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Kinetics of iron oxidation-reduction in hydrous silicic melts

FABRICE GAILLARD,* BRUNO SCAILLET, AND MICHEL PICHAVENT

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ABSTRACT

The kinetics of Fe oxidation-reduction in two hydrous rhyolitic melts, one metaluminous and the other peralkaline, have been studied at 800 °C, 2kb, for melt water contents from ~5 wt% to saturation and fO₂ between NNO-2 and NNO + 3 (NNO = nickel-nickel oxide redox buffer). The metaluminous melt (~1 wt% FeO₂) reached redox equilibrium after 10 hours and the peralkaline one (~3 wt% FeO₂) after 3 hours. The kinetics of Fe oxidation and reduction are similar and unaffected by the presence or absence of a hydrous fluid phase. No redox front is observable in the glass as the Fe³⁺/Fe²⁺ ratio evolves, implying that the Fe redox kinetics in hydrous silicic melts is rate-limited neither by the diffusion of H₂ nor by the mobilities of divalent cations, as observed for anhydrous basaltic melts. We propose a two-step reaction mechanism that involves: (1) virtually instantaneous diffusion of H₂ in the sample, followed by (2) slower structural/chemical reorganizations around Fe atoms. The overall redox process involving iron and hydrogen in Fe-poor, H₂O-rich melts is thus reaction-limited and obeys a first-order logarithmic rate law. The relatively slow kinetics of oxidation/reduction explains why melt Fe³⁺/Fe²⁺ can be readily quenched in laboratory experiments. Simulation of oxidation of magmas due to H₂ exchange with wall rocks is performed using these new kinetics laws and two ΔfG values extracted from the literature. We demonstrate that the metasulminate composition is not significantly modified whereas the peralkaline composition undergoes important and fast changes of Fe³⁺/Fe²⁺ during short processes such as ascent prior to Plinian-style eruptions.

INTRODUCTION

Because Fe is the most abundant heterovalent component of terrestrial magmas, the Fe³⁺/Fe²⁺ ratio of melts or glasses is commonly used as a measure of redox conditions or oxygen fugacity (fO₂). To constrain fO₂ precisely from glass Fe³⁺/Fe²⁺, numerous experimental studies have characterized the dependence of silicate melt Fe³⁺/Fe²⁺ on fO₂, melt composition, T, and P (Sack et al. 1980; Kilinc et al. 1983; Mysen and Virgo 1989; Kress and Carmichael 1991; Moore et al. 1995; Nikolaev et al. 1996; Baker and Rutherford 1996; Gaillard et al. 2001). The Fe-redox ratio of erupted lavas is commonly regarded as a window on the redox state of the magma source region (Carmichael 1991). However, the oxidation state of Fe in natural silicate melts may also reflect the interplay of numerous complex processes such as interaction with C-O-H-S fluids (Brandon and Draper 1996), change of melt composition, crystallization, and chemical exchange with host rocks. The ability of these different processes to affect the melt Fe³⁺/Fe²⁺ ratio depends on, among other factors, their kinetics. So, to understand and model the redox behavior of natural magmas, kinetic constraints on Fe³⁺-Fe²⁺ reaction in silicate melts are required. These kinetic constraints must involve redox couples and cover conditions that reflect those of the Earth’s interior.

Currently, experimental studies aimed at characterizing the kinetics of the Fe-redox equilibrium have been conducted at 1 atm, on H-free melts (Naney and Swanson 1984; Wendlandt 1991; Cooper et al. 1996; Cook and Cooper 2000). Under these conditions, the redox half-couples involved are O₂/O²− or CO₂/C, and Fe³⁺/Fe²⁺. Cooper et al. (1996) have demonstrated that Fe²⁺ oxidation is rate-limited by the mobilities of divalent cations and electron holes. The mobile cations, which include Ca, Mg, and Fe, migrate from the oxidizing front toward the glass-air interface, decreasing locally the cation/oxygen ratio, whereas a second wave composed of univalent cations (mainly Na) migrates from the reduced inner part of the sample toward the oxidizing front. As stressed by Cook and Cooper (2000), because O²− ions are relatively immobile in silicate melts, the fastest reaction path acting to level out redox gradients in dry silicate melts involves cation diffusion toward the glass-air interface.

The mechanisms described above may apply to the oxidation of Fe-bearing melts at the Earth’s surface where O₂ is a major component. However, in the Earth’s interior, free O₂ is almost negligible amounts. Furthermore, most terrestrial magmas contain C-O-H-S volatile phases (Stolper 1982b; Blank and Brooker 1994), which introduce additional redox half-couples (H₂/H₂O, S²−/S⁶⁺, CO/CO₂; Candela 1986; Baker and Rutherford 1996; Moore et al. 1995; Gaillard et al. 2001). Water is by far the most abundant volatile species in magmas (Johnson et al. 1994), so that the two main redox half-couples in hydrous silicate melts are H₂/H₂O and FeO/FeO₂. In such systems, the mobilities of H₂ and H₂O (or other H-bearing species) and of electron holes are most likely to dominate the kinetics and mechanisms of Fe oxidation-reduction. H₂O mobility has been investigated mostly in low-Fe rhyolitic melts. However, the relationship between H₂O diffusion and Fe³⁺/Fe²⁺ has not yet been elucidated (Zhang et al. 1991).

