The solubility of quartz in H₂O in the lower crust and upper mantle

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Abstract—The solubility of quartz in H₂O has been determined experimentally from 5 to 20 kb and 500 to 900°C. The results double the pressure range over which the molality of aqueous silica (msio₂) has been determined and lead to more accurate estimates of quartz solubility in H₂O below 5 kb because of the rapid-quench methods employed. At constant temperature, log msio₂ increases with increasing pressure and (δ log msio₂/δP) T decreases with increasing pressure. Comparison of the new data with previous low-pressure experiments demonstrates that isothermal values of log msio₂ increase linearly with increasing log pH₂O between 200 and 900°C. This observation was used to derive the following expression for the equilibrium constant (K) of the reaction quartz = SiO₂aq:

\[ \log K = 4.2620 - \frac{5764.2}{T} + \frac{1.7513 \times 10^6}{T^2} - \frac{2.2869 \times 10^8}{T^3} + \left[ 7.8454 \right] \log m_{H_2O}, \]

where \( \log K = \log m_{SiO_2} \). The equation agrees well with previous results, while accurately reproducing measured quartz solubilities over a much wider range in pressure and temperature, from 25°C and 1 bar to the conditions of this study. If the isothermal variation of log m_{SiO_2} with log p_{H_2O} is assumed to be linear, the results can be extrapolated to >20 kb. The equation allows evaluation of aqueous silica transport in Barrovan metamorphic belts, subduction zones, and metasomatized magma source-regions in the mantle.

INTRODUCTION

The solubility of quartz in aqueous solutions is of fundamental importance to geochemistry and petrology. Concentrations of aqueous Si in equilibrium with quartz increase with pressure and temperature from ~0.1 wt% in pure H₂O at conditions of low-grade metamorphism and geothermal systems to >5 wt% at conditions of high-grade metamorphism in the middle crust (Kennedy, 1950; Morey and Hesselgesser, 1951; Wyart and Sabatier, 1955; Khitarov, 1956; Kitahara, 1960; Van Lier et al., 1960; Morey et al., 1962; Siever, 1962; Weil and Fye, 1964; Anderson and Burnham, 1965, 1967; Heitmann, 1965; Crerar and Anderson, 1971; Hemley et al., 1980; Walter and Orville, 1983; Rimstidt, 1984). The magnitude and variation of quartz solubility in H₂O results in a strong potential for local to large-scale redistribution of Si during fluid-rock interaction in diverse geologic environments (e.g., Wood and Walter, 1986).

Because quartz solubility increases with pressure at a given temperature, Si metasomatism is particularly important in metamorphic and igneous processes at high pressures. However, virtually all experimental studies of quartz solubility have been carried out below ~4 kb. The seminal study of Anderson and Burnham (1965) has provided the only data with which to evaluate the pressure and temperature dependence of quartz solubility in H₂O up to ~10 kb, but solubility constraints at higher pressure are required to assess Si mass transfer by H₂O during fluid-rock interaction in, for example, subduction zones or magma source regions in the mantle. In addition, slow experimental quench times (8–10 min) may have led Anderson and Burnham (1965) to underestimate quartz solubilities at some conditions below 10 kb. Thus, accurate evaluation of the scale and magnitude of aqueous Si transport during fluid-rock interaction in mid-crustal to upper mantle environments requires new measurements of Si concentrations in H₂O in equilibrium with quartz at high pressures and temperatures.

Here, I present results of rapid-quench experiments on quartz solubility in H₂O at 500–900°C and 5–20 kb. The new determinations double the pressure range over which quartz solubility has been measured. Moreover, although quartz solubility is a complex function of solvent composition (e.g., Sommerfield, 1967; Shetkel, 1974; Novgorodov, 1975; Hemley et al., 1980; Fournier et al., 1982; Walter and Orville, 1983; Saccocia and Seyfried, 1990; Xie and Walter, 1993), results in the simple system SiO₂–H₂O constrain the thermodynamic properties of aqueous silica needed for model phase equilibrium and transport calculations to upper-mantle pressures. This can be seen by writing the equilibrium between aqueous SiO₂ and quartz as

\[ \text{SiO}_2^{\text{(quartz)}} = \text{SiO}_2^{\text{(aq)}} \]  

