

Semi-empirical Gibbs free energy formulations for minerals and fluids for use in thermodynamic databases of petrological interest

Journal Article

Author(s):

Gerya, Taras; Maresch, W.V.; Podlesskii, K.K.; Perchuk, L.L.

Publication date:

2004-09

Permanent link:

https://doi.org/10.3929/ethz-b-000160343

Rights / license:

In Copyright - Non-Commercial Use Permitted

Originally published in:

Physics and chemistry of minerals 31(7), https://doi.org/10.1007/s00269-004-0409-8

ORIGINAL PAPER

T. V. Gerya K. K. Podlesskii L. L. Perchuk W. V. Maresch

Semi-empirical Gibbs free energy formulations for minerals and fluids for use in thermodynamic databases of petrological interest

Received: 13 October 2003/Accepted: 24 May 2004

Abstract The P-T partition function in statistical thermodynamics can be used to derive semi-empirical formulations of the Gibbs free energy G for minerals and fluids. Parameterization of these equations includes simultaneous regression of experimental heat capacity and molar volume data, allowing fitting, appraisal and optimization of various data sources, as required in the construction of internally consistent petrological data bases. This approach can also be extended to minerals with λ -transitions and to fluids by considering the Gibbs free energy as a function of pressure P, temperature T and an ordering parameter X_{α} , so that accurate modelled representation and extrapolation of the thermodynamic properties of large numbers of petrologically significant minerals and coexisting fluids can be attained. The ordering parameter is chosen to denote the equilibrium mole fraction (thermodynamic probability) of ordered clusters (structural units) in a substance when G(T,P), X_{α}) = min. The procedure is tested on existing experimental data for the system MgO-SiO₂-H₂O. The proposed Gibbs free energy formulation permits thermodynamic properties of minerals, fluids and phase equilibria to be described and extrapolated over a wide range of pressure (0-800 kbar) and temperature

T. V. Gerya · W. V. Maresch Institut für Geologie, Mineralogie und Geophysik, Fakultät für Geowissenschaften, Ruhr-Universität Bochum, 44780 Bochum, Germany

T. V. Gerya · K. K. Podlesskii · L. L. Perchuk Institute of Experimental Mineralogy, Russian Academy of Sciences Chernogolovka, Moscow district, 142432, Russia

Present address: T. V. Gerya (☒) Geologisches Institut - ETH - Zürich, CH-8092 Zurich, Switzerland e-mail: taras.gerya@erdw.ethz.ch

Tel.: +41-1-6322336 Fax: +41-1-6321030

L. L. Perchuk

Department of Petrology, Geological Faculty, Moscow State University, Vorobievy Gory, Moscow, 119899 Russia

(20–3000 K), thus allowing effective use in thermodynamic data bases of petrological interest.

Keywords Equations of state · Thermodynamic data bases · Partition function · Ordering—disordering processes · Molecular clusters in fluids

Introduction

Internally consistent thermodynamic databases (e.g. Karpov et al. 1976; Helgeson et al. 1978; Dorogokupets and Karpov 1984; Berman 1988; Grevel 1995; Holland and Powell 1990, 1998; Chatterjee et al. 1994, 1998; Berman and Aranovich 1996; Gottschalk 1997; Saxena et al. 1993) are widely used for the construction of petrologically relevant phase diagrams, as well as for the modelling of the compositional and physical properties of rocks and minerals at the high pressures and temperatures. In petrological systems, where reactions at depth in the Earth are of foremost interest, pressure, temperature and composition generally are the independent variables of choice, making the Gibbs free energy the most useful stability criterion. In thermodynamic data bases the Gibbs free energy is usually cast in the form

$$G_{P,T} = \Delta H_{o} - T \cdot S_{o} + \int_{T_{o}}^{T} [Cp_{o(T)}] dT$$
$$- T \cdot \int_{T_{o}}^{T} [Cp_{o(T)}/T] dT + \int_{P_{o}}^{P} [V_{(P,T)}] dP, \tag{1}$$

where P is pressure, T is temperature, $G_{P,T}$ is the molar Gibbs free energy at given P and T; $\Delta H_{\rm o}$ and $S_{\rm o}$ are the enthalpy of formation and third-law entropy, respectively, of a substance at standard pressure $P_{\rm o}$ and temperature $T_{\rm o}$; $Cp_{{\rm o}(T)}$ is the heat capacity as a function of temperature at standard pressure $P_{\rm o}$; $V_{(P,T)}$ is the molar volume of a substance as a function of pressure and temperature.

Equation (1) can lead to a direct representation of the Gibbs potential as a function of P and T when $Cp_{o(T)}$ and $V_{(P,T)}$ data are available for integration. Such data are still a major problem, however, and various empirical and semi-empirical power expansion series for $Cp_{o(T)}$ and $V_{(P,T)}$ have been suggested (see, for example, the reviews of Richet et al. 1992 and Saxena et al. 1993) that attempt to systematize the available, usually limited experimental data. Of particular importance is the fact that the form of the fitted polynomial should ideally have a physical significance (e.g. Dubrovinskaya et al. 1997), in order to allow realistic extrapolation beyond the range of the available data base. As discussed in detail by Richet et al. (1992), Saxena et al. (1993) and Dubrovinskaya et al. (1997), this limitation is particularly critical with respect to $Cp_{o(T)}$. While experimental calorimetric measurement leads to isobaric heat capacity $Cp_{o(T)}$, existing models for calculating calorimetric properties of minerals based on lattice dynamics lead instead to the isochoric heat capacity Cv. In addition, the vibration spectra of many of the minerals necessary for a useful petrological data base are still extremely complex and poorly understood. Thus, these models are limited in their effectiveness for providing a physical description of the temperature dependence of Cpo. All available polynomial fits for $Cp_{o(T)}$ of either experimental data or analogy-based estimates (cf. Richet et al. 1992), although widely used in petrological circles (Berman 1988; Holland and Powell 1990, 1998), are at present entirely empirical.

Theoretical approaches for modelling C_V turn to spectroscopic data and statistical thermodynamics. Molar volume and temperature are taken as independent variables and the Helmholtz free energy $F_{V,T}$ is chosen as the equilibrium potential for minerals (e.g. Kieffer 1985; Polyakov and Kuskov 1994; Dubrovinskaya et al. 1997) and fluids (e.g. Stewart and Jacobson 1989; Saul and Wagner 1989; Hill 1990). Theoretically derived equations for the Helmholtz free energy of solids (e.g. Kieffer 1985; Polyakov and Kuskov 1994; Dubrovinskaya et al. 1997) are primarily based on a statistical mechanical approach employing the canonical partition function in terms of molar volume and temperature that treats minerals as systems of quantum oscillators (e.g. Kubo 1965; Toda et al. 1992; Landau and Lifshitz 1959). Parameterization of these equations is based on either vibration spectra (e.g. Polyakov and Kuskov 1994) or data on isochoric heat capacity and volume (e.g. Dubrovinskaya et al. 1997) at different pressure and temperature. Analytical expressions for any other thermodynamic parameters (e.g. isobaric heat capacity, Gibbs free energy, entropy etc.) can then be derived using standard thermodynamic relations. Thus the Gibbs free energy of a substance at given V and T can be calculated as $G_{V,T} = F_{V,T} - V \cdot \partial F_{V,T} / \partial V$. However, this approach does not lead to a direct formulation of the Gibbs free energy as a function of P and T, as is preferable for a system of petrological interest, nor does it provide a basis for directly fitting experimental $Cp_{o(T)}$ data.

It has recently been shown that an analogous statistical approach (Kut'in and Pyadushkin 1998; Gerya et al. 1998) can be based on a partition function in pressure and temperature (e.g. Kubo 1965; Toda et al. 1992) to obtain expressions for the Gibbs free energy of solids as a function of P and T rather than V and T. Parameterization includes simultaneous regression of $Cp_{o(T)}$ and V(P,T) data, allowing fitting and optimization of various data sources, as required in the construction of an internally consistent petrological data base. This semi-empirical approach can also be extended to minerals with λ -transitions (e.g. Gerya et al. 1998) and to fluids (Gerya and Perchuk 1997), by considering the Gibbs free energy G as a function of pressure, temperature and an ordering parameter, so that accurate modelled representation and extrapolation of the thermodynamic properties of large numbers of petrologically significant minerals and coexisting fluids can be attained. The present paper is intended to summarize this new approach and to test it against well-established data on both the thermodynamic properties of some critical phases of petrological interest and on phase equilibria at high pressures and temperatures.

Summary of the basic method

Derivation of basic equations

For a substance in a crystalline state considered as a system of independent quantum oscillators (e.g. Landau and Lifshitz 1959), the Gibbs free energy can be expressed using the P-T partition function of statistical thermodynamics (Toda et al. 1992). Using a set of approximations for the calculation of the statistical sum over states, the following expression of the molar Gibbs free energy can be obtained (Appendix 1)

$$G_{\rm s} = H_{\rm s} - TS_{\rm s}^{\rm o} + RT \sum_{i}^{n} c_i \ln\{1 - \exp[\Delta H_{\rm si}/RT]\},$$
 (2)

where R is the gas constant, $H_{\rm s}$ and $S_{\rm s}^{\rm o}$ are, respectively, molar enthalpy and entropy for a state of zero-point vibrations, n is the number of groups of oscillators considered, c_i is a statistical weight of the i-th group equal to the number of oscillators in the i-th group divided by $N_{\rm A}$ (Avogadro's number), and $\Delta H_{\rm si}$ is the system enthalpy change related to transition between neighbouring energy levels in the i-th group (taken for $N_{\rm A}$ oscillators). The effects of pressure on $H_{\rm s}$ and $\Delta H_{\rm si}$ are given by

$$H_{\rm s} = H_{\rm s}^{\rm o} + \int\limits_{P_o}^{P} V_{\rm s} \mathrm{d}P, \tag{3}$$

$$\Delta H_{\rm si} = \Delta H_{\rm si}^{\rm o} + \int_{P_o}^{P} \Delta V_{\rm si} dP, \tag{4}$$

where P_o is standard pressure, H_s^o is molar enthalpy at 0 K and P_o , ΔH_{si}^o is the ΔH_{si} value at P_o , V_s is molar volume at 0 K as a function of pressure, ΔV_{si} is the system volume change due to transition between neighbouring energy levels for the *i*-th group of oscillators as a function of pressure (taken for N_A oscillators). Theoretical forms of the pressure dependence of V_s and ΔV_{si} are unknown, and thus some well-established semi-empirical functions must be employed. One of the possible choices is the Murnaghan equation, which demonstrates good extrapolation properties in a wide region of pressure at a given temperature (e.g. Zharkov and Kalinin 1971; Saxena et al. 1993):

$$V_{\rm s} = V_{\rm s}^{\rm o} (P_{\rm o} + \phi)^{1/5} / (P + \phi)^{1/5}, \tag{5}$$

$$\Delta V_{si} = \Delta V_{si}^{o} (P_o + \phi)^{1/5} / (P + \phi)^{1/5}, \tag{6}$$

where $V_{\rm s}^{\rm o}$ is molar volume at 0 K and $P_{\rm o}$, $\Delta V_{\rm si}^{\rm o}$ is the $\Delta V_{\rm si}$ value at $P_{\rm o}$, and ϕ is an empirical parameter approximating the mean effect of attractive interaction between the atoms in the whole pressure range. From Eq (3)–(4), the effect of pressure on $H_{\rm s}$ and $\Delta H_{\rm si}$ is given by

$$H_{\rm s} = H_{\rm s}^{\rm o} + V_{\rm s}^{\rm o} \Psi, \tag{7}$$

$$\Delta H_{\rm si} = \Delta H_{\rm si}^{\rm o} + \Delta V_{\rm si}^{\rm o} \Psi, \tag{8}$$

where
$$\Psi = 5/4(P_0 + \phi)^{1/5}[(P + \phi)^{4/5} - (P_0 + \phi)^{4/5}].$$

Standardizing Eq. (2) relative to T_o and P_o with a reduced number of empirical terms ($n \le 5$, Gerya et al. 1998), and taking into account Eq. (7)–(8), yields

$$G_{s} = H_{ToPo} - TS_{ToPo} + V_{s}^{o}\Psi + \sum_{i=1}^{n \le 5} c_{i} [RT \ln(1 - e_{i}) - \Delta H_{si}^{o}(1 - T/T_{o})e_{oi}/(1 - e_{oi}) - RT \ln(1 - e_{oi})],$$
(9)

$$S_{ToPo} = -(\partial G_s / \partial T)_{ToPo}$$

$$= S_s^o + \sum_{i}^{n} c_i [\Delta H_{si}^o / T_o e_{oi} / (1 - e_{oi}) - R \ln(1 - e_{oi})],$$
(9a)

$$H_{ToPo} = H_s^o + \sum_{i}^{n} c_i \Delta H_{si}^o \mathbf{e}_{oi} / (1 - \mathbf{e}_{oi}),$$
 (9b)

$$e_i = \exp(-\Delta H_{si}/RT) \tag{9c}$$

and

$$e_{oi} = \exp(-\Delta H_{si}^o/RT_o), \tag{9d}$$

where G_{ToPo} , H_{ToPo} , and S_{ToPo} are, respectively, the molar Gibbs free energy, enthalpy and entropy of the substance at standard T_o and P_o . If T_o is taken to be 0 K, the terms in Eq. (9) containing T_o are equal to zero. Equation (9) directly expresses the Gibbs potential of a condensed substance as a function of P and T.

An empirical parameterization of Eq. (9) can now be done using experimental data on isobaric heat capacities and volume measurements. Analytical expressions for Cp and V can be obtained from Eq. (9) as follows

$$C_p = -T(\partial^2 G_s / \partial T^2) = \sum_{i=1}^{n \le 5} c_i \Delta H_{si}^2 / R T^2 [e_i / (1 - e_i)^2],$$
(10)

$$V = \partial G_s / \partial P = \left[\sum_{i=1}^{n \le 5} c_i \Delta V_{si}^o \mathbf{e}_i / (1 - \mathbf{e}_i) + V_s^o \right] (P_o + \phi)^{1/5} / (P + \phi)^{1/5}.$$
(11)

Calculated examples

As an example of the applicability of Eqs. (9)–(11) for the description of the thermodynamic properties of crystalline substances, data for periclase, brucite, coesite and stishovite (Table 1) are considered. With n=3, $S_s^o=0$ (according to Nernst's law), $P_o=1$ bar and $T_o=298.15$ K, an equation of molar Gibbs free energy for these minerals takes the form (Gerya et al. 1998):

$$G_s = H_{298,1} + V_s^o \Psi + \sum_{i=1}^3 c_i [RT \ln(1 - e_i) - \Delta H_{si}^o e_{oi} / (1 - e_{oi})],$$
(12)

where
$$e_i = \exp[-(\Delta H_{si}^o + \Delta V_{si}^o \Psi)RT]$$
, $e_{oi} = \exp(-\Delta H_{si}^o/R298.15)$, $\Psi = 5/4(P_o + \phi)^{1/5}[(P + \phi)^{4/5} - (1 + \phi)^{4/5}]$.