First-order constraints on H₂ diffusivity in silicate melts were
provided by Chekhmir et al. (1985). They concluded that H₂ is, by far, the fastest H-bearing diffusing species in melts (\(D_{H2} \sim 2.5 \log \text{units faster than } D_{O2}\)). The work of Chekhmir et al. (1985) involved H₂ diffusion in either graphite- or Mn-doped nominally anhydrous albite melts. No equivalent information is presently available for Fe-bearing melts. In this paper, we present the results of an experimental study aimed at determining the kinetics of Fe oxidation-reduction in hydrous rhyolitic melts. We demonstrate in particular that the mechanism of Fe oxidation-reduction in water-bearing melts does not result from the progression of a redox front, in strong contrast to what is observed at 1 atm in air.

**EXPERIMENTAL AND ANALYTICAL METHODS**

Starting products

The experiments were performed on two rhyolitic compositions. The first one is a synthetic glass having the composition of the matrix glass of the June 15, 1991, Mt. Pinatubo dacite (Scaillet and Evans 1999; Gaillard et al. 2001). This glass was synthesized using a standard gel method (see Pichavant 1987). The gel was melted at 1400 °C, 1 bar in air (\(f_{O2} = 0.2 \text{ atm}\)) and then quenched to a glass, which was analyzed with the electron microprobe. The glass is slightly peralkaline (Table 1). It contains 1 wt% FeO (total iron expressed as FeO) and has a FeO concentration (analyzed by titration) of 0.34 wt%, yielding a molar FeO/FeO ratio of 0.97. The second sample studied is a nearly aphryic natural peralkaline obsidian from Ascension Island (Harris 1983) that contains ~3 wt% FeO (Table 1). To test the influence of the Fe⁴⁺/Fe⁵⁺ ratio of the starting glass, both reduced and oxidized glasses were prepared from the Ascension obsidian. The reduced glass was obtained by reacting the powdered Ascension glass under pure H₂ at 800 °C during 6 hours (Fe₂O₃/FeO = 0.09, Table 2). For the oxidized glass, the obsidian was held at 1300 °C in air during 3 hours (Fe₂O₃/FeO = 1.54, Table 2). Glasses ground in a agate mortar to a mesh size of 20 μm were loaded in either Au or Ag₉₄Pd₆ capsules (i.d. = 2.5 mm; o.d. = 2.9 mm) together with distilled and deionized water. Most experimental charges consisted of ~80 mg of glass plus 6.5 μL of H₂O, which corresponds to water-saturated conditions at 800 °C and 2 kbar. For three charges, the amount of H₂O added was 4.5 μL, corresponding to water-undersaturated conditions. Each capsule was subsequently welded shut with a graphite arc-welder and placed inside the pressure vessel.

**Experimental apparatus**

All experiments were performed in three René 41 cold-seal pressure vessels (CSPV) equipped with a modified semi-permeable H₂ Shaw-type membrane made of an Ag₉₄Pd₆ alloy (Scaillet et al. 1992; Schmidt et al. 1995). Temperature was monitored using unsheathed external type-K thermocouples. Each vessel-furnace pair was calibrated under pressure (see the method of Pichavant 1987) using internal sheathed dual type-K thermocouples calibrated against the melting points of NaCl and LiCl (Schmidt et al. 1995). Overall, the temperature is known to within ±7 °C. Total pressure (~2 kbar in all experiments) was measured by a high-pressure transducer (Asco Instrument PR 851), calibrated against a 7 kbar Heise tube gauge, and is known within ±20 bar. The H₂ membrane and line are connected to a H₂ tank allowing H₂ pressures up to 70 bar to be applied. Conversely, low H₂O pressures were measured with a Bourdon XM801 transducer calibrated against a Protai tube gauge and are known to within ±0.1 bar.

In practice, the vessel is first loaded with a given pressure of Ar. A known H₂ fugacity is then applied to the H₂ membrane and line, and temperature is increased up to 800 °C. The duration of the heating period is about 20 mn. During that period, H₂ starts to transfer across the H₂ semi-permeable membrane toward the vessel. At equilibrium, usually attained a few minutes after thermal equilibration (see below for more information on the kinetics of H₂ transfer from membrane to capsule),

| Table 1. Electron microprobe analysis of the starting glasses* |
|-----------------|-----------------|-----------------|
|                  | Pinatubo matrix glass | Ascension rhyolite |
| SiO₂             | 78.44            | 73.57           |
| Al₂O₃            | 12.59            | 12.32           |
| FeO              | 1.00             | 3.21            |
| MnO              | 0.03             | 0.09            |
| MgO              | 0.21             | 0.21            |
| CaO              | 1.33             | 0.32            |
| Na₂O             | 3.31             | 5.54            |
| K₂O              | 2.89             | 4.68            |
| TiO₂             | 0.17             | 0.24            |
| P₂O₅             | 0.02             | 0.03            |
| Total†           | 99.91            | 99.5            |
| A/CNK‡           | 0.95             | 0.80            |

* Compositions recalculated on an anhydrous basis.
† Original total is reported.
‡ A/CNK = Al₂O₃/(Na₂O + K₂O + CaO) in moles.

