

CrossFire[™] Gmelin Data Fields Reference Guide





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1 General

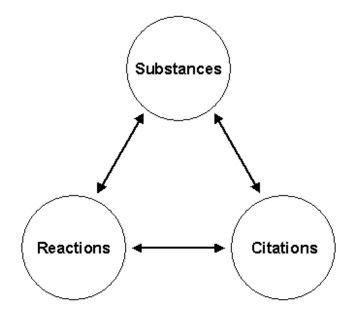
1.1 Introduction

The GMELIN File is a database, which contains information about structural, chemical, and physical information on inorganic and organometallic compounds.

This is an implementation under CrossFire. The MDL CrossFire system is a complete chemical information system comprising of the Gmelin Database, the CrossFire search and retrieval server software, and the Commander client.

When using CrossFire, the Commander is configured with a Structure and Reaction editor, a Fact editor and the Display Hits module. These are all linked together through the Commander to give one coherent interface. The Commander provides the means to access the CrossFire databases; allowing searches to be carried out, results displayed, and the database to be browsed - thus giving you the ability to navigate through the world of inorganic and organometallic chemistry.

The CrossFire system provides a new way of handling chemical information. The emphasis is on content, not on form. There are no artificial barriers between the different forms of viewing information. CrossFire allows you to search and display chemical information as you prefer, whether in terms of Substances (structures and properties), Reactions, or Citations and provides hyperlinks between the different forms.





Substances:

The database contains a broad variety of chemical and physical parameters for approximately 2 million inorganic and organometallic compounds. You can access structures and their associated chemical, physical, and bibliographic information. A specified substance is directly accessible via its unique GMELIN Registry Number, which is assigned during registration of new input. Hyperlinks are available to reactions and literature references, as well as, to other referenced compounds such as precursors or components.

Reactions:

The database contains over 1.5 million reactions, which are searchable with CrossFire as reaction substructures. You define the role of a structure or substructure in the reaction mode. Furthermore it is possible to do a factual reaction search or a search combining structural and factual information. Hyperlinks are provided to substances and to literature references.

Properties:

The database contains a very broad variety of chemical and physical parameters. The facts of a substance are shown in the field availability list. Hyperlinks are provided to the fact details. You can search for the properties using predefined forms or tabular queries.

Citations:

These are stored in a separate domain of the database. This facilitates citation searching and a citation orientated display. It is also possible to search for titles and abstracts. You can display, in one document, all of the compounds and reactions that have been abstracted from the original reference. Hyperlinks are provided to substances, components and reactions.

Hyperlinks:

These are provided for substances, reactions and citations. Thus, for example, when you are viewing a reaction one click of the hyperlink calls up the substance record for one of the reaction partners. Similarly, when viewing some properties, simply click the citation hyperlink to call up the citation record showing all the compounds and reactions abstracted from that reference.

Display Context:

You can carry out structure searches (on the whole database or on subsets), factual searches, and combined searches. The Display Context of the search, which is usually automatically set at search time (a search for reactions will result in a display in reaction context), can be subsequently changed for either single compound, lists of compounds (or reactions or citations) or for the whole hit set. By using combinations of searches, changes of context and hyperlinks, navigation through the database results in not only solutions to inorganic chemistry problems but can also lead to discoveries by "chance".



1.2 Database Contents

There are two different sources for the Gmelin database:

- 1. The first source of information is the Gmelin Handbook of Inorganic and Organometallic Chemistry, from 1817 to 1975, including all main volumes and supplements. It presents critically evaluated facts based on original literature and systematically arranges for a given substance.
- 2. The second source of information is a series of 110 scientific journals dealing with inorganic, organometallic, and physical chemistry, as well as, with physics. From these journals inorganic and organometallic compounds, together with their relevant facts, are extracted. Information from this source is from 1975 to the present.

This data source is continuously evaluated and updated to provide data for all data fields.

1.3 Acceptance Criteria for Substances

The GMELIN database is a factual database on inorganic and organometallic compounds.

Each substance abstracted (together with its possible components) is assigned a unique number, the GMELIN Registry Number (GRN), during registration procedure. This registry number is the link between structure and data.

The following substances are designated as Gmelin substances:

- all compounds without carbon
- compounds, which contain carbon, containing at least one 'Gmelin element'
- all elements, including carbon (including all modifications and fullerenes)
- alloys and multi-components with carbon components
- carbides and carbideoxides
- carbonic acids and the thio- and seleno-analoges as well as their salts with 'Gmelin cations'
- compounds, in which carbon is present exclusively in the form of cyanide or isocyanide, e.g. HCN, HOCN, HSCN, their isoacids and salts, dicyan, polycyan, cyanamide
- and small molecules like CO, CS, CO₂, CS₂, COS, or phosgene.



"Gmelin" elements are shaded green in the periodic table:

Н																	He
Li	Be											В	С	N	0	F	Ne
Na	Mg											AL	Si	Ρ	S	CI	Ar
K	Ca	Sc	Ti	V	Cr	Mn	Fe	Со	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr
RЬ	Sr	Y.	Zr	NЬ	Мо	Тс	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Te	1	Xe
Cs	Ba		Hf	Ta	W	Re	Os	- Ir	Pt	Au	Hg	TI	РЬ	Bi	Po	At	Rn
Fr	Ra																
		La	Се	Pr	Nd	Pm	Sm	Eu	Gd	ТЬ	Dy	Ho	Er	Tm	Yb	Lu	
		Ac	Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr	

All further substances belong to the "organic" domain and can be found in CrossFire Beilstein.

Reference compounds:

If references are made in the preparation or the chemical behavior to starting materials or products, which are defined as Beilstein compounds, and which are necessary for the correct description of the reaction (e.g. iron(III) chloride with pyridine) then the formula and structure of the Beilstein compound is abstracted. However their properties are included in the Beilstein database.

The Gmelin database distinguishes between single- and multi-component compounds.



1.4 Single-Component Compounds

These are "classical" compounds having a stoichiometric formula. Substances of this type may consist of one (e.g. oxygen, ferrocene) or more fragments (e.g. salts, adducts).

Substances of this type are

Elements	e.g. H ₂ , Fe, Cu
Simple Molecules	e.g. H ₂ O, CO ₂ , HCI
Inorganic Salts	e.g. NaCl, Na ₂ SO ₄ , KMnO ₄ , KOH, CaTiO ₃
Adducts	e.g. Na₂SO₄*10H₂O, CoCl₂*4NH₃
Coordination	e.g. $Fe(C_5H_5)_2$, $Mo(CO)_6$, $[Co(NH_3)_6]Cl_3$,
Compounds	$[Co(NH_3)_5(SCN)]Cl_2$
Polymers	e.g. $(HF)_n$, $\{Pt(CO)_2\}_n$, $\{Fe(C_5H_4)_2\}_n$
Solid Solutions (exact formula)	e.g. Na _{0.5} K _{0.5} Cl, Ca _{0.2} Y _{0.8} Ba ₂ Cu ₃ O _{6.78}
Minerals	e.g. dolomite $CaMg(CO_3)_2$, pyrite FeS_2
Alloys (as formula)	e.g. $Fe_{0.66}Pd_{0.34}$, $Ni_{45}Zr_{55}$, Cu_2TiAl
Glasses / Ceramic	
Materials	e.g. BaTiO ₃ , SiO ₂ , YBa ₂ Cu ₃ O _(x)
(as formula)	

Single-component compounds are divided into fragments, if they consist of discrete polyatomic ions or molecules. The individual fragments are separated from one another by asterisks (*) in a linearized molecular formula. Rules on whether a compound is fragmented or not:

1. Solid state structures, ionic compounds:

 Substances are divided into formal fragments if the compounds contain at least one discrete polyatomic ion or molecule in the crystal, the gas phase, the liquid phase, or in solution. The structures of these discrete ions or molecules are available.
 e.g. 2Na⁽¹⁺⁾*SO4⁽²⁻⁾=Na2SO4
 e.g. K⁽¹⁺⁾*MnO4⁽¹⁻⁾=KMnO4
 e.g. 2Na⁽¹⁺⁾*SO4⁽²⁻⁾*10H2O=Na2SO4*10H2O
 e.g. Ca⁽²⁺⁾*Mg⁽²⁺⁾*2CO3⁽²⁻⁾=CaMg(CO3)2



 Substances are not divided into fragments, if the compounds consist exclusively of single-atomic ions (including OH, SH⁻, SeH⁻, and TeH⁻), e.g. NaCl, KOH, or have no discrete polyatomic ion or molecule, e.g. CaTiO₃

2. Organometallic and coordination compounds:

- Organometallic compounds and coordination compounds with known manner of ligand coordination are stored with the complete formula and structure.
 e.g. Fe(C₅H₅)₂
- Organometallic salts and ionic coordination compounds with known manner of ligand coordination are divided into cations and anions. The complete structures of the ions are available.
 e.g. [Co(NH₃)₆]⁽³⁺⁾*3Cl⁽¹⁻⁾=[Co(NH₃)₆]Cl₃
- Organometallic compounds and coordination compounds with unknown manner of ligand coordination are divided into fragments by separating all ligands from the metal center.

e.g. If there is an unknown coordination of thiocyanate to $Co^{(3+)}$ in $[Co(NH_3)_5(SCN)]Cl_2$ (coordination by N or S atom possible), the formula is given as $Co^{(3+)*}5NH_3*SCN^{(1-)*}2Cl^{(1-)}=[Co(NH_3)_5(SCN)]Cl_2$.

1.5 Multi-Component Compounds

Multi-component compounds consists of two or more single-component compounds having no stoichiometrical relationship to each other. Each single-component compound may consist of one or more fragments.

For multi-component compounds the linearized formulas of individual compounds are separated from one another by a hush (#).

Substances of this type are:

Description	Gmelin formula	Components		
(Multi-Component) System	ms			
e.g. zinc oxide and titanium oxide	ZnO#TiO2	ZnO and TiO $_2$		
e.g. lead sulfate, lead oxide and silicium dioxide	PbSO4#PbO#SiO2	$PbSO_4$ and PbO and SiO_2		
Solutions				



aqueous solution of sodium chloride	NaCl#H2O	NaCl and H_2O
aqueous solution of lithium nitrate	LiNO3#H2O	LiNO $_3$ and H $_2O$
solution of aluminium chloride in phosgene Doped Compounds	AICI3#COCI2	$AICI_3$ and $COCI_2$
silicon doped with		
boron nitride	Si#dotBN	Si and BN
tin-doped indium(III)		
oxide	In2O3#dotSn	In_2O_3 and Sn
cadmium		
titaniumhexafluoride		CdTiF₅*6H₂O and Mn ⁽²⁺⁾
hexahydrate doped	CdTiF6*6H2O#dotMn(2+)	
with manganese(2+)		
Solid Solutions (with form	nula range or undefined formula)	
	•	
solid solution of zinc sulfide (range x=0-015) and zinc selenide	ZnS(x)Se(1-x) x=0-0.15	ZnSe and ZnS _{0.15} Se _{0.85}
sulfide (range x=0-015)	•	ZnSe and $ZnS_{0.15}Se_{0.85}$ Na ₂ SO ₄ and Cs ₂ SO ₄
sulfide (range x=0-015) and zinc selenide solid solution of sodium sulfate and cesium	ZnS(x)Se(1-x) x=0-0.15	
sulfide (range x=0-015) and zinc selenide solid solution of sodium sulfate and cesium sulfate solid solution of potassium chloride and potassium bromide note: the description of solid solutions is dependent on the author's declaration	ZnS(x)Se(1-x) x=0-0.15 Na2SO4#Cs2SO4	Na ₂ SO ₄ and Cs ₂ SO ₄

and 5 weight% nickel



Glasses / Ceramic Materials (in percentage)

silicium dioxide glass containing 10-15 weight% aluminium oxide and 10-15 weight% boron oxide

Al2O3(10-15),B2O3(10-15), O2Si (W)

 Al_2O_3 and B_2O_3 and SiO_2

According to the fragmentation rules of single-component compounds, the following examples mentioned in the table are divided into fragments. Each single-component compound is viewed isolated on fragmentation.

 $\label{eq:pb} \begin{array}{l} Pb^{(2+)*}SO_4^{(2-)} = PbSO4\#PbO\#SiO2\\ Li^{(1+)*}NO_3^{(1-)} = LiNO3\#H2O\\ Cd^{(2+)*}TiF_6^{(2-)*}6H_2O = CdTiF6*6H2O\#dotMn^{(2+)}\\ 2Na^{(1+)*}SO_4^{(2-)} = Na2SO4\#2Cs^{(1+)*}SO4^{(2-)} = Cs2SO4\\ \end{array}$

Note:

Multi-component compounds have no overall molecular formulas, because there is no stoichiometric relationship between the single components.

1.6 Registry Citeria

There can be more than one GMELIN Registry Number for the same Molecular Formula.

The following critera are used to assign a compound its own GMELIN Registry Number:

- Molecular Formula
- Structure STR comprising the connectivity of atoms, total charge of fragments or compounds, and stereo descriptors
- The types of substance
 - glasses or ceramic materials
 - isomorphous or diadochous compounds, solid solutions
 - isotopes or isotope containing compounds
 - minerals
 - polymers
- Modification



Note:

The family names alloys and coordination compounds are not criteria for registration The chemical name of a substance is not a criterion for registration The substance physical state is only displayed within the relevant facts: DIC, ELCO, ST, COM, MVOL, DEN, RI, VP, VPFI, TCND, TEC, CV, CVFI, CP, CPFI; GFOR, GFFI, HFOR, HFFI, SFOR, SFFI, DYVI, and KIVI.

1.7 Acceptance Criteria for Attributes

Acceptable data for the Gmelin database falls into the following categories:

- Characterization / Identification
- General Information
- Physical Properties
- Properties of / in Systems
- Quantum Chemical Calculations
- Electrochemistry
- Chemical Properties
- Bibliographic Information

The general rules for accepted data will be discussed in the relevant chapters. Subjects are sorted in hierarchical order. To find information for a specific topic, use the Help index or the Help Find function.

1.8 Database Searching

CrossFire Gmelin offers you different ways of searching:

- searching for compounds either via characterization criteria or via structure
- searching for reactions either via factual fields or via structure
- searching for physical and chemical properties
- searching for bibliographic information

All of these topics can be combined with each other.



Structures:

Full structures, substructures, generic structures, and stereoisomers can be searched for. You can use global query options such as "Free Sites on All Atoms", or set options on specific atoms and bonds. Query atom types such as "X", to designate any halogen can be set as well as Atom lists (e.g. A1= Fe, Co, Ni). Predefined generic groups, e.g. ALK (alkyl), and user defined generic groups are easily attached.

Note:

You should be aware that only compounds or compound parts, which are formed by discrete molecules or ions, have a structure. Compounds with a solid-state structure will not have a structure in the Gmelin database.

Reactions:

The role of a substance (reactant or product) can be defined in the reaction mode of the structure editor.

Thus you can search for the following reaction types:

? => B	"how do I make this type of compound"
A => ?	"what do these starting materials give"
A => B	"general or specific transformations".

Properties:

A very broad variety of chemical and physical data can be searched for as numeric values and ranges, strings, and keywords. The CrossFire database contains more than 6 million reports on chemical and physical properties. You can search for molecular formulas, reaction conditions, electric, or magnetic data. All of these are stored in discrete fields.



1.9 Searching for Facts

The database contains numeric and textual data in more than 800 searchable fields. Searching is carried out using the Fact Editor, where the matrix form, field name lists, and index expand buttons make the query formulation straightforward. High flexibility is given by option to carry out numeric, text, field availability, and hierarchical searching. A successful search retrieves records, each of which contain all the information of a particular substance, reaction, or citation in the database. For information on structure and reaction searching, and displaying hit lists, see the relevant chapters.

- Substance Factual Search
- Reaction Factual Search
- Citation Factual Search
- Numerical Search
- Text Searching
- Keyword Searching
- Truncation
- Operators
- Field Availability Search
- Hierarchical Group Code Search

1.9.1 Substance Factual Search

This is a search specific for substance related facts in the database. Whether numeric, numeric range, string, or keywords, you will retrieve all substances in the database which contain the matching terms. The display context is automatically set to Substance. This type of search applies to most of the fields in the database, with the exceptions of reactions and citations.

See also: Basic Index for Substances

1.9.2 Reaction Factual Search

This is a search specific for graphical reaction related facts. The display context will automatically be set to Reactions.

See also: Basic Index for Reactions.



1.9.3 Citation Factual Search

This is a search specific to citation related fields. The display context will automatically be set to Citation.

See also: Basic Index for Citations

1.9.4 Numerical Search

The GMELIN file contains more than 100 search fields with numeric data for chemical and physical properties of substances.

In numeric fields searching can be carried out for single value or ranges, as well as, greater than/less than a certain value. The values of physical properties and parameters are stored in the units and dimensions of the CrossFire Database, these are the units and dimensions most commonly used in inorganic chemistry and are, therefore, not necessarily SI units.

For searching a single value, e.g. a melting point (field code mp) at 223°C, the following values and ranges may be employed:

- the exact temperature 223°C
- the range from e.g. 220 225°C (temperature is within the range)
- the ranges from e.g. 220 223°C and 223 225°C (temperature is either upper or lower limit of the range)

When searching for a range, e.g. melting point mp=220-225, all records will be retrieved where the value falls within this range, no matter whether they are derived from a single value or a range.

Numerical values are also present in some string fields, where they are searchable as strings.

Scientific Notation:

Numeric values of physical properties may cover a large range of numbers, for example, from $1.88E-53 \text{ mol}^2/l^2$ for the solubility product of the alpha modification of mercury(II) sulfide in water at 25°C to 5.309E13 Hz for the molecular rotation constant about a particular axis of the Ar*HCONH2 molecule. For this reason the scientific notation for numbers is used throughout to display Numeric values.



1.9.5 Text Search

Searching for strings or partial strings can be carried out in text fields. To search for a partial string use the wild card symbols. It is highly recommended to look in the index and use "List" to check the spelling of terms before carrying out a search. Index terms can be directly copied into the Fact Editor for searching.

The text fields are indexed in several ways:

String (Phrase)

The complete string is indexed without splitting:

e.g. Molecular Formula, Keyword Fields

String (Wordwise)

The entry is split into words using the following symbols:

.,;:()<>[]{}/=?*+-#!\$% space

These symbols will not be found in the index, and should therefore not be searched for.

1.9.6 Keyword Search

Many topics in the Gmelin database have fields of the type Description. These fields contain entries, which usually derive from a controlled vocabulary. These keywords are indexed as phrases and can be searched as a typical text field.

It is recommended always to use the List function of the CrossFire system before starting a search. The keywords can be copied out of the list into the edit mask for searching.

1.9.7 Truncation

All text fields can be searched by using truncation (wild card) symbols.

- * any number of characters (including no character)
- ? any single character
- ?? any two characters

Both right, left, and middle truncation are allowed.



Left truncation is useful for phrase fields where the searched for term occurs within the phrase.

Right truncation is useful for free text fields, e.g. Abstracts, to make sure that all occurrences of a term are found. Thus, to retrieve the records with both photochemistry and photochemical the following should be input:

ab=photochem*

It is also particularly useful for the Authors fields, due to the different rules used by journals. Author's first names are often written differently in various journals. Right truncation can be used to ensure that all articles of a particular author are found.

1.9.8 Field Availability Search

This is a search for the occurrence of a fact in the database. Thus you can search for all compounds with physiological information by entering PHYB in the Field Name column and leaving the Value column empty.

Field name	Relation	1
Physiological Behavior(PHYB)	exists	-

1.9.9 Hierarchical Group Code Search

This is a search for the presence of data in any specified group of fields. Data fields are grouped together according to the hierarchy of the Gmelin Database. The group fields, e.g. Spectroscopic Information (SPE) (which identifies all particular data fields concerned with spectoscopic data information) are searchable using their group codes.

Thus you can carry out a search for compounds for which at least one piece of information from any type of spectral data, e.g. NMR, IR, etc., is available by entering SPE in the field name column.

Operator	(Field name	Relation	n	Field content	List
		Spectroscopic Information(SPE)	exists	•		•



1.9.10 Operators

In the operator column of the CrossFire Commander Fact Editor as well as in the fact query window of the Commander itself the Boolean operators, AND, NOT, OR and PROXIMITY can be input. For search in text fields the two additional operators NEAR and NEXT are also available:

AND Operator OR Operator NOT Operator PROXIMITY Operator NEAR Operator NEXT Operator

1.9.10.1 AND Operator

With this Boolean operator the two connected search terms will only be retrieved as a hit if they are present in the same record.

e.g. mp > 1000 AND tec < 0

This search will result in a list of compounds, where the compounds melt above 1000°C and have negative expansion coefficients.

When using this operator with hit lists, e.g. from structure searches, the resulting list will only contain those records present in both original lists.

e.g. .q1 AND .q2

1.9.10.2 OR Operator

This Boolean operator retrieves only hit lists in which the individual records contain at least one of the search terms.

e.g. mp > 1000 OR tec < 0

This search will result in a list of compounds, where either the compounds melt above 1000°C or have negative expansion coefficients (including records with both properties).



When using this operator with hit lists, e.g. from structure searches, the resulting list will contain all records from both original lists.

e.g. .q1 OR .q2

1.9.10.3 NOT Operator

This Boolean operator retrieves only those records from the first hit set not present in the second hit set.

e.g. type = "alloy" NOT (elc = "Fe" OR elc = "Ni")

This example will retrieve all alloys without the elements iron or nickel.

When using this operator with hit lists, e.g. from structure searches, the resulting list will contain all records from the first list minus all records from the second list.

e.g. .q1 NOT .q2

1.9.10.4 PROXIMITY Operator

This operator is a version of the AND operator, whereby the two connected search terms will only be retrieved as a hit if they are present in the same occurrence of a fact. This operator is very important when searching for facts that contain sub-fields or parameter fields.

Thus when searching for a boiling point measured at a particular pressure, the PROXIMITY operator should be used to ensure that both retrieved terms come from the same occurrence of the Boiling Point fact.

e.g. bp = 120 PROXIMITY bp.p = 750-760

This example will retrieve all compounds having a boiling point of 120 ° C measured in the pressure range of 750 to 760 Torr.

1.9.10.5 NEAR Operator

This operator retrieves hit lists in which the individual records always contain both of the search terms and are adjacent to each other.

e.g. ab=crystal* NEAR ab=structure*



1.9.10.6 NEXT Operator

This operator retrieves hit lists in which the individual records always contain both of the search terms and where the first search term will appear before the second search term.

e.g. ab=solid* NEXT ab=state*

1.10 Description of Data Fields

The description of data fields will be illustrated using the field MSUS as an example:

Description:

The volumetric magnetic susceptibility is the difference in the magnetic permeability of a substance and of vacuum, divided by the permeability of vacuum. The specific (or mass) magnetic susceptibility of a substance is the quotient of its volumetric susceptibility and its density. The molar magnetic susceptibility of a substance is the product of its volumetric susceptibility and its molar volume. The <u>Magnetic Susceptibility</u> field contains the values for a given substance in the literature. The search field is a Numeric range search field and is linked to the associated temperature MSUS.T. Further information about the magnetic susceptibility is given as diagram or equation using the parameter field codes MSUS.DGM and MSUS.EQN respectively. Information about pretreatment of samples, or concentration of solutions, is given in the field remarks MSUS.MR.

Note:

Searching for magnetic susceptibility diagrams or equation field codes requires the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
MSUS	Molar Magnetic Susceptibility	Numeric	ml mol ⁻¹
MSUS.VALA	Specific Magnetic Susceptibility	Numeric	l kg⁻¹
MSUS.T	Temperature	Numeric	°C
MSUS.DGM	Diagram	String (phrase)	
MSUS.EQN	Equation	String (phrase)	
MSUS.MR	Remarks	Wordwise	



The GMELIN File offers approximately 800 textual, numeric, and flag fields. In general, flag fields can be searched by their group codes. However, if the flag field is part of further data fields of a group code, it is indexed as a String with a controlled term. The corresponding field value is given in the note. For an easy search you can generally use wildcards ("*").

Physical properties can appear as molar or specific and are stored in individual fields without the extension property (molar) and property.vala (specific).



2 Data Field Description

2.1 Data Field Description

The GMELIN Database is organized hierarchically. Click one of the hyperlinks to follow the hierarchy of the datastructure and to get more detailed information. The following is the top level of the hierarchy:

Basic Indexes (BI)

Bibliographic Information (BIB)

Characterization (ID)

General Information (GEN)

Physical Properties (PHY)

Properties of / in Systems (MCS)

Quantum Chemical Calculations (QCC)

Electrochemistry (EC)

Chemical Properties (CHE)

2.2 Basic Indexes

2.2.1 Basic Indexes (BI)

Basic Indexes are collective indexes, which allow convenient searching without thinking about field codes. The Gmelin database offers the following Basic Indexes:

All GRN's (AGRN) All Chemical Names (ACN) All Chemical Name Segments (ACNS) Substance Basic Index (BISUB) Reaction Basic Index (BIREA) Citation Basic Index (BICIT)



2.2.2 Substance Basic Index (BISUB)

Description:

This is the basic index for substances. If a user enters keywords and operators, such as "and", "or" or "not", in the text search area of the Commander, without specifying a field name and with staying in the substance context, the keywords will automatically be searched in this index. The substance basic index belongs to the substance context and a search will result in a substance/structure hitset.

