A proposal of an XML format for TDB files

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It seems there is a general agreement that an XML format is beneficial for Calphad databases, the discussion is how detailed it should be. The proposal here is intended to make it possible to edit an XTDB file as easily as the current TDB format while providing better facilities to handle new features implemented in the different softwares using TDB files. This may require modifications in the XML format already adopted in some software.

1 XTDB version 0.0.1

The following proposal is for a common definition of an XTDB file format based on XML. It can hopefully become an SGTE standard acceptable by all software which develop or use Calphad databases.

As XML is extendable each software may add their own flavour in a controlled way and it will be easier to convert an XTDB file from one software to another.

The "note" for the attributes for an XML element in the tables below is "+" for a compulsory attribute, "-" for an optional one or a letter A-Z for special cases explained in section 1.9.

All **Id** attributes are case insenitive as in the TDB file. Except for phase names the "Id" cannot be abbreviated when used in attributes.

An XML parser is convenient for reading XML files but to simplify the life for those reading the XTDB file manually or using a sequential read operation in a software, it is recommended that the attributes to an XML element is provided in the order listed for each element below.

The order of XML elements is free but to simplify for a human reading and editing the XTDB file a recommended order is **Defaults**, **DatabaseInfo**, **SoftSpecific** followed by **Element**, **Species**, **Model** and **Phases** and **TPfun** and **Parameter** for the pure elements. After that **TPfun** and **Parameter** for binary, ternary and higher order systems, normally arranged for each subsystem.

An XML element referenced in the text below is written in **bold**.

1.1 The system XTDB elements

Element	Attributes	Note	Explanation			
XTDB	>>>		Containing XML elements for an XTDB database.			
	Version	+	Version of XTDB for this file.			
	Software	+	Name of software generating the database.			
	Date	+	Year/month/day the database was written or edited			
	Signature	_	Name/email of person or organisation generating the database.			
Defaults	>>>		Provides default values of attributes in other XML elements.			
	LowT	_	Default value of low T limit. Default value of high T limit.			
	HighT	_				
	Refid	_	Default bibliographic reference.			
	Elements	_	VA and/or /- (the electron).			
	Model	_	Any model applicable for the system, for example EEC			
DatabaseInfo	o >>>		Free text.			
	Date	_	Last update.			
SoftSpecific	>>>>		Free text explaining software specific XML elements and attributes used.			

1.2 The chemical element and species element

Element	Attributes	Note	Explanation			
Element	>>>		Specifies a chemical element in the database. The vacancy is denoted			
			"VA" and the electron "/-".			
	Id	+	Chemical element symbol, for example FE, H			
	Refstate	+	Name of a phase, for example GAS			
	Mass	+	Mass in g/mol			
	H298	_	Enthalpy difference between 0 and 298.15 K in reference state.			
	S298	_	Entropy difference between 0 and 298.15 K in reference state.			
Species	>>>		Specifies a molecular like aggregate used as constituent of phases. The			
			elements, except the electron, are also species.			
	Id	A	Species name max 24 letters. For note see 1.9.			
	Stoichiometry	r В	One or more elements each followed by an unsigned real representing the			
			stoichiometric ratio. For note see 1.9. See also appendix A.1.			
	MQMQA	\mathbf{C}	For a constituent in the MQMQA model. For note see 1.9.			
	UNIQUAC	D	For a constituent in the UNIQUAC model. For note see 1.9.			

1.3 The function and temperature range elements

There is at present no way to handle several pressure ranges. The value of a function, as well as its first and second derivative with respect to T, must be continuous across a change of T range.

Element	Attributes	Note	Explanation			
TPfun	>>>		Defines a T, P expression to be used in parameters or other functions.			
	Id	\mathbf{E}	Function name, max 16 characters. The name can be used in the "Expr"			
			attribute of other functions or parameters. For note see 1.9.			
	LowT	_	If the default low T limit applies.			
	Expr	+	Simple mathematical expression terminated by ;. Use the Trange ele-			
			ment if several ranges. See section 1.10			
	HighT	_	If the default high T limit applies.			
Trange	>>>		Only inside a TPfun or Parameter element for an expression with			
			several T ranges.			
	Expr	+	Simple mathematical expression terminated by ;. See section 1.10.			
	HighT	_	If the default high T limit applies.			