Entropy S_{ToPo} is absent from Eq. (12) since $S_s^o = 0$ and all other terms in Eq. (9a) are cancelled by corresponding terms in Eq. (9). Third-law entropy can be calculated according to Eq. (9a) assuming that low-temperature heat capacity (20–298 K) is accurately represented by Eq. (10).

The results for the parameters of Eq. (12) are listed in Table 2. The derivation of these internally consistent thermodynamic data was performed by a non-linear least-squares method. Processing of the experimental data was done in cyclic fashion with quality control of the refined parameters. The measured parameters are reproducible within experimental error. To illustrate this, the calculated results for isobaric heat capacity and volume are plotted against the original experimental data in Figs. 1–4. It is seen that by comparison to the power expansion formulations used in petrological data bases (e.g. Berman 1988; Holland and Powell 1998), Eq. (12) allows a better fit of isobaric heat capacities and volumes over a very wide region of pressure (0–800 kbar) and temperature (20–3000 K). Equation (2) requires that c_i in Eq. (12) should sum to 3v, where v is the number of atoms in the formula unit. This is fulfilled for coesite and stishovite (the sum is 9), but not quite for brucite (the sum is 14.6 instead of 15) and periclase (the sum is 6.6 instead of 6). These discrepancies are due to the relatively low value of n = 3chosen in Eq. (12), which may not be sufficient to

Table 1 Source of experimental data used in this study

| Starting data | References | | | |
|--|--|--|--|--|
| Isobaric heat capacity of quartz | Gurevitch and Khlyustov (1979); Watanabe (1982); Hemingway (1987); Grønvold et al. (1989); Hemingway et al. (1991); Akaogi et al. (1995) | | | |
| Molar volume of quartz | Jay (1933); Ackermann and Sorell (1974); Danielsson et al. (1976); Lager et al. (1982) Olinger and Halleck (1976); Jorgensen (1978); d'Amour et al. (1979); Levien et al. (1980); Hazen et al. (1989); Kihara (1990); Vaidya et al. (1973); Glinnemann et al. (1992) | | | |
| Adiabatic modulus of quartz, $(\partial T/\partial P)_S$ | Boehler et al. (1979); Boehler (1982) | | | |
| Relative molar enthalpy of quartz | Richet et al. (1982) | | | |
| Isobaric heat capacity of coesite | Holm et al. (1967); Akaogi et al. (1995) | | | |
| Molar volume of coesite | Bassett and Barnett (1970); Levien and Prewitt (1981) Galkin et al. (1987); Smyth et al. (1987) | | | |
| Relative molar enthalpy of coesite Isobaric heat capacity of stishovite | Holm et al. (1967); Akaogi and Navrotsky 1984; Akaogi et al. (1995) Holm et al. (1967); Akaogi et al. (1995) | | | |
| Molar volume of stishovite | Bassett and Barnett (1970); Liu et al. (1974); Olinger (1976); Sato (1977); Endo et al. (1986); Doroshev et al. (1987); Tsuchida and Yagi (1989); Ross et al. (1990) Suito et al. (1996) | | | |
| Enthalpy of phase transition quartz-coesite | Holm et al. (1967); Akaogi and Navrotsky (1984); Kuskov et al. (1991); Akaogi et al. (1995) | | | |
| Phase transition α -quartz - β -quartz | Gibson (1928); Yoder (1950); Cohen and Klement (1967); Koster van Groos and Ten Heege (1973); Mirwald and Massone (1980b); Shen et al. (1993) | | | |
| Phase transition quartz—coesite | Boyd and England (1960); Kitahara and Kennedy (1964); Mirwald and Massone (1980a,b); Akella (1979); Bohlen and Boettcher (1982); Ishbulatov and Kosyakov (1990); Bose and Ganguly (1995) | | | |
| Phase transition coesite-stishovite | Akimoto and Syono (1969); Yagi and Akimoto (1976); Suito (1977); Zhang et al. (1993), (1996); Serghiou et al. (1995) | | | |
| Water: tabulated data on molar Gibbs free energy, molar entropy and molar volume of water at $T=0$ –1000 °C and $P=1$ –10 000 bar; tabulated P – V – T data on liquid-gas transition | Saul and Wagner (1989) | | | |
| Isobaric heat capacity of periclase | Giauque and Archibald (1937); Barron et al. (1959); Glushko et al. (1979); Krupka et al. (1979); Watanabe (1982) | | | |
| Molar volume of periclase | Reeber et al. (1995); Fiquet et al. (1999); Fei (1999) Dewaele et al., (2000); Zhang (2000) | | | |
| Adiabatic modulus of periclase $(\partial T/\partial P)_S$ Relative molar enthalpy of periclase Isobaric heat capacity of brucite Molar volume of brucite | Boehler (1982) Richet and Fiquet (1991) Giauque and Archibald (1937); Glushko et al. (1979) Chakoumakos et al. (1997); Catti et al. (1995); Parise et al. (1994); Fei and Mao (1993); Nagai et al. (2000); Xia et al. (1998) | | | |
| Phase transition brucite = periclase + water | Franz (1982); Schramke et al. (1982); Johnson and Walker (1993); Aranovich and Newton (1996) | | | |

reflect the actual number of different groups of oscillators.

Modification to include minerals with λ -transitions

Derivation of basic equations

Some petrologically important phases such as quartz demonstrate specific changes in thermodynamic properties that are related to a λ -transition not accounted for in Eq. (9). Most approaches attempt to describe this effect in terms of the formalism of Landau theory (e.g. Carpenter et al. 1998; Dove 1997; Holland and Powell 1998). Landau theory (e.g. Landau and Lifshitz 1959) suggests that the Gibbs potential of a crystalline phase in the region of the λ -transition can be considered as a function of P, T and an order parameter η . The equilibrium order parameter corresponds to a minimum of the Gibbs free energy $G(T,P,\eta) = \min$. This general

formalism is consistent with thermodynamic concepts of homogeneous equilibria in crystals (Thompson 1969) widely developed for phases of petrological interest (e.g. Holland and Powell 1996). A similar approach has recently been applied to the λ -transition in quartz (Gerya et al. 1998).

A thermodynamic process related to a λ -transition in a one-component mineral can be viewed as an internal ordering reaction (transition)

$$aA = \alpha,$$
 (13)

where A is a disordered cluster (structural unit), $\alpha = A_a$ is an ordered cluster (structural unit), a is the stoichiometric coefficient. The thermodynamic equilibrium for Reaction (transition) 13 is given by

$$\Delta G_{\text{ord}} = RT \ln(K_{\text{ord}}) + \Delta G_{\text{ord}}^{\text{o}} + \Delta G_{\text{ord}}^{\text{e}} = 0, \tag{14}$$

where $K_{\rm ord} = X_{\alpha}/(X_{\rm A}^a)$ and $X_{\rm A} + X_{\alpha} = 1$. $X_{\rm A}$ and X_{α} are the mole fractions (thermodynamic probabilities) of the disordered and ordered clusters (structural units),

Table 2 Calculated parameters of Eqs. (12), (20) and (30) for the molar Gibbs free energy of minerals and water

| Parameter | Coesite | Stishovite | Brucite | Periclase | Quartz | Water |
|---|------------------------------|------------------------------|------------------------------|------------------------------|------------------------------|-------------------------|
| H _{298,1} , J | -907051.35 | -870123.92 | -925267.08 | -601500.00 ^a | -910712.59 ^a | -286831.56 ^a |
| S_{2981} , J K ⁻¹ | _ | _ | _ | _ | _ | 65.188 |
| $V_{\rm s}^{\rm o}$, J bar ⁻¹ | 2.05333 | 1.40010 | 2.43162 | 1.12228 | 2.25888 | 1.71382 |
| ϕ , bar | 217664 | 612340 | 98040 | 301795 | 71662 | 6209 |
| c_1 | 1.44913 | 0.28816 | 1.87663 | 1.96612 | 1.42569 | 7.23576 |
| c_2 | 3.71385 | 2.73177 | 8.55012 | 4.12756 | 3.50283 | 0.31482 |
| c_3 | 3.83702 | 5.98007 | 4.16639 | 0.53690 | 4.07148 | _ |
| $\Delta H_{\rm s1}^{\rm o}$, J | 1435.64 | 1179.01 | 1637.57 | 2966.88 | 1278.08 | 4586.46 |
| $\Delta H_{\rm s2}^{\rm o}$, J | 4099.77 | 3845.99 | 4403.89 | 5621.69 | 3911.05 | _ |
| $\Delta H_{\rm s3}^{\rm s2o}$, J | 10479.16 | 8042.74 | 11614.44 | 27787.19 | 10469.21 | _ |
| $\Delta V_{\rm s1}^{\rm o}$, J bar ⁻¹ | 0.00158600 | 0.00233400 | 0.00541752 | 0.00352971 | 0.00312581 | 0.04310884 |
| $\Delta V_{\rm s2}^{\rm o}$, J bar ⁻¹ | $=\Delta V_{\rm s1}^{\rm o}$ | _ |
| $\Delta V_{\rm s3}^{\rm o}$, J bar ⁻¹ | $=\Delta V_{s1}^{o}$ | $=\Delta V_{s1}^{\circ}$ | 0.07026587 | 0.19849568 | $=\Delta V_{\rm s1}^{\rm o}$ | _ |
| $\Delta H^{\rm o}_{\rm ord}$, J | _ | - " | _ | _ | -2636.90 | -44838.80 |
| $\Lambda S^{o} \longrightarrow I K^{-1}$ | _ | _ | _ | _ | _ | -122.397 |
| $\Delta Cp^{\circ}_{\text{ord}}$, J K ⁻¹ | _ | _ | _ | _ | _ | 21.486 |
| $\Delta V_{\rm ord}^{\rm o}$, J bar ⁻¹ | _ | _ | _ | _ | -0.127829 | |
| W_{1}^{H}, J^{-1} | _ | _ | _ | _ | 14255.08 | -28793.19 |
| $\Delta Cp^{\circ}_{\text{ord}}$, J K ⁻¹ $\Delta V^{\circ}_{\text{ord}}$, J bar ⁻¹ W^{H}_{1} , J ⁻¹ W^{S}_{1} , J K ⁻¹ | _ | _ | _ | _ | _ | -11.704 |
| $W^{\text{Cp}}_{1}, \text{ J K}^{-1}$ | _ | _ | _ | _ | _ | 5.086 |
| W_1^{V} , J bar ⁻¹ | _ | _ | _ | _ | 0.633504 | _ |
| $\Delta H_{\rm s\lambda}^{\rm ro}$, J | _ | _ | _ | _ | 12947.71 | _ |
| $\Delta V_{\rm s\lambda}^{\rm so}$, J bar ⁻¹ | _ | _ | _ | _ | 0.242077 | _ |

^aThe $H_{298,1}$ value has been adjusted to obtain the enthalpies of formation at 298.15 K and 1 bar, as recommended by CODATA (1978)

respectively. $\Delta G^{\rm o}_{\rm ord}$ and $\Delta G^{\rm e}_{\rm ord}$ are changes of the standard and excess Gibbs free energy, respectively, due to Reaction (13). According to our formalism, ordered/disordered clusters need not always be static (long-lived) structural features, but may reflect dynamic (short-lived) variations due to the thermal vibration of atoms (e.g. Kihara 1990). This does not preclude a uniform thermodynamic treatment, but instead changes the interpretation of the ordering parameter X_{α} . For static features this parameter represents the mole fraction of ordered clusters, while for dynamic variations it stands for the thermodynamic probability of finding the given structural units in the ordered state. The additional molar Gibbs free energy relative to the ordered state $(X_{\alpha} = 1)$ is defined as (Gerya et al. 1998):

$$G_{\alpha} = \{RT[(1 - X_{\alpha}) \ln(1 - X_{\alpha}) + X_{\alpha} \ln(X_{\alpha})] - \Delta G_{\text{ord}}^{o}(1 - X_{\alpha})/a + \Delta G_{\text{ord}}^{e}\}/[1 + X_{\alpha}(a - 1)],$$

where
$$\Delta G^{o}_{ord} = \Delta H^{o}_{ord} + \Delta V^{o}_{ord} \Psi$$
, $G^{e}_{ord} = W^{G}(1 - X_{\alpha})$, $W^{G} = W^{H} + W^{V}\Psi$.

 $G^{\rm e}_{\rm ord}$ is the integral excess Gibbs free energy of mixing of ordered and disordered clusters expressed using the Margules formalism (e.g. Thompson 1969), with the expressions for $\Delta G^{\rm o}_{\rm ord}$ and $W^{\rm G}$ formulated by analogy to Eq. (7).

