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To cite this version:

Fabrice Gaillard, Michel Pichavant, Stephen Macwell, Rémi Champallier, Bruno Scaillet, et al.. Chemical transfer during redox exchanges between H-2 and Fe-bearing silicate melts.. American Mineralogist, Mineralogical Society of America, 2003, 88, pp.(2-3) 308-315. <hal-00069454>

HAL Id: hal-00069454
https://hal-insu.archives-ouvertes.fr/hal-00069454
Submitted on 31 May 2006

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Chemical transfer during redox exchanges between $H_2$ and Fe-bearing silicate melts

FABRICE GAILLARD, MICHEL PICHAVANT, STEPHEN MACKWELL, RÉMI CHAMPALLIER, BRUNO SCAILLET, AND CATHERINE MCCAMMON

INTRODUCTION

In concentrations ranging from parts per million to weight percent, hydrogen is always present in natural minerals or melts in the form of a variety of species with different redox states ($H^+$, $OH^-$, $H_2O$, or $H_2$) (Ingrin and Skogby 2000; Johnson et al. 1994; Schmidt et al. 1998; Stolper 1982). The identification of the nature and the mobility of the different H species is a key for understanding the Earth because H incorporation in mineral or melt dramatically modifies their physical properties. In nominally anhydrous minerals (NAM), the kinetics and mechanisms of H incorporation or extraction have been and are still being studied intensively. Water-derived species incorporation/extraction in NAM recently has been revealed to evolve through a two-stage process (Kohlstedt and Mackwell 1998). The fastest stage was demonstrated to be related to redox processes involving H motion as protons and electronic defects related to $Fe^{2+}$-$Fe^{3+}$ exchange (Kohlstedt and Mackwell 1998; Hercule and Ingrin 1999). Therefore, both the H content and $Fe^{2+}$/Fe$^{3+}$ ratio in NAM are strongly interdependent parameters. The slower stage does not involve redox exchanges and seems to be associated with intrinsic defect mobility (Kohlstedt and Mackwell 1998).

With respect to silicate melts, it is generally accepted that H-bearing species diffuse as $H_2O$ molecules (e.g., Zhang et al. 1991; Behrens and Nowak 1997). However, at low water contents (<0.8 wt%) the mobile water-derived species may be NaH$_2$O$^-$, H$_2$O$^-$, or $H^+$ (Stanton et al. 1990). Schmidt et al. (1998) have also identified the possibility of hydrogen incorporation as $H_2$ molecules in Fe-free silicate melts. In natural melts, which all contain H and Fe, redox interactions occur between $H_2$-$H_2O$ and $Fe^{2+}$-$Fe^{3+}$ (Gaillard et al. 2002). Currently, the mechanisms and the mobilities of the species involved in these interactions are still not known accurately.

Gaillard et al. (2002) studied the kinetics of Fe redox reactions at 2 kb in $H_2O$-rich (5–6 wt%), Fe-poor (1–3 wt% FeO$_{tot}$) melts that occur in response to variations of hydrogen fugacity ($f_{H_2}$). No redox fronts were observed. The kinetic data were interpreted in terms of a two-step reaction mechanism that involves first a virtually instantaneous diffusion of $H_2$ in the sample and then slower structural/chemical reorganizations, involving slower interactions of water-derived species with Fe in the melt. Oxidation-reduction of Fe in these low-Fe, high-$H_2O$ rhyolitic melts is reaction-limited, in contrast to the diffusion-limited process identified by Cooper et al. (1996) for the oxidation of dry basaltic melts.