1.4 The phase element

Normally the software requires information of sublattices, constrituents and models for the phase in order to create the appropriate datastructures. The **Parameter** and some models, see section 1.8.3 can appear anywhere in the XTDB file.

The **Splitphase** model (also known as the disordered fraction model) combines the Gibbs energies of an ordered and a disordered phase where the disordered phase has fewer sublattices than the ordered and the constituent fractions on its first sublattice are the average of the fractions summed over several sublattices in the ordered phase as specified by the attribute **Sum**. The **SplitPhase** element must appear within the ordered **phase** element before any other nested XML elements such as **AmendPhase**, **Parameter**, **KohlerModel** or **ToopModel** elements. The **Parameter**, **ToopModel** and **KohlerModel** elements may appear anywhere in the XTB file.

Some software may not have a separate disordered phase but arrange all **Parameter** elements for the ordered and disordered phase within the same phase.

Element	Attributes	Note	Explanation			
Phase	>>>		All data belongs to a phase.			
	Id	\mathbf{F}	Phase name, see 1.9 note F.			
	Configuration	ı G	Model for the configurational entropy. For note see 1.9.			
	State	_	G for gas phase, L for liquid phase. Needed if EEC is used.			
Sublattices	>>>		Only inside a Phase element.			
	Number-of	+	Number of sublattices, an integer value 1-10			
	Ratios	+	Sites on the sublattices, as many reals as sublattices separated by a space.			
Constituents	>>>		Only inside a Phase element.			
	Sublattice	+	Indicates the sublattice for the list of species.			
	List	+	Species Id, separated by a space, in the sublattice.			
SplitPhase	>>>		Only inside the Phase element of the ordered phase.			
	Refid	+	Where the model is described.			
	Disordered	K	Name of disordered phase, for note see 1.9.			
	Sum	+	Number of sublattices in the ordered phase to be summed for the disor-			
			dered phase constitution. All sublattices summed must have the some			
	~ .		constituents.			
	Subtract	L	Must have value Y if the Gibbs energy of ordered phase as disordered			
			should be subtracted, for note see 1.9.			
AmendPhase			Only inside a Phase element.			
	Model	+	List of models "Id", separated by a space, for this phase. The "Id" is			
			specified in a model element below.			

1.5 The simple parameter element

All thermodynamic data are defined by the parameter elements. They can be arranged inside a phase element or separately for each binary, ternary etc. subsystem. See also section 1.6 and appendix A.4.

Element	Attributes	Note	Explanation		
Parameter	>>>		Specifies the T, P expression of a model parameter for a set of con-		
			stituents.		
	Id	+	As in TDB file, for example G(LIQUID,A,B:VA;2). See also the Param-		
			eter2 element. May include the Trange element for parameters with		
			multiple ranges.		
	LowT	_	If the default low T limit applies.		
	Expr	+	Simple mathematical expression terminated by ;. If several ranges use a		
			Trange element. See section 1.10.		
	HighT	_	If the default high T limit applies.		
	Bibref	+	Bibliographic reference.		

1.6 The elaborate parameter element

An alternative more elaborated XML element can be used for parameters which may be preferred by software. It is striaghtforward to convert from one to another.

Element	Attributes	Note	Explanation			
Parameter2	>>>		A more detailed parameter element preferred by software.			
	Id	_	As in TDB file, for example G(LIQUID,A,B:VA;2). See also the Param-			
			eter element.			
	MPID	Н	Model parameter identifier, for example G or TC. For note see 1.9.			
	Phase	_	Can be omitted inside a phase element, otherwise the phase name.			
	LowT	_	If the default low T limit applies.			
	Expr	+	Simple mathematical expression terminated by ;. If several ranges use a			
			Trange element. See section 1.10.			
	HighT	_	If the default high T limit applies.			
	Bibref	+	Bibliographic reference.			
ConstArray	>>>		Only inside a parameter2 element.			
	Degree	I	Can be omitted if zero, for note see 1.9.			
Constituent	>>>		Only inside a ConstArray element.			
	Sublattice	+	Sublattice of constituent.			
	Species	+	Name of a constituent.			

