The S content of silicate melts at sulfide saturation: New experiments and a model incorporating the effects of sulfide composition

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**ABSTRACT**

The extent to which sulfur dissolves in silicate melts saturated in an immiscible sulfide phase is a fundamental question in igneous petrology and plays a primary role in the generation of magmatic ore deposits, volcanic degassing, and planetary differentiation. In igneous systems, sulfide melts can be described as FeS-NiS-CuS$_{0.5}$ solutions with Fe/(Fe+Ni+Cu) significantly less than 1. Despite the presence of Ni and Cu in the sulfide, however, most experimental studies to date have concentrated on the effects of silicate melt composition on sulfur solubility and have used essentially pure FeS as the sulfide liquid.

We have carried out 49 new experiments at pressures of 1.5–24 GPa and temperatures of 1400 to 2160 °C to investigate the effects of sulfide composition on sulfur solubility as well as extending the pressure and temperature ranges of the available data on sulfide saturation. We find that in the compositional range of most sulfide melts [Fe/(Fe+Ni+Cu) > 0.6] sulfur solubility decreases linearly with Fe content such that at Fe/(Fe+Ni+Cu) of 0.6 the sulfur content at saturation is 0.6 times the value at pure FeS saturation. At lower values of Fe/(Fe+Ni+Cu), however, deviations from this ideal solution relationship need to be taken into consideration. We have treated these non-idealities by assuming that FeS-NiS-CuS$_{0.5}$ liquids approximate ternary regular solutions.

We have fitted our data, together with data from the literature (392 in total), to equations incorporating the effects of silicate melt composition, sulfide liquid composition, and pressure on the solubility of sulfur at sulfide saturation ([S]$_{SCSS}$). The temperature dependence of [S]$_{SCSS}$ was assumed either to be an unknown or was taken from 1 bar thermodynamic data. The most important best-fit silicate melt compositional term reflects the strongly positive dependence of [S]$_{SCSS}$ on the FeO content of the silicate melt. The best-fit value of this parameter is essentially independent of our assumptions about temperature dependence of [S]$_{SCSS}$ or the solution properties of the sulfide.

All natural compositions considered here exhibit positive dependences of [S]$_{SCSS}$ on temperature and negative dependences on pressure, in accord with previous studies using smaller data sets.

**Keywords:** Sulfur solubility, silicate melt, sulfide, MORB

**INTRODUCTION**

The solubility of sulfur in silicate melts is a subject that attracts the attention of metallurgists interested in the conditions under which immiscible sulfide mattes segregate from silicate slags (e.g., Fincham and Richardson 1954) and of geologists concerned with the behavior of sulfur and chalcophile elements during igneous processes. In the former case, the work of Fincham and Richardson led to a quantitative model for the solubility of S (as S$_{2}$) and its dependence on oxygen and sulfur fugacities. In the latter the dependence of sulfur solubility on silicate melt composition and temperature control volcanic degassing (Fischer et al. 1998; Wallace 2005) and the formation of magmatic sulfide ore deposits (Li and Ripley 2005; Mungall 2007). Sulfide liquid precipitates during the differentiation of MORB (Peach et al. 1990), thereby controlling the concentrations of chalcophile elements such as Cu, Ag, Tl, and the PGEs in the crystallizing silicate melts. Precipitation of immiscible sulfide and sulfur solubility may also be important during planetary accretion and differentiation (Holzheid and Grove 2002; Wood et al. 2014). For these reasons there have been numerous experimental investigations of the processes that control the incorporation of sulfur in naturally occurring silicate melts (e.g., Shima and Naldrett 1975; Mavrogenes and O’Neill 1999; O’Neill and Mavrogenes 2002; Li and Ripley 2005; Liu et al. 2007).

At relatively low oxygen fugacities [i.e., below that of the fayalite-magnetite-quartz (FMQ) buffer], Fincham and Richardson (1954) proposed that sulfur dissolves in silicate melts as S$_{2}$ and that it substitutes for oxygen on the anion sublattice via the reaction:

$$O^{2-} + \frac{1}{2}S_{2} = S^{2-} + \frac{1}{2}O_{2}.$$  

(1)

In silicate melts, the concentrations of O$_{2}$ are generally two or more orders of magnitude greater than those of S$_{2}$, even at sulfide saturation. Given this constraint we can take the O$_{2}^{-}$ concentration on the anion sublattice to be constant and rearrange the equilibrium constant for reaction 1 to yield the Fincham-Richardson relationship (Fincham and Richardson 1954):

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\[ \ln C_s = \ln[S] + \frac{1}{2} \ln \left( \frac{f_{O_2}}{f_{S_2}} \right). \] (2)

In Equation 2, \( C_s \) is the sulfide capacity of the melt (analogous to the equilibrium constant), and \([S]\) is the concentration of sulfur, usually in parts per million. Fincham and Richardson (1954) experimentally verified the relationship of Equation 2 by measuring sulfur contents of silicate melts in the system CaO-Al₂O₃-SiO₂ at fixed values of \( f_{O_2} \) and \( f_{S_2} \). In the geologic literature, most interest has been on the conditions of sulfide saturation and precipitation of sulfides from basaltic and related liquids (e.g., Haughton et al. 1974; Katsura and Nagashima 1974; Wallace and Carmichael 1992). Nevertheless, O’Neill and Mavrogenes (2002) broadened the scope of study by measuring the concentrations of \( S \) in 19 melts in the geologically relevant system CaO-MgO-Al₂O₃-SiO₂±TiO₂±FeO. They showed that, at 1400 °C and known \( f_{O_2} \) and \( f_{S_2} \) between –3.36 and 1.59, and –10.92 and –6.78, respectively, all 19 melts obey the Fincham-Richardson relationship.

