

## Part 5. Multisite mixing models built-in functions in GEM-Selektor v.3.3 code

Treatment of condensed symmetric non-electrolyte solutions in GEM-Selektor code is based on the default expression for the chemical potential of a  $j$ -th end member:

$$\nu_j = \frac{g_{j,T,P}}{RT} + \ln x_j + \ln f_j \quad (5.1-1),$$

where  $g_{j,T,P}$  is the molar apparent Gibbs energy function at  $T, P$  of interest;  $x_j = \frac{n_j^{(x)}}{\sum n_k^{(x)}}$  is the end member mole fraction ( $n_j$  is the primal mole quantity of  $j$ -th species and  $n_k = \sum x_j$ ,  $j \in L_k$  is the total mole quantity of the  $k$ -th phase), and  $f_j$  is a ‘real’ activity coefficient.

If all  $f_j = 1$  at any composition then eqn (5.1-1) reduces to the Raoult’s law of simple ideal mixing, where the activity of  $j$ -th end-member equals its mole fraction. This definition of ideal mixing, corresponding to the configurational entropy of random mixing on a single type of structural sites, is rather narrow. In a broader sense, ideal solutions can be defined as having zero excess entropy, zero enthalpy and zero volume of mixing (Oates, 1966; Price, 1985; Anderson and Crerar, 1993). Microscopic moieties (usually ions or atoms) can substitute each other independently on one or several types of structural sites, often called ‘sublattices’ (Hillert, 1998; Lucas et al., 2007). Sublattices formally represent long-range ordering (LRO).

Eq (5.1-1) corresponds to the simplest case where two or more moieties substitute each other on one site type, and there is one moiety per formula of end member. This is the case e.g. for carbonate (A,B,...)CO<sub>3</sub> or sulfate (A,B,...)SO<sub>4</sub> solid solutions, but it is clearly insufficient for most of rock-forming silicate or oxide minerals where mixing occurs on several sublattices simultaneously, and can be complicated by short-range ordering (SRO) effects. In such ideal ‘multisite mixing’, the activity  $a_j$  of  $j$ -th end member is not simply its mole fraction  $x_j$ , but a more complex function of ‘site fractions’  $y_{s,m}$  of  $m$ -th substituting moiety on  $s$ -th site type in  $k$ -th solution phase. This corresponds to a more complex form of the configurational entropy, and causes the ‘reciprocal’ relations between end members and their standard-state properties.

Here we need to introduce a set  $\Xi_k$  of indexes of sublattices (structural site types) in the  $k$ -th phase, where  $n(\Xi_k)$  is the number of site types indexed with  $s \in \Xi_k$ . Further, we introduce a set  $M_s$  of indexes of  $n(M_k)$  moieties that can appear on  $s$ -th site in  $k$ -th phase structure; moieties will be indexed with  $m \in M_k$ . Let  $\eta_{j,s,m}$  be a ‘moiety-site multiplicity’, i.e. the number of moles of  $m$ -th moiety on  $s$ -th site in one mole of  $j$ -th end member at standard state ( $j \in L_k$ ), and  $\eta_s$  be the number of  $s$ -th sites in one formula unit of any end member of the  $k$ -th phase ( $s \in \Xi_k$ ) (‘site multiplicity’). Note that the same species (e.g. ion M<sup>+</sup>), if it appears on two different sublattices, must be considered as two different moieties; the same applies to vacancies.

### 5.1. Ideal part of the end-member activity term

Now, the approximation of chemical potential of  $j$ -th end member of  $k$ -th phase takes the form

$$\nu_j = \frac{g_{j,T,P}}{RT} + \ln a_j^{(con)} + \ln f_j \quad j \in L_k \quad (5.1-2),$$

where  $f_j$  is the ‘macroscopic’ end member activity coefficient, and  $a_j^{(con)}$  is the ideal part of activity related to the partial molar configurational entropy:

$$-R \ln a_j^{(con)} = \Delta \bar{S}_j^{(con)} - \Delta \bar{S}_j^{(con),o} \quad (5.1-3).$$

Following the derivation in (Price, 1985),

$$\Delta\bar{S}_j^{(con)} = -R \sum_s \sum_m \eta_{j,s,m} \ln y_{s,m} \quad m \in M_k \quad s \in \Xi_k \quad j \in L_k \quad (5.1-4),$$

where the site fraction of  $m$ -th moiety on  $s$ -th site in  $k$ -th phase is

$$y_{s,m} = \frac{n_{s,m}}{\sum_m n_{s,m}} \quad m \in M_k \quad (5.1-5).$$

here  $n_{s,m}$  is the number of moles of  $m$ -th moiety on  $s$ -th sublattice sites in the ‘real’ state.