The **Parameter2** element is preferred by software but for manual editing **Parameter** is simpler, see the appendix A.4. Both elements can be used in the XTDB file. Software can convert a **Parameter** element to a **Parameter2** element. If an "Id" is present in a **parameter2** element the software should check that the "Id" is identical to the long form and report an error if not.

The use of wildcards, "*", for species in a parameter must be defined, see section 1.11.

1.7 Bibliography for parameters and models.

For each parameter there must be a bibliographic reference. This can be the same for all parameters from the same source. The models also have a reference which should indicate a paper where the model is explained, see Appendix A.5.

Element	Attributes	Note	Explanation	
Bibliography	y >>>		Contains bibliographic references	
Bibitem	>>>		Only inside a Bibliography element.	
	Id	+	Used as bibref attribute for a parameter or model, normally a paper or	
			a comment by the database manager.	
	Text	+	Reference to a paper or comment	
	Date	_	 Date when the bibref element was created 	
	Sign	_	Signature of the one adding the reference	

1.8 XTDB model elements

These XML elements for generally accepted models can be on a separate file but the model "Id" is the important and used in the **AmendPhase** element for each phase which has the model. Most model has one or more model parameter identifiers (MPID) used in the **Parameter** element. See also appendix B. Software specific models should be explained in this XML element.

Element	Attributes	Note	Explanation
Models	>>>		Contains model descriptions and identifiers used in "AmendPhase" ele-
			ments for phases. The models usually specify one or more model param-
			eter identifiers needed by the model.

The following XML elements should only be used inside a **Model** element and are used to specify the model parameter identifiers associated with the model.

1.8.1 XTDB generally accepted model elements

Element	Attributes	Note	Explanation				
Magnetic	>>>		There are several magnetic models.				
	Id	+	This Id is used in AmendPhase element.				
	Refid	+	Where the model is described.				
	MPID1	+	Specifies a magnetic model parameter identifier (MPID) for parameters.				
	MPID2	+	Specifies a magnetic model parameter identifier (MPID) for parameters.				
	MPID3	+	Specifies a magnetic model parameter identifier (MPID) for parameters.				
Volume	>>>		Specifies the model for volume of a phase.				
	Id	+	This Id is used in the AmendPhase element.				
	Refid	+	Where the model is described.				
	MPID1	+	Specifies volume model parameter identifier (MPID) for parameters.				
	MPID2	+	Specifies volume model parameter identifier (MPID) for parameters.				
	MPID3	+	Specifies volume model parameter identifier (MPID) for parameters.				
FCC4Perm	>>>		For the FCC 4 sublattice tetrahedron model identical permutations of a				
			parameter are listed only once.				
	Id	+	This Id is used in AmendPhase element.				
	Refid	+	Where the model is described.				
BCC4Perm	>>>		For the BCC 4 sublattice tetrahedron model identical permutations of a				
			parameter are listed only once.				
	Id	+	This Id is used in AmendPhase element.				
	Refid	+	Where the model is described.				

1.8.2 New unary models

There is also a new magentic model.

Element	Attributes	Note	Explanation			
Einstein	>>>		The low T vibrational model.			
	Id	+	This Id is used in AmendPhase element.			
	Refid	+	Where the model is described.			
	MPID1	+	Specifies the Einstein model parameter identifier (MPID) for parameters.			
Liquid2state	>>>		The liquid 2-state model.			
	Id	+	This Id is used in AmendPhase element.			
	Refid	+	Where the model is described.			
	MPID1	+	Specifies liquid model parameter identifier (MPID) for parameters.			
	MPID2	+	Specifies Einstein model parameter identifier (MPID) for parameters.			
EEC	>>>		Specifies that the Equi-entropy model applies to current database			
	Refid	+	Where the model is described.			

1.8.3 XTDB elements for constituent specific models

These models are related to specific constituents of a phase and they must be specified explicitly and cannot be included in the **AmendPhase** element for the phase because they include additional information. The Toop

and Kohler ternary extrapolation models are most likely specified together with the parameters for a specific ternary. The Muggianu ternary model is the default extrapolation model.