Because of its geologic importance and the large number of data currently available, our study has been primarily concerned with the conditions of sulfide saturation in silicate melts of different composition over wide ranges of pressure and temperature. In this context we begin by assuming that the Fincham-Richardson relationship applies to all melts of geologic interest. Equilibrium between sulfide and silicate melt may then be described in terms of the reaction:

\[ \text{FeO}_{\text{silicate}} + \frac{1}{2} S_2 = \text{FeS}_{\text{sulfide}} + \frac{1}{2} O_2. \] (3)

for which, at equilibrium we have:

\[ \Delta G^o = -RT \ln \left( \frac{f_{O_2}^{\text{FeS}} f_{S_2}^{\text{FeS}}}{f_{O_2}^{\text{FeO}} f_{S_2}^{\text{FeO}}} \right) \] (4)

where \( \Delta G^o \) is the standard state free energy change of reaction 3 and activities and fugacities of the four components have their usual symbols. Rearranging Equation 4 we obtain:

\[ \frac{\Delta G^o}{RT} = \ln \frac{a_{\text{FeS}}^{\text{silicate}}}{a_{\text{FeS}}^{\text{FeO}}} - \ln \frac{a_{\text{FeO}}^{\text{silicate}}}{a_{\text{FeO}}^{\text{FeS}}}. \] (5)

We may now substitute from Equation 2 for the ratio of oxygen to sulfur fugacity as follows:

\[ \ln[S]_{\text{SCSS}} = \frac{\Delta G^o}{RT} + \ln C_s - \ln a_{\text{FeS}}^{\text{silicate}} + \ln a_{\text{FeO}}^{\text{FeS}}. \] (6)

In Equation 6, \([S]_{\text{SCSS}}\) refers to the sulfur content of the silicate melt at sulfide saturation and \( a_{\text{FeS}}^{\text{FeO}} \) and \( a_{\text{FeO}}^{\text{FeS}} \) to the activities of FeS and FeO components in sulfide and silicate melts, respectively.

Inspection of Equation 6 enables us to consider, qualitatively, the important influences on the sulfur content of any particular melt at sulfide saturation. The standard state free energy change of the reaction \( \Delta G^o \) depends on pressure and temperature, which requires that \([S]_{\text{SCSS}}\) is also \( P-T \) dependent. Measurements of \( C_s \), and of sulfur concentrations at sulfide saturation have demonstrated that \( C_s \) is composition dependent, most notably varying strongly and positively with the FeO content of the silicate melt. In contrast, the term \( a_{\text{FeO}}^{\text{FeS}} \) requires an increase in S content with decreasing FeO content of the silicate melt. There is thus a trade-off between the positive contribution of FeO to \([S]_{\text{SCSS}}\) and the negative effect of \( a_{\text{FeO}}^{\text{FeS}} \) on sulfur solubility that leads to a theoretical U-shape of a plot of \([S]_{\text{SCSS}}\) vs. FeO content (O’Neill and Mavrogenes 2002 Fig. 21). Thus, most attempts to model \([S]_{\text{SCSS}}\) have emphasized the major compositional terms for the silicate melt. Pressure effects have also been shown to be very important (e.g., Holzheid and Grove 2002; Mavrogenes and O’Neill 1999). Much less attention has been paid to the composition of the sulfide melt, however, as represented by \( a_{\text{FeS}}^{\text{silicate}} \) in Equation 6 (Ariskin, et al. 2013). Instead, almost all experimental measurements to date have used pure FeS as the sulfide phase, with the implicit assumption that lowering \( a_{\text{FeS}}^{\text{silicate}} \) by diluting Fe with other cations has no effect on sulfur solubility. Equation 6 indicates, however, that S solubility must decrease as FeS activity decreases, meaning that, in general it must be lower than is implied by pure FeS saturation. Immiscible droplets of sulfide in basaltic contain up to 18 and 20 wt% Ni and Cu, respectively (Francis 1990; Patten et al. 2013), which means that the sulfide is only about 60% FeS by mole. Despite these observations, and the appreciable effort made, to date, to determine S solubility in silicate melts, there are few data enabling the effects of dilution of Fe by other cations to be evaluated. One major goal of the present experimental study is to address this fundamental question and to quantify the effects of Ni and Cu substitutions on FeS activity and hence on \([S]_{\text{SCSS}}\) in relevant natural compositions. To this end we have performed 49 new experiments at 1.5 to 24 GPa and 1400 to 2160 °C with immiscible sulfide melt compositions ranging from pure FeS to nearly pure NiS and CuS. The new data enable us to quantify the effects of Ni and Cu substitution for Fe as well as providing the basis, together with literature data, for determining the effects of pressure and temperature and silicate melt composition on \([S]_{\text{SCSS}}\).