**Table 2. Time series experiments on the Ascension composition**

<table>
<thead>
<tr>
<th>Starting products</th>
<th>Reduced</th>
<th>Oxidized</th>
<th>Reduced</th>
<th>Oxidized</th>
<th>Reduced</th>
<th>Oxidized</th>
<th>Reduced</th>
<th>Oxidized</th>
<th>Reduced</th>
<th>Oxidized</th>
<th>Reduced</th>
<th>Oxidized</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duration (h)</td>
<td>0</td>
<td>0</td>
<td>3</td>
<td>3</td>
<td>7</td>
<td>7</td>
<td>24</td>
<td>24</td>
<td>48</td>
<td>48</td>
<td>96</td>
<td>96</td>
</tr>
<tr>
<td>H₂O*</td>
<td>-0</td>
<td>0</td>
<td>4.68</td>
<td>5.00</td>
<td>5.62</td>
<td>6.05</td>
<td>6.85</td>
<td>6.81</td>
<td>6.67</td>
<td>6.85</td>
<td>6.37</td>
<td>6.41</td>
</tr>
<tr>
<td>FeO†</td>
<td>3.21</td>
<td>3.21</td>
<td>2.45</td>
<td>2.25</td>
<td>2.47</td>
<td>2.33</td>
<td>2.60</td>
<td>2.40</td>
<td>2.71</td>
<td>2.55</td>
<td>3.07</td>
<td>3.11</td>
</tr>
<tr>
<td>FeO‡</td>
<td>2.80</td>
<td>0.82</td>
<td>1.90</td>
<td>1.45</td>
<td>1.77</td>
<td>1.64</td>
<td>1.85</td>
<td>1.73</td>
<td>1.90</td>
<td>1.81</td>
<td>2.00</td>
<td>2.05</td>
</tr>
<tr>
<td>FeO³§</td>
<td>2.73</td>
<td>0.79</td>
<td>1.69</td>
<td>1.17</td>
<td>1.57</td>
<td>1.39</td>
<td>1.69</td>
<td>1.51</td>
<td>1.78</td>
<td>1.64</td>
<td>2.00</td>
<td>2.05</td>
</tr>
<tr>
<td>(X_{FeO}^{liq}/X_{FeO}^{cr})</td>
<td>0.09</td>
<td>1.54</td>
<td>0.22</td>
<td>0.46</td>
<td>0.34</td>
<td>0.27</td>
<td>0.30</td>
<td>0.26</td>
<td>0.28</td>
<td>0.27</td>
<td>0.26</td>
<td></td>
</tr>
</tbody>
</table>

* Note: All experiments were performed at \(P = 2 \text{ kb}, T = 800 °C, t_o = 12 \text{ bar}, \text{ and water saturation.}
† Water in the glass determined by difference on 100% of electron microprobe analysis (EMPA).
‡ Total iron in the glass expressed as wt% FeO determined by EMPA.
§ Bulk FeO (wt%) analyzed by wet chemistry.
∥ Molar Fe⁴⁺/Fe⁵⁺ ratio of the glass. Fe₂O₃ calculated as (FeO₃·FeO³). 1.1113
Equation 2 shows that, at constant temperature and pressure, \( f_{\text{O}_2} \) is directly related to \( f_{\text{H}_2} \) if \( f_{\text{H}_2O} \) is constant. In our experiments, a steady \( f_{\text{H}_2} \) can be imposed and maintained during long durations because the Shaw membrane and H\(_2\) line have a large volume (46 cm\(^3\)) compared with the free volume of the vessel (~8 cm\(^3\)). Therefore, the H\(_2\) membrane acts as an H\(_2\) buffer. Another important aspect is that our experimental setup allows \( f_{\text{H}_2} \) to be varied in situ, at constant temperature and under nearly constant \( P_{\text{total}} \) (changing \( f_{\text{H}_2} \) in the membrane slightly changes pressures), which allows either reduction or oxidation cycles to be performed.

Quenching was performed by removing the vessel from the furnace and is relatively slow (about 3–5 °C/s). Gaillard et al. (2001) have demonstrated that glasses quenched at 300 °C/s and at 0.5–1 °C/s have the same Fe-redox ratios. Therefore, no modification of the Fe\(^{3+}/\text{Fe}^{2+}\) ratio is expected at the quench rates used in this study, and the Fe-redox ratios measured in quenched glasses reflect the speciation of Fe in the melts at 800 °C and 2 kbar.

**Analytical methods**

Major element compositions of experimental products (glasses, magnetite, hematite, and plagioclase crystals) were obtained with a Cameca-Camebax electron microprobe at the CNRS-BRGM-UO laboratories at Orléans. For crystals, the operating analytical conditions were: 15 kV accelerating voltage, 6 nA sample current, 10 s counting time on peak, and a beam diameter of 1–2 μm. For glasses, a beam diameter of 10 to 25 μm was used to minimize the migration of alkalies (e.g., Pichavant 1987; Devine et al. 1995). Concentrations of alkalies were corrected using secondary hydrous glass standards. Water concentrations of glass were estimated using the by-difference method (Devine et al. 1995) employing, as secondary standards, a set of 4 hydrous rhyolitic glasses whose water contents (2.0–6.38 wt% H\(_2\)O) have been measured by Karl-Fischer titration.

For selected samples (numbers 1–5), the homogeneity of water concentrations (OH+H\(_2\)O) was checked using a Nicolet 760 Magna FTIR spectrometer on doubly polished glass wafers. Four to five spectra were collected for each analyzed glass using a beam diameter of ~100 μm. Analytical procedures are similar to those described in Gaillard et al. (2001).