Element	Attributes	Note	Explanation
ToopModel	>>>		Subset of 3 constituents for which the Toop ternary extrapolation model
			should be used. By default the Muggianu extrapolation is used.
	Refid	+	Where the model is described.
	Phase	+	Can be omitted if used inside a Phase element. Otherwise the phase for
			which the model should be used.
	Constituents	s +	Specifies 3 constituents, the Toop constituent first.
KohlerMode	l >>>		Subset of 3 constituents for which the Kohler ternary extrapolation model
			should be used. By default the Muggianu extrapolation is used.
	Refid	+	Where the model is described.
	Phase	+	Can be omitted if used inside a Phase element. Otherwise the phase for
			which the model should be used.
	Constituents	s +	Specifies 3 constituents in any order.

1.9 Notes

Note that the XML elements and attributes are case sensitive whereas the names of elements, species, functions and other text are case insensitive. For example a chemical element written FE, Fe, fe and fE is the same element. Thus CO is Cobalt and C1O is carbon monoxide.

- **A:** A species name or **Id** must start with letter A-Z and can contain letters, digits and the special characters "_", "/", "-" and "+". It must not be abbreviated when used as constituent in a **Phase** or **Parameter** element.
 - The letters "-" and "-" may be treated as identical?
- **B:** The species stoichiometry is a sequence of one or more element names (case insensitive) followed by a real number specifying the stoichiometric ratio. Following the TDB standard an element with a two letter name does not need a stoichiometric ratio equal to unity, for example MGO for MG1O1. A final stoichiometry unity can also be ignored. No parenthesis are allowed. For MQMQA clusters this can be omitted. ... maybe more needed ...
- C: The MQMQA attribute should contain two or more element names separated a comma or a sublattice separator ":" and followed by equal number of unsigned reals representing the bond fractions, For example: CS,LA:F 9.0 6.0 4.0
 - See also section A.1. According to the model the stoichometry is calculated as 6.0 divided by the bond fractions. For endmember clusters (with one species in each sublattice) a real representing the SNN/FNN ratio, usually 2.4, must be supplied.
- **D:** For UNIQUAC species. Contains two reals representing volume and area of the species in m².
- **E:** A TPfun name must start with a letter A-Z and may contain letters, digits and the special character "_". It must not be longer than 16 characters. It cannot be abbreviated.
- **F:** A phase name must start with a letter A-Z and have no more than 24 characters. It can contain letters, numbers and the special character "_". In some software "(" and ")" may be used in phase names but they can be replaced by "_" in other software.
 - A phase name may be abbreviated in parameters and some other cases and thus a phase name must be unique and not an abbreviation of another phase.

- **G:** Configuration model, for example CEF, I2SL, MQMQA etc. Maybe also "RKM" for a Redlich-Kister-Muggianu model with a single lattice and one site?
- **H:** An initial set of model parameter identifies (MPID) must be defined. At present G is the Gibbs energy, TC Curie T, BMAGN the Bohr magneton number etc. The MPID must not be abbreviated. See section 1.12
- **I:** The degree can be omitted if zero.
- **K:** The disordered phase within a **SplitPhase** element can be omitted if the parameters for the disordered phase are provided within the ordered phase, using a suffix "D" after the MPID name and with the appropriate reduced number of sublattices.

For example GD(SIGMA,CR) can be used as **Id** in a **Parameter** element for the lattice stability of Cr in SIGMA if the SIGMA phase is modelled with 3 or more sublattices and has the **SplitPhase** element. This avoids the need for a meaningless disordered DIS_SIGMA phase.

L: A BCC or FCC phase with a disordered fraction set can have the "ordered" part "as disordered" subtracted in order to allow the modeling of the disordered part separately. If so the value of the **Subtract** attribute should be "Y".

For phases which never disorder, for example the sigma phase, one can also have a "disordered" part with a single sublattice containing the pure elements as endmembers (and possibly some long range interaction parameters). In Thermo-Calc terminology this is the "NEVER" model. In order not to subtract the ordered part as disordered in XTDB the **Subtract** attribute can be omitted or given any other value than "Y".

1.10 The mathematical expression used in Expr

The mathematical expressions for T and P used in **TPfun**, **Trange** and **Parameter** are the same as in TDB files. It is very limited because some software must calculate first and and second derivatives with respect to T, P (and constitution). A more extended mathematical expression could be allowed for expressions which are not used for database parameters. In OC it is allowed to enter more complex expressions for post processing in batch/macro files.