At standard state, the  $s$ -th site fraction of  $m$ -th moiety in  $j$ -th end member is

$$y_{j,s,m}^o = \frac{\eta_{j,s,m}}{\eta_s} \quad (\text{‘site multiplicity’ } \eta_s = \sum_m \eta_{j,s,m}) \quad (5.1-6),$$

and the end member has a configurational entropy term itself:

$$\Delta\bar{S}_j^{(con),o} = -R \sum_s \sum_m \eta_{j,s,m} \ln y_{j,s,m}^o \quad (5.1-7)$$

Combination of eqs (5.1-4, 5.1-7 and 5.1-3) yields

$$\ln a_j^{(con)} = \sum_s \sum_m \eta_{j,s,m} \cdot \ln \left( \frac{y_{s,m}}{y_{j,s,m}^o} \right) \quad m \in M_k \quad s \in \Xi_k \quad j \in L_k \quad (5.1-8)$$

The site fraction  $y_{s,m}$  (eq 5.1-5) in the  $k$ -th phase can be expressed through mole fractions of end members  $x_j$  because

$$n_{s,m} = \sum_{j_k} \eta_{j_k,s,m} n_{j_k} = n_k \sum_{j_k} \eta_{j_k,s,m} x_{j_k} \quad j_k \in L_k \quad (5.1-9),$$

where  $n_{j_k}$  is the primal mole amount of  $j_k$ -th end member, and  $n_k = \sum_{j_k} n_{j_k}$  (here, summations

run over all end members in  $k$ -th phase). Using  $\eta_s = \sum_m \eta_{j,s,m}$ , eq (5.1-5) takes the form:

$$y_{s,m} = \frac{\sum_{j_k} \eta_{j_k,s,m} x_{j_k}}{\sum_m \sum_{j_k} \eta_{j_k,s,m} x_{j_k}} = \frac{1}{\eta_s} \sum_{j_k} \eta_{j_k,s,m} x_{j_k} \quad j_k \in L_k \quad m \in M_k \quad s \in \Xi_k \quad (5.1-10).$$

Substituting this and eq (5.1-6) into eq (5.1-8) results in the ideal activity of  $j$ -th end member

$$\ln a_j^{(con)} = \sum_s \sum_m \eta_{j,s,m} \{ \ln(\sum_{j_k} \eta_{j_k,s,m} x_{j_k}) - \ln \eta_{j,s,m} \} \quad j_k \in L_k \quad (5.1-11).$$

Now, eq (5.1-2) for the end member chemical potential takes a more general form:

$$\nu_j = \frac{g_{j,T,P}}{RT} + \sum_s \sum_m \eta_{j,s,m} \{ \ln(\sum_{j_k} \eta_{j_k,s,m} x_{j_k}) - \ln \eta_{j,s,m} \} + \ln f_j \quad (5.1-12).$$

At  $n(\Xi) = 1$  and all  $\eta_{j,s,m}$  numbers equal to unity, eq (5.1-11) reduces to  $\ln x_j$  – the simple ideal single-site mixing activity contribution – and eq (5.1-12) reduces to eq (5.1-1). In principle, any multi-site mixing model can be put on top of the  $\ln x_j$  term in eq (5.1-1) by introducing a ‘configurational entropy’ term  $\lambda_j$ , also called ‘fictive activity coefficient’:

$$\nu_j = \frac{g_{j,T,P}}{RT} + \ln x_j + \ln \lambda_j + \ln f_j \quad (5.1-13)$$

where

$$\ln \lambda_j = \sum_s \sum_m \eta_{j,s,m} \left\{ \ln \left( \sum_{j_k} \eta_{j_k,s,m} x_{j_k} \right) - \ln \eta_{j,s,m} \right\} - \ln x_j \quad (5.1-14).$$

