## Part 1. Temperature corrections of standard molar thermodynamic properties of condensed substances and gases using data in DComp records of GEM-Selektor

This calculation is derived from integration of the heat capacity equation Cp = f(T) which has the following form:

$$Cp = a_0 + \sum_i a_i T^{n_i}, \quad i = 1, ..., 9$$
 (1-1)

where  $a_i$  are the empirical coefficients, and power coefficients  $n_i$  are:  $n_0 = 0$ ;  $n_1 = 1$ ;  $n_2 = -2$ ;  $n_3 = -0.5$ ;  $n_4 = 2$ ;  $n_5 = 3$ ;  $n_6 = 4$ ;  $n_7 = -3$ ;  $n_8 = -1$ ;  $n_9 = 0.5$ . Some of these terms are used in most published databases. The corresponding isobaric heat capacity equation is:

$$Cp = a_0 + a_1 T + a_2 T^{-2} + a_3 T^{-0.5} + a_4 T^2 + a_5 T^3 + a_6 T^4 + a_7 T^{-3} + a_8 T^{-1} + a_9 T^{0.5}.$$
 (1-2).

As known from chemical thermodynamics [e.g. Anderson and Crerar, 1993], the increment of standard molar Gibbs energy  $G^o$  at the reference pressure  $P_r = 1$  bar from reference temperature  $T_r$  to temperature of interest T (in K) can be found by integrating the absolute molar entropy  $S^o$ :

$$G_T^o - G_{T_r}^o = -\int_{T_r}^T S_T^o dT (1-3),$$

where  $S_T^o = \int_0^T \frac{Cp}{T} dT = S_{T_r}^o + \int_{T_r}^T \frac{Cp}{T} dT$ . Equation (1-3) can be transformed as follows:

$$G_{T}^{o} - G_{T_{r}}^{o} = -S_{T_{r}}^{o} (T - T_{r}) - \int_{T_{r}}^{T} \int_{T_{r}}^{T} Cp / T dT dT = -S_{T_{r}}^{o} (T - T_{r}) - T \sum_{i} M_{n_{i}} a_{i}$$
(1-4),

where the  $M_{n_i}$  terms stand for the Temkin-Shwarzman functions of temperature [Dorogokupets et al., 1988]:

$$M_{n_i} = \frac{T^{n_i}}{n_i(n_i+1)} + \frac{T_r^{n_i+1}}{T(n_i+1)} - \frac{T_r^{n_i}}{n_i}.$$

Using these functions and coefficients  $n_i$  from eqn. (1-1), separate terms can be defined:

$$\begin{split} F_{G0} &= TM_o = T \ln \frac{T}{T_r} - T + T_r; \\ F_{G1} &= TM_1 = \frac{1}{2} (T - T_r)^2; \\ F_{G2} &= TM_{-2} = (T - T_r)^2 / (2T \cdot T_r^2); \\ F_{G3} &= TM_{-0.5} = 2 \left( \sqrt{T} - \sqrt{T_r} \right)^2 / \sqrt{T_r}; \\ F_{G4} &= TM_2 = \left( T^3 + 2T_r^3 - 3T \cdot T_r^2 \right) / 6; \end{split}$$

$$\begin{split} F_{G5} &= TM_3 = \left(T^4 + 3T_r^4 - 4T \cdot T_r^3\right)/12; \\ F_{G6} &= TM_4 = \left(T^5 + 4T_r^5 - 5T \cdot T_r^4\right)/20; \\ F_{G7} &= TM_{-3} = \left(T_r^3 - 3T^2T_r + 2T^3\right)/\left(6T^2T_r^3\right); \\ F_{G8} &= TM_{-1} = \frac{T}{T_r} - 1 - \ln \frac{T}{T_r}; \\ F_{G9} &= TM_{0.5} = \frac{2}{3}\left(2T\sqrt{T} - 3T\sqrt{T_r} + T_r\sqrt{T_r}\right). \end{split}$$

Now, the standard molar apparent Gibbs energy function  $g^o_T$  of a substance can be calculated easily at reference pressure  $P_r = 1$  bar and temperature of interest  $T \neq T_r$  (usually, reference temperature  $T_r = 298.15$  K):

$$g_T^o = G_{T_r}^o - S_{T_r}^o (T - T_r) - T \sum_i M_{n_i} a_i =$$

$$= G_{T_r}^o - S_{T_r}^o (T - T_r) - a_0 F_{G0} - a_1 F_{G1} - a_2 F_{G2} - a_3 F_{G3} - a_4 F_{G4} -$$

$$- a_5 F_{G5} - a_6 F_{G6} - a_7 F_{G7} - a_8 F_{G8} - a_9 F_{G9}$$
(1-5)