In the 1990 definition of the TDB file the type of expression allowed consists of "simple terms" such as:

```
[signed real number] * [TPfun name] ** [power] *T** [power] *P**[power]
```

where [power] is an integer, a negative power must be surrounded by parenthesis. No spaces allowed in a simple term. A "complex term" is a simple term multiplied with a math function of a simple term such as:

```
[simple term] *LN( [simple term] )
```

An Expr attribute in **TPfun**, **Trange** or **Parameter** consists one or more terms. A positive sign of the first term can be omitted.

The following general math functions are allowed in OC:

 $\exp(), \ln(), \log(), \operatorname{erf}()$. Note that $\log()$ and $\ln()$ is the same and $\operatorname{erf}()$ is the error function. The number of math functions can be extended.

```
The following math function is needed for the unary project: GEIN(\theta) to calculate 1.5R\theta + 3RT \ln(1 - \exp(-\theta/T));
```

It is not allowed to group several terms together using parenthesis.

1.11 The use of wildcards for constituents in parameters

In some parameters a wildcard, "*", is used to indicate the parameter is independent of the constituent in this sublattice. For example:

• A parameter G(sigma,GA:GA:*;0)=100000; means that the Gibbs energy of formation of "phase" with GA in first and second sublattice is independent of the constituent in the third. If one later finds that for example the Gibbs energy for sigma with TA in the third sublattice should be just +70000 then, in OpenCalphad, one must set G(sigma,GA:GA:TA)=-30000; because OC stores the two parameters separately and will add them when calculating.

This may be handled differently in other software and requires attention.

• The EBEF model, using the same notation for parameters as in CEF, replaces a large number of Gibbs endmembers energies by fewer **Parameter** elements, representing bond energies, using wildcards in one or more sublattices. The bond energies specify constituents in only 2 or 3 sublattices in phases with 3 or more sublattices. Using the **SplitPhase** model and provided there two endmember parameters GD(SIGMA,CR) and GD(SIGMA,FE) (or a separate disordered phase with these parameters) a parameter G(SIGMA,CR:FE:*:*:*) represent the bond energy between CR in first and FE in second sublattice, independent on the constituents in the other sublattices (note that G(SIGMA,FE:CR:*:*:*) have another value).

Using such parameters (fitted for example to DFT calculated endmember energies) the number of parameters can be reduced by more than an order of magnitudes and, in addition, extrapolations can be improved!

In order to avoid a phase representing the "disordered part" in EBEF one can introduce a set of endmember parameters such as G(SIGMA,CR:*:*), G(SIGMA,*:CR:*), G(SIGMA,*:*:CR) and G(SIGMA,FE:*:*), G(SIGMA,*:FE:*) and G(SIGMA,*:*:FE) (the so called "wild" model) which is ingenious but likely confusing to students.

1.11.1 A missing wildcard in the I2SL model parameters

An I2SL model may exist with only neutrals in the anion sublattice. For example the elements C and S can be neutrals in the I2SL model and their parameters are in Thermo-Calc are written as G(I2SL,C), omitting the cation sublattice. In OpenCalphad they are written G(I2SL,*:C) which may be a bit more consistent as it indicate that the constituent on the cation sublattice is irrelevant. It is suggested that the XTDB format adopts the OC way to specify a constituent in each sublattice in order to be consistent. The same for interactions between neutrals.

Note that an interaction between neutrals with a single specific cation is forbidden in the I2SL model.

1.12 Model parameter identifiers, MPID

An MPID must start with a letter A-Z and contain letters and digits and not exceed 8? charactes. They cannot be abbreviated. There are already some defined but there should be an extensive list of future MPIDs to avoid that different software use the same for (slightly) different things. See appendix C.

The letter "&", frequently used for mobilities, is forbidden in XML but it can be replaced by some other character, for example "@". Using "&" seems unnecessary and quite clumsy.

1.13 CVM and the cluster site model

These should eventually be included also.

