Solubility of Anhydrite, CaSO₄, in NaCl–H₂O Solutions at High Pressures and Temperatures: Applications to Fluid–Rock Interaction

ROBERT C. NEWTON AND CRAIG E. MANNING

DEPARTMENT OF EARTH AND SPACE SCIENCES, UNIVERSITY OF CALIFORNIA LOS ANGELES, LOS ANGELES, CA 90095-1567, USA

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INTRODUCTION

Sulfate minerals are common daughter crystals in saline fluid inclusions in rocks of deep-seated origin. Anhydrite occurs in fluid inclusions in minerals of carbonatites (Samson et al., 1995), alkaline igneous rocks (Hansteen & Burke, 1990; Belkin et al., 1997), Cu–Mo ores of magmatic origin (Roedder, 1984), eclogite veins in the Western Alps (Philippot & Selverstone, 1991), ultra-high-pressure coesite-bearing schists in the same region (Philippot et al., 1995), and in volatile-rich melt inclusions in xenoliths in mantle-derived basalts of alkaline affinity (McInnes & Cameron, 1994). Anhydrite also forms during sulfur-rich magmatism (Luhr et al., 1984; Bernard et al., 1991; Streck & Dilles, 1998; Barth & Dorais, 2000; Parat et al., 2002; Audétat et al., 2004).

An outstanding characteristic of these occurrences is a high oxidation state, as recorded in the high Fe₂O₃/FeO ratios of the volcanic rocks (McInnes & Cameron, 1994) and in the Fe₂O₃-rich ilmenites of the Pinatubo dacites (Hattori, 1993). The great range of valence states possible in sulfur, from ‑6 to ‑2, suggests that sulfur minerals may provide an important monitor of oxidation state and, in some circumstances, may even help to control oxygen fugacity.

Several well-studied granulate–facies terranes exhibit high oxidation states on a regional scale (Griffin et al., 1978; Currie & Gittins, 1988; Harlov et al., 1997). Harlov et al. (1997) ascribed intense metamorphic oxidation of
granulites of the Archean Shevaroy Hills, southern India, to the infiltration of oxidizing fluids, which left vein networks of pyrite, pyrrhotite and magnetite and caused grain-boundary alkali exchange in feldspars. They suggested migrating brines as alkali-exchanging and oxidizing agents. Widespread sulfate–carbonate scapolite in the Furua (Tanzania) granulite complex (Coolen, 1980) may indicate the action of sulfate-bearing metamorphic fluids. These rocks are also very oxidized, with Fe$_2$O$_3$-rich ilmenite present in most rocks and discrete hematite in a few. Cameron & Hattori (1994) suggested regional infiltration of SO$_2$-rich fluids of juvenile magmatic origin as a general cause of deep-crustal oxidation.

Disseminated anhydrite is common and sometimes abundant in the alteration zones of the Cu–Mo sulfide porphyry ore deposits of the western United States, and anhydrite is a common daughter mineral in the ultra-saline fluid inclusions in quartz associated with the highest-temperature, near-magmatic ore veins of these bodies (usually, roof zones of small granitic stocks: Roedder, 1971). The deposits are invariably highly oxidized, with ubiquitous dispersed hematite. Analogy with the Mt Pinatubo dacites suggests that widespread sulfate mineralization and high oxidation state are closely related phenomena. These sub-volcanic associations could have been influenced in their oxidation states by thermally driven downward circulation of surface waters (Norton, 1979), but several workers have emphasized that the S-rich volcanic products like those of Mt Pinatubo may result from exsolution of S-bearing gases from underlying primitive magmas (e.g. Luhr, 1990; Hattori, 1993; Hattori & Keith, 2001).

Large-scale transfer of oxidized sulfur by aqueous fluids flowing through rocks requires that sulfate must be at least moderately concentrated in the fluid. A major problem thus arises in that the solubility of anhydrite in H$_2$O at elevated $P$ and $T$ is very low. Morey & Hesselgesser (1950) found the solubility of anhydrite to be only 0.002 molal (m) at 1 kbar and 500°C. Blount & Dickson (1969) measured anhydrite solubility to 1 kbar and 450°C in pure H$_2$O and in NaCl solutions up to 26 wt % [NaCl mole fraction ($X_{NaCl}$) of 0-1]. They found substantial enhancement of solubility with salinity, about 100-fold at 26 wt % NaCl. Furthermore, the strong negative temperature coefficient of anhydrite in pure H$_2$O is moderated and becomes slightly positive above 300°C at the highest salinity investigated. Large increases might also be expected at high $P$, based on the solution behavior of calcite (Newton & Manning, 2002). This is because the high degree of ionization in alkali chloride solutions at $>5$ kbar ($H_2O$ densities $>0.7$ g/cm$^3$; Aranovich & Newton, 1996) may facilitate the formation of the solute ions CaCl$^+$ and NaSO$_4$, which could form by reaction of CaSO$_4$ and NaCl. Furthermore, the anhydrous join CaSO$_4$–NaCl shows a deep eutectic at 1 bar with CaSO$_4$ amounting to 37 wt % of the liquid (Bergman & Golubeva, 1953). Because concentrated brines at high $P$ and $T$ behave more like fused salts than aqueous solutions (Aranovich & Newton, 1996), the binary fused-salt eutectic may be expected to persist into H$_2$O-bearing systems. Determination of a high CaSO$_4$ carrying capacity of alkali chloride solutions over a range of $P$, $T$ and salinity provides insight into sulfidation, scapolitization and oxidation processes in the deeper parts of the crust and in the upper mantle.