**EXPERIMENTAL AND ANALYTICAL PROCEDURES**

**Experimental methods**

Starting materials consisted of mixtures of ~50% (Fe,Ni,Cu)S and ~50% synthetic silicate, by weight. The sulfide component consisted of mixtures of analytical grade FeS, NiS, and CuS. The silicate constituent was in many cases a composition close to the 1.5 GPa eutectic composition in the system anorthite–diopside–forsterite (Anₓ,Diₓ,Fo₁₋ₓ) (Presnall et al. 1978) with variable FeₓNi₁₋ₓ added, but a range of basaltic, andesitic, and komatiitic compositions was also employed. All components were added to these mixtures either as analytical-grade oxides (SiO₂, TiO₂, Al₂O₃, MgO, FeO, MnO, P₂O₅) or as carbonates (Na₂CO₃, K₂CO₃, CaCO₃). The silicate mixtures were pelleted and decaarbonated at 950 °C for 2 h. After that the FeO was added, mixtures were reground, pelleted, and reduced in a CO-CO₂ atmosphere for 2 h at 1000 °C and an oxygen fugacity approximately 2 log units above the IW buffer. Silicate and sulfide constituents were intimately mixed in approximately 50:50 proportions prior to each experiment. Additional Fe (as Fe₃C)O was added to some experiments to increase FeO activity. The starting mixtures were dried at 110 °C immediately before the experiment.

Most experiments were performed at 1.5 and 2.5 GPa using a 12.7 mm diameter Boyd-England type piston-cylinder apparatus at the University of Oxford. The sample cell employed an outer sleeve of pressed CaF₂ graphite heater of 8 mm outside and 6 mm inside diameter and internal parts of machineable MgO. Most experiments were performed in 3.0 mm O.D., 1 mm I.D. graphite capsules, a subset of which were sealed in Pt outer capsules. A few experiments were performed in SiO₂ glass capsules. Experiment durations were fixed at times substantially longer than those required to reach sulfide-silicate and metal-silicate partitioning equilibrium in capsules of 1 mm I.D. (Tuff et al. 2011; Kiseeva and Wood 2013). For piston-cylinder experiments at 1500 °C and higher temperatures

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the outer calcium fluoride sleeve was replaced by an outer thin-walled BaCO₃ cylinder with an inner sleeve of SiO₂ glass. All temperatures were monitored and controlled using W₆Re₂-W₅Re₃ thermocouples housed in alumina sheaths and separated from the capsule by a 0.6 mm thick alumina disk. Experiments at 5.5 and 7 GPa employed a Walker-type multi-anvil apparatus and cast MgO-based octahedra. The furnace assemblies consisted of straight graphite heaters inside ZrO₂ sleeves and the capsules were, as before fabricated from graphite. One experiment was performed at 24 GPa/2160 °C at the Bayerisches Geoinstitut, Universität Bayreuth. In this case the capsule was made of single-crystal MgO. As for the piston-cylinder experiments, all multi-anvil experiments employed W-Re thermocouples, in these cases in direct contact with the capsule. Experimental run conditions and starting compositions are given in Table 1.

**Analytical techniques**

Experimental products were analyzed using a JEOL JXA8600 electron microprobe housed in the Department of Archaeology at the University of Oxford. Silicate glasses were analyzed by wavelength-dispersive spectroscopy (WDS) employing a 15 kV accelerating voltage and 12 ± 20 nA beam current. The spot was typically defocused to 10 μm diameter. Standards used for silicate glass analysis include natural wollastonite (Si, Ca), natural jadeite (Na, Al), synthetic periclase (Mg), rutile (Ti), hematite (Fe), NdPO₃ (P), orthoclase (K), and galena (S). Natural almandine and natural S-bearing glasses VG-1 and L17 were used as secondary standards for S. Counting times were as follows: 30 s and 15 s background for major elements (Si, Al, Ca, Mg, Fe); 60 s peak and 30 s background for minor elements (Na, K, Ti, P); 180 s peak and 90 s background for S. Analysis of sulfides by WDS used a 15 kV accelerating voltage, a 20 nA beam current and a defocused beam, generally of 10 μm, but occasionally of 15 μm diameter for the most heterogeneous quenched liquids. Standards for sulfide analysis consisted of Hematite (Fe, O), galena (S), Ni metal (Ni), and Cu metal (Cu). Count times were 30 s peak and 15 s background for Fe and S, and 60 s peak and 30 s background for Ni and Cu. Oxygen in the sulfide was measured using the Kα peak and a LDE pseudozircon (Kiseeva and Wood 2013) with count times of 100 s peak and 50 s background. There was no evidence of strong heterogeneity in oxygen contents, as might have been expected if there were large grains of quenched oxide within the sulfide blobs. Electron microprobe analyses of silicate and sulfide run products are provided in Supplemental Table S1.

**Table 1. Experimental run conditions**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Starting composition</th>
<th>Temperature (°C)</th>
<th>Pressure (GPa)</th>
<th>Capsule</th>
<th>Duration (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N1-1</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
<td>1400</td>
<td>1.5</td>
<td>Graphite</td>
<td>120</td>
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<tr>
<td>N1-2</td>
<td>Basalt + 6% FeO + 42% FeS + 8% CuS</td>
<td>1400</td>
<td>1.5</td>
<td>SiO₂</td>
<td>90</td>
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<tr>
<td>N1-3</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
<td>1400</td>
<td>1.5</td>
<td>SiO₂</td>
<td>120</td>
</tr>
<tr>
<td>N1-4</td>
<td>Basalt + 5% FeO + 25% FeS + 25% NiS</td>
<td>1400</td>
<td>1.5</td>
<td>SiO₂</td>
<td>120</td>
</tr>
<tr>
<td>N1-5</td>
<td>Basalt + 5% FeO + 25% FeS + 25% NiS</td>
<td>1400</td>
<td>1.5</td>
<td>SiO₂</td>
<td>120</td>
</tr>
<tr>
<td>N1-6</td>
<td>Basalt + 5% FeO + 25% FeS + 25% NiS</td>
<td>1400</td>
<td>1.5</td>
<td>SiO₂</td>
<td>120</td>
</tr>
<tr>
<td>F1-1</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
<td>1400</td>
<td>1.5</td>
<td>SiO₂</td>
<td>90</td>
</tr>
<tr>
<td>F1-2</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
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<td>SiO₂</td>
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<td>F1-3</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
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<td>SiO₂</td>
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<td>F1-4</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
<td>1400</td>
<td>1.5</td>
<td>SiO₂</td>
<td>120</td>
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<tr>
<td>F1-5</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
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<td>1.5</td>
<td>SiO₂</td>
<td>120</td>
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<tr>
<td>F1-6</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
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<td>SiO₂</td>
<td>120</td>
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<tr>
<td>F1-7</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
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<tr>
<td>F1-8</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
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<tr>
<td>F1-9</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
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<td>120</td>
</tr>
<tr>
<td>F1-10</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
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<td>1.5</td>
<td>SiO₂</td>
<td>120</td>
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<tr>
<td>F1-11</td>
<td>An₁₋₁ Di₂₋₂ Fe₂₋₂ S</td>
<td>1400</td>
<td>1.5</td>
<td>SiO₂</td>
<td>120</td>
</tr>
</tbody>
</table>