where \( \text{SiO}_2^{\text{(aq)}} \) refers to solvated aqueous silica independent of hydration state (Walter and Helgeson, 1977). Adopting standard states of unit activity (\( a \)) for pure quartz and H₂O at any pressure and temperature and unit activity for SiO₂ in a hypothetical one molal solution referenced to infinite dilution at any pressure and temperature, the mass-action relation for Eqn. 1 reduces to

\[ \log K_{eq} = \log a_{\text{SiO}_2^{\text{(aq)}}} \]
where $K_{(1)}$ is the equilibrium constant for equilibrium 1. WALTHER and HELGESON (1977) and WALTHER and ORVILLE (1983) noted that aqueous silica exists as a neutral hydrated monomer species in pure H$_2$O, so the activity coefficient of SiO$_2$(aq) can be taken as unity, leading to
\[
\log K_{(1)} = \log m_{\text{SiO}_2\text{(aq)}}.
\]

and
\[
\Delta G_{\text{SiO}_2\text{(aq)}}^0 = \Delta G_{\text{quart}}^0 - 2.303 RT \log m_{\text{SiO}_2\text{(aq)}},
\]

where $m$ is molality, $\Delta G^0$ is the change in the apparent standard molar Gibbs free energy at constant pressure and temperature, $R$ is the gas constant, and $T$ is temperature in Kelvin. Thus, when combined with Eqns. 1-4 and appropriate derivatives, the new solubility measurements allow prediction of concentrations and thermodynamic properties of SiO$_2$(aq) in pure H$_2$O over a wider range of conditions than has previously been possible: from the Earth's surface to its upper mantle.

### Experimental Methods

Two types of solubility experiments were conducted (Table 1). Initial experiments involved double encapsulation of crushed quartz grains following the methods of ANDERSON and BURNHAM (1965).
These experiments are referred to as crushed-quartz experiments. Modifications of the capsule and furnace design (MANNING and BOETTCHER, 1994) allowed solubilities to be determined from single crystals. Experiments employing the new techniques are referred to as single-crystal experiments.

All experiments used inclusion-free Brazilian quartz. For crushed-quartz experiments, a large crystal was crushed and sieved. Grains from 0.35 to 0.42 mm in size were washed in H2O in an ultrasonic bath and inspected for adhering ultrafines by optical and scanning-electron microscopy (Fig. 1a). In each crushed-quartz experiment, 20–40 μL distilled deionized H2O and a crimped Ag0Pd0.1 1.6 mm O.D. capsule with ~10 mg quartz were placed in 3 mm O.D. Pt capsules. The outer Pt capsule was then carefully crimped, weighed, sealed by arc welding, and weighed again. The replicate weightings identified water loss during each step. ANDERSON and BURNHAM (1965) used a perforated inner capsule. Exploratory experiments showed that this did not influence measured solubilities (Table 1), so inner capsules were left unperforated.

For single-crystal experiments, 3–4 mm thick slabs were cut normal to c from a large, subhedral crystal of the same Brazilian quartz used in the crushed-quartz experiments. Cylinders weighing 5–20 mg were drilled parallel to c with a 1.8 mm I.D. diamond coring bit (Fig. 1b). The microcores were cleaned with dilute HNO3, placed in 2 mm O.D. Au capsules, and then loaded in a 5 mm O.D. Pt capsule together with 50–75 μL H2O. Though not required for the measurement of solubility, the inner capsule prevented the microcore from breaking if it wedged into a corner of the outer capsule during an experiment. An end-loaded piston-cylinder apparatus with 2.54 cm-diameter pistons and NaCl-graphite furnace assemblies was used for all experiments (BOHLEN, 1984; MANNING and BOETTCHER, 1994). Pressures were measured using Heise gauges calibrated against the melting of alkali halides (BOHLEN, 1984) and albite-jadeite-quartz equilibrium at 600°C (HAYS and BELL, 1973; MANNING and BOETTCHER, 1994). Pressure and temperature were held within 10°C of values listed in Table 1. All experiments employed Pt-PtRh10 thermocouples resting on the capsules throughout the run. Capsules for crushed-quartz experiments were placed horizontally in the furnace assembly. For this geometry, the temperature gradients in the sample were <5°C (BOHLEN, 1984). Capsules used in single-crystal experiments had vertical lengths of 4–5 mm, but temperature gradients were similar to those in crushed-quartz experiments because thick graphite heaters with long hotspots were employed (MANNING and BOETTCHER, 1994). Since the magnitude of the effects of pressure, pressure gradients, and temperature gradients on emf are poorly known, uncorrected temperatures read from thermocouples are reported here. Note that following this standard convention likely introduces uncertainty of no more than 2.5–10°C at the conditions of the experiments (GETTING and KENNEDY, 1970; CHENG et al., 1975).