From these two studies, it is clear that the redox mechanisms in silicate melts strongly depend on the presence or absence of hydrogen (as $H_2$ and OH/$H_2O$). In addition, another major difference concerns the Fe concentration of the studied melts. Both factors make direct comparisons between the two studies difficult.
In this work, reduction experiments have been performed in a near-pure H₂ atmosphere on a nearly anhydrous Fe-rich silicate melt at low pressures to help elucidate the role of H₂ on redox mechanisms in the low-water concentration range. We show that, in contrast to previous observations for Fe-poor and high-H₂O melts, a redox front is clearly identified. We characterize the rate law for the progression of this redox front and the associated chemical mass transfers in the melt. We demonstrate that the process that rate-limits reduction is much slower than rates of molecular H₂ diffusion in the Fe-free system, but is slightly faster than molecular H₂O diffusion. Therefore, this study brings evidence that different water-derived species with different apparent mobilities may exist in silicate melts.

**EXPERIMENTAL TECHNIQUES**

**Starting glass**

Crystal-free, natural peralkaline obsidian was used as a starting material (Eburru, Kenya; MacDonald and Bailey 1973). The composition of the glass, including Fe and Fe₂O₃ (25% of iron as Fe³⁺) and water content (0.25 wt% H₂O), is provided in Table 1. A series of 11 water content determinations, performed by FTIR (see Analyses), on different wafers yielded 0.25 ± 0.01 wt% H₂O, suggesting a homogeneous distribution of water-derived species in the starting glass.

The strong peralkalinity of this obsidian places the glass transition temperature at 400–500 °C, and allows us to perform low-temperature experiments under nearly dry conditions in the melt stability field without crystallization. A dry glass of the albite-orthoclase-quartz (AOQ) system was also used in one experiment as an Fe-free reference (Table 1).

**Materials and strategy**

Cylinders of the Eburru glass were placed in tubes of pure gold (ID = 5 mm, OD = 5.4 mm; the glass cylinder was cored using a drill of 5 mm ID). These assemblies, with one end of the capsule arc-welded and the other end open, were placed in cold-seal pressure vessels at 800 °C under 50 bars of pure H₂ for various durations (see Table 2). One additional experiment was performed using a platinum-capsule with one end sealed by an AOQ glass cap and the other end open to H₂ (Fig. 1a). The AOQ glass cap was formed by melting glass powder in the platinum-capsule at 1400 °C and 1 bar for 6 hours. A cylinder of Eburru glass was then placed in the same capsule in contact with the AOQ glass. This assembly was heated to 1200 °C at 1 bar for 7 minutes to make an airtight contact between the two glasses and between the samples and the capsule wall. The end of the capsule with the AOQ glass was cut so that the AOQ glass was directly exposed to the H₂ gas in the experiment, and thus behaved as a screen between one end of the Eburru glass and the H₂ atmosphere. The other end of the Eburru glass was in direct contact with H₂.

The vessels used in the experiments allow rapid heating and quenching to be performed by mechanically moving the sample rapidly while at pressure between the cool and hot parts of the vessel. Once the experimental temperature was attained (~30 min), the samples were moved to the hot part of the vessel. Using this strategy, each sample was heated from ambient temperatures to 800 °C within less than 5 min. After the experiment, the sample was quenched rapidly by moving it to the cool part of the vessel. Subsequently, the samples assemblies were cut into slices and polished for optical observation, electron microprobe analysis (EMPA), and infrared spectroscopy (FTIR). Wet-chemical analyses and Mössbauer spectroscopy were also performed on selected samples.

**Analyses**

EMPA was performed using an SX-50 Cameca electron microprobe under the following conditions: 15 kV accelerating potential, 6 nA beam current, 10–20 μm beam size, 10 s counting time on element peak positions and 5 s counting time on the background. Multi-element chemical profiles were performed with a 20 μm step increment across the sample.

FTIR analysis was performed on doubly polished samples using a Nicolet 760 Magna spectrometer equipped with an infrared microscope. A CaF₂ beamsplitter was used with a visible light source and liquid N₂ cooled MCT/A detector. The mean beam size was 80–100 μm. Total H₂O content was estimated from the height of the 3600 cm⁻¹ stretching O-H band. The extinction coefficient of the 3600 cm⁻¹ absorption band was calibrated against measurements using Karl-Fisher titration performed on the starting glass (Table 1). The extinction coefficient was determined to be 40 L/g/cm.