The Substance Basic Index contains the following fields:

RN	CAS Registry Number
.CN/.EDN/ .PRO	Substance Names
.PA/.PHAS*	Chemical Partners
.TYP	Specification to Facts
.MET	Methods
.MR	Remarks

Type of Indexing:

Strina	(wordwise))
Cunig	(1001000)	

Except: CAS Registry Number: String (phrase)

Example1:

Search for all substances with the information "triphos":

Field Name	Value
------------	-------

BISUB triphos

The hits show the ligand itself, as well as, compounds containing the ligand.

You should be aware that only those substances are retrieved having a name assigned.



Example2:

Search for all substances with the information "pressure":

Field Name Value

BISUB pressure

Substances are retrieved having properties depending on pressure.

2.2.3 Reaction Basic Index (BIREA)

Description:

This is the basic index for reactions. If a user enters keywords and operators, such as "and", "or" or "not", in the text search area of the CrossFire Commander without specifying a field name and with staying in the reaction context the keywords will automatically be searched in this index.

The Reaction Basic Index contains the following fields:

Field Code	Name
RX.RCT	Reactant (name)
RX.PRO	Product (name)
RX.RGT	Reagent
RX.BPRO	Byproduct
RX.INT	Intermediate (name)
RX.CAT	Catalyst (name)
RX.SOL	Solvents
RX.SCON	Special Conditions
RX.GCON	General Conditions
RX.ISOL	Purification, Isolation



Type of Indexing:

String (wordwise)

Example1:

Search for irradiation reactions:

Field Name	Value
------------	-------

BIREA irradiation

Example2:

Search for all reactions with the information "mechanism":

Field Name Value

BIREA mechanism

The hits show reactions with descriptions of reaction mechanism in the articles.

2.2.4 Citation Basic Index (BICIT)

Description:

This is the basic index for citations. If a user enters keywords and operators, such as "and", "or" or "not", in the text search area of the Commander without specifying a field name and with staying in the citation context the keywords will automatically be searched in this index.

The Citation Basic Index contains the following fields:

Field Code	Name
AU	Author
JT	Journal Title
JTT	Journal Title (Translation)
ТІ	Article Title



AB Article Abstract

AB.KW Article Abstract Keywords

The citation basic index belongs to the citation context and a search will result in a citation hitset (reference list).

Type of Indexing:

String (wordwise)

Example1:

Search for all documents with the information "enzym*":

Field Name Value

BICIT enzym*

The hits show documents containing phrases like enzymes, enzymatic activity, enzymatic methods, and enzymatic reactions in titles, keywords, and abstracts.

Example2:

Search for all documents with the information "industr*":

Field Name Value

BICIT industr*

Documents are retrieved containing phrases like chemical industry, industrial processes, industrial applications, and industrial tools in titles, keywords, and abstracts.



2.2.5 All GRN's (AGRN)

Description:

This is the basic index for Gmelin Registry Numbers (GRN's).

This basic index belongs to the substance context and a search will result in a substance/structure hitset.

The basic index contains the following fields:

Field Name	Name
GRN	Gmelin Registry Number
FRAGRN	Fragment Gmelin Registry Number
CGRN	Component Gmelin Registry Number
MULT.GRN	System Component Registry Number
AZE.PG	Azeotropes (GRN)
POT.EGRN	Electrochemical Coreactant(s) (Registry Number)
POT.PGRN	Electrochemical Product(s) (Registry Number)

Notes:

The AGRN index is useful where referenced compounds need to be found.

Type of Indexing:

Numeric

Example1:

Search for all substances containing GRN 171518 (= KH(HCO₂)₂):

Field Name Value

AGRN 171518

The first hits show the single-component compound $KH(HCO_2)_2$. The second hit is a multi-component compound, the solution of $KH(HCO_2)_2$ in HCO_2H having $KH(HCO_2)_2$ as component.



Example2:

Search for all substances containing GRN 8908 (= BaTiO₃):

Field Name Value

AGRN 8908

The hits show all substances having $BaTiO_3$ as component, e.g. doped compounds, ceramic materials, or systems. Furthermore, hits are retrieved with $BaTiO_3$ as system component.



2.3 Characterization

Search Fields:

Identification of Substance (IDE) Formula Search Data (FOR) Composition (CMPS) Ligands around Metals (LIG)

2.3.1 Identification

2.3.1.1 Substance Identification (IDE)

GMELIN Registry Number (GRN) Search MF Range (MOFO) Alloy Search Field (ALLOY) Linearized Structure Formula (LSF) Molecular Formula (MF) Fragment Molecular Formula (FRAGMF) Fragment GMELIN Registry Number (FRAGRN) Ligand Molecular Formula (LIGMF) Ligand Formulas (LIGFO) Ligand Formula Count (LIGFC) Number of Components (NC) Number of Fragments (NF) Number of Structures (NSTRUC) Molecular Weight (MW) CAS Registry Number (RN) Chemical Name (CN) Chemical Name Segment (CNS) Type of Substance (STYPE)



Modification (MODTXT) Entry Date (ED) Update Date (UPD)

See also:

Ligands around Metals (LIG) Formula Search Data (FOR) Composition (CMPS)

2.3.1.2 GMELIN Registry Number (GRN)

Description:

The numeric search field GMELIN Registry Number contains the accession number, which is assigned at the registration of a substance. This number is unique for a given substance and depends on the formula, structure, modification, and the family names isotope or isotope containing compounds, mineral, polymer, isomorphous or diadochous compound, solid solution, and glass or ceramic material.

Note:

The chemical name of a substance is not used as a criterion for the assignment of the GMELIN Registry Number.

Type of Indexing:

Numeric

See also:

AGRN



Example:

Search for substance with the Gmelin Registry Number 3385

3385

Field Name Value

GRN

2.3.1.3 Search MF Range (MOFO)

Description:

The MOFO field is a non-display field that allows the search of compounds by normal chemical formula using chemical symbols in capital and small letters as well as the search of ranges with molecular formula. It is not possible to have a look at this field via the List function, because it is calculated "on the fly" during the search.

Tips and Hints:

 If you want to restrict searches to compounds with a specific number of components or fragments it is necessary to combine the query with restrictions in the field NC or NFRAG, respectively. Otherwise, multi-component compounds with your compound as one component will also be found.
 For example, to search for a one-component compound, NC = 1 has to be set additionally.
 This is a powerful search code for formulas, especially when searching for inorganic compounds without molecular structures.

- 2. You can search for molecular formulas with stoichiometric range (see example (e)).
- 3. The symbols greater than/less than (<>) are not allowed, instead you must enter a large range; for example, more than three Oxygen atoms can be input as in (f).
- 4. You can search for certain molecular formulas by typing an asterisk at the end of a value (see example (g)).
- 5. Please note that no charged compounds can be searched. For this the search fields MF or FRAGMF should be used.

Type of Indexing:

String (phrase)



Examples:

	Search for	Field Name	Value
а	sodium chloride	MOFO	NaCl
b	sodium sulfate	MOFO	Na_2SO_4
С	iron(III) chloride with 6 ammonia	MOFO	FeC ₅*6NH₃
d	tris(tetrabutylammonium) hexakis-	MOFO	$(N(C_4H_9)_4)_3[Os(SCN)_6]$
u	(thiocyanato) osmate(III)		
е	zirconium oxides with defined zirconium	MOFO	Zr ₍₁₋₃₎ O ₍₁₋₅₎
C	and oxygen range Zr:1-3, O:1-5		L (1-3) (1-5)
f	yttrium barium copper oxides with	MOFO	YBa ₂ Cu ₃ O ₍₄₋₉₉₎
•	variable O content (>3)		1 2 4 2 0 4 3 0 (4-33)
g	any dodecamolybdates	MOFO	Mo ₁₂ O ₄₀ *

2.3.1.4 Alloy Search Field (ALLOY)

Description:

The alphanumerical search field Alloy Search Field allows the search of alloys, glasses and ceramic materials specified by percentages. The content of each constituent, an element or compound, can be specified by a percentage value or range. Additionally the form of the percentage information must be specified as

- A atom or mol percent
- V volume percent
- W weight percent
- X indefinite percent.

Note:

Compounds have to be entered using "Hill order".

Constituents in traces are specified as 0 %.

Type of Indexing:



String (phrase)

Example:

Search for a silicium dioxide glass with 10-15 weight% Al₂O₃ and 10-15 weight% B₂O₃.

Component Formula: (e.g. Fe or Fe203)	Percentage: (Number or range: 20 or 20-40
Al203	10-15
6203	10-15
025i	- _
[
[
[ī [.

being equivalent to query: alloy="Al₂O₃(10-15),B₂O₃(10-15),O₂Si (W)"

2.3.1.5 Linearized Structure Formula (LSF)

Description:

The alphanumerical search field Linearized Structure Formula contains a linear representation of the structure of a compound. The formula may contain isotopes.

Compounds are divided into fragments, if discrete polyatomic ions, molecules, or ligands with unknown coordination exist. For multi-fragment compounds the molecular formulas of individual fragments are separated from one another by asterisks (*).

For systems, the formulas of individual compounds are separated from one another by a hush mark (#). Each compound can consist of several fragments.

For doped compounds, the formulas of the dopants are separated from the basis compound by the designation '#dot'.

Note:



If a compound includes an adduct, which has undefined prefix-factor, this adduct is formulated with the prefix-factor 99.

Tips and Hints:

Searches for compounds with isotopes or multi-fragment compounds with their reported stoichiometric multipliers as well as for doped materials and systems can be carried out using this field.

Type of Indexing:

String (phrase)

Examples:

	Search for	Field Name	Value
(a)	sodium carbonate	LSF	2Na(1+)*CO3(2-)=Na2CO3
(b)	hydrated nickel acetate	LSF	Ni(2+)*2CH3COO(1-)*99H2O= Ni(CH3COO)2*99H2O
(c)	boron doped silicium	LSF	Si#dotB
(d)	aqueous solution of sodium chloride	LSF	NaCl#H2O
(e)	system of zinc oxide and titanium dioxide	LSF	ZnO#TiO2

2.3.1.6 Molecular Formula (MF)

Description:

The alphanumerical search field Molecular Formula contains the molecular formula of a substance. Within multi-component compounds, each component is searchable separately.



Note:

The Molecular Formula has to be entered using "Hill order".

Fragment formulas are arranged as follows:

- Carbon containing compounds come first
- number of carbon atoms
- number of hydrogen
- alphabetical order of elements
- Non carbon containing compounds follow
- alphabetical order of elements

Type of Indexing:

String (phrase)

Examples:

 $\begin{array}{l} C_{6}H_{3}N_{3}O_{7} \ ^{*} \ C_{5}H_{5}N \\ 2C_{6}H_{7}N \ ^{*} \ C_{6}H_{6}CI_{6} \ ^{*} \ 2CIH \end{array}$

Searching a specific compound in the MF field excludes all isotope labeled compounds. Searching an isotope molecular formula exactly hits the isotopic labeled compounds specified in the query.

	Search for	Field Name	Value
(a)	sodium acetate	MF	C2H3NaO2
(b)	sodium chloride	MF	CINa
(c)	sodium sulfate	MF	Na2O4S
(d)	hafnium	MF	Hf
(e)	hydrofluoric acid	MF	FH



2.3.2 Fragment Molecular Formula (FRAGMF)

Description:

The search field fragment molecular formula contains the molecular formula of a fragment. It can be used to retrieve all compounds containing a given fragment (and the fragments themselves).

Note:

The fragment molecular formula has to be entered using "Hill order". Fragment formulas are arranged as follows:

- Carbon containing compounds come first
- number of carbon atoms
- number of hydrogen
- alphabetical order of elements
- Non carbon containing compounds follow
- alphabetical order of elements

Examples:

 AI_2O_3 , O_2Si , OZn, O_5P_2

Tips and Hints:

The fragment molecular formula is combined with the order of the corresponding component. Therefore, a common search should be done by combination of the fragment molecular formula with an asterisk (*).

Type of Indexing:

String (phrase)

	Search for	Field Name	Value
(a)	substances containing acetate fragments	FRAGMF	C2H3O2(1-)*
(b)	substances containing acetate fragments (in single-component compounds or as first component of multi-component	FRAGMF	C2H3O2(1-):1



compounds) substances containing acetate fragments (c) (as second component of multi-FRAGMF C2H3O2(1-):2 component compounds) (d) substances containing FRAGMF O12P4(4-)* cyclotetraphosphate fragments substances containing (e) FRAGMF C6FeN6(3-)* hexacyanoferrate(III) fragments

2.3.2.1 Fragment GMELIN Registry Number (FRAGRN)

Description:

The numeric search field, Fragment GMELIN Registry Number, contains the accession number which is assigned at the registration of a fragment. This number is an unambiguous identifier for the different fragments present in a multi-fragment compound.

Type of Indexing:

Numeric

Example:

Search for	Field Name	Value
substances with the Fragment Gmelin Registry Number 2120	FRAGRN	2120

2.3.2.2 Ligand Molecular Formula (LIGMF)

Description:

The alphanumerical search field <u>Ligand Molecular Formula</u> contains the abstract formula of the coordination centers and all ligand codes of a compound.



The ordering of the formula is as follows:

- all metal centers (ordered alphabetically)
- the ligand codes (see LIGFO) are ordered in the following manner: A, CN, CNO, CNR, CNS, CO, CS, D, L, Q, X
- within the same ligand types, they are ordered on ascending denticity and the frequency is not taken into consideration

Note:

Charges are not taken into account.

Tips and Hints:

Searches for compounds using the GMELIN Ligand Search System are very powerful to look for a class of complexes with special connectivity.

Type of Indexing:

String (phrase)

Related Information:

LIG.LIGMF

	Search for	Field Name	Value
(a)	hexa carbonyl chromium	LIGMF	Cr{CO}6
(b)	tetra halogeno palladium compounds	LIGMF	Pd{X}4
(c)	copper complexes with two halogenides and two ligands connecting by chalcogen atoms	LIGMF	Cu{Q}2{X}2
(d)	Platinum complex with two substituted isocyanide, one ligand connecting by pnictide atom and one ligand connecting by carbon	LIGMF	Pt{CNR}2{D}{L}
(e)	cobalt complexes with three chelate ligands connecting by chalcogen atoms	LIGMF	Co{(2)Q}3
(f)	nickel complex with two tridentate ligands connecting by one pnictide and two chalcogenide atoms	LIGMF	Ni{D(2)Q}2
(g)	ferrocene and substituted ferrocene compounds	LIGMF	Fe{(5)L}2



(h)	bridged ferrocenyl compounds	LIGMF	Fe{(10)L}
(i)	divanadium nonahalogenides	LIGMF	V2{X}9
(k)	molybdenum(6) halogenide(14) cluster	LIGMF	Mo6{X}14
(I)	molybdenum(4) tungsten(2) halogenide(14) cluster	LIGMF	Mo4W2{X}14

2.3.2.3 Ligand Formulas (LIGFO)

Description:

The search field, <u>Ligand Formula</u>, contains the ligand codes according to the Gmelin Ligand Search System, which describes the elements (in groups) and the number of the atoms that are connected to the central metal atom(s). A ligand formula is given for all structured compounds or fragments which contain at least one metal atom and at least one coordinating ligand atom of the classes "L, A, D, Q, X" or a special ligand.

The codes include the following elements:

Code Atom L С А В Si Ge D Ν Ρ As Sb Q 0 S Se Те Н F Х CI At Br Т

There are the following "special" ligands:

Code CO CS CN CNS CNO CNR*)

*) The substituent R has no further bond to metal atoms.

If a ligand is described by a combination of the codes, they are ordered alphabetically and the denticity of the code is put in parentheses in front of the code.

Note:

The ligand formula describes only the denticity of the ligand not the connectivity to the metal center(s).



Type of Indexing:

String (phrase)

Related Information:

LIG.LIGFO

Examples:

	Search for	Field Name	Value
(a)	halogenid or hydride ligand, μ-halogenide or μ- hydride ligand	LIGFO	Х
(b)	ligand connecting by five C atoms to one or more metal centers e.g. $\eta^{\text{5}}\text{-cyclopentadienyl}$	LIGFO	(5)L
(c)	ligand connecting by two chalcogenides e.g. η^2 - or μ -acides (like acetic acid), oxalate (bidentate), di- oles (like ethylene glycol), salicylates, acetylacetonates, dithiols, μ -bisulfido	LIGFO	(2)Q
(d)	ligand connecting by four chalcogenides e.g. silicate, phosphate, 12-crown-4, tetra-thiocyclohexadecane, oxalate (tetradentate)	LIGFO	(4)Q
(e)	ligand connecting by one pnictide und two chalcogenide atoms e.g. diole-amines, imino-diacetates	LIGFO	D(2)Q
(f)	ligand connecting by two pnictide und four chalcogenide atoms e.g. ethylenediaminetetraacetate	LIGFO	(2)D(4)Q

Further examples are available in chapter Ligand Search System.

2.3.2.4 Ligand Formula Count (LIGFC)

Description:

In the field <u>Ligand Formula Count</u>, the frequency of a certain ligand within a compound is indexed.

For ligand codes see LIGFO.



Tips and Hints:

Different ligand formula counts can be combined with the "and" operator. To restrict the search to one complex or complex anion – to one structure – the number of fragments (nfrag) should be set to 1 additionally. Because of this restriction no multifragment compounds are found. However you can find these compounds by the Fragment GMELIN Registry Number.

Type of Indexing:

String(phrase)

Related Information:

LIG.LIGFC

Example1:

	Search for	Field Name	Value
(a)	complex containing one halogenid or hydride, µ- halogenide or µ-hydride ligand	LIGFC	X:1
(b)	complex with six carbonyl ligands	LIGFC	CO:6
(c)	complex with three ligands connecting by two pnictides e.g. ethylenediamine, bipyridyl	LIGFC	(2)D:3



Example2:

Search for complexes containing one η^5 -C ligand, one ligand connecting by two pnictide atoms and three halogenide (or hydride) ligands

Operator	Field Name	Value
	LIGFC	(5)L:1
and	LIGFC	(2)D:1
and	LIGFC	X:3
and	NF	1

The hit set contains mono- as well as poly-nuclear complexes.

2.3.2.5 Number of Components (NC)

Description:

The search field <u>Number of Components</u> contains the total number of components of a compound.

A number of components of 1 (nc=1) means a one-component compound. These are substances like the elements, molecules, salts, complexes, adducts, mixed crystals (in simple formula description), alloys, glasses and ceramic materials (in formula description).

e.g. Au, H2O, NaCl, Fe(C5H5)2, Na2SO4*10H2O, Nb0.8Ta0.2C, Fe0.55Pd0.45, SiO2, YBa2Cu3O(7-x)

A number of components greater than 1 (nc>1) means a multi-component compound. These are heterogeneous compounds like systems or solutions of two or more compounds. Additionally, some homogeneous compounds, like doped compounds, mixed crystals (in complex formula description), alloys, glasses and ceramic materials (in tabular description) belong (not exclusive) to this category.

Type of Indexing:

Numeric



Tips and Hints:

You can restrict the hit sets to one-component compounds, if nc=1 is combined with your query.

Examples:

Example multi- component compounds	potential (linearized) formula	Component 1	component 2
system iron – carbon	Fe#C	Fe	С
solution of sodium chloride in water	NaCl#H2O	NaCl	H ₂ O
tin doped In ₂ O ₃	In2O3#dotSn	In ₂ O ₃	Sn
mixed crystal of zirconia and titanium oxide	(Zr,Ti)O2 ZrO2#TiO2	ZrO ₂	TiO ₂
brass with 30% zinc	Cu(b),Zn(30)(X%)	Cu – 70%	Zn – 30%
ceramics of 8 wt% yttrium oxide and zirconia	ZrO2(b),Y2O3(8)(W%)	$ZrO_2 - 92$ wt%	$Y_2O_3 - 8$ wt%

Multi-component compounds have the fact "composition" showing information to each component.

2.3.2.6Number of Fragments (NF)

Description:

The search field <u>Number of Fragments</u> contains the total number of fragments of a compound.

A number of fragments of 1 (nf=1) means a component without fragmentation. These are substances like the elements, neutral molecules, neutral complexes (with known structure) and compounds without molecular units (e.g. solid-state compounds).

e.g. Au, H2O, Fe(C5H5)2, NaCl



For these compounds the number of components is 1.

A number of fragments greater than 1 (nf>1) means a multi-fragment compound. Compounds with more than one fragment are separated in the linearized formula by an asterisk. A one-component compound can consist of several fragments, e.g. as adducts, salts (of at least one ion with molecular unit)

Example multi-fragment compounds	number of fragments	number of components
FeCl3*6NH3	NF=2	NC=1
2Na(1+)*SO4(2-)=Na2SO4	NF=2	NC=1
Cu(2+)*SO4(2-)*5H2O=CuSO4*5H2O	NF=3	NC=1

All multi-component compounds are multi-fragment compounds, too. The number of fragments of multi-component compounds is the sum of the number of fragments of the single components.

Rules to determine whether a compound is fragmented or not are as follows

1. Solid state structures, ionic compounds:

 Substances are divided into formal fragments, if the compounds contain at least one discrete polyatomic ion or molecule in the crystal, the gas phase, the liquid phase, or in solution. The structures of these discrete ions or molecules are available.

e.g. 2Na(1+)*SO4(2-)=Na2SO4 (nf=2, nc=1)

- Substances are not divided into fragments, if the compounds consist exclusively
 of one-atomic ions (including OH(1-), SH(1-), SeH(1-), and TeH(1-)) or have no
 discrete polyatomic ion or molecule.
 e.g. NaCl, KOH, CaTiO3 (nf=1, nc=1)
- 2. Organometallic and coordination compounds:
 - Organometallic compounds and coordination compounds with known manner of ligand coordination are stored with the complete formula and structure.
 e.g. Fe(C5H5)2, [Co(NH3)3Cl3] (nf=1, nc=1)
 - Organometallic salts and ionic coordination compounds with known manner of ligand coordination are divided into cations and anions. The complete structures of the ions are available.
 e.g. [Pd(CH3CN)4](2+)*2BF4(1-)=[Pd(CH3CN)4](BF4)2 (nf=2, nc=1)



Organometallic compounds and coordination compounds with unknown manner of ligand coordination are divided into fragments by separating all ligands from the metal center.
 e.g. If there is an unknown coordination of thiocyanate to Co(3+) in [Co(NH3)5(SCN)]Cl2 (coordination by N or S atom possible), the formula is given as Co(3+)*5NH3*SCN(1-)*2Cl(1-)=[Co(NH3)5(SCN)]Cl2; (nf=4, nc=1).

The number of fragments can be determined only for compounds in formula description. Compounds given in percentage composition get nf=0.

Tips and Hints:

You can restrict the hit sets to one fragment, if nf=1 is combined with your query.

Type of Indexing:

Numeric

2.3.2.7 Number of Structures (NSTRUC)

Description:

The search field <u>Number of Structures</u> contains the total number of structures of a compound.

Example	Number of Structures	
elements, simple salts like NaCl, solid state compounds like CaTiO3, alloys, glasses, ceramic materials	NSTRUC=0	no structure available
molecules like H2O, C60, neutral complexes like Fe(C5H5)2	NSTRUC=1	
molecules, complexes, ionic compounds with two fragments like 2Na(1+)*SO4(2-), [Ru(py)2Cl2]*H2O, Fe(C5H5)2*C60	NSTRUC=2	
molecules, complexes, ionic compounds with three fragments like Cu(2+)*SO4(2-)*5H2O, Zn(2+)*TiF6(2-)*6H2O	NSTRUC=3	



The number of structures of compounds is the sum of the number of structures of the single fragments.

The number of structures of multi-component compounds is the sum of the number of structures of the single components.

Notes:

You should be aware that only molecular structures are available in the database.

Type of Indexing:

Numeric

2.3.2.8 Molecular Weight (MW)

Description:

The search field <u>Molecular Weight</u> contains the molecular weight of a substance. The molecular weight is defined as the sum of the relative atomic masses of all atoms in a compound. It is related to single compounds.

The molecular weight of compounds with more than one component, and compounds with undefined indices in the molecular formula, is indexed as zero.

Tips and Hints:

Searches with exact values requires the input of the digits behind the point, e.g. 19.9994 (D2O). A molecular weight of exact 19 are compounds with isotopes only, e.g. (19)F, (11)B(2)H4(1-), (7)Li(6)Li2, (4)He4(3)He.

Type of Indexing:

Numeric

	Search for	Field Name	Value
(a)	compounds with molecular weight between 186.01 – 186.15	MW	186.05-186.15
(b)	compounds with molecular weight lower 100	MW	<100



(c)	complex with molecular weight		
	greater 5000	MW	>5000

2.3.2.9 CAS Registry Number (RN)

Description:

The search field Registry Number contains the CAS accession number in the Registry file.

Notes:

The CAS Registry Number present in the Gmelin database has been mainly extracted from the Gmelin handbook. There are only few CAS Registry Number available in primary literature that are available.

Type of Indexing:

String (phrase) with fixed format

	Search for	Field Name	Value
(a)	substance with CAS Registry Number 102-54-5	RN	000102-54-5
(b)	substance with CAS Registry Number 77321-08-5	RN	077321-08-5
(c)	substance with CAS Registry Number 126543-78-0	RN	126543-78-0



2.3.2.10 Chemical Name (CN)

Description:

The search field <u>Chemical Name</u> contains the chemical names of the substances from the original literature.

Compounds can have multiple entries for chemical names such as systematic, trivial, and trade names. Additionally incomplete names are available describing only a part of the compound (e.g. the ligand) or steric information.

Notes:

In the GMELIN database not all compounds have a name. The names given do not necessarily conform to the IUPAC nomenclature.

Tips and Hints:

Name searches are not recommended to identify compounds, because names are ambiguous or non-systematic and are not available in many cases. In some cases name fragments might be useful to search for substance classes or complex compounds with same ligand. However there are more powerful methods available such as structure, substructure searches, or formula searches.