In earlier versions of GEMS, this could be done only in Phase definition scripts which contained all the specific  $\eta_{j,s,m}$  numbers for a particular sublattice mixing model. This slowed calculations down and precluded such phase definitions from being used in GEMS3K kernel and, thus, in any reactive mass transport or other coupled codes. This work represents an extension that enables built-in ideal or non-ideal multi-site mixing models also in GEMS3K. For this extension in TSolMod class, it is necessary to use an array of  $\eta_{j,s,m}$  numbers arranged as a sequence of  $n(L_k)$  matrices  $\|\eta_{s,m}\|_j$ , each having  $n(\Xi_k)$  rows and  $n(M_k)$  columns. These matrices will be arranged in (part of) a MoiSN array (to be created in the MULTI data structure) then passed as a pointer into TSolMod class instance. The moieties will be arranged in a common list (without duplicate entries) for the solution phase. If the same cation or anion occurs in different sublattices then it must be considered as a different moiety on each sublattice site. For example, in ferrites, divalent and trivalent cations may be redistributed between tetrahedral and octahedral sites, e.g. in Ni-ferrite:  $(Ni_{1-z}Fe^{III}_z)_{Tet}(Ni_zFe^{III}_{2-z})_{Oct}O_4$ , where  $0 < z < 1$  is the extent of inversion which depends on temperature. At given temperature, e.g. setting  $z = 0.8$ , the Fe-Ni ferrite system may be represented using two end members:  $(Ni_{0.2}Fe^{III}_{0.8})_{Tet}(Ni_{0.8}Fe^{III}_{1.2})_{Oct}O_4$  and  $(Fe^{II}_{0.2}Fe^{III}_{0.8})_{Tet}(Fe^{II}_{0.8}Fe^{III}_{1.2})_{Oct}O_4$ . Note that Ni cation on the Tet sublattice is not the same moiety as Ni cation on the Oct sublattice. If vacancies are considered on different sites, the same applies to vacancies.

The most versatile way of providing the information about moieties occupancies on sites in the Phase definition seems to be through the formulae of end members. To do this, the formulae syntax needs to be extended. For example, the usual formulae of feldspar end members are

KAlSi3O8 (microcline);    NaAlSi3O8 (albite);    CaAl2Si2O8 (anorthite) etc.

This format does not contain information about sites and moieties that substitute on them. However, this information can be incorporated e.g. in the following way:

{K}:{Al}{Si}3:08    {Na}:{Al}{Si}3:08    {Ca}:{Al}2{Si}2:08

Essential feature here is that each moiety (incl. vacancy  $v_a$ ) is taken in the {} braces; the colon ‘:’ separates sublattice sites which must follow the same order in all end members. The  $\eta_{j,s,m}$  numbers appear just as stoichiometry coefficients after moieties. Upon parsing the formula, the number of sublattices is counted by encountered colons ‘:’ (two in the above example), and the moieties are collected in a list along with their site-moiety multiplicity  $\eta$  numbers and the site index. This list can then be merged into the overall list of  $n(M_k)$  moieties for the phase, and the  $\eta$  numbers then assigned to  $\|\eta_{s,m}\|_j$  tables, the sequence of which forms the

MoiSN array. The same ion can be represented as different moieties by adding the moiety index after species name. In the above example of ferrite with cation inversion, the formulae of end members will look like this (default valence of IC Ni is 2 and that of IC Fe is 3):

```
{Ni}0.2{Fe}0.8:{Ni}0.8{Fe}1.2:04
{Fe|2|}0.2{Fe|3|}0.8:{Fe|2|}0.8{Fe|3|}1.2:04
```

In the former end member, Ni is present on both sites, so the program creates two moieties: [Ni]0 and [Ni]1; it also creates two moieties for divalent Fe: [Fe]0 and [Fe]1. By analyzing

the latter end member,  $[Fe|3|]0$  and  $[Fe|3|]1$  will be added to the list of moieties, and so on.

Chemical formulae of DCs are parsed in GEM-Selektor using the syntax that is now extended, as shown below in boldface:

```

<formula> ::= <fterm> | <fterm> <charge>
<fterm>   ::= <icterm> | <fterm> <icterm> |
               <site_term> : | <fterm> <site_term> :
<site_term> ::= <moiety> | <site_term> <moiety>
<moiety>  ::= {<elem>} | {<elem>} <elem_st_coef>
<icterm>   ::= <elem> | <elem> <elem_st_coef>
<elem>     ::= (<fterm>) | [<fterm>] |
               <isotope_mass><icsymb><valence> |
               <isotope_mass><icsymb> |
               <icsymb><valence> | <icsymb>
<charge>   ::= +<real> | -<real> | + | - | @
<elem_st_coef> ::= <real>
<icsymb>   ::= <Capital_letter> | <icsymb><lcase_letter>
               | <icsymb>_ | va
<valence> ::= |-<integer>| | +<integer>| | <integer>|
<isotope_mass> ::= /<integer>/
<integer>  ::= <num>
<num>       ::= <digit> | <num><digit>
<digit>    ::= 0 | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9
<real>     ::= <num>.<num> | <num>. | .<num> | <num>

```

In the **<elem>** term, up to three levels of nesting is allowed. The vertical bar "|" means "or", i.e. one selection from several possible values or terms. The "va" in **<icsymb>** means "vacancy" and should be ignored everywhere except where moieties and site terms are analyzed. When the formula parser is called for checking the DComp or ReacDC formulae, all the { and } brackets must be replaced with ( and ) respectively, and colons : removed.