If the λ -transition leads to significant changes in the thermal vibrations of atoms, this must be taken into account in the vibrational part of the Gibbs potential. The corresponding formulation derived from the P-T partition function by analogy to Eq. (9) takes the form (see Appendix 2)

$$G_{s} = H_{ToPo} - TS_{ToPo} + V_{s}^{o}$$

$$+ \sum_{i=1}^{n \leq 5} c_{i} \{RT \ln\{(1 - e_{\alpha i})(1 - e_{\beta i}) / [(1 - e_{\beta i}) + e_{\lambda i}(e_{\beta i} - e_{\alpha i})]\} - \Delta H_{si\alpha}^{o}(1 - T/T_{o}) e_{o\alpha i} / (1 - e_{o\alpha i})$$

$$- RT \ln(1 - e_{o\alpha i})\}, \qquad (16)$$
where $S_{ToPo} = -(\partial G_{s} / \partial T)_{ToPo}$

$$= S_{s}^{o} + \sum_{i}^{n} c_{i} [\Delta H_{si}^{o} / T_{o} e_{o\alpha i} / (1 - e_{o\alpha i}) - R \ln(1 - e_{o\alpha i})],$$

$$H_{ToPo} = H_{s}^{o} + \sum_{i}^{n} c_{i} \Delta H_{si}^{o} e_{o\alpha i} / (1 - e_{o\alpha i}),$$

$$e_{\alpha i} = \exp(-\Delta H_{si\alpha} / RT), e_{\beta i} = \exp(-\Delta H_{si\beta} / RT),$$

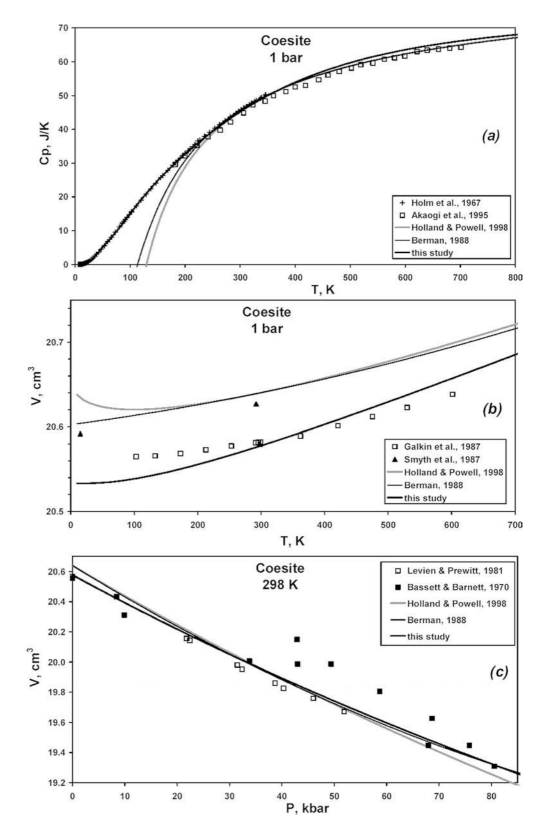
$$e_{\lambda i} = \exp(-\Delta H_{si\alpha} / RT), e_{o\alpha i} = \exp(-\Delta H_{si\beta} / RT_{o}),$$

$$\Delta H_{si\alpha} = \Delta H_{si\alpha}^{o} + \Delta V_{si\alpha}^{o} \Psi, \Delta H_{si\beta} = \Delta H_{si\beta}^{o} + \Delta V_{si\beta}^{o} \Psi,$$
and
$$\Delta H_{si\lambda} = \sum_{k=0}^{2} (\Delta H_{si\lambda k}^{o} + \Delta V_{si\lambda k}^{o} \Psi) (X_{\alpha})^{k}.$$

 $\Delta H_{si\alpha}$ and $\Delta H_{si\beta}$ are values of ΔH_{si} for ordered and disordered substances, respectively; $\Delta H_{si\lambda}$ is the apparent enthalpy parameter dependent on X_{α} and related to the order–disorder transition for the *i*-th group of oscillators.

Under isothermal-isobaric conditions, an equilibrium in a non-reacting phase is given by

Fig. 1a-c Comparison of calculated isobaric heat capacity (a) and molar volume (b,c) of coesite with measured data



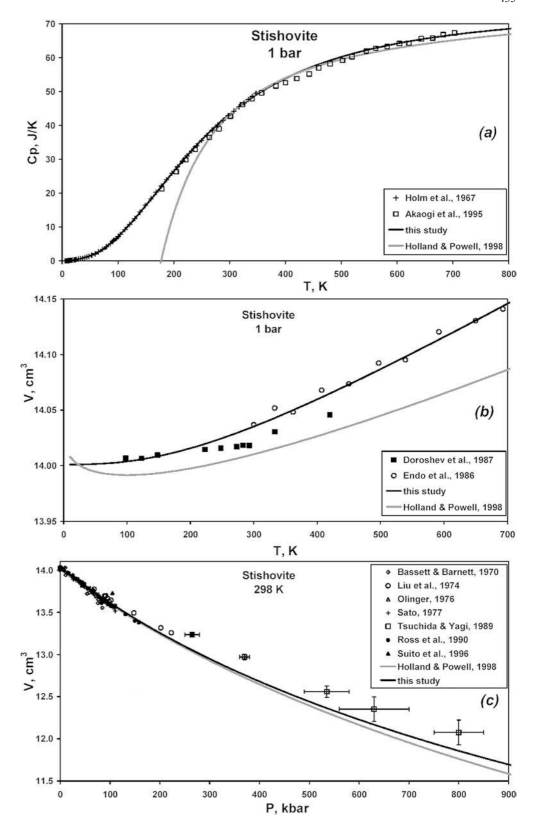
$$G_{s+\alpha} = G_s + c_\alpha G_\alpha = \min, \tag{17}$$

$$\partial G_{s+\alpha}/\partial X_{\alpha} = 0, \tag{18}$$

where $G_{s+\alpha}$ is the molar Gibbs free energy of a substance with a λ -transition, G_s is given either by Eq. (7)

when changes in thermal vibrations are insignificant, or by Eq. (16), when these changes must be taken into account, and c_{α} is the number of degrees of freedom corresponding to the ordering Reaction (13). X_{α} can be considered to be equivalent to the ordering parameter η

Fig. 2a-c Comparison of calculated isobaric heat capacity (a) and molar volume (b,c) of stishovite with measured data

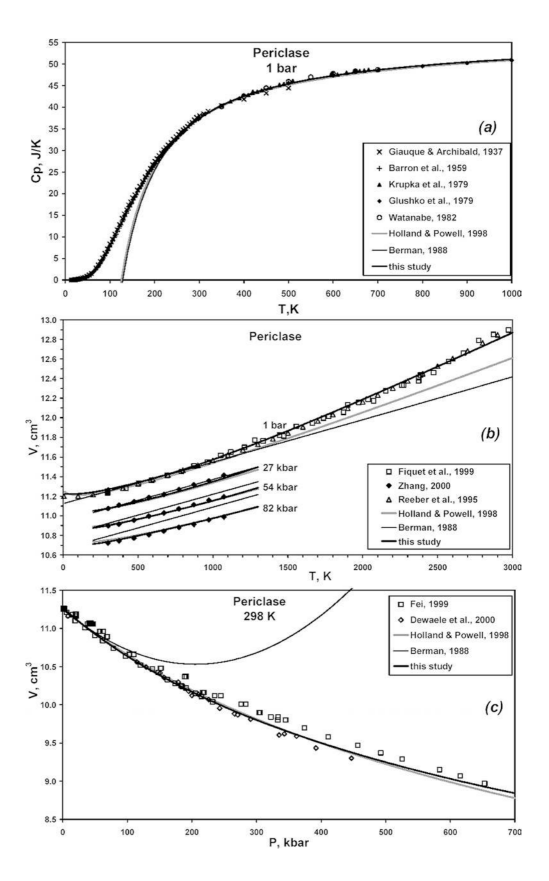


above, which tends to a maximum $(X_{\alpha}^{\max} \leq 1)$ in the ordered phase with decreasing temperature, while, with increasing temperature, it tends to a minimum $(X_{\alpha}^{\min} \geq 0)$ in the disordered phase.

Application to quartz

As an example of the applicability of Eqs. (13)–(18), to the description of the thermodynamic properties of

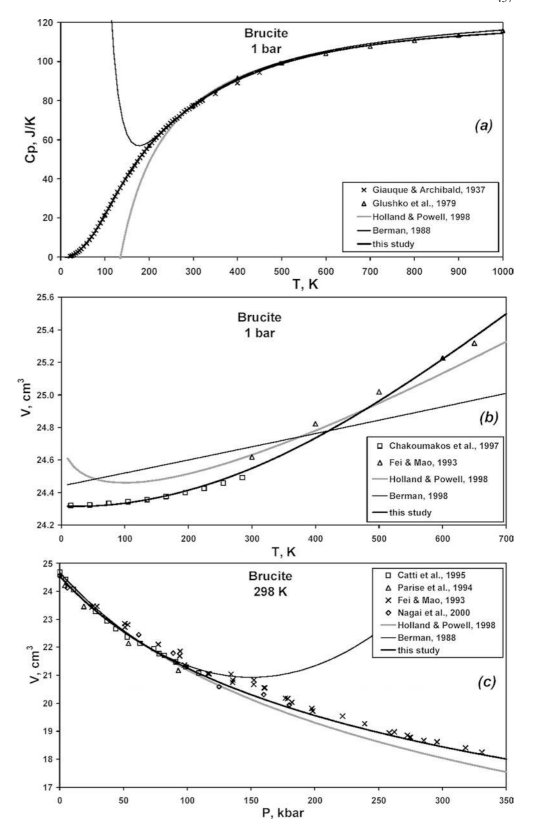
Fig. 3a-c Comparison of calculated isobaric heat capacity (a) and molar volume (b,c) of periclase with measured data



minerals with a λ -transition, the results of their parameterization can be considered for quartz.

studied over a wide range of P-T conditions and manifest themselves in anomalous behaviour of heat Ordering-disordering phenomena in quartz are well capacity and thermal expansion, thus providing tight

Fig. 4a—c Comparison of calculated isobaric heat capacity (a) and molar volume (b,c) of brucite with measured data



constraints on the proposed model (e.g. Dorogokupets 1995). The data for quartz were treated using Eqs. (15)–(18), with ordering represented by the following homogeneous ordering reaction

$$3SiO_2(Qtz_{\beta}) = Si_3O_6(Qtz_{\alpha}), \tag{19}$$

where Qtz_{β} and Qtz_{α} are, respectively, disordered and ordered quartz clusters (e.g. Castex and Madon 1995;

Spearing et al. 1992). Using $c_{\alpha} = 1$ in Eq. (17) and n = 3, $V_{si\alpha}^{\circ} = V_{si}^{\circ}$, $V_{si\beta}^{\circ} = 0$, $S_{s}^{\circ} = 0$ (according to Nernst's law), $P_{o} = 1$ bar and $T_{o} = 298.15$ K in Eq. (16), the Gibbs free energy equation for quartz takes the following semi-empirical form (Gerya et al. 1998)

$$G_{s+\alpha} = H_{298,1} + V_s^{\text{o}} \Psi$$

$$+ \sum_{i=1}^{3} c_i \left\{ RT \ln\{(1 - e_{\alpha i})(1 - e_{\beta i}) / [(1 - e_{\beta i}) + e_{\lambda}(e_{\beta i} - e_{\alpha i})] \right\} - \Delta H_{si}^{\text{o}} e_{\text{o}i} / (1 - e_{\text{o}i}) \right\}$$

$$+ \left\{ RT[X_{\alpha} \ln(X_{\alpha}) + (1 - X_{\alpha}) \ln(1 - X_{\alpha})] - (\Delta H_{\text{ord}}^{\text{o}} + \Delta V_{\text{ord}}^{\text{o}} \Psi)(1 - X_{\alpha}) / 3 + (W_1^{\text{H}} + W_1^{\text{V}} \Psi)(1 - X_{\alpha}) X_{\alpha} \right\} / (1 + 2X_{\alpha}), \tag{20}$$

where $e_{\alpha i}=\exp[-(\Delta H_{si}^{\ o}+\Delta V_{s}^{\ o}\Psi)/RT],\ e_{\beta i}=\exp(-\Delta H_{si}^{\ o}/RT),\ e_{\alpha i}=\exp(-\Delta H_{si}^{\ o}/R298.15),\ e_{\lambda}=\exp[-(\Delta H_{s\lambda}^{\ o}X_{\alpha}^{\ 2}+\Delta V_{s\lambda}^{\ o}\Psi)/RT],\ \Psi=5/4(P_{o}+\phi)^{1/5}[(P+\phi)^{4/5}-(1+\phi)^{4/5}].\ X_{\alpha}$ is the mole fraction of ordered Si₃O₆ clusters. The empirical parameters obtained for Eq. 20 are presented in Table 2.

Equation (20) allows an accurate description of both experimental heat capacity and volume of α - and β -quartz over a wide region of P and T that includes the λ -transition (Fig. 5). By comparison with the formulations of Berman (1988), Dorogokupets (1995) and Holland and Powell (1998), Eq. 20 demonstrates a superior fit of the available experimental data (Fig. 5).

Gibbs potential of fluids

Derivation of basic equations

Fundamental, precise descriptions of the thermodynamic properties of fluid over a wide P-T interval, including the two-phase region, are traditionally based on the Helmholtz potential and related P = f(V,T)equations of state (e.g. Stewart and Jacobson 1989; Saul and Wagner 1989; Hill 1990). However, it was recently shown (Gerya and Perchuk 1997) that a similar description can be based on the Gibbs potential. The phenomenological approach suggested by Gerya and Perchuk (1997) uses the concept of an internal reaction of association of molecules to form dynamic (shortlived) clusters in fluids. Such an approach is well developed theoretically for describing the hydrogen bonding in supercritical fluids (e.g. Luck 1980; Gupta et al. 1992) and its potential applicability to any fluid has been demonstrated (Barelko et al. 1994; Gerya and Perchuk 1997).