Notes: An = anorthite, Di = diopside, Fo = forsterite, Qz = quartz, Wo = wollastonite, Per = periclase, Cor = corundum, And = andalusite, BCR-2 = USGS Columbia River Basalt.
RESULTS

O’Neill and Mavrogenes (2002) followed Haughton et al. (1974) in treating the measured sulfide capacity as a parameter with simple dependence on the mole fractions $X_M$ of the single metal oxide components (SiO$_2$, AlO$_1.5$, MgO, etc.)

$$\ln C_S = A_\theta + \sum_M X_M A_M / T. \quad (7)$$

The rationale for this approach is that the $A_M$ are related to the differences between the standard state free energies $\mu^\circ_M$ of oxide and sulfide components of the cations of interest (i.e., MgO, MgS, SiO$_2$, Si$_2$O, Si$_3$S etc.). This gives the following theoretical form for $C_S$:

$$\ln C_S = -\ln \gamma_S + \sum_M X_M \mu^\circ_{M,S} / RT. \quad (8)$$

In Equation 8, $\left(\mu^\circ_{M,S} - \mu^\circ_{M,O}\right)$ is the standard state free energy difference between oxide and sulfide components of M and $\gamma_S$ is the activity coefficient of S in the silicate melt. Replacing $C_S$ in Equation 6 with the form of Equation 7 leads to:

$$\ln [S]_{SCSS} = \frac{\Delta G^\circ}{RT} + A_\theta + \sum_M X_M A_M / T + \ln \alpha_{FeO}^{\text{silicate}} - \ln \alpha_{FeO}^{\text{oxide}}. \quad (9)$$

O’Neill and Mavrogenes (2002) used tabulated thermodynamic data to obtain $\Delta G^\circ$ at 1 bar then regressed their $C_S$ data at 1400 °C and 1 bar to an equation of similar form to Equation 8 except, since their experiments were isothermal, the dependence of the compositional $X_M A_M$ terms on temperature ($X_M A_M / T$) was ignored. If we use their results at face value, all of the regressed $A_M$ terms are positive and $\Delta G^\circ$ has a negative temperature dependence. This means that $[S]_{SCSS}$ should decrease with increasing temperature irrespective of whether or not we treat the $X_M A_M$ terms as having dependences on reciprocal temperature. All observations to date, however, including our own, demonstrate that $[S]_{SCSS}$ increases with increasing temperature. This means either that the temperature dependence of $\Delta G^\circ$ derived from tabulated thermodynamic data is profoundly in error or that the $X_M A_M$ terms of Equation 9 are predominantly negative. Although we consider that the former is unlikely, we have treated the data in two fundamentally different ways to resolve this issue. First, we treated $\Delta G^\circ$ and its temperature and pressure dependences as unknowns. In this case we divide $\Delta G^\circ / RT$ into enthalpy ($\Delta H^\circ$), entropy ($\Delta S^\circ$), and molar volume terms ($\Delta V^\circ$) as follows:

$$\frac{\Delta G^\circ}{RT} = \frac{\Delta H^\circ}{RT} - \frac{\Delta S^\circ}{R} + \frac{\Delta V^\circ}{RT} = A_\theta + B + \frac{CP}{T}. \quad (10)$$

Combining Equations 9 and 10 yields

$$\ln [S]_{SCSS} = \frac{A_\theta}{T} + B + \frac{CP}{T} + \sum_M X_M A_M / T + \ln \alpha_{FeO}^{\text{silicate}} - \ln \alpha_{FeO}^{\text{oxide}} \quad (11)$$

where $B'$ is the sum of the entropy term in Equation 10 and $A_\theta$ from Equation 7. We now use Equation 11 as one of the bases for our regression of $\ln [S]_{SCSS}$ as a function of compositional terms $A_M$, pressure, and temperature.

