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Experimental determination of activities of FeO and Fe₂O₃ components in hydrous silicic melts under oxidizing conditions

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Abstract—The critical role of iron on crystal-silicate liquid relationships and melt differentiation is mainly controlled by the redox conditions prevailing in magmas, but the presently available database merely constrains the thermodynamic properties of iron-bearing components in strongly reduced and anhydrous molten silicate where iron is in the ferrous form. This paper provides new standard states for pure ferrous (FeO^{Iiq}) and ferric $(Fe_2O_3^{Iiq})$ molten iron oxides and extends the experimental database towards oxidizing and water-bearing domains. Iron-iridium, iron-platinum alloys, magnetite or hematite were equilibrated with synthetic silicic liquids at high temperature and high pressure under controlled oxygen fugacity (fO_2) to determine activity-composition relationships for FeO^{liq} and Fe₂O₃^{liq}. Between 1000 and 1300°C, the fO₂ ranges from that in air to 3-log units below that of the nickel-nickel oxide buffer (NNO). Experiments were performed on both anhydrous and hydrous melts containing up to 6-wt.% water. Incorporation of water under reducing conditions increases the activity coefficient of FeO^{liq} but has an opposite effect on $Fe_2O_3^{-liq}$. As calcium is added to system, the effect of water becomes weaker and is inverted for Fe₂O₃^{liq}. Under oxidizing conditions, water has a negligible effect on both activities of FeO^{liq} and $Fe_2O_3^{liq}$. In contrast, changes in redox conditions dominate the activity coefficients of both FeO^{Iiq} and $Fe_2O_3^{Iiq}$, which increase significantly with increasing fO₂. The present results combined with the previous work provide a specific database on the energetics of iron in silicate melts that cover most of the condition prevailing in natural magmas. Copyright © 2003 Elsevier Ltd

1. INTRODUCTION

The construction of thermodynamic models for multicomponent melts and magmas represents an important direction of research in petrology. Calibration of such models requires determination of both standard states and mixing properties for the melt components. Presently, the most abundant source of data constraining the mixing properties of silicate melts is phase equilibria (Berman and Brown, 1987), as illustrated by several models (e.g., Berman and Brown, 1984; Eriksson and Pelton, 1993; Ghiorso and Sack, 1995; Kirschen and Pichavant, 2001). To test and verify predictions of these models and to provide supplementary constraints for the calibration of the mixing relationships of multicomponent silicate melts, additional measurements such as determinations of activities of the melt components are needed.

Among the different components to be considered, the iron oxide melt components are critical because both ferric and ferrous iron exist in different proportions in magmas, depending on fO_2 , P, T and chemical composition (e.g., Thornber et al., 1980; Mysen and Virgo, 1989; Kress and Carmichael, 1991; Gaillard et al., 2001). In the literature of material science, numerous studies on the energetics of iron in molten silicate were performed on synthetic systems that are, however, very different from natural magmas compositions (Matsuzaki et al., 1998). Surprisingly, despite the fact that iron is the most abundant multivalent element present in magmas, the thermo-

dynamic behavior of other easily reducible cations such as Ni or Co has retained more attention in the geological literature (O'Neill and Eggins, 2002). Also, the presently available database is mostly devoted to the thermodynamic properties of iron in strongly reduced and anhydrous molten silicate where iron is in the ferrous form (Doyle and Naldrett, 1986). Changes in ferric-ferrous ratio have important effects on Fe-Ti oxide saturation curves (Osborn, 1959; Hamilton et al., 1964). The position of Fe-Ti oxides in the crystallization sequence controls the iron content (and ferric-ferrous ratio) of the residual melt, and consequently the stability and composition of major silicate phases (Sisson and Grove, 1993; Martel et al., 1999; Pichavant et al., 2002). Fe-Ti oxides can be near-liquidus phases in basalt for fO2 above the nickel-nickel oxide buffer (NNO) under hydrous conditions (Hamilton et al., 1964; Sisson and Grove, 1993; Martel et al., 1999; Pichavant et al., 2002). Conversely, under anhydrous and reducing conditions, Fe-Ti oxides crystallize near the solidus (Lapin et al., 1985; Snyder et al., 1993; Toplis and Carroll, 1995). For modeling Fe-Ti oxide saturation and liquid lines of descent, an accurate calibration of the thermodynamic properties of ferrous and ferric iron in multicomponent melts is therefore needed. The phase equilibria calculated from models presently available do not reproduce accurately experimental Fe-Ti oxide saturation curves (Ghiorso and Sack, 1995; Toplis and Carroll, 1996; Ariskin, 1999). This problem demonstrates the need for additional specific constraints on the ferrous and ferric melt components under relevant redox conditions. This paper provides new measurement of the activities of liquid FeO^{liq} and Fe₂O₃^{liq} in SiO₂-rich melt, in the presence of significant amounts of Fe³⁺ and under both anhydrous and hydrous conditions.