According to Barelko et al. (1994), the dynamic process of association of molecules in any supercritical fluid can be considered as a chain clusterization reaction

$$A_1 + A_n = A_{n+1}, (21)$$

where A_1 stands for a monomer, and A_n and A_{n+1} denote clusters of dimensions n and n+1. The thermodynamic equilibrium of Reaction (21) is given by

$$\Delta G_{\text{cls}} = RT \ln[K_{\text{cls}(n)}] + \Delta G_{\text{cls}(n)}^{\text{o}} + \Delta G_{\text{cls}(n)}^{\text{e}} = 0, \qquad (22)$$

where $K_{\operatorname{cls}(n)} = X_{n+1}/(X_1 \, X_n)$ and $\sum_{n=1}^{\infty} X_n = 1$; X_1 , X_n , and X_{n+1} are the mole fractions (thermodynamic probabilities) of clusters of dimension 1 (i.e. monomers), n and n+1, respectively; $\Delta G^{\operatorname{o}}_{\operatorname{cls}(n)}$ and $\Delta G^{\operatorname{e}}_{\operatorname{cls}(n)}$ are changes of the standard and excess Gibbs free energy, respectively, due to Reaction (21). The additional molar Gibbs free energy relative to the associated $(X_1=0,X_\infty=1)$ state is given by

$$G_{\alpha} = \sum_{n=1}^{\infty} X_{n} [RT \ln(X_{n}) + n(G_{1}^{o} + G_{1}^{e})$$

$$+ \sum_{m=1}^{n-1} (\Delta G_{\operatorname{cls}(m)}^{o} + \Delta G_{\operatorname{cls}(m)}^{e})] / \sum_{n=1}^{\infty} nX_{n} - [G_{1}^{o} + \sum_{n=1}^{\infty-1} \Delta G_{\operatorname{cls}(m)}^{o} / \infty],$$
(23)

where G_1° and G_1^{e} are standard and excess Gibbs free energy of monomers, respectively. When the equilibrium constant of Reaction (22) is assumed to be independent of n, summation in Eq. (23), using Eq. (22), yields

$$G_{\alpha} = \{RT[(1 - X_1) \ln(1 - X_1) + X_1 \ln(X_1)] - \Delta G_{cls}^{o} X_1 + G^{e}\},$$
(24)

where $\Delta G^{\rm o}_{\rm cls}$ is the change of standard Gibbs free energy in Reaction (21) (independent of n), $G^{\rm e}$ is the excess molar energy of mixing, related to the presence of different clusters in the fluid. A similar equation was obtained by Gerya and Perchuk (1997) by treating the dynamic association process as a simple, short-term "ordering" transition

$$Gas = Liq, (25)$$

where Gas stands for a molecule in the "gas-like" (free, "disordered") mode, and Liq denotes a molecule in the "liquid-like" (associated, "ordered") mode. According to Reaction (25) the additional molar Gibbs free energy relative to the associated ($X_{\rm Gas}=0,~X_{\rm Liq}=1$) state is given by

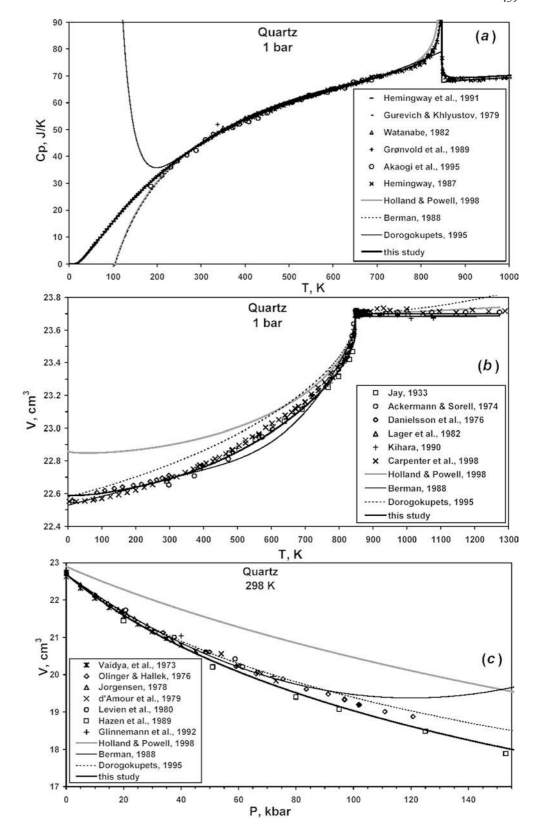
$$G_{\alpha} = RT[X_{\text{Liq}} \ln(X_{\text{Liq}}) + X_{\text{Gas}} \ln(X_{\text{Gas}})] - \Delta G_{\text{ord}}^{\text{o}} X_{\text{Gas}} + G^{\text{e}},$$
(26)

where $\Delta G^{\circ}_{\text{ord}}$ is a change of standard Gibbs free energy in "ordering" Reaction (25). Thus, from the viewpoint of thermodynamics, both considerations are equivalent and yield similar results.

A semi-empirical formulation of the last two terms in Eq. (26) yields (see Appendix 3)

$$G_{\alpha} = RT[X_{\text{Liq}} \ln(X_{\text{Liq}}) + (1 - X_{\text{Liq}}) \ln(1 - X_{\text{Liq}})] + (1 - X_{\text{Liq}})RT \ln[(P + \phi X_{\text{Liq}}^{2})/P_{\text{o}}] - (1 - X_{\text{Liq}})\{\Delta H_{\text{ord}}^{\text{o}} - T\Delta S_{\text{ord}}^{\text{o}} + \Delta C_{P \text{ ord}}^{\text{o}}[T - T_{\text{o}} - T \ln(T/T_{\text{o}})]\} + \{W_{1}^{\text{H}} - TW_{1}^{\text{s}} + W_{1}^{\text{Cp}}[T - T_{\text{o}} - T \ln(T/T_{\text{o}})]\}X_{\text{Liq}}(1 - X_{\text{Liq}}),$$
(27)

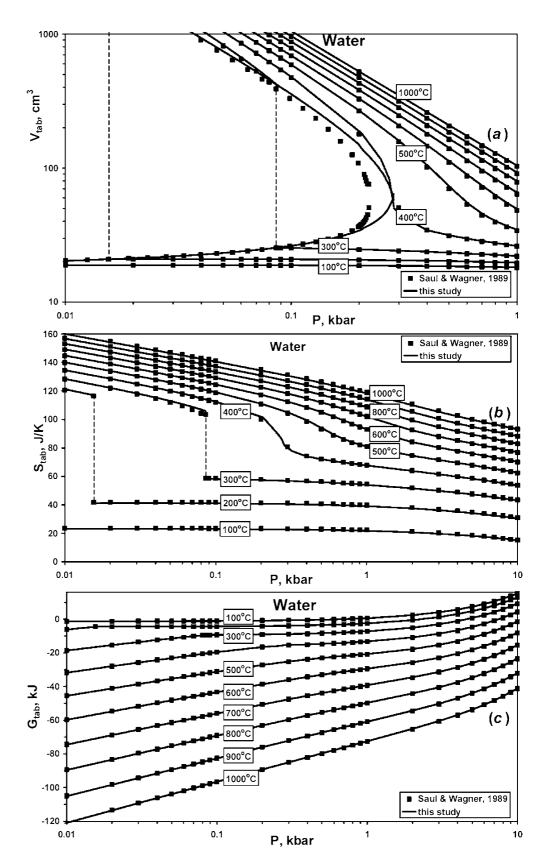
Fig. 5a-c Comparison of calculated isobaric heat capacity (a) and molar volume (b,c) of quartz with measured data



where $\Delta H^{\rm o}_{\rm ord}$, $\Delta S^{\rm o}_{\rm ord}$, $\Delta C_{P}^{\rm o}_{\rm ord}$ are standard enthalpy, entropy and heat capacity effects of Reaction (25), $W_1^{\rm H}$, $W_1^{\rm S}$, $W_1^{\rm Cp}$ are enthalpy, entropy and heat capacity Margules parameters.

By analogy with minerals, a standard Gibbs free energy, G_s , of fluid in fully associated "liquid-like" state ($X_{\rm Gas}=0$, $X_{\rm Liq}=1$), considered as a system of quantum oscillators, can be approached by Eq. (9). Under

Fig. 6a–c Comparison of calculated volume (a), entropy (b) and Gibbs energy (c) of water with tabulated data (Saul and Wagner 1989)



isothermal-isobaric conditions, equilibrium in a onephase fluid is given by

$$G_{s+\alpha} = G_s + G_\alpha = \min$$

$$\partial G_{\alpha}/\partial X_{\text{Liq}} = 0.$$
 (29)

where $G_{s+\alpha}$ is the molar Gibbs free energy of a fluid, and (28) G_s is given by Eq. 9.

Thermodynamic properties of aqueous fluid

As an example of the applicability of Eqs. (26)–(29) for describing the thermodynamic properties of compressed gases, the results of parameterizing an aqueous fluid can be considered. The tabulated data for water (Saul and Wagner 1989) in the experimentally well-studied region T=0–1000 °C and P=1–10 000 were treated using Eqs. (9) as well as (27)–(29) on the basis of a non-linear least-squares method.

Using n = 2 in Eq. (9), $P_o = 1$ bar and $T_o = 298.15$ K in Eqs. (9) and (27), the Gibbs free energy equation for aqueous fluid takes the following semi-empirical form, which is also applicable to other gases (e.g. Ar, CO_2 , CH_4 ; Gerya and Perchuk 1997)

$$\begin{split} G_{\text{s}+\alpha} &= H_{298,1} - TS_{298,1} + V_{\text{s}}^{\text{o}} \Psi + RT[c_1 \ln(1-e_1) \\ &+ c_2 \ln(1-e_2)] - (c_1 + c_2) [\Delta H_{\text{s}1}^o (1-T/T_{\text{o}}) e_{\text{o}}/(1-e_{\text{o}}) \\ &+ RT \ln(1-e_{\text{o}})] + RT[(1-X_{\text{Liq}}) \ln(1-X_{\text{Liq}}) \\ &+ X_{\text{Liq}} \ln X_{\text{Liq}}] + (1-X_{\text{Liq}}) RT \ln[\phi X_{\text{Liq}}^2 + P] \\ &- (1-X_{\text{Liq}}) \{\Delta H_{\text{ord}}^o - T\Delta S_{\text{ord}}^o + \Delta C_{P \text{ord}}^o [T-298.15 \\ &- T \ln(T/298.15)]\} + \{W_1^H - TW_1^S \\ &+ W_1^{CP} [T-298.15 - T \ln(T/298.15)]\} X_{\text{Liq}} (1-X_{\text{Liq}}), \end{split}$$

where $e_1 = \exp[-(\Delta H_{\rm s1}^{\rm o} + \Delta V_{\rm s1}^{\rm o} \Psi)/RT]]$, $e_2 = \exp[-\Delta H_{\rm s1}^{\rm o}/RT]]$, $e_0 = \exp[-\Delta H_{\rm s1}^{\rm o}/R298.15]]$, $\Psi = 5/4(P_{\rm o} + \phi)^{1/5}$ $[(P + \phi)^{4/5} - (1 + \phi)^{4/5}]$. Table 2 lists the calculated parameters of Eq. (30). This equation allows an accurate, continuous description of the thermodynamic properties of water in both the two-phase and supercritical P-V-T region (Fig. 6), i.e. without the need to

Fig. 7 Comparison of calculated molar volume of water at high pressure and temperature with thermodynamic data (Wagner and Pruss 1997) recommended for scientific use (IAPWS-95)

separate the P-V-T field into sub-regions (e.g. Holland and Powell 1998). The most significant errors were detected for the molar volume of the aqueous gas in the vicinity of the critical point (Fig. 6a). Problems in describing the molar volumes of fluids near the critical region have long been known, and empirical switch functions are commonly used to improve the description in this region without disturbing the continuity of the far-field P-V-T equations (e.g. Hill 1990). Taking into account that the related errors in the Gibbs free energy are not large (Fig. 6c), we decided not to introduce this additional complexity to our model. Figure 7 shows a comparison of the molar volumes of water, calculated according to Eq. (30), with those recommended by IAPWS-95 for scientific use (Wagner and Pruß 1997). The comparison shows that our equation demonstrates more consistency with internationally approved tabulated properties of water than other calibrations (Duan et al. 1996; Holland and Powell 1998), especially at high P and T.

Figure 8 shows the results of extrapolating the molar volumes of water to high *P-T*. It is seen that Eq. (30) shows good extrapolation properties in terms of both pressure and temperature, and better coincides with the results of independent shock-wave and volumetric experiments than calibrations suggested by Duan et al. (1996) and Holland and Powell (1998).

Combined applications: phase equilibria in the system $MgO-SiO_2-H_2O$

Phase equilibria in the MgO–SiO₂– H_2O system have been experimentally studied to very high P and T, thus providing the opportunity to test the equations of

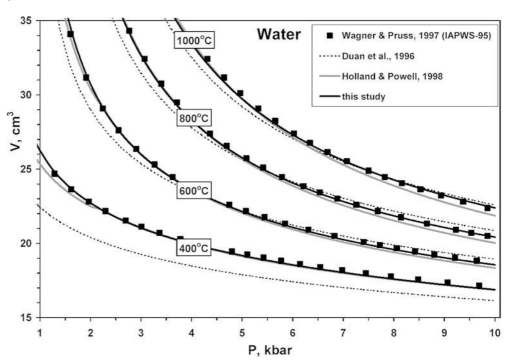
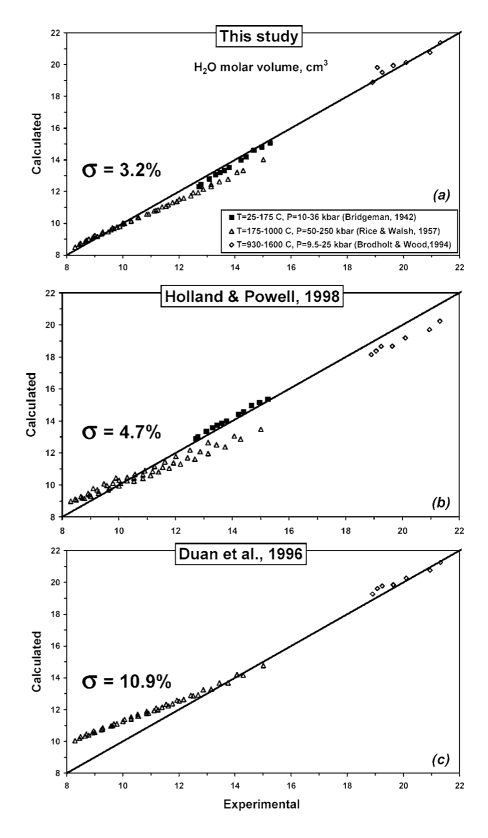


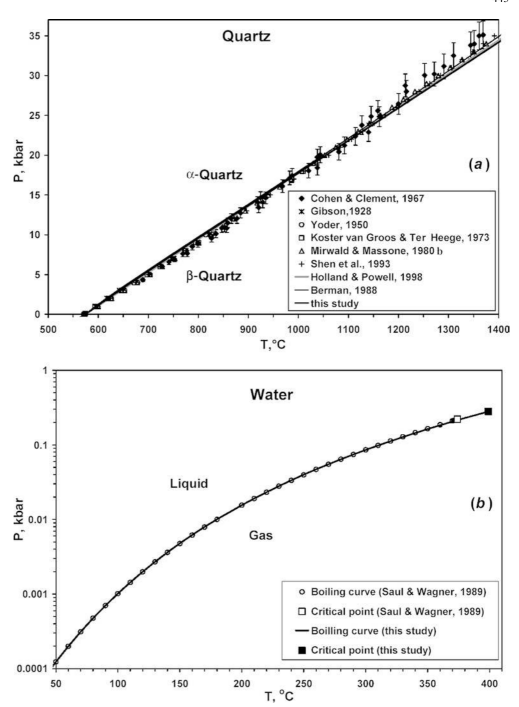
Fig. 8a–c Comparison of the extrapolated molar volumes of water with results of the independent volumetric (Bridgeman 1942; Brodholt and Wood 1994) and shockwave (Rice and Walsh 1957) experiments. Different diagrams show the volumes calculated using Eq. (30) (a), and PVT formulations suggested by Holland and Powell (1998) (b) and Duan et al. (1996) (c)



Gibbs free energy described here over a wide range of pressures and temperatures. The extrapolation properties of the above equations can be simultaneously tested for both minerals and the coexisting aqueous fluid. All calculations are inherently based on internally consistent data on the basis of non-linear least-squares analysis.

