42	SiS2 (C42)	C42	oI12	(72, Ibam)		1	(SiS2)1
AL2S3	alpha-Al2S3		hP30	(169, P6_1)		1	(Al2S3)1
CUS_B18	Covellite (CuS, B18)	B18	hP12	(194, P6_3/mmc)		2	(CU)1(S)1
CU2S_ALPHA	Cu2S-alpha		mP144	(14, P2_1/c)		2	(CU)2(S)1
CU2S_BETA	Cu2S-beta		hP16	(194, P6_3/mmc)		2	(CU)2(S)1
CU31S16	Cu31S16		mP376	(14, P2_1/c)		2	(CU)1.93(S)1
CU7S4	Cu7S4		oP44	(62, Pnma)		2	(CU)1.75(S)1
CO9S8	Co9S8 (D89)	D89	cF68	(225, Fm-3m)		2	(CO, FE, NI)9(S)8
P2S5	P2S5		aP28	(2, P-1)		2	(P)2(S)5
Ti2S	Ta2P		oP36	(58, Pnnm)		2	(TI)2(S)1
Ti8S3	Ti8S3		mS88	(12, C2/m)		2	(TI)8(S)3
TiS3	ZrSe3		mP8	(11, P2_1/m)		2	(TI)1(S)3
Ti8S10	Ti0.81S		hP20	(194, P6_3/mmc)		2	(TI)8(S)10
Ti8S9	TiS		hR18	(166, R-3m)		2	(TI)8(S)9
PENTLANDITE	Co9S8 (D89)	D89	cF68	(225, Fm-3m)		3	(FE, NI)8(FE, NI)1(S)8
THIOSPINEL	Spinel (Co3O4, D72)	D72	cF56	(227, Fd-3m)	This is a sulphur	3	(FE, NI)1(CR, NI)2(S)4

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
					spinel: (Cu,Fe,Mn)Cr ₂ S ₄ , Co ₃ S ₄ , FeNi ₂ S ₄ and Ni ₃ S ₄ .		
CE7Ni3_D102	Fe3Th7 (D102)	D102	hP20	(186, P6_3mc)		2	(CE)0.7(NI)0.3
CEZN3	CeZn3		oS16	(63, Cmcm)		2	(CE)0.25(ZN)0.75
CE13ZN58	Gd13Zn58		hP142	(194, P6_3/mmc)		2	(CE)0.183(ZN)0.817
CE3ZN22	Ce3Zn22		tI100	(141, I4_1/amd)		2	(CE)0.12(ZN)0.88
CENI_B33	CrB (B33)	B33	oS8	(63, Cmcm)		2	(CE)0.5(NI)0.5
CENi3	CeNi3		hP24	(194, P6_3/mmc)		2	(CE)0.25(NI)0.75
CE2Ni7	Ce2Ni7		hP36	(194, P6_3/mmc)		2	(CE)0.22222(NI)0.77778
MG41M5	Ce5Mg41		tI92	(87, I4/m)		2	(MG)41(CE)5
MG3M_D03	BiF3 (D03)	D03	cF16	(225, Fm-3m)		2	(MG)3(CE, MG)1
MG17R2	CeMg10		hP44	(194, P6_3/mmc)		2	(MG)17(CE)2
MG12R_D2B	Mn12Th (D2b)	D2b	tI26	(139, I4/mmm)		2	(MG)12(CE)1
CECO5_D2D	CaCu5 (D2d)	D2d	hP6	(191, P6/mmm)		2	(CE)1(CO, CU, NI, ZN)5
CE5CO19	Ce5Co19		hR24	(166, R-3m)		2	(CO)19(CE)5
CE2CO7	Co7Gd2		hR18	(166, R-3m)		2	(CO)7(CE)2