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		Standard state	Thermodynamic data for		
Reference	Method	for the liquid	the liquid	Equilibria	ΔG° (1 bar,J/mol)*
		Ferrous	iron oxide component		
Schumann and Ensio (1951), Bodsworth (1959)	Pure Fe-metal equilibration	Unspecified	Unspecified	$\mathrm{Fe} + ^{1\!\!/_{2}} \mathrm{O}_{2} \Leftrightarrow \mathrm{FeO}^{\mathrm{liq}}$	-230585 + 44.72. T
Roeder (1974)		XX7 / 1 1			247450 + 40.24 T
Doyle and Naldrett (1986)	Pure Fe-metal equilibration	Wustite liquid	Stull and Prophet (1971)	$Fe + \frac{1}{2} O_2 \Leftrightarrow FeO^{liq}$	-247450 + 49.24. T
Doyle (1988; 1989)	Pure Fe-metal equilibration	Wustite liquid	Coughlin (1954)	$Fe + \frac{1}{2} O_2 \Leftrightarrow FeO^{liq}$	-238771 + 48.53. T
Holzheid et al. (1997)	Fe-Ni-Co alloy equilibration	Wustite liquid	Barin (1989)	$Fe + \frac{1}{2} O_2 \Leftrightarrow FeO^{liq}$	-244118 + 115.56. T
					– 8.474. T. ln T
Snyder and Carmichael (1991)	Fe-Ni alloy equilibration	Wustite solid	Coughlin (1954)	$\begin{array}{c} 0.947\text{Fe} + \frac{1}{2} \text{ O}_2 \Leftrightarrow \\ \text{Fe}_{0.947} \text{O}^{\text{sol}} \end{array}$	-265020 + 64.8. T
Matsuzaki et al (1998)	Pure Fe-metal equilibration	Wustite liquid	Banya et al. (1980)	$Fe + \frac{1}{2}O_2 \Leftrightarrow FeO^{liq}$	-215853 + 35.65. T
	1	FeO liquid	Calculated from their study	$\mathrm{Fe} + \frac{1}{2} \mathrm{O}_2 \Leftrightarrow \mathrm{FeO}^{\mathrm{liq}}$	-224393 + 42.79. T
O'Neill and Eggins (2002)	Pure Fe-metal equilibration	Wustite liquid	Barin (1989)	$Fe + \frac{1}{2} O_2 \Leftrightarrow FeO^{liq}$	-244118 + 115.56. T
					– 8.474. T. ln T
This study	Fe-Ir alloy equilibration	FeO liquid	See appendix	$Fe + \frac{1}{2} O_2 \Leftrightarrow FeO^{liq}$	-226244 + 42.49. T
This study	Hematite equilibration	FeO liquid	See appendix	$ \begin{array}{c} \operatorname{Fe_2O_3^{sol}} \Leftrightarrow 2 \ \operatorname{FeO^{liq}} \\ + \frac{1}{2} \ \operatorname{O_2} \end{array} $	998615-342.43.T
	1	Ferric i	ron oxide component	-	
Snyder and Carmichael (1991)	Fe-Ni alloy equilibration	Hematite solid	Robie et al. (1978)	$Fe + \frac{2}{3} O_2 \Leftrightarrow Fe_2 O_3^{sol}$	-806222 + 248.65. T
This study	Fe-Ir alloy equilibration	Fe ₂ O ₃ liquid	This study	$Fe + \frac{2}{3} O_2 \Leftrightarrow Fe_2 O_3^{liq}$	$-0.0562. T^{2}+374.59.$ T-846564
This study	Hematite equilibration	Fe ₂ O ₃ liquid	This study	$\operatorname{Fe_2O_3}^{sol} \Leftrightarrow \operatorname{Fe_2O_3}^{liq}$	-0.0562. T ² +128.59. T-38275

Table 1. Summary of previous determinations of activities of iron oxides in silicate melts.

* Gibbs free energy of formation of the iron oxide melt component

2. BACKGROUND AND PREVIOUS WORK

Activities of FeO in a wide range of silicate melt compositions (a_{FeO}^{Iiq}) have been determined by geoscientists and material scientists (ref. in Table 1). The different approaches are summarized in Table 1. In most studies, activities of FeO^{Iiq} were determined by equilibrating pure metallic iron with a silicate melt at 1 atm with fO₂ slightly below the iron-wustite (IW) buffer. Under these conditions, iron is present in the melt almost solely as Fe²⁺. The corresponding equilibrium may be written as:

$$Fe + \frac{1}{2} O_2 \Leftrightarrow FeO \tag{1}$$

metal gas melt

Application of the law of mass action yields:

$$a_{FeO}^{liq} = \exp(\Delta G^{o}_{(1)}/RT) a_{Fe} (fO_2)^{1/2}$$
 (2)

where a_{FeO}^{liq} is the activity of FeO in the melt, a_{Fe} is the activity of iron in metal, $\Delta G^{\circ}_{(1)}$ is the Gibbs free energy change associated with equilibrium (1), R is the gas constant and T the temperature (see Table 1). Note that, at 1 bar, $\Delta G^{\circ}(1)$ is simply the Gibbs free energy of formation of the ferrous iron oxide component in the melt, i.e., FeO in Eqn. 1. From Eqn. 2, at equilibrium between melt and pure Fe metal (i.e., $a_{Fe}^{metal} = 1$), the activity of FeO^{liq} is known if fO₂ and temperature are fixed.