We considered several experimentally studied equilibria involving aqueous fluid, quartz, coesite, stishovite, periclase, and brucite. Figure 9 show the results

Fig. 9a, b Comparison of phase boundaries calculated for quartz (a) and water (b) with experimental data



obtained for the α – β transition in quartz (Fig. 9a) and the liquid–gas transition in water (Fig. 9b), calculated with the use of Eqs. (20) and (30), respectively. Along the phase boundaries, mole fractions of the ordered/associated clusters (X_{α} / $X_{\rm Liq}$) in the coexisting phases correspond to the conditions of equilibrium for both the internal "ordering" transition in each phase and the heterogeneous reaction between them. This is exemplified by Fig. 10, showing the Gibbs potential of quartz (Fig. 10a) and water (Fig. 10b) in the region of the phase transitions. Along these curves, two

analogous minima occur in both cases, corresponding to the two stable phases differing in the degree of ordering/association. Figure 9 shows that the derived Gibbs energy equations allow an accurate description of experimental data. However, the critical point of water calculated using Eq. (30) appears to be 25 K higher than the experimentally determined datum (Fig 9b). This discrepancy is related to the difficulties discussed above in describing the continuous thermodynamic properties of fluid in the region of the critical point.

Fig. 10a, b Diagram illustrating the method for determining the equilibrium ordering parameter corresponding to the Gibbs energy minimum. The diagrams are calculated for quartz under pressure of 10 kbar (a) and water under a pressure of 86 bar (b). *Arrows* relate minima of the Gibbs energy at a given temperature

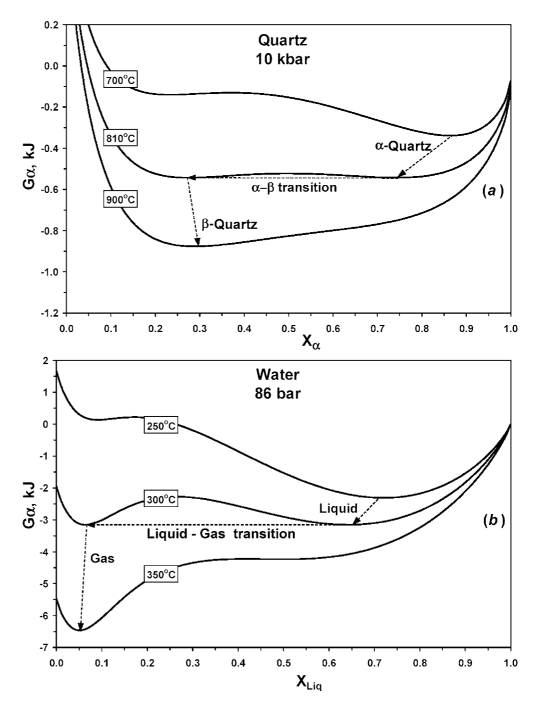


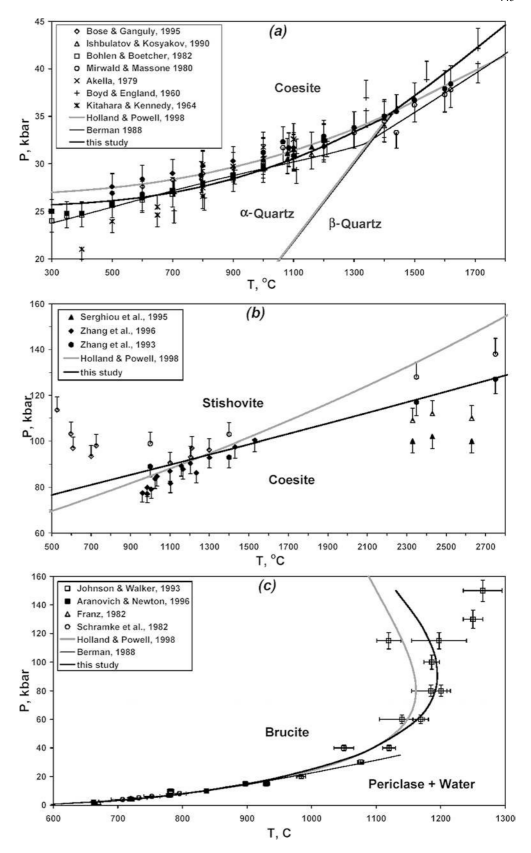
Figure 11 shows some mineral equilibria calculated using the data of Table 2. Comparison of the calculated phase boundaries with experimental data shows that the equations of Gibbs free energy derived here lead to accurate descriptions of the various phase boundaries, including the dehydration reaction brucite = periclase + water to very high pressures and temperatures (1250 °C, 150 kbar).

Discussion and conclusions

The equations for Gibbs free energy described here are sufficiently accurate to describe and extrapolate

thermodynamic properties and phase equilibria on the basis of experimental data on isobaric heat capacity and molar volume. This implies that they can be effectively used in thermodynamic data bases of petrological interest. Moreover, they are relatively simple and lead to a uniformity of representation of the Gibbs potential as a direct function of P and T for very different phases (the software implementing the equations discussed in this paper for thermodynamic calculations is available by request from the corresponding author). It should be noted that the proposed model employs not more than 16 adjustable parameters. By comparison, the Helmholtz free energy equations that allow reasonable

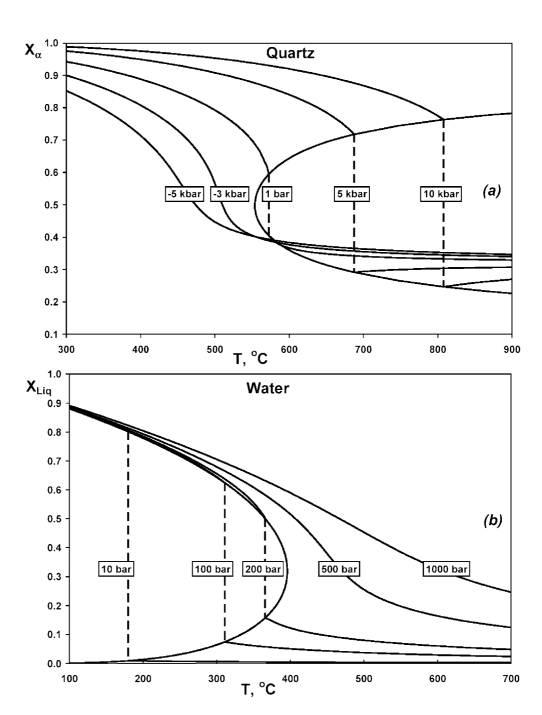
Fig. 11a-c Comparison of selected calculated phase transitions in the system MgO-SiO₂-H₂O with the experimental data



descriptions of a one-component fluids over a wide P-T range, including the two-phase region, normally contain 30–80 empirical parameters (e.g. Altunin 1975; Sychev

et al. 1979; Saul and Wagner 1989; Stewart and Jacobson 1989; Hill 1990). The Gibbs free energy equation for quartz proposed by Dorogokupets (1995) for a

Fig. 12a, b Isobars of the themperature dependence of ordering parameters for quartz (a) and water (b)

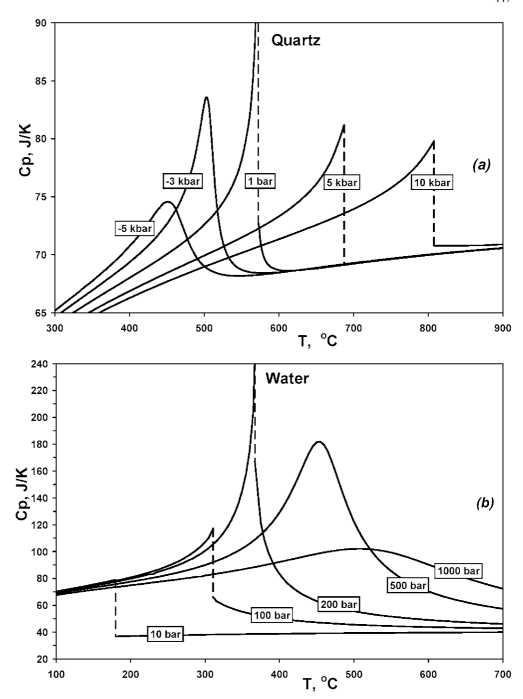


description of the λ -transition contains 26 parameters. In the case of simple phases, Equation (12) with 10–12 adjustable parameters allows an accurate description of the heat capacity and volume over a wide range of P–T conditions (20–3000 K, 0–800 kbar), including the important low-temperature (20–200 K) region relevant for correct representation of the third-law entropy according to Eq. (9a).

On the other hand, the approach presented here also leads to novel insights into the structure of crystalline phases with λ -transitions and fluids. The derived Gibbs free energy equations for such phases require the calculation of an equilibrium ordering parameter to satisfy Eqs.

(19) and (21), as illustrated by Fig. 10. Figure 12 shows the calculated P and T dependence of the ordering parameter for quartz (Fig.12a) and for water (Fig.12b). For quartz, the calculated miscibility gap widens with increasing pressure (the calculated critical point corresponds to T = 553 °C, P = -880 bar). A hypothetical homogeneous silica phase, with the ordering parameter X_{α} changing continuously with temperature, would exist below this point (Fig. 12a). These relationships are very similar to those in the aqueous fluid above the critical point (Fig. 12b). Accordingly, the heat capacity and coefficient of thermal expansion show maxima at the maximum value of the derivative $\partial X/\partial T$ (Figs. 13, 14).

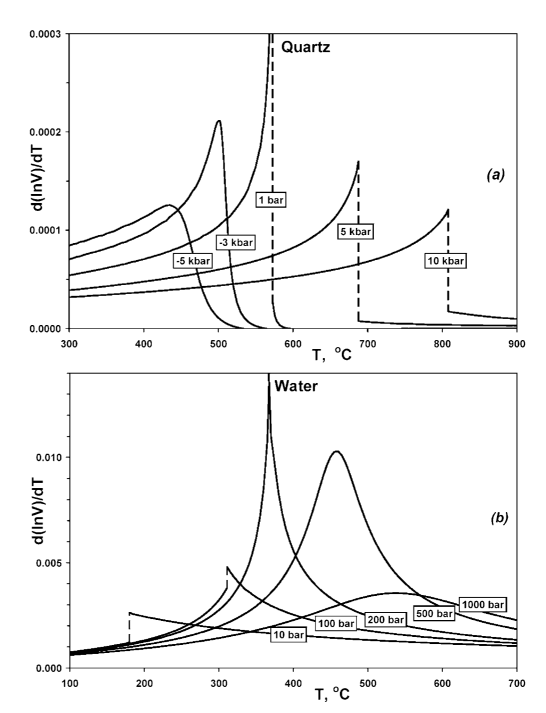
Fig. 13a, b Isobars of the temperature dependence of isobaric heat capacities for quartz (a) and water (b)



Comparison of the thermodynamic functions and ordering parameters of quartz and those of aqueous fluid (Figs. 13, 14) thus reveal a clear analogy, demonstrating the thermodynamic similarity of ordering phenomena in fluid and crystalline states (Landau and Lifshitz 1959). It should also be noted that the possible existence of a critical point for a crystalline substance with a λ -transition was predicted theoretically by Landau (e.g. Landau and Lifshitz 1959). Our test calculations have also shown that a similar thermodynamic approach can be applied to treat order/disorder transitions in binary alloys with pronounced λ -anomaly.

An important question concerns the physical meaning of the ordering parameters X_{α} and $X_{\rm Liq}$ used for the description of the Gibbs free energy of quartz and aqueous fluid, respectively. For both water and quartz, this parameter cannot be directly obtained from physical measurement, and therefore appears entirely abstract. However, a correspondence between $X_{\rm Liq}$ and the number of clusters follows from the similarity of Eqs. (22) and (24). Therefore, $X_{\rm Gas} = 1 - X_{\rm Liq}$ may be interpreted as an apparent mole fraction (thermodynamic probability) of the monomers, $X_{\rm I}$, in the fluid, and $X_{\rm Liq}$ is the bulk mole fraction (thermodynamic

Fig. 14a, b Isobars of the temperature dependence of the coefficient of thermal expansion for quartz (a) and water (b)

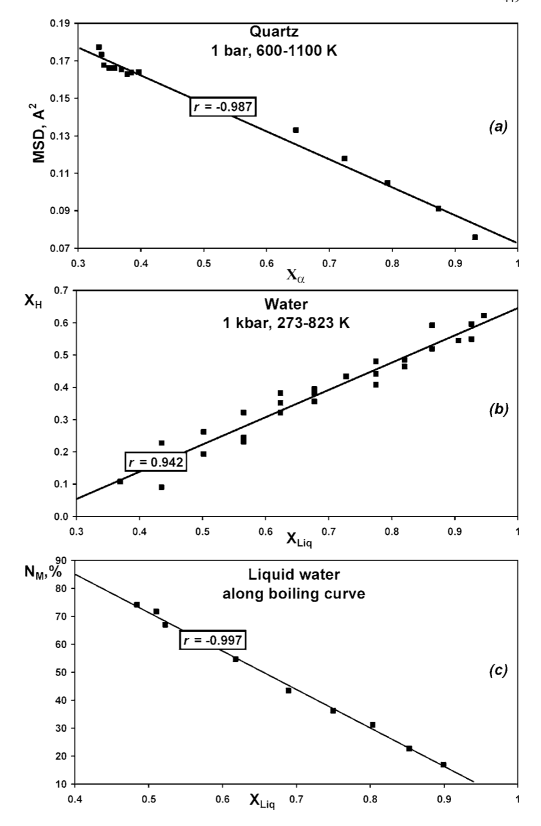


probability) of the clusters with $n \ge 2$. The concept of the existence of two distinct states of molecules in a fluid appears consistent with investigations of the water structure by Gorbaty and Demianets (1983), who demonstrated that the near-critical fluctuations are preserved at elevated P-T conditions above the critical point to a pressure of 1 kbar and temperatures of 500 °C. In the supercritical region, there are short-term space fluctuations that correspond to the liquid-like and gas-like states. Presence of clusters of water in the supercritical region is found in molecular dynamics experiments, even at densities of less than 0.2 gcm^{-3}

(Churakov and Kalinichev 1999), thus validating our thermodynamic formalism.

Further examples of the correspondence between the calculated ordering parameters and physical measurements are demonstrated in Fig. 15. Figure 15a presents the negative linear correlation of the estimated ordering parameter X_{α} with the sum of mean-squares displacements (MSD) of the oxygen atoms in the quartz structure, as determined for the λ -transition by Kihara (1990). The high correlation coefficient (r = -0.987) shows that the calculated decrease of X_{α} with increasing temperature reflects the real mechanism of disordering.