**Experimental methods**

The starting material was large limpid natural anhydrite from the UCLA mineral collection (No. MS 1608). X-ray diffraction confirmed the identity of anhydrite with no extraneous reflections. The orthorhombic unit cell constants were: $a_0 = 0.6996(2)$ nm; $b_0 = 0.7002(2)$ nm; $c_0 = 0.6239(2)$ nm. These constants agree, within uncertainties, with the ASTM data (No. 37-1496) for synthetic anhydrite. Energy dispersive analyses indicated that
the only detectable departure from CaSO₄ chemistry was SrO < 0.5 wt %. The experimental procedure used in this study was nearly identical to that used by Newton & Manning (2002) for calcite solubility. Briefly, crystals of a natural anhydrite sample were heated in air at 400°C for 20 min to purge any volatiles. Anhydrite grains, weighed amounts of deionized H₂O and reagent NaCl were then sealed by arc-welding into Pt capsules of 3 mm diameter and 0.15 mm wall thickness. The anhydrite was encased in an inner Pt tube of 1.5 mm diameter and 2 mm length, lightly crimped at the ends and perforated with 6–10 small holes to allow access of fluids. The solubility at the highest temperatures and most concentrated NaCl solutions proved to be too high for this simple technique, and either larger unencapsulated crystals were used, or additional powdered anhydrite was added to the outer capsule to presaturate the fluid. The latter method assumes that finely divided anhydrite dissolves much faster than the large, Pt-encased single crystals—an assumption that appears to be justified by the solubility behavior.

All experiments were done with the 1.91 cm diameter piston–cylinder apparatus with NaCl pressure medium and graphite heater sleeve. The pressure chamber was highly polished WC with a no-tolerance piston fit. The Heise Bourdon tube pressure gauge was calibrated against a highly precise 1000 bar Heise gauge of 40 cm dia diameter. The two gauges read the same in the range 0–300 bar (equivalent to 0–9 kbar sample pressure) to within 1 bar, with no systematic discrepancy or hysteresis. Temperatures were controlled automatically and measured with calibrated matched pairs of W–3%Rh vs W–25%Rh thermocouples. A typical thermoelement was compared with a certified Pt vs Pt–10%Rh thermocouple in a test piston–cylinder assembly identical to those of the solubility experiments. The two thermocouple junctures were less than 0.2 mm apart, though mutually insulated. The test was made at 10 kbar and 600–800°C. The Pt–Rh thermocouple read consistently 1°C lower at 800°C than the W–Re thermocouple, which was the control element. Precision of measurements is conservatively estimated at ±300 bars sample pressure and ±3°C.

Solubility determinations were made from the weight losses of inner Pt capsules or unencapsulated anhydrite crystals. Weights were determined on a Mettler M3 microbalance at a precision of 2 μg (1σ) based on repeated weighings over the course of this study. Quenched and cleaned sample capsules generally showed negligible weight changes from the starting values, indicating no loss of material from the capsules during the experiments. The H₂O of the charges was reetermined by a puncture and drying technique, which confirmed the weighing-in values within 0.3%.

The drying weight loss was taken to be the H₂O content of the experiment.

Previous investigations of anhydrite solubility have not considered a possible effect of variable oxygen fugacity (fO₂). The measurements of Blount & Dickson (1969) were self-buffered, in the sense that the solute sulfate would have reacted with H₂O to form small amounts of H₂S, with the release of O₂, resulting in a rather high, but unknown, fO₂. In the present study, in addition to the unbuffered runs, we measured the solubility of anhydrite in initially pure H₂O at 10 kbar and 800°C using internal buffer capsules containing hematite + magnetite, NiO + Ni metal, or MnO₂ + H₂O (10–12 mg solid and 2–3 mg H₂O) enclosed in small welded Pt tube segments. The buffer capsules were sealed with encapsulated anhydrite crystals and about 36 mg H₂O in the outer Pt jackets. The buffered runs were made over periods of 5–40 h to test the degree of equilibration of the fluids with the internal buffers, and to investigate the variation in solubility with fO₂.

### EXPERIMENTAL RESULTS

Experimental results are given in Table 1. Measurable solubility was detected at all P, T and X_{NaCl} investigated. A single experiment (No. 14, Table 1) was performed to control for S loss to Pt encapsulating materials by loading an empty inner capsule along with anhydrite powder and H₂O. The empty inner capsule showed no detectable weight change, indicating that measured weight changes can be ascribed entirely to anhydrite dissolution.

Large weight losses in experiments on NaCl-rich compositions indicated very high solubilities. These experiments yielded complex textures of quench material along with the residual crystal(s). Halite–anhydrite intergrowth textures in the dried former fluid phase were examined with the polarizing microscope, scanning electron microscope and X-ray diffraction. A regular box-work of large, thin anhydrite quench-crystals and interstitial halite (Fig. 1a) shows that the anhydrite formed first in the quench, with subsequent co-precipitation of anhydrite and halite. X-ray diffractograms of the dried quench material showed only halite and anhydrite. D-spacings of the quench anhydrite were identical to those of the starting material. Residual starting crystals retrieved from NaCl-rich experiments were sometimes solution-rounded flat bars (Fig. 1b) which could be easily separated from the quench material for weighing, but often there was recrystallization into many smaller crystals, which were harder to identify and separate. For this reason, the majority of the NaCl-rich experiments used encapsulated single crystals together with external powdered anhydrite to presaturate the fluid phase.
### Table 1: Experimental results