Type of Indexing:

String (phrase)

Related Information:

CNS

	Search for	Field Name	Value
(a)	ferrocene	CN	ferrocene
(b)	Alum	CN	alum
(c)	adduct of (r)-binaphthol with tin tetrachloride	CN	(r)-binaphthol*tin tetrachloride



2.3.2.11 Chemical Name Segment (CNS)

Description:

The search field <u>Chemical Name Segment</u> uses the same source as the field Chemical Name. However, it is indexed differently allowing fast searching for segments of chemical names without entering truncation symbols.

In some cases name fragments might be useful to search for substance classes or complex compounds with the same ligands.

Type of Indexing:

String (wordwise)

Related Information:

CN

Example1:

	Search for	Field Name	Value
(a)	compounds containing the name term ferrocene	CNS	ferrocene
(b)	compounds containing the name term hexacyanoferrate	CNS	hexacyanoferrate
(c)	compounds containing the name term binaphthol	CNS	binaphthol

Example2:

Search for compounds containing the name terms trans, chloro and iron

Operator	Field Name	Value
	CNS	trans
AND	CNS	chloro
AND	CNS	iron



2.3.2.12 Type of Substance (STYPE)

Description:

The search field <u>Type of Substance</u> contains the chemical classification of the substances.

Following classes are searchable:

- Alloy
- Coordination Compound
- Dopant
- Glass or Ceramic Material
- Isomorphous or Diadochous Compound, Solid Solution
- Isotope or Isotope containing Compound
- Mineral (assigned to natural minerals only)
- Polymer

Type of Indexing:

String (phrase)

2.3.2.13 Modification (MODTXT)

Description:

The search field <u>Modification</u> contains the modification of a substance. In many cases (e.g. minerals) the modification is identical to the chemical name.

Modifications can be:

Mineral name	e.g. calcite
Color	e.g. yellow
Crystal system	e.g. orthorhombic
Bravais type	e.g. face centered cubic
Space group (Hermann-Mauguin symbol)	e.g. p21/c
Crystal structure type	e.g. rutile
Numbers and Greek letters	e.galpha.
Qualitative temperature and pressure references	e.g. high temperature
ortho / para	



singlet / triplet

Notes:

A modification can consist of several designations, which are separated by a comma. In multi-component systems the modification is assigned to the single component.

Tips and Hints:

The description of modifications is ambiguous or non-systematic in literature. Therefore it is possible that the same modification is recorded by several designations or combinations.

Type of Indexing:

String (phrase)

2.3.2.14 Entry Date (ED)

Description:

The search field <u>Entry Date</u> contains the date (year/month/day) when this compound was first entered into the database. Dates have to be searched with a fixed format: yyyy/mm/dd.

Type of Indexing:

String (phrase)

2.3.2.15 Update Date (UPD)

Description:

The search field <u>Update Date</u> contains the date (year/month/day) when the last update of this compound was entered into the database. Dates have to be searched with a fixed format: yyyy/mm/dd.

Type of Indexing:

String (phrase)



2.3.3 Formula Search Data

2.3.3.1 Formula Search Data (FOR)

Number of Atoms (NA) Number of Elements (NE) Element Symbols (ELS) Element Count (ELC)

2.3.3.2 Number of Atoms (NA)

Description:

The numeric search field <u>Number of Atoms</u> contains the total number of atoms in a molecular formula MF. Thus NA will be multiple for multi-component compounds.

Note:

NA is a numeric value that can be searched for with exact values or open and close ranges.

Compounds with a negative number of atoms (NA = -1) have components with undefined stoichiometric indexes, like YBa2Cu3O(7-x), or adducts with an undefined prefix factor, like CuBr2*99H2O.

Tips and Hints:

The Number of Atoms must not be confused with the Number of Elements (NE) or the Element Count (ELC).

This search field is useful especially in combination with other search fields of this topic, especially for searching simple inorganic or solid-state compounds.

Type of Indexing:

Numeric

Search for	Field Name	Value
compounds with 190 to 200 atoms	NA	190-200



2.3.3.3 Number of Elements (NE)

Description:

The numeric search field <u>Number of Elements</u> contains the number of different elements present in a molecular formula MF. Thus NE will be multiple for multi-component compounds.

Isotopes, including deuterium and tritium, are not treated as different elements.

Note:

NE is a numeric value that can be searched for with exact values or open and close ranges.

Type of Indexing:

Numeric

Tips and Hints:

The Number of Elements must not be confused with the Element Count (ELC) or the Number of Atoms (NA).

This search field is useful especially in combination with other search fields of this topic, especially for searching simple inorganic or solid-state compounds.

Example:

Search for	Field Name	Value
compounds more than ten elements	NE	>10

2.3.3.4 Element Symbol (ELS)

Description:

The search field <u>Element Symbol</u> contains the symbol of each distinct element in the molecular formula.

Isotopes, including deuterium and tritium, are not treated as different elements.



Note:

Elements with atomic number greater 103 are available by the element symbols recommended by IUPAC (e.g. element #104 as Unq).

The term Qx represents positive charged compounds, the term Qy represents negative charged compounds.

Tips and Hints:

The Element Symbol must not be confused with the Element Count (ELC), the Number of Elements (NE) or the Number of Atoms (NA).

This search field is useful especially in combination with other search fields of this topic, especially for searching simple inorganic or solid-state compounds.

Type of Indexing:

String (phrase)

Examples:

Search for zirconium and zirconium compounds

Operator	Field Name	Value
	ELS	Zr

Search for phosphorus nitride compounds (of any stoichiometry)

Operator	Field Name	Value
	ELS	Р
AND	ELS	Ν
AND	NE	2
AND	NC	1

The hit set contains phosphorus nitride compounds without further elements. Search for barium copper oxides with three up to four atoms



Operator	Field Name	Value	
	ELS	Ва	
AND	ELS	Cu	
AND	ELS	0	
AND	NA	3-4	
AND	NC	1	

The hit set contains barium copper oxides with various oxygen content and barium copper oxides with one further element as well as mixed crystals with further elements.

2.3.3.5 Element Count (ELC)

Description:

The numeric search field <u>Element Count</u> contains the respective counts of the distinct elements in a component molecular formula MF.

Isotopes, including deuterium and tritium, are not treated as different elements.

Note:

An element count is defined as the element symbol and the number of its occurrences per molecule.

Odd-numbered values result from compounds like mixed crystals, alloys and glasses or ceramic materials.

Negative element counts indicate compounds with the respective element in undefined stoichiometry.

The element count is fixed to two decimal places, even if the compound has more.

Tips and Hints:

The Element Symbol must not be confused with the Element Symbol (ELS), the Number of Elements (NE) or the Number of Atoms (NA).

Element counts can be used to restrict searches on the basis of number of elements. It is possible to exclude elements by ELC search combinations using the Boolean Operator NOT.



This search field is useful especially in combination with other search fields of this topic as well as with structure fragments to search for complexes being fragmented.

Type of Indexing:

String (phrase)

Examples:

Search for compounds containing three ruthenium atoms

Operator	Field Name	Value
	ELC	Ru3.00

Search for barium copper oxides with two Ba and three copper atoms per formula

Operator	Field Name	Value
	ELC	Ba2.00
AND	ELC	Cu3.00
AND	ELS	0
AND	NC	1

The hit set contains compounds like ABa2Cu3(O,X), where A and/or X can be further element(s) or not available.

Search for a ruthenium complex containing four sulfur atoms but no phosphorus with maximal five different elements



Operator	Field Name	Value	
	ELC	Ru1.00	
AND	ELC	S4.00	
NOT	ELS	Р	
AND	NE	5	
AND	NC	1	

The hit set contains a number of mononuclear ruthenium complexes, as well as, some polynuclear complexes. There are also fragmented complexes with unknown ligand coordination.

2.3.4 Composition

2.3.4.1 Composition (CMPS)

Percentage (PERC) Percentage Type (PERT)

2.3.4.2 Percentage (PERC)

Description:

The search field <u>percentage</u> contains the percentage values of alloys, glasses or ceramic materials. PERC is a numeric value that can be searched for with exact values or open and close ranges.

Note:

Constituents can be combined by search field MF using the proximity operator.

Tips and Hints:

Searches using the percentage field can be tailored individually because of the numeric percentage values and the separation from the percentage type.

Type of Indexing:

Numeric

Related Information:



ALLOY

Example:

Search for alloys with more than 70 weight% palladium and 2-10 weight% rhodium				
Operator Field Name Value		Value		
	MF	Pd		
PROXIMITY	PERC	>70		
AND	MF	Rh		
PROXIMITY	PERC	2-10		
AND	PERT	weight%		
The hit est is r	act restricted to the number	of constituents		

The hit set is not restricted to the number of constituents.

2.3.4.3 Percentage Type (PERT)

Description:

The search field <u>percentage type</u> contains the form of the percentage information of alloys, glasses or ceramic materials. This search field is useful in combination with the percentage search field.

The following types are available:

% indefinite percent

atom% atom or mol percent

volume% volume percent

weight% weight percent

Type of Indexing:

String (phrase)



2.3.5 Ligands around Metals

2.3.5.1 Ligands around Metals (LIG)

Ligand Molecular Formula of one Center (LIG.LIGMF) Ligand Formulas of one Center (LIG.LIGFO) Ligand Formula Count of one Center (LIG.LIGFC)

2.3.5.2 Ligand Molecular Formula of one Center (LIG.LIGMF)

Description:

The alphanumerical search field <u>Ligand Molecular Formula</u> of one Center contains the abstract formula of the single coordination centers and the corresponding ligand codes of a compound.

The ordering of the formula is as follows:

1. metal center

- 2. the ligand codes (see LIG.LIGFO) are ordered in the following manner:
 - A, CN, CNO, CNR, CNS, CO, CS, D, L, Q, X
 - within the same ligand types, they are ordered on ascending denticity and the frequency is not taken into consideration

The details to each different metal center are displayed separately in field Ligands around Metals, respectively.

Note:

Charges are not taken into account.

In contrast to the search field Ligand Molecular Formula only one center is described here even if the whole compound contains several metal centers. For a mononuclear complex both ligand molecular formulas are identical.

Tips and Hints:

Searches for compounds using the GMELIN Ligand Search System are very powerful when looking for a class of complexes with special connectivity.

Type of Indexing:

String (phrase)



Related Information:

LIGMF

Examples:

	Search for	Field Name	Value
(a)	hexa carbonyl chromium	LIG.LIGMF	Cr{CO} 6
(b)	tetra halogeno palladium compounds	LIG.LIGMF	Pd{X}4
(c)	nickel complex with two tridentate ligands connecting by one pnictide and two chalcogenide atoms	LIG.LIGMF	Ni{D(2)Q}2

The hit set contains mono- as well as poly-nuclear complexes.

2.3.5.3 Ligand Formulas of one Center (LIG.LIGFO)

Description:

The search field <u>Ligand Formula of one Center</u> contains the ligand codes according to the Gmelin Ligand Search System, which describes the elements (in groups) and the number of the atoms that are connected to one central metal atom. A ligand formula is given for all structured compounds or fragments which contain at least one metal atom and at least one coordinating ligand atom of the classes "L, A, D, Q, X" or a special ligand.

The codes include the following elements:

Code	Atom					
L	С					
А	В	Si	Ge			
D	Ν	Ρ	As	Sb		
Q	0	S	Se	Те		
Х	Н	F	CI	Br	Ι	At



There are the following "special" ligands:

Code CO CS CN CNS CNO CNR *) *) The substituent R has no further bond to metal atoms.

If a ligand is described by a combination of the codes, they are ordered alphabetically and the denticity of the code is put in parentheses in front of the code.

Note:

The ligand formula describes only the denticity of the ligand not the connectivity to the metal center.

In contrast to the search field Ligand Formula only one center is described here, even if the whole compound contains several metal centers. For a mononuclear complex both ligand formulas are identical.

Type of Indexing:

String (phrase)

Related Information:

LIGFO

	Search for	Field Name	Value
(a)	halogenid or hydride ligand μ-halogenide or μ-hydride ligand	LIG.LIGFO	х
(b)	ligand connecting by two halogenide or hydride atoms e.g. η²-hydrogen, η²-BH₄	LIG.LIGFO	(2)X
(c)	(monodentate) carbonyl	LIG.LIGFO	СО
(d)	ligand connecting by C and chalcogen e.g. μ-η²-carbonyl (by C and O)	LIG.LIGFO	LQ
(e)	ligand connecting by four pnictide atoms e.g. porphyrins, phthalocyanines, tetra-aza-cyclam	LIG.LIGFO	(4)D



(f)	ligand connecting by five C atoms to one or more metal centers e.g. η ⁵ -cyclopentadienyl	LIG.LIGFO	(5)L
(g)	ligand connecting to metal(s) by C e.g. μ -carbonyl (only by C), bridged alkyl-, η^2 alken-, η^2 -alkin ligands	LIG.LIGFO	(2)L

The hit set contains mono- as well as poly-nuclear complexes.

2.3.5.4 Ligand Formula Count of one Center (LIG.LIGFC)

Description:

In the field <u>Ligand Formula Count of one Center</u> the frequency of a certain ligand, relating to one metal center within a compound, is indexed. For ligand codes see LIG.LIGFO.

Note:

In contrast to the search field Ligand Formula Count only one center is described here, even if the whole compound contain several metal centers. For a mononuclear complex both ligand formula counts are identical.

Tips and Hints:

Different ligand formula counts can be combined with the "proximity" operator. This is a very powerful search code to tailor coordination compounds.

Type of Indexing:

String (phrase)

Related Information:

LIGFC

Example:

Search for complexes containing one η^5 -C ligand, one ligand connecting by two pnictide atoms and three halogenide (or hydride) ligands

Operator Field Name Value



	LIG.LIGFC	(5)L:1
PROXIMITY	LIG.LIGFC	(2)D:1
PROXIMITY	LIG.LIGFC	X:3

The hit set contains mono- as well as poly-nuclear complexes.



2.4 General Information

2.4.1 General Information (GEN)

Occurrence in Nature (OCN) Purification (PUR) Use of the Substance (USE) Handling (HDLG) Health Protection (HEP) Ecological Data (ECOL) Physiological Behavior (PHYB) History (HIST) Economic Data (ECON)

2.4.2 Occurence in Nature (OCN)

Description

This field identifier is a group code indicating the <u>Occurrence in Nature</u> of a compound, like the occurrence in plants, animals, in the atmosphere, hydrosphere, lithosphere or in the universe.

Type of Indexing:

Group Code

2.4.3 Purification (PUR)

Description:

This field identifier is a group code indicating the <u>Purification</u> of a compound. Here the isolation or enrichment of a compound from a mixture, as well as, general purification methods are considered. However, the "working up" of a reaction mixture is indexed in the Reaction Details Purification / Isolation (RX.ISOL).

Type of Indexing:



2.4.4 Use of the Substance (USE)

Description:

This field identifier is a group code indicating the <u>Use of the Substance</u>. Here the application of the compound in laboratory as well as in (chemical) industry is considered. The use of a substance as catalyst is not indexed here, but under the group code Chemical Behavior in field Behavior as Catalyst (CAT).

Type of Indexing:

Group Code

2.4.5 Handling (HDLG)

Description:

This field identifier is a group code indicating the <u>Handling</u>, storage or disposal of a compound. Here, aspects like handling during transport, special storage conditions or removal, or waste disposal, are considered.

Type of Indexing:

Group Code

2.4.6 Health Protection (HEP)

Description:

This field identifier is a group code indicating the protection of persons from the effects of a compound or the degree of toxicity of a compound. Here information about MAC values, lethal doses, prophylaxis or therapeutic procedures can be searched.

Type of Indexing:



2.4.7 Ecological Data (ECOL)

Description:

This field identifier is a group code indicating the effects of a compound on the environment.

Type of Indexing:

Group Code

2.4.8 Physiological Behavior (PHYB)

Description:

This field identifier is a group code indicating the <u>Physiological Behavior</u> of a compound including effects on men, animals, plants, microorganisms, or metabolism of a compound.

Type of Indexing:

Group Code

2.4.9 History (HIST)

Description:

This field identifier is a group code indicating the History of a compound.

Type of Indexing:

Group Code

2.4.10 Economic Data (ECON)

Description:

This field identifier is a group code indicating the Economic or commercial data of a compound.

Type of Indexing:



2.5 Physical Properties

2.5.1 Physical Properties (PHY)

Description:

The topic Physical Properties contains data for single-component systems. This field identifier is a group code for Field Availability searches in the hierarchical system of the CrossFire database.

Acceptance Criteria:

Data determined by experimental measurements in the abstracted article are indexed as values or corresponding keywords.

Values which have been calculated by classical (not quantum mechanical) methods or which come from experimentally measured data have also been accepted, e.g. rotational constants, enthalpy of vaporization, thermal expansion coefficient etc.

Note:

Data to multi-component systems are recorded in the topic Properties of / in Systems (MCS). Values, which have been calculated by quantum mechanical methods, are indicated in the topic Quantum Chemical Calculations (QCC).

The topics on physical data follow the hierarchy described below:

Molecular Properties (MOL) Information on Phase Transitions (PHT) Information on Condensed Phase (CRY) Mechanical Properties (MEC) Thermal Properties (THD) Transport Phenomena (TRA) Thermodynamic Data (TDD) Electric Properties (ELE) Magnetic Properties (MAG) Optical Properties (OPT) Spectroscopic Information (SPE)



2.5.2 Molecular Properties

2.5.2.1 Molecular Properties (MOL)

Conformation and Bonding Models (CONF) Symmetry / Point Group (PGR) Intramolecular Distances and Angles (IDA) Molecular Vibration (VIB) Molecular Rotation (MCR) Electronic States (EST) Bond Dissociation Energy (EDIS) Polarizability (ELP) Dipole Moment (DM) Ionization Potential (IP) Potential Energy Function (PEF)

2.5.2.2Conformation and Bonding Models (CONF)

Description:

The field Conformation and Bonding Models gives indications of conformation and bonding models or descriptions of molecular structure. The information indicated is more detailed than the structure shown in the database.

Type of Indexing:

Group Code

2.5.2.3 Symmetry / Point Group (PGR)

Description:

The field Symmetry / Point Group contains the Schoenflies symbol of the point group, which the molecule exhibits.



Note:

Avoid confusion of the symmetry point group of a molecule with its crystal space group recorded in the search field CST.SGROUP.

Type of Indexing:

String (phrase)

2.5.2.4 Intramolecular Distances and Angles (IDA)

Description:

This field identifier is a group code for Field Availability searches in the hierarchical system of the CrossFire database. The group code Intramolecular Distances and Angles gives indications of intramolecular distances, bond angles and dihedral angles of structure. You can search for the determination method using the associated field code IDA.MET

Type of Indexing:

Group Code

Associated Parameter Fields:

Field CodeFull NameType of IndexingIDA.METMethod of DeterminationString (phrase)

2.5.2.5 Method of Determination (IDA.MET)

Description:

Intramolecular Distances and Angles (IDA) gives indications of intramolecular distances, bond angles and dihedral angles of structure. You can search for the following determination methods:

- Electron Diffraction
- Empirical Calculations. (Phys. Properties)
- ESR Spectroscopy
- EXAFS Spectroscopy
- Fluorescence Spectroscopy



- Infrared Spectroscopy
- Microwave Spectroscopy
- Neutron Diffraction
- NMR Spectroscopy
- Powder X-Ray Diffraction
- Raman Spectroscopy
- Single Crystal X-Ray Diffraction
- Ultraviolet Spectroscopy
- X-Ray Diffraction
- X-Ray Spectroscopy

Type of Indexing:

String (phrase)

2.5.2.6 Molecular Vibration (VIB)

Description:

Indication of fundamental vibrations, force constants, anharmonicity constants and vibration-rotation-interaction constants is given in the fields VIB.FUND, VIB.FORCE, VIB.ANHRM and VIB.ROTA.

Note:

Searching for the special field codes requires the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
VIB.FORCE	Force Constants	String (phrase)
VIB.FUND	Fundamental Vibrations	String (phrase)
VIB.ANHRM	Anharmonicity Constants	String (phrase)
VIB.ROTA	Vibration-Rotation- Interaction Constants	String (phrase)



2.5.2.7 Molecular Rotation (MRC)

Description:

The Molecular Rotation constant is inversely proportional to the moment of inertia of a molecule rotating about a particular axis. The search field contains the rotation constants for the vibrational ground state in the electronic ground state (Ao, Bo, Co). Indication of centrifugal distortion or Coriolis coupling is given in the search field MRC.CENT.

Note:

Searching for centrifugal distortion require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
MRC.CONA	Rotational Constant A(0)	Numeric	Hz
MRC.CONB	Rotational Constant B(0)	Numeric	Hz
MRC.CONC	Rotational Constant C(0)	Numeric	Hz
MRC.CENT	Centrifugal Distortion	String (phrase)	

2.5.2.8 Electronic States (EST)

Description:

The field Electronic States gives indications of properties of electronic energy levels e.g. term values, lifetimes, fine or hyperfine structure.

Note:

Assignments of UV bands to transitions between electronic states are indicated in the UV.ASSIG search field.

Type of Indexing:



2.5.2.9 Bond Dissociation Energy (EDIS)

Description:

The field Bond Dissociation Energy is the amount of energy needed to break a single bond of one type in a molecule. The field Bond Dissociation Energy gives indications of bond energy in the electronic ground state.

Type of Indexing:

Group Code

2.5.2.10 Polarizability (ELP)

Description:

The electric Polarizability of a molecule is the induced electric dipole moment per unit field strength. If the polarizability of an ion is determined the whole compound containing the ion is stated by formula or chemical name. You can search for this associated information using the parameter field code ELP.COFO.

Unit:

The default unit in the CrossFire database for this fact is ml.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
ELP.COFO	Complete Formula	String (phrase)

Type of Indexing:

Numerical (range)

2.5.2.11 Dipole Moment (DM)

Description:

The dipole moment is the electric moment of the charge density in an asymmetrical molecule. You can search for the determination method using the associated field code DM.MET.



Unit:

The default unit in the Gmelin database for this fact is D. D = Debye = $3.33 \ 10-30 \ C \ m$

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing

DM.MET Method of Determination String (phrase)

Type of Indexing:

Numerical (range)

2.5.2.12 Method of Determination (DM.MET)

Description:

The dipole moment is the electric moment of the charge density in an asymmetrical molecule.

Following determination methods can be searched:

- Electric Constant, Permittivity
- Dielectric Relaxation
- IR Spectroscopy
- Microwave Spectroscopy
- Molecular Beam Electric Resonance

E.g the method dielectric constant means that the dipole moment value is calculated from the dielectric constant.

Type of Indexing:

String (phrase)



2.5.2.13 Ionization Potential (IP)

Description:

The Ionization Potential is the energy required to remove an electron from the neutral molecule bond infinity.

Please find valid determination methods in the field IP.MET and possible types of the ionization process in the field IP.TYP.

Generally, only the first ionization potential is recorded numerically. Further ionization potentials are indicated in the field IP.POT.

Indication of additional information on ion formation (like ionization cross section, electron affinity) is given in the field IP.IONFORM.

Unit:

The default unit in the CrossFire database for this fact is eV. $eV = 1.6021 \ 10^{-19}$ Joule

Note:

Searching for higher ionization potentials require the field value "available", formation of ions the field value "described", or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
IP.MET	Method of Determination	String (phrase)
IP.TYP	Kind of Ionization	String (phrase)
IP.POT	Higher Ionization Potentials	String (phrase)
IP.IONFORM	Formation of Ions	String (phrase)

Type of Indexing:

Numerical (range)



2.5.2.14 Method of Determination (IP.MET)

Description:

The Ionization Potential is the energy required to remove an electron from the neutral molecule bond infinity. Valid determination methods in the field IP.MET are as follows:

Electron Impact

- Mass Spectroscopy
- Photoelectron Spectroscopy
- Photoemission Yield Spectroscopy
- Photoionization

Type of Indexing:

String (phrase)

2.5.2.15 Kind of Ionization (IP.TYP)

Description:

The Ionization Potential is the energy required to remove an electron from the neutral molecule bond infinity.

Possible types of the ionization process in the field IP.TYP are:

- Adiabatic
- Vertical
- Not Given

Type of Indexing:

String (phrase)



2.5.2.16 Potential Energy Function (PEF)

Description

Indication of Potential Energy Function is given in this field. You can search for information given as diagram or equation using the associated parameter fields PEF.DGM respectively PEF.EQN.

Note:

Searching for potential energy diagrams or equations, requires the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
PEF.DGM	Diagram	String (phrase)
PEF.EQN	Equation	String (phrase)

2.5.3 Information on Phase Transition

2.5.3.1 Information on Phase Transition (PHT)

State Diagram (STAT) Triple Point (TP) Critical Point (CRPO) Transition Point Solid / Solid (TSS) Transition Enthalpy Solid / Solid (HTSS) Transition Entropy Solid / Solid (STSS) Transition Gibbs Energy Solid / Solid (GTSS) Melting Point (MP) Melting Enthalpy (HFUS) Melting Entropy (SFUS) Melting Gibbs Energy (GFUS) Melting Point (Further Information) (MPFI)



Sublimation Point (SP) Sublimation Enthalpy (HSUB) Sublimation Entropy (SSUB) Sublimation Gibbs Energy (GSUB) Sublimation Point (Further Information) (SPFI) Boiling Point (BP) Boiling Enthalpy (HVAP) Boiling Entropy (SVAP) Boiling Gibbs Energy (GVAP) Vapor Pressure (VP) Vapor Pressure (Further Information) (VPFI)

2.5.3.2 State Diagram (STAT)

Description:

The field State Diagram indicates a phase (state) diagram as a plot of pressure versus temperature for a certain substance (one component). Phase diagrams of multi-component systems are indicated in the search field META.