Wet-chemical techniques were used to determine the FeO content of selected samples. The method and its precision are detailed in Gaillard et al. (2001). Fe₂O₃ contents were obtained by subtraction of FeO from total Fe analyzed by EMPA. As wet chemistry is a bulk technique, only the samples interpreted as chemically homogeneous based on their uniform color were analyzed (samples Eb/0, Eb/6). In addition, analyses using the Mössbauer milliprobe (McCammon et al. 1991; McCammon 1994) were performed on sample no. Eb/5 with a spot size of ~300 μm to determine Fe³⁺/Fe⁴⁺ in the two regions of the sample with different colors.

**RESULTS**

**Optical observation**

The glasses quenched from all time-series experiments showed a sharp contrast in color, having translucent-green rims and inner regions with the same dark color as the starting glasses (see Fig. 2). The boundary between translucent-green and dark

### Table 1. EMP analyses of the starting Eburru Obsidian and the AOQ glass

<table>
<thead>
<tr>
<th>Sample</th>
<th>SiO₂</th>
<th>TiO₂</th>
<th>Al₂O₃</th>
<th>MgO</th>
<th>FeO</th>
<th>MnO</th>
<th>NiO</th>
<th>CaO</th>
<th>Na₂O</th>
<th>K₂O</th>
<th>SO₂</th>
<th>P₂O₅</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eburru</td>
<td>70.13</td>
<td>0.28</td>
<td>8.16</td>
<td>0.02</td>
<td>6.84</td>
<td>0.23</td>
<td>0.04</td>
<td>0.21</td>
<td>6.48</td>
<td>4.40</td>
<td>0.01</td>
<td>0.02</td>
<td>96.81</td>
</tr>
<tr>
<td>Eburru</td>
<td>FeO = 5.31 wt%, Fe₂O₃ = 1.7 wt%, H₂O = 0.25 wt%</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>AOQ</td>
<td>70.10</td>
<td>0.02</td>
<td>12.10</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>4.40</td>
<td>5.10</td>
<td>0</td>
<td>0</td>
<td>98.75</td>
</tr>
</tbody>
</table>

**Notes:** Wet chemistry was used for FeO and Fe₂O₃ contents and H₂O was determined by Karl Fisher titration. All data are given in wt%.
zones moved toward the inner part of the sample as the experiment duration increased. The rate of progression of this boundary was characterized optically. Table 2 summarizes the 6 observations from the 6 different run durations. After a run of 2916 min (~49 h), the sample was completely translucent-green (Eb/6). Micro-crystals were observed within the glasses after the runs. Using optical techniques we identified feldspar crystals, but these were not analyzed with the EMP.

Fe$^{3+}$/Fe$^{tot}$

Wet chemistry results, expressed in terms of Fe$^{3+}$/Fe$^{tot}$, are presented in Table 2. The starting obsidian (Eb/0) has Fe$^{3+}$/Fe$^{tot}$ = 0.23 ± 0.05, whereas sample no. Eb/6 was characterized as an essentially Fe$^{3+}$-free glass (Fe$^{3+}$/Fe$^{tot}$ = 0.01 ± 0.01). The two micro-Mössbauer analyses were performed on sample no. Eb/5 at the positions shown in Figure 2. The dark zone yielded Fe$^{3+}$/Fe$^{tot}$ = 0.25 ± 0.05, similar to the starting obsidian (Eb/0),
whereas no Fe\(^{3+}\) was measurable in the green zone (Table 2). This observation gives credence to the argument that the difference of color between the green and dark zone indicates a strong difference of Fe redox state. Given the strongly reducing conditions of the experiment and the homogeneity of color within the green zone, we propose that the Fe\(^{3+}\) content is constant in this region and nearly equal to 0 for all runs (Table 2).