Fig. 15a-c Comparison of the ordering parameters with experimentally determined structural parameters for quartz (a) and water (b),(c) **a** A negative correlation (r = -0.987) between X_{α} and sum of mean-squares displacements (MSD) of oxygen atoms along axes 1-3 (Kihara 1990) at P = 1 bar and T = 600-1100 K. **b** A positive correlation (r =0.942) between X_{Liq} and mole fraction of hydrogen bonds $X_{\rm H}$ (Gorbaty and Kalinichev 1995) at a P = 1kbar and T = 0-550 °C. (c) A negative correlation (r = -0.997) between X_{Liq} and content of monomers $N_{\rm M}$ (mol%) (Luck 1980) in the liquid phase along the boiling curve



In Fig. 15b, the mole fraction of molecules in the liquidlike state $X_{\rm Liq}$ is compared to the mole fraction of hydrogen bonds $X_{\rm H}$, calculated by Gorbaty and Kalinichev (1995) from experimental data (the association of

molecules in an aqueous fluid is mainly determined by hydrogen bonds). Figure 14c presents a comparison of the mole fraction of molecules in the liquid-like state $X_{\rm Liq}$ with the mole percent of monomers $N_{\rm M}$ (Luck

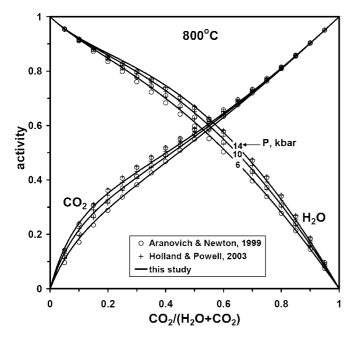


Fig. 16a–c Activity–composition plot for H₂O–CO₂ at 800 °C and 6–14 kbar calculated from eq. (27) with parameters linearly dependent on fluid composition (see text for discussion). Binary models of Aranovich and Newton (1999) and Holland and Powell (2003) are shown for comparison

1980). High coefficients of linear correlation between $X_{\rm H}$ and $X_{\rm Liq}$ (r=0.942) and between $N_{\rm M}$ and $X_{\rm Liq}$ (r=-0.997) suggest that $X_{\rm Liq}$ quantitatively characterizes the degree of association of molecules in an aqueous fluid. Thus, it can be concluded that our relatively abstract ordering parameter reflects physical processes in minerals and fluids.

Figure 16 shows the results of modelling the mixing properties of an H₂O-CO₂ fluid with Eq. (27) and parameters linearly dependent on fluid composition, i.e. $B_{\rm H_2O-CO_2} = B_{\rm CO_2} X_{\rm CO_2} + B_{\rm H_2O} X_{\rm H_2O}$, where $X_{\rm CO_2} = {\rm CO_2}$ /(CO₂ + H₂O), $X_{\rm H_2O} = {\rm H_2O}$ /(CO₂ + H₂O); $B_{\rm H_2O-CO_2}$ are parameters of Eq. (27) for the binary fluid of given composition; $B_{H,O}$ and B_{CO_2} are, respectively, empirical parameters of Eq. (27) for pure H₂O (Table 2) and pure CO₂ ($\Delta H^{o}_{ord} = -20608.153 \text{ J}, \Delta S^{o}_{ord} = -138.42315 \text{ J/K},$ $\Delta C_{P}^{o}_{ord} = 10.0415201 \text{ J/K}, W^{H} = -20859.049 \text{ J},$ $W^{S} = 36.8658607 \text{ J/K}, W^{Cp} = 0, \text{ and } \phi = 6551.312 \text{ bar},$ Gerya and Perchuk 1997). Our results coincide well with the empirical mixing models of Aranovich and Newton (1999) and Holland and Powell (2003) based on experimental data. This suggests that simple linear mixing rules can be used to extend our model on fluid mixtures without introducing any additional parameters. This also implies that non-ideality of mixing in multicomponent fluids is primarily related to changes in the degree of association of fluid molecules (X_{Liq}) with changing fluid composition, as directly accounted for by Eq. 27.

Acknowledgements The authors are grateful to V.B. Polyakov and O.L. Kuskov, Y.V. Petrov and A.G.Kalinichev for discussions on the details of the formulation and for fruitful consultations. We appreciate the detailed comments on an earlier version of this

manuscript by M.S. Ghiorso, Z. Duan, N. Dubrovinskaya and S.K. Saxena, who stimulated its substantial revision. The study was supported by the Russian Foundation for Basic Research (grants 02-05-64025, 03-05-64633, 1645-2003-5) and an Alexander von Humboldt Foundation Research Fellowship to Gerya. Helpful reviews by two anonymous reviewers are appreciated.

References

Ackermann RJ, Sorell CA (1974) Thermal expansion and the high low transformation in quartz, I. High-temperature X-ray studies. J Appl Crystallogr 1: 461–467

Akaogi M, Navrotsky A (1984) The quartz-coesite-stishovite transformations: new calorimetric measurements and calculation of phase diagrams. Phys Earth Planet Interiors 36: 124–134

Akaogi M, Yusa H, Shiraishi K, Suzuki T (1995) Thermodynamic properties of α-quartz, coesite, and stishovite and equilibrium phase relations at high pressures and high temperatures. J Geophys Res (B)100: 22337–22347

Akella J (1979) Quartz-coesite transition and the comparative friction measurement in piston-cylinder apparatus using talcalsimag-glass (TAG) and NaCl-high pressure cells. N Jb Mineral Monatsh 217–224

Akimoto S, Syono Y (1969) Coesite-stishovite transition. J Geophys Res 74: 1653–1659

Altunin VV (1975) Thermophysical properties of carbon dioxide (in Russia). Standards, Moscow, 551 pp

Aranovich LY, Newton RC (1996) H₂O activity in concentrated NaCl solutions at high pressures and temperatures measured by the brucite–periclase equilibrium. Contrib Mineral Petrol 125: 200–212

Aranovich LY, Newton RC (1999) Experimental determination of CO₂–H₂O activity–composition relations at 600–1000 °C and 6–14 kbar by reversal decarbonation and dehydration reactions. Am Mineral 84: 1319–1332

Barelko VV, Zakirov IV, Pechatnikov EL (1994) On the dependence of the molecular clusterization in a one-component gas medium on parameters of its equation of state (in Russian). Khimicheskaya Fizika 13: 42—50

Barron THK, Berg WT, Morrison JA (1959) On the heat capacity of crystalline magnesium oxide. Proceedings Roy Soci London 250(A): 70–83

Bassett WA, Barnett JD (1970) Isothermal compression of stishovite and coesite up to 85 kilobars at room temperature by X-ray diffraction. Phys Earth Planet Interiors 3: 54–60

Berman RG (1988) Internally consistent thermodynamic data for minerals in the system Na₂O-K₂O-CaO-MgO-FeOFe₂O₃-Al₂O₃-SiO₂-TiO₂-H₂O-CO₂. J Petrol 29: 445-522

Berman RG, Aranovich LY (1996) Optimized standard state and solution properties of minerals,1. Model calibration for olivine, orthpyroxene, cordierite, garnet, ilmenite in the system FEO–MGO-CaO-Al2O3-TiO3-SiO2. Contrib Mineral Petrol 126: 1-24

Boehler R (1982) Adiabats of quartz, coesite, olivine, and magnesium oxide to 50 kbar and 1000 K, and the adiabatic gradient in the Earth's mantle. J Geophys Res 87: 5501–5506

Boehler R, Skoropanov A, O'Mara D, Kennedy GC (1979) Gruneisen parameter of quartz, quartzite, and forsterite. J Geophys Res 84: 3527–3531

Bohlen SR, Boettcher AL (1982) The quartz-coesite transformation: a precise determination and the effects of other components. J Geophys Res 87: 7073–7078

Bose K, Ganguly J (1995) Quartz-coesite transition revisited: reversed experimental determination at 500–1200 °C and retrieved thermochemical properties. Am Mineral 80: 231–238 Boyd FR, England JL (1960) The quartz-coesite transition. J Geophys Res 65: 749–756

Bridgeman PW (1942) Freezing parameters and compressions of 21 substances to 50 000 kgcm⁻². Proc Am Acad Arts Sci 74: 399–424

- Brodholt JP, Wood BJ (1994) Measurements of the *PVT* properties of water to 25 kbar and 1600 °C from synthetic fluid inclusions in corundum. Geochim Cosmochim Acta 58: 2143–2148
- Carpenter MA, Salje EKH, Graeme-Barber A, Wruck B, Dove MT, Knight KS (1998) Calibration of excess thermodynamic properties and elastic constant variations associated with the α ←→ β phase transition in quartz. Am Mineral 83: 2–22
- Castex J, Madon M (1995) Test of the vibrational modelling for the lambda-type transitions—application to the α -B quartz transition. Phys Chem Miner 22: 1–10
- Catti M, Farraris G, Hull S, Pavese A (1995) Static compression and H disorder in brucite, Mg(OH)₂, to 11 GPa: a powder neutron diffraction study. Phys Chem Miner 22: 200–206
- Chakoumakos BC, Loong C-K, Schultz AJ (1997) Low-temperature structure and dynamics of brucite. J Phys Chem B101: 9458–9462
- Chatterjee ND, Miller K, Olbricht W (1994) Bayes estimation—a novel approach to derivation of internally consistent thermodynamic data for minerals, their uncertainties, and correlations ,2. Application. Phys Chem Miner 21: 50–62 1994
- Chatterjee ND, Kruger R, Haller G, Olbricht W (1998) The Bayesian approach to an internally consistent thermodynamic database: theory, database, and generation of phase diagrams. Contrib Mineral Petrol 133: 149–168
- Churakov SV, Kalinichev AG (1999) Size and structure of molecular clusters in supercritical water. J Structural Chem 40: 548–553
- CODATA task group on key values for thermodynamics. (1978) CODATA recommended key values for thermodynamics 1977. CODATA Bull 28: 1–17
- Cohen LH, Klement WK Jr (1967) High-low quartz inversion: determination to 35 kilobars. J Geophys Res 72: 4245–4251
- d'Amour H, Denner W, Schulz H (1979) Structure determination of α-quartz up to 68·10⁸ Pa. Acta Crystallogr 35: 550–555
- Danielsson S, Grenthe I, Oskarsson A (1976) A low-temperature apparatus for single-crystal diffractometry: the unit-cell dimensions of α-quartz in the temperature range 86–298 K. J Appl Crystallogr 9: 14–17
- Dewaele A, Fiquet G, Andrault D, Hausermann D (2000) P-V-T equation of state of periclase from synchrotron radiation measurements. J Geophys Res 105: 2869–2878
- Dorogokupets PI (1995) Equation of state for lambda transition in quartz. J Geophys Res (B)100: 8489–8499
- Dorogokupets PI, Karpov IK (1984) Thermodynamics of minerals and mineral equilibria (in Russian). Nauka, Novosibirsk, 185 pp
- Doroshev AM, Galkin VM, Kuznetzov GN (1987) Thermal expansion of stishovite in temperature interval 98-420 K (in Russian). Geokhimiya 10: 1463–1467
- Dove MT (1997) Theory of displacive phase transitions in minerals. Am Mineral 82: 213–244
- Duan Z, Møller N, Weare J (1996) A general equation of state for supercritical fluid mixtures and molecular dynamics simulation of mixture PVTX properties. Geochim Cosmochim Acta 60: 1209–1216
- Dubrovinskaya NA, Dubrovinsky LS, Saxena SK (1997) Systematics of thermodynamic data on solids: thermochemical and pressure–volume–temperature properties of some minerals. Geochim Cosmochim Acta 61: 4151–4158
- Endo S, Akai T, Akahama Y, Wakatsuki M, Nakamura T, Tomai Y, Koto K, Ito Y, Tokonami M (1986) High-temperature X-ray study of single-crystal stishovite synthesized with Li₂WO₄ as flux. Phys Chem Miner 13: 146–151
- Fei Y (1999) Effects of temperature and composition on the bulk modulus of (Mg,Fe)O. Am Mineral 84: 272–276
- Fei Y, Mao HK (1993) Static compression of Mg(OH)₂ to 78 GPa at high temperature and constraints on the equation of state of fluid H₂O. J Geophys Res 98: 11875–11884
- Fiquet G, Richet P, Montagnac G (1999) High-temperature thermal expansion of lime, periclase, corundum and spinel. Phys Chem Miner 27: 103–111
- Franz G (1982) The brucite-periclase equilibrium at reduced H₂O activities: some information about the system H₂O-NaCl. Am J Sci 282: 1325–1339