Our second approach was to assume that the tabulated thermodynamic data are correct and to adopt the 1-bar values of $\Delta G^\circ$ given by O’Neill and Mavrogenes (2002). This gives the following equation:

$$\ln [S]_{SCSS} = \frac{14695}{T} - 9.656 + 1.02\ln T + \frac{B' + \frac{CP}{T} + \sum_M X_M A_M / T + \ln \alpha_{FeO}^{\text{silicate}} - \ln \alpha_{FeO}^{\text{oxide}}}{T}. \quad (12)$$

In Equation 12 the only part of $\Delta G^\circ$ that is treated as unknown is the pressure-dependence, with fit parameter $C$. The parameter $B'$ in this case corresponds to $A_\theta$ of Equation 7

Linear least-squares regression

We have a total of 392 experimental data (Supplemental Table S2), 343 from previous studies and 49 from this work as the basis of our fits to Equations 11 and 12. These include data on hydrous melts containing up to 8.5 wt% H$_2$O equilibrated under conditions where the FeS-rich sulfide was a liquid. Note that we have explicit terms for the effect of H on [S]$_{SCSS}$ in our fits of Equations 11 and 12 (Table 3). We started with the assumption that $\alpha_{FeO}^{\text{oxide}}$ is equal to the mole fraction of FeO in the silicate melt on a single cation basis using components FeO, SiO$_2$, AlO$_1.5$, and so on. In practice, FeO has an activity coefficient close to 1 in silicate melts over a fairly wide compositional range (Wood and Wade 2013). The ideal approximation should, therefore, be adequate for our needs, particularly since any compositional dependence of $\alpha_{FeO}^{\text{silicate}}$ will be absorbed by the $A_M$ terms of Equations 11 and 12. An exception to this are experiments done at highly reducing conditions (i.e., below the iron-wüstite oxygen buffer). Silicate melts in equilibrium with sulfide melt under these conditions contain less than 1 wt% FeO. Based on sulfide/silicate trace-element partitioning (Wood and Kiseeva 2015) at these low FeO concentrations $\alpha_{FeO}^{\text{silicate}}$ decreases by at least an order of magnitude and our assumption that $\alpha_{FeO}^{\text{silicate}}$ is unity is no longer valid. To avoid these unusual compositions biasing our results we have therefore excluded from the regression six experiments with FeO$_{sil}$/FeS$_{sil}$ ratios of less than 0.01. These are nevertheless shown in Figure 1 for comparison with the remaining 392 data.

Kiseeva and Wood (2013, 2015) have shown that, to a good approximation, sulfide liquids in igneous systems can be treated as ideal FeS-NiS-CuS$_{0.5}$ solutions when trace-element partitioning between sulfide liquids and silicate melts is considered. We therefore began by approximating $\alpha_{FeO}^{\text{oxide}}$ by $X_{FeO}^{\text{silicate}}$ where $X$ is equal to Fe/(Fe+Ni+Cu). Following Kiseeva and Wood (2013) we took no explicit account of the presence of oxygen in the sulfide. Kiseeva and Wood (2013) found that the FeO content of the sulfide (in weight percent) is approximately the same as the FeO content of the silicate in weight percent. Nevertheless, they found that making explicit provision for the entropy of mixing of O$^-$ into the S$^2$ sublattice in FeS liquids using a Temkin-like solution model generated a worse approximation for $\alpha_{FeO}^{\text{silicate}}$ than the simpler Fe/(Fe+Ni+Cu). We therefore began by using this simple “ideal” solution model.

Step-wise linear-regression of the data set to Equations 11 and 12 was carried out using the statistics package SPSS with the requirement that fit parameters pass the F-test at F of 0.05. The results of the regression are presented in Table 3.

If we treat $\Delta G^\circ$ and its temperature and pressure dependences as unknowns (Eq. 11) then we find that $\Delta G^\circ / RT$ has a negative temperature dependence that is compensated-for, as predicted, by negative values of almost all the $A_M$ parameters. Thus, $[S]_{SCSS}$...
increases with increasing temperature because the negative $A_M/T$ terms become smaller with increasing temperature. Note that we tested the possibilities of adding cross-terms of the form $X_iX_jA_{ij}$ but found that only the $X_0X_F$ term is significant. It is this term that controls the positive effect of FeO on S solubility. Note also that we have data for silicate melts containing between 0.3 and 40.1 wt% FeO, so the effect of FeO on S solubility is very well constrained.

Our second set of regressions assumed that $\Delta G^o$ derived from thermodynamic data is correct and involved the fitting of Equation 12 to the same 392 data as before. In this case (Table 3), we have a slightly improved $r^2$ (0.977 instead of 0.963) and the negative $A_M$ terms are all smaller than those obtained from the fit to Equation 11. This is because the temperature dependence of $\Delta G^o$ obtained from thermodynamic data is much less negative than that obtained by treating $\Delta H^o$ and $\Delta S^o$ as unknowns. Hence, the magnitudes of the negative terms required to compensate for the temperature dependence of $\Delta G^o$ and to produce the observed positive temperature dependence of $\langle S \rangle_{SCSS}$ are smaller if $\Delta G^o$ is fixed at the tabulated value from thermodynamic data. As one might expect, the pressure term is, within uncertainty the same in both cases. Interestingly, the positive $X_0X_F$ term is also the same in both cases within uncertainty. We consider therefore that this large positive cross-term is well-constrained and realistic since it does not depend significantly on the assumptions made about the standard state free energy change of the sulfur dissolution reaction.

Figure 1 shows the results of the regression plotted as calculated ln[$\langle S \rangle_{SCSS}$] vs. the observed value for all 392 data used in the regression as well as the six experiments done under highly reducing conditions. Of 398 calculated $\langle S \rangle_{SCSS}$ values, 351 fall within the range of 0.667–1.5 times the observed S concentrations, which we consider an excellent result considering the ranges of pressure (1 bar to 24 GPa), temperature (1150–2160 °C), FeO content of silicate (0.3–40.1%) sulfide composition covering the entire range of the FeS-NiS-CuS$_{0.5}$ system employed in the regression.