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
CECO3	Ni3Pu		hR12	(166, R-3m)		2	(CO)3(CE)1
CE24CO11	Ce24Co11		hP70	(186, P6_3mc)		2	(CO)11(CE)24
AL4CE_D13	Al4Ba (D13)	D13	tI10	(139, I4/mmm)		2	(AL)0.8(CE)0.2
AL11CE3	Ni3Sn (D019)	D019	hP8	(194, P6_3/mmc)		2	(AL)0.7857(CE)0.2143
AL3CE_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)		2	(AL)0.75(CE)0.25
AL3CE_D019	Ni3Sn (D019)	D019	hP8	(194, P6_3/mmc)		2	(AL)0.75(CE, Y)0.25
ALCE1	AlCe		oS16	(63, Cmcm)		2	(AL)0.5(CE)0.5
ALCE3_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)		2	(AL)0.25(CE)0.75
ALCE3_D019	Ni3Sn (D019)	D019	hP8	(194, P6_3/mmc)		2	(AL)0.25(CE)0.75
CE2FE17	Ni17Th2		hP38	(194, P6_3/mmc)		2	(CE)2(FE, ZN)17
A_R2O3_D52	La2O3 (D52)	D52	hP5	(164, P-3m1)		3	(CE+2, CE+3)2(O-2)2(O-2, VA)1
H_R2O3	H-La2O3		hP10	(194, P6_3/mmc)		3	(CE+2, CE+3)2(O-2)2(O-2, VA)1
X_R2O3	Nd2O3		cl26	(229, Im-3m)		2	(CE+2, CE+3)2(O-2, VA)3
CE7O12	Pr7O12		hR19	(148, R-3)		2	(CE)7(O)12
CE1S2	Ce10Se19		tP58	(86, P4_2/n)		2	(CE)1(S)2
CE2S3	Th3P4 (D73)	D73	cl28	(220, I-43d)		2	(CE)2(S)3
CE3S4_D73	Th3P4 (D73)	D73	cl28	(220, I-43d)		2	(CE)3(S)4

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
CE2O12S3	Unknown Structure					3	(CE)2(O)12(S)3
CE2O2S1	Ce2O2S		hP5	(164, P-3m1)		3	(CE)2(O)2(S)1
CEC2_BETA	CaC2		cF36	(225, Fm-3m)		2	(CA, CE)1(C)2
CEC2_C11A	CaC2-I (C11a)	C11a	tI6	(139, I4/mmm)		2	(CA, CE, Y)1(C)2
CECU6	CeCu6		oP28	(62, Pnma)		2	(CU)0.857(CE)0.143
CECU4	CeCu4		oP20	(58, Pnnm)		2	(CU)0.8(CE)0.2
CECU_B27	FeB (B27)	B27	oP8	(62, Pnma)		2	(CU)0.5(CE)0.5
CE2C3_D5C	Pu2C3 (D5c)	D5c	cl40	(220, I-43d)		2	(CE)0.4(C)0.6
MGB4	MgB4		oP20	(62, Pnma)		2	(MG)1(B)4
MGB7	MgB7		ol64	(74, Imma)		2	(MG)1(B)7
MG2C3	Mg2C3		oP10	(58, Pnnm)		2	(MG)2(C)3
CANI3	Ni3Pu		hR12	(166, R-3m)		2	(CA)0.25(NI)0.75
CANI5_D2D	CaCu5 (D2d)	D2d	hP6	(191, P6/mmm)		2	(CA)1(NI, ZN)5
COP3	Skutterudite (CoAs3, D02)	D02	cl32	(204, Im-3)		2	(CO)1(P)3
CR3RU_A15	Cr3Si (A15)	A15	cP8	(223, Pm-3n)		2	(CR)0.685(RU)0.315
MG3N2_D53	Bixbyite (Mn2O3, D53)	D53	cl80	(206, Ia-3)		2	(CA, MG)3(N)2
MO2S3	Mo2S3		mP10	(11, P2_1/m)		2	(MO)2(S)3
MOS2_C7	Molybdenite (MoS2, C7)	C7	hP6	(194, P6_3/mmc)		2	(MO, W)1(S)2