2 Points of discussions

- 1. Should mathematical expressions be terminated by a ";"? It is not necessary because there is a final double quote of the expression but it may be nice.
- 2. The GEIN function is used in the **Parameter** for a pure element modeled with several Einstein θ . Only one of these is selected to vary with composition using the LNTH model parameter identifier (MPID).
- 3. Should the sign of Redlich-Kister terms depend on the alphabetrical order of the constituents or as the order the constituents are written in the parameter Id? Currently the TDB file use alphabetical.
- 4. Automatic reation of multiple composition sets? For example in Fe-Ti-C there will normally be an MC carbide with the FCC model parameters. Could such a composition set have a different name? It may be useful to provide handling of carbides or ordering in application software.
- 5. Using a "#digit" to identify a composition set? #1 is the default (first) composition set.

3 Summary

There are certainly many more things to take care of but I think it is more important to agree on a minimum common XML format which can make the thermodynamic databases more useful both for experimentalists, assessments, database maintenance and thermodynamic software in particular for the development of new models and applications. We have to take one step at a time.

Appendix A Examples

A few examples of the XTDB format. It may not be completely consistent with the definitions above as I have taken them from my previous conversion software.

A.1 Elements and species.

The way to define MQMQA constituents and stoichiometry is tentative.

```
<Element Id="/-" Refstate="ELECTRON_GAS" Mass=" 0.000000E+00" />

<Element Id="VA" Refstate="VACUUM" Mass=" 0.000000E+00" />

<Element Id="AL" Refstate="FCC_A1" Mass=" 2.698200E+01" H298=" 4.577300E+03" S298=" 2.832200E+01" />

<Element Id="FE" Refstate="BCC_A2" Mass=" 5.584700E+01" H298=" 4.489000E+03" S298=" 2.728000E+01" />

...

<Species Id="VA" Stoichiometry="VA" />

<Species Id="AL" Stoichiometry="AL" />

<Species Id="FE" Stoichiometry="FE" />

<Species Id="ALFE" Stoichiometry="AL0.5FE0.5" />
...

<Species Id="LA/F" MQMQA="LA:F 6.0 2.0 2.4" />

<Species Id="LACS/F" MQMQA="LA,CS:F 9.0 6.0 4.0" />
```

The stoichiometry of the MQMQA species is calculated from the bond ratios according to the model. The **MQMQA** attribute for an "endmember" also include a factor needed in the configurational entropy expression.

Note that species cannot be abbreviated as constituents in phases or parameters and thus one can have species which are abbreviations of another species. For example "O" and "O2".

For phases one can abbreviate its name and thus one cannot have a phase with a name which is an abbreviation of another phase name. Phase names can be abbreviated for each part separated by an underscore, "_". A phase name "AL_X" is thus an abbreviation of "AL2_X".

A.2 Defaults, TPfun and Trange

<TPfun Id="GAL3FE" Expr="+3*UFALFE+9000;" />
<TPfun Id="GAL2FE2" Expr="+4*UFALFE;" />

Using default T limits the function are not much more complex than in the TDB file.

A.3 Phase

Entering phases in the XTDB file is a bit more complex but we have get rid of the TYPE_DEFINITION. It is not so nice to use CEF for the liquid, that is why I think RKM might be a nice option for a liquid model without sublattices and maybe **Sublattices** can be omitted?

```
<Phase Id="LIQUID" Configuration="RKM" state="L" >
   <Sublattices Number_of="1" Ratios="1" >
     <Constituents sublattice="1" list="AL C" />
   </Sublattices>
   <AmendPhase model="LIQ2STATE" />
</Phase>
<Phase id="A2_BCC" Configuration="CEF" state="S" >
   <Sublattices Number_of="2" Ratios="1 3" >
     <Constituents sublattice="1" list="AL FE" />
     <Constituents sublattice="2" list="VA" />
   </Sublattices>
   <AmendPhase model="IHJBCC" />
</Phase>
<Phase id="AL8FE5_D82" Configuration="CEF" state="S" >
   <Sublattices Number_of="2" Ratios="8 5" >
     <Constituents sublattice="1" list="AL FE" />
     <Constituents sublattice="2" list="AL FE" />
   </Sublattices>
</Phase>
<Phase id="BCC_4SL" Configuration="CEF" state="S" >
   <Sublattices Number_of="5" Ratios="0.25 0.25 0.25 0.25 3" >
     <Constituents Sublattice="1" List="AL FE" />
     <Constituents Sublattice="2" List="AL FE" />
     <Constituents Sublattice="3" List="AL FE" />
     <Constituents Sublattice="4" List="AL FE" />
     <Constituents Sublattice="5" List="VA" />
   <SplitPhase Disordered="A2_BCC" Sum="4" Subtract="Y" Refid="09Sun" />
   <AmendPhase model="IHJBCC BCC4Perm" />
 </Phase>
```