**Effects of temperature, pressure, and silicate melt composition**

Figure 2a shows the effects of FeO content on S concentration at FeS saturation for melts ranging in composition from komatiitic to andesitic. Silicate melt compositions are given in Table 4. As anticipated from Equation 6 calculated $\langle S \rangle_{SCSS}$ is a u-shaped function of FeO content because of the trade-off between the positive coupling between S and FeO in the silicate and the negative influence of $a_{FeO}^{\text{silicate}}$ on S solubility (O’Neill and Mavrogenes 2002). Our model predicts that the solubility of S in silicate melts should reach a minimum at approximately 5 wt% FeO, which is broadly consistent with the work of Li and Ripley (2005). The result disagrees, however, with the expressions of Holzheid and Grove (2002) and Fortin et al. (2015), which do not take account of the term in $a_{FeO}^{\text{silicate}}$ required by Equation 6.

As can be seen in Figure 3a all compositions show a positive dependence of $\langle S \rangle_{SCSS}$ on temperature. Note that results are, in some cases, extrapolated to temperatures below those of the silicate liquidi. Figure 3b shows a comparison of our results for $\langle S \rangle_{SCSS}$ with literature models of S contents at FeS saturation for the MORB composition of Table 4. The latter are based on more limited data sets than those available to us. As can be seen, our results for MORB are in very good agreement with the model of Fortin et al. (2015) but that of Li and Ripley (2005) predicts much higher concentrations than those observed. The results of O’Neill and Mavrogenes (2002) are close to ours at their experimental temperature of 1400 °C, but their equation does not provide for the correct form of the temperature extrapolation, as noted above.

Figures 4a and 4b shows the calculated effects of pressure on the solubility of sulfur in the silicate melt compositions of Table 4 at a fixed temperature of 1400 °C. All melts of concern exhibit predicted declines in $\langle S \rangle_{SCSS}$ with increasing pressure, consistent with previously published results (Holzheid and Grove 2002). Figure 4b shows, consistent with Figure 3, that the best agreement with previous results is with the model of Fortin et al. (2015).

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**Table 3. Results of linear least-squares regression of experimentally determined sulfur solubilities in silicate melts**

<table>
<thead>
<tr>
<th></th>
<th>Ideal $\Delta G^o$ (this study)</th>
<th>Ideal $\Delta G^o$ (O+M 2002)</th>
<th>Non-ideal $\Delta G^o$ (O+M 2002)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coefficient</td>
<td>Std. Error</td>
<td>Coefficient</td>
<td>Std. Error</td>
</tr>
<tr>
<td>$A$</td>
<td>-14683 (452)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>$B'$</td>
<td>8.03 (0.25)</td>
<td>9.087 (0.25)</td>
<td>9.352 (0.25)</td>
</tr>
<tr>
<td>$C$</td>
<td>-265.80 (24.07)</td>
<td>-269.40 (24.17)</td>
<td>-264.85 (23.68)</td>
</tr>
<tr>
<td>$A_0$</td>
<td>-27561 (500)</td>
<td>-27996 (500)</td>
<td></td>
</tr>
<tr>
<td>$A_1$</td>
<td>16430 (1465)</td>
<td>-11220 (1424)</td>
<td>-10715 (1398)</td>
</tr>
<tr>
<td>$A_2$</td>
<td>9295 (811)</td>
<td>-18450 (794)</td>
<td>-19000 (768)</td>
</tr>
<tr>
<td>$A_{Fe}$</td>
<td>13767 (515)</td>
<td>-13970 (627)</td>
<td>-14512 (627)</td>
</tr>
<tr>
<td>$A_{Cu}$</td>
<td>19893 (737)</td>
<td>-7831 (856)</td>
<td>-8832 (871)</td>
</tr>
<tr>
<td>$A_{Ni}$</td>
<td>-7080 (2082)</td>
<td>-34274 (2376)</td>
<td>-34895 (2330)</td>
</tr>
<tr>
<td>$A_{Na}$</td>
<td>14197 (1441)</td>
<td>-13247 (1414)</td>
<td>-13713 (1388)</td>
</tr>
<tr>
<td>$A_{Si}$</td>
<td>-29015 (2962)</td>
<td>-28584 (2900)</td>
<td></td>
</tr>
<tr>
<td>$A_{Fe}$</td>
<td>-2799 (560)</td>
<td>-17495 (561)</td>
<td>-17766 (553)</td>
</tr>
<tr>
<td>$A_{Fe}$</td>
<td>117827 (5474)</td>
<td>116568 (6066)</td>
<td>117816 (5943)</td>
</tr>
<tr>
<td>$E$</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Effect of sulfide composition

One of the major aims of our study was to investigate the effects of sulfide liquid composition on $[S]_{\text{SCSS}}$. As can be seen from Figure 1 and Supplemental Table S1, the assumption of ideal FeS-NiS-CuS$_{0.5}$ solution produces very good fits to the experimental data over most of the composition range investigated and implies that $[S]_{\text{SCSS}}$ declines almost linearly as Fe/(Fe+Ni+Cu) declines. It is known, however, that neither FeS-NiS (Fleet 1989) nor FeS-CuS$_{0.5}$ (Eric and Timucin 1981) melt solutions are perfectly ideal and these deviations from ideality likely cause the deviations of $[S]_{\text{SCSS}}$ from the predicted values at low Fe/(Fe+Ni+Cu) (Supplemental Table S1).