Experiments were quenched to ~100°C in ~30 s by cutting power to the apparatus. Pressure decreased to ~20–25% of run pressure during the temperature drop and was then lowered to 1 atm over several minutes. Total time from quench to quartz weighing was typically, ~15 min. Final aqueous solutions ranged from clear to milky, evidently depending on the degree of formation of colloidal silica on quench. Dried material from fluid in the outer capsule was isotropic and had an index of refraction <1.52, consistent with vapor quench. Occasionally, up to three or four small quartz crystals could be found in this material. It is unclear whether they sifted out from the inner capsule or precipitated on quench, but their weight was undetectable and thus, too small to affect solubility determinations.

Quartz from the inner capsule was carefully inspected for adhering quench material using optical and scanning-electron microscopy. In crushed-quartz experiments (Fig. 1c), the crystals showed subhedral terminations and a wider range in grain size than the starting material, suggesting substantial dissolution and reprecipitation during the experiments. This is consistent with an abundance of two-phase fluid inclusions in the product grains. In single-crystal experiments, optical inspection readily revealed any adhering particles or signs of breakage. In some cases, well-developed growth facets formed at one or both ends of the cylinder (Fig. 1d).

Quartz solubility was determined from the weight loss of quartz grains by

\[ m_{SiO_2\,soln} = \frac{\left( w_{dry\,quench} - w_{dry\,in} \right) \times 1000}{60.048 \times w_{H_2O}}, \]

where \( w_{dry\,quench} \) and \( w_{dry\,in} \) are the initial and final weights of quartz, the number 60.048 is the molecular weight of SiO2, and \( w_{H_2O} \) is the weight of H2O in the experiment. ANDERSON and BURNHAM (1965) noted that SiO2 precipitates from the solution trapped in the inner capsule in crushed-quartz experiments. This problem was eliminated by subtracting from \( w_{dry\,quench} \) the weight of the trapped solution as measured by the difference between wet and dry weighings. Trapped solution averaged 5 wt% of the fluid.

Run durations required for equilibration were assessed at 700°C. Measured solubilities for experiments lasting from 15.75 to 90 h do not vary with time (Fig. 2). Accordingly, all experiments at ~700°C were 2–16 h in duration. ANDERSON and BURNHAM (1965) reported that 50 to 60 h were required for equilibrium at 500°C. These were taken as minimum durations for experiments at 500 and 600°C.

**Fig. 2.** Log \( m_{SiO_2\,soln} \) vs. experiment duration at 700°C and 8.8 kbar. Solid line corresponds to the mean of the six experiments (log \( m_{SiO_2\,soln} = -0.181 \)) and error bars are 2σ relative to mean (±0.016).
WALther and ORVille (1983) showed that, at low pressures, experiments approaching equilibrium from both undersaturation and supersaturation gave results consistent with previous experiments in which equilibrium was approached from undersaturation. In this study, all but one of the experiments approached equilibrium from undersaturation.

RESULTS

Table 1 and Fig. 3 give experimentally determined quartz solubilities in H$_2$O along isotherms between 500 and 900°C. The same type of apparatus was used for all experiments and all solubilities were determined by weight loss, so it can be assumed that sources of random error are similar for all measurements. Uncertainties can therefore be established by pooling repeated experiments through the equation (e.g., SKOOG et al., 1994), where $\sigma$ is the sample standard deviation, $m_n$ is the $n$th solubility measurement at a given $P$ and $T$, and $N_1$ and $N_2$ are the mean measured solubility and number of experiments at the first $P$ and $T$, and so on for $N_i$ distinct experimental conditions. Nine sets of two to six replicates (Table 1) yield a pooled sample standard deviation of 0.020 mdat. Uncertainty levels in Fig. 3 are 2$\sigma$ from pooled replicate experiments (see text). Results of ANDERSON and BURNHAM (1965) are shown for comparison. Fill patterns vary to allow distinction of isothermal datasets.