<table>
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<th>Expt. number</th>
<th>Temperature (°C)</th>
<th>Pressure (kbar)</th>
<th>Duration (h)</th>
<th>Anhydrite (mg)</th>
<th>Capsule (mg)</th>
<th>NaCl (mg)</th>
<th>H$_2$O (mg)</th>
<th>$X_{\text{NaCl}}$</th>
<th>$X_{\text{H}_2\text{O}}$</th>
<th>Anhydrite out (mg)</th>
<th>Capsule out (mg)</th>
<th>Solubility (molal)</th>
<th>Notes</th>
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<td><strong>Unbuffered experiments</strong></td>
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<td>6</td>
<td>800</td>
<td>10</td>
<td>18</td>
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<td>69-175</td>
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<td>32-212</td>
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<td>32</td>
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<td>10</td>
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<td>38-454</td>
<td>0.000</td>
<td>38-567</td>
<td>71-951</td>
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<td>White rosettes (gypsum?) in inner capsule</td>
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<td>17</td>
<td>800</td>
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<td>20</td>
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<td>13-088</td>
<td>33-902</td>
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<td>33-700</td>
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<td>72-651</td>
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<td>16</td>
<td>4-525 + 2-709$^+$</td>
<td>74-496</td>
<td>10-994</td>
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<td>5</td>
<td>800</td>
<td>10</td>
<td>39</td>
<td>8.167</td>
<td>u</td>
<td>29-459</td>
<td>20-138</td>
<td>0.311</td>
<td>20-171</td>
<td>0.000</td>
<td>&gt;2.974</td>
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<tr>
<td>7</td>
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<td>20</td>
<td>20.540</td>
<td>u</td>
<td>33-955</td>
<td>24-411</td>
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<td>24-541</td>
<td>2-423</td>
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<tr>
<td>14</td>
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<td>10</td>
<td>38</td>
<td>4-526</td>
<td>31-951</td>
<td>0.000</td>
<td>34-469</td>
<td>0.000</td>
<td>33-535</td>
<td>31-955</td>
<td>Blank test for sulfidation of Pt</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

| Buffered experiments | | | | | | | | | | | | | | |
| 28 | 800 | 10 | 20 | 2.251 | 51-101 | 0.000 | 35-766 | 0.000 | 36-206 | 50-349 | 0.153 | NNO buffer; strong H$_2$S odor |
| 29 | 800 | 10 | 5 | 1.088 | 73-572 | 0.000 | 35-734 | 0.000 | 35-784 | 73-499 | 0.397 | 0.015 | HM buffer |
| 27 | 800 | 10 | 19 | 2.418 | 47-365 | 0.000 | 34-319 | 0.000 | 34-339 | 47-280 | 2.365 | 0.018–0.021 | HM buffer |
| 33 | 800 | 10 | 43 | 0.918 | 68-851 | 0.000 | 36-376 | 0.000 | 36-413 | 68-773 | 0.830 | 0.016–0.028 | HM buffer |
| 51 | 800 | 10 | 18 | 0.432 | 70-837 | 0.000 | 35-721 | 0.000 | 35-721 | 70-768 | 0.014 | MnO buffer |

Explanation: ‘in’ and ‘out’ refer respectively to weights before and after experiment. H$_2$O out was used in all solubility calculations (see text). ‘u’ signifies that anhydrite crystal was unencapsulated. Blank entries indicate that measurement could not be made. 1σ uncertainty in solubility determinations is 0.001 molal, based on propagated weighing errors of ±0.002 mg per weighing step; 1σ in $X_{\text{NaCl}}$ < 2.3 × 10$^{-4}$. HM, hematite–magnetite; MnO, Mn$_2$O$_3$–MnO$_2$ buffer; NNO, Ni–NiO. *First weight refers to anhydrite single crystal, second weight to anhydrite powder added to outer capsule. †Range in solubility due to minimum from capsule weight change and maximum from crystal weight change (see text).
Quenched NaCl was dissolved from the inner capsules by soaking in water at 90 °C for periods of 10 min followed by drying, until a capsule came to constant weight. This treatment did not remove the small amount of quench anhydrite from the inner capsule, so that the final weight gave a lower limit to the solubility. The inner capsule was cut open and the contents were transferred to a Pt weighing pan. The quench anhydrite was texturally distinct and could be separated with a needle, but this treatment usually resulted in the loss of a small amount of granular anhydrite. The weight of the residual crystals therefore returned an upper limit for solubility. The lower and upper limits so determined define a range within which there is uniform probability of finding the solubility of anhydrite.

Figure 2 plots the solubilities from the HM-buffered and unbuffered runs with initially pure H2O. The time required for equilibration with the buffers was at most 10 h. The run of 18 h with the MnO2 buffer (run 31, Table 1) gave the lowest solubility. The capsules of buffering material retained H2O without detectable weight loss and all members of the buffering assemblages were still present. The MnO2 buffer material was coarsely recrystallized to dominant pyrolusite and subsidiary bixbyite (Mn2O3). Extrapolation of low-T thermodynamic data for pyrolusite suggests that fO2 in these experiments was above HM.