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
NB3RU5	Rh5Ge3		oP16	(55, Pbam)		2	(NB, RU)0.375(RU)0.625
RU4Si3	Ru4Si3		oP28	(62, Pnma)		2	(RU)4(SI)3
RU2Si3	Ge3Ru2		oP40	(60, Pbcn)		2	(RU)2(SI)3
FEAL2S4	ZnIn2S4		hR7	(160, R3m)		3	(FE)1(AL)2(S)4
M4Si1_G3	AlAu4		cP20	(198, P2_13)		4	(CR, FE, NI)3(NI)1(SI)1(C, VA)1
NI31Si12	Ni31S12		hP42	(150, P321)		2	(FE, NI)5(SI)2
CO2Si_C37	Co2Si (C37)	C37	oP12	(62, Pnma)		2	(CO, FE, NI, RU)2(SI)1
M2Si_TETA	AlNi2		hP6	(164, P-3m1)		3	(FE, NI)1(FE, NI, VA)1(SI)1
NI3Si2	Ni3Si2		oP80	(36, Cmc2_1)		2	(FE, NI)3(SI)2
CR5Si3_D8M	W5Si3 (D8m)	D8m	tl32	(140, I4/mcm)		3	(CR, FE)4(CR, FE)1(SI)3
AL6FE6ZR_D2B	Mn12Th (D2b)	D2b	tl26	(139, I4/mmm)		2	(AL, FE)12(ZR)1
FE3SiZR2	MgZn2 Hexagonal Laves (C14)	C14	hP12	(194, P6-3/mmc)		3	(FE)3(SI)1(ZR)2
FE2Si2ZR_D13	Al4Ba (D13)	D13	tl10	(139, I4/mmm)		3	(FE)2(SI)2(ZR)1
FE5Si12ZR3	Unknown Structure					3	(FE)5(SI)12(ZR)3
FE2Si3ZR3	Hf3Ni2Si3		oS32	(63, Cmcm)		3	(FE)2(SI)3(ZR)3
FE4Si2ZR1	ZrFe4Si2		tP14	(136, P4-2/mnm)		3	(FE)4(SI)2(ZR)1
FE29Si42ZR29	Unknown Structure					3	(FE)29(SI)42(ZR)29

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
NBFEZN12	Unknown Structure					3	(NB)0.071(FE)0.071(ZN)0.857
NB1ZN1	Unknown Structure					2	(NB)1(ZN)1
NB2ZN3_D85	Fe7W6 (D85) mu-phase	D85	hR13	(166, R-3m)		2	(NB)2(ZN)3
NBZN7	Unknown Structure					2	(NB)1(ZN)7
ALH3	AlH3		oP24	(58, Pnnm)		2	(AL)1(H)3
DELTA_TIH2	Fluorite (CaF2, C1)	C1	cF12	(225, Fm-3m)		2	(NB, Ti, V, Y, ZR)1(H, VA)2
NBH_BETA	Ta2H		oS8	(21, C222)		2	(NB)1(H, VA)1.1
TA2H_EPSILON	Unknown Structure					2	(TA)2(H, VA)1
TI2NIH	Ti2NiH		cF128	(227, Fd-3m)		3	(Ti)2(NI)1(H)1
V3H2	Unknown Structure					2	(V)0.6(H)0.4
YH3_EPSILON	H3Ho		hP24	(165, P-3c1)		2	(Y)1(H, VA)3
DIS_BETA	V2H1.1		tI56	(141, I4_1/amd)		2	(H, VA)0.5(V)0.5
BETA_PHASE	V2H		mS6	(12, C2/m)		3	(H, VA)0.25(H, VA)0.25(V)0.5
CAH2_C37	Co2Si (C37)	C37	oP12	(62, Pnma)		2	(CA)1(H)2
CAH2_BETA	Unknown Structure					2	(CA)1(H)2
MGH2_C4	Rutile (TiO2, C4)	C4	tP6	(136, P4_2/mnm)		2	(MG)1(H)2
CEH3_EPSILON	CeH3		cF44	(225, Fm-3m)		2	(CE)1(H, VA)3
FE5SN3_B82	Ni2In (B82)	B82	hP6	(194, P6_3/mmc)		2	(CO, CR, FE, MN)5(SN)3