Otherwise, the site index should be incremented upon encountering the colon ':'. The newly encountered moieties (including va) is put into square brackets, appended with the sublattice index, and then added to the list of moieties, if not already present there, and the moiety index and site index are then used for inserting the respective occupancy number (if given after '}', 1 by default) into the occupancy table, which must exist for all end members in the phase. The strict order of site terms is essential. The total of moiety occupancies (the site multiplicity) for a given site is checked to be the same for this site term in all end members (see eq 5.1-6).

## 5.2. Non-ideal site-interaction part of the end-member activity term

after T.Wagner 21.06.11

To represent intra-site non-ideal interactions for substitutions of moieties on sublattices, the approach of Berman and Brown (1984), Berman (1990) is applied because of its relative simplicity and compatibility with ‘petrologic’ style of defining end members with more than one moiety on a sublattice site. So far, this approach ignores cross-site interactions (which at least partially can be accounted for using the reciprocal terms).

In our notation, the ideal contribution to the end member activity used in Berman’s models,

$$a_j^{(con)} = \prod_s \prod_m \left( y_{s,m} \frac{\eta_s}{\eta_{j,s,m}} \right)^{\eta_{j,s,m}} \quad m \in M_k \quad s \in \Xi_k \quad j \in L_k \quad (5.2-1)$$

is shown to be equivalent to eq (5.1-8) because from eq (5.1-6),  $\frac{1}{y_{j,s,m}^o} = \frac{\eta_s}{\eta_{j,s,m}}$ .

The non-ideal contribution to end-member activity (see eq 5.1-2) is defined as the sum of

$$RT \ln f_j = \sum_s f_{j,s} \quad s \in \Xi_k \quad j \in L_k \quad (5.2-2)$$

Furthermore,

$$\begin{aligned} f_{j,s} &= RT \sum_m \ln \gamma_{j,s,m} \\ RT \ln \gamma_{j,s,m} &= y_{j,s,m}^o \sum_t y_d y_e y_f W_{def,s} \left( Q_{s,m} \frac{y_{j,s,m}^o}{y_{s,m}} - \Theta \right) \quad t \in W_s \end{aligned} \quad (5.2-3)$$

where the index  $m$  refers only to moieties that are present on  $s$ -th site in  $j$ -th end member;  $Q_{s,m}$  is the number of  $d, e, f$  subscripts equal to  $m$  (0, 1 or 2);  $\Theta$  is the order of interaction (symmetric: 1; asymmetric: 2); and  $W_s$  is the subset of indexes  $p$  of entries in the list of interaction parameters that refer to  $s$ -th sublattice. The  $d, e$ , or  $f$  subscript can be an index  $m$  of any moiety that exists on the  $s$ -th site. Hence,  $W_{def,s}$  is the interaction parameter for a particular combination of moieties substituting each other on the  $s$ -th site. The  $T,P$  dependence of the interaction parameter is represented, as usual, by three coefficients  $a, b$ , and  $c$  (J,K,bar):

$$W = a - b \cdot T + c \cdot P \quad (5.2-4)$$

In the case of symmetric interaction, the  $f$  subscript does not exist, and eq (5.2-3) takes the form

$$RT \ln \gamma_{j,s,m} = y_{j,s,m}^o \sum_t y_d y_e W_{de,s} \left( Q_{s,m} \frac{y_{j,s,m}^o}{y_{s,m}} - 1 \right) \quad (5.2-5)$$

where  $y_{j,s,m}^o = \frac{\eta_{j,s,m}}{\eta_s}$  (eq 5.1-6). For the asymmetric interactions,

$$RT \ln \gamma_{j,s,m} = y_{j,s,m}^o \sum_t y_d y_e y_f W_{def,s} \left( Q_{s,m} \frac{y_{j,s,m}^o}{y_{s,m}} - 2 \right) \quad (5.2-6)$$

where either  $d$  and  $e$  or  $e$  and  $f$  indexes refer to the same moiety on  $s$ -th site in the case of a pseudo-binary interaction, and all three indexes are different in a ternary interaction.

In addition to tables  $\|\eta_{s,m}\|_j$  and  $\|y_{s,m}\|$  taken over from the calculation of ideal contributions to end member activities (see Section 5.1), another set of tables  $\|f_{j,s}\|_j$  is needed in the algorithm to collect the contributions to terms  $RT \ln \gamma_{j,s,m}$  (see eq 5.2-3).