Solubilities in experiments with initially pure H2O at 10 kbar are low, in the range 0.01–0.03 m, and increase slightly with T (Table 1). Runs at 800 °C and 10 kbar with HM and MnO2 buffers show a minor, probably real, decrease of solubility with increase of fO2, though the uncertainty limits of the measurements overlap. The NNO experiment (run 28, Table 1) showed greatly

Fig. 1. Back-scattered electron microscope images of quenched charges. (a) Portion of dried fluid phase from run at 800 °C, 10 kbar, \( X_{NaCl} = 0.3 \) (run 7, Table 1)—large-bladed anhydrite quench crystals in parallel arrays with interstitial halite and smaller anhydrite. Halite partially dissolved away in upper portion. (b) Residual anhydrite starting material (run 22, 800 °C, 10 kbar, \( X_{NaCl} = 0.2 \))—originally two angular cleavage fragments, now grown together and solution-rounded. The oval striations are solution-etched cleavage planes that define ‘contours’ on a domed surface. (c) Bead of quenched molten anhydrite from run 28 (800 °C, 10 kbar, initially pure H2O, Ni–NiO buffer). (d) Magnification of C, showing anhydrite quench-laths.
different behavior. The capsule emitted a strong H$_2$S odor when punctured. The inner capsules and the inside of the outer capsule were darkened, probably indicating reaction to PtS. The content of the inner sample capsule was a rounded bead (Fig. 1c) showing the imprint of machining striations from the Pt. Higher-magnification examination (Fig. 1d) showed the bead to be a mass of skeletal crystals; the anhydrite had been melted. The tabular habit of the anhydrite quench crystals was quite similar to that quenched from the NaCl-rich runs. The solubility in the NNO-buffered run was substantially higher than in the high-$fO_2$ experiments, although some of the apparently high solubility may result from loss of S to the Pt capsule.

Figure 3 shows results of 10 kbar NaCl-bearing experiments. Run durations of $>$60 h at 600°C to $>$20 h at 800°C were assumed to be adequate to ensure equilibrium in view of results in pure H$_2$O (Fig. 2), and the fact that Blunt & Dickson (1969) demonstrated attainment of equilibrium in 24 h at 400°C. The experiments show that anhydrite solubility increases significantly with $X_{NaCl}$ at all $T$. For example, anhydrite is quite insoluble in pure H$_2$O at 800°C and 10 kbar, but 30 mol % NaCl in the fluid raises the solubility by a factor of nearly 200.

This anhydrite-saturated solution has the composition 27 wt % CaSO$_4$, 42 wt % NaCl, 31 wt % H$_2$O.

The solubility data of the NaCl-present experiments from 6 to 14 kbar (Table 1) can be systematized by a general expression of the form $\log [m(CaSO_4) - m_0(CaSO_4)] = a + b \log X(\text{NaCl})$, where $a$ and $b$ are functions of $P$ and $T$ (in kbar and Kelvins, respectively) and $m_0$ is the molality in pure H$_2$O. An excellent fit to the data could be achieved by functions of the forms $a = a_1 + a_2 T$ and $b = b_1 + b_2 T + b_3 (P - 10)$. Regression of the experimental data gave the constants $a_1 = -1.533$, $a_2 = 0.00291$, $b_1 = 1.441$, $b_2 = 0.000160$, $b_3 = -0.0413$.

Figure 3 also shows regressed and extrapolated solubility curves.

**DISCUSSION**

The solubility behavior of anhydrite is analogous to that of calcite (Newton & Manning, 2002; Caciagli & Manning, 2003) in several respects. Both solutions show marked decrease of solubility with rising temperature at low pressure (Fig. 4), with vanishingly small solubilities at 1 kbar and $>$600°C. Solubility increases by several orders of magnitude with pressures to 10 kbar in this
temperature range, and the temperature coefficient of solubility becomes positive for both CaCO$_3$ and CaSO$_4$. Both oxysalts show great solubility enhancement in NaCl solutions, which undoubtedly follows from the high solubility of CaCl$_2$ as a reaction product.

Caciagli & Manning (2003) showed that aqueous CO$_2$ is the dominant carbon species when calcite dissolves at high pressure. By analogy, the neutral species H$_2$S and SO$_2$ are likely to be the dominant sulfur products of anhydrite dissolution in initially pure H$_2$O. These volatile species are generated by the reactions

$$\text{CaSO}_4 + 2\text{H}_2\text{O} = \text{H}_2\text{S} + \text{Ca}^{2+} + 2\text{OH}^- + 2\text{O}_2$$  \hspace{1cm} (1)

and

$$2\text{CaSO}_4 + 2\text{H}_2\text{O} = 2\text{SO}_4^{2-} + 2\text{Ca}^{2+} + 4\text{OH}^- + \text{O}_2.$$  \hspace{1cm} (2)

These reactions reveal that the solubility of anhydrite (i.e. the concentration of Ca$^{2+}$), as well as the H$_2$S/SO$_4^{2-}$ ratio in the solution, depends strongly on the oxygen fugacity.

Figure 5 shows equilibrium concentrations of solute species in the system CaSO$_4$–H$_2$O as calculated at 5 kbar and 800°C by simultaneous solution of mass action, mass balance, charge balance and activity coefficient (Davies, 1962) equations for the species shown. Only species present at >10$^{-6}$ molal were considered (except H$^+$). It can be seen that total Ca and S concentrations (i.e. anhydrite solubility) decrease with increasing f$_{O_2}$, as indicated by experimental results at higher P (Table 1). Figure 5 also shows a representation of the present 800°C, 10 kbar solubility data at a range of f$_{O_2}$. Translation of the anhydrite solubility curve from 5 to 10 kbar suggests that the unbuffered runs had f$_{O_2}$ above the H$_2$S–SO$_2$ equivalence point but below HM. The unbuffered measurements indicate that CaSO$_4$ is a strong oxidizing agent in a nominally closed system with H$_2$O.