In a similar way, the standard molar entropy  $S_T^o$  at  $P_r$  and T can be calculated

$$S_{T}^{o} = \int_{0}^{T} \frac{Cp}{T} dT = S_{T_{r}}^{o} + \int_{T_{r}}^{T} \frac{Cp}{T} dT = S_{T_{r}}^{o} + a_{0} \ln \frac{T}{T_{r}} + a_{1}(T - T_{r}) + \frac{1}{2} a_{2} \left( \frac{1}{T_{r}^{2}} - \frac{1}{T^{2}} \right) + 2a_{3} \left( \frac{1}{\sqrt{T_{r}}} - \frac{1}{\sqrt{T}} \right) + \frac{1}{2} a_{4} \left( T^{2} - T_{r}^{2} \right) + \frac{1}{3} a_{5} \left( T^{3} - T_{r}^{3} \right) + \frac{1}{4} a_{6} \left( T^{4} - T_{r}^{4} \right) + \frac{1}{3} a_{7} \left( \frac{1}{T_{r}^{3}} - \frac{1}{T^{3}} \right) + \frac{1}{4} a_{8} \left( \frac{1}{T_{r}} - \frac{1}{T} \right) + 2a_{9} \left( \sqrt{T} - \sqrt{T_{r}} \right)$$

$$(1-6),$$

as well as the standard molar apparent enthalpy function  $i_T^o$ :

$$i_{T}^{o} = H_{T_{r}}^{o} + \int_{T_{r}}^{T} Cp \, dT = H_{T_{r}}^{o} + a_{0}(T - T_{r}) + \frac{1}{2}a_{1}(T^{2} - T_{r}^{2}) + a_{2}(\frac{1}{T_{r}} - \frac{1}{T}) + \\
+ 2a_{3}(\sqrt{T} - \sqrt{T_{r}}) + \frac{1}{3}a_{4}(T^{3} - T_{r}^{3}) + \frac{1}{4}a_{5}(T^{4} - T_{r}^{4}) + \frac{1}{5}a_{6}(T^{5} - T_{r}^{5}) + \\
\frac{1}{2}a_{7}(\frac{1}{T_{r}^{2}} - \frac{1}{T^{2}}) + a_{8}\ln\frac{T}{T_{r}} + \frac{2}{3}a_{9}(T\sqrt{T} + T_{r}\sqrt{T_{r}})$$
(1-7)

For any compound substance, the apparent molar  $g_T^o$  and  $i_T^o$  can be converted into standard molar Gibbs energy  $G_{f,T}^o$  and enthalpy  $H_{f,T}^o$  of formation from elements, respectively:

$$G_{f,T}^{o} = g_{T}^{o}(\text{compound}) - \sum g_{T}^{o}(\text{elements});$$
 (1-8)

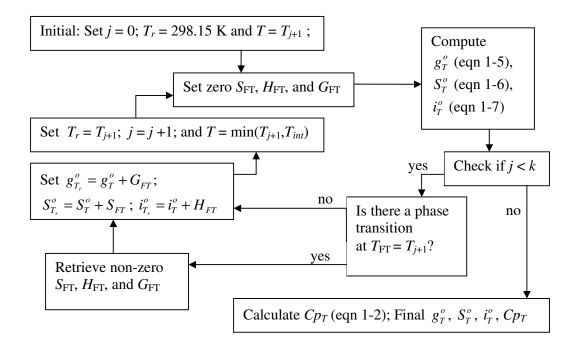
$$H_{f,T}^o = i_T^o \text{ (compound)} - \sum i_T^o \text{ (elements)}.$$
 (1-9)

It follows that at  $T_r$  =298.15 K,  $g_{298}^o = G_{f,298}^o$  and  $i_{298}^o = H_{f,298}^o$  because, for any element at its standard state,  $g_{T_r}^o = G_{f,T_r}^o = 0$  and  $i_{T_r}^o = H_{f,T_r}^o = 0$  by convention. Note that in Selektor codes (as well as in SUPCRT92 [Johnson et al., 1992]), apparent molar functions  $g_T^o$  and  $i_T^o$ , but not the formation functions  $G_{f,T}^o$  and  $H_{f,T}^o$  are calculated throughout at elevated temperatures.

It is possible that, within a wide temperature range, the heat capacity function Cp = f(T) is approximated by more than one set of coefficients of the form eqns (1-1) and (1-2). In the DComp data format, up to 10 such sets are allowed, and each set of coefficients must be supplied with a temperature applicability interval  $[T_{min}; T_{max}]$ . No break is allowed within the whole temperature range covered by n(Q) sets; i.e.  $T_{min}^{(q)} = T_{max}^{(q-1)}$ , q > 0, q = 0, 1, ..., n(Q).

For simplicity, let  $T_0$  be the lowest temperature for the first Cp approximation interval;  $T_1$  be the upper temperature for the latter ( $T_0$  to  $T_1$ ) and at the same time – the lower boundary for the next interval ( $T_1$  to  $T_2$ ), and so on. Usually,  $T_0$  equals 273.15 or 298.15 K, and some of the intermediate temperature boundaries refer also to temperatures of phase transitions at  $P = P_r$ .