In the TSolMod class implementation, the setup of interaction parameters (IP) is done in the Phase definition record. Upon calculation of equilibria, the IP list is passed automatically into the TSolMod instance for calculation of end-member activity coefficients. In the microscopic non-ideality model, the IP list needs to be arranged in the following format:

IP index	Site index	Moiety indexes	IP coefficients (J, K, bar)				
<i>p</i>	<i>s</i>	<i>d</i>	<i>e</i>	<i>f</i>	<i>a</i>	<i>b</i>	<i>c</i>
0	0	0	1	-1	-6000	-1.3	0.4
1	1	3	3	4	17000	5.1	0.1
...	...	...	...	...	...	...	...

IP coefficients refer to eq (5.2-4),  $W_{def,s} = a - b \cdot T + c \cdot P$  (note the minus sign at  $bT$  term),  $T$  in K,  $P$  in bar,  $a$  in J/mol.

### Example: Muscovite (white mica) solid solution (Parra et al. 2002)

End-members:

muscovite (mu), celadonite (cel), ferrous celadonite (fcel), Na-celadonite (ncel), Na-ferrous celadonite (nfcel), paragonite (pa), and pyrophyllite (prl).

Moiety- sublattice site allocation table (encoded in end-member formulae).

Sublattice End member	A (0)	M23 (1)	T2 (2)	T1 (3)	End-member formula (as written in DComp record)
(0) fcel	K	Al Fe	Si <sub>2</sub>	Si <sub>2</sub>	{K}:{Al}{Fe}:{Si}2:{Si}2:O10(OH)2
(1) cel	K	Al Mg	Si <sub>2</sub>	Si <sub>2</sub>	{K}:{Al}{Mg}:{Si}2:{Si}2:O10(OH)2
(2) mu	K	Al <sub>2</sub>	Al Si	Si <sub>2</sub>	{K}:{Al}2:{Al}{Si}:{Si}2:O10(OH)2
(3) nfcel	Na	Al Fe	Si <sub>2</sub>	Si <sub>2</sub>	{Na}:{Al}{Fe}:{Si}2:{Si}2:O10(OH)2
(4) ncel	Na	Al Mg	Si <sub>2</sub>	Si <sub>2</sub>	{Na}:{Al}{Mg}:{Si}2:{Si}2:O10(OH)2
(5) pa	Na	Al <sub>2</sub>	Al Si	Si <sub>2</sub>	{Na}:{Al}2:{Al}{Si}:{Si}2:O10(OH)2
(6) prl	Va	Al <sub>2</sub>	Si <sub>2</sub>	Si <sub>2</sub>	{Va}:{Al}2:{Si}2:{Si}2:O10(OH)2

Microscopic interaction parameters (J, K, bar).

$W_{KKNa,A} = 19460 - 1.65 \cdot T - 0.46 \cdot P$
$W_{KNaNa,A} = 12230 - 5.0 \cdot T + 0.67 \cdot P$
$W_{KKVa,A} = 35000 + 25.0 \cdot T - 0.85 \cdot P$
$W_{KVaVa,A} = 45000 + 10.0 \cdot T - 0.85 \cdot P$
$W_{NaNaVa,A} = 40000 + 5.0 \cdot T$
$W_{NaVaVa,A} = 40000 + 5.0 \cdot T$
$W_{KNaVa,A} = 95480 + 19.13 \cdot T - 0.745 \cdot P$
$W_{AlMg,M23} = -30500 + 15.0 \cdot T + 0.78 \cdot P$
$W_{AlFe,M23} = -5500 + 15.0 \cdot T + 0.65 \cdot P$
$W_{MgFe,M23} = 0$
$W_{AlSi,T2} = 0$

In GEMS Phase window (Page 3), the recalculation results in the following allocation table:

End member- moiety multiplicity number table (multiplicity numbers can be non-integer).

m	0	1	2	3	4	5	6	7	8
j	{K}0	{Al}1	{Fe}1	{Si}2	{Si}3	{Mg}1	{Al}2	{Na}0	{Va}0
0	1	1	1	2	2	0	0	0	0
1	1	1	0	2	2	1	0	0	0
2	1	2	0	1	2	0	1	0	0
3	0	1	1	2	2	0	0	1	0
4	0	1	0	2	2	1	0	1	0
5	0	2	0	1	2	0	1	1	0
6	0	2	0	2	2	0	0	0	1

Indexes m and j are shown for the reader's convenience

To set up interaction parameters, remake the Phase record and set the Berman (B) built-in (S) model with total number of input interaction parameters (in this example, 11). On Page 3 of Phase window, you will see a combined list filled with zeros.

As shown below, for each interaction parameter, the indexes of sublattice s and moieties d,e,f should be entered into four ipxT columns (for a symmetric interaction parameter, index f should be set to -1). For each moiety, its sublattice index can be found right after its name, e.g. {K}0. For example, the parameter  $W_{KKNa,A} = 19460 - 1.65 \cdot T - 0.46 \cdot P$  will be set as shown below:

s	d	e	f	a	b	c
ipxT[0]	ipxT[1]	ipxT[2]	ipxT[3]	ph_cf[0]	ph_cf[1]	ph_cf[2]
0	0	0	7	19460	-1.65	-0.46
...	...	...	...	...	...	...