The present pure H$_2$O data are not sufficient to define anhydrite solubility as a function of f$_{O_2}$ at all P and T studied. However, the enhancement effect of NaCl dwarfs any effect of f$_{O_2}$, as shown in Fig. 3. The 10 kbar isothermal solubility curves are approximately quadratic, implying a solvent dissolution reaction similar to

$$\text{CaSO}_4 + 2\text{NaCl} = \text{CaCl}_2 + \text{Na}_2\text{SO}_4,$$  \hspace{1cm} (3)

anhydrite.
Fig. 4. Comparison of the solubility of anhydrite at ≥600°C and ≥6 kbar (squares; this study) with previous results at 1 kbar and lower T (triangles; Morey & Hesselgesser, 1950; Blount, 1965; Blount & Dickson, 1969) at $X_{\text{NaCl}} = 0$ (a) and 0.1 (b). Partly filled squares indicate that value is minimum (see Fig. 3 caption). In (a), all lines are fitted by eye. The solubility behavior of CaSO$_4$ in pure H$_2$O is similar to that of CaCO$_3$ (circles) at 1 kbar (Walther & Long, 1986) and at 10 kbar (Caciagli & Manning, 2003). In (b), the continuous lines connecting data from the present study are from the empirical formula given in the text. Our extrapolated curve at 1 kbar and $X_{\text{NaCl}} = 0.1$ (dashed line) agrees well with the 1 kbar data of Blount (1965) and Blount & Dickson (1969) at lower temperatures (fitted by eye).
The salts on the right-hand side are very soluble, which tends to drive the reaction to the right. The above reaction is analogous to that deduced by Newton & Manning (2002) for calcite solution in NaCl solutions, though the NaCl enhancement for CaSO$_4$ is five times as great as for CaCO$_3$ at 10 kbar. The temperature coefficient of solubility at constant $X_{\text{NaCl}}$ is very large, as for calcite. Figure 3 shows the predicted 10 kbar
solubility curves at 500 and 900°C. The solubility at 900°C is so large that the fluids are best regarded as supercritical hydrosaline melts or fused salt mixtures. For instance, an anhydrite-saturated fluid at 10 kbar, 900°C and $\Delta N_{NaCl} = 0.5$ is predicted to contain 38 wt % CaSO$_4$, 36 wt % NaCl and 26 wt % H$_2$O.

PETROLOGICAL APPLICATIONS

High-S volcanic eruptions

The andesite–dacite eruption of Mt Pinatubo, Philippines, in 1991, released great quantities of oxidized sulfur to the atmosphere. The finding of anhydrite in the dacite tephra ejected by this eruption (Bernard et al., 1991) has led to much discussion of the role of oxidized sulfur in subduction zone volcanism. Studies of anhydrite occurrence at Mt Pinatubo (Hattori, 1993) and in the tephra of the recent eruptions of El Chichón, Mexico (Luhr et al., 1984) and Nevado del Ruiz, Colombia (Melson et al., 1990) have emphasized their similarities, especially with respect to the high oxidation states shown by the tephra. Experimental studies on the stability of anhydrite in the crystallization of andesite (Luhr, 1990) and dacite (Scaliet et al., 1998) indicate that quite high oxygen fugacity, from 1 to 3 log units above the nickel–nickel oxide buffer, is required to stabilize anhydrite relative to sulfides at magmatic temperatures (750–1000°C) and upper-crustal pressures (1–4 kbar).

The source of the sulfur in high-S volcanism could be ophiolite sulfide deposits or anhydrite-bearing evaporite beds, either in the source regions of the magmas, or scavenged by the magmas in transit. The former source does not provide an oxidation mechanism and the latter source, proposed by Rye et al. (1984) for El Chichón on the basis of evaporite beds of Mesozoic age underlying this volcano, would not be appropriate for Mt Pinatubo, which has no underlying evaporite stratigraphy (Imai et al., 1996). The most probable source of sulfur in these eruptions was primitive mafic magmas which transferred sulfur to more evolved magmas and crustal melts (El Chichón: Luhr, 1990; Nevado del Ruiz: Melson et al., 1990; Pinatubo: Pallister et al., 1992). Pallister et al. (1992) postulated a “basalt trigger” for the eruption process.

Luhr et al. (1984) pointed to a mass-balance problem arising from the high-S contents of the El Chichón tephra: experiments by Carroll & Rutherford (1987) on S solubility in a basaltic magma at 2 kbar show a maximum of about 800 ppm, and this occurs at very high oxygen fugacity. According to Luhr et al. (1984), the >2 wt % of S of some of the El Chichón rocks could never have been dissolved in a magma of the rock composition, but must have been introduced by a fluid phase that interacted with the partially solidified magma and that deposited anhydrite. Hattori (1993) came to the same conclusion for the anhydrite-bearing dacite tephra of Mt Pinatubo. Those workers and Gerlach et al. (1996) envisioned a C–O–H–S magmatic gas as the agent of sulfur transfer. A fluid-transport mechanism for anhydrite emplacement in the Mt Pinatubo tephra was confirmed recently by scanning electron microscopy, which revealed etch pits and growth features diagnostic of fluid deposition (Jakubowski et al., 2002).

The present work suggests an alternative mechanism of fluid-mediated S transfer. The experiments of Klink & Burnham (1972) on Cl solubility in granitic melts and those of Webster et al. (1999) in basaltic and andesitic magmas show that, at pressure greater than near-surface conditions, a saline solution, rather than a Cl$_2$-bearing gas, would probably be evolved during crystallization of a hydrous Cl-bearing basalt magma. Concentrated brines have been identified as primary fluid inclusions in alkali-rich volcanic and plutonic rocks (Frost & Touret, 1989; Lowenstern, 1994). At 2–4 kbar and 1000°C, a NaCl solution could contain up to a few weight percent of sulfate, depending on the salinity. For instance, a mildly saline aqueous fluid having 5 mol % NaCl would have, at 1000°C and 4 kbar, an anhydrite saturation CaSO$_4$ moleality of 0.495, or 5 wt %, based on the present data. Because of the large temperature coefficient of anhydrite solubility, cooling to 800°C (0.143 m) would precipitate most of the CaSO$_4$. Thus, a substantial amount of anhydrite could be delivered to the host rocks, even for small fluid/rock ratios.