### Major elements

EMP chemical profiles were performed on each experimental charge. In the run products no. Eb/1, no. Eb/2, and no. Eb/3, no clear major-element concentration variations were observed across the glasses. Comparisons with the starting glass revealed that Na content decreased by about 7% for all of these run products (Table 2). For samples no. Eb/4 and no. Eb/5, gradients in the concentration of Na at the sample rims were observed (Fig. 3). The maximum in Na concentration located at the gas-melt interface was equal to the Na content of the starting glass. Surprisingly, the position of this chemical gradient does not match that of the redox boundary. Rather, it is located behind the reduction front. Thus, for run no. Eb/5 (Figs. 2 and 3), the reduction front is at 480 μm from the sample surface, whereas the first evidence of Na motion was identified at 150–180 μm. No motion of other major elements was detected by EMPA (Fig. 3). For sample no. Eb/6, the Na\(_2\)O distribution within the sample was homogeneous and matched that of the starting glass.

### H\(_2\)O

FTIR profiles were performed for several samples (see Table 2, and Figs. 2 and 3). In all samples analyzed, we observed water to be present mainly in the form of OH groups due to the dominance of the 4500 cm\(^{-1}\) O-H band over the 5200 cm\(^{-1}\) H\(_2\)O band. The green zone has significantly higher water contents than the dark zone (0.73–0.43 vs. 0.25 wt% H\(_2\)O, respectively). In the dark zone, the water content was similar to that of the starting glass. Between the green and the dark zones, the water content seems to decrease sharply although the spatial resolution of the FTIR analysis does not allow precise determination of the shape of the water profile at this boundary. The H\(_2\)O content of the green zone is not homogeneous. Rather, a maximum H\(_2\)O concentration is reached near the green/dark boundary (0.73–0.43 vs. 0.25 wt% H\(_2\)O, respectively). For the longest experiment, no. Eb/6, the H\(_2\)O content is homogeneous. The uniform H\(_2\)O content of 0.42 wt% is similar to the values measured at the edges of the other samples (Table 2).

### H\(_2\) mobility in melts

Experiment no. Eb/1, where the sample adjoined the AOQ glass, was characterized by growth of reduced green layers that are similar in width on the sides directly exposed to H\(_2\) and in contact with the AOQ glass (Fig. 1). FTIR spectra were col-

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### Table 2. Summary of the run conditions and descriptions of starting and run products

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Eb/0*</th>
<th>Eb/1</th>
<th>Eb/2</th>
<th>Eb/3</th>
<th>Eb/4</th>
<th>Eb/5</th>
<th>Eb/6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Run duration (Minutes)</td>
<td>0</td>
<td>30</td>
<td>65</td>
<td>90</td>
<td>150</td>
<td>180</td>
<td>2916</td>
</tr>
<tr>
<td>Capsule</td>
<td>–</td>
<td>Pt+AOQ</td>
<td>Au</td>
<td>Au</td>
<td>Au</td>
<td>Au</td>
<td>Au</td>
</tr>
<tr>
<td>Optical Advancement of the reduction front in μm</td>
<td>0</td>
<td>180</td>
<td>302</td>
<td>355</td>
<td>430</td>
<td>480</td>
<td>&gt;1800</td>
</tr>
</tbody>
</table>

**Fe\(^{3+}\)/Fe\(_{\text{tot}}\)** (Wet chemical analyzes no. Eb/0 and Eb/6; Mössbauer analyzes no. Eb/5, see figure 2)

| Green zone | none | n.a | n.a | n.a | n.a | 0 | 0 |
| Dark zone | 0.23 | n.a | n.a | n.a | n.a | 0.25 | none |

**FTIR (H\(_2\)O wt%)**

| Green zone | 0.25 | 0.23 | 0.24 | n.a | 0.26 | 0.25 | none |
| Dark zone (homogeneous) | 0.25 | 0.23 | 0.24 | n.a | none |

**Notes:** n.a. = Non analyzed; None = Not visible within the sample.

* Starting glass.
† Position relative to the sample geometry (see figure 2 for the location).
‡ 6 FTIR points were measured on this sample indicating a homogeneous H\(_2\)O content (±0.04 wt%).