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
FE3Sn2	Fe3Sn2		hR10	(166, R-3m)		2	(CR, FE)3(SN)2
FESN_B35	CoSn (B35)	B35	hP6	(191, P6/mmm)		2	(CO, CR, FE, MN, ZN)1(SN)1
FESN2_C16	Khatyrkite (Al2Cu, C16)	C16	tI12	(140, I4/mcm)		2	(CO, FE, MN)1(SN)2
NI3Sn2	Ni3Sn2		oP20	(62, Pnma)		2	(CO, FE, MN, NI)3(SN)2
SNS_B16	GeS (B16)	B16	oP8	(62, Pnma)		2	(SN)1(S)1
SN3Y1	GdSn2.75		oS16	(38, Amm2)		2	(SN)3(Y)1
SN5Y2	Shcherbinaita (V2O5) (Revised)		oP14	(59, Pmmn)		2	(SN)5(Y)2
SN2Y_C49	ZrSi2 (C49)	C49	oS12	(63, Cmcm)		2	(SN)2(Y)1
SN10Y11	Ge10Ho11		tI84	(139, I4/mmm)		2	(SN)10(Y)11
SN3Y5_D88	Mavlyanovite (Mn5Si3, D88)	D88	hP16	(193, P6_3/mcm)		3	(Y, ZR)5(SN)3(SN, VA)1
SN2ZR_C54	TiSi2 (C54)	C54	oF24	(70, Fddd)		2	(SN)2(ZR)1
SNZr4_A15	Cr3Si (A15)	A15	cP8	(223, Pm-3n)		2	(SN, ZR)1(SN, ZR)3
CU41Sn11	Cu41Sn11		cF416	(216, F-43m)	delta	2	(CU)0.788(SN)0.212
CU10Sn3	Cu10Sn3		hP26	(173, P6_3)	zeta	2	(CU)0.769(SN)0.231
CU3Sn_L	Cu3Sn		oS80	(63, Cmcm)	epsilon	2	(CU)0.75(SN)0.25
CU6Sn5_L	Cu6Sn5		mS44	(15, C2/c)		2	(CU)0.545(SN)0.455
CU6Sn5_B81	NiAs (B81)	B81	hP4	(194, P6_3/mmc)		2	(CU)0.545(SN)0.455

Name	Prototype	Strukturbericht	Pearson Symbol	Space Group	Info	Sublattices	Formula Unit
NI3SN_D019	Ni3Sn (D019)	D019	hP8	(194, P6 ₃ /mmc)		2	(FE, MN, NI)3(SN)1
NI3SN_D03	BiF3 (D03)	D03	cF16	(225, Fm-3m)		1	(NI, SN)1
NI3SN4	delta-Ni3Sn4 (D7a)	D7a	mS14	(12, C2/m)		2	(NI)3(SN)4
FESNZR_N	Unknown Structure					3	(FE)16(SN)16(ZR)12
FE6SN6ZR	MgFe6Ge6		hP13	(191, P6/mmm)		3	(FE)6(SN)6(ZR)1
FE3SN9ZR10	Unknown Structure					3	(FE)3(SN)9(ZR)20
FESNZR_Y	Unknown Structure					3	(FE)18(SN)9(ZR)5
CA2SN_C37	Co2Si (C37)	C37	oP12	(62, Pnma)		2	(CA)2(SN)1
CA36SN23	Sn23Yb36		tP118	(127, P4/mbm)		2	(CA)36(SN)23
CA31SN20	Pu31Rh20		tl204	(140, I4/mcm)		2	(CA)31(SN)20
CA7SN6	Ca7Sn6		oP52	(62, Pnma)		2	(CA)7(SN)6
CASN3_L12	Bogdanovite (Cu3Au, L12)	L12	cP4	(221, Pm-3m)		2	(CA)1(SN)3

TCFE14 Properties Data

Model Descriptions

For more information about the various thermophysical, thermomechanical, and properties 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 [Steels and Fe-Alloys Databases](#) page on our website where you can access a Validation and Calculation Examples Collection and the Technical Information plus learn more about the compatible kinetic database. Also explore further [applications of Thermo-Calc to iron and steels](#) including links to resources such as publications, webinars, videos, and more.

TCFE: TCS Steels/Fe-Alloys Database Revision History

Current Database Version

Database name (acronym):	TCS Steel and Fe-alloys Database (TCFE)
Database owner:	Thermo-Calc Software AB
Database version:	14
First release:	TCFE1 was released in 1992

Changes in the Most Recent Database Release

TCFE13.1 to TCFE14

Software release version 2025a (January 2025)

ELASTIC PROPERTIES ADDED

- Elastic constants (C11, C12, C13, C33, and C44) are added for FCC_A1, BCC_A2, and HCP_A3 phases. The elastic constants description can be used to derive the elastic moduli (bulk modulus, shear modulus, and Young's modulus) for a single-phase microstructure.
- Unary systems assessed:
 - FCC_A1, BCC_A2, HCP_A3: Al, Ca, Ce, Co, Cr, Cu, Fe, Mg, Mn, Mo, Nb, Ni, P, Ru, S, Si, Sn, Ta, Ti, V, W, Y, Zn, Zr
- Binary systems assessed:
 - Fe-X (X = Al, Co, Cr, Mg, Mn, Mo, Nb, Ni, Si, Ti, V, W)
 - Mo-X (X = Nb, Sn, Ta, Ti, W, Zr)
 - Nb-X (X = Sn, Ta, Ti, V, W, Zr)
 - Sn-X (X = Ta, Ti, Zr)
 - Ta-X (X = Ti, W, Zr)
 - Ti-X (X = Al, Cr, V, W, Y, Zr)
 - W-X (X = Cr, V, Zr)
 - Cr-Ni