An important consideration is whether the anhydrite of the Mt Pinatubo dacites is simply a monitor of generally high oxidation state of the magmatism or whether fluid-transported sulfur undergoes spontaneous changes in oxidation state in cooling and reaction with country rocks such that it becomes an oxidizing agent after release from a basaltic magma source. The latter point of view was argued by Hattori (1993), who pointed to a greatly increased stability of H$_2$S in cooling of a C–O–H–S fluid from basalt magma temperatures. Oxygen liberated from the conversion of SO$_2$ to H$_2$S would oxidize ferrous iron in the partially molten dacite envelope and stabilize anhydrite.

Hattori (1993) envisioned a reduced and relatively dry basaltic source of the sulfur-bearing fluid. However, a sufficiency problem remains with this hypothesis: sulfur solubility in basalt magma would be at a minimum at low oxygen fugacity and low H$_2$O activity, which would therefore require that the external S-rich dacites react with very large volumes of fluid expelled from the basalts, the majority of the S remaining in the fluid as H$_2$S. If, on the other hand, the basaltic source magma was originally hydrous and highly oxidized, then, according to the measurements of Luhr (1990), the magma could contain a high sulfate content: 0.5 wt % S at 1000°C and 4 kbar, steadily increasing with pressure. A subduction-zone-derived basaltic magma could have
been initially very oxidized and sulfur-rich. The fluid separating from this magma would have been initially in oxidative equilibrium with the magma and therefore also very oxidized. High oxidation state would favor the partitioning of S into the evolving fluid phase. From this point of view, emplacement of anhydrite in the surrounding rocks would seem to be more an indicator of a fundamentally high oxidation state of the magmatism than an oxidizing mechanism. It should be pointed out, however, that SO₂- or sulfate-bearing fluids can conceivably increase somewhat in oxidation state by sulfidation reactions in country rocks or derivative magmas that the fluids traverse while cooling. Hattori (1993) found petrographic evidence of pyrrhotite emplacement in the Pinatubo dacites at an early stage of metasomatic alteration. Conversion of SO₂ to H₂S in the fluid could account for some additional oxidation of host rocks if fluid fluxes were large.

Pasteris et al. (1996) discovered brine inclusions in quartz crystals from Mt Pinatubo tephra. Some of these inclusions contain halite and anhydrite daughter crystals, indicating that extremely concentrated, S-rich saline fluids coexisted with partially molten dacites at a pre-eruptive stage. This information, together with Luhr’s (1990) experimental evidence for the existence of more primitive parental magmas, rich in sulfur and highly oxidized, with the experimental evidence for the important role of Cl in the early outgassing of mafic magmas (Webster et al., 1999), and with the present finding of the high carrying capacity for sulfate of Cl-rich aqueous fluids at high T and P, leads to the conclusion that concentrated and oxidized brine, expelled from deep-seated mafic magmas, should be considered as a medium for anhydrite deposition and oxidation in S-rich volcanism.

**Oxidized granulites**

Several workers have suggested that the source of heat for granulite-facies metamorphism may be commonly or principally deep-crustal intrusions of mantle-derived ('underplated') magmas (Bohlen, 1987; Harley, 1989). Until recently, advocates of underplating have not examined the possibility that infiltration of magma-derived Cl-rich and perhaps S-rich supercritical fluids, analogous to those discussed in the preceding section on S-rich volcanism, could have been a factor in the metamorphism.

A key consideration is the high oxidation states exhibited by some large granulite-facies terranes. The principal signatures of such granulites are high Fe₂O₃ contents of ilmenite (over 50 vol. % exsolved hematite in the Wilson Lake, Labrador and Shevaroy Hills, South India areas: Currie & Gittens, 1988; Harlov et al., 1997), high Mg/(Mg + Fe) ratios in the mafic minerals, indicative of Fe depletion of silicates, and the presence of Fe-deficient pyrrhotite and/or pyrite with magnetite. The Shevaroy Hills rocks are permeated with veins of pyrite + magnetite, and these two minerals without pyrrhotite accompany the highest Fe₂O₃ contents in ilmenite and the most Fe-depleted mafic silicates (Harlov et al., 1997), indicating that the high oxidation state shown by many of the Shevaroy Hills granulites is a near-peak metamorphic feature. Arima et al. (1986), on the other hand, inferred that the great oxidation of the Wilson Lake rocks was inherited from a highly oxidized sedimentary protolith. This hypothesis could not apply to the South Indian Archean Craton, in which orthogneiss greatly exceeds metasedimentary components, especially in view of the reducing sedimentary environments that prevailed in the Late Archean (Hattori & Cameron, 1986).