A wet-chemical technique was used to determine the FeO iron content of experimental products. Glass chips totaling 50–60 mg, ground under acetone to 10–20 μm mesh size, were used for each analysis. FeO iron was determined by titration with K\(_2\)Cr\(_2\)O\(_7\). The uncertainty in FeO titration, calculated through standard error propagation, ranges from 0.04 to 0.08 wt% depending on the FeO content of the sample [see Gaillard et al. (2001) for details on the reproducibility and accuracy of the technique].

\[ H_2 + 1/2 O_2 \leftrightarrow H_2O \quad (1) \]

where

\[ 1/2 \log f_{O_2} = \log f_{H_2O} - \log f_{H_2} - \log K_{H_2} \quad (2) \]

**Experimental strategy**

In this study, both constant and variable \( f_{\text{H}_2} \) experiments were performed at 800 °C and 2 kbar total pressure.

**Constant \( f_{\text{H}_2} \) experiments**

Ten time-series experiments were performed with a \( f_{\text{H}_2} \) of 12 bar (~NNO-0.2) using both the oxidized and reduced Ascension glasses (Table 2). These experiments provide a test of reversibility and serve to establish the kinetics of attainment of steady-state glass Fe\(^{3+}/\text{Fe}^{2+}\) ratios (interpreted as equilibrium value) for a given \( f_{\text{H}_2} \). In addition, four experiments were carried out on the Ascension obsidian for \( f_{\text{H}_2} \) between 50 and 1.8 bar, and five on the Pinatubo glass for \( f_{\text{H}_2} \) between 50 and 1 bar (Table 3). These experiments serve to establish the equilibrium relationship between glass Fe\(^{3+}/\text{Fe}^{2+}\) and \( f_{\text{H}_2} \) at constant \( T, P \) and \( f_{\text{H}_2O} \). To avoid magnetite crystallization observed in the time-series experiments, these experiments were systematically started by annealing at \( f_{\text{H}_2} = 50 \) bar (~NNO-1.5, see Table 3) for 48 hours. Then, \( f_{\text{H}_2} \) was brought to the desired value and conditions were kept constant for another 24 to 48 hours (Table 3).

**Variable \( f_{\text{H}_2} \) experiments (oxidation/reduction cycles)**

In variable \( f_{\text{H}_2} \) experiments, either oxidation or reduction cycles were performed in situ (i.e., during a given experiment) and the Fe-redox ratio of the quenched melt was monitored as a function of time. For oxidation cycles, the experiments were started by an annealing step at \( f_{\text{H}_2} = 50 \) bar for 48 hours. Then, \( f_{\text{H}_2} \) was dropped in the membrane and H\(_2\) line by evacuating H\(_2\) with the vacuum pump (\( f_{\text{H}_2} < 0.1 \) bar) while the temperature was kept constant. The experiments were then maintained under these new conditions for durations ranging between 0.1 and 27.5 hours, and then quenched (Table 4). Under our experimental \( P-T \) conditions, changing \( f_{\text{H}_2} \) from 50 to <0.1 bar corresponds to a change in \( f_{\text{H}_2} \) from NNO-1.5 to ~NNO+3. For reduction cycles (performed on the Pinatubo composition only, Table 4), after the initial annealing step at \( f_{\text{H}_2} = 50 \) bar, H\(_2\) was evacuated and a \( f_{\text{H}_2} < 0.1 \) bar was maintained for a duration known to produce a crystal-free oxidized melt on the basis of the previous results (Table 4). Then, a \( f_{\text{H}_2} \) of 50 bar was re-established within the Shaw membrane and H\(_2\) line. Conditions were kept constant for durations of between 0.66 and 6.25 hours, and the experiments were quenched (Table 4).

It must be stressed that the charge is not affected instantaneously by the \( f_{\text{H}_2} \) change that results from varying \( f_{\text{H}_2} \) in the membrane and H\(_2\) line. The rate of change of redox conditions inside the capsule is controlled by H\(_2\) permeability in noble metals (both capsule and membrane) and also by the free volume of the vessel. We have calculated (see Scaillet et al. 1992) the time necessary for H\(_2\) to equilibrate in the Au capsule following an isothermal \( f_{\text{H}_2} \) drop in the membrane such as imposed in the oxidation cycles. Results range between 4 and 7 min depending on the method of estimation of the number of moles of H\(_2\) to be transferred. This time-scale is a much shorter than that required to attain a steady-state melt Fe-redox ratio under fixed \( f_{\text{H}_2} \) (see below). However, this time-scale is of the same order of magnitude as the minimum duration necessary to observe detectable changes in melt Fe\(^{3+}/\text{Fe}^{2+}\) ratios after a change in \( f_{\text{H}_2} \) (see below). Therefore, experiments with differ-
ent capsule materials have been performed to test the effect of the rate of \( \text{H}_2 \) transfer.