Type of Indexing:

Group Code

2.5.3.3 Triple Point (TP)

Description:

The Triple Point is the thermodynamic state at which three phases of a substance exist in equilibrium recorded in the fields temperature TP and pressure TP.P. The search field phases contains the description of the three phases existing at the triple point. You can search for this associated information using the parameter field code TP.PHAS.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
TP.PHAS	Phases	String (phrase)	
TP.P	Pressure	Numeric	Torr



TΡ

Temperature

Numeric

°C

Type of Indexing:

Group Code

2.5.3.4 Critical Point (CRPO)

Description:

The Critical Point CRT is the temperature above which a gas cannot be liquified, regardless of the pressure. The critical pressure CRP of a substance is the minimum pressure required for liquefaction at the critical temperature. The critical density CRD is the density exhibited by a substance at critical temperature and critical pressure. The critical volume CRPV (CRPO.MVOLA) of a substance is the molar (specific) volume at critical pressure and critical temperature.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
CRP	Pressure	Numeric	Torr
CRT	Temperature	Numeric	°C
CRD	Density	Numeric	g ml ⁻¹
CRV	Molar Volume	Numeric	ml mol ⁻¹
CRPO.MVOLA	Specific Volume	Numeric	l kg ⁻¹

Type of Indexing:

Group Code

2.5.3.5 Transition Point Solid / Solid (TSS)

Description:

The (crystal) Transition Point Solid / Solid is the thermodynamic state at which two crystalline modifications are in equilibrium. The TSS field contains the temperature at which a phase transition between two specified crystalline phases occurs.



It is linked to the associated pressure by parameter field codes TSS.P. The crystalline modifications are displayed in the field TSS.PHAS1 and TSS.PHAS2.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
TSS.PHAS1	Solid Phase 1	String (phrase)	
TSS.PHAS2	Solid Phase 2	String (phrase)	
TSS.P	Pressure	Numeric	Torr
TSS	Temperature	Numeric	°C

Type of Indexing:

Group Code

2.5.3.6 Transition Enthalpy Solid / Solid (HTSS)

Description:

The (specific) enthapy of phase transition is the change in enthalpy for conversion of one mole (gram) of a substance from one solid phase 1 into phase 2 without change in temperature. The transition enthalpy is linked to the associated temperature and pressure by parameter field codes HTSS.T and HTSS.P. The crystalline modifications are displayed in the field HTSS.PHAS1 and HTSS.PHAS2.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
HTSS.PHAS1	Solid Phase 1	String (phrase)	
HTSS.PHAS2	Solid Phase 2	String (phrase)	
HTSS.T	Temperature	Numeric	°C
HTSS.P	Pressure	Numeric	Torr
HTSS	Molar Transition Enthalpy	Numeric	J mol⁻¹
HTSS.ENTHA	Specific Transition Enthalpy	Numeric	J g⁻¹



Type of Indexing:

Group Code

2.5.3.7 Transition Entropy Solid / Solid (STSS)

Description:

The (specific) entropy of phase transition is the change in entropy for conversion of one mole (gram) of a solid phase 1 into a different phase 2 without change in temperature. The transition entropy is linked to the associated temperature and pressure by parameter field codes STSS.T and STSS.P. The crystalline modifications are displayed in the field STSS.PHAS1 and STSS.PHAS2.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
STSS.PHAS1	Solid Phase 1	String (phrase)	
STSS.PHAS2	Solid Phase 2	String (phrase)	
STSS.P	Pressure	Numeric	Torr
STSS.T	Temperature	Numeric	°C
STSS	Molar Transition Entropy	Numeric	J mol ⁻¹ K ⁻¹
STSS.ENTRA	Specific Transition Entropy	Numeric	J g ⁻¹ K ⁻¹

Type of Indexing:

Group Code

2.5.3.8 Transition Gibbs Energy Solid / Solid (GTSS)

Description:

The (specific) Gibbs energy of phase transition is the change in Gibbs energy for conversion of one mole (gram) of a solid phase 1 into a different phase 2 without change in temperature. The transition Gibbs enthalpy is linked to the associated temperature and pressure by parameter field codes GTSS.T and GTSS.P. The crystalline modifications are displayed in the field GTSS.PHAS1 and GTSS.PHAS2.



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
GTSS.PHAS1	Solid Phase 1	String (phrase)	
GTSS.PHAS2	Solid Phase 2	String (phrase)	
GTSS.P	Pressure	Numeric	Torr
GTSS.T	Temperature	Numeric	°C
GTSS	Molar Gibbs Energy of Transition	Numeric	J mol⁻¹
GTSS.FRENA	Specific Gibbs Energy of Transition	Numeric	J g⁻¹

Type of Indexing:

Group Code

2.5.3.9 Melting Point (MP)

Description:

The Melting Point or freezing point of a substance is the temperature at which the solid and liquid phases are in thermodynamic equilibrium. The MP search field is linked to the associated pressure by parameter field code MP.P and to the solvent, from which the substance was crystallized, by parameter field code MP.SOL. Decomposition of the substance during melting is indicated in associated parameter field MP.DEC.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
MP	Temperature	Numeric (range)	°C
MP.P	Pressure	Numeric (range)	Torr
MP.DEC	Decomposition	String (phrase)	
MP.SOL	Solvent	String (phrase)	

Type of Indexing:



2.5.3.10 Melting Enthalpy (HFUS)

Description:

The (specific) enthalpy of fusion is the change in enthalpy when one mole (gram) of a solid substance is converted into liquid without a change in temperature. The Melting Enthalpy is linked to the associated temperature and pressure by parameter field codes HFUS.T and HFUS.P and to the solvent from which the substance was crystallized by parameter field code HFUS.SOL.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
HFUS.T	Temperature	Numeric	°C
HFUS.P	Pressure	Numeric	Torr
HFUS.SOL	Solvent	String (phrase)	
HFUS	Molar Enthalpy of Melting	Numeric	J mol⁻¹
HFUS.ENTHA	Specific Enthalpy of Melting	Numeric	J g⁻¹

Type of Indexing:

Group Code

2.5.3.11 Melting Entropy (SFUS)

Description:

The (specific) entropy of fusion is the change in entropy for the transition of one mole (gram) of solid substance into liquid at constant temperature. The Melting Entropy is linked to the associated temperature and pressure by parameter field codes SFUS.T and SFUS.P and to the solvent from which the substance was crystallized by parameter field code SFUS.SOL.



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
SFUS.T	Temperature	Numeric	°C
SFUS.P	Pressure	Numeric	Torr
SFUS.SOL	Solvent	String (phrase)	
SFUS	Molar Entropy Of Melting	Numeric	J mol ⁻¹ K ⁻¹
SFUS.ENTRA	Specific Entropy Of Melting	Numeric	J g⁻¹ K⁻¹

Type of Indexing:

Group Code

2.5.3.12 Melting Gibbs Energy (GFUS)

Description:

The (specific) Gibbs energy of fusion is the change in Gibbs energy when one mole (gram) of a solid substance is converted into liquid without change in temperature. The Melting Gibbs Energy is linked to the associated temperature and pressure by parameter field codes GFUS.T and GFUS.P and to the solvent from which the substance was crystallized by parameter field code GFUS.SOL.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
GFUS.T	Temperature	Numeric	°C
GFUS.P	Pressure	Numeric	Torr
GFUS.SOL	Solvent	String (phrase)	
GFUS	Molar Gibbs Energy of Melting	Numeric	J mol ⁻¹
GFUS.FRENA	Specific Gibbs Energy of Melting	Numeric	J g⁻¹

Type of Indexing:



2.5.3.13 Melting Point (Further Information) (MPFI)

Description:

The search field Melting Point (Further Information) indicates if further information about the melting point is given as diagram, table, or equation.

Note:

Searching for the special field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
MPFI.DGM	Diagram	String (phrase)
MPFI.TAB	Table	String (phrase)
MPFI.EQN	Equation	String (phrase)

Type of Indexing:

Group Code

2.5.3.14 Sublimation Point (SP)

Description:

The Point of Sublimation is the temperature at which the vapor pressure of a solid substance is equal to the external pressure. The SP search field is linked to the associated pressure by parameter field code SP.P and to the solvent from which the substance was crystallized by parameter field code SP.SOL.



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
SP	Temperature	Numeric	°C
SP.P	Pressure	Numeric	Torr
SP.SOL	Solvent	String (phrase)	

Type of Indexing:

Group Code

2.5.3.15 Sublimation Enthalpy (HSUB)

Description:

The (specific) enthalpy of sublimation is the change in enthalpy for conversion of one mole (gram) of a solid substance into vapor without change in temperature. The Sublimation Enthalpy is linked to the associated temperature and pressure by parameter field codes HSUB.T and HSUB.P and to the solvent from which the substance was crystallized by parameter field code HSUB.SOL.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
HSUB.T	Temperature	Numeric	°C
HSUB.P	Pressure	Numeric	Torr
HSUB.SOL	Solvent	String (phrase)	
HSUB	Molar Sublimation Enthalpy	Numeric	J mol⁻¹
HSUB.ENTHA	Specific Sublimation Enthalpy	Numeric	J g⁻¹

Type of Indexing:



2.5.3.16 Sublimation Entropy (SSUB)

Description:

The (specific) entropy of sublimation is the change of entropy for the transition of one mole (gram) of a solid substance to saturated vapor at constant temperature and under equilibrium conditions. The Sublimation Entropy is linked to the associated temperature and pressure by parameter field codes SSUB.T, SSUB.P and to the solvent from which the substance was crystallized by parameter field code SSUB.SOL.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
SSUB.T	Temperature	Numeric	°C
SSUB.P	Pressure	Numeric	Torr
SSUB.SOL	Solvent	String (phrase)	
SSUB	Molar Sublimation Entropy	Numeric	J mol ⁻¹ K ⁻¹
SSUB.ENTRA	Specific Sublimation Entropy	Numeric	J g⁻¹ K ⁻¹

Type of Indexing:

Group Code

2.5.3.17 Sublimation Gibbs Energy (GSUB)

Description:

The (specific) Gibbs energy of sublimation is the change in Gibbs energy for conversion of one mole (gram) of a solid substance into vapor without change in temperature. The Sublimation Gibbs Energy is linked to the associated temperature and pressure by parameter field codes GSUB.T, GS UB.P and to the solvent from which the substance was crystallized by parameter field code GSUB.SOL.



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
GSUB.T	Temperature	Numeric	°C
GSUB.P	Pressure	Numeric	Torr
GSUB.SOL	Solvent	String (phrase)	
GSUB	Molar Gibbs Energy of Sublimation	Numeric	J mol⁻¹
GSUB.FRENA	Specific Gibbs Energy of Sublimation	Numeric	J g⁻¹

Type of Indexing:

Group Code

2.5.3.18 Sublimation Point (Further Information) (SPFI)

Description:

The search field Sublimation Point (Further Information) indicates if further information about the melting point is given as diagram, table, or equation.

Note:

Searching for the special field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
SPFI.DGM	Diagram	String (phrase)
SPFI.TAB	Table	String (phrase)
SPFI.EQN	Equation	String (phrase)

Type of Indexing:

Group Code

2.5.3.19 Boiling Point (BP)



Description:

The Boiling Point is the temperature at which the vapor pressure of a liquid equals the external pressure. The normal boiling point is the temperature at which the vapor pressure equals the normal atmospheric pressure. The BP search field is linked to the associated pressure by parameter field code BP.P. Decomposition of the substance during boiling is indicated in associated parameter field BP.DEC.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
BP	Temperature	Numeric (range)	°C
BP.P	Pressure	Numeric (range)	Torr
BP.DEC	Decomposition	String (phrase)	

Type of Indexing:

Group Code

2.5.3.20 Boiling Enthalpy (HVAP)

Description:

The (specific) enthalpy of vaporization is the enthalpy change required to convert one mole (gram) of liquid substance into saturated vapor at constant temperature. The Boiling Enthalpy is linked to the associated temperature and pressure by parameter field codes HVAP.T and HVAP.P.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
HVAP.T	Temperature	Numeric	°C
HVAP.P	Pressure	Numeric	Torr
HVAP	Molar Vaporization Enthalpy	Numeric	J mol⁻¹
HVAP.ENTHA	Specific Vaporization Enthalpy	Numeric	J g⁻¹

Type of Indexing:



2.5.3.21 Boiling Entropy (SVAP)

Description:

The (specific) entropy of vaporization is the change in entropy for the transition of one mole (gram) of liquid substance into vapor at constant temperature and under equilibrium conditions. The Boiling Entropy is linked to the associated temperature and pressure by parameter field codes SVAP.T and SVAP.P.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
SVAP.T	Temperature	Numeric	°C
SVAP.P	Pressure	Numeric	Torr
SVAP	Molar Vaporization Entropy	Numeric	J mol ⁻¹ K ⁻¹
SVAP.ENTRA	Specific Vaporization Entropy	Numeric	J g ⁻¹ K ⁻¹

Type of Indexing:

Group Code

2.5.3.22 Boiling Gibbs Energy (GVAP)

Description:

The (specific) Gibbs energy of vaporization is the change in Gibbs energy when one mole (gram) of a liquid substance is converted into vapor. The Boiling Gibbs Energy is linked to the associated temperature and pressure by parameter field codes SVAP.T and SVAP.P.



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
GVAP.T	Temperature	Numeric	°C
GVAP.P	Pressure	Numeric	Torr
GVAP	Molar Gibbs Energy of Vaporization	Numeric	J mol⁻¹
GVAP.FRENA	Specific Gibbs Energy of Vaporization	Numeric	J g⁻¹

Type of Indexing:

Group Code

2.5.3.23 Vapor Pressure (VP)

Description:

The Vapor Pressure of a pure solid or liquid substance is the pressure exerted by its vapor under equilibrium conditions.

You can search for the associated temperature using the parameter field code VP.T. The vapor pressure is linked to the forms of state of the substance by parameter field code VP.S.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
VP	Temperature	Numeric	°C
VP.T	Pressure	Numeric	Torr
VP.S	Forms of State	String (phrase)	

Type of Indexing:



2.5.3.24 Vapor Pressure (Further Information) (VPFI)

Description:

The search fields Vapor Pressure (Further Information) indicates if information about vapor pressure is given as a diagram, table or equation. This field is linked to the forms of state of the substance by parameter field code VPFI.S.

Note:

Searching for the special field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
VPFI.DGM	Diagram	String (phrase)
VPFI.EQN	Table	String (phrase)
VPFI.TAB	Equation	String (phrase)
VPFI.S	Forms of State	String (phrase)

Type of Indexing:



2.5.4 Information on Condensed Phase

2.5.4.1 Information on Condensed Phase (CRY)

Color / Habit (COHA) Crystal Growth (CGRO) Crystal Bonding (CBND) Crystal Defects (CDEF) Superlattice (SLAT) Crystal Energy (CEN) Crystal Structure (CST)

2.5.4.2 Color / Habit (COHA)

Description:

The search field contains general information on color and habit of a substance in the condensed phase. Information in the field COHA.NOTAT refers to the color of the substance in the corresponding solvent. Habits can be searched in the field COHA.HABIT

The crystal habit may be strongly dependent upon the conditions of crystallization.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
COHA.COLOR	Color	String (phrase)
COHA.NOTAT	in Solvent	String (phrase)
COHA.HABIT	Habit	String (phrase)

Type of Indexing:



2.5.4.3 Habit (COHA.HABIT)

Description:

The search field contains information on habit of a substance in the condensed phase. The following habits can be searched in the field COHA.HABIT:

- bipyramids
- blocks
- bricks
- columns
- cubes
- dodecahedrons
- fibres
- flakes
- leaves
- needles
- octahedrons
- parallelepipeds
- parallelograms
- pinacoids
- plates
- prisms
- pyramids
- rhombododecahedrons
- rhombohedrons
- rods
- skalenohedrons
- slabs
- tetrahedrons

Type of Indexing:

String (phrase)

2.5.4.4 Crystal Growth (CGRO)

Description:

Indication of crystallization mechanisms or preparation of single crystals, e.g. by the application of various methods, is given in the field CGRO.CRYS. Information on the growth of crystals, e.g. twin formation, or epitaxy is indicated in the field CGRO.FORM.



Note:

Searching for the special field codes requires the field value "described" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
CGRO.CRYS	Crystallization / Preparation. of Single Crystals	String (phrase)
CGRO.FORM	Growth Forms / Epitaxy	String (phrase)

Type of Indexing:

Group Code

2.5.4.5 Crystal Bonding (CBND)

Description:

The field Crystal Bonding gives indications on bond states, bond types or crystal structure of crystalline compounds, e.g. layer structures.

Type of Indexing:

Group Code

2.5.4.6 Crystal Defects (CDEF)

Description:

Indication on crystal defects; e.g. color centers, line defects, displacements, disorders, twinning, vacancies or surface defects, is given in this field.

Type of Indexing:



2.5.4.7 Superlattice (SLAT)

Description:

The search field Superlattice indicates a solid system in which there exists a regular array of atoms of one species interspersed among the lattice structure of another species related by their crystal symmetry.

Type of Indexing:

Group Code

2.5.4.8 Crystal Energy (CEN)

Description:

The lattice energy is the energy associated with the construction of a crystal lattice relative to the energy of all constituent atoms separated by infinite distances. A characteristic term of the lattice energy is the Madelung constant determined solely by the geometry of an ionic crystal. The field Crystal Energy gives indications on lattice energy as well as Madelung constants.

Type of Indexing:

Group Code

2.5.4.9 Crystal Structure (CST)

Description:

The Crystal Structure type of a solid is the name of an isostructural prototype. The crystal type search field CST. TYP contains terms describing the crystal structure types for the substances in the literature.

According to the translational lattice, crystals may be divided into seven crystal systems and 14 Bravais types. The crystal system search field CST.SYST contains the crystal system names for the substances in the literature. The Bravais types are given in the Bravais type search field CST.BRAV.

The unit cell of a crystal lattice requires three vectors for its description. The set of the three axes for the lattice length (a, b, c) is given in CST.LENA(B, C) and the three lattice angles (alpha, beta, gamma) are given in CST.ANGA(B; C). The measuring temperature is given in CRYST.T and the molecules per unit cell in CST.ZVAL.



Methods of determination can be searched with the search field code CST.MET.

The symmetry group of a three-dimensional crystal pattern is called its space group. There are 230 crystallographic space groups. They are classified in 7 crystal systems (see CST.SYST) and 32 crystallographic point groups. The crystal space group search field CST.SGROUP contains the space group for a given substance in the literature.

Note:

Avoid confusion of the crystallographic space group with the point group of a molecule recorded in the search field PGR.

Indication on atomic coordinates, distances, angles or interplanar spacings is given in the field CST.ATPOS.

Searching for the field CST.ATPOS require the field value "available" or more general "*".

Field Code	Full Name	Type of Indexing	Unit
CST.TYP	Crystal Type	String (phrase)	
CST.SYST	Crystal System	String (phrase)	
CST.BRAV	Bravais Type	String (phrase)	
CST.LENA	Lattice Length A	Numeric	Angstroem
CST.LENB	Lattice Length B	Numeric	Angstroem
CST.LENC	Lattice Length C	Numeric	Angstroem
CST.ANGA	Lattice Angle .Alpha	Numeric	
CST.ANGB	Lattice Angle .Beta	Numeric	
CST.ANGC	Lattice Angle .Gamma	Numeric	
CST.T	Temperature	Numeric	°C
CST.MET	Method of Determination	String (phrase)	
CST.ZVAL	Molecules per Unit Cell	Numeric	dimensionless
CST.DEN	X-Ray Density	Numeric	g ml ⁻¹
CST.MVOL	Molar Volume	Numeric	m /mol ⁻¹
CST.MVOLA	Specific Volume	Numeric	l kg ⁻¹
CST.SGROUP	Space Group(s)	Numeric	
CST.ATPOS	Atomic Positions	String (phrase)	

Associated Parameter Fields:



Type of Indexing:

Group Code

2.5.4.10 Crystal Type (CST.TYP)

Description:

The Crystal Type search field CST.TYP contains terms describing the crystal structure types for the substances in the literature.The following terms are valid:

Antifluorite, Aragonite (CaCO₃), Calcite (CaCO₃), CdCl2,Cdl2 (Pbl2), Cooperite (PtS), Corundum (Al₂O₃), CsCl, Cuprite (Cu₂O), Cu, Diamond, Fluorite (CaF₂), Graphite, Ilmenite (FeTiO₃), Inverted Spinel (FeIII)(MgFe(III)O₄), Mg, NaCl, NiAs, PbO, Perowskite (CaTiO₃), Pyrite (FeS₂), Rutile (TiO₂), Silicates, Skutterudite (CoAs₃), Sphalerite, Spinel (MgAl₂O₄), Wurtzite (ZnS), W, Zinc blende (ZnS)

Type of Indexing:

String (Phrase)

2.5.4.11 Crystal System (CST.SYST)

Description:

According to the translational lattice, crystals may be divided into seven crystal systems and 14 Bravais types. The crystal system search field CST.SYST contains the crystal system names for the substances in the literature.

The following names are given:

- cubic
- hexagonal
- trigonal (including rhombohedral)
- tetragonal
- rhombic (=orthorhombic)
- monoclinic
- triclinic

Type of Indexing:

String (Phrase)



2.5.4.12 Bravais Type (CST.BRAV)

Description:

According to the translational lattice, crystals may be divided into seven crystal systems and 14 Bravais types.

The following Bravais types are given:

- all-faces-centered orthorhombic
- body-centered cubic
- body-centered monoclinic
- body-centered orthorhombic
- body-centered tetragonal
- face-centered cubic
- one-face-centered orthorhombic
- simple cubic
- simple hexagonal
- simple monoclinic
- simple orthorhombic
- simple rhombohedral
- simple tetragonal
 simple triclinic
- simple triclinic

Type of Indexing:

String (Phrase)

2.5.4.13 Method of Determination (CST.MET)

Description:

The following methods can be searched:

- electron diffraction
- neutron diffraction
- powder X-ray diffraction
- single crystal X-ray diffraction
- X-ray diffraction

Type of Indexing:

String (Phrase)



2.5.5 Mechanical Properties

2.5.5.1 Mechanical Properties (MEC)

Mole Volume (VOL) Elastic Behavior (ELAS) Compressibility (COMP) Plastic Behavior (PLAS) Strength (STRE) Hardness (HARD) Density (DEN)

2.5.5.2 Mole Volume (VOL)

Description:

Indication of molar volume, molecular volume, atomic volume or ionic volume is given in the field Mole Volume. This field is linked to the forms of state of the substance by parameter field code VOL.S.

Associated Parameter Field:

Field Code	Full Name	Type of Indexing
VOL.S	Forms of State	String (phrase)

Type of Indexing:

Group Code

2.5.5.3 Elastic Behavior (ELAS)

Description:

The elasticity of a substance is the response to stress, indicating the degree to which strain disappears from the substance when the stress has been removed. The field Elastic Behavior indicates information on elastic behavior of a substance, e.g. elasticity coefficients, elastic modulus, torsional rigidity or shearing force.



Type of Indexing:

Group Code

2.5.5.4 Compressibility (COMP)

Description:

The compressibility of a substance is the extent to which a material reduces its volume when it is subjected to compressive stresses. The field Compressibility indicates information on compressible behavior of a substance. This field is linked to the forms of state of the substance by parameter field code COMP.S.

Associated Parameter Field:

Field Code	Full Name	Type of Indexing
COM.S	Forms of State	String (phrase)

Type of Indexing:

Group Code

2.5.5.5 Plastic Behavior (PLAS)

Description:

The plasticity of a substance is the property of a substance by which it undergoes plastic deformation when the applied stress exceeds a certain value, known as the yield value. The field Plastic Behavior indicates information on the plastic behavior of a substance, e.g. plastic limit, superplasticity, or creep behavior.

Type of Indexing:



2.5.5.6 Strength (STRE)

Description:

The strength of a substance is its resistance to a force producing or tending to produce permanent deformation, especially as described by its stress level at which it fractures or fails. The Strength search field is a numeric range searchable field and is linked to the associated temperature STRE.T and to the kind of strength STRE.TYP.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
STRE	Value of Strength	Numeric	N mm ⁻²
STRE.TYP	Kind of Strength	String (phrase)	
STRE.T	Temperature	Numeric	°C
STRE.MR	Remarks	Wordwise	

Type of Indexing:

Group Code

2.5.5.7 Kind of Strength (STRE.TYP)

Description:

The strength of a substance is its resistance to a force producing or tending to produce permanent deformation, especially as described by its stress level at which it fractures or fails.

Following strength types are valid:

- Bend Strenght
- Compression Strenght
- Creep Strenght
- Fatigue Strenght
- Fracture Strenght
- Impact Strenght
- Notch Impact Strenght
- Shear Strenght
- Tensile Strenght
- Torsion Strenght
- Impact Bending



Yield Point

Type of Indexing:

String (phrase)

2.5.5.8 Hardness (HARD)

Description:

The Hardness of a solid substance is its resistance to indentation, scratching, abrasion, or cutting. Based on different methods of measurement there are several scales of hardness numbers:

- Identation Hardness: Brinell-, Knoop-, Vickers-, Micro-, Rockwell Hardness
- Scratch Hardness: Mohs Hardness
- Scleroscope Hardness

Please find valid harness Types in HARD.TYP. Information about pretreatment of samples is given in the field remarks HARD.MR.