Let k be the index of the temperature interval containing the temperature of interest  $T_{int}$ . Obviously, if k = 0 and  $T_0 \le T_r < T_1$  then  $T_0 \le T_{int} \le T_1$  and eqns (1-2) and (1-5) to (1-7) apply directly using the first set of Cp = f(T) coefficients. If k > 0 then the value of molar Cp at  $T = T_{int}$  where  $T_k \le T_{int} \le T_{k+1}$  is still found from eqn (1-2) applied directly to the k-th set of Cp coefficients. However, the values of  $g_T^o$ ,  $i_T^o$  and  $S_T^o$  must be calculated incrementally in k loops, as shown in the flow chart below, which also includes accounting for optional molar increments  $S_{FT}$ ,  $H_{FT}$ ,  $G_{FT}$  at phase transitions. It is required that the temperature of any phase transition  $T_{FT}$  (at  $P_r$ ) always co-insides with one of the boundary temperatures  $T_{j+1}$ .



Note that usage of GEM solvers of equilibria does not really require representing phase transitions in a single DComp record as for a single dependent component. Polymorph modifications, as well as solid, liquid and gaseous states of a substance can be represented as different phases or phase components (i.e. separate DComp records), because GEM will always select the most stable one at T,P of interest. Besides, this also allows circumventing the problem of pressure dependence of the phase transition temperature.

## **References to Part 1**

Anderson G.M. and Crerar D.A. (1993): *Thermodynamics in geochemistry: The equilibrium model*. Oxford Univ. Press, N.Y.,Oxford.

Dorogokupets P.I., Karpov I.K., Lashkevich V.V., Naigebauer V.A. and Kazmin L.A. (1988): Isobaric-isothermal potentials of minerals, gases and aqueous species in the "Selektor" program package. In: *Physicochemical models in geochemistry* (eds. Kiselev A.I. and Karpov I.K.), Nauka Publ., Novosibirsk, p. 124-147 (in Russian).

Johnson J.W., Oelkers E. and Helgeson H.C. (1992): SUPCRT92: A software package for calculating the standard molal thermodynamic properties of minerals, gases, aqueous species, and reactions from 1 to 5000 bar and 0 to 1000 °C. *Computers and Geosciences* **18**, 899-947.

**Appendix 1-1.** Built-in options for calculation of temperature and pressure correction to standard molar (partial molal) properties of substances and reactions using data in DComp records (GEM-Selektor v.2.2.0 PSI).

Code	Model	Input parameters
CSC	integration of heat capacity, constant molar volume	empirical $Cp$ coefficients $a_0$ to $a_9$ entered into <b>aiCpT</b> array; optional phase transition parameters entered into <b>FT_p</b> array
CSK	integration of heat capacity, molar volume from empirical $Vm = f(T,P)$ function	empirical $Cp$ coefficients $a_0$ to $a_9$ entered into <b>aiCpT</b> array; 5 empirical coefficients for molar volume entered into <b>aiVTP</b> array; optional phase transition parameters entered into <b>FT_p</b> array
CSE	integration of heat capacity, molar volume from HP Birch-Murnaghan equation	empirical <i>Cp</i> coefficients $a_0$ to $a_9$ entered into <b>aiCpT</b> array; isothermal compressibility and isobaric expansibility into <b>BetAlp</b> fields
CHE	integration of heat capacity with Landau contribution, molar volume from HP Birch-Murnaghan equation	empirical $Cp$ coefficients $a_0$ to $a_9$ entered into <b>aiCpT</b> array; isothermal compressibility and isobaric expansibility into <b>BetAlp</b> fields; $S$ and $T$ of Lambda transition into <b>LamST</b> fields; $Vx$ into <b>ab[0]</b> field
CSB	integration of heat capacity, molar volume from Birch- Murnaghan equation	empirical $Cp$ coefficients $a_0$ to $a_9$ entered into aiCpT array; coefficients of BM equation into aiODc array
css	integration of heat capacity, <i>VdP</i> contribution from Churakov-Gottschalk fluid EoS	empirical <i>Cp</i> coefficients $a_0$ to $a_9$ entered into aiCpT array; coefficients of CG fluid EoS into Cemp_s array
CSR	integration of heat capacity, <i>VdP</i> contribution from Peng-Robinson-Stryjek-Vera fluid EoS	empirical <i>Cp</i> coefficients $a_0$ to $a_9$ entered into aiCpT array; coefficients of PRSV fluid EoS into CritPg array
HWF	water properties from Haar-Gallagher-Kell (HGK) model	only standard state data need to be provided; paramters of HKF model in aihkf array are all set to zero
HKF	aqueous species properties from revised Helgeson-Kirk- ham-Flowers (HKF) model	coefficients of revised HKF model into aihkf array (in their original calorie-based units)
НКР	prediction of HKF parameters of complexes using Parcor correlation algorithm	DComp record is automatically created from ReacDC calculation in PRONSPREP/Parcor mode. Only recalculation of DComp is required to obtain correct $Cp^{\circ}$ , $V^{\circ}$ and $H^{\circ}$ . Upon this, the code will be automatically changed to <b>HKF</b> .