NEW SYSTEMS AND PHASES

- 18 new ternary systems (total of 335)
- 4 new quaternary systems (total of 84)
- 30 new phases (total of 465)

IMPROVED SOLUBILITY

- BN in liquid and solid Fe: B-N, Fe-B, Fe-B-N
- Mg in cast irons: C-Fe-Mg, C-Fe-Mg-Si
- Si in cementite phase: C-Fe-Si
- Sn in corundum phase: O-Sn, Fe-O-Sn

IMPROVED SYSTEMS

- S systems
 - S-X (X= Ce, Co, Cr, Fe, Ni, Ru, Ti, V)
 - Fe-S-X (X=Ce, Cr, Mo, Ni, Ta, Ti, V, Zn)
 - Cr-Fe-Ni-S
- Ti systems
 - Ti-X (X=Al, Fe, Si)
 - Fe-Ti-X (X= Al, Cr, Nb, Ni, Si, V)

IMPROVED EQUILIBRIA

Improved $\alpha/\sigma/\gamma$ equilibria for:

- Fe-Ni-V, Cr-Ni-V, Cr-Fe-Ni-V
- Mo-V, Fe-Mo-V

IMPROVED DESCRIPTIONS

- Cubic carbides: C-Nb, C-X-Y (X-Y= Cr-Ti, Cr-V, Fe-Ti, Mo-Nb, Mo-Ti, Mo-V, Nb-Ti, Nb-V, Ti-V)
- TiN inclusions in steel melts: Cr-Fe-Ni-Ti-N
- Ni₃X (X=Mo,Ti) precipitates in maraging steels: Mo-Ni, Mo-Nb-Ni, Mo-Ni-Ti, Nb-Ni-Ti

IMPROVED THERMOPHYSICAL PROPERTIES

- Thermal conductivity for cemented carbide systems improved

Previous Releases

TCFE13.0 to TCFE13.1

Software release 2023b Update 1 (September 2023)

- Thermal conductivity (THCD) and electrical resistivity (ELRS): Updated for C- and N-containing solution phases.
- Ternary system Fe-Nb-P re-assessed.

TCFE12.0 to TCFE13.0

Software release 2023b (June 2023)

- The full gas description is included.
- Assessed Fe-Mn-Ti system.
- Revised Cr-Fe-Ni-Si subsystems (Cr-Fe-Si, Cr-Ni-Si) to increase the accuracy of phase equilibria involving the FCC/BCC/LIQUID phases.
- Reassessed THCD of FeTi-B2 and added THCD ELRS of BN_BK.
- Updated the Diamond molar volume and corrected wrong volume parameters.
- Updated Ca-Fe to a more recent description.
- Removed the HIGH_SIGMA phase from the Cr-Mn binary.
- Revised S-Zr and Fe-S-Zr.
- Estimated Fe-Nb-S.
- Revised the liquid in Fe-O-Y and C-Ni-Si.
- The surface tension was re-assessed based on the Redlich-Kister-Muggianu (R-K-M) sub-regular solution model.

TCFE11.0 to TCFE12.0

Software release 2022a (December 2021/January 2022)

ADDED TIN (Sn)

- 26 Binaries Sn-X (X=Al, B, C, Ca, Ce, Co, Cr, Cu, Fe, Mg, Mn, Mo, Nb, Ni, O, P, Ru, S, Si, Sn, Ta, Ti, V, W, Y, Zn, Zr)
- 13 Ternaries: Fe-Sn-X (X=C, Cr, Cu, Mn, Nb, Ni, O, S, Si, W, Zn, Zr), and Al-C-Sn

NEW THERMOPHYSICAL PROPERTIES

- Electrical resistivity is modeled for crystalline phases and liquid. The descriptions can be used for deriving electrical conductivity.
- Thermal conductivity is modeled for crystalline phases and liquid. The descriptions can be used for deriving thermal resistivity as well as thermal diffusivity (by combining with our density and heat capacity data).