---

**Figure 2.** Transmitted light photomicrograph showing the change in color of the glass associated with reduction of Fe\(^{3+}\). The two large circles indicate the location of the Mössbauer milliprobe analyses listed in Table 2. The five small circles illustrate the location of FTIR analyses from Table 2. Charge no. Eb/5.
lected on the AOQ sample before and after the experiment. The water content was about 0.01 wt% before and 0.015 wt% H₂O after the experiment (see Fig. 1). A striking peak at 4115 cm⁻¹ was observed everywhere in the AOQ glass. This peak, which is always observed after annealing under very high f₂O conditions (Schmidt et al. 1998), is due to molecular H₂O dissolved in the glass. This observation establishes unambiguously that, within a very short time, molecular H₂ had permeated the entire AOQ glass whose thickness is about 4 mm. No H₂ peak was observed for the Eburru glass in this experiment.

**DISCUSSION**

In these experiments, several reactions operate as suggested by the decoupling between H₂ mobility, Na migration, and the reduction rate of Fe³⁺. For clarity, hereafter we discuss separately the different processes.

**Constraints on the Fe³⁺ → Fe²⁺ rate constant**

Figure 4 shows a plot of the square of the reduction front position as a function of time. The linear relationship suggests that the reduction of Fe³⁺ to Fe²⁺ is a diffusion-limited process. The calculated slope gives the parabolic rate constant for Fe³⁺ → Fe²⁺ at 800 °C, \( K_{FeO}^{1/2} \rightarrow Fe₂O \) = \( 2 \times 10^{-3} \) m²/s. Chekhmir et al. (1985) also observed the progression of a reduction front in albite melt (claimed to be Mn⁷⁺ → Mn²⁺), which they postulated to be controlled by diffusion and internal decomposition of molecular H₂. We calculated a parabolic rate constant from their single experiment and we find \( K_{MnO}^{1/2} \rightarrow Mn₂O \) = \( 2 \times 10^{-10} \) m²/s at 1000 °C, which seems reasonably consistent with our lower temperature results. However, because Chekhmir et al. (1985) did not perform systematic time-series experiments, the comparison is tenuous. It is interesting to note that, in both studies, the parabolic rate constant is close to but slightly higher than the diffusion of molecular water \([\sim 0.5 \text{ to } 0.8 \log \text{ unit higher, } D_{H₂O} \text{ calculated after Zhang et al. (1991)}])

**Constraints on H₂ mobility from no. Eb/1**

The presence of molecular H₂ in the AOQ glass, and the similarity between the widths of the reduction rim at the H₂-melt and melt-AOQ interfaces, suggest that H₂ has diffused very rapidly through the AOQ melt (run no. Eb/1). We excluded the possibilities of H₂ migration at the AOQ-Eburru boundary because we consider the interface to be welded tightly in the previous anneal. We estimate that if H₂ had taken more than 5 min to cross the 4 mm length of the AOQ cap, we should have been able to see asymmetry in the width of the reduction rim. Therefore, we can calculate a minimum diffusion coefficient for molecular H₂ in the AOQ melt using the relationship \( x = (D·t)^{1/2} \), where \( x \) is the diffusion length (4 mm), \( D \) the diffusion coefficient, and \( t \) the minimum duration required for diffusion across the cap. We estimate \( D_{H₂} \geq 10^{-8} \) m²/s at 800 °C. This value is significantly higher than the diffusion coefficient proposed by Chekhmir et al. (1985) for diffusion of molecular H₂ in amorphous albite under essentially the same conditions.