**RESULTS**

**Constant \( f_{\text{H}_2} \) (equilibrium) experiments**

Results for the time-series experiments are detailed in Table 2. Magnetite is present in all experiments with durations of \(<48\) hours, irrespective of the redox state of the starting glass (either oxidized or reduced). In the two longest experiments from this series (96 hours), magnetite is not found. This magnetite thus nucleates and grows during heating-up but is not stable at the final \( P-T-f_{\text{H}_2}-f_{\text{H}_2O} \) conditions. The FeO content of the glass is obtained from the bulk FeO content of the charge (analyzed by titration) by subtracting the contribution of magnetite whose composition is known from electron microprobe analysis and whose proportion in run products is calculated by mass balance (Table 2). Results show that Fe-redox equilibrium is closely approached in the melt after experimental durations of \(~10\) hours (Fig. 1). A steady-state glass \( \frac{\text{Fe}^{3+}}{\text{Fe}^{2+}} \) ratio is attained after \(~20\) hours. The equilibrium glass \( \frac{\text{Fe}^{3+}}{\text{Fe}^{2+}} \) is approached from both sides (i.e., oxidized and reduced) and at similar rates.

Results of the equilibrium experiments are presented as a function of \( f_{\text{H}_2} \) and \( f_{\text{O}_2} \) in Table 3. All data concern crystal-free
glasses. No Fe loss from sample to capsule was detected (see total FeO contents in Table 3). Glasses have homogeneous major element compositions and water contents. Observation of the samples under an optical microscope shows that the color of the glasses is systematically related to its Fe-redox ratio. For the Ascension composition, glasses synthesized at \( f_{O_2} < NNO \) are green-blue and progressively change their color to brown-black for \( f_{O_2} > NNO \). For the Pinatubo composition, glasses synthesized at \( f_{O_2} < NNO \) are transparent and progressively change their color to black for \( f_{O_2} > NNO \). Analyses of different chips of the same homogeneously colored glasses (synthesized from both the Ascension and Pinatubo compositions) were performed by Gaillard et al. (2001). They yielded identical FeO contents, demonstrating that color homogeneity implies constant glass \( Fe^{3+}/Fe^{2+} \) ratio. For all samples of this study, the homogeneity of the Fe-redox state is therefore inferred from the lack of color variation across the sample. Note that experimental durations at the equilibrium \( f_{H_2} \) (i.e., following the annealing step at \( f_{H_2} = 50 \) bar) range between 22 and 30 hours (Table 3), and are therefore longer than durations necessary for attainment of steady-state \( Fe^{3+}/Fe^{2+} \) ratios (Fig. 1).

Glass \( Fe_2O_3/FeO \) values are plotted vs. \( f_{O_2} \) and \( f_{H_2} \) in Figure 2. For both starting samples, glass \( Fe_2O_3/FeO \) progressively decreases with increasing \( f_{H_2} \) (or decreasing \( f_{O_2} \)). For a given \( f_{H_2} \), glass \( Fe_2O_3/FeO \) differ between the two compositions studied. The experimental data from this study are compared with the empirical equation of Kress and Carmichael (1991) in Figure 2a. As observed in previous studies (Baker and Rutherford 1996; Gaillard et al. 2001), there are significant differences between the measured and calculated glass \( Fe_2O_3/FeO \), particularly for the Ascension composition in the present case.

### Variable \( f_{H_2} \) experiments (oxidation-reduction cycles)

Results of these experiments are detailed in Tables 4–5. Charges are generally crystal-free, except for the long-duration oxidation experiments (Table 4). As in the previous series, glasses are chemically homogeneous (major-element compositions and water contents, Table 4). One critical point to be stressed is the absence of any color gradient in the glasses. This feature was checked carefully through observation of doubly polished glass wafers across the entire sample section. This homogeneity is true even for glasses that have \( Fe_2O_3/FeO \) different from equilibrium values (the case for most glasses from FIGURE 2).

### Table 5. Results of a reduction cycle at 800 °C and 2 kb performed on the Pinatubo composition

<table>
<thead>
<tr>
<th>Run number</th>
<th>( f_{H_2} ) membrane (in bar)</th>
<th>Duration at ( f_{H_2} \sim 55 ) bar (min)</th>
<th>( FeO ) (wt%)†</th>
<th>( Fe_2O_3 ) (wt%)‡</th>
<th>( Fe_3O_4 ) (wt%)§</th>
<th>Phases</th>
</tr>
</thead>
<tbody>
<tr>
<td>23</td>
<td>&lt;0.1</td>
<td>0</td>
<td>0.96</td>
<td>0.62</td>
<td>0.38</td>
<td>gl, fl</td>
</tr>
<tr>
<td>24</td>
<td>53.1</td>
<td>40</td>
<td>0.96</td>
<td>0.72</td>
<td>0.27</td>
<td>gl, fl</td>
</tr>
<tr>
<td>25</td>
<td>51.9</td>
<td>75</td>
<td>0.97</td>
<td>0.79</td>
<td>0.20</td>
<td>gl, fl</td>
</tr>
<tr>
<td>26</td>
<td>51.3</td>
<td>375</td>
<td>0.97</td>
<td>0.89</td>
<td>0.09</td>
<td>gl, fl</td>
</tr>
</tbody>
</table>

Note: gl = glass; fl = vapor.  
† Total iron in the glass expressed as wt% FeO determined by EMPA.  
‡ Bulk FeO (wt%) analyzed by wet chemistry.  
§ Bulk \( Fe_2O_3 \) (wt%) \( Fe_3O_4 \) = \((FeO_{tot}-FeO)\times1.1113\).