Figure 3 shows that the solubility of quartz in H$_2$O increases with increasing pressure along each isotherm. Compared with the results of ANDERSON and BURNHAM (1965), the present study illustrates that $\left(\frac{\partial \log m_{SiO_2}}{\partial P}\right)_T$ is greatest below 5 kb and decreases strongly with increasing pressure. The results also confirm that, at constant pressure, quartz solubility increases with temperature at high pressures.

The solubilities reported by ANDERSON and BURNHAM (1965) are similar to those measured here at 500°C, but their results are systematically near the lower limit of my 2$\sigma$ uncertainties at $\leq$600°C (Fig. 3). The most likely explanation is the different quench times used in each study. ANDERSON and BURNHAM (1965) used a gas apparatus and reported quench times of 8–10 min from the experimental temperature to $<100°C$. Similar temperature drops required $\approx$30 s in the piston-cylinder apparatus used here. I investigated the effect of these different quench times at 700°C and 8.8 kb. After sufficient time for equilibration, temperatures were decreased linearly to 100°C over 0.5 to $\sim$20 min. Figure 4 illustrates that apparent solubility increases with decreasing quench times. Because apparent solubility decreases most strongly between 0 and 2 min quench times (Fig. 4), rapid quenching is required for accurate solubility determinations where concentrations are high. Aware of this problem, ANDERSON and BURNHAM (1965) attributed decreasing precision at high solubility (Fig. 3) to reprecipitation of quartz on quench. Figures 3 and 4 imply that their solubilities may be too low by up to 10% at $\geq$600°C.

DISCUSSION

Equation for Quartz Solubility to High Pressures

Though accurate for most mid- to upper-crustal conditions, previous equations for computing the solubility of quartz in pure H$_2$O can not be used at the higher pressures of the present study. WALTHER and HELGESON (1977) predicted quartz solubilities to 550°C and 5 kb. McKENZIE and HELGESON (1984) extended these results to 900°C at 2 kb. Subsequent modifications to the Helgeson-Kirkham-Flowers (HKF: HELGESON et al., 1981) equation of state for aqueous species (TANGER and HELGESON, 1988) allowed SHOCK et al. (1989) to improve and extend the predicted thermodynamic properties of SiO$_2$$_{(aq)}$ to 1000°C at $\leq$5 kb. The HKF model has the advantage that its parameters describe specific solvation
and nonsolvation contributions to thermodynamic properties of aqueous species. However, the solvation contributions depend in part on the static dielectric constant of H$_2$O (e$_{\text{H}_2\text{O}}$) and its derivatives. Despite apparently reliable estimates of e$_{\text{H}_2\text{O}}$ at water densities below 1.0 g/cm$^3$ (e.g., Pitzer, 1983; Walther, 1991), the lack of experimental data or adequate predictive models for e$_{\text{H}_2\text{O}}$ at higher densities currently limits use of the HKF model to ≤ 5 kb. Fournier and Potter (1982) described quartz solubility to 10 kb using a correlation equation based on temperature and the specific volume of H$_2$O, for which experimental data exist within the fit range. However, their equation is strongly weighted by the data of Anderson and Burnham (1965). Moreover, extrapolations beyond ~10 kb do not reproduce the solubilities measured in the present study.

At constant temperature, values of $\log K$ for the ionization of H$_2$O and a variety of other aqueous species are linear in the logarithm of the density (p) of H$_2$O over a range in pressure (Franck, 1956, 1961; Quist, 1970; Marshall and Quist, 1967; Marshall, 1970, 1972; Sweeton et al., 1974; Marshall and Franck, 1981; Marshall and Mesmer, 1984; Mesmer et al., 1988, 1989). As noted by Walther (1991), this linear behavior is probably a consequence of the linear correlations between temperature and 1/$\rho_{\text{H}_2\text{O}}$ at constant water density. Marshall and Franck (1981) showed that the ionization of H$_2$O to 1000°C and 10 kb could be parameterized by an equation of the form

$$\log K = A + \frac{B}{T^2} + \frac{C}{T^3} + \frac{D}{T^4} + \left[ \frac{E + F + G}{T} \right] \log \rho_{\text{H}_2\text{O}}.$$  (7)

where the terms $A$ through $G$ are regressed constants. This equation has two parts: a third-order polynomial fit to $(\partial \log K/\partial (1/T))$ at $\rho_{\text{H}_2\text{O}} = 1.0$ g/cm$^3$ (terms A-D), and a second-order polynomial fit to $(\partial \log K/\partial \log \rho_{\text{H}_2\text{O}})_T$ vs. 1/$T$ (terms F-G). Equation 7, or variants, have been explored for geologic conditions by Eugster and Baumgartner (1987) and Anderson et al. (1991). Its dependence only on temperature and $\rho_{\text{H}_2\text{O}}$ makes Eqn. 7 ideal for calculating thermodynamic properties of aqueous species at high pressures, provided that $\rho_{\text{H}_2\text{O}}$ is known and that linear dependence of $\log K$ on $\log \rho_{\text{H}_2\text{O}}$ can be demonstrated.