Note:

The displayed unit [N mm⁻²] is not valid for Mohs, Rockwell and Scleroscope hardness, because they have no dimension in the GMELIN database.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
HARD	Value of Hardness	Numeric	N mm ⁻²
HARD.VALU	Hardness without Unit		
HARD.TYP	Kind of Hardness	String (phrase)	
HARD.T	Temperature	Numeric	°C
HARD.MR	Remarks	Wordwise	

Type of Indexing:

Group Code

2.5.5.9 Kind of Hardness (HARD.TYP)

Description:



The Hardness of a solid substance is its resistance to indentation, scratching, abrasion, or cutting.

The following Hartness Types are valid:

- Brinell Hardness
- Knoop Hardness
- Microhardness
- Moh's Hardness
- Rockwell Hardness
- Scleroscope Hardness
- Vickers Hardness

Type of Indexing:

String (phrase)

2.5.5.10 Density (DEN)

Description:

The density of a substance is the mass per unit volume at defined temperature and pressure. The Density search field is a numeric range searchable field and is linked to the associated temperature DEN.T, the reference temperature DEN.RT, and to the pressure DEN.P. Information about pretreatment of samples is given in the field remarks DEN.MR. Further information about the density is given as a diagram or equation using the parameter field codes DEN.DGM respectively DEN.EQN. The field is linked to the forms of state of the substance by parameter field code DEN.S.

Note:

Searching for density diagram or equation field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
DEN	Densitiy	Numeric	g ml ⁻¹
DEN.T	Temperature	Numeric	°C
DEN.RT	Reference Temperature	Numeric	°C
DEN.P	Pressure	Numeric	Torr
DEN.MR	Remarks	Wordwise	



DEN.DGM Diagram

DEN.EQN Equation

DEN.S Forms of State

String (phrase) String (phrase) String (phrase)

Type of Indexing:



2.5.6 Thermal Properties

2.5.6.1 Thermal Properties (THD)

Thermal Expansion (TEC) Thermal Conductivity (TCND)

2.5.6.2 Thermal Expansion (TEC)

Description:

The Thermal Expansion coefficient is the ratio of the change in length per unit length or change in volume per unit volume to the change of temperature. The coefficient field is linked to the associated temperature TEC.T and to the kind of expansion TEC.LIN or TEC.CUB, as well as, to the forms of state of the substance by parameter field code TEC.S.

Note:

Searching for the kind of expansion requires the field value "yes" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
TEC	expansion coefficient	Numeric	K ⁻¹
TEC.LIN	linear expansion	String (phrase)	
TEC.CUB	cubic expansion	String (phrase)	
TEC.T	temperature	Numeric	°C
TEC.MR	remarks	Wordwise	
TEC.S	forms of state	String (phrase)	

Type of Indexing:



2.5.6.3 Thermal Conductivity (TCND)

Description:

The Thermal Conductivity of a substance is the time rate of heat transfer by conduction, across unit area, through unit thickness of substance, for unit difference of temperature. The coefficient field is linked to the associated temperature TCND.T and to the forms of state of the substance by parameter field code TCND.S.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
TCND	thermal conductivity	Numeric	cal m ⁻¹ s ⁻¹ K ⁻¹
TCND.T	temperature	Numeric	°C
TCND.MR	remarks	Wordwise	
TCND.S	forms of state	String (phrase)	

Type of Indexing:



2.5.7 Transport Phenomena

2.5.7.1 Transport Phenomena (TRA)

Dynamic Viscosity (DV) Kinematic Viscosity (KV) Non-Newtonian Behavior (NONT)

2.5.7.2 Dynamic Viscosity (DV)

Description:

The Dynamic Viscosity is the ratio of the shearing stress to the shear of the motion of a fluid, melt, or suspension. It is independent of the velocity distribution and the dimensions of the system. For a gas it is independent of pressure except at very low pressures.

The dynamic viscosity field contains the values for the dynamic viscosity of substances given at specified temperatures. You can search for this associated information using the parameter field code DV.T.

The search field remarks contain concentration details of solutions, as well as, information to further specialize conditions relating to the dynamic viscosity values. You can search for this associated information using the parameter field code DV.MR. The forms of state of the substance can be restricted by the associated parameter field code DV.S

Unit:

The default unit in the CrossFire database for this fact is P.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
DV	Dynamic Viscosity	Numeric	Ρ
DV.T	Temperature	Numeric	°C
DV.MR	Remarks	Wordwise	
DV.S	Forms of State	String (phrase)	



Type of Indexing:

Group Code

2.5.7.3 Kinematic Viscosity (KV)

Description:

The Kinematic Viscosity of a substance is the ratio of the dynamic viscosity to its density. The values are given at specific temperatures. These temperatures can be searched using the associated parameter field code KV.T. The search field remarks contain concentration details of solutions, as well as, information to further specialize conditions relating to the dynamic viscosity values. You can search for this associated information using the parameter field code KV.MR. The forms of state of the substance can be restricted by the associated parameter field code KV.S

Unit:

The default unit in the CrossFire database for this fact is Stokes=cm²s⁻¹.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
KV	Kinematic Viscosity	Numeric	St
KV.T	Temperature	Numeric	°C
KV.MR	Remarks	Wordwise	
KV.S	Forms of State	String (phrase)	

Type of Indexing:

Group Code

2.5.7.4 Non-Newtonian Behavior (NONT)

Description:

Compounds show Non-Newtonian Behavior if their viscosity at a given temperature is not a constant, but depends on the shear velocity or the shear stress.

Type of Indexing:



2.5.8 Thermodynamic Data

2.5.8.1 Thermodynamic Data (TDD)

Specific Heat at Constant Pressure (CP) Cp (Further Information) (CPFI) Specific Heat at Constant Volume (CV) Cv (Further Information) (CVFI) Enthalpy of Formation (HFOR) Enthalpy of Formation (Further Information) (HFFI) Gibbs Energy of Formation (GFOR) Gibbs Energy of Formation (Further Information) (GFFI) Entropy of Formation (SFOR) Entropy of Formation (Further Information) (SFFI) Equation of State (EQST) Acentric Factor (ACEN)

2.5.8.2 Specific Heat at Constant Pressure (CP)

Description:

The Specific (Molar) Heat Capacity at Constant Pressure is the enthalpy necessary to raise the temperature of one mole (one mass unit) of substance one degree at constant pressure. The coefficient field is a numeric, range searchable field and is linked to the associated temperature CP.T and pressure CP.P, as well as, to the forms of state of the substance by parameter field code CP.S.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
СР	Cp (molar)	Numeric	J mol ⁻¹ K ⁻¹
CP.VALA	Cp (specific)	Numeric	J g⁻¹K ⁻¹
CP.T	Temperature	Numeric	°C
CP.P	Pressure	Numeric	Torr
CP.MR	Remarks	Wordwise	



CP.S Forms of State String (phrase)

Type of Indexing:

Group Code

2.5.8.3 Cp (Further Information) (CPFI)

Description:

The search field CP (Further Information) indicates if further information about the specific heat at constant pressure is given as a diagram, table, or equation in the fields CPFI.DGM, CPFI.TAB and CPFI.EQN. The search field is linked to the forms of state of the substance by parameter field code CPFI.S.

Note:

Searching for the special field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
CPFI.DGM	Cp as Diagram	String (phrase)
CPFI.TAB	Cp as Table	String (phrase)
CPFI.EQN	Cp as Equation	String (phrase)
CPFI.S	Forms of State	String (phrase)

Type of Indexing:

Group Code

2.5.8.4 Specific Heat at Constant Volume (CV)

Description:

The Specific (Molar) Heat Capacity at Constant Volume is the internal energy required to raise the temperature of one mole (one gram) of substance one degree at constant volume. The coefficient field is a Numeric, range searchable field and is linked to the associated temperature CV.T and pressure CV.P as well as to the forms of state of the substance by parameter field code CV.S.



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
CV	CV molar	Numeric	J mol ⁻¹ K ⁻¹
CV.VALA	CV specific	Numeric	J g⁻¹K ⁻¹
CV.T	Temperature	Numeric	°C
CV.P	Pressure	Numeric	Torr
CV.MR	Remarks	Wordwise	
CV.STATE	Forms of State	String (phrase)	

Type of Indexing:

Group Code

2.5.8.5 Cv (Further Information) (CVFI)

Description:

The search field CV (Further Information) indicates if further information about the specific heat at constant volume is given as a diagram, table, or equation in the fields CVFI.DGM, CVFI.TAB and CVFI.EQN. The search field is linked to the forms of state of the substance by parameter field code CVFI.S.

Note:

Searching for the special field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
CVFI.DGM	Cv as Diagram	String (phrase)
CVFI.TAB	Cv as Table	String (phrase)
CVFI.EQN	Cv as Equation	String (phrase)
CVFI.S	Forms of State	String (phrase)

Type of Indexing:



2.5.8.6 Enthalpy of Formation (HFOR)

Description:

The (specific) Enthalpy of Formation is the change in enthalpy associated with the formation of one mole (one gram) of a substance from its elements in their standard states at specified temperature and pressure. The coefficient field is a Numeric, range searchable field and is linked to the associated temperature HFOR.T and pressure HFOR.P as well as to the forms of state of the substance by parameter field code HFOR.S.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
HFOR	Molar Enthalpy of Formation	Numeric	J mol⁻¹
HFOR.VALA	Specific Enthalpy of Formation	Numeric	J g⁻¹
HFOR.T	Temperature	Numeric	°C
HFOR.P	Pressure	Numeric	Torr
HFOR.S	Forms of State	String (phrase)	

Type of Indexing:

Group Code

2.5.8.7 Enthalpy of Formation (Further Information) (HFFI)

Description:

The search field Enthalpy of Formation (Further Information) indicates if further information about the enthalpy of formation is given as a diagram, table, or equation in the fields HFFI.DGM, HFFI.TAB and HFFI.EQN. The search field is linked to the forms of state of the substance by parameter field code HFFI.S.

Note:

Searching for the special field codes require the field value "available" or more general "*".



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
HFFI.DGM	Diagram	String (phrase)
HFFI.TAB	Table	String (phrase)
HFFI.EQN	Equation	String (phrase)
HFFI.S	Forms of State	String (phrase)

Type of Indexing:

Group Code

2.5.8.8 Gibbs Energy of Formation (GFOR)

Description:

The (specific) Gibbs Energy of Formation is the change in Gibbs energy associated with the isothermal formation of one mole (one gram) of the substance from the elements in their standard states at specified temperature and pressure. The coefficient field is a Numeric, range searchable field and is linked to the associated temperature GFOR.T and pressure GFOR.P as well as to the forms of state of the substance by parameter field code GFOR.S.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
GFOR	Molar Gibbs Energy of Formation	Numeric	J mol⁻¹
GFOR.VALA	Specific Gibbs Energy of Formation	Numeric	Jg⁻¹
GFOR.T	Temperature	Numeric	°C
GFOR.P	Pressure	Numeric	Torr
GFOR.S	Forms of State	String (phrase)	

Type of Indexing:



2.5.8.9 Gibbs Energy of Formation (Further Information) (GFFI)

Description:

The search field Gibbs Energy of Formation (Further Information) indicates if further information about the Gibbs energy of formation is given as a diagram, table, or equation in the fields GFFI.DGM, GFFI.TAB and GFFI.EQN. The search field is linked to the forms of state of the substance by parameter field code GFFI.S.

Note:

Searching for the special field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
GFFI.DGM	Diagram	String (phrase)
GFFI.TAB	Table	String (phrase)
GFFI.EQN	Equation	String (phrase)
GFFI.S	Forms of State	String (phrase)

Type of Indexing:

Group Code

2.5.8.10 Entropy of Formation (SFOR)

Description:

The (specific) Entropy of Formation is the change in entropy associated with the formation of one mole of a substance from its elements in their standard states at specified temperature and pressure. The entropy of formation (specific: SFOR:VALA, molar: SFOR) search field is a Numeric, range searchable field and is linked to the associated temperature SFOR.T and pressure SFOR.P, as well as, to the forms of state of the substance by parameter field code SFOR.S.



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
SFOR	Molar Entropy of Formation	Numeric	J mol ⁻¹ K ⁻¹
SFOR.VALA	Specific Entropy of Formation	Numeric	Jg⁻¹K ⁻¹
SFOR.T	Temperature	Numeric	°C
SFOR.P	Pressure	Numeric	Torr
SFOR.S	Forms of State	String (phrase)	

Type of Indexing:

Group Code

2.5.8.11 Entropy of Formation (Further Information) (SFFI)

Description:

The search field Entropy of Formation (Further Information) indicates if further information about the Gibbs energy of formation is given as a diagram, table, or equation in the fields SFFI.DGM, SFFI.TAB and SFFI.EQN. The search field is linked to the forms of state of the substance by parameter field code SFFI.S.

Note:

Searching for the special field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
SFFI.DGM	Diagram	String (phrase)
SFFI.TAB	Table	String (phrase)
SFFI.EQN	Equation	String (phrase)
SFFI.S	Forms of State	String (phrase)

Type of Indexing:

Group Code

2.5.8.12 Equation of State (EQST)



Description:

The equation of state describes the state of a substance in terms of the relationship of the basic physical quantities of volume, pressure, and temperature. The field Equation of State indicates a thermodynamic equation of state.

Type of Indexing:

Group Code

2.5.8.13 Acentric Factor (ACEN)

Description:

The Pitzer Acentric Factor is a measure of nonsphericity or acentricity of a molecule's potential force field.

Associated Parameter Field:

Field Code	Full Name	Type of Indexing
ACEN.VAL	Acentric Factor	Numeric

Type of Indexing:



2.5.9 Electric Properties

2.5.9.1 Electric Properties (ELE)

Electronic Band Structure (ELSR) Electric Conductivity (ELCO) Superconductivity (SPCO) Photoconductivity (PHCO) Thermoelectric Effects (THEL) Dielectric Constant (DIC) Piezoelectric Effects (PIEZ) Pyroelectric Effects (PYRO) Ferroelectricity (FREL) Antiferroelectricity (AFREL)

2.5.9.2 Electronic Band Structure (ELSR)

Description:

The field Electronic Band Structure indicates discussion or calculation of (electron) band models, like band theory, valence band, conduction band, and Brillouin zone.

Type of Indexing:

Group Code

2.5.9.3 Electric Conductivity (ELCO)

Description:

The Electric Conductivity of a substance is its time rate of electricity flowing across a unit area per unit potential gradient. The electric conductivity search field is a numeric range searchable field and is linked to the associated temperature ELCO.T. Information about pretreatment of samples is given in the field remarks ELCO.MR. Further information about the electric conductivity is given as a diagram or equation using the parameter field codes ELCO.DGM respectively ELCO.EQN. Indication of the electric charge mobility or the conduction mechanism is given in the parameter field ELCO.MOB.



The search field is linked to the forms of state of the substance by parameter field code ELCO.S.

Note:

Searching for electric conductivity diagrams or equation field codes require the field value "available", the parameter field ELCO.MOB the field value "described" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
ELCO	Electric Conductivity	Numeric	S cm ⁻¹
ELCO.T	Temperature	Numeric	°C
ELCO.MR	Remarks	Wordwise	
ELCO.DGM	Diagram	String (phrase)	
ELCO.EQN	Equation	String (phrase)	
ELCO.MOB	Conduction Mechanism	String (phrase)	
ELCO.S	Forms of State	String (phrase)	

Type of Indexing:

Group Code

2.5.9.4 Superconductivity (SPCO)

Description:

The superconductivity is the abrupt increase of electric conductivity of some materials at low temperatures. Indication on the general description of Superconductivity is given in the field SPCO.FLG. Each substance has a transition temperature (onset-, midpoint-, zero-resistance temperature) Tc above which it is a normal conductor. The superconductivity SPCO.T field contains this critical temperature for a given substance in the literature.

Note:

Searching for superconductivity field codes require the field "described" or more general "*".



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
SPCO.FLG	superconductivity	String (phrase)	
SPCO.T	critical temperature	Numeric	°C

Type of Indexing:

Group Code

2.5.9.5 Photoconductivity (PHCO)

Description:

The photoconductivity is the increase in electrical conductivity of nonmetallic substances when exposed to light. The production of voltage in semiconductor materials by light or other electromagnetic radiation is called photovoltaic effect. The field Photoconductivity indicates information on photoconductivity or photovoltaic effects of a substance.

Type of Indexing:

Group Code

2.5.9.6 Thermoelectric Effects (THEL)

Description:

The thermoelectric effect is the production of current or voltage (thermal electromotive force) at a junction between two dissimilar metals or at a temperature gradient along a metal rod. The value depends on the metal, as well as, the temperature or temperature gradient, respectively. The field Thermoelectric Effects indicates thermoelectric phenomena, e.g. Benedicks effect, Thomson effect, or Bridgman effect.

Type of Indexing:



2.5.9.7 Piezoelectric Effects (PIEZ)

Description:

The piezoelectric effect is the electrical response of a substance induced by mechanical forces, and the mechanical response to an electrical force. The occurrence of piezoelectricity depends on the crystal structure. The crystal must belong to one of the non-centrosymmetric point groups. The field Piezoelectric Effects indicates piezoelectric substances.

Type of Indexing:

Group Code

2.5.9.8 Dielectric Constant (DIC)

Description:

The Dielectric Constant is the ratio of the capacity of a condenser with the given substance as dielectric to the capacity of the same condenser with a vacuum as dielectric. The dielectric constant search field is a numeric range searchable field and is linked to the associated fields temperature DIC.T, frequency DIC.F and pressure DIC.P. Information about pretreatment of samples is given in the field remarks DIC.MR. The search field is linked to the forms of state, of the substance, by the parameter field code DIC.S.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
DIC	Dielectric Constant	Numeric	
DIC.T	Temperature	Numeric	°C
DIC.F	Frequency	Numeric	Hz
DIC.P	Pressure	Numeric	Torr
DIC.MR	Remarks	Wordwise	
DIC.S	Forms of State	String (phrase)	

Type of Indexing:



2.5.9.9 Pyroelectric Effects (PYRO)

Description:

The pyroelectric effect is the development of a spontaneous electrical polarization and is temperature dependent. This is mainly because the thermal expansion that occurs on heating changes the sizes (i.e. lengths) of the dipoles. The occurrence of pyroelectricity depends on the crystal structure. The crystal must belong to one of the non-centrosymmetric point groups. The field Pyroelectric Effects indicates pyroelectric substances.

Type of Indexing:

Group Code

2.5.9.10 Ferroelectricity (FREL)

Description:

Ferroelectricity is due to parallel ordering of electric dipole moments in a substance in the absence of an applied electrical field. In ferroelectrics, domain structures are formed because adjacent dipoles tend to align themselves parallel to each other. Above a critical temperature ferroelectricity often disappears and a paraelectric state is generated. The field Ferroelectricity indicates ferroelectric substances.

Type of Indexing:

Group Code

2.5.9.11 Antiferroelectricity (AFREL)

Description:

Antiferroelectricity is due to antiparallel ordering of electric dipole moments in a substance in the absence of an applied electrical field. The substance is built from two sublattices having equal and opposite electric polarization. Above a critical temperature antiferroelectricity disappears and a paraelectric state is generated. The field Antiferroelectricity indicates the antiferroelectric substances.

Type of Indexing:



2.5.10 Magnetic Properties

2.5.10.1 Magnetic Properties (MAG)

Magnetostriction (MAGS) Galvanomagnetic Effects (GALV) Thermomagnetic Effects (THMG) Magnetic Susceptibility (MSUS) Curie Weiss Temperature (CUWE) Neel Temperature (NEEL) Curie Temperature (CURI) Magnetic Moment (MM) Magnetic Property (MPRO)

2.5.10.2 Magnetostriction (MAGS)

Description:

The field Magnetostriction indicates information on the volume change of substances under the influence of an external magnetic field.

Type of Indexing

Group Code

2.5.10.3 Galvanomagnetic Effects (GALV)

Description:

The galvanomagnetic effect is any thermal or electric effect occurring in an electronic conductor or semiconductor in the presence of a magnetic field. The field Galvanomagnetic Effects indicates information on

- Hall effect (formation of a transversal potential gradient)
- Ettingshausen effect (formation of a transversal temperature gradient)
- Galvanomagnetic Thomson effect (change of transversal resistivity)
- Nernst effect (formation of a longitudinal temperature gradient)



Type of Indexing:

Group Code

2.5.10.4 Thermomagnetic Effects (THMG)

Description:

Indication of Thermomagnetic Effects including all effects caused by a heat flow in electronic conductors or semiconductors in the presence of a magnetic field, e.g.:

- Righi-Leduc effect (formation of a transversal temperature gradient)
- Maggi-Righi-Leduc effect (change of thermal conductivity)
- Ettingshausen-Nernst effect (formation of a potential gradient)

Type of Indexing:

Group Code

2.5.10.5 Magnetic Susceptibility (MSUS)

Description:

The volumetric magnetic susceptibility is the difference in the magnetic permeability of a substance and of a vacuum, divided by the permeability of the vacuum. The specific (or mass) magnetic susceptibility of a substance is the quotient of its volumetric susceptibility and its density. The molar magnetic susceptibility of a substance is the product of its volumetric susceptibility and its molar volume. The Magnetic Susceptibility field contains the values for a given substance in the literature. The search field is a numeric range searchable field and is linked to the associated temperature MSUS.T. Further information about the magnetic susceptibility is given as a diagram or equation using the parameter field codes MSUS.DGM, and MSUS.EQN respectively. Information about pretreatment of samples or concentration of solutions is given in the field remarks MSUS.MR.

Note:

Searching for magnetic susceptibility diagram or equation field codes require the field value "available" or more general "*".



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
MSUS	Molar Magnetic Susceptibility	Numeric	ml mol ⁻¹
MSUS.VALA	Specific Magnetic Susceptibility	Numeric	l kg⁻¹
MSUS.T	Temperature	Numeric	°C
MSUS.DGM	Diagram	String (phrase)	
MSUS.EQN	Equation	String (phrase)	
MSUS.MR	Remarks	Wordwise	

Type of Indexing:

Group Code

2.5.10.6 Curie Weiss Temperature (CUWE)

Description:

The Curie-Weiss temperature is the temperature at which a plot of the reciprocal molar magnetic susceptibility against the absolute temperature T intersects the T-axis. The Curie-Weiss temperature can adopt positive, as well as, negative values. The Curie-Weiss Temperature CUWE.T search field contains the values for a given substance in the literature and is a numeric range searchable field. Information about pretreatment of samples is given in the field remarks CUWE.MR.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
CUWE.T	Curie Weiss Temperature	Numeric	К
CUWE.MR	Remarks	Wordwise	

Type of Indexing:



2.5.10.7 Neel Temperature (NEEL)

Description:

The Neel temperature is the temperature at which a substance experiences a phase transition from the antiferromagnetic to the paramagnetic state. The Neel temperature is also known as antiferromagnetic Curie point. The Neel Temperature search field contains the values for a given substance in the literature and is a numeric range searchable field. Information about pre-treatment of samples is given in the field remarks NEEL.MR.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
NEEL.T	Neel Temperature	Numeric	К
NEEL.MR	Remarks	Wordwise	

Type of Indexing:

Group Code

2.5.10.8 Curie Temperature (CURI)

Description:

The Curie temperature is the temperature above which a substance experiences a phase transition from the ferromagnetic to the paramagnetic state. The Curie Temperature search field contains the values for a given substance in the literature and is a numeric range searchable field. Information about pretreatment of samples is given in the field remarks CURI.MR.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
CURI.T	Curie Temperature	Numeric	К
CURI.MR	Remarks	Wordwise	

Type of Indexing:



2.5.10.9 Magnetic Moment (MM)

Description:

The magnetic dipole moment of a molecule is the moment due to the spin of electrons and nuclei. The Magnetic Dipole Moment is a numeric range searchable field.

Associated Parameter Field:

Field Code	Full Name	Type of Indexing	Unit
MM.VAL	Magnetic Moment	Numeric	A cm ²

Type of Indexing:

Group Code

2.5.10.10 Magnetic Property (MPRO)

Description:

The search field to Magnetic Property indicates if information about magnetic effects is available for a given substance in the literature. Indication on magnetic phenomena is given in the field MPRO.MPH. Indication of the anisotropy of the magnetic susceptibility is given in the field MPRO.MAN. Indication of the dependence of the spatial arrangement and orientation of the atomic magnetic moments in a crystal or indication of magnetic space groups (also designated as Schubnikow groups) is given in the field MPRO.MST. Indication of the magnetization (magnetic dipole moment per unit volume) is given in the field MPRO.MAG. Indication of the relationship between the magnetization or magnetic induction and the effective field strength given as a diagram is given in the field MPRO.MMD. Indication of the relationship between the magnetization M, the magnetic field H, or the magnetic induction B and the temperature given as a diagram is available in the field MPRO.MPD.

Note:

Searching for magnetic anisotropy field codes require the field value "described", the field codes magnetic structure / space group, magnetization, magnetization diagram, and magnetic phase diagram require "available" or more general "*".