Cameron & Hattori (1994) suggested a link between highly oxidized granulites and S-rich volcanism, in that both phenomena might result from the action of oxidizing late-stage magmatic fluids. The possibility that SO₂ in fluids derived from underplated magmas could have played a role in oxidation of the Shevaroy Hills granulites was suggested by Harlov et al. (1997, p. 714), based on the abundance of sulfide–oxide veins in these rocks. The present work introduces a modification of these ideas, in that sulfate ion, dissolved in a concentrated brine, possibly of magmatic origin, could be effective in sulfidation and oxidation reactions during deep-crustal metamorphism. Interaction of sulfate-bearing solutions with deep-crustal (i.e. granulite facies) assemblages can be modeled with the reaction

\[
\text{CaSO}_4\text{aq} + 2\text{FeSiO}_3 = \text{FeS} + \text{CaFeSi}_2\text{O}_6 + 2\text{O}_2.
\]

Reaction (4) implies that the assemblage orthopyroxene (opx) + clinopyroxene (cpx) + pyrrhotite (po) generates, under granulite-facies P-T conditions and high oxygen fugacity, a high activity of CaSO₄, which, if a fluid is present, would correspond to a CaSO₄,aq concentration approaching anhydrite saturation. Conversely, an infiltrated metamorphic fluid with a high sulfate content, which would necessarily be highly oxidized, could serve as an agent of oxygen metasomatism. In order to coexist with the anhydrous assemblage of reaction (4) at 800°C and 10 kbar, the H₂O activity could be no higher than about 0-5, which could correspond to a NaCl mole fraction of near 0.3 (Aranovich & Newton, 1996). This solution could be very rich in CaSO₄, perhaps as much as 20 wt %, before anhydrite saturation would occur.

Figure 6 shows reaction (4) at 8 kbar, assuming that \(X_{\text{NaCl}} = 0.3\), \(X_{\text{Fe}}\) in ferrosilite is 0.5, \(X_{\text{Fe}}\) in hedenbergite is 0.2, pyroxene activity is the square of mole fraction, and that pyrrhotite activity is 0.875. These compositions are representative of the highest grade (Zone D) part of
the Bamble Sector, South Norway, or the Shevaroy Hills terrane, South India (Harlov, 1992; Harlov et al., 1997). The curve was constructed with thermodynamic data for the minerals from SUPCRT92 (Helgeson et al., 1978; Johnson et al., 1992). Contours of CaSO$_4$ concentration are shown as percentages of anhydrite saturation. The calculation is quite robust, in that an uncertainty of as much as $\pm 20\text{kJ}$ in $\Delta G$ of the reaction has only a small effect on the position of the curves. CaSO$_4$ activities are taken proportional to the squares of their concentrations, in keeping with reaction (3). Also shown in Fig. 6 are calculated oxygen fugacity and temperature data for individual rocks from the two terranes (Harlov, 1992; Harlov et al., 1997).

The indication of Fig. 6 is that the $fO_2$–$T$ conditions inferred for Bamble Zone D are compatible with CaSO$_4$ concentrations in aqueous fluids ranging from 1 to 10% of anhydrite saturation, and for the Shevaroy Hills of 15–40% of anhydrite saturation. The distinctly different oxidation signatures of the two terranes are not likely to result from a systematic difference in protolith oxidation, in view of the expected local variability of initial oxidation state, and in view of the more reduced nature of surficial conditions in the Archean (Hattori & Cameron, 1986), but more probably result from more consistent controls which characterized the high-grade metamorphism in the two areas. We propose that the difference in oxidation state may have resulted from unique sulfur and chlorine concentrations and oxidation states of underplated magmas in the two areas. As with Mt Pinatubo-type volcanism, the basaltic source of oxidizing volatiles must itself been very oxidized in order to accomplish major host–rock oxidation with moderate amounts of expelled fluid.

Small fluid/rock ratios are indicated by the often limited amounts of metasomatism and stable isotope homogenization observed in some granulites (Valley et al., 1990). The question of whether small amounts of fluids could have accomplished the large-scale oxidation observed in some granulites is addressed quite simply by calculating the amounts of infiltrated fluids necessary to transport enough oxygen to oxidize a significant

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**Fig. 6.** CaSO$_4$ in metamorphic fluids at 8 kbar, expressed as a percentage of anhydrite saturation in a saline fluid of $X_{NaCl} = 0.3$ (dashed lines). The array of such curves is limited at high $fO_2$ by anhydrite saturation [reaction (4), text]. All curves calculated using SUPCRT92 (Helgeson et al., 1978; Johnson et al., 1992). Anh, anhydrite; Fs, ferrosilite; Hd, hedenbergite; Po, pyrrhotite. Also plotted are the self-consistent $fO_2$–$T$ points for rocks from the Bamble (South Norway; Harlov, 1992) and Shevaroy Hills (South India; Harlov et al., 1997) terranes. If solute CaSO$_4$ accounts for the observed high oxidation states, the metamorphic fluids would be 1–10% CaSO$_4$-saturated for Bamble, and 15–40% CaSO$_4$-saturated for Shevaroy Hills.
amount, say half, of the FeO in an intermediate granulite to Fe₂O₃. For a typical Shevaroy Hills orthopyroxene-bearing orthogneiss of 5 wt % FeO, 0.56 wt % of additional oxygen must be introduced. For a NaCl-rich brine 25% saturated in anhydrite at 800°C and 8 kbar, the amount of introduced fluid of $X_{\text{NaCl}} = 0.3$ would have to be only 9-2 wt % of the rock mass it interacts with, assuming that reaction (4) goes to completion. This small quantity is consonant with the limited amount of fluid necessary to break down biotite to orthopyroxene (Aranovich & Newton, 1998), but would be too small to produce measurable oxygen isotope shifts in rocks that are volumetrically 90% oxygen, even if the infiltrating fluid is initially far removed from isotopic equilibrium with the host rock. On the other hands, the amounts of infiltrated pure H₂O or CO₂ that would be necessary to oxidize a substantial amount of the ferrous iron in an intermediate orthogneiss would be improbably large, as the amounts of O₂ such fluids can release by dissociation at high $f_O₂$ are minuscule (Frost, 1991). Thus, a concentrated sulfate-rich brine may be a uniquely effective vector for large-scale oxidation in the deep crust.