---

**Figure 1.** Change in glass \( X_{Fe_2O_3}/X_{FeO} \) with experimental duration for time-series experiments performed on the Ascension starting composition. Data are listed in Table 2.
the oxidation-reduction cycles). The lack of color variation in these glasses suggests that they have uniform, though non-equilibrated, Fe$_2$O$_3$/FeO ratios.

Because Fe loss to the capsule was not detected, the evolution of the Fe-redox state upon either oxidation or reduction cycles can be represented by the change of the glass FeO concentration with time (Figs. 3–5). For an oxidation cycle (Fig. 3), glasses from both starting compositions have FeO concentrations that decrease progressively with experimental duration at $f_{\text{H}_2}<0.1$ bar (Table 4). For the Pinatubo composition, no significant modification of the FeO concentration was detected in a glass quenched 20 min after imposing the new $f_{\text{H}_2}$ (< 0.1 bar) in the membrane. In contrast, for the Ascension composition, a drop in FeO concentration was detected in the glass only 6 min after the imposition of the new $f_{\text{H}_2}$ value. For both compositions, steady-state bulk FeO concentrations (0.54 wt% for Pinatubo and 1.20–1.26 wt% for Ascension) were attained after ~10 and ~3 hours respectively at < 0.1 bar $f_{\text{H}_2}$. The results consistently show that the Ascension composition reacts more rapidly to $f_{\text{H}_2}$ changes and suggest a dependence of the kinetics of Fe oxidation on melt composition. Partial crystallization due to high Fe$_2$O$_3$/FeO ratio of the charge is observed for the two longest experiments in both series (Table 4). For the Pinatubo composition, magnetite appears together with plagioclase, being joined by hematite in the longest experiment (Table 4). Coexisting Fe-Ti oxide compositions yield a $f_{\text{O}_2}$ slightly higher than the MNO buffer (NNO+2.8, corresponding to $f_{\text{H}_2} \sim 10^{-3}$/bar), as calculated using the empirical calibration of Scaillet and Evans (1999). For the Ascension composition, only magnetite was observed to crystallize.

The effect of excess H$_2$O on the kinetics of Fe oxidation was tested by comparing the kinetic response of water-saturated (H$_2$O = 6.3 ± 0.8 wt%) and slightly water-undersaturated (H$_2$O = 4.7 ± 0.6 wt%) charges (Table 4). The rates of change of glass FeO concentrations following a $f_{\text{H}_2}$ drop <0.1 bar, as seen on Figure 4, are identical within analytical uncertainty for water-saturated and water-undersaturated melts (both for the Pinatubo composition), suggesting that they have identical kinetics of Fe oxidation. Similarly, using Ag$_{40}$Pd$_{60}$ capsules instead of Au results in no difference in the rate of change of
glass FeO concentrations (Table 4; Fig. 4), despite H₂ permeability being two orders of magnitude faster in Ag₆₄Pd₃₆ than in Au under our experimental conditions (Chou 1986). Therefore, the kinetics of Fe oxidation is not influenced by the kinetics of H₂ transfer through the capsules.

Results of a reduction cycle performed on the Pinatubo composition are shown in Figure 5. Samples (after being annealed under 50 bar \( f_{H_2} \)) were kept under <0.1 bar \( f_{H_2} \) for 330 min. At this point, the FeO concentration of the glass was 0.62 wt\% (Tables 4 and 6). Following the re-establishment of an \( f_{H_2} \) of ~50 bar in the membrane, glass FeO concentrations progressively increased with time. The highest FeO concentration (0.89 wt\%), attained after 375 min at ~50 bar \( f_{H_2} \) (Table 6), is within the measured range of equilibrium FeO concentrations for 50 bar \( f_{H_2} \) (0.89–0.92 wt\% FeO, Tables 3 and 5). As for the oxidation cycles, the kinetic response is characterized by an initially rapid variation of the FeO concentration of the glass, followed by a slower variation as equilibrium is approached progressively. Patterns shown by oxidation and reduction cycles are symmetrical. No difference exists between the kinetics of oxidation and reduction of Fe and the process is fully reversible (Figs. 5 and 6, see also Fig. 1).

**DISCUSSION**

Among the results presented above, the strong compositional dependence of the rate of change of the Fe\(^{3+}/\text{Fe}^{2+}\) ratio, together with the lack of influence of the capsule material, suggest that the main factor controlling the Fe-redox kinetics is the nature of the silicate liquids (and not the kinetics of the experimental set-up used to vary the \( f_{H_2} \) at \( P \) and \( T \)). Our results thus demonstrate that at constant \( f_{H_2} \), varying \( f_{H_2} \) at equilibrium with a \( \text{H}_2\text{O} \)-, Fe-bearing silicate melt leads to changes in the redox state of iron. At 800 °C, the time needed to establish a new equilibrium melt Fe\(^{3+}/\text{Fe}^{2+}\) ratio after imposition of a new \( f_{H_2} \) is on the order of several to ten hours, which is evidence for the relatively slow kinetics of the Fe-redox reaction in rhyolitic melts. This result is consistent with previous experimental findings by Gaillard et al. (2001), who demonstrated that the Fe-redox ratio of hydrous silicate melts is unchanged when varying the quench rate between 300 and 0.5–1 °C/s, suggesting that the Fe\(^{3+}/\text{Fe}^{2+}\) ratio of such synthetic and natural melts is readily quenchable.