**Mechanisms of reduction: Ionic vs. molecular migrations**

Macroscopically, reduction of a melt results from an increase of the cation/oxygen ratio. This can be accomplished by a loss of oxygen anions or by incorporation of cations (see Fig. 5 for an illustration of all the likely scenarios). The growth of reduced rim is accompanied by an abrupt increase of the OH content, which is consistent with an incorporation of protons (increase of cation/oxygen ratio). Mechanism 1, shown in Figure 5, involves inward migration of cations as a mirror image of the oxidation-reaction path operating under dry conditions (Cooper et al. 1996). As no migration of divalent cations was observed, this mechanism does not operate here.

From a microscopic point of view, there are several ways to incorporate these protons into the Eburru melt. If molecular hydrogen (H₂) solubility in the melt is sufficiently high, proton incorporation can be achieved by H₂ dissolution at the vapor-melt boundary together with H₂ diffusion within the melt plus H₂ breakdown by reduction of Fe³⁺ at the reduction front, resulting in the formation of two Fe²⁺ and two protons (see Fig. 5). In a peralkaline melt such as Eburru, Fe³⁺ (in tetrahedral coordination) is charge-compensated structurally by alkalis (Dickenson and Hess 1986). Therefore, to account for the in-
creases of OH content accompanying reduction of Fe$^{3+}$ by H$_2$, we write [in the notation of Hess (1980)]:

$$
H_2 + 2NaFe^{3+}O_2 + 2SiOSi \rightarrow 2NaOH + 4(Fe^{2+})_{Si}OSi
$$

accompanied by a structural partitioning of water:

$$
NaOH + SiOSi \rightarrow NaOSi + HOSi.
$$

If the rate of reduction is controlled by solubilization of molecular H$_2$ at the vapor-melt interface, the growth of the reacted layer should depend linearly on time. However, as the process shows a parabolic dependence on time, diffusion within the melt must be rate limiting. Diffusion of molecular H$_2$ in AOQ is 3 to 4 orders of magnitude faster than the growth rate of the reduction rim in Eburru. Either: (1) H$_2$ diffusion in the AOQ and Eburru melts are very different; (2) H$_2$ diffusion is rate-limiting, but additional processes delay the reduction rate of the Eburru melt; or (3) H$_2$ is not directly involved in the reaction and diffusion of H$_2$ is not rate limiting.

In the first case, no specific investigation on the compositional dependence of $D_{H_2}$ is available. Hence, we cannot argue in favor of variable diffusion rates for hydrogen. However, many studies devoted to diffusion properties of neutral molecules in silicate melts [He, Ne, Ar, H$_2$O... see Watson (1994)] showed that the smaller the molecule is, the weaker is the compositional dependence of the diffusion coefficient. For He whose size is close to that of H$_2$, $D_{He}$ varies by much less than one log unit between basalt and rhyolite (Watson 1994) whereas we observed a difference of 3 log unit between $D_{H_2}$ in AOQ and the reaction rate in Eburru. We therefore consider that even if $D_{H_2}$ can be different in the AOQ and Eburru melts, this difference cannot explain the contrast in velocity between $D_{H_2}$ in AOQ and the reaction rate in Eburru.

For the second case (Fig. 5, mechanism 2), Crank (1975) gave an approximate solution for the growth of the reaction layer controlled by the dissolution rate of a reactive gas.

$$
X = [(2D_{H_2}S_{H_2}t)/(2C_{Fe^{3+}})]^{1/2}
$$

where $D$, $S$, and $C$ are, respectively, the diffusion coefficient, solubility, and concentration of the subscripted species, and $X$ gives the thickness of the reacted layer as a function of time, $t$. This equation may provide an explanation for the difference between the reaction rate and the diffusivity of the mobile species. In the present case, if the ratio $S_{H_2}/2C_{Fe^{3+}} << 1$ but still large enough that transport of molecular H$_2$ dominates the reduction process, the growth of the reduced rim can be much slower than diffusion of H$_2$ while still obeying a diffusion-lim-
ite law. The lack of an H₂ signature in the infrared spectra of the Eburru glass is consistent with this solubility criterion.