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
MPRO.MPH	Magnetic Phenomena	String (phrase)
MPRO.MAN	Magnetic Anisotropy	String (phrase)
MPRO.MST	Magnetic Structure / Space Group	String (phrase)
MPRO.MAG	Magnetization	String (phrase)
MPRO.MMD	Magnetization Diagram	String (phrase)
MPRO.MPD	Magnetic Phase Diagram	String (phrase)

Type of Indexing:

Group Code

2.5.10.11 Magnetic Phenomena (MPRO.MPH)

Description:

Indication on the following magnetic phenomena is given in this field:

- antiferromagnetic
- diamagnetic
- ferrimagnetic
- ferrimagnetic domains
- ferromagnetic
- ferromagnetic domains
- paramagnetic

Type of Indexing:

String (phrase)



2.5.11 Optical Properties

2.5.11.1 Optical Properties (OPT)

Refractive Index (RI) Circular Dichroism (CDIC) Optical Anisotropy (OA) Optical Rotation (OR) Magnetooptical Effects (MAGO) Electrooptical Effects (ELOP)

2.5.11.2 Refractive Index (RI)

Description:

The refractive index is the ratio of the velocity of light in a vacuum to the velocity of light in the substance. The Refractive Index is a numeric range searchable field and is linked to the associated temperature RI.T and wave length RI.W. Indication of birefringence is given in the field RI.BR. Information about pretreatment of samples is given in the field remarks RI.MR. The refractive index is linked to the forms of state of the substance by parameter field code RI.S.

Note:

Searching for birefringence field code requires the field value "yes" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
RI	Refractive Index	Numeric	
RI.T	Temperature	Numeric	°C
RI.W	Wave Length	Numeric	nm
RI.MR	Remarks	Wordwise	
RI.BR	Birefringence	String (phrase)	
RI.S	Forms of State	String (phrase)	

Type of Indexing:



Group Code

2.5.11.3 Circular Dichroism (CDIC)

Description:

Circular dischroism is due to the phenomen that planar polarization converts to elliptic polarization when linearly polarized incident light passes through an active medium, e.g. an optically active substance. Optically active substances absorb left and right polarized light unequally. The field Circular Dichroism indicates information on circular dichroism or optical rotary dispersion of a substance.

Type of Indexing:

Group Code

2.5.11.4 Optical Anisotropy (OA)

Description:

The behavior of a substance whose effect on electromagnetic radiation varies with the direction of the propagation is designated as optical anisotropy. The field Optical Anisotropy indicates information on optical anisotropic behavior of substances.

Type of Indexing:

Group Code

2.5.11.5 Optical Rotation (OR)

Description:

The optical rotation is the rotation of the plane of polarization in a beam of light as it passes through an optically active substance. The field Optical Rotation indicates the optical rotary power of an optical active substance.

Type of Indexing:

Group Code

2.5.11.6 Magnetooptical Effects (MAGO)



Description:

Magnetooptic effects relate to the effect of magnetism on the propagation of light. The field Magnetooptical Effects indicates information on magnetooptical effects of substances, e.g. Faraday effect, magneto-optic Kerr effect, magneto-optical-rotational-dispersion (MORD) or Zeeman effect.

Type of Indexing

Group Code

2.5.11.7 Electrooptical Effects (ELOP)

Description:

Electrooptic effects are the alteration of the optical behavior, e.g. refractive properties, by an applied electric field. The field Electrooptical Effects indicates information on electrooptical effects of substances, e.g. Kerr effect, Stark effect, or Pockels effect.

Type of Indexing:



2.5.12 Spectroscopic Information

2.5.12.1 Spectroscopic Information (SPE)

This field identifier is a group code for Field Availability searches concerned with spectral information in the hierarchical system of the Gmelin database.

NMR Spectroscopy (NMR) IR Spectroscopy (IR) NQR Spectroscopy (NQR) ESR Spectroscopy (ESR) Mass Spectrometry (MS) UV/VIS Spectroscopy (UV) Moessbauer Spectroscopy MOE) Rotational Spectroscopy (ROT) Raman Spectroscopy (RAMAN) UV Photoelectron Spectroscopy (UPS) X-Ray Photoelectron Spectroscopy (XPS) X-Ray Spectroscopy (XSPE) Auger Spectroscopy (AUG) Fluorescence Spectroscopy (FLU) Phosphorescence Spectroscopy (PHO) Luminescence Spectroscopy (LUM) Electron Energy Loss Spectroscopy (EEL)

Type of Indexing:

Group Code

2.5.12.2 NMR Spectroscopy (NMR)

Description:

NMR spectroscopy is based on resonance of nuclei (nuclear spin I>0) with a magnetic field and an electromagnetic field. Usually the frequency of the electromagnetic field is swept.



The NMR Spectroscopy search field is linked to the associated search fields solvent NMR.SOL and temperature parameters NMR.T. The corresponding spin-spin coupling atoms are searchable in NMR.NUI. Indications about a shown spectrum is given in the parameter field code NMR.DGM, as well as, further information about signals and intensities (NMR.SIGNAL), linewidths or dynamic effects (NMR.LINDYN), and description of spectra (NMR.DESCR).

Note:

Searching for spectrum diagram, signals / intensities, linewidths, or spectrum description field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
NMR.NUC	Nucleus	String (phrase)	
NMR.DGM	Spectrum Diagram	String (phrase)	
NMR.NUI	Spin-Spin Coupling to Nuclei	String (phrase)	
NMR.SOL	Solvent	String (phrase)	
NMR.T	Temperature	Numeric	°C
NMR.SIGNAL	Signal / Intensities	String (phrase)	
NMR.LINDYN	Linewidths	String (phrase)	
NMR.DESCR	Description	String (phrase)	

Type of Indexing:

Group Code

2.5.12.3 IR Spectroscopy (IR)

Description:

The IR absorption spectrum of a substance is produced by its interaction with electromagnetic radiation (usually in the range 10-10000 cm⁻¹) due to vibrational transitions. The IR Spectroscopy search fields are linked to the associated search fields solvent IR.SOL and temperature parameters IR.T. The wavelength range is searchable with IR.SPEC. Indications about a shown spectrum is given in the parameter field code IR.DGM as well as further information about bands and intensities (IR.BANDS), and description of spectra (IR.DESCR).

Note:



Searching for spectrum diagram, bands / intensities, or spectrum description field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
IR.DGM	Spectrum Diagram	String (phrase)	
IR.SOL	Solvent	String (phrase)	
IR.SPEC	Spectral Range	Numeric	cm⁻¹
IR.T	Temperature	Numeric	°C
IR.BANDS	Bands / Intensities	String (phrase)	
IR.DESCR	Description	String (phrase)	

Type of Indexing:

Group Code

2.5.12.4 NQR Spectroscopy (NQR)

Description:

The NQR-Spectrum is based on resonance between the nuclear quadrupole moment and a variable electric field gradient. The nuclei are searchable with the parameter field NQR.NUC. Indications about a shown spectrum is given in the parameter field code NQR.DGM as well as further information about quadrupole splitting (NQR.SPLIT), and description of spectra (NQR.DESCR).

Note:

Searching for spectrum diagram, quadrupole splitting, or spectrum description field codes require the field value "available" or more general "*".



Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
NQR.NUC	Nucleus	String (phrase)
NQR.DGM	Spectrum Diagram	String (phrase)
NQR.SPLIT	Quadrupole Splitting	String (phrase)
NQR.DESCR	Description	String (phrase)

Type of Indexing:

Group Code

2.5.12.5 ESR Spectroscopy (ESR)

Description:

ESR spectroscopy is determined by measuring the paramagnetic properties of atoms or molecules in a magnetic field. Resonance absorption occurs on irradiation with electromagnetic waves of appropriate frequency. Hyperfine coupling is due to interaction of the unpaired electrons with nuclei possessing a magnetic moment. The ESR Spectroscopy search fields contain the coupling nuclei ESR.NUI, g-factors ESR.GFAC, and solvent ESR.SOL and temperature parameters ESR.T. Indications about a shown spectrum is given in the parameter field code ESR.DGM, as well as, description of spectra in ESR.DESCR.

Note:

Searching for spectrum diagram, g-factors, or spectrum description field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of	Unit
		Indexing	
ESR.DGM	Spectrum Diagram	String (phrase)	
ESR.GFAC	G-Factors	String (phrase)	
ESR.NUI	Hyperfine Coupling To Nuclei	String (phrase)	
ESR.SOL	Solvent	String (phrase)	
ESR.T	Temperature	Numeric	°C



ESR.DESCR Description

String (phrase)

Type of Indexing:

Group Code

2.5.12.6 Mass Spectrometry (MS)

Description:

Method based on sending an ionized beam of molecules or molecule fragments through an inhomogeneous magnetic field to achieve a separation depending on the mass/charge ratio of the particles. Indications about a shown spectrum is given in the parameter field code MS.DGM, as well as, further information of a given molecular peak (MS.PEAK), the fragmentation (MS.FRAG), and description of spectra (MS.DESCR).

Note:

Searching for the special field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
MS.DGM	Spectrum Diagram	String (phrase)
MS.PEAK	Molecular Peak	String (phrase)
MS.FRAG	Fragmentation Pattern	String (phrase)
MS.DESCR	Description	String (phrase)

Type of Indexing:



2.5.12.7 UV / VIS Spectroscopy (UV)

Description:

The UV and VIS absorption spectrum of a substance is produced by its interaction with electromagnetic radiation (usually in the range 200-800 nm) due to electronic transitions. The UV / VIS Spectroscopy search field contains solvent (UV.SOL) and spectral range parameters (UV.SPEC) for ultraviolet and visible spectra. Indications about a shown spectrum is given in the parameter field UV.DGM as well as further information of assignments (UV.ASSIG), and description of spectra (MS.DESCR).

Note:

Searching for spectrum diagrams, assignment, or spectrum description field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
UV.DGM	Spectrum Diagram	String (phrase)	
UV.SOL	Solvent	String (phrase)	
UV.SPEC	Spectral Range	Numeric	nm
UV.ASSIG	Assignment	String (phrase)	
UV.DESCR	Description	String (phrase)	

Type of Indexing:

Group Code

2.5.12.8 Moessbauer Spectroscopy (MOE)

Description:

Gamma-ray nuclear resonance absorption spectroscopy, with frozen Moessbauer nuclides, is searchable in the parameter field MOENUC. Indications about a shown spectrum are given in the parameter field code MOE.DGM, as well as, further information of chemical shifts (MOE.SHIFTS), quadrupole splitting (MOE.SPLIT), and description of spectra (MOE.DESCR).



Note:

Searching for spectrum diagrams, chemical shifts, quadrupole splitting, or spectrum description field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
MOE.NUC	Nucleus	String (phrase)
MOE.DGM	Spectrum Diagram	String (phrase)
MOE.SHIFTS	Chemical Shifts	String (phrase)
MOE.SPLIT	Quadrupole Splitting	String (phrase)
MOE.DESCR	Description	String (phrase)

Type of Indexing:

Group Code

2.5.12.9 Rotational Spectroscopy (ROT)

Description:

The field Rotational Spectroscopy indicates a molecular spectrum produced by rotational-level transition. It is produced in the far-infrared region between 50 and 200 μ m and in the microwave region.

Type of Indexing:

Group Code

2.5.12.10 Raman Spectroscopy (RAMAN)

Description:

Raman spectroscopy is a method in which the sample is illuminated with monochromatic light, usually generated by a laser. The scattered light produced by the sample is first scatter with exactly the same energy and wavelength as the incident light and second Raman scatter, which is usually much less intense than the first one and emerges at either longer or shorter wavelength than the incident light.



The field Raman Spectroscopy indicates the application of, or information to, a vibrational (Raman) spectrum of a substance.

Type of Indexing:

Group Code

2.5.12.11 UV Photoelectron Spectroscopy (UPS)

Description:

UV photoelectron spectroscopy (UPS) is a method in which a surface is bombarded with photons in the ultraviolet energy range and the resulting electronic emission spectrum is analyzed for information about molecular orbital characteristics or occupied and unoccupied density of states of substances. The field UV Photoelectron Spectroscopy indicates the application of, or information to, a UV photoelectron spectrum of a substance.

Type of Indexing:

Group Code

2.5.12.12 X-Ray Photoelectron Spectroscopy (XPS)

Description:

X-Ray photoelectron spectroscopy (XPS) is a method in which a substance is irradiated with X-rays and the energies of the photoelectrons emitted from the substance surface are determined. The field X-Ray Photoelectron Spectroscopy indicates the application of, or information to, an X-ray photoelectron spectrum of a substance.

Type of Indexing:

Group Code

2.5.12.13 X-Ray Spectroscopy (XSPE)

Description:

X-ray spectroscopy measures the absorption of X-rays by samples, especially at energies in the region of absorption edges. The field X-Ray Spectroscopy indicates the application of X-ray spectroscopy e.g. EXAFS, XANES or EDAX.



Type of Indexing:

Group Code

2.5.12.14 Auger Spectroscopy (AUG)

Description:

Auger Spectroscopy is a method to analyze the energy of secondary (Auger) electrons ejected from a solid surface upon irradiation with accelerated electrons or X-ray photons. The field Auger Spectroscopy indicates the application of, or information to, an Auger spectrum of a substance.

Type of Indexing:

Group Code

2.5.12.15 Fluorescence Spectroscopy (FLU)

Description:

Fluorescence spectrum is produced by the emission of radiation following absorption of and excitation by radiant energy. The field Fluorescence Spectroscopy indicates the application of, or information to, a fluorescence spectrum of a substance.

Type of Indexing:

Group Code

2.5.12.16 Phosphorescence Spectroscopy (PHO)

Description:

The phosphorescence spectroscopy is a method based on the emission of light to spinforbidden decay of electrons from excited states to the ground state. The field Phosphorescence Spectroscopy indicates the application of, or information to, a phosphorescence spectrum of a substance.

Type of Indexing:



2.5.12.17 Luminescence Spectroscopy (LUM)

Description:

Luminescence spectroscopy is a method based on the emission of light from a non thermal excitation. Luminescence is obtained under a variety of excitation sources, e.g. electromagnetic radiation, particles (like electron beam), electric field. The field Luminescence Spectroscopy indicates the application of, or information to, a luminescence spectrum of a substance.

Type of Indexing:

Group Code

2.5.12.18 Electron Energy Loss Spectroscopy (EEL)

Description:

An electron energy loss spectrum is a method in which a substance is bombarded with monochromatic electrons and the distribution of the energy loss by scattered electrons is determined. The field Electron Energy Loss Spectroscopy indicates the application of or information to an electron energy loss spectrum of a substance.

Type of Indexing:



2.6 Properties of / in Systems

2.6.1 Properties of / in Systems (MCS)

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Sorption (SOR)
Diffusion (DIF)
Azeotropes (AZE)
Distribution (DIST)
System Components (MULT)
Multicomponent System Data (META)
Eutectic Temperature (ETEC)
Peritectic Temperature (PTEC)
Solubility In Systems (MULS)
Vapor Pressure of Solution (VPR)
Solubility (SLB)
PL-Value (SLBP)
Surface Tension (ST)
```

2.6.2 Sorption (SOR)

Description:

The Sorption search field contains the availability of references about adsorption, absorption, desorption and chemisorption. It is divided into the sorption of a substance, or a title compound.

Sorption to Title Compound (SPTO)

Sorption of Title Compound (SPOF)

2.6.2.1 Sorption to Title Compound (SPTO)

Description:

The Sorption to Title Compound search field contains the availability of references about adsorption, absorption, desorption and chemisorption to a substance, a title compound. The title compound is the adsorbent. The other component, the adsorbate, is searchable in the associated field SPTO.PA.



Further information about the sorption is given as a diagram, table or equation using the parameter field codes SPTO.DGM, SPTO.TAB respectively SPTO.EQN. Further data are given in the field SPOF.

Note:

Searching for a diagram, table or equation field require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
SPTO.PA	Other Component	String (phrase)
SPTO.DGM	Diagram	String (phrase)
SPTO.TAB	Table	String (phrase)
SPTO.EQN	Equation	String (phrase)

Type of Indexing:

Group Code

2.6.2.2 Sorption of Title Compound (SPOF)

Description:

The Sorption of Title Compound search field contains the availability of references about adsorption, absorption, desorption and chemisorption of a substance, a title compound. The title compound is the adsorbate. The other component, the adsorbent, is searchable in the associated field SPOF.PA. Further information about the sorption is given as a diagram, table or equation using the parameter field codes SPOF.DGM, SPOF.TAB respectively SPOF.EQN. Further data are given in the field SPTO.

Note:

Searching for a diagram, table or equation field codes require the field value "available" or more general "*".

Field Code	Full Name	Type of Indexing
SPOF.PA	Other Component	String (phrase)
SPOF.DGM	Diagram	String (phrase)



SPOFTAB Table SPOF.EQN Equation String (phrase) String (phrase)

Type of Indexing:

Group Code

2.6.3 Diffusion (DIF)

Description:

The Diffusion search field contains the availability of references about diffusion of a substance into a second one without the influence of external forces. It is divided into the diffusion of a substance, or a title compound.

Diffusion of Title Compound (DIFOF)

Diffusion into Title Compound (DIFIN)

2.6.3.1 Diffusion of Title Compound (DIFOF)

Description:

The diffusion coefficient is the ratio of the mass flux to the concentration gradient in a system, which is free of external force fields and is homogeneous with respect to temperature and pressure. The Diffusion of Title Compound search field contains references to the diffusion of a title compound into a second compound and eventually the diffusion coefficient of a given substance (gas, liquid, but also solid) in the literature through the second one. The field is linked to the associated temperature DIFOF.T and to the compound (other phase) DIFOF.PA through which the given substance diffuses.

Field Code	Full Name	Type of Indexing	Unit
DIFOF.PA	Other Component	String (phrase)	
DIFOF	Diffusion Coefficient	Numeric	cm ² s ⁻¹
DIFOF.T	Temperature	Numeric	°C



Type of Indexing:

Group Code

2.6.3.2 Diffusion into Title Compound (DIFIN)

Description:

The diffusion into title compound is the diffusion of a substance into the title compound without influence of external forces. The Diffusion into Title Compound search field contains references to the diffusion of a substance into the title compound for a given substance in the literature. The field is linked to the compound (other phase) DIFOF.PA which diffuses through the given substance.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
DIFIN.PA	Other Component	String (phrase)

Type of Indexing:

Group Code

2.6.4 Azeotropes (AZE)

Description:

An azeotrope is a solution with an extremum boiling point at which composition of liquid and vapor phases are equal; therefore, there is no change in composition on boiling. The azeotrope field contains the azeotropy of the title compound with another substance in the literature.

Field Code	Full Name	Type of Indexing
AZE.PG	Azeotropes (GRN)	Numeric
AZE.PA	Azeptropes	String (phrase)



Type of Indexing:

Group Code

2.6.5 Distribution (DIST)

Description:

A substance distributes itself between two immiscible solvents so that the ratio of its concentrations in the two solvents is approximately a constant. The distribution coefficient gives the ratio of the concentrations in the two phases at a specified temperature at the equilibrium state. The two phases can be specified in the search fields DIST.PHAS1 and DIST.PHAS2. The Distribution search field contains references to the distribution of a given substance in the literature between two phases and eventually the distribution coefficient. The distribution coefficient search field DIST is a numeric search field and is linked to the associated temperature DIST.T.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
DIST.PHAS1	Phase 1	String (phrase)	
DIST.PHAS2	Phase 2	String (phrase)	
DIST	Distribution Coefficient	Numeric	
DIST.T	Temperature	Numeric	°C

Type of Indexing:

Group Code

2.6.6 System Components (MULT)

Description:

In the Gmelin database a multi-component system describes the characteristics to heterogeneous mixtures of two or more single-component compounds. The linearized formula for such a system is given by the formula of the individual single-component compounds, the "corner compounds", linked to each other by a hush (#), e.g. BaO#CuO. Phases that are formed in such a system (system components) can be retrieved in the field MULT by GRN or chemical name



Associated Parameter Fields:

	Field	Code	Full N	lame
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MULT.GRN Component Registry Number

MULT.PA Component Name(s)

Type of Indexing:

Group Code

2.6.7 Multicomponent System Data (META)

Description:

Indication of phase, solubility or melting diagrams of heterogeneous multicomponent systems is given in the field META.DGM. Indication on metastable phases of heterogeneous multicomponent systems is given in the field META.PHAS.

Note:

Searching for a diagram field code requires the field value "available", for metastabile phases the field value "described" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
META.DGM	Phase / Solubility / Melting Diagram	String (phrase)
META.PHAS	Metastable Phases	String (phrase)

Type of Indexing:

Group Code

2.6.8 Eutectic Temperature (ETEC)

Description:

The Eutectic Temperature is the temperature on which co-crystallization of two phases occurs. It is the lowest possible liquid temperature in a system containing two stable phases.

Type of	Indexing
Numeric	

String (phrase)



The liquid curve forms a singularity at this temperature in the melting diagram of a heterogeneous multicomponent system. The eutectic temperature field ETEC.T contains the values for a given substance in the literature. The concentration of the phases at the eutectic point is given in the remark field ETEC.MR.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
ETEC.T	Eutectic Temperature	Numeric	°C
ETEC.MR	Remarks	Wordwise	

Type of Indexing:

Group Code

2.6.8.1 Peritectic Temperature (PTEC)

Description:

The Peritectic Temperature is the temperature at which the melt of a heterogeneous multicomponent system begins to react with the solid phase to form a new compound. Normally this temperature is identical with the incongruent melting point of this compound. The peritectic temperature field contains the values for the hetereogeneous multicomponent system in the literature. The concentration of the phases at the peritectic point is given in the remark field PTEC.MR.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
PTEC.T	Peritectic Temperature	Numeric	°C
PTEC.MR	Remarks	Wordwise	

Type of Indexing:



2.6.9 Solubility in Systems (MULS)

Description:

The search field Solubility in Systems indicates information on solubility or miscibility in heterogeneous multicomponent systems.

Type of Indexing:

Group Code

2.6.10 Vapor Pressure of Solution (VPR)

Description:

Indication of the vapor pressure of a heterogeneous multicomponent system as a diagram, table or equation is given in the fields VPR.DGM, VPR.TAB or VPR.EQN.

Note:

Searching for special field codes require the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
VPR.DGM	Diagram	String (phrase)
VPR.TAB	Table	String (phrase)
VPR.EQN	Equation	String (phrase)

Type of Indexing:

Group Code

2.6.11 Solubility (SLB)

Description:

The Solubility of a substance is the maximum concentration of a compound in a binary mixture at a given temperature forming a homogeneous solution. Entries with the solubility unit mol/l can be found in the field SLB.



Entries with solubility units other than mol/l can be found in the field SLB.VALU. The solvent is given in the associated search field SLB.SOL. The solubility is linked to the associated temperature by the parameter field code SLB.T. A qualitative description of the solubility of a compound can be retrieved in the field SLB.TXT.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
SLB.TXT	Description of Solubility	String (phrase)	
SLB	Solubility (molar)	Numeric	mol I ⁻¹
SLB.VALA	Solubility (weight)	String (phrase)	
SLB.VALU	Solubility (other)	String (phrase)	
SLB.T	Temperature	Numeric	°C
SLB.SOL	Solvent	String (phrase)	
SLB.UNIT	Solubility Unit	String (phrase)	

Type of Indexing:

Group Code

2.6.12 PL-Value (SLBP)

Description:

The solubility product is the product of the concentrations of the ions of a substance in a saturated solution. It is contained in the search field SLBP.SOLPRO. The corresponding PL Value is given in the search field SLBP.VAL. The field SLBP.TXT additionally displays a qualitative description on solubility. The temperature at which the solubility product was measured is searchable in the parameter field SLBP.T

Field Code	Full Name	Type of Indexing	Unit
SLBP.TXT	Description of Solubility	String (phrase)	
SLBP.VAL	PL-Value	Numeric	
SLBP.SOLPRO	Solubility Product	Numeric	mol l ⁻¹
SLBP.T	Temperature	Numeric	°C
SLBP.SOL	Solvent	String (phrase)	



Type of Indexing:

Group Code

2.6.13 Surface Tension (ST)

Description:

The surfaces tend to contract to a minimum area, the surface tension is caused by cohesive forces between the molecules. It is measured as a force per unit length along the surface or the work, which must be done to extend the surface by a unit area. The Surface Tension search field contains the values for a given substance in the literature and is linked to the associated temperature ST.T. Further information about the samples is given in the field remarks ST.MR. The surface tension is linked to the forms of state of the substance by the parameter field code ST.S.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
ST	Surface Tension	Numeric	erg cm ⁻²
ST.T	Temperature	Numeric	°C
ST.MR	Remarks	Wordwise	
ST.S	Forms of State	String (phrase)	

Type of Indexing:



2.7 Quantum Chemical Calculations

2.7.1 Quantum Chemical Calculations (QCC)

Description:

The topic Quantum Chemical Calculations contains data derived from quantum mechanical methods. This field identifier is a group code for Field Availability searches in the hierarchical system of the CrossFire database.

Quantum Chemical Calculations (QUAN)

Note:

Values which have been calculated by classical (not quantum mechanical) methods or which come from experimentally measured data are not recorded here, but under the corresponding property.

2.7.2 Quantum Chemical Calculations (QUAN)

Description:

The Quantum Chemical Calculations QCC.PROP search field refers to quantum chemical calculations performed for a substance. The calculated properties are available in this field together with a classification of the corresponding quantum chemical method of calculation QCC.MET.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
QUAN.PROP	Calculated Properties	String (phrase)
QUAN.MET	Method of Calculation	String (phrase)

Type of Indexing:



2.7.3 Method of Calculation (QUAN.MET)

Description:

Following calculation methods are valid:

- ab initio calculation
- electron correlation and CI calculation
- empirical method
- further quantum calculation
- other semiempirical calculations
- relativist. calculation
- semiempirical NDO calculation
- statist. model calculation
- valence bond calculation

Type of Indexing:

String (phrase)



2.8 Electrochemistry

2.8.1 Electrochemistry (EC)

Electrolysis (ELYS) Transference Number (TRAN) Ionic Mobility (ION) Behavior As Electrode (ELTR) Electrochemical Cell (CELL) Electrochemical Reaction (POT) Electrolytic Conductivity (ELTC)

2.8.2 Electrolysis (ELYS)

Description:

Indication on a detailed description of Electrolysis, with general conditions and phenomena arising from the flow of current, is given in the field ELYS.FLG. Further data are given in the Numeric search field decomposition potential ELYS.POT.

Note:

Searching for the electrolysis field code requires the field value "described" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
ELYS.FLG	Electrolysis	String (phrase)	
ELYS.POT	Decomposition Potential	Numeric	V

Type of Indexing:



2.8.3 Transference Number (TRAN)

Description:

The transference number is the fraction of the electrical current carried by a single ion in an electrolytic solution. The field Transference Number indicates information on transference number of compounds or ions.

Type of Indexing:

Group Code

2.8.4 Ion Mobility (ION)

Description:

The ionic mobility is the migration velocity of ions in an electric field of homogenous field strength. The Ion Mobility search field is a numeric range searchable field and is linked to the associated temperature ION.T. Further information about the samples is given in the field remarks ION.MR.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
ION	Ionic Mobility	Numeric	$m^2 V^{-1} s^{-1}$
ION.T	Temperature	Numeric	°C
ION.MR	Remarks	Wordwise	

Type of Indexing:

Group Code

2.8.5 Behavior as Electrode (ELTR)

Description:

The field Behavior as Electrode indicates detailed information on compounds as electrodes or parts of an electrode.



This includes all processes, which change the electrode mechanically, optically, chemically, or physically, as well as, a description of the use and stability of the electrode, information to material composition or quality, polarization or adsorption phenomena, and overvoltage.

Type of Indexing:

Group Code

2.8.6 Electrochemical Cell (CELL)

Description:

The Electrochemical Cell potential field CELL.POT contains the values for cells using substances of the file as electrode material, as electrolyte, or as a compound of the electrolyte. The electrochemical cell potential field is a numeric range searchable field and is linked to the associated temperature CELL.T. A pair of electrodes dipping into an ionic medium called electrolyte (solid, liquid or gas) and connected by an external electric conductor constitutes an electrochemical cell. The description of cell field CELL.DESCR contains the description of cells using substances of the file as electrode material, as a compound of the electrolyte.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
CELL.DESCR	Description of Cell	String (phrase)	
CELL.POT	Cell Potential	Numeric	V
CELL.T	Temperature	Numeric	°C

Type of Indexing:

Group Code

2.8.7 Electrochemical Reaction (POT)

Description:

An electrochemical reaction is a reaction where electrons between the reaction partners and external sources are exchanged. You can search for the reaction partners by GRN or chemical name and the associated potential value and kind of potential (POT.PTYP).



The field is linked to the associated parameters temperature (POT.T), solvent (POT.SOL), method of determination (POT.MET), the reference electrode (POT.REL), and the supporting electrolyte (POT.ELYT). Further information is indicated in the search field potential diagram (POT.DGM).

Note:

Searching for a potential diagram field code requires the field value "available" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
POT.EGRN	Coreactant(s) (Registry Number)	String (phrase)	
POT.EDN	Coreactant(s) (Name)	String (phrase)	
POT.PGRN	Product(s) (Registry Number)	String (phrase)	
POT.PRO	Product(s) (Name)	String (phrase)	
POT	Potential Value	Numeric	V
POT.PTYP	Kind of Potential	String (phrase)	
POT.T	Temperature	Numeric	°C
POT.SOL	Solvent	String (phrase)	
POT.MET	Method of Determination	String (phrase)	
POT.REL	Reference Electrode	String (phrase)	
POT.DGM	Potential Diagram	String (phrase)	
POT.ELYT	Supporting Electrolyte	String (phrase)	

Type of Indexing:

Group Code

2.8.8 Electrolytic Conductivity (ELTC)

Description:

The molar (specific) electrolytic conductivity is the reciprocal of the resistivity, per mole (gram) per unit volume of solution.

The equivalent electrolytic conductivity is the molar electrolytic conductivity per number of equivalents. The field is linked to the associated parameters temperature (ELTC.T),



solvent (ELTC.SOL), and kind of conductivity (ELTC.TYP). Further information about the samples is given in the field remarks ELTC.MR.

Note:

 $S \text{ cm}^2 \text{ mol}^{-1} = 10^{-3} \text{ S cm}^{-1} \text{ I mol}^{-1}$

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
ELTC	Molar Electrolytic Conductivity	Numeric	SI cm ⁻¹ mol ⁻¹
ELTC.VALA	Specific Electrolytic Conductivity	Numeric	SIcm ⁻¹ g ⁻¹
ELTC.AEVAL	Equivalent Conductivity	Numeric	cm ² ? ⁻¹ val ⁻¹
ELTC.T	Temperature	Numeric	°C
ELTC.SOL	Solvent	String (phrase)	
ELTC.TYP	Kind Of Conductivity	String (phrase)	
ELTC.MR	Remarks	Wordwise	

Type of Indexing:



2.9 Chemical Properties

2.9.1 Chemical Properties (CHE)

Description:

The group Chemical Data contains information about:

Chemical Behavior (CHB) Reaction with Substance Classes (RCLA) Surface Reaction (SURF) Reaction (RX) Reaction Details (RX)

Tips and Hints:

The Reactions are stored in the facts Reaction Identification Data and Reaction Details. These reactions have been registered and hence assigned a Reaction Identification Number.

2.9.1.1 Chemical Behavior (CHB)

Decomposition (DP) Information on Stability (STAB) Flash Point (FLAP) Behavior as Catalyst (CAT) Behavior as Inhibitor (BINH) Behavior as Acid / Base (ACBS) Colloid Chemical Behavior (COLL) Polymerisation (POLY) Association (ASS) Solvation / Hydration (SOHY)



2.9.1.2 Decomposition (DP)

Description:

The decomposition point is the temperature at which a substance decomposes thermally. The Decomposition Point search field is a numeric range searchable field. Decomposition in solutions is linked to the associated solvent by the DP.SOL search field.

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing	Unit
DP	Temperature	Numeric	°C
DP.SOL	Solvent	String (phrase)	

Type of Indexing:

Group Code

2.9.1.3 Information on Stability (STAB)

Description:

The search field Information on Stability displays a description of the stability, as well as, the behavior of a substance influenced by light, moisture or air.

Associated Parameter Field:			
Field Code	Full Name	Type of Indexing	
STAB	Description of Stability	String (phrase)	

Type of Indexing:



2.9.1.4 Flash Point (FLAP)

Description:

The Flash Point is the lowest temperature at which a liquid gives off sufficient vapor to form an ignitable mixture with air near the surface of the liquid or within the vessel used.

Unit: °C

Type of Indexing:

Numeric

2.9.1.5 Behavior as Catalyst (CAT)

Description:

The field Behavior as Catalyst indicates information and references of a compound catalysing organic or inorganic reactions.

Note:

You can search for the corresponding inorganic reaction, which is catalysed, using the field RX.CGRN or RX.CAT.

Type of Indexing:

Group Code

2.9.1.6 Behavior as Inhibitor (BINH)

Description:

The field Behavior as Inhibitor indicates the behavior of a compound as an inhibitor. They can be designated as reaction inhibitors, negative catalysts, or radical scavengers.

Type of Indexing:



2.9.1.7 Behavior as Acid / Base (ACBS)

Description:

The field Behavior as Acid / Base indicates the behavior of a compound as an acid or base. The acid-base behavior is the object of the article, or how the pH, pK_{s} , or pK_{B} values are determined.

Type of Indexing:

Group Code

2.9.1.8 Colloid Chemical Behavior (COLL)

Description:

The field Colloid Chemical Behavior indicates information on colloids, gels, sols, aerosols smoke or mist.

Type of Indexing:

Group Code

2.9.1.9 Polymerisation (POLY)

Description:

The field Polymerisation indicates information on the polymerisation or oligomerisation of a compound. Only the polymerisation of a substance with itself is recorded, no co-polymerisation.

Type of Indexing:

Group Code

2.9.1.10 Association (ASS)

Description:

The field Association indicates information on the association of the molecules of a compound.



Type of Indexing:

Group Code

2.9.1.11 Solvation / Hydration (SOHY)

Description:

The field Solvation / Hydration indicates information on the solvation or hydration of a compound. The focus is the substance without solvate molecules.

Type of Indexing:



2.9.2 Reaction with Substance Classes (RCLA)

Description:

Indication on the chemical behavior, against classes of compounds, is given in the fields RCLA.REAC and RCLA.NOREAC. The classes of substances are specified in the field RCLA.CLASS. The product substance classes are specified in the field RCLA.PROD.

Note:

Searching for "does react" and "does not react" field codes requires the field value "yes" or more general "*".

Associated Parameter Fields:

Field Code	Full Name	Type of Indexing
RCLA.REAC	does react	String (phrase)
RCLA.NOREAC	does not react	String (phrase)
RCLA.CLASS	reaction with	String (phrase)
RCLA.PROD	reaction product	String (phrase)

Type of Indexing:

Group Code

2.9.3 Surface Reaction (SURF)

Description:

The field Surface Reaction indicates chemical changes (reactions) of the surface of a substance. The formation of surface compounds is described. The focus is on the substance where the surface reaction takes place.

Type of Indexing:



2.9.4 Reaction (RX)

Description:

All chemical syntheses and reactions of inorganic or organometallic compounds can be retrieved in the RX fields. The Reaction Identification Data are stored in six fields. These contain the keys to the reactants and products, which can be searched for by GRN, name, or by structure (if available). The parameter data to any reaction, such as solvent, catalyst, and further conditions, is stored in the corresponding fact Reaction Details.

Reaction Identification Fields:

Field Code	Full Name	Type of Indexing
RX.ID	Reaction ID	Numeric
RX.RGRN	Reactant(s) (GRN)	Numeric
RX.RCT	Reactant(s) (Name)	String (phrase)
RX.PGRN	Product(s) (GRN)	Numeric
RX.PRO	Product(s) (Name)	String (phrase)
RX.NVAR	Number of Reaction Details	Numeric
RX.BCODE	Reaction Code (broad)	Numeric
RX.MCODE	Reaction Code (medium)	Numeric
RX.NCODE	Reaction Code (narrow)	Numeric
RX.ED	Reaction Entry Date	String (phrase)
RX.UPD	Reaction Update Date	String (phrase)

Note

Phase transformations are stored in Transition Point Solid / Solid.

2.9.4.1 Reaction ID (RX.ID)

Description:

The Reaction ID is assigned when a chemical reaction is registered in the Gmelin Information System. The Reaction ID is an unambiguous identifier of a reaction. The Reaction ID is the accession number of the reaction record in CrossFire. It is an integer, which does not contain further information.

Type of Indexing:



Numeric

2.9.4.2 Reactant(s) (GRN) (RX.RGRN)

Description:

Reactants (starting materials) are defined as compounds that form the product(s). Inorganic starting materials are registered compounds and are therefore present as title compounds with data. Organic starting materials have usually no further data in the Gmelin database; they will be present and have minimal "identification" data in addition to the structure. Starting materials are searchable either by GRN (in the RX.RGRN field) or by names (in the RX.RCT field).

Type of Indexing:

Numeric

Example:

Searching for substances that have been prepared from copper(II) sulfide (GRN unknown).

Step 1: Searching for GRN of copper(II) sulfide in substance context:

Operator	Field Name	Value
	MOFO	CuS
AND	NC	1

Display of Identification Data including the GRN. The first hit describes the usual CuS compound; GRN = 13690.

Step 2: Searching for reactions with copper(II) sulfide as a starting material in reaction context:

Operator	Field Name	Value
	RX.RGRN	13690

Tips and Hints:

Searching for reactions by GRN is useful for compounds having no molecular or no completely defined molecular structure. So you can tailor all reactions of a title compound into preparations and reactions.



You can perform reaction searching for compounds with full structures directly in the structure editor using the reaction editmode. Here you can search with full structures or substructures.

2.9.4.3 Reactant(s) (Name) (RX.RCT)

Description:

Reactants (starting materials) are defined as compounds that form the product(s). Inorganic starting materials are registered compounds and are therefore present as title compounds with data. Organic starting materials have usually no further data in the Gmelin database, they will be still present and have minimal "identification" data in addition to the structure. Starting materials are searchable either by GRN (in the RX.RGRN field) or by names (in the RX.RCT field). The search field contains formulas and, if present, chemical names.

Tips and Hints:

Name searches are not recommended to identify compounds, because names are ambiguous or non-systematic and complete chemical names are not available in many cases. In some cases, name fragments might be useful to search for substance classes. However there are more powerful methods available such as structure or substructure searches, or GRN search.

Type of Indexing:

String (phrase)

2.9.4.4 Product(s) (GRN) (RX.PGRN)

Description:

Reaction products are registered compounds and are therefore present as title compounds with data. The main focus of the Gmelin database is the formation of inorganic and organometallic compounds. Reaction products are searchable either by GRN (in the RX.PGRN field) or by names (in the RX.PRO field).

Type of Indexing:

Numeric

Example:

Searching for preparations of Bi₂Sr₂CaCu₂O₈ (GRN unknown). Step 1: Searching for GRN of calcium titanate in substance context:



Operator	Field Name	Value
	MOFO	Bi2Sr2CaCu2O8
AND	NC	1

Display of Identification Data including the GRN. The first hit is $Bi_2Sr_2CaCu_2O_8$ without further specification; the GRN is 24127.

Step 2: Searching for reactions with copper(II) sulfide as a starting material in reaction context:

Operator	Field Name	Value
	RX.PGRN	24127

Tips and Hints:

Searching for reactions by GRN is useful for compounds having a partially defined or undefined molecular structure. You can tailor all reactions of a title compound into preparations and reactions.

You can perform reaction searching for compounds with full structures directly in the structure editor using the reaction editmode. Here you can search with full structures or substructures.

2.9.4.5 Product(s) (Name) (RX.PRO)

Description:

Reaction products are registered compounds and are therefore present as title compounds with data. The main focus of the Gmelin database are the formation of inorganic and organometallic compounds. Reaction products are searchable either by GRN (in the RX.PGRN field) or by names (in the RX.PRO field).

Note:

The search field contains formulas and, if present, chemical names additionally.

Tips and Hints:

Name searches are not recommended to identify compounds, because names are ambiguous or non-systematic and complete chemical names are not available in many cases. In some cases, name fragments might be useful to search for substance classes. However there are more powerful methods available such as structure or substructure searches, or GRN search.



Type of Indexing:

String (phrase)

2.9.4.6 Number of Details (RX.NVAR)

Description:

This field gives the number of reaction details.

Note:

At the moment the RX.NVAR field is always 1, because the reaction details are linked to one registered reaction.

Type of Indexing:

Numeric

2.9.5 Reaction Details (RX)

Reaction Detail Fields:

Field Code	Full Name	Type of Indexing
RX.NYD	Yield (Numeric)	Numeric
RX.YD	Yield	String (phrase)
RX.RGT	Reagent	String (phrase)
RX.IGRN	Intermediate(s) (GRN)	Numeric
RX.INT	Intermediate(s) (Name)	String (phrase)
RX.BPRO	Byproduct	String (phrase)
RX.SOL	Solvent	String (phrase)
RX.CGRN	Catalyst(s) (GRN)	Numeric
RX.CAT	Catalyst(s) (Name)	String (phrase)
RX.RCS	Index Reagents/Catalysts/Solvents	word and phrase



RX.IND	Industrial Synthesis	String (phrase)
RX.SCON	Special Conditions	String (phrase)
RX.GCON	General Conditions	String (phrase)
RX.ISOL	Purification / Isolation	String (phrase)
RX.KIN	Kinetics	String (phrase)
RX.DED	Entry Date	String (phrase)

2.9.5.1 Yield (Numeric) (RX.NYD)

Description:

The yields of products are stored in this field. Only the absolute yield for the pure product quoted by the author is accepted. In this field the yield is indexed as a numeric value. You can search with single values or ranges, as well as, greater than / less than a certain value.

Type of Indexing:

Numeric Examples:

Search for reactions with product yields greater than 98%

Field Name	Value

RX.NYD >98

Search for reactions with product yields between 84 and 85 %

Field Name	Value
RX.NYD	84-85

Search for reactions with product yields less than 5 %

Field Name	Value
RX.NYD	<5

Note:



The yield 0 % means that the desired product was not formed. If only a trace of a product is formed the yield is stored as <1 %. Quantitative yields are expressed as >99 % yield.

Related Information:

RX.YD

2.9.5.2 Yield (RX.YD)

Description:

The yields of products are stored in this field. Only the absolute yield for the pure product quoted by the author is accepted. In this field the yield is indexed as a String. This means the reaction yield has to match exactly with the entered String for a hit.

Type of Indexing:

String (phrase)

Examples:

Search for reactions with product yields greater than 98%

Field Name	Value
RX.YD	>98

Search for reactions with product yields between 84 and 85 %

Field Name	Value
RX.YD	84-85

Search for reactions with product yields less than 5 %Field NameValue

RX.NYD <5

Note:

The yield 0 % means that the desired product was not formed. If only traces of a product is formed the yield is stored as <1 %. Quantitative yields are expressed as >99 % yield. If mathematic signs are added for a search, the input has to be set in quotation marks.



Related Information:

RX.NYD

2.9.5.3 Reagent (RX.RGT)

Description:

Reagents are compounds, which take part in the reaction but do not make a contribution to the product, e.g. compounds for pH adjustment, reducing agents, or oxidizing agents. Further undefined compounds, which cannot be expressed by formula but only by name, are also available in this field, like air, charcoal, or Rh-catalyst supported on SiO2 (Rh/SiO2). Reagents have no GRN.

Note:

Alternative agents are linked by "or".

Type of Indexing:

String (phrase)

2.9.5.4 Intermediate(s) (GRN) (RX.IGRN)

Description:

Intermediates are stored as non-isolated compounds formed in a one-pot reaction, or compounds of intermediate steps for which measured data are available. Intermediates are searchable either by GRN (in the RX.IGRN field) or by names (in the RX.INT field).

Type of Indexing:

Numeric



2.9.5.5 Intermediate(s) (Name) (RX.INT)

Description:

Intermediates are stored as non-isolated compounds formed in a one-pot reaction, or compounds of intermediate steps for which measured data are available. Intermediates are searchable either by GRN (in the RX.IGRN field) or by names (in the RX.INT field).

Tips and Hints:

Name searches are not recommended to identify compounds, because names are ambiguous or non-systematic and complete chemical names are not available in many cases. In some cases name fragments might be useful to search for substance classes.

Type of Indexing:

String (phrase)

2.9.5.6 Byproduct (RX.BPRO)

Description:

By-products are compounds, which are produced during the reaction but have no focus or are not characterized. By-products can be precipitated salts like AgCl, formed gases like CO_2 , or organic compounds.

Type of Indexing:

String (phrase)

2.9.5.7 Solvent (RX.SOL)

Description:

This field contains information on the solvent, or the solvent mixture, used in the reaction. If solvents are also reaction partners, they are stored as reactants. There are some special solvent descriptions available:

For reactions in melt:	melt
For reactions without solvent: e.g. at higher temperatures	neat (no solvent)
Solid-state reactions:	neat (no solvent, solid phase)



Gas-phase reactions:

neat (no solvent, gas phase)

Type of Indexing:

String (phrase)

2.9.5.8 Catalyst(s) (GRN) (RX.CGRN)

Description:

Catalysts are compounds that do not appear in the stoichiometric equation of the reaction. They only influence the rate of the reaction. In the Gmelin database catalysts are inorganic or organometallic compounds and neither identical with a starting material nor an acid-base catalyst. Catalysts are searchable either by GRN (in the RX.CGRN field) or by names (in the RX.CAT field).

Type of Indexing:

Numeric

2.9.5.9 Catalyst(s) (Name) (RX.CAT)

Description:

Catalysts are compounds that do not appear in the stoichiometric equation of the reaction. They only influence the rate of the reaction. In the Gmelin database catalysts are inorganic or organometallic compounds and neither identical with a starting material nor an acid-base catalyst. Catalysts are searchable either by GRN (in the RX.CGRN field) or by names (in the RX.CAT field).

Tips and Hints:

Name searches are not recommended to identify compounds, because names are ambiguous or non-systematic and complete chemical names are not available in many cases. In some cases name fragments might be useful e.g. to search for substance classes.

Type of Indexing:



2.9.5.10 Industrial Synthesis (RX.IND)

Description:

The field Industrial Synthesis indicates the corresponding reaction as an industrial reaction.

Note:

Searching for industrial synthesis field code requires the field value "yes" or more general "*".

Type of Indexing:

String (phrase)

2.9.5.11 Special Conditions (RX.SCON)

Description:

This field contains information on special reaction conditions related to a list of controlled terms.

Following special conditions are valid:

- electric arc
- electrochemical process
- electrolysis
- high pressure
- irradiation (uv/vis)
- other radiation
- sonication

Type of Indexing

String (phrase)

2.9.5.12 General Conditions (RX.GCON)

Description:

The field General Conditions contains a short description of the performance of the reaction. Details like protective gas, reaction time, temperature, pressure, or special performance of a reaction are described.



Type of Indexing:

wordwise

2.9.5.13 Purification / Isolation (RX.ISOL)

Description:

The field Purification / Isolation contains a short description of the purification/isolation of the reaction products. Details to methods of separation and purification are described. Further elemental analysis or purity details of the products are indicated. Methods of identification are noted especially for compounds, which are not isolated.

Type of Indexing:

wordwise

2.9.5.14 Kinetics (RX.KIN)

Description:

The field Kinetics indicates descriptions of kinetics of the reaction, e.g. rate constants, reaction order or activation energies.

Note:

Searching for kinetic field code requires the field value "available" or more general "*".

Type of Indexing:

String (phrase)

2.9.5.15 Reaction Update Date (RX.UPD)

Description:

The search field Reaction Update Date contains the date (year/month/day) when the last update of this reaction was entered into the database. Dates have to be searched with a fixed format: yyyy/m m/dd.

Type of Indexing:



2.10 Bibliographic Information

2.10.1 Bibliographic Information (BIB)

Citation Number (CNR)

Citation (CIT)

Abstract (AB)

2.10.1.1 Citation Number (CNR)

Description:

Every reference in the Gmelin Database has been assigned a citation number, which can be used to access a certain citation directly.

Type of Indexing:

Numeric

2.10.1.2 Citation Entry Date (CED)

Description:

The search field Citation Entry Date contains the date (year/month/day) when the citation was first entered into the database. Dates have to be searched with a fixed format: yyyy/mm/dd.

Type of Indexing:



2.10.1.3 Citation Update Date (CUPD)

Description:

The search field citation Update Date contains the date (year/month/day) when the last update of this citation was entered into the database. Dates have to be searched with a fixed format: yyyy/mm/dd.

Type of Indexing:

String (phrase)

2.10.2 Citation

2.10.2.1 Citation (CIT)