**Comparison between H-free and hydrous melts**

From a kinetic point of view, it is interesting to compare the oxidation-reduction kinetics observed in this study with previous results for dry melts. The kinetics of Fe-oxidation in dry rhyolitic melts (SiO₂ = 73.4 wt\%, FeO tot = 1 wt\%) was studied by Naney and Swanson (1984) at \( T = 1343 \) °C in air. They found that approximately 40 hours were needed to reach Fe\(^{3+}/\text{Fe}^{2+}\) equilibrium for sample volumes similar to ours. In contrast, for compositionally equivalent but H₂O-bearing melts at 800 °C, the present study shows that equilibrium is attained in only 3–10 hours (Figs. 1 and 3). Yet, the melt viscosities in both studies are similar despite the large temperature difference: the rhyolite melt investigated by Naney and Swanson (1984) has a viscosity of \( 1.7 \times 10^2 \) Pa·s at 1343 °C, very close to that of the Pinatubo rhyolite under our experimental \( T \)-\( \text{H}_2\text{O} \) conditions (4.38 \times 10^4 Pa·s, calculated from Hess et al. 1996). Therefore, the melts in both studies have similar relaxation timescales (Dingwell and Webb 1990) even though the kinetics of attainment of Fe-redox equilibrium differ from each other (40 vs. 3–10 hours).

In terms of reaction mechanisms, there is a major difference between our results and previous observations made on oxidation of dry Fe-bearing melts (Cooper et al. 1996; Cook and Cooper 2000). The hydrous glass samples from this study bear evidence neither for a redox front that progresses from the capsule-sample interface toward the core of the sample, nor for cation migration, as observed in the dry systems. Another difference to be noted concerns the delay of several minutes between the application of a new \( f_{H_2} \) and the response in terms of the glass Fe\(^{3+}/\text{Fe}^{2+}\) ratio, which is most clearly seen for the Pinatubo composition (Table 4). Therefore, mechanisms of oxidation-reduction in H-bearing melts appear to differ from those identified by Cooper and co-workers in H-free melts.

**Mechanisms of Fe oxidation-reduction in hydrous silicic melts**

The absence of a redox front and redox heterogeneity in the glasses, despite gradual changes of FeO/FeO, is the major observation of this study. This finding suggests that the redox potentials within each part of the melt are equal but out of equilibrium. Consequentially the redox process appears to be reaction-limited rather than diffusion-limited. Molecular H₂ can be transported across the sample either as a dissolved melt species (see Schmidt et al. 1998) or as a fluid species (quenched melts contain bubbles of trapped aqueous fluids). Given the high diffusion rates of H₂ in melts (Chekhmir et al. 1985), communication of \( f_{H_2} \) is nearly instantaneous at the sample scale. Partial equilibrium of the chemical potential of H₂ (\( \mu_{H_2} \)) should thus be attained virtually instantaneously within the whole sample. In contrast, the delay and rate of FeO/FeO changes after application of a new \( f_{H_2} \) indicates that Fe\(^{3+} \leftrightarrow \text{Fe}^{2+}\) transformations are much slower and thus rate-limit the overall redox processes.

The microscopic phenomena driving the Fe\(^{3+} \leftrightarrow \text{Fe}^{2+}\) trans-
formations are not accessible through our results. However, we note that structural (compositional) controls apply because, for the two compositions studied, two different rates of Fe$^{3+} \leftrightarrow$ Fe$^{2+}$ transformations due to $f_{H_2}$ changes have been measured. This difference can be attributed either to the peralkaline nature (i.e., lower viscosity) or, to a higher Fe concentration in, the Ascension glass compared to the Pinatubo composition.

Mechanisms of interactions between H$_2$ and Fe-bearing silicate minerals such as pyroxene, olivine, and garnet show that hydrogenation-dehydrogenation and Fe$^{3+}$-Fe$^{2+}$ reactions are coupled (Hercule and Ingrin 1999; see Ingrin and Skogby 2000, for a review). The kinetics of these reactions have been proposed to be rate-limited either by the mobility of the hydrogen atoms (at high Fe concentrations) or by the mobility of electron holes (i.e., jumps between Fe$^{2+}$ and Fe$^{3+}$, at low Fe concentrations).

In this study, we have observed that the most Fe-rich melt (Ascension) has the fastest kinetics of Fe-redox equilibration, which could be consistent with the kinetic model proposed for silicate minerals. However, the analogy with minerals cannot be pursued any further because, in contrast to what was observed in NAM, there is evidence neither for mass transfer by diffusion of hydrogen-bearing species as illustrated by the homogeneous H$_2$O concentrations (Table 4), nor for apparent redox heterogeneity in experimental products as would be expected in the case of migration of electron holes.