The complete list for this example is shown below. The interaction parameters (rows) can be given in an arbitrary sequence. Two rows with zero IPs can, in principle, be dropped (by re-

making the Phase record and setting the number of rows to 9 instead of 11). However, the sequence of end members and moieties is essential; if it changes, the indexation in ipxT columns must be revised according to the new moiety allocation table.

Interaction parameters setup in Phase record for the Muscovite example.

ipxT[0]	ipxT[1]	ipxT[2]	ipxT[3]	ph_cf[0]	ph_cf[1]	ph_cf[2]
0	0	0	7	19460	-1.65	-0.46
0	0	7	7	12230	-5	0.67
0	0	0	8	35000	25	-0.85
0	0	8	8	45000	10	-0.85
0	7	7	8	40000	5	0
0	7	8	8	40000	5	0
0	0	7	8	95840	19.13	-0.745
1	1	5	-1	-30500	15	0.78
1	1	2	-1	-5500	15	0.65
1	5	2	-1	0	0	0
2	6	3	-1	0	0	0

Worked test examples of multi-site solid solution models with Berman microscopic non-ideality will be provided in the BermanMSS test project and its documentation (TBD).

### 5.3. Reciprocal part of the end-member activity term

The typical multi-site solid solution with two or more sublattices and two or more moieties substituted on each sublattice is called *reciprocal*. Meaning of this term is best illustrated using the system {A,B}{X,Y} (e.g. Wood and Nicholls, 1978; Hillert, 2001). All possible compositions can be represented on a ‘composition square’ formed by four possible end members: AX, BX, AY, BY. There are four end members, but only three are needed to describe any bulk composition of this solid solution phase. For instance, the center of the square can be obtained by mixing equal amounts of either AX and BY or AY and BX, hence the name ‘reciprocal’. Any three of four end members can be declared as ‘independent’, while the remaining one will be ‘dependent’ because it’s composition and thermodynamic properties can be obtained from that of ‘independent’ end members and that of a *reciprocal reaction*



with a standard molar Gibbs energy effect of reaction

$$\Delta^oG_{(A,B)(D,E)} = G^o_{AX} + G^o_{BY} - G^o_{AY} - G^o_{BX}$$

For example, if the BE end member is declared as ‘dependent’ then it’s standard molar Gibbs energy can be obtained from eq (5.3-1) as  $G^o_{BY} = \Delta^oG_{(A,B)(X,Y)} + G^o_{AY} + G^o_{BX} - G^o_{AX}$ , provided that the Gibbs energy effect of the reciprocal reaction is known (e.g. fitted from the experimental data). There is also a condition of internal equilibrium:  $\mu_{AX} + \mu_{BY} = \mu_{AY} + \mu_{BX}$  (Wood and Nicholls, 1978). Following the derivations in this paper, assuming random ideal mixing on both sublattices, the chemical potentials of end members in this system are:

$$\mu_{AX} = G^o_{AX} + RT\ln(y_{0,A}y_{1,X}) - y_{0,B}y_{1,Y}\Delta G^o_{(A,B)(X,Y)}$$

$$\mu_{BY} = G^o_{BY} + RT\ln(y_{0,B}y_{1,Y}) - y_{0,A}y_{1,X}\Delta G^o_{(A,B)(X,Y)}$$

$$\begin{aligned}\mu_{AY} &= G_{AY}^o + RT \ln(y_{0,A}y_{1,Y}) + y_{0,B}y_{1,X}\Delta G_{(A,B)(X,Y)}^o \\ \mu_{BX} &= G_{BX}^o + RT \ln(y_{0,B}y_{1,X}) + y_{0,A}y_{1,Y}\Delta G_{(A,B)(X,Y)}^o\end{aligned}$$

Comparing these with the general form of the chemical potential, we see that the terms like  $y_{0,B}y_{1,Y}\Delta G_{(A,B)(X,Y)}^o$  stay in place of one more activity coefficient like  $\gamma_{AX}$ :

$$\begin{aligned}RT \ln \gamma_{AX} &= -y_{0,B}y_{1,Y}\Delta G_{(A,B)(X,Y)}^o \\ RT \ln \gamma_{BY} &= -y_{0,A}y_{1,X}\Delta G_{(A,B)(X,Y)}^o \\ RT \ln \gamma_{AY} &= +y_{0,B}y_{1,X}\Delta G_{(A,B)(X,Y)}^o \\ RT \ln \gamma_{BX} &= +y_{0,A}y_{1,Y}\Delta G_{(A,B)(X,Y)}^o\end{aligned}$$