Use of Eqn. 7 to predict quartz solubilities from the surface of the Earth to the upper mantle requires accurate calculation of the density of H$_2$O from 1 bar to >100 kb and 25 to ≥1000°C, such that density is a continuous function of temperature and pressure. Many equations of state and molecular-dynamics simulations compute the thermodynamic properties of H$_2$O for different geologic conditions (Rice and Walsh, 1957; Helfgott and Kirkham, 1974; Holloway, 1977; Delany and Helgeson, 1978; Kerrick and Jacobs, 1981; Halbach and Chatterjee, 1982; Haar et al., 1984; Saxena and Fei, 1987a,b; Saul and Wagner, 1989; Brodholt and Wood, 1990; Hill, 1990; Stixrude and Bukowinski, 1990; Belonoshko and Saxena, 1991; Holland and Powell, 1991). But, as noted by Holland and Powell (1991), most available equations are not applicable over the wide pressure-temperature range desired. Exceptions are the equations of Halbach and Chatterjee (1982), Saul and Wagner (1989), and Holland and Powell (1991). The empirical Redlich-Kwong formulation of Halbach and Chatterjee (1982) was used in the present study to calculate $\rho_{\text{H}_2\text{O}}$ because it accurately represents the properties of water to 50 kb and 1000°C (e.g., Rice and Walsh, 1957; Bulakh, 1979), it can be extrapolated to higher pressures, and it is simple in form.

The many investigations of the solubility of quartz in H$_2$O provide a set of experimental constraints that range widely not only in pressure and temperature, but also in accuracy and precision. Nevertheless, as illustrated in Fig. 5, the variation in $\log m_{\text{H}_2\text{O}(q)}$ with $\log \rho_{\text{H}_2\text{O}}$ (Halbach and Chatterjee, 1982) along isotherms from selected experiments is remarkably linear. Previous studies also noted this behavior over more limited density intervals (e.g., Mosebach, 1957). Although Fournier and Potter (1982) suggested that isotherms become nonlinear at high temperatures below the critical density ($\rho_{\text{H}_2\text{O}} < 0.322$ g/cm$^3$), Fig. 5 implies that the assumption of linear behavior for most geologic conditions is justified.

In the regression analysis, I combined the new measurements with the results of Hemley et al. (1980) and Walther and Orville (1983) to ensure that experiments of both high and low quality were not averaged. These studies were used for two reasons. First, they avoided quench problems through rapid quenching (Hemley et al., 1980) or solution extraction at pressure and temperature (Walther and Orville, 1983). Also, the two studies could be combined to generate isothermal solubility measurements from the boiling curve of H$_2$O to 2 kb at 200–550°C needed for the fit procedure. To constrain Eqn. 7 below 200°C, solubilities were calculated to 5 kb along isotherms of 25, 50, 100, and 150°C from Fournier and Potter (1982).

Regression of the experimental data gave

$$\log K(1) = 4.2620 - \frac{5764.2}{T} + \frac{1.7513 \times 10^6}{T^2}$$

$$- \frac{2.2869 \times 10^8}{T^1} + \left[ \frac{2.8454 - 1006.9}{T} + \frac{3.5689 \times 10^7}{T^2} \right] \log \rho_{\text{H}_2\text{O}}.$$  (8)

The correlation coefficient ($R$) for the polynomial describing the variation in $\log K(1)$, with 1/T at $\rho_{\text{H}_2\text{O}} = 1.0$ gm/cm$^3$ is 0.9989. A lower value of $R = 0.6942$ was obtained for the fit to $(\partial \log K/\partial \log \rho_{\text{H}_2\text{O}})_T$ as a function of 1/T. This is because of fewer data from Hemley et al. (1980) and Walther and Orville (1983) along some isotherms, which results in lower precision of calculated changes in the slopes of isotherms with temperature. However, the low $R$ does not appear to affect significantly the accuracy of predicted solubilities.