A.4 Parameters

```
<Parameter Id="G(A2_BCC,FE:VA;0)" Expr="+GHSERFE;" Bibref="91Din" />
    <Parameter Id="TC(A2_BCC,FE:VA;0)" Expr="1043;" Bibref="91Din" />
    <Parameter Id="BMAGN(A2_BCC,FE:VA;0)" Expr="2.22;" Bibref="91Din" />
    <Parameter Id="G(AL8FE5_D82,AL:AL;0)" Expr="+13*GALBCC;" Bibref="08Sun" />
    <Parameter Id="G(BCC_4SL,AL:AL:FE:FE:VA;0)" Expr="+GB2ALFE;" Bibref="08Sun" />
    <Parameter id="G(BCC_4SL,AL;AL;FE:*:*:VA;1)" Expr="-634+0.68*T;" Bibref="08Sun" />
```

In the last parameter above the "wildcard" or asterisk, "*", is used for species in three of the sublattices and it means that the parameter is independent of the constituents in these sublattices. See the discussion in section 1.11 how this is treated.

Using the **Parameter2** element the last two parameters above are:

```
<Parameter2 Id="G(BCC_4SL,AL:AL:FE:FE:VA;0)" Expr="+GB2ALFE;" Bibref="08Sun" MPID="G" Phase="BCC_4SL" >
   <ConstArray Degree="0">
      <Constituent Sublattice="1" Species="AL" />
      <Constituent Sublattice="2" Species="AL" />
       <Constituent Sublattice="3" Species="FE" />
       <Constituent Sublattice="4" Species="FE" />
       <Constituent Sublattice="5" Species="VA" />
   </ConstArray>
/Parameter2>
<Parameter2 id="G(BCC_4SL,AL,FE:*:*:VA;1)" Expr="-634+0.68*T;" Bibref="08Sun" MPID="G" Phase="BCC_4SL" >
   <ConstArray Degree="1">
      <Constituent Sublattice="1" Species="AL" />
      <Constituent Sublattice="1" Species="FE" />
      <Constituent Sublattice="2" Species="*" />
      <Constituent Sublattice="3" Species="*" />
      <Constituent Sublattice="4" Species="*" />
       <Constituent Sublattice="5" Species="VA" />
   </ConstArray>
/Parameter2>
```

A.5 Models

There should be a separate global library file with the "Id" and description of the models.

```
<Models>
  <Magnetic Id="IHJBCC" MPID1="TC" MPID2="BMAGN" Bibref="82Her" />
  <Magnetic Id="IHJREST" MPID1="TC" MPID2="BMAGN" Bibref="82Her" />
  <Einstein Id="GLOWTEIN" MPID1="LNTH" bibref="01Qing" />
  <Liquid2state Id="LIQ2STATE" MPID1="G2" MPID2="LNTH" bibref="14Becker" >
        Unified model for the liquid and the amorphous state treated as an Einstein solid
  </Liquid2state>
  <Volume Id="VOLOWP" MPID1="VO" MPID2="VA" MPID3="VB" bibref="05Lu" >
        The volume of a phase is described as function of T, P and its constitution.
  </Volume>
...
  </Models>
...
  <ToopModel Phase="Liquid" Constituents="C Fe Cr" />
```

The **ToopModel** or **KohlerModel** elements can appear anywhere in the XTDB file, either within the **Phase** element or, in order to simplify editing a large XTDB file, more likely together with all the **Parameter** elements from an assessment of the ternary C-Fe-Cr system. Either way may be a bit complicated to handle by the calculating software reading the XTDB file.

The **SplitPhase** in example A.3 must appear within a **Phase** element for the ordered phase.

Appendix B Proposal of Id for agreed models.

The agreed models need only to added by their idenfiers to the phases.