Non-ideality in the sulfide solution can be treated in several different possible ways. We could use previously measured activity coefficients from the studies mentioned above. This would require re-fitting Equations 11 and 12 to derive new values of the $A_M$ parameters. Since, however, our results indicate relatively small deviations from FeS-NiS-CuS$_{0.5}$ ideality, we have opted to use a simple non-ideal solution model and to treat the non-ideality parameters as unknowns. In that case, applying a ternary symmetrical solution model for $A_{\text{FeS}}$ (e.g., Wood and Fraser 1976) we add compositional terms as follows to that involving $A_{\text{FeS}}$

$$\ln a_{\text{FeS}} = \ln X_{\text{FeS}} + \frac{W_{\text{NiFe}}}{RT} X_{\text{NiS}} + \frac{W_{\text{CuFe}}}{RT} X_{\text{CuS}} - W_{\text{NiCu}}.$$  (13)

In Equation 12, the $W_{ij}$ parameters are the interaction parameters for $i-j$ pairs and are obviously 0 if the solution is ideal. Treating the $W_{ij}/R$ as fit parameters and adding them to Equation 12 yields:

**TABLE 3.** Silicate melt compositions used for modeling of SCSS in Figures 2, 3, and 4

<table>
<thead>
<tr>
<th>Melt Type</th>
<th>SiO$_2$ (wt%)</th>
<th>TiO$_2$ (wt%)</th>
<th>Al$_2$O$_3$ (wt%)</th>
<th>FeO (wt%)</th>
<th>MgO (wt%)</th>
<th>CaO (wt%)</th>
<th>Na$_2$O (wt%)</th>
<th>K$_2$O (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N-MORB$^a$</td>
<td>50.42</td>
<td>1.53</td>
<td>15.13</td>
<td>9.81</td>
<td>7.76</td>
<td>11.35</td>
<td>2.83</td>
<td>0.14</td>
</tr>
<tr>
<td>Martian Basalt$^b$</td>
<td>45.50</td>
<td>0.60</td>
<td>6.70</td>
<td>17.90</td>
<td>14.30</td>
<td>7.65</td>
<td>0.70</td>
<td>0.05</td>
</tr>
<tr>
<td>Andesite$^c$</td>
<td>56.52</td>
<td>1.08</td>
<td>17.54</td>
<td>7.65</td>
<td>14.30</td>
<td>7.40</td>
<td>3.94</td>
<td>1.31</td>
</tr>
<tr>
<td>Komatiite$^d$</td>
<td>46.80</td>
<td>0.26</td>
<td>4.50</td>
<td>11.00</td>
<td>29.60</td>
<td>5.21</td>
<td>0.28</td>
<td>0.14</td>
</tr>
</tbody>
</table>

$^a$ Average N-MORB (Gale et al. 2013).
$^b$ Basaltic shergottite Dhofar019 (Bridges and Warren 2006).
$^c$ Average andesite (Wilkinson 1986).
We fitted Equation 13 to the same 392 $[\text{S}]_{\text{SCSS}}$ data as those used previously using, once more, stepwise linear regression. This approach led to $D$ and $F$ parameters that are not statistically significant and an $E$ parameter of 546 K (Table 2). The latter reflects Fe-Cu interactions and its inclusion leads to improvements in calculated $[\text{S}]_{\text{SCSS}}$ for sulfur-rich compositions (Supplemental Table S1). The overall improvement in $r^2$ is, however, very small, increasing from 0.977 to 0.978. An important point to note is that the fitted compositional $A_M$ parameters (Table 2) are identical within uncertainty to those derived by assuming ideal sulfide solution.

Figures 5a and 5b shows the calculated $[\text{S}]_{\text{SCSS}}$ for the MORB composition of Table 4 at saturation in FeS-NiS-CuS$_{0.5}$ sulfide of the indicated composition. In the ideal solution case of Figure 5a, sulfur solubility declines linearly at fixed ratio to $X_{\text{FeS}}$ as discussed above. The non-ideal case of Figure 5b shows little difference from the ideal solution calculation at $X_{\text{FeS}} > 0.5$, but differences between the two increase with decreasing $X_{\text{FeS}}$. At present we consider that the ideal solution model of Table 3 and Figure 5a is adequate for most geologic situations and certainly in those cases of $X_{\text{FeS}} > 0.5$. The important point that we wish to reiterate is that $[\text{S}]_{\text{SCSS}}$ is extremely dependent on sulfide composition and should not be treated as a constant at fixed $P$, $T$, and silicate composition. Supplemental Table 3 comprises a spreadsheet for calculation of $[\text{S}]_{\text{SCSS}}$ using our regressed parameters for both ideal and non-ideal assumptions.

**Application to natural systems**

Based on Fe$^{2+}$/Fe$^{3+}$ measurements the $f_O^2$ of MORB has been estimated to be around that of the fayalite-magnetite-quartz (FMQ) oxygen buffer (Cottrell and Kelley 2013). Under these conditions sulfur will be present dominantly as $S^2–$, suggesting that the Fincham-Richardson relationship should apply to MORB melts. Although chemical and textural evidence (Peach et al. 1990; Patten et al., 2013) suggests that MORBs are sulfide saturated throughout their crystallization histories, most...
previous models for SCSS, imply, in contrast, that MORBs are sulfide undersaturated during much of crystallization (O’Neill and Mavrogenes 2002; Li and Ripley 2005, 2009; Fortin et al. 2015) (Fig. 6a).