UPDATED ZINC CORNER

As well as systems related for the galvanization process:

- Remodeled and unified all gamma-brass D82 phases stable in Fe-Zn, Cu-Zn, Ni-Zn, Mn-Zn, Co-Zn with a 4SL FE3ZN7_D82.
- Added Al-Fe-Mg, Cu-Fe-Zn, Fe-Ni-Zn, Fe-Si-Zn, Fe-Mg-Si, Fe-Mg-Ni, Fe-Mn-Zn, Mg-Si-Zn

OTHER UPDATES

- Unified several phases with the same crystallography
- Corrected the low melting pseudobinary in Fe-O-Si
- Extended a ternary phosphide: FEMP_C37 (CO,CR,FE,NB,NI,TI,V,W)1(CO,CR,FE,NB,NI,TI,V,W)1P1
- Revised C-Ca and estimated C-Ca-Fe

TCFE10.1 to TCFE11.0

Software release version: 2021a (December 2020/January 2021)

THERMOPHYSICAL PROPERTIES

- Added surface tension of the metallic liquid.
- Updated molar volume of the liquid.

NEW ELEMENTS AND SYSTEMS

- Added binary M-H systems where M=Al, Ca, Ce, Co, Cr, Cu, Fe, Mg, Mn, Mo, Nb, Ni, Ru, Si, Ta, Ti, V, W, Y, Zn, Zr.
- Liquid, FCC, BCC and HCP solutions and important hydrides are included. In addition, ternary C-Fe-H, Fe-H-Si, Cr-Fe-H, Al-Fe-H, Fe-H-Mo, Fe-H-Ni, Fe-H-V, Fe-H-W, Al-H-Ti, H-Ni-Ti, H-Ti-Zr systems and quaternary C-Fe-Mn-H, Cr-Fe-H-Ni systems were verified against the experimental data.
- Al-Fe-Nb, Fe-Nb-Ni, Co-Fe-Nb, Al-Cr-Nb

- Al-Fe-Zr, Cr-Fe-Zr, Cu-Fe-Zr
- Ca-P, Ca-Ti, Ca-V, Ca-W
- Ce-Nb, Ce-Ta, Ce-Ti, Ce-W
- Ca-N, N-Y, Ce-N
- Nb-Zn, Ta-Zn, W-Zn, Fe-Nb-Zn
- P-V, Fe-P-V
- S-Ta

UPDATED SYSTEMS AND PHASES

- Fe-N and C-Fe-N are updated to a more recent modeling work including the non-stoichiometric composition of Fe_4N nitride.
- Fe-Si and C-Fe-Si are updated to avoid a high temperature miscibility gap in the liquid.
- Cr-Si, Cr-Fe-Si, and C-Cr-Si are updated to a more recent modeling work.
- FeNbP , FeTiP , NbCrP , NbNiP , CrNiP are modeled as FEMP_C23 phase.
- Co-Nb, Cr-Nb, and Cr-Nb-Ni are updated to separate C15 and C14 laves phases.
- Fe-Y is updated to a more recent modeling work.
- Fe-Zr and Fe-Si-Zr are updated to a more recent modeling work.
- Fe-Nb is updated with an updated formation energy of MU_D85 phase.
- Nb-N is updated to a more recent modeling work.
- Ni-S is updated to assess the solubility of S in FCC nickel.

TCFE8.1 to TCFE8.2 and TCFE9.2 to TCFE9.3

A simultaneous release as part of software release version: 2021a (December 2020/January 2021)

- For users who have TCFE8 and TCFE9 and also use the Steel Model Library, there is an update to correct the volume of liquid Zn.

TCFE10.0 to TCFE10.1

Software release version: 2020b (June 2020)

There was a simultaneous update to both TCFE9 and TCFE10 that is bundled with the 2020b software release and needed for the Steel Model Library Martensite Temperature Model.

- Revised parameters of BCC_A2 phase in Fe-Mn-N system to avoid the appearance of a high N composition set at low temperatures.
- Removed a previously added ad-hoc modification to the lattice stability of BCC_A2 Nickel at temperatures below 450K.

TCFE9.1 to TCFE9.2

Software release version: 2020b (June 2020)

There was a simultaneous update to both TCFE9 and TCFE10 that is bundled with the 2020b software release and needed for the Steel Model Library Martensite Temperature Model.

- Removed a previously added ad-hoc modification to the lattice stability of BCC_A2 Nickel at temperatures below 450K.

TCFE9.0 to TCFE10.0

Software release version: 2020a (January 2020)

The TCFE10 database is significantly improved compared to TCFE9. Below is a summary of the major updates. Read more detail on the [TCFE history section](#) on our website.