**Comparison to Previous Work**

Figure 5 compares calculated solubilities (Eqn. 8) with selected experiments. It is evident in Fig. 5 that there are no systematic deviations between calculated solubilities and those experiments used in the derivation of Eqn. 8. Also, predicted solubilities are somewhat greater than those of Anderson and Burnham (1965), consistent with the inference of minor reprecipitation on quench in their experiments.
Equation 8 predicts that $\log K_{11} = -4.003$ at 25°C and 1 bar. This agrees well with the values of $-4.001$ of MOREY et al. (1962) and $-3.992$ of SUPCRT92 (JOHNSON et al., 1992). The degree to which Eqn. 8 reproduces other experimental data at elevated pressures and temperatures can be assessed using average differences, $\Delta$, between experimental and calculated solubilities, where $\Delta = \log m_{\text{experimental}} - \log m_{\text{calculated}}$. The sign of $\Delta$ signifies whether, on average, calculated solubilities underestimate (positive $\Delta$) or overestimate (negative $\Delta$) those measured; the average of the absolute values of $\Delta$, $|\Delta|$, is a measure of the precision of the calculated values relative to the experimental values. Table 2 compares values of $\Delta$ and $|\Delta|$ generated using equations of FOURNIER and POTTER (1982), the data of SHOCK et al. (1989) in SUPCRT92 (JOHNSON et al., 1993), and Eqn. 8. Calculations were confined to experiments within the stated pressure and temperature limits of each study: $\leq 5$ kbar in the case of SHOCK et al. (1989) and $\leq 10$ kbar in the case of FOURNIER and POTTER (1982). Values in Table 2 differ slightly from those given by FOURNIER and POTTER (1982), because of different equations of state for H$_2$O. The same magnitudes of $|\Delta|$ for all three studies imply a similar level of precision for each. Values of $\Delta$ vary somewhat between the three studies, indicating that predicted quartz solubilities are optimized at different pressures and temperatures. There is good agreement between Eqn. 8 and the results of KENNEDY (1950), MOREY and HESSELGESSER (1951), WYART and SABATIER (1955), WEILL and FYFE (1964), and CRERAR and ANDERSON (1971). Solubilities measured by KHI TAROV (1956), KITAHARA (1960), and ANDERSON and BURNHAM (1965) are systematically low relative to Eqn. 8, probably because of quench problems. VAN LIER et al. (1960), MOREY et al. (1962), and SIEVER (1962) measured higher solubilities than those predicted by Eqn. 8. FOURNIER and POTTER (1982), who also underestimated these solubilities, suggested that the higher surface energies of finely ground quartz used in these experiments caused fluid supersaturation with respect to quartz. The results of RIMSTIDT (1984) are also greater than those predicted by Eqn. 8, as well as by FOURNIER and POTTER (1982) and SHOCK et al. (1989). Because ultrafine problems were carefully avoided, it is unclear why his results are inconsistent with all three equations for quartz solubility.

Figure 6 illustrates graphically the success of Eqn. 8 in predicting quartz solubilities from liquid-vapor saturation to 2 kb over a range in temperatures. With the exception of the data of HEMLEY et al. (1980) and WAL ther and ORVILLE (1983), none of the additional ~200 solubility determinations by other workers were used to derive Eqn. 8. Of particular note is the sharp predicted decrease in quartz solubility within several °C of the critical point (Fig. 6a), a feature that has previously proved difficult to reproduce (e.g., WAL ther and HELGESON, 1977; SHOCK et al., 1989).

![Graph](image)