To address this apparent discrepancy, we have taken an average of primitive MORB glass compositions (MgO > 9.3 wt%) from Jenner and O’Neill (2012) and determined the liquid line of descent from 1230 to 1170 °C using Petrolog3 (Danyushchevsky and Plechov 2011). We then applied our model for SCSS assuming ideal solution in the immiscible sulfide, a pressure of 0.3 GPa and a Fe/(Fe+Ni+Cu) of 0.7 common to sulfides found in MORB (Patterson et al. 2013; Peach et al. 1990).

The results for SCSS along the liquid line of descent of MORB based on the model presented here agree with the high sulfur contents of MORB glasses measured by Jenner and O’Neill (2012) over the entire range of MgO contents. In contrast, as discussed above, most previous [S]_{SCSS} expressions suggest that, over a significant range of MgO concentrations MORB melts are undersaturated with respect to S (O’Neill and Mavrogenes 1999; Holzheid and Grove 2002; Li and Ripley 2005, 2009; Fortin et al. 2015) as shown in Figure 6a. The two exceptions to this are the models of Mavrogenes and O’Neill (1999) and Holzheid and Grove (2002), which substantially underpredict the S contents of the MORB glasses. Although there is a pronounced cluster of S analyses along our predicted trend at sulfide saturation (Fig. 6) we note that several S concentrations fall below those we calculate. Lower sulfur concentrations are plausibly the result of degassing of the melt during eruption (Fig. 6b). We therefore conclude, based on our measurements that the MORB source region is sulfide saturated and the melts remain at sulfide saturation throughout their crystallization histories.

**Implications and Conclusions**

We have demonstrated that the solubility of sulfur in silicate melt at sulfide saturation [S]_{SCSS} depends, at fixed pressure, temperature, and silicate melt composition, on the composition of the sulfide liquid. This dependence, to a good approximation, leads to [S]_{SCSS} being a linear function of the mole fraction of FeS in the sulfide liquid, defined as \([S]_{FeS} = \text{Fe}/(\text{Fe}+\text{Ni}+\text{Cu})\]. Departures from linear behavior at [S]_{FeS} below ~0.5 are consistent with known non-idealities in the FeS-NiS and FeS-CuS<sub>0.5</sub> liquid systems.

We took both ideal and non-ideal sulfide solution models and fit [S]_{SCSS} to the available data using and equation of similar form to that of O’Neill and Mavrogenes (2002):

\[
\ln [S]_{SCSS} = \frac{\Delta G^0}{R T} + A_M + \sum_{j} X_{j} A_{j} \frac{1}{T} + \ln a_{FeS} - \ln a_{FeO}.
\]

In this equation \(\Delta G^0\) is the standard state free energy change for the reaction:

\[
\text{FeO}_{\text{silicate}} + \frac{1}{2} S_2 = \text{FeS}_{\text{sulfide}} + \frac{1}{2} O_2.
\]

The expression for [S]_{SCSS} incorporates compositional parameters \(A_M\) in terms dependent on \(X_M\), the mole fraction of the oxide of M in the silicate melt on a single cation basis. The activities \(a_{FeS}^{\text{silicate}}\) and \(a_{FeO}^{\text{sulfide}}\) were assumed equal to the mole fractions of FeO in the silicate and FeS in the sulfide, respectively except where the latter was treated as a (non-ideal) ternary symmetrical solution.

We fitted [S]_{SCSS} to the 392 available data points to derive best-fit values of \(A_M\) and the \(A_M\) together with the pressure dependence of \(\Delta G^0\). The standard free energy change \(\Delta G^0\) was adopted from O’Neill and Mavrogenes (2002). Since [S]_{SCSS} is observed to have a positive dependence on temperature while \(\Delta G^0\) decreases with increasing temperature, the \(A_M\) parameters are required to be mostly negative. We tested this conclusion by treating \(\Delta G^0\) and its temperature dependence \(\Delta S^0\) as unknowns and fitted Equation 11, which has these extra unknowns, to the data. As before, the best-fit \(\Delta G^0\) has a negative temperature dependence and the fit parameters are all negative except for a cross-term \(A_{FeS}^{\text{SiFeS}}\), which takes account of a positive correlations between [S]_{SCSS} and the product \(X_{FeS}^{\text{SiFeS}}\).

We find that the cross-term \(A_{FeS}^{\text{SiFeS}}\) is robust in that it has, within uncertainty, the same value (~113 000 K) in all three cases considered: (1) \(\Delta G^0\) treated as a \(P-T\) dependent fit parameter, with ideal FeS-NiS-CuS<sub>0.5</sub> sulfide solution; (2) \(\Delta G^0\) at 1 bar adopted from thermodynamic data and assuming ideal FeS-NiS-CuS<sub>0.5</sub> sulfide solution.
solution; and (3) $\Delta G^\circ$ at 1 bar adopted from thermodynamic data and assuming non-ideal Fe$_7$Ni$_5$Cu$_3$S$_8$ sulfide solution.

The large positive term in $X_{Fe}X_{Ni}$ (on a single cation basis) means that $[S]_{SCSS}$ is predominantly dependent on the FeO content of the silicate melt decreasing (for FeS saturation at 1 GPa, 1400 °C) from 4692 ppm for a Martian basalt with 17.9 wt% FeO to 1084 ppm for Andesite with 7.7 wt% FeO.

In agreement with previous versions of the effects of pressure and temperature on $[S]_{SCSS}$, all natural silicate melt compositions considered exhibit positive dependences on temperature and negative dependences of $[S]_{SCSS}$ on pressure. Our results are, however, in best agreement with those of Fortin et al. (2015).

Finally, application of our results to the MORB glasses analyzed by Jenner and O’Neill (2012) indicates that MORB are sulfide saturated throughout their crystallization paths.

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