Thus, the complete form of the end-member chemical potential will be

$$\mu_j = \mu_j^o + RT \ln x_j + RT \ln \lambda_j + RT \ln f_j + RT \ln \gamma_j \quad (5.3-2)$$

where  $RT \ln \gamma_j$  is the reciprocal activity coefficient term, which turns to zero (ideal behaviour) only if the Gibbs energy effect of the reciprocal reaction (5.3-1)  $\Delta G_{(A,B)(X,Y)}^o = 0$ . This formalism can be extended for the case when several moieties substitute on each of two sublattices (Wood and Nicholls, 1978; Aranovich, 1991). The number of end members in this case is  $M_0 M_I$ ; the number of independent end members is  $M_0 + M_I - 1$ ; and the number of all possible reciprocal reactions is the number of combinations  $C(M_0 M_I, M_0 + M_I - 1)/4$ .

Unfortunately, it is very difficult to generalize this “reciprocal reaction” formalism to solid solutions with simultaneous substitutions in three or more sublattices. For the general case, Sundman and Ågren (1981) suggested a method related to the site fractions matrix Y made of the  $y_{s,m}$  elements (see eq 5.1-5). The total Gibbs energy per mole of k-th phase is represented as

$$G_k = G_k^{ref} - TS_{k,id}^{conf} + G_k^{EX} \quad (5.3-3)$$

where  $G_k = \sum_j \mu_j x_j$ ;  $S_{k,id}^{conf} = \sum_j x_j \Delta \bar{S}_j^{(con)}$  where  $\Delta \bar{S}_j^{(con)}$  is given by eq (5.1-4);  $G_k^{EX} = RT \sum_j x_j \ln f_j$  where  $\ln f_j$  is given by eq (5.2-2); and  $G_k^{ref}$  is the so-called frame-of-reference Gibbs energy (Sundman and Ågren, 1981; Hillert, 1998).

For the substitution on one sublattice,  $G_k^{ref} = \sum_{m_0}^{M_0} y_{0,m_0} G_{m_0}^o = \sum_j x_j G_j^o$ .

For simultaneous substitutions on two sublattices,  $G_k^{ref} = \sum_{m_0}^{M_0} \sum_{m_1}^{M_1} y_{0,m_0} y_{1,m_1} G_{m_0 m_1}^o$  where  $G_{m_0 m_1}^o$  is the standard-state Gibbs energy of the end-member compound made of a moiety  $m_0$  on sublattice 0 and moiety  $m_1$  on sublattice 1.

For three sublattices,  $G_k^{ref} = \sum_{m_0}^{M_0} \sum_{m_1}^{M_1} \sum_{m_2}^{M_2} y_{0,m_0} y_{1,m_1} y_{2,m_2} G_{m_0 m_1 m_2}^o$ , and so on.

Sundman and Ågren (1981) suggested the following general notation:

$$G_k^{ref} = \sum_I G_I^o \wp_I(Y) \quad (5.3-4)$$

where  $I$  is the “component array” that defines one moiety on each sublattice, and  $\wp_I(Y)$  stands for the corresponding product of site fractions from the Y matrix. The CEF (compound energy formalism) eq (5.3-3) assumes that in any end member, there is exactly one moiety on each sublattice. In the formalism considered in Sections 5.1 and 5.2, end members with fractional site occupation are allowed, i.e with two or more moieties on one sublattice with end

member site fractions  $y_{j,s,m}^o = \frac{\eta_{j,s,m}}{\eta_s}$  (see eq 5.1-6). To account for this case, we assume that  $\wp_{I0}(Y) = \wp_j(Y)$  is the product of sums  $y_{s,\sigma}^{(j)}$ :

$$y_{s,\sigma}^{(j)} = \sum_m y_{j,s,m}^o y_{s,m} = \frac{1}{\eta_s} \sum_m \eta_{j,s,m} y_{s,m}. \quad (5.3-5)$$

Clearly, the CEF simply means that there is only one m-th non-zero element  $\eta_{j,s,m} = \eta_s$  per j-th end member in s-th sublattice, so the  $y_{s,\sigma}^{(j)}$  turns into  $y_{s,m_j}$ , the site fraction of moiety  $m_j$  which is present on s-th site in j-th end member formula.