**Table 2. Comparison of calculated and measured quartz solubilities in H$_2$O**

| Used to derive (eq. 8) | P range (kb) | T range (°C) | $\Delta^*$ | $|\Delta|^*$ | $\Delta$ | $|\Delta|$ | $\Delta$ | $|\Delta|$ |
|-----------------------|-------------|-------------|------------|-------------|----------|----------|----------|----------|
| This Study            | 5.20        | 500-900     | -0.010     | 0.031       | 0.033    | 0.040    | 0.031    | 0.043    |
| Hemley et al. (1980)  | 1-2         | 350-550     | -0.011     | 0.013       | 0.014    | 0.022    | 0.012    | 0.036    |
| Walther & Orville (1983) | 1-2       | 200-500     | -0.010     | 0.031       | 0.033    | 0.040    | 0.031    | 0.043    |
| Independent of (eq. 8) |             |             |            |             |          |          |          |          |
| Anderson & Burnham (1965) | 1-9        | 500-900     | -0.033     | 0.042       | -0.012   | 0.029    | -0.059   | 0.059    |
| Creer & Anderson (1971) | 1-7         | 179-329     | -0.051     | 0.054       | 0.013    | 0.034    | 0.006    | 0.023    |
| Kennedy (1950)         | 1-7.5       | 160-610     | -0.059     | 0.069       | 0.005    | 0.066    | 0.038    | 0.070    |
| Khaturov (1956)        | 1-4         | 400         | -0.038     | 0.038       | -0.057   | 0.057    | -0.026   | 0.026    |
| Kitahara (1960)        | 1-4         | 140-370     | -0.095     | 0.102       | -0.060   | 0.127    | -0.057   | 0.098    |
| Morey & Hesselfeisser (1951) | 1-2   | 500-600     | -0.017     | 0.085       | 0.033    | 0.086    | 0.006    | 0.072    |
| Morey & Hesselfeisser (1951) | 0.5-2     | 500-600     | -0.017     | 0.085       | 0.033    | 0.086    | 0.006    | 0.072    |
| Morey et al. (1962)    | 1           | 25-300      | 0.057      | 0.077       | 0.032    | 0.054    | 0.062    | 0.083    |
| Rimstidt (1984)        | 1           | 50-96       | 0.126      | 0.126       | 0.092    | 0.092    | 0.080    | 0.080    |
| Sierer (1962)          | 1           | 125-182     | 0.094      | 0.094       | 0.110    | 0.110    | 0.084    | 0.084    |
| Van Lier et al. (1960) | 1           | 70-100      | 0.121      | 0.121       | 0.121    | 0.121    | 0.107    | 0.107    |
| Weill & Frye (1964)    | 1-4         | 400-550     | -0.005     | 0.013       | 0.029    | 0.051    | -0.015   | 0.020    |
| Wyart & Sabatier (1955) | 0.5-2      | 400         | -0.051     | 0.051       | -0.017   | 0.019    | -0.037   | 0.037    |

* $\Delta = $ average difference between experimental and calculated log $m_{\text{SiO}_2(aq)}$ (see text).

** $|\Delta| = $ average of the absolute value of the difference between experimental and calculated log $m_{\text{SiO}_2(aq)}$ (see text).

† $P_{\text{sat}}$ data only (see FOURNIER and POTTER, 1982).
Aqueous solubility of quartz at high pressure

The System SiO₂-H₂O to High Pressures

The observation that log $K_{(1)}$ is linear with respect to log $\rho_{H₂O}$ to 20 kb provides the basis for extrapolation of the results beyond 20 kb. At constant temperature, log $\rho_{H₂O}$ increases only slightly with increasing pressure at high pressures. Thus, application of Eqn. 8 to higher pressures probably will not lead to substantial errors in predicted log $K_{(1)}$, even if future experiments demonstrate that it is not linear in log $\rho_{H₂O}$ above 20 kb.

Figure 7 shows phase relations in the system SiO₂-H₂O assuming saturation with respect to the stable SiO₂ polymorph and linear extrapolation of log $K_{(1)}$ with log $\rho_{H₂O}$ above 20 kb. Polymorphic transformations of SiO₂ are superimposed on the diagram from data of Berman (1988), Mirwald and Massone (1980), and Bohlen and Boettcher (1982). Values of log $m_{SiO₂(aw)}$ along the H₂O-saturated melting curve of SiO₂ are from Kennedy et al. (1962), whereas those in equilibrium with H₂O ices were derived from Eqn. 8, using $\rho_{H₂O}$ calculated along the stable phase boundaries reported by Bridgman (1937) and Pistorius et al. (1963). Figure 7 illustrates phase relations from 25°C to water-saturated melting and from 1 bar to > 50 kb. As noted above, log $K_{(1)}$ varies linearly with log $\rho_{H₂O}$ along isotherms. In contrast, isobaric variation in log $K_{(1)}$ is not linear. Log $K_{(1)}$ increases with temperature along low-pressure isobars (< 1 kb) to a maximum between 300 and 350°C and then decreases, producing the well-known "reverse-solubility" behavior of quartz (Fig. 6a). At higher pressures, log $K_{(1)}$ increases with temperature at constant pressure and $\partial^2 \log K_{(1)}/\partial (\log \rho_{H₂O})^2$ decreases. Also, Fig. 7 shows that $\partial^2 \log K_{(1)}/\partial (\log \rho_{H₂O})^2$ decreases significantly with increasing pressure.