```
Author (AU)
CODEN (CO)
Journal Title (JT)
Publication Year (PY)
Language (LA)
CODEN (Translation) (COT)
Journal Title (Translation) (JTT)
Publication Year (Translation) (PYT)
Language (Translation) (LAT)
```

2.10.2.2 Author (AU)

Description:

The Author field contains the surname of the author, the first name or abbreviations as they were written in the original publication. For publications with more than one author each author name will be separately indexed. Use LIST to look in the index and see exactly how a particular author has been cited. The surname is always separated by a comma and a space from first names or affixes e.g. "van der".



Titles such as "Dr.", "Sir" or "Prof." are ignored. The five names mentioned in the article are given, further names are displayed as "et al."

Authors can be indexed as follows:

Adams, Richard D. Arduengo, Anthony J. III Baudler, Marianne Cotton, F. Albert Schnering, H. G. von Simon, A.

Tips and Hints:

To ensure that all references from a particular author are found it is recommended that the LIST function is used to check the way the author has been cited. Thus it is often useful to use truncation when author searches are carried out and restrict the search to surname only.

The following examples shows the effect of truncation on the search functionality ot authors names:

Type of Indexing:

String (phrase)

Example1:

Search for articles published by authors with surname Adams

Field Name	Value
AU	Adam,*

Search for articles published by all authors beginning with Adam

Field Name	Value
AU	Adam*

Search for articles published by Richard Adams

Field Name	Value
AU	Adams, Richard



Search for articles published by Richard Adams including other first names

Field Name	Value
AU	Adams, Richard*

Search for articles published by R. Adams – including Richard Adams – or other first names starting with R

Field Name	Value
AU	Adams, R*

Example2:

Using the PROXIMITY operator will only retrieve those publications of F. Albert Cotton from Polyhedron (PLYHDE):

Operator	Field Name	Value
	AU	Cotton, F. Albert
PROXIMITY	СО	PLYHDE

Related Information:

PY, CO, JT, LA

2.10.2.3 CODEN (CO)

Description:

Journal titles may be searched for using the CASSI CODEN. CASSI CODENs are unique, unambiguous, six-character codes assigned to journals and other serial and non-serial publications. They are used instead of the abbreviated journal titles to search for references from a particular publication.

Type of Indexing:



Example:

Search for substances published in the journals Inorg. Chim. Acta and J. Mol. Catal. A Chem.:

Operator	Field Name	Value	
	CO	ICHAA3	
OR	CO	JMCCF2	

Related Information:

AU, PY, JT, LA

2.10.2.4 Journal Title (JT)

Description:

The title of the journal is present in this field if it could be assigned via the CODEN. You can request a list of the journal titles if you use the LIST function in the citation query input mask.

Type of Indexing:

String (phrase)

Related Information:

AU, PY, CO, LA

2.10.2.5 Publication Year (PY)

Description:

The content of the Publication Year field is the publication year of the article cited. It should not be confused with the entry date.

Type of Indexing:

Numeric



Example:

Search for all substances reported in 2000 in the journal J. Orgmet. Chem. (JORCAI):

Operator	Field Name	Value
	PY	2000
PROXIMITY	CO	JORCAI

Related Information:

AU, CO, JT, LA

2.10.2.6 Language (LA)

Description:

The Language of the original publication is contained in this field. You can see the valid languages by using the LIST function. All data in the Gmelin database is in English. This field will tell you the language of the original article.

Type of Indexing:

String (phrase)

Related Information:

AU, PY, CO, JT

2.10.2.7 CODEN (Translation)(COT)

Description:

The Gmelin database contains also some journals, which are originally published in a different language than English, e.g. Russian. These journals are excerpted with their translation, usually an English translation.

For these journals the field CODEN (Translation) contains the CASSI CODEN of the translation. The CASSI CODEN of the original journals is listed in the field CODEN (CO).

Type of Indexing:



Example:

Searching for substances published in the journal Inorganic Materials (Transl. of Neorg. Mater.):

Operator	Field Name	Value
	СОТ	INOMAF

Related Information:

CO, JTT, PYT, LAT

2.10.2.8 Journal Title (Translation) (JTT)

Description:

The Gmelin database contains also some journals, which are originally published in a different language than English, e.g. Russian. These journals are excerpted with their translation, usually an English translation. Then the field Journal Title (Translation) contains the title of a journal of the translation. The journal title of the original is listed in the field Journal Title (JT).

Type of Indexing:

String (phrase)

Related Information:

JT, COT, PYT, LAT

2.10.2.9 Publication Year (Translation)(PYT)

Description:

The field Publication Year (Translation) refers to the publication year of an article in a journal processed as translation.

Type of Indexing:

Numeric



Related Information:

PY, COT, JTT, LAT

2.10.2.10 Language (Translation)(LAT)

Description:

This field contains the language of a journal, which is processed as translation. Usually the language of a translated journal is English.

Type of Indexing:

String (phrase)

Related Information:

LA, COT, JTT, PYT

2.10.3 Abstract

2.10.3.1 Abstract (AB)

Description:

The content of the abstract field is taken from the original literature from 1995 to the present. However, there are also sometimes abstracts available from journals before 1995.

Tips and Hints:

The Boolean Operators AND, OR, NOT can be used to find occurrences of the indicated terms anywhere in the abstract. In the case of PROXIMITY the entire abstract is considered as information unit. Therefore, the PROXIMITY operator works like AND. You can link words using the Operators NEAR or NEXT.

Search Fields:

Title (TI) Abstract (AB) Abstract Language (ALA) Keywords (AB.KW)



2.10.3.2 Title (TI)

Description:

The content of the Title field is taken from the original literature from 1995 to the present. However, there are also titles available from journals before 1995.

Tips and Hints:

The Boolean Operators AND, OR, NOT can be used to find occurrences of the indicated terms anywhere in the abstract. In the case of PROXIMITY the entire abstract is considered as information unit. Therefore, the PROXIMITY operator works like AND. You can link words using the operators NEAR or NEXT.

Type of Indexing:

String (wordwise)

Example1:

Search for all documents where bioactive substances are determined:

Field Name	Value
------------	-------

TI Bioactive

Example2:

Search for documents where a relationship between the structure of drugs has been published:

Operator	Field Name	Value
	ТІ	structur*
AND	ТΙ	drug*

2.10.3.3 Abstract (AB)

Description:

The content of the Abstract field is taken from the original literature from 1995 to the present. However, there are also abstracts available from journals before 1995.

Tips and Hints:



The Boolean Operators AND, OR, NOT can be used to find occurrences of the indicated terms anywhere in the abstract. In the case of PROXIMITY the entire abstract is considered as information unit. Therefore, the PROXIMITY operator works like AND. You can link words using the operators NEAR or NEXT.

Type of Indexing:

String (wordwise)

Examples:

Search for all documents where anticancer or antimetastatic substances are described:

Operator	Operator Field Name Value	
	AB	Anticancer
OR	AB	antimetastat*

Search for documents with the phrase catalytic activity in the abstract:

Operator	Field Name	Value
	AB	catalyt*
AND	AB	Activity

Search for documents concerning with p-type semiconductors:

Operator	Field Name	Value
	AB	Р
NEXT	AB	Туре
AND	AB	semiconductor*

2.10.3.4 Abstract Language (ALA)

Description:

The Abstract Language field contains the language of the literature abstract. In general, the abstract is available in English.

Type of Indexing:



2.10.3.5 Keywords (AB.KW)

Description:

This field contains keywords, index headings or similar items, which are present in an article.

Type of Indexing:

String (Wordwise)

Example1:

Search for all documents with keyword photochemistry or similar phrases:

Field Name	Value

AB.KW photochem*

Example2:

Search for documents dealing with kinetics and catalysis (including variations):

Operator	Field Name	Value	
	AB.KW	kinetic*	
AND	AB.KW	*catal*	



3 Gmelin Special Features

3.1 Gmelin - Special Features

3D Structures Stereochemical Information Polyhedra Gmelin Ligand Search System

3.2 3D Structures

The Gmelin database contains a large collection of inorganic and organometallic compounds. Most of them are available with structures. However, you should be aware that only compounds or compound parts, which are formed by discrete molecules or ions, have a structure; e.g. all molecules or coordination compounds. Compounds with a solid-state structure, alloys, glasses, or ceramic materials do not have a structure in the Gmelin database.

Many of the structures in the Gmelin database have been entered with three dimensional coordinates. This is necessary to describe stereochemistry in coordination compounds correctly. Because of the more complex stereocenters in inorganic chemistry, which cannot be expressed by normal up and down bonds, the stereochemical description in the Gmelin database is different from the Beilstein database.

Certain inorganic and organometallic structures are displayed three-dimensionally. To get a real impression of a 3D structure you need to be able to rotate this structure along the three axis of the coordination system. This is possible with the structure editor (CrossFire Structure Editor or ISIS/Draw). Copy a structure from the Results tab to the Query tab of the Commander and launch the structure editor.

CrossFire Structure Editor: Select the menu item "3D Rotate" from Editmode menu. Use the Rotate tool from the tool bar to rotate the selected objects.

ISIS/Draw: Select the 3D Rotate tool from the tool bar to rotate an object. MDL/Draw: 3D Rotate feature not yet available.



3.3 Stereochemical Information

The structures of fragments or compounds in the Gmelin file are defined by the connectivity of atoms, the total charge and by stereodescriptors which mark a certain polyhedron. Two principal different stereochemical aspects can be searched for:

Stereochemical information on bonds:

Specified double bond information (cis or trans) can be searched for if the bond attribute "Double Steric" is chosen in the structure editor, and the structure is drawn as desired. Additionally, stereoinformation on single bonds (eclipsed and staggered-configuration) is stored in the database, e.g. for dipyridyl derivates. However, these descriptors are not searchable.

Stereochemical information on centers:

Stereochemical information on centers can be searched for by adding stereo center search attributes on a chiral atom. These attributes are expressed by polyhedra. In the case of trigonal pyramidal, tetrahedral, or octahedral centers, a stereodescriptor is set if one, or more than one, possible configuration of a stereocenter can be assigned. In all other cases the mentioning of a polyhedral center within a compound in the original literature is sufficient for the designation of a polyhedral descriptor. The polyhedral descriptor fix also the ligand configuration. If the configuration is unknown, no descriptor is set.

All stereocenters are drawn three-dimensionally. Fragments with stereochemical information are assigned either an absolute or a racemic configuration. Further, chiral isomers with at least a known sign of optical rotation, but unknown polyhedral geometry, are distinguished. However, these descriptors are not searchable. They are displayed as (+)- or (-)-enantiomers, respectively.

Stereochemical Search:

It is recommended to use the CrossFire Structure Editor for stereochemical search. In the structure editor, coordination polyhedras can be specified if the edit mode is set to "Inorganic". A exact stereochemical search can only be done with structures built up by polyehedra from the template "polyatom". These 3D polyehedra have a fixed numbering of the ligand positions. Relating to these reference polyehedra the desired isomer can be structured. In addition the attribute "stereo" (absolute, relative, racemic) in the "query options" menu had to be chosen to perform stereo searching."



3.4 Polyhedra

Another feature of the structure editor and the CrossFire server, in combination with the Gmelin database, is the capability to enter stereo center search attributes for coordination compounds. These attributes are called "polyeder". They can be entered in the atom attribute dialog box of the structure editor, if the edit mode is set to "Inorganic". The following descriptors are available to describe polyhedra in the Gmelin file:

Simple Polyhedra

Descriptor	Polyhedron
P-3	trigonal pyramid
T-4	tetrahedron
SP-4	square planar
SP-5	square pyramid
TB-5	trigonal bipyramid
OC-6	octahedron
TP-6	trigonal prism
PB-7	pentagonal bipyramid
CU-8	cube
SA-8	square antiprism
DD-8	dodecahedron
HB-8	hexagonal bipyramid
TPS-9	tricapped trigonal prism
HB-9	heptagonal bipyramid
OC-6-PI	octahedron (with p ligand)

These polyhedra can be used for a stereochemical search in combination with structures built up from the polyatom template.



Two polyhedra, linked by a chain of bonds

Descriptor	Polyhedron	
22E	two triangles	eclipsed
22S	two triangles	staggered
23E	triangle plus tetrahedron	eclipsed
23S	triangle plus tetrahedron	staggered
33E	two tetrahedra	eclipsed
33S	two tetrahedra	staggered
24E	triangle plus square pyramid	eclipsed
24S	triangle plus square pyramid	staggered
25E	triangle plus octahedron	eclipsed
25S	triangle plus octahedron	staggered
44E	two square pyramids	eclipsed
44S	two square pyramids	staggered
45E	square pyramid plus octahedron	eclipsed
45S	square pyramid plus octahedron	staggered
55E	two octahedra	eclipsed
55S	two octahedra	staggered

In contrast to the "simple polyhedra" these polyhedra are only displayable.



3.5 Gmelin Ligand Search System

The fact editor allows the formulation of queries, which gives you fast and easy access to classes or coordination compounds via ligand codes. This Gmelin Ligand Search System describes the elements (in groups) in combination to the number of the atoms that are connected to the central metal atom(s).

Complete ligand molecular formulas are available for all structured compounds or fragments which contain at least one metal atom and at least one coordinating ligand atom of the classes "L, A, D, Q, X" or a "special" ligand. Single ligands are indexed for complete compounds (LIGFO) and also for single centers in multinuclear compounds (LIG.LIGMF).

The codes include the following elements:

Code	Atom					
L	С					
А	В	Si	Ge			
D	Ν	Ρ	As	Sb		
Q	0	S	Se	Те		
Х	Н	F	CI	Br	Ι	At

There are the following "special" ligands:

Code CO CS CN CNS CNO CNR *) *) The substituent R has no further bond to metal atoms.

If a ligand is described by a combination of the codes, they are ordered alphabetically and the denticity of the code is put in parentheses in front of the code. The ligand formula describes only the denticity of the ligand not the connectivity to the metal center(s).



Examples:

Search for	Field Name	Value
halogenid or hydride ligand μ-halogenide or μ-hydride ligand	LIGFO	Х
ligand connecting by two halogenide or hydride atoms e.g. η^2 -hydrogen, η^2 -BH ₄	LIGFO	(2)X
(monodentate) carbonyl	LIGFO	CO
ligand connecting by C and chalcogen e.g. $\mu\text{-}\eta^2\text{-}carbonyl$ (by C and O)	LIGFO	LQ
ligand connecting to metal(s) by C e.g. μ -carbonyl (only by C), bridged alkyl-, $\eta^{\rm 2}$ alken-, $\eta^{\rm 2}$ -alkin ligands	LIGFO	(2)L
ligand connecting by three C atoms e.g. $\eta^{\scriptscriptstyle 3}\mbox{-allyl}, \eta^{\scriptscriptstyle 1}\mbox{-}y\mbox{-}\eta^{\scriptscriptstyle 2}\mbox{-}$ alkenyl	LIGFO	(3)L
ligand connecting by four C atoms e.g. η ³ -butadiene, norbornadiene, cycloocta1,5-diene	LIGFO	(4)L
ligand connecting by five C atoms to one or more metal centers e.g. η ^₅ -cyclopentadienyl	LIGFO	(5)L
ligand connecting by six C atoms e.g. $\eta^{\mbox{\tiny 6}}\mbox{-cyclobenzene}$	LIGFO	(6)L
ligand connecting by ten C atoms to one or more metal centers e.g. two bridged $\eta^{\mbox{\tiny 5}}\mbox{-cyclopentadienyl}$	LIGFO	(10)L
ligand connecting by twelve C atoms e.g. $\eta^{\scriptscriptstyle 6}\!,\eta^{\scriptscriptstyle 6}\!\!-\!$ cyclophanes	LIGFO	(12)L
ligand connecting by two chalcogenides e.g. η^2 - or μ -acides (like acetic acid), oxalate (bidentate), di-oles (like ethylene glycol), salicylates, acetylacetonates, dithiols, μ -bisulfido	LIGFO	(2)Q
ligand connecting by three chalcogenides e.g. tri-ole, µ-carbonato, bis(alkylthio)-sulfide, 2,6-diformyl-phenolate	LIGFO	(3)Q
ligand connecting by four chalcogenides e.g. silicate, phosphate, 12-crown-4, tetra-thiocyclohexadecane, oxalate (tertadentate)	LIGFO	(4)Q



ligand connecting by six chalcogenide atoms e.g. 18-crown-6, bridged TeOն, SbOն, IOն	LIGFO	(6)Q
ligand connecting by two pnictide atoms e.g. ethylenediamine, η ² -nitrogen, bipyridine, di-substituated phosphinoalkans (like dppm, dppe), di-substituated arsine (like diars)	LIGFO	(2)D
ligand connecting by three pnictide atoms e.g. terpyridine, tri-aza-cyclononane, tris-pyrazolyl-borates, triphos	LIGFO	(3)D
ligand connecting by four pnictide atoms e.g. porphyrins, phthalocyanines, tetra-aza-cyclam	LIGFO	(4)D
ligand connecting by five pnictide atoms e.g. cobalamin	LIGFO	(5)D
ligand connecting by one pnictide und one C atom e.g. benzylamine, phenylpyridine	LIGFO	DL
ligand connecting by one pnictide und two chalcogenide	LIGFO	D(2)Q
atoms e.g. diole-amines, imino-diacetates		D(2)Q
ligand connecting by two pnictide und two chalcogenide atoms e.g. disalicylidene-diiminato ligands (like salen)	LIGFO	(2)D(2)Q
ligand connecting by two pnictide und four chalcogenide atoms e.g. ethylenediamine-tetraacetate	LIGFO	(2)D(4)Q
ligand connecting by two pnictide und four chalcogenide atoms e.g. cryptand (222)	LIGFO	(2)D(6)Q
ligand connecting by three group III and two C atoms e.g. carboranes (like close- $C_2B_9H_{11}$)	LIGFO	(3)A(2)L

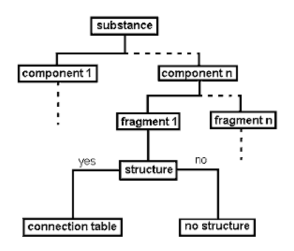


4 Guidelines for Searching

4.1 Guidelines for the Searching of Compounds in the Gmelin Database with CrossFire - Registration of Compounds

In order to get good retrieval results on searching for compounds in the GMELIN database with CrossFire, the user should have some knowledge of how these compounds are stored.

Substance Hierarchy in the GMELIN file:



General Information on Searching for Compounds Compound Formulas with more than one Component Compound Formulas with more than one Fragment General Information on Structures Stereochemistry Identical Molecular Formula - New Compound?



4.2 General Information on Searching for Compounds

In general compounds can be searched using compound formulas, formula data, or structures. Combined searches are also possible.

Searching for compound formulas:

- by code MOFO. Here the formula can be entered in the usual chemical description, e.g. NaCl, K2SO4, FeCl3*6NH3, Fe(C5H5)2. Index ranges can also be used in formulas, e.g. BaCuO(1-3), PN(1-10). Asterisks at the end of a formula increase the formula search by further elements, e.g. Ba2Cu3O7*. Combinations are also possible, e.g. Y(0-1)Ba2Cu3O(6-8)*. It is not possible to display this field via the expand command.
- by code MF. Here the molecular formula has to be entered in the "Hill order", i.e. "carbon" first, "hydrogen" next, and then all further elements in alphabetic order, e.g. C10H10Fe for ferrocene. Formulas of compounds without "carbon" are always entered in alphabetic order, e.g. CINa for sodium chloride, K2O4S for potassium sulfate. Consequently, an input "hf" means the element "hafnium"; hydrofluoric acid must be entered as "fh".
- by code LSF. Compounds can consist of more than one fragment being separated in the linearized structure formula field by an asterisk (see Compound Formulas with more than one Fragment)

Note:

To tailor the search to a one component compound the formula search should additionally be restricted by the number of components (nc=1).

Searching for formula data:

- by code ELS. The symbol of each distinct element of the molecular formula can be entered.
- by code ELC. The respective count of the distinct elements in a molecular formula can be fixed.
- by code NE. The total number of elements of the molecular formula can be restricted.
- by code NA. The total number of atoms of the molecular formula can be restricted.

Note:

To tailor the search to a one component compound the formula search should additionally be restricted by the number of components (nc=1).



Searching for full / sub structures:

- Structures even complex structures are easily drawn using the structure editor.
- Predefined templates on the menu bar or in menu "file" are available.
- Atom attributes can be specified, like atom symbols, atom charge, atom mass.
- Free sites on atoms vary the bondings to this atom. If "free sites" remain zero, no further bondings are considered as drawn. If the number of "free sites" is changed, the coordination sphere of the atom is increased further by the respective count.
- If the edit mode is set to "Inorganic", coordination polyhedras can be specified.
- In the "query options" menu structure related attributes can be set and stereo searching can be performed.

4.3 Compound Formulas with more than one Component

These are heterogeneous multi-component compounds, like systems or solutions of two or more compounds. There are also homogeneous compounds, like alloys, glasses, ceramic materials, solid solutions, minerals, and doped compounds, that belong (not exclusive) to this category. All these compounds have the number of components greater than one (NC>1).

- First, compounds can be stored in a kind of tabular form containing the components of alloys, glasses and ceramics together with their atomic, volume, weight, or undefined percentages. On searching for these compounds the search fields ALLOY, PERC and PERT are recommended. Example: Pd(b),Rh(2-3) (W%), this means a palladium based alloy containing 2-3 weight % rhodium. Al2O3(10-15),B2O3(10-15),O2Si (W%), this means a silicium dioxide (Hill order) glass containing 10-15 weight % aluminium oxide and 10-15 weight % boron oxide.
- Second, and this is the normal case, compounds are stored with their line formula. These compounds can consist of one, or more, component, and each component can consist of one or more than one fragment, and each fragment may or may not have a structure. Each component can be searched by its molecular formula, e.g. by MOFO or MF. The single fomulas can be combined. Example: MF=BaO and MF=CuO and MF=O3Y2 and NC=3, this means a threecomponent substance of the compounds BaO, CuO and Y2O3.
- Compounds with more than one component can be given in the field LSF (linearized structure formula) in the following style:
 - ABC#XYZ the single components ABC and XYZ are separated by a hash sign (#)
 - (A,B)Z the single component parts are separated by a comma, giving the components AZ and BZ



- A(x)B(1-x)C x:0.9-0.95; gives the components A0.95B0.05C and A0.9B0.1C.
- Formulas of type 2 and 3 often refer to solid solutions, minerals or diadochous compounds. Type 1 often refers to homogeneous or heterogeneous multicomponent compounds.

4.4 Compound Formulas with more than one Fragment

Compounds with more than one fragment are separated in the linearized structure formula field by an asterisk. Example: 2Na(1+)*SO4(2-)=Na2SO4 Normally, the formula is repeated in the conventional manner after the '=' symbol as shown in the example above. Rules whether a compound is fragmented or not are as follows:

Solid state structures and ion lattices:

- Substances, which contain in the crystal, the gas phase, the liquid phase, or in solution discrete polyatomic ions are divided into formal fragments if there is at least one structurable fragment, e.g. 2Na(1+)*SO4(2-)=Na2SO4.
- on lattices, which consist exclusively of one-atomic ions (including OH, SH, SeH-, and TeH-), or solid state structures without discrete ions or molecules, have no structure in the database; the formula is given without fragmentation, e.g. NaCl, KOH, CaTiO3.

Organometallics and Coordination Compounds:

- Organometallic compounds, and coordination compounds with known manner of ligand coordination, are stored with the complete formula and structure, e.g. Cr(CO)6, Fe(C5H5)2, (Co(NH3)3Cl3).
- Organometallic salts, and coordination compounds with known manner of ligand coordination, are divided into cations and anions. The complete structure of the ions are available, e.g. (Pd(CH3CN)4)(2+)*2BF4(1-)=(Pd(CH3CN)4)(BF4)2
- Organometallic compounds, and coordination compounds with unknown manner of ligand coordination, are divided into fragments by separating all ligands from the metal center. E.g., if there is an unknown coordination of the dianion of oxalic acid to Fe(2+) in [Fe(OOCCOO)2](2-) (mono- or bidentate), the formula is given as Fe(2+)*2OOCCOO(2-). E.g. if there is an unknown coordination of thiocyanate to Co(2+) in [Co(NH3)5(SCN)](2+) (coordination by N or S atom), the formula is given as Co(2+)*5NH3*SCN(1-). Additional the complete structure of the complex ion (Co(NH3)5(SCN))(2+) can be available (in this case twice with Co-SCN and Co-NCS coordination).

Note:

Clusters with unknown ligand coordination are not divided further; hydrogen atoms are not separated from the cluster units; the cluster cores get no structure; the structures of the ligands are given e.g. if the coordination of CO in the cluster H2Os3(CO)10 is unknown, the formula is given as H2Os3*10CO; the cluster core gets no structure.



4.5 General Information on Structures

You should be aware that only molecular structures are available, for instance all coordination compounds or molecules.

Single bonds, double bonds and triple bonds are available as bond types in the database, but note, the bond type is no criterion for registration. If one is not sure about the bond order, the bond type 'any bond' should be preferred in the search query. There are no tautomeric or aromatic bonds. In the case of keto/enol tautomerism, the ketonic form is always stored. Different tautomers are stored explicitly. All pi-interactions within organometallic pi-complexes are drawn as single bonds.

Structures of fragments are identical if the connectivity of the chosen atoms and the overall charge are the same. Only additional stereodescriptors can differentiate them further.

Main group organometallic compounds:

In the gray area between covalent and ionic bonds, usually both formulations occur in the original literature for main group organometallic compounds. From the point of view of input conventions the following rules are generally favored by the Gmelin file, but one may find both alternatives, and it is recommended that you search for both structures to ensure completeness:

- Lithiumorganic compounds are stored with a covalent Li-C single bond
- Sodium and potassiumorganic compounds are stored in ionic form (more than one fragment)
- Grignard compounds are stored in the covalent form
- Organometallic compounds with metals of the main groups III-VI are stored in the covalent form

Transition metal organometallic compounds:

Transition metal organometallic compounds are input with the connectivity of atoms given in the original literature. If the connectivity is unknown, the compounds are separated formally into the central metal atoms and the surrounding ligands.

4.6 Stereochemistry

Stereochemistry is a further identification criterion of the structures in the Gmelin file. Two principal different stereochemical aspects can be searched for: Stereochemical information on bonds and on centers.

Specified double bond information (cis or trans) can be searched for if the bond attribute "Double Steric" is chosen in the structure editor, and the structure is drawn as desired.



All stereocenters are drawn three-dimensionally. Fragments with stereochemical information are assigned either an absolute or a racemic configuration.

In the case of trigonal pyramidal, tetrahedral, or octahedral centers, a stereodescriptor is set if one, or more than one, possible configuration of a stereocenter can be assigned. This also applies to all pi-descriptors and all descriptors linking two polyhedra by a chain of bonds.

In all other cases the mentioning of a polyhedral center within a compound in the original literature is sufficient for the designation of a polyhedral descriptor. In the case of sterical isomers the polyhedral descriptor fix also the ligand configuration. If the configuration is unknown, no descriptor is set.

GMELIN polyhedron descriptors:

Descriptor	Polyhedron
P-3	trigonal pyramid
T-4	tetrahedron
SP-4	square planar
SP-5	square pyramid
TB-5	trigonal bipyramid
OC-6	octahedron
TP-6	trigonal prism
PB-7	pentagonal bipyramid
CU-8	cube
SA-8	square antiprism
DD-8	dodecahedron
HB-8	hexagonal bipyramid
TPS-9	tricapped trigonal prism
HB-9	heptagonal bipyramid
OC-6-PI	octahedron (with p ligand)

Note:

Additional to isomerism on double bonds stereoinformation on single bonds (eclipsed and staggered-configuration) is stored in the database. Further chiral isomers with at least known sign of optical rotation but unknown polyhedral geometry are distinguished. However, these descriptors are not searchable.



4.7 Identical Molecular Formula - New Compound?

Searching for a certain molecular formula may often give more than one hit. The following criteria are responsible for a further registration of a compound with molecular formula identical to another:

- different charge
- isotope marked atoms in the formula
- the substance types
 - glass or ceramic material
 - Isomorphous, diadochous compounds, solid solutions
 - Isotope or isotope containing compounds
 - Minerals
 - Polymer
- modifications of inorganic compounds
- different connectivity of atoms in the structure
- additional stereodescriptors



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