Kinetics of Fe oxidation-reduction in hydrous silicic melts

Kinetic constants were extracted from the data obtained on each studied composition. For the Ascension composition, results from the oxidation cycles were used whereas for the Pinatubo composition, data from both the oxidation and reduction cycles were used. The best and simplest fit of the experimental data was obtained by considering a first-order (logarithmic) rate law (Lasaga 1998):

$$\ln(C_{eq} - C)/\ln(C_{eq} - C_0) = k.t$$  \hspace{1cm} (3)

where $C_{eq}$ is the equilibrium Fe$^{3+}$/Fe$^{2+}$ concentration, $C_0$ the initial Fe$^{3+}$/Fe$^{2+}$ concentration, $C_t$ the Fe$^{3+}$/Fe$^{2+}$ concentration at time $t$, and $k$ the first-order rate constant. The fitting procedure is explained in the caption of Figure 6. The extracted values yield $k = 0.003$/min for the Pinatubo and $k = 0.0152$/min for the Ascension composition (Fig. 6). The fact that the data obey first-order logarithmic kinetics confirms that the overall process is rate-limited by the Fe$^{3+} \leftrightarrow$ Fe$^{2+}$ transformations, and not by the diffusion of H$_2$, which can be considered instantaneous.

Geological application

This study shows that $f_{O_2}$ changes can be driven by $f_{H_2}$ changes. Hereafter, we model changes in melt Fe$^{3+}$/Fe$^{2+}$ that result from H$_2$ exchanges between a silicic magma (assumed to have the composition of the Pinatubo and Ascension samples) and its host rocks. The principle of the simulation is to impose an initial $f_{H_2}$ difference between the two reservoirs (melt and host rocks), and to calculate, for a given duration, the gradient in $f_{H_2}$ and the associated change in Fe$^{3+}$/Fe$^{2+}$ that results from diffusive exchange of H$_2$. Only lateral transfer of H$_2$ is considered (Figs. 7A and 7B). The simulations may best apply to magma ascent into a conduit during a volcanic eruption. The starting $f_{H_2}$ of the magma is fixed at 20 bar, and the host rock is considered as an infinite reservoir with a constant $f_{H_2} = 0.05$ bar. H$_2$ transfer thus takes place from the magma toward the host rock. The $f_{H_2}$ variation in the magma is computed using Equation 18 of Watson (1994), which applies to the diffusion of a volatile species in a melt in contact with a semi-infinite medium. Temperature is fixed to 800 °C and the magma is assumed to be water-saturated.

To illustrate the importance of the mobility of H$_2$, two values of H$_2$ diffusion rates were tested. The first, equal to 10$^{-9}$ m$^2$/s, represents the lower limit of H$_2$ diffusion rate that can be inferred from our data [using $X = (D.H)^{0.5}$ with $X$ as the capsule diameter and $t$, the minimum duration for observation of Fe$^{3+}$/Fe$^{2+}$ change in the Ascension composition, see Table 4, no. 16]. This value is identical to that determined by Chekhmir et al. (1985) in viscous (10$^{10}$Pa.s) molten albite at 800 °C. Chekhmir et al. (1985) have shown that $D_{H_2}$ is inversely correlated with viscosity (see also Watson 1994). According to their data, in a melt with a viscosity of 10$^{10}$Pa.s (corresponding to the viscosity of our experimental melts), $D_{H_2} \sim 10^{-6}$ m$^2$/s. We therefore adopt 10$^{-5}$ m$^2$/s as an upper limit value for $D_{H_2}$ in hydrous silicic melts at 800 °C.

Calculations were done for an imposed duration of 10 hours, which is realistic for Plinian eruptions. The results are sensibly dependent on the value taken for the H$_2$ diffusivity. In Figure 7A and 7B, variations of $f_{H_2}$ and melt Fe$^{3+}$/Fe$^{2+}$ are presented as a function of distance from the wall rock. Calculated
values of melt $X_{Fe^3+/Fe^{2+}}$ takes into account the kinetic laws identified from Equation 3 (Fig. 6) for both compositions. For $\log D_{H_2} = -9$ m$^2$/s, a zone of ~3 cm is affected by H$_2$ loss and the portion of melt whose Fe$^{3+}$/Fe$^{2+}$ is significantly affected remains negligible (<2 cm for both compositions, Fig. 7A). Note that the two curves nearly overlap. For $\log D_{H_2} = -5$ (Fig. 7B), the $f_{O_2}$ modifications propagate up to 2 m within the magma. For the Pinatubo composition, calculated changes in Fe$^{3+}$/Fe$^{2+}$ propagate in the melt within the first 20 cm corresponding to a fraction of magma affected of 4% for a conduit diameter of 10 m. As a consequence, during a Plinian-type eruption of a metaluminous rhyolite with ~1 wt% FeO, melt $X_{Fe^3+/Fe^{2+}}$ is not significantly affected even for the highest $D_{H_2}$ considered because the kinetics of the Fe$^{3+}$ ↔ Fe$^{2+}$ reaction are not fast enough in such melts. Under these conditions, glasses are therefore good indicators of pre-eruptive $f_{O_2}$. In contrast, similar simulations for the Ascension composition show that the melt Fe$^{3+}$/Fe$^{2+}$ is significantly modified over the first 100 cm (Fig. 7B), corresponding to a fraction of magma affected of 20% for a conduit diameter of 10 m. In such a case, special care should be taken when using Fe$_2$O$_3$/FeO as indicator of pre-eruptive $f_{O_2}$ conditions. These calculations illustrate the strong compositional control on the rate of Fe$^{3+}$ ↔ Fe$^{2+}$ changes during ascent, and stress the need for a more precise knowledge of $D_{H_2}$ for a correct interpretation of the Fe$^{3+}$/Fe$^{2+}$ ratios of erupted lavas.

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