- Galkin VM, Doroshev AM, Babich YV (1987) Thermal expansion of coesite (in Russian). Geokhimiya 11: 1645–1647
- Gerya TV, Perchuk LL (1997) Equations of state of compressed gases for thermodynamic databases used in petrology. Petrology 5: 366–380
- Gerya TV, Podlesskii KK, Perchuk LL, Swamy V, Kosyakova NA (1998) Equations of state of minerals for thermodynamic databases used in petrology. Petrology 6: 511–526
- Giauque WF, Archibald RC (1937) The entropy of water from the third law of thermodynamics. The dissociation pressure and calorimetric heat of the reaction Mg(OH)₂ = MgO + H₂O. The heat capacities of Mg(OH)₂ and MgO from 20 to 300 K. J Am Chem Soc 59: 561–569
- Gibson RE (1928) The influence of pressure on the high-low inversion of quartz. J Phys Chem 32: 1197–1210
- Glinnemann J, King HE Jr, Schulz H, Hahn T, LaPlaca SJ, Dacol F (1992) Crystal structures of the low-temperature quartz-type phases of SiO₂ and GeO₂ at elevated pressure. Z Kristallogr 198: 177–212
- Glushko VP, Gurvich LV, Bergman GA, Weiz IV, Medvedev VA, Khachkuruzov GA, Yungman VS (eds) (1979) Themodynamic properties of individual substances, vol 2, (in Russian). Moscow: Nauka, 784 pp
- Gorbaty YE, Demianets YN (1983) The pair correlation functions of water at a pressure of 1000 bar in temperature range 25–500 °C. Chem Phys Lett 100: 450–453
- Gorbaty YE, Kalinichev AG (1995) Hydrogen bonding in supercritical water, 1. Experimental results. J Phys Chem 99: 5336
- Gottschalk M (1997) Internally consistent thermodynamic data for rock-forming minerals in the system SiO₂–TiO₂–Al₂O₃–Fe₂O₃–CaO–MgO–FeO–K₂O–Na₂O–H₂O–CO₂. Eur J Mineral 9: 175–223
- Grevel K-D (1995) Internally consistent thermodynamic data for high-pressure phases in the system CaO–MgO–Al₂O₃–SiO₂–H₂O. Bochumer Geol Geotech Arb 44: 64–69
- Grønvold F, Stølen S, Svendsen SR (1989) Heat capacity of α-quartz from 298.15 to 847.3 K, and of β-quartz from 847.3 to 1000 K—transition behavior and reevaluation of the thermodynamic properties. Thermochim Acta 139: 225–243
- Gupta RB, Panayiotou CG, Sanchez IC, Johnston KP (1992)
 Theory of hydrogen bonding in supercritical fluids. American Institute of Chemical Engineers Journal J 38: 1243–1253
- Gurevich VM, Khlyustov VG (1979) A calorimeter for measurements of low-temperature heat capacity of quartz at 9-300 K (in Russian). Geokhimiya 6: 829–839
- Hazen RM, Finger LW, Hemley RJ, Mao HK (1989) High-pressure crystal chemistry and amorphization of α -quartz. Solid State Commun 72: 507–511
- Helgeson HC, Delany JM, Nesbitt HW, Bird DK (1978) Summary and critique of the thermodynamic properties of rock-forming minerals. Am J Sci 278A: 229 pp
- Hemingway BS (1987) Quartz: heat capacities from 340 to 1000 K and revised values for the thermodynamic properties. Am Mineral 72: 273–279
- Hemingway BS, Robie RA, Evans HT Jr, Kerrick DM (1991) Heat capacities and entropies of sillimanite, fibrolite, andalusite, kyanite, and quartz and the Al₂SiO₅ phase diagram. Am Mineral 76: 1597–1613
- Hill TL (1956) Statistical mechanics. Principles and selected applications, McGraw-Hill New York, 432 pp
- Hill PG (1990) An unified fundamental equation for the thermodynamic properties of H₂O. J Phys Chem Ref Data 19: 1233–1274
- Holland TJB, Powell R (1990) An enlarged and updated internally consistent thermodynamic data set with uncertainties and correlations: the system K₂O-Na₂O-CaO-MgO-FeO-Fe₂O₃-Al₂O₃-TiO₂-SiO₂-C-H₂-O₂. J Metamorph Geol 8: 309–343
- Holland TJB, Powell R (1996) Thermodynamics of order–disorder in minerals, I. Symmetric formalism applied to minerals of fixed composition. Am Mineral 81: 1413–1424
- Holland TJB, Powell R (1998) Internally consistent thermodynamic data set for phases of petrological interest. J Metamorph Geol 16: 309–344

- Holland TJB, Powell R. (2003) Activity composition relations for phases of petrological calculations: an asymmetric multicomponet formulation. Contrib Mineral. Petrol. 145: 492–501
- Holm JL, Kleppa OJ, Westrum EF (1967) Thermodynamics of polymorphic transformations in silica. Thermal properties from 5 to 1070 K and pressure-temperature stability fields for coesite and stishovite. Geochim Cosmochim Acta 31: 2289–2307
- Ishbulatov RA, Kosyakov AV (1990) Problems of pressure estimation in high-temperature experiments using solid media apparatus; pressure calibration with reference to breakdown of albite and quartz–coesite transformation (part II). Proc Indian Acad Sci 99: 133–140
- Jay AH (1933) The thermal expansion of quartz by X-ray measurements. Proc Roy Soc London 142A(846): 237–247
- Johnson MC, Walker D (1993) Brucite [Mg(OH)₂] dehydration and the molar volume of H_2O to 15 GPa. Am Mineral 78: 271-284
- Jorgensen JD (1978) Compression mechanisms in α -quartz structures—SiO₂ and GeO₂. J Appl Phys 49: 5473–5478
- Karpov IK, Kiselev AI, Letnikov FA (1976) Computer modeling of natural mineral formation (in Russian). Nedra, Moscow, 256 pp
- Kieffer SW (1985) Heat capacity and entropy: systematic relations to lattice vibrations. Mineralogical Society of America, Washington DC, Rev Mineralogy, vol 14: pp 65–126
- Kihara K (1990) An X-ray study of the temperature dependence of the quartz structure. Eur J Mineral 2: 63–77
- Kitahara S, Kennedy GC (1964) The quartz-coesite transition. J Geophys Res 69: 5395–5400
- Koster van Groos, AF, Ter Heege JP (1973) The high-low quartz transition up to 10 kilobars pressure. J Geol 81: 717–724
- Krupka KM, Robie RA, Hemingway BS (1979) High-temperature heat capacities of corundum, periclase, anorthite, CaAl₂Si₂O₈ glass, muscovite, pyrophyllite, KAlSi₃O₈ glass, grossular, and NaAlSi₃O₈ glass. Am Mineral 64: 86–101
- Kubo R (1965) Statistical mechanics. North-Holland, Amsterdam, 425 pp
- Kuskov OL, Zidikova AP, Semenov YV, Babich YV, Fabrichnaya OV (1991) Thermodynamics of polymorph modyfications of silica (in Russian). Geokhimiya 8: 1175–1185
- Kut'in AM, Pyadushkin DV (1998) Analytical approximation of thermodynamic functions of solids at the basis of phenomenological statistics of interaction nodes (in Russian). Z Fizicheskoy Khimii 72: 1735–1740
- Lager GA, Jorgensen JD, Rotella FJ (1982) Crystal structure and thermal expansion of α-quartz at low temperatures. J Appl Phys 53: 6751–6756
- Landau LD, Lifshitz EM (1959) Statistical physics. Pergamon, London, 484 pp
- Levien L, Prewitt CT (1981) High-pressure crystal structure and compressibility of coesite. Am Mineral 66: 324–333
- Levien L, Prewitt CT, Weidner DJ (1980) Structure and elastic properties of quartz at pressure. Am Mineral 65: 920–930
- properties of quartz at pressure. Am Mineral 65: 920–930 Liu LG, Basset WA, Takahashi T (1974) Effect of pressure on the
- lattice parameters of stishovite. J Geophys Res 79: 1160–1164 Luck WAP (1980) A model of hydrogen-bonded liquid. Angew Chem Int Ed Eng 19: 28
- Mirwald PW, Massone H-J (1980a) Quartz–coesite transition and the comparative friction measurement in piston-cylinder apparatus using talc-alsimag-glass (TAG) and NaCl high-pressure cells: a discussion. N Jb Mineral Monatsh 10: 469–477
- Mirwald PW, Massone H-J (1980b) The low-high quartz and quartz-coesite transition to 40 kbar between 600 and 1600 °C and some reconnaissance data on the effect of NaAlO₂ component on the low quartz-coesite transition. J Geophys Res 85: 6983–6990
- Nagai T, Hattori T, Yamanaka T (2000) Compression mechanism of brucite: an investigation by structural refinement under pressure. Am Mineral 85: 760–764
- Olinger B (1976) The compression of stishovite. J Geophys Res 81: 5341–5343
- Olinger B, Halleck PM (1976) The compression of α -quartz. J Geophys Res 81: 5711–5714

- Parise JB, Leinenweber K, Weidner DJ, Tan K, Von Dreele RB (1994) Pressure-induced H bonding: neutron diffraction study of brucite, Mg(OH)₂, to 9.3 GPa. Am Mineral 79: 193–196
- Polyakov VB, Kuskov OL (1994) Self-consistent model for calculation of thermoelastic and caloric properties of minerals (in Russian). Geokhimiya 7: 1096–1122
- Prigogine I, and Defay R (1954) Chemical thermodynamics. Longmans Green, London
- Reeber RR, Goessel K, Wang K (1995) Thermal expansion and molar volume of MgO, periclase, from 5 to 2900 K. Eur J Mineral 7: 1039–1047
- Rice MH, Walsh JM (1957) Equation of state of water to 250 Kbars. J Chem Phys 26: 824–830
- Richet P, Fiquet G (1991) High-temperature heat capacity and premelting of minerals in the system MgO–CaO–Al₂O₃–SiO₂. J Geophys Res 96: 445–456
- Richet P, Bottinga Y, Denielou L, Petitet JP, Tequi C. (1982) Thermodynamic properties of quartz, cristoballite and amorphous SiO2: drop calorimetry measurements between 1000 and 1800 K and a review from 0 to 2000 K. Geochim Cosmochim Acta 46: 2639–2658
- Richet P, Gillet P, Fiquet G (1992) Thermodynamic properties of minerals: macroscopic and microscopic approaches. Adv Phys Geohem 10: 98–131
- Ross NL, Shu J-F, Hazen RM, Gasparik T (1990) High-pressure crystal chemistry of stishovite. Am Mineral 75: 739–747
- Sato Y (1977) Pressure-volume relationship of stishovite under hydrostatic compression. Earth Planet Sci Lett 34: 307–312
- Saul A, Wagner W (1989) A fundamental equation for water covering the range from melting line to 1273 K at pressure up to 25 000 Mpa. J Phys Chem Ref Data 18: 1537–1563
- Saxena SK, Chatterjee N, Fei Y, Shen G (1993) Thermodynamic data on oxides and silicates: an assessed data set based on thermochemistry and high-pressure phase equilibrium. Springer, Berlin, Heidelberg, New York, 428 pp
- Serghiou G, Zerr A, Chudinovskih LT, Boehler R (1995) The coesite–stishovite transition in a laser-heated diamond cell. Geophys Res Lett 22: 441–444
- Schramke JA, Kerrick DM, Blencoe JG (1982) The experimental determination of the brucite–periclase + water equilibrium with a new volumetric technique. Am Mineral 67: 269–276
- Shen AH, Basset WA, Chou I-M (1993) The α - β quartz transition at high temperatures and pressures in a diamond-anvil cell by laser interferometry. Am Mineral 78: 694–698
- Smyth JR, Smith JV, Artioli G, Kvick A (1987) Crystal structure of coesite, a high-pressure form of SiO₂, at 15 and 298 K from single-crystal neutron and X-ray diffraction data: test of bonding models. J Phys Chem 91: 988–992
- Spearing DR, Farnan I, Stebbins JF (1992) Dynamics of the α-β phase transitions in quartz and cristobalite as observed by insitu high-temperature ²⁹Si and ¹⁷O NMR. Phys Chem Miner 19: 307–321
- Stewart RB, Jacobson RT (1989) Thermodynamic properties of argon from the triple point to 1200 K with pressures to 1000 MPa. J Phys Chem Ref Data 18: 639–798
- Suito K (1977) Phase relations of pure Mg₂SiO₄ up to 200 kilobars. In: Manghnani MH, Akimoto S (eds) High-pressure research: application to geophysics. Academic Press, San Diego, pp 365–371
- Suito K, Miyoshi M, Onodera A, Shimomura O, Kikegawa T (1996) Thermal expansion studies of stishovite at 10.5 GPa using synchrotron radiation. Phys Earth Planet Interiors 93: 215–222
- Sychev VV, Wasserman AA, Zagoruchenko VA, Kozlov AD, Spiridonov GA, Tsimarny VA (1979) Thermodynamic properties of methane (in Russian). Standards, Moscow; 348 pp
- Thompson JB Jr (1969) Chemical reactions in crystals. Am Mineral 54: 341–375
- Toda M, Kubo R, Saito N (1992) Statistical physics, I. Equilibrium statistical mechanics (2nd ed). Springer, Berlin Heidelberg, New York, 252 pp

Tsuchida Y, Yagi T (1989) A new, post-stishovite high-pressure polymorph of silica. Nature 340: 217–220

Vaidya SN, Bailey S, Pasternack T, Kennedy GC (1973) Compressibility of 15 minerals to 45 kilobars. J Geophys Res 78: 6893–6898

Wagner W, Pruss A (1997) Die neue internationale Standard-Zustandsgleihung für Wasser für den allgemeinen und wissenschaftlichen Gebrauch, Jahrbuch 97. VDI-Gesllst Verfahrenstechnik und Chemieingenieurwesen, VDI-Verlag, Düsseldorf, pp 134–156

Watanabe H (1982) Thermochemical properties of synthetic high pressure compounds relevant to the Earth's mantle. In: Akimoto S, Manghnani M (eds) High-pressure research in geophysics. Center for Academic Publications, Tokyo, pp 441–464

Xia X, Weidner DJ, Zhao H (1998) Equation of state of brucite: single-crystal Brillouin spectroscopy study and polycrystalline pressure-volume-temperature measurement. Am Mineral 83: 68-74

Yagi T, Akimoto S (1976) Direct determination of coesite–stishovite transition by in situ X-ray measurements. Tectonophysics 35: 259–270

Yoder HS (1950) High-low quartz inversion up to 10 000 bars. Transactions American Geophysical Union, Washington DC, 31: 827–835

Zhang J (2000) Effect of pressure on the thermal expansion of MgO up to 8.2 Gpa. Phys Chem Miner 27: 145–148

Zhang J, Liebermann RC, Gasparik T, Herzberg CT, Fei Y (1993) Melting and subsolidus relations of SiO₂ at 9–14 GPa. J Geophys Res (B)98: 19785–19793

Zhang J, Li B, Utsumi W, Liebermann RC (1996) In situ X-ray observations of the coesite-stishovite transition: reversed phase boundary and kinetics. Phys Chem Miner 23: 1–10

Zharkov VN, Kalinin VA (1971) Equations of state of solids at high pressures and temperatures. Consultant Bureau, New York, 257 pp

Appendix 1. Derivation of Eq. (2)

The derivation of Eq. (2) follows standard logic of statistical mechanics (e.g. Hill 1956; Kubo 1965; Toda et al. 1992; Landau and Lifshitz 1959), but is based on an ensemble not treated in standard textbooks on statistical mechanics.

Let us first consider a substance in a condensed state, solid or liquid, at a relatively low temperature, when its atoms are close to each other and vibrate mainly around equilibrium positions. A system with N atoms will have 3N vibrational degrees of freedom, and, from the viewpoint of mechanics, can be referred to as a system of 3N independent quantum oscillators (e.g. Landau and Lifshitz 1959). The Gibbs free energy of a system of quantum oscillators in equilibrium at constant P and T can also be expressed with the P-T partition function (Toda et al. 1992)

$$G = -k_B T \ln(Y), \tag{A1}$$

$$Y = \sum_{i} \int_{0}^{\infty} \exp\{-[E_{i}(V) + PV]/k_{B}T\} dV$$
 (A2)

where Y is the statistical sum over states, k_B is Boltzmann's constant, V is the system volume and $E_i(V)$ is the system energy for the i-th quantum state as a function of the volume. Theoretical expressions for $E_i(V)$ are

unavailable and empirical approximations are used instead. For integration of Eq. (A2) we take into account that at given P and T the fluctuation of volume of condensed substance is insignificant. If for an i-th state at a given P only one system volume $V_i(P)$ refers to a non-zero probability, then

$$\int_{0}^{\infty} \exp\{-[E_{i}(V) + PV]/k_{B}T\}dV = \exp[-H_{i}(P)/k_{B}T],$$
(A3)

where $H_i(P) = E_i(P) + PV_i(P)$ and $E_i(P) = E_i[V_i(P)]$.