For models that are not agreed a full description of the model should be provided in the XTDB file. Note the model parameter identifiers may be different in each software.

```
**Stagnetic id="HIRBCC" MPIDI="TC" MPID2="BMADN" anti-ferromagnetic_factors" -1.00"
f_below_TC="-1-0.905299383+Tab=(-1)-0.153008346+Tab=3-0.005207859854+Tab=-0.0153008346+Tab=+15;"
f_abov_TC="-1-0.905299383+Tab=(-1)-0.153008346+Tab=-0.0153008346+Tab=+15;"
f_abov_TC="-1-0.905299383+Tab=(-1)-0.1530934740**(-15)-0.00427820805+Tab=+0.00524740**(-25); blbref="82Her" > in G=f(TAD)+LN(BETA+1) where TAD=T/TC

**Magnetic id="HIRBST" MPID1="TC" MPID2="MMADN" anti-ferromagnetic_factors" -3.00"
f_below_TC="-1-0.806038755+Tab=(-1)-0.1749124*Tab=-3.00775516524*Tab=-9.001749124*Tab=+15;"
f_abov_TC="-1-0.806038755+Tab=(-1)-0.1749124*Tab=-3.00775516524*Tab=-9.001749124*Tab=+15;"
f_abov_TC="-1-0.806038755+Tab=(-1)-0.1749222256*Tab=-3.00774409825*Tab=(-1)-0.00**

**Magnetic id="HIRDX" MPID1="CT" MPID2="MT" MPID3="BMADN" anti-ferromagnetic_factors" 0.00"
f_below_TC=-1-0.8028049633*Tab=(-1)-0.17424222256*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-3.00774409825*Tab=-0.001742422225*Tab=-0.001742422225*Tab=-0.001742422225*Tab=-0.001742422225*Tab=-0.001742422225*Tab=-0.001742422225*Tab=-0.00174242225*Tab=-0.001742422225*Tab=-0.001742422225*Tab=-0.001742422225*Tab=-0.001742422225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=-0.00174242225*Tab=
```

Appendix C Proposal of initial set of Model Parameter Identifiers, MPID

The 32 model parameter identifiers, MPID, defined in OC as shown in Table 1. In OC a parameter for a disordred part of a **SplitPhase** use the same phase name but the MPID has a suffix "D".

Table 1: Current set of model parameter identifiers in OC. For each parameter it is indicated if it can depend on T, P or have an extra constituent specification. Most of them have no associated code.

Indx	Ident	Т	Р	Specification	Status	Note
	G	Т		-	0	Gibbs Energy
2	TC	-	P			Combined Curie/Neel T
3	BMAG	-	_		1	Average Bohr magneton number
4	CTA	-	P			Curie temperature
5	NTA	-	P		2	Neel temperature
6	IBM	-	P	<pre>&<constituent#sublattice>;</constituent#sublattice></pre>	12	Individual Bohr magneton number
7	LNTH	-	P		2	Einstein temperature
8	VO	-	-		1	Volume at TO, PO
9	VA	T	-		4	Thermal expansion
10	VB	T	P		0	Bulk modulus
11	VC	T	P		0	Alternative volume parameter
12	VS				0	Diffusion volume parameter
13				<pre>&<constituent#sublattice>;</constituent#sublattice></pre>	10	Mobility activation energy
14	MF	T	P	<pre>&<constituent#sublattice>;</constituent#sublattice></pre>	10	RT*ln(mobility freq.fact.)
15	MG	T	P	& <constituent#sublattice>;</constituent#sublattice>	10	Magnetic mobility factor
16	G2	T	P		0	Liquid two state parameter
17	THT2	-	P		2	Smooth step function T
18	DCP2				2	Smooth step function value
19	LPX	T	P		0	Lattice param X axis
20	LPY	T	P		0	Lattice param Y axis
21	LPZ	T	P		0	Lattice param Z axis
22	LPTH	T	P		0	Lattice angle TH
	EC11					Elastic const C11
	EC12				0	Elastic const C12
	EC44				-	Elastic const C44
		T	P	<pre>&<constituent#sublattice>;</constituent#sublattice></pre>		UNIQUAC residual parameter (OC)
	RHO	T	-		0	Electric resistivity
	VISC					Viscosity
	LAMB					Thermal conductivity
	HMVA					Enthalpy of vacancy formation (MatCalc)
	TSCH					Schottky anomaly T (OC)
	CSCH					Schottky anomaly Cp/R. (OC)
33	NONE	Т	P		0	Unused