In the third case, if we consider that molecular H₂ is so insoluble in the melt that there are insufficient H₂ species available to reduce the Fe³⁺, other water-derived species must be rate limiting. Thus, molecular H₂ may react with Fe³⁺ to produce Fe²⁺ and OH groups at the gas-melt interface, as shown in Equation 1. Then the redox potential is conveyed toward the interior of the sample by coupled transport of protons and electronic species (see Fig. 5, mechanism 3). In Equation 1, H₂ should then be replaced by H⁺ and electrons e⁻ or electron holes h⁺. Because Fe-bearing glasses and melts are polaron-type semiconductors (Cooper et al. 1996, and references therein), such migration of cations in a redox gradient is charge-compensated by electronic migration. Diopside (Hercule and Ingrин 1999) and olivine crystals (Kohlstedt and Mackwell 1998) have been shown to facilitate incorporation of protons coupled to outward migration of electron holes caused by reduction of Fe³⁺. In that case, the growth rate of the reduced rim is controlled either by proton or electronic mobilities. According to Schmalzried (1981), the width of the reacting layer is directly proportional to the square root of time, t, to the diffusion coefficient of the rate-limiting species i (H⁺ or electronic species), and to the Gibbs free energy of the ongoing reaction normalized by the temperature, T:

\[ X = \left[ \frac{2 D_i \Delta G}{(RT) t} \right]^{1/2} \]  

(4)

In general, proton diffusion in SiO₂-rich silicate melts is not regarded as a likely mechanism (Zhang et al. 1991; Nowak and Behrens 1997). Although Behrens and Nowak (1997) ruled out the possibility of proton migration in Fe-free SiO₂-rich melt, incorporation of Fe may introduce electron holes (Fe⁴⁺) that provide charge compensation for proton jumps. Also, according to Stanton et al. (1990) (see also the review of Dingwell 1995), a mobile, positively charged H-bearing species can dominate the transport of water-derived species in low-H₂O melts (<0.8 wt% H₂O; Stanton et al. 1990). This mobile, water-derived species can be NaH₂O³⁻, NaH₂O⁻, or H⁺. We can exclude NaH₂O in our experiments because the chemical profile for Na does not extend all the way to the redox front. A massbalance calculation, based on the Mössbauer data (Fe³⁺/Fe₄⁰⁺) and FTIR spectra (OH) close to the reduction front, indicates variation of Fe³⁺ and OH concentration that reasonably matches with proton rather than H₂O⁻ incorporation (one OH band created for the reduction of one Fe³⁺). Therefore, the parabolic rate constant extracted from Figure 4 should reflect either proton or electron migration rates, depending on which is the slowest.

Na migration

The migration of Na seems to be decoupled kinetically from the redox interaction between H₂ and iron. A rapid decrease in the Na content (of ~7%) is observed within the first 30 min of exposure to an H₂ gas. We interpret these results as the consequence of the redox partitioning of Na between melt and gas already observed under H-free conditions (see Georges et al. 2000). The fₒ₂ during the beginning of the runs is very reducing so that it causes a displacement toward the RHS of the following equilibria:

\[ \text{Na}_2\text{O}_{\text{melt}} \leftrightarrow \text{Na}_{\text{gas}}^{\text{gas}} + \frac{1}{2}\text{O}_2^{\text{gas}} \]  

(5)