Combining Eqns. 4 and 8 with $\Delta G_{aq}$ (Berman, 1988) allows prediction of the solubility of coesite in H₂O. Figure 7 illustrates that log $m_{SiO₂(aw)}$ in H₂O in equilibrium with coesite decreases with increasing water density below ~800°C.

Kennedy et al. (1962) determined a critical endpoint in the system SiO₂-H₂O at 1080°C and 9.7 kb. Because this defines complete miscibility in the system, log $m_{SiO₂(aw)}$ should approach infinity at this pressure and temperature. Thus, the slopes of isotherms should approach infinity near the temperature of the endpoint. Although calculated slopes of isotherms increase slightly above 500°C (Fig. 7) and are accurate

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**Fig. 6.** Log $m_{SiO₂}$ vs. temperature along steam saturation curve and selected isobars. Published experimental solubility determinations shown in squares (Kennedy, 1950; Morey and Hesselsgesser, 1951; Wyart and Sabatier, 1955; Kitahara, 1960; Van Lier et al., 1960; Morey et al., 1962; Siever, 1962; Weill and Pyfe, 1964; Anderson and Burnett, 1963; Creem and Anderson, 1971; Hemley et al., 1980; Walton and Orville, 1983; Rimstidt, 1984). Solid curves predicted from Eqn. 8.

**Fig. 7.** Phase diagram for the system SiO₂-H₂O showing wt% SiO₂ and log $m_{SiO₂}$ in H₂O in equilibrium with the stable SiO₂ polymorph at any given pressure and temperature. Phase boundaries (heavy lines) superimposed on diagram from Bridgman (1937), Kennedy et al. (1962), Pistorius et al. (1963), and Berman (1988). Light dashed and solid curves are respectively isotherms and isobars calculated from Eqn. 8.
at the temperatures and pressures investigated experimentally, Eqn. 8 does not reproduce this critical behavior. Experiments on quartz solubility at pressures and temperatures near the critical endpoint would allow refinement of Eqn. 8 to account for the results of KENNEDY et al. (1962).

Figure 7 provides a basis for comparing aqueous SiO₂ concentrations from different geologic environments. For example, the formation of eclogite from hydrous metabasaltic assemblages in subduction zones liberates water that may play an important role in metasomatism and melt generation in the mantle wedge. These reactions proceed at or near quartz saturation over a range in pressure and temperature of ~12-15 kb and 500-600°C (e.g., PEACOCK, 1993). Figure 7 shows that log \( n_{SiO2} \) in aqueous pore fluids in equilibrium with quartz at these conditions is ~0.7 to ~0.2, or 0.3 to 0.6 mol/ kg. By comparison, peak metamorphic conditions in Barrovian metamorphic belts may reach 600-700°C and 6-8 kb. Si concentrations in aqueous pore fluids in equilibrium with quartz in this environment would be 0.4-0.6 mol/kg (Fig. 7). The capacity for silica transport in and above subducting oceanic crust is therefore broadly similar to that in the deeper portions of orogenic belts, assuming pure H₂O. Thus, use of Fig. 7 and Eqn. 8 allows simple comparisons as these are for a wide range of geologic settings, as well as more detailed analyses of Si mass transfer in previously inaccessible metamorphic environments. Moreover, the single equation describing log \( K_1 \) over a wide range of pressures and temperatures ensures internal consistency for comparing Si metasomatism between environments or along pressure-temperature paths.

CONCLUSIONS

New experiments on the solubility of quartz in H₂O provide the first systematic data on the thermodynamic properties of SiO₂ in H₂O between 10 and 20 kb. Because of the rapid-quench methods used, the results also lead to more accurate estimation of quartz solubilities at lower pressures. By combining these experiments with previous high-quality results, a simple equation correlating quartz solubility with the density of H₂O allows calculation of the concentrations of SiO₂ in aqueous fluids from the surface of the Earth to the upper mantle. These results can be used to model Si metasomatism in magma source regions in the mantle, in subducting slabs and the overlying mantle wedge, and in Barrovian metamorphic belts.

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