 $H_i(P)$, $E_i(P)$ and $V_i(P)$ are enthalpy (or, more specifically, its microscopic analogue), energy and volume of the system, respectively, in an *i*-th state as functions of P. If the apparent volume $V_i(P)$ and corresponding $H_i(P)$ and $E_i(P)$ in Eq. (A3) are valid for a case of the i-th quantum state at given P with insignificant fluctuations of the system volume V relative to $V_i(P)$, then Eq. (A2) can be simplified to

$$Y = \sum_{i} \exp[-H_i(P)/k_B T]. \tag{A4}$$

Equation (A4) differs from equations normally given in statistical mechanics (e.g. Hill 1956) in that the volume integral is taken before summation of the quantum states of the oscillators.

Normalizing the partition function relative to zeropoint vibrations gives

$$G = G_0(P) - k_B T \ln(Y_0), \tag{A5}$$

where $Y_0 = \sum \exp[-\Delta H_i(P)/\ k_BT]$, $G_0 = H_0(P) - TS_0$, $\Delta H_i(P) = H_i(P) - H_0(P) = \Delta E_i(P) + P\Delta V_i(P), H_0(P) = E_0(P) + PV_0(P)$, $\Delta E_i(P) = E_i(P) - E_0(P)$, $\Delta V_i(P) = V_i(P) - V_0(P). Y_0$ is the statistical sum normalized relative to the state of zero-point vibrations. $G_0(P)$, $H_0(P)$, $E_0(P)$ and $V_0(P)$ are, respectively, the pressure-dependent Gibbs free energy, enthalpy, energy and volume of the system for the state of zero-point vibrations. S_0 is entropy of the system for the state of zero-point vibrations (in case of ordered crystalline phases, it is equal to zero according to Nernst's law). $\Delta H_i(P)$, $\Delta E_i(P)$ and $\Delta V_i(P)$ are, respectively, the pressure-dependent changes of enthalpy, energy and volume due to transition from the state of zero-point vibrations to the *i*-th quantum state. Taking into account 3N independent oscillators:

(A1)
$$Y_0 = \prod_{i}^{3N} Y_{0i},$$
(A2) where $Y_{0i} = \sum_{j}^{M} \exp[-\Delta H_{ij}(P)/k_B T],$

$$\Delta H_{ij}(P) = \Delta E_{ij}(P) + P\Delta V_{ij}(P).$$

 Y_{0i} is the statistical sum for the *i*-th oscillator normalized relative to the state of zero-point vibrations, M is the number of energy levels for the *i*-th oscillator, $\Delta H_{ij}(P)$ $\Delta E_{ii}(P)$ and $\Delta V_{ij}(P)$ are, respectively, pressure-dependent

changes in enthalpy, energy and volume due to transition of the *i*-th oscillator from the state of zero-point vibrations to the *j*-th energy level. To calculate Y_{0i} , it is assumed that every oscillator has an infinite number of energy levels and that $\Delta H_{ij}(P)$, $\Delta E_{ij}(P)$ and $\Delta V_{ij}(P)$ are proportional to *j* (e.g. Landau and Lifshitz 1959). Then

$$\Delta H_{ij}(P) = j\Delta H_{i0}(P),\tag{A7}$$

$$Y_{0i} = 1/\{1 - \exp[-\Delta H_{i0}(P)/k_B T]\},\tag{A8}$$

$$\Delta H_{i0}(P) = \Delta E_{i0}(P) + \Delta V_{i0}(P). \tag{A9}$$

$$G = H_0(P) - TS_0 + k_B T \sum_{i=1}^{3N} \ln\{1 - \exp[-\Delta H_{i0}(P)/k_B T]\},$$

where $\Delta H_{i0}(P)$, $\Delta E_{i0}(P)$, and $\Delta V_{i0}(P)$ are, respectively, pressure-dependent changes in enthalpy, energy and volume due to transition of the *i*-th oscillator between neighbouring energy levels. The form of Eq. (A10) is similar to a standard equation for the Helmholtz free energy of the system of independent quantum oscillators (e.g. Landau and Lifshitz 1959; Kubo 1965; Toda et al. 1992), and the only difference is that the statistical sum over states is expressed in terms of enthalpy changes (not energy changes) related to the transitions of the oscillators.

If $N = vN_A$ (N_A is Avogadro's number, v is the number of atoms in a substance molecule) and functions $\Delta H_{i0}(P)$ have similar values for n big groups of oscillators, then Eq. (A9) can be rewritten in the form of Eq. (2)

Appendix 2. Derivation of Eq. (16)

The derivation of Eq. (16) is also based on the P-T partition function (Toda et al. 1992) and follows the same logic as the derivation of Eq. 2 (see Appendix 1).

If the λ -transition leads to notable changes in the thermal vibrations of atoms, this must be taken into account in a description of the energy transitions of oscillators; i.e. G_s in Eq. (2) depends on X_α . Suppose that for a certain oscillator energy level m, whose energy depends on X_α , the enthalpy effect of the transition ΔH_{si} changes. Then, the statistical sums Y_{0i} in Eq. (A6) take the form

$$Y_{0i} = \{ [1 - \exp(-\Delta H_{i\beta}/k_B T)] + \exp(-\Delta H_{i\lambda}/k_B T) \times [\exp(-\Delta H_{i\beta}/k_B T) - \exp(-\Delta H_{i\alpha}/k_B T)] \} / \{ [1 - \exp(-\Delta H_{i\alpha}/k_B T)] \cdot [1 - \exp(-\Delta H_{i\beta}/k_B T)] \},$$
(A11)

where $\Delta H_{i\alpha}$ is the enthalpy change of the energy transitions (ΔH_{i0}) beginning with the zero level to the level (m-1) in the ordered phase, $\Delta H_{i\beta}$ is the enthalpy change of the energy transitions beginning with the level m in the disordered phase, $\Delta H_{i\lambda}$ is the enthalpy change of the transition from the zero level to the level m as a function of X_{α} and can be expressed by a power series expansion:

$$\Delta H_{i\lambda} = \sum_{k=0}^{u} \Delta H_{i\lambda k} (X_{\alpha})^{k}, \tag{A12}$$

where $\Delta H_{i\lambda k}$ denotes the expansion coefficients. Then Eq. (A10) can be transformed to:

$$G_{s} = H_{s} - TS_{s}^{o} + \sum_{i=1}^{n} c_{i}RT\ln\{(1 - e_{\alpha i})(1 - e_{\beta i})/[(1 - e_{\beta i})]\}$$

$$+ e_{\lambda i}(e_{\beta i} - e_{\alpha i})]\}, \tag{A13}$$

where $e_{\alpha i} = \exp(-\Delta H_{si\alpha} / RT)$, $e_{\beta i} = \exp(-\Delta H_{si\beta} / RT)$, $e_{\lambda i} = \exp(-\Delta H_{si\lambda} / RT)$, $e_{oi} = \exp(-\Delta H_{si\alpha}^{\circ} / RT_{o})$, $\Delta H_{si\alpha} = \Delta H_{si\alpha}^{\circ} + \Delta V_{si\alpha}^{\circ} \Psi$, $\Delta H_{si\beta} = \Delta H_{si\beta}^{\circ} + \Delta V_{si\beta}^{\circ} \Psi$, and $\Delta H_{si\lambda} = \sum_{k=0}^{u} (\Delta H_{si\lambda k}^{\circ} + \Delta V_{si\lambda k}^{\circ} \Psi)(X_{\alpha})^{k}$.

 $\Delta H_{\text{s}i\alpha}$, $\Delta H_{\text{s}i\beta}$, and $\Delta H_{\text{s}i\lambda}$ are values of $\Delta H_{i\alpha}$, $\Delta H_{i\beta}$ and $\Delta H_{i\lambda}$ taken for N_A oscillators. Equation (16) is derived from Eq. (A13) by using u = 2, $n \le 5$ and Eqs. (5)–(6).

Appendix 3. Derivation of Eq. (27)

The last two terms in Eq. (26) can be expressed as

$$-X_{\text{Gas}}\Delta G_{\text{ord}}^o + G^e = G_P + G_T, \tag{A14}$$

where
$$G_P = \int_{P_O}^P (-\Delta V_{\text{ord}}^o X_{\text{Gas}} - V^e) dP$$
, and $G_T = -X_{\text{Gas}}$
 $(\Delta H_{\text{ord}}^o - T\Delta S_{\text{ord}}^o) + H^e - TS^e + \int_{T_o}^T [-\Delta C_P^o(T)_{\text{ord}} X_{\text{Gas}} + C_P^e(T)] dT - T \int_T^T [(-\Delta C_P^o(T)_{\text{ord}} X_{\text{Gas}} + C_P^e(T))/T] dT$.

 $\Delta V_{\rm ord}^o$ and $V^{\rm e}$ are the standard volume change of short-term "ordering" transition 25 (Gas = Liq) and the excess volume of the fluid, respectively, both being functions of P and T, $\Delta H_{\rm ord}^o$ and $\Delta S_{\rm ord}^o$ are the standard enthalpy and entropy changes of Reaction (25) at $P_{\rm o}$ and $T_{\rm o}$, $H^{\rm e}$ and $S^{\rm e}$ are, respectively, contributions to excess enthalpy and entropy not related to $V^{\rm e}$ at $P_{\rm o}$ and $T_{\rm o}$, ΔC_P^o (T)_{ord} and C_P^e (T) are, respectively, the standard heat capacity change of "ordering" transition (25) and the excess heat capacity of the fluid, both dependent on temperature at $P_{\rm o}$.

To quantify G_P from Eq. (A14) it can be assumed that the volume of a phase that contains only molecules in the gas-like (free) state can be expressed as two terms, namely, the conventional volume of the molecules $V_{\rm s}$ and the free volume $V_{\rm f}$ (Prigogine and Defay 1954). Then

$$V_{\text{Gas}}^o = V_{\text{s}} + V_{\text{f}} = V_{\text{s}} + RT/P.$$
 (A15)

The volume of one mole of a substance that contains only molecules in the liquid-like (associated) state is solely determined by the conventional volume of the molecules

$$V_{\text{Liq}}^o = V_{\text{s}}.\tag{A16}$$

Hence, the standard volume change of Reaction (25) is $\Delta V_{\rm ord}^{\rm o} = V_{\rm Liq}^{\rm o} - V_{\rm Gas}^{\rm o} = -{\rm R}T/P$.

Considering Eqs. (A15) and (A16), the total volume V of a mixture consisting of molecules in both the liquidlike and the gas-like states

$$V = X_{Gas}V_{Gas}^{o} + X_{Liq}V_{Liq}^{o} - V^{e} = V_{s} - X_{Gas}\Delta V_{ord}^{o} - V^{e}$$

= $V_{s} + X_{Gas}RT/P - V^{e}$. (A17)

The negative sign of V^{e} corresponds to a decrease in the free volume $V_{\rm f}$ caused by attraction between molecules of any type. Then, in line with the van der Waals equation

$$P = \mathbf{R}T/(V - \beta) - \phi_{\mathbf{a}},$$

where β is the correction for molecular volume, i.e. β $V_{\rm s}$, and $\phi_{\rm a}={\rm const}/V^2$ is the correction for the attractive interaction between the molecules. Equation (A17) can be rewritten as

$$V = V_{\rm s} + X_{\rm Gas} RT/P - V^{\rm e} = V_{\rm s} + X_{\rm Gas} RT/(P + \phi_{\rm a}).$$

Apparently, ϕ_a depends on the average distance between molecules in a fluid. Taking into account the volume change of Reaction (25), this distance is a function of $X_{\rm Lig}$. As the external pressure decreases, the volume of the system approaches that of the ideal gas, RT/P, and both X_{Liq} and ϕ_a tend to zero. As the pressure increases, $X_{\rm Liq}$ and $\phi_{\rm a}$ also increase. Hence, $\phi_{\rm a}$ can be expressed as a function of $X_{\rm Liq}$ in the form

$$\phi_{\rm a} = \phi_{\rm a1} X_{\rm Liq} + \phi_{\rm a2} X_{\rm Liq}^2 + \dots + \phi_{\rm aq} X_{\rm Liq}^q,$$
 (A19)

where $\phi_{a1}, \phi_{a2}, \ldots$, and ϕ_{aq} are coefficients independent of $X_{\rm Liq}$. Evidently, with $X_{\rm Liq}$ approaching 1, $\phi_{\rm a}$ approaches ϕ (Eq. 5), because both corrections characterize the same

attractive interaction between molecules. Then, in accordance with Eq. (A19)

$$\phi = \phi_{a1} + \phi_{a2} + \dots + \phi_{aq}. \tag{A20}$$

For a transition from the liquid-like state to the gas-like one, when distances between the molecules increase significantly, Eqs. (A18)–(A19) provide an additional correction for the attractive interaction. Taking for the standard state $\phi_a = 0$, integration of Eq. (A17) in accordance with Eq. (A14) yields the following equation for G_P

$$G_P = X_{\text{Gas}} R T \ln[(P + \phi_a)/P_o,$$
 (A21)
where $\phi_a = \phi_{a1} X_{\text{Liq}} + \phi_{a2} X_{\text{Liq}}^2 + \dots + \phi_{aq} X_{\text{Liq}}^q$.

Using the Margules expansion to express the integral excess Gibbs free energy of mixing of liquid-like and gaslike molecules in a fluid and assuming $\Delta C_{Pord}^{o}(T)$ and $C_{\rm P}^{\rm e}(T)$ constant, G_T in Eq. (A14) can be defined as

$$G_{T} = -X_{Gas} \{ \Delta H_{ord}^{o} - T \Delta S_{ord}^{o} + \Delta C_{Pord}^{o} [T - T_{o} - T \ln(T/T_{o})] \} + \sum_{i=1}^{m} W_{j}^{G} X_{Gas} X_{Liq}^{j},$$
 (A22)

where $W_j^{\rm G}=W_j^{\rm H}-TW_j^{\rm S}+W_j^{\rm Cp}[T-T_{\rm o}-T\ln(T/T_{\rm o})].$ $C_{P{\rm ord}}^{\rm o}$ is the standard heat capacity effect of Reaction (25) at $P_{\rm o}$ and $T_{\rm o}$, and the $W{\rm s}$ are the Margules parameters.

Equation (27) is derived from Eqs. (A14), (A21) and (A22) using q = 2, m = 1 and $X_{Gas} + X_{Lig} = 1$.