- 53 new binary and 13 new ternary systems are included.
- More than 15 binary and 14 ternary systems , and many quaternary, are either completely updated or partially modified to improve the database's robustness and predictability.
- Addition of Ru and 19 Ru-X binary systems (X=Al, B, C, Ca, Co, Cr, Cu, Fe, Mg, Mn, Mo, Nb, Ni, S, Si, Ta, Ti, V, Zn). In addition, the Cr-Fe-Ru is assessed at the Fe corner.
- Better predictive capacity for nitrogen alloyed duplex stainless steels: The systems relevant for nitrogen alloyed duplex stainless steels were systematically reviewed and revised as necessary.
 - The database predictivity is improved in these critical areas: Equilibria at liquidus - solidus; equilibria of matrix phases ($\alpha+\gamma$) at high temperature; and equilibria of secondary phases at intermediate temperature.
- TCFE10 contains 31 new phases among which the so called η -silicide is named as ETA_M5SiN; and τ_1 -silicide (Fe_{2.5}Ni_{1.5}Si) and π -silicide (Cr_{1.5}Ni_{2.5}Si) are modeled as M4Si1_G3.
- The viscosity of metallic liquid alloys is modeled in TCFE10.
- Split description of ordered phases: The ordered phases BCC_B2 and FCC_L12 are described based on the order/disorder partitioning model as described by Lukas et al. [2007].

TCFE9.0 to TCFE9.1

Software release version: 2019b (June 2019)

The changes include:

- Revision of C-Fe-S system.
- Revision of Cr-Fe-Nb and Fe-Nb-Si system and the addition of 15 new silicide phases.
- Revision of the Laves phase description in Fe-Nb-W and Cr-Mo-Nb systems.
- Updates to the molar volumes of Liquid Mn, CEMENTITE, Fe-Si-B ternary phases, MNS, and several sulfides.
- Correction of the magnetic properties of CBCC_A12 phase.
- Removing the pressure dependent parameters from Fe for compatibility with GES6.

TCFE8.0 to TCFE9.0

Software release version: 2017a (March 2017)

Below is a summary of the major updates. Read more detail on the [TCFE history section](#) on our website

The database is developed and validated for simulation of the solidification process, the relative stability of matrix phases (austenite and ferrite), precipitation of secondary phases such as sulfides, borides, oxides, phosphides, carbides, nitrides, carbonitrides, and also intermetallic phases such as the sigma and laves phases.

The TCFE9 database is significantly improved compared to the previous TCFE8 version, with 32 new binary and 35 new ternary systems added to the database. In addition, more than 50 binary and 39 ternary, and many quaternary, systems are either completely updated or partially modified to improve the robustness and predictability of the database.

TCFE8.0 to TCFE8.1

Software release version 2016b (November 2016)

- A bug fix that previously resulted in a faulty estimation of molar volume of borides.
- The change of description of Fe-B and Fe-Ti-B systems due to incompatibility

TCFE7.0 to TCFE8.0

Software release version: 2015a (June 2015)

In order to increase the predictive capability of the database, several significant re-assessments have been performed by Thermo-Calc Software AB which are incorporated in the new release.

The elements Zn and Y and relevant phases associated with these elements have been added in the new TCFE8.

The element Zn has been added mainly for the purpose of galvanization process with the focus on the Zn corner of Al-Cr-Fe-Zn system for, but several other binaries and ternaries are also included. The element Y has been added mainly for the purpose of oxide dispersion strengthened (ODS) steels and the Al-Cr-Cu-Fe-Mn-Ni-O-Si-Y-Zr has been included which contains many assessed oxygen containing binary and ternary systems.

The description of Cu containing systems are improved with addition of Co-Cu, Co-Cu-Fe and Cu-Fe-Si data.

The FE2SITI_L21 phase has been added in the database which is important for the Fe-Si-Ti precipitation hardening steels.

The LIQUID, MU_PHASE, LAVES_PHASE_C14, SIGMA and L12_FCC phases have been improved due to the modifications for the LIQUID in the Mn-O-S, Mo-Si, C-Fe-Si and C-Fe-O systems, MU_PHASE in the Co-Nb and Co-Ta systems, LAVES_PHASE_C14 in Co-Mo and Cu-Fe systems, SIGMA phase in the Co-Cr, Al-Nb, Mn-Ta, Mo-V, Nb-V, Ta-Ti and Ta-V systems, and L12_FCC in Co-V system.