As we did not observe any Na gradient in sample no. Eb/1, the above equilibria should be achieved very rapidly consistent with very high Na diffusion rate in peralkaline melts (Henderson et al. 1985). Molecular O₂⁻ is not stable, as H₂O will be produced instead at the melt/gas interface by oxidation of H₂ by Na₂O. This oxidation may effect a small, transient change in the fₒ₂ of the gas near the surface of the melt. In sample no. Eb/4 and no. Eb/5 (Fig. 3), as well as dehydration of the melt, we observed chemical migration of Na. The release of water into the gas phase should increase the fₒ₂/fₒ₂ ratio and therefore increase the fₒ₂ in the gas. Equation 5 is therefore displaced to its LHS and Na migration from the gas toward the melt occurs. Figure 3 shows that the position of both Na migration and dehydration fronts nearly corresponds to the calculated position of a front of molecular H₂O migration [calculated after Zhang et al. (1991), see caption of Fig. 3]. Therefore, we anticipate that migration of Na from the gas to the melt operated at a rate controlled by melt dehydration (i.e., H₂O mobility) and not by Na diffusion in the melt.

Comparison with previous work

The kinetics of Fe oxidation-reduction in anhydrous basaltic melts has been demonstrated to be rate-limited by diffusion of divalent cations (Cooper et al. 1996; Cook and Cooper 2000). In contrast, we provide clear evidence that, for low H₂O melts, molecular hydrogen or coupled proton-electronic species are the mobile species controlling the advance of the reduction front. The kinetics of change in Fe³⁺/Fe²⁺ is thus much faster in H-bearing systems than in anhydrous systems (representing the several orders of magnitude difference between Ca²⁺ and H₂, H⁺ diffusion rates). Given that H-bearing species are always present in natural magmas, modification of the Fe redox state is most likely to occur following the mechanisms we have identified in this study. It should be noted, however, that for high-H₂O, low-Fe melts, Gaillard et al. (2002) have not detected any coupled proton/electronic species motion. The reason for this discrepancy may be due to reduced electronic conductivity in Fe-poor melts and/or high concentration of water in the melts that may affect both diffusion and solubility of H₂.

CONCLUDING REMARKS AND PERSPECTIVES

The major points from this study are: (1) reduction of Fe³⁺ in the melt by H₂ operates by the progression of a redox front that is accompanied by an increase in the concentration of OH-groups; (2) H₂ diffusion in silicate melts is much more rapid than the observed rate of reduction of Fe³⁺; (3) the velocity of the redox front within the melt is faster than the mobility of molecular H₂O, suggesting that various hydrogen-bearing species with different diffusivities may exist in a silicate melt; and (4) exchange of Na between the gas and the melt occur at a very high rate and are extremely sensitive to the redox conditions.

Understanding the response of a magma to changes in its environment, such as during degassing or exchanges during mixing or interaction with surrounding solids requires the...
knowledge of the identity and the mobility of all H-derived species. In this study, we have shown that redox exchanges between H and Fe causes incorporation of OH groups at a rate different from H2O migration. We have also shown that the H content and Fe3+/Fe2+ ratios in melt are strongly correlated. We should thus question the robustness of Fe3+/Fe2+ of melts as an indicator of pre-eruptive oxygen fugacity because these melts may have undergone degassing (Christie et al. 1986). Further studies, however, are clearly needed to quantify the relationship between Fe redox state and H incorporation/extraction for a range of melts compositions and experimental conditions that permit direct comparison with natural magma processes.

ACKNOWLEDGMENTS

Interpretations of the experimental data were greatly enhanced by discussions with Reid Cooper and Rebecca Everman. Julian Mecklenburgh made a useful informal review of the final version. Ray MacDonald, who kindly supplied the Eburru glass, is also thanked. Hans Keppler and Max Wilke are acknowledged for their reviews. Mike Toplis is greatly thanked for the editorial work.

REFERENCES CITED


MANUSCRIPT RECEIVED NOVEMBER 6, 2001
MANUSCRIPT ACCEPTED OCTOBER 8, 2002
MANUSCRIPT HANDLED BY MICHAEL TOPLIS