It is concluded that saline fluids, plausibly expelled from underplated mafic magmas, should be considered as an additional important factor in deep-crustal metamorphism. The highly variable volatile-element characteristics of mantle-derived magmas may in part account for the extremes of oxidation states observed in granulites, from the hematite-bearing granulites of Wilson Lake to the reduced granulites of the Ivrea Zone of North Italy, for which underplated magmas and their derivative saline solutions have been assigned major roles in the metamorphism (Franz & Harlov, 1998).

### Scapolite granulites

The high-sulfur scapolites of some mafic and intermediate granulites are confined to terranes where paleopressures of 9 kbar or more are recorded: Nilgiri Hills, South India (Srikantappa et al., 1986); Fiordland, New Zealand (Bradshaw, 1989); Kohistan Arc, Pakistan (Yoshino & Satish-Kumar, 2001); Furua Complex, Tanzania (Coolen, 1980); and in deep-crustal xenoliths in alkali basalts (Lovering & White, 1964). All of these occurrences are sulfide mizzonites of the approximate formula $3$An$0.67$Ab$0.33$C$0.3$C₃ and $15$ kbar, the restricted mizzonite in orthogneisses to high-pressure granulites may be in part because the endmember sulfate scapolite 3An-Cs (sulfate meionite) is stable only at pressures above 10 kbar at 800°C (Goldsmith & Newton, 1977) and partly because coexistence with orthopyroxene may be limited in pressure by a decarbonation reaction similar to

$$\text{meionite}(3\text{An} \cdot \text{Cc}) + \text{orthopyroxene} = \text{garnet} + \text{clinopyroxene} + \text{CO}_2.$$  

(Yoshino & Satish-Kumar, 2001).

The occurrence of some deep-crustal scapolites in veins (Bradshaw, 1989; Yoshino & Satish-Kumar, 2001) suggests that carbonate and sulfate were infiltrated volatiles, and carbon isotopes of these scapolites were interpreted by those workers to infer a juvenile magmatic or subcrustal origin of the volatiles. It is postulated as a consequence of the present experimental work that infiltration of sulfate and carbonate into the deep crust in a concentrated alkali chloride solution formed scapolite by reaction with intermediate plagioclase. The complex salt solution plausibly emanated from underplated basaltic magmas, which supplied the heat for metamorphism.

Existing experimental work on the stability of sulfate-carbonate mizzonite relative to plagioclase is not comprehensive enough to allow assessment of the CaSO₄ and CaCO₃ activities necessary to form scapolite in mafic to intermediate granulites at deep-crustal conditions. Goldsmith & Newton (1977) synthesized a mizzonite of equimolar Cc and Cs composition at 1100°C and 15 kbar in the presence of excess calcite and anhydrite. It has not yet been demonstrated that a low-Cl mizzonite can form from a concentrated NaCl solution undersaturated in calcite and anhydrite. Equilibria of feldspars with complex brines will be an important avenue of investigation for the future.

### Porphyry Cu–Mo ore deposits

Many of the world’s largest Cu–Mo sulfide ore deposits occur in the roof zones of small alkaline granite or monzonite plutons (e.g. White et al., 1981). Many of the lower-temperature alteration zones have disseminated to massive anhydrite with or without hematite. Several students of these deposits believed that ore-forming fluids must have encountered evaporite sequences in order to acquire CaSO₄; a minority preferred orthomagmatic saline fluids as a source of CaSO₄. Roedder (1971) found that several of the largest porphyry ore deposits of the western United States show very concentrated brine inclusions in vein quartz from the highest-temperature, near-magmatic deposits. The ultrasonic fluid inclusions typically contain abundant halide, carbonate, sulfate, sulfide and oxide daughter minerals. Roedder (1984, table 15-5) listed many worldwide occurrences of this type of ore deposit, with descriptions of the fluid inclusions. The majority of the occurrences have highly saline inclusions containing anhydrite and hematite, often with other sulfate and sulfide minerals. Roedder (1984) explained the highly oxidized nature of
the fluid inclusions by a process of H₂ leakage out of the inclusions after entrainment, with residual oxidation.

The preceding applications of the present experimental work have emphasized a juvenile/magmatic origin of highly oxidized brines containing large amounts of sulfate. We offer an analogous interpretation for the intrusion-related sulfide ore deposits—an interpretation that explains the highly oxidized fluid inclusions as a primary, rather than a secondary, effect. The very high solubility of CaSO₄ in concentrated NaCl solutions at near-magmatic temperatures (>750°C) indicates that a large proportion of solute sulfur in a mafic magma would partition into an exhalative saline fluid as sulfate. Subsequent interaction of this fluid with country rocks could provoke hydrous melting to produce hypabyssal granitic stocks or Pinatubo-type dacites. The solubility of Cl is much lower in granitic magmas than in mafic magmas (Webster et al., 1999), hence the salt components (and metals) of the postulated fluid will concentrate during fluid–rock interactions as H₂O is extracted into granitic liquids. Precipitation of sulfate minerals would further increase the oxygen fugacity of the already highly oxidized fluids. An intimate connection between Pinatubo S-rich volcanism and young western Luzon porphyry ore deposits in the same area was inferred by Imai et al. (1996). We postulate that both phenomena could be the result of the same basalt-derived saline fluid processes manifest at different crustal levels. This conjecture is similar in principle to processes of porphyry copper mineralization envisioned by Hattori & Keith (2001).

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