The chemical potential of j-th end member can be obtained as a partial derivative of eq (5.3-3) on the amount of j-th end member,

$$\mu_j = G_k + \sum_s \left( \frac{\partial G_k}{\partial y_{s,m_j}} - \sum_m \frac{\partial G_k}{\partial y_{s,m}} \right)_{T,P,n_{l \neq j}} \quad (5.3-6).$$

(Sundman and Ågren, 1981). Because of the additivity of  $G_k$  function, one can write

$$RT \ln \gamma_j = G_k^{ref} + \sum_s \left( \frac{\partial G_k^{ref}}{\partial y_{s,m_j}} - \sum_m y_{s,m} \frac{\partial G_k^{ref}}{\partial y_{s,m}} \right) - G_j^o, \quad s \in \Xi_k, \quad m \in M_k \quad (5.3-7)$$

where

$$\frac{\partial G_k^{ref}}{\partial y_{s,m_j}} = \sum_l \wp_l(Y) \cdot \frac{\delta_{j,s,m_j}}{y_{s,m}} \cdot G_l^o, \quad l \in L_k \quad (5.3-8)$$

$$\frac{\partial G_k^{ref}}{\partial y_{s,m}} = \sum_l \wp_l(Y) \cdot \frac{\delta_{l,s,m}}{y_{s,m}} \cdot G_l^o, \quad l \in L_k \quad (5.3-9)$$

Here,  $\delta_{j,s,m}$  is the special form of Kronecker's delta:  $\delta_{j,s,m} = 1$  if  $\eta_{j,s,m} \neq 0$ , else  $\delta_{j,s,m} = 0$ .

There are two limitations of the formalism eqs (5.3-4) to (5.3-9): (i) partially-substituted end members are not allowed; (ii) all possible end members must be included into the model. If these two conditions are not satisfied then the reciprocal energy contribution terms cannot be computed correctly and must be disabled.

From eq. (5.3-3) it is evident that the configurational entropy contribution is the same in Berman approach and in CEF model, whereas the excess energy contribution in Berman model is not the same as in CEF. Also, the reciprocal energy contributions were ignored in most known applications of Berman's model (maybe except those for systems with two sublattices, cf. Aranovich 1991). Therefore, in TSolMod library, two models of this kind are implemented: Berman (coded 'B') and CEFmod (coded '\$'), see Appendix 5.2.

#### 5.4. Non-ideal excess energy part of the end-member activity term in CEFmod model

<under construction>

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**Appendix 5.2.** Built-in options for activity coefficients for end members of condensed (solid and liquid) non-electrolyte solution phases (GEM-Selektor v. 3.3).

Code	Model	Equations	Interaction parameters (to enter into <b>ph_cf</b> array)
<b>G</b>	Redlich-Kister model (binary)	4.2-36 to 4.2-38	three constant binary interaction parameters
<b>M</b>	Subregular model (binary)	4.2-10 to 4.2-13	two binary interaction parameters, each represented with 3 coefficients as $a - b \cdot T + c \cdot P$
<b>T</b>	Regular model (ternary)	4.2-21 to 4.2-24	one binary interaction parameter per pair and 1 ternary interaction parameter, each represented with 3 coefficients as $a - b \cdot T + c \cdot P$
<b>R</b>	Regular model (multicomponent)	4.2-29, 4.2-30	one binary interaction parameter per pair of components, represented as $a + b \cdot T + c \cdot P$
<b>V</b>	Van Laar model (multicomponent)	4.2-31 to 4.2-35	one binary interaction parameter per pair of components, represented as $a + b \cdot T + c \cdot P$ one size parameter per component (end member) in <b>dc_cf</b> array
<b>B</b>	Berman multisite (multicomponent)	5.1-14 to 5.2-6	one binary or ternary interaction parameter for moieties on a sublattice site, represented as $a - b \cdot T + c \cdot P$
<b>\$</b>	CEF multisite (multicomponent)	5.1-14; 5.3-6; 5.4-	one binary, ternary or quaternary interaction parameter for moieties on a sublattice site, represented as $a + b \cdot T + c \cdot T \cdot \ln(T) + d \cdot P$
<b>K</b>	Redlich-Kister (multicomponent)	4.2-41, 4.2-42	four binary interaction parameters per pair of components, represented as $a + b \cdot T + c \cdot T \cdot \ln(T) + d \cdot P$
<b>L</b>	NRTL model (multicomponent)	4.2-50, 4.2-51	two normalized binary interaction energies per pair of components, represented as $A + B/T + C \cdot T + D \cdot \ln(T)$ 1 parameter $\alpha$ per pair of components: $E + F \cdot (T - 273.15)$
<b>W</b>	Wilson model (multicomponent)	4.2-59, 4.2-60	two binary interaction energies per pair of components, represented as $\exp[A + B/T + C \cdot T + D \cdot \ln(T)]$
<b>U</b>	User-defined		Must be provided as Phase scripts and parameters