The description of C-Co-Cr system has also been added for cemented carbide applications.

All necessary volume data (including molar volume and thermal expansion) has been updated for all phases in this new release. However, the molar volume data incorporated has no pressure dependence.

Some of the major improvements to the TCFE8 database include improved/added thermodynamic descriptions for several binary, ternary and quaternary systems.

TCFE6.0 to TCFE7.0

TCFE7 released in 2012.

In order to increase the predictive capability of the database, several significant re-assessments have been performed by Thermo-Calc Software AB which are incorporated in the new release. The elements Ta and Zr and relevant phases associated with these elements have been added. Three more B containing phases and two more Mn nitride phases have been also implemented into the new TCFE7 database. All oxide phases for the Fe-Al-Ca-Cr-Mg-Mn-Ni-Si-O system have been updated or added, which includes the addition of many new oxide phases. Within the latter system all lower-order sub-systems have been evaluated.

In version 4 (TCFE4), all necessary volume data (including molar volume and thermal expansion) for various alloy phases were incorporated; such volume data has been updated for all phases in this new release. However, the molar volume data incorporated has no pressure dependence.

Some of the major improvements to the TCFE7 database include improved/added thermodynamic descriptions for the following binary, ternary and quaternary systems (with many sub-systems also re-assessed).

TCFE5.0 to TCFE6.0

TCFE6 released in 2008.

In order to increase the predictive capability of the database, several significant re-assessments have been performed by Thermo-Calc Software AB and incorporated in the new release. The element Ca has been added and the alloying ranges for the elements C, Co, Cu, N, Ti and V have been extended. In version 4 (TCFE4), all necessary volume data (including molar volume and thermal expansion) for various alloy phases were incorporated; such volume data has been updated for all phases in this new release. However, the molar volume data incorporated has no pressure dependence.

Some of the major improvements to the TCFE6 database include improved thermodynamic descriptions for the following ternary and quaternary systems (with many sub-systems also re-assessed).

TCFE4.0 to TCFE5.0

TCFE5 released in 2007.

This release builds on to the previous version of the database, i.e. TCFE4, and additionally includes complete reassessments of several binary and ternary systems performed under a three year collaborative program within the framework of the CCT-Applied project for stainless steels. CCT (Centre of Computational Thermodynamics) is a collaborative effort between the Royal Institute of Technology (Stockholm, Sweden), Kimab and Swedish industries.

Some of the major improvements to the TCFE5 database include improved thermodynamic descriptions for the following ternary and quaternary systems:

- C-Cr-Si
- Cr-Ni-N
- Fe-Cr-Cu
- Fe-Cr-Si
- Fe-Ni-N
- Fe-Ni-Si
- Fe-Cr-Mo-C
- Fe-Cr-C-N
- Fe-Cr-Mn-N
- Fe-Cr-Mo-N
- Fe-Cr-Ni-N
- Fe-Cr-Si-N
- Fe-Cr-Ni-Al
- Nb-V-C-N

TCFE3.0 to TCFE4.0

TCFE4 released in 2005-6.

This release builds on the previous version of the database, i.e. TCFE3, and additionally includes a complete assessment of molar volume data for all phases typically present in steels.

The benefit from having molar volume data included in the database is the possibility to calculate and plot directly within Thermo-Calc:

- Volume fraction of phases.
- Volume for a phase or for the whole system.
- Density for a phase or for the whole system.
- Lattice parameters (for phases having cubic structures).
- Relative length change and thermal expansion coefficients.

TCFE2.0 to TCFE3.0

TCFE3 released in 2002-3.

TCFE3 is the successor of the well-known TCFE2 (i.e., TCFE-2000 or TCFE2K) thermodynamic database for steels and Fe-alloys. A number of improvements have been made in order to increase the predictive capability of the TCFE3 database.

Some of the major improvements are:

- Improved data for the important δ -phase in stainless steels.
- Addition of data for the binary Nb-Ni system.
- Improved data for the ternary Fe-Cr-Mo system.
- Improved data for the ternary Cr-Mo-Ni system.
- Improved data for the fcc Ti(C,N) carbonitride.

TCFE1.0 to TCFE2.0

- TCFE 2.0 was released in 1999/2000 and was also called TCFE2000.
- TCFE1.5 was released in 1996 and also called FEDAT.