Curves of constant a_{FeO}^{Iiq} in a silicate melt equilibrated with pure Fe metal are shown on a T-log fO₂ plot (Fig. 1, see note on Fig. caption). The $a_{FeO}^{Iiq} - X_{FeO}^{Iiq}$ relations have been determined with this approach for mafic to intermediate ferric iron-free melts at 1 bar and under anhydrous conditions (Roeder, 1974; Doyle and Naldrett, 1986; Doyle, 1988, 1989). In the concentration range 0 to 17–18 wt.% FeO, activities of FeO^{Iiq} were found to follow Henry's law, with the Henry's constant a function of melt composition.

The pure Fe metal-silicate equilibration method is, however, limited to a T-fO2 domain restricted to that below the ironwustite buffer (Fig. 1), which corresponds to conditions much more reducing than most terrestrial magmas (Carmichael, 1991). Two studies reported activities of FeO above IW and, consequently, in presence of significant amounts of ferric iron in the melt (Snyder and Carmichael, 1991; Holzheid et al., 1997). For these works, Fe^{3+} in the melt was not measured but calculated using the equation of Kress and Carmichael (1991). To extend the T-fO₂ field of application of the metal-silicate equilibration method, iron-bearing metallic alloys were used instead of pure Fe. Alloying of iron with other metallic elements decreases a_{Fe}^{metal} and thus shifts metal-silicate equilibrium to conditions more oxidizing than the IW buffer (Fig. 1). Snyder and Carmichael (1991) and Holzheid et al. (1997) used, respectively, Fe-Ni and Fe-Ni-Co alloys to monitor aFeO^{liq} in

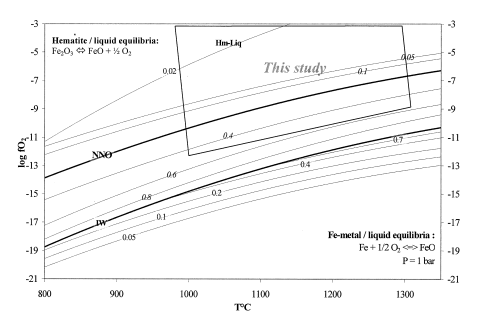


Fig. 1. T-fO₂ field of application of metal-liquid and hematite-liquid equilibria. The two bold curves are NNO (upper) and IW (lower) redox buffer calculated after Robie (1979). The black full lines below the IW buffer (lower part of the graph) are iso- a_{FeO}^{IIq} lines defined by the pure Fe-silicate melt equilibrium calculated using eq. (3). Above the IW redox buffer, the dashed lines defined the T-fO₂ equilibrium between Fe-Ir alloys with different X_{Fe}^{Ir-Fe} (values labeled on curves, calculated after Schwartzendruber, 1984) and silicate melts saturated with respect to pure magnetite (Ghiorso and Sack, 1991). The numbers shown on the dashed curves therefore represent the maximum of Fe that can be found in a Fe-Ir alloys as fO₂ increases. In the upper part of the graph, a curve labeled Hm-Liq define the T-fO₂ domain of equilibrium between pure hematite and a silicate liquid with $a_{FeO}^{Iiq} = 0.02$ (calculated using eq. 9). The box indicates the T-fO₂ conditions explored in this study.

silicate melts at 1 atm for fO₂ up to ~ 5 log units above the IW buffer. Although ferric iron is a minor component (Fe₂O₃ < 1 wt.%) under these conditions, the results of Snyder and Carmichael (1991) clearly show a positive dependence of the activity coefficient of FeO^{1iq} (γ_{FeO}^{Iiq}) with the melt Fe³⁺/Fe²⁺. However, relatively few data points were obtained in the T-fO₂ field where Fe³⁺ is present as a major component. The a_{FeO}^{1iq} database presently available (450 data points) is, therefore, still biased toward ferric iron-free compositions. Snyder and Carmichael (1991) also reported data on the activity of Fe₂O₃ in the liquid but the very low proportion of iron present as Fe₂O₃ (<10% of total Fe). The Fe₂O₃ content was estimated from the empirical regression of Kress and Carmichael (1991) and not directly measured in the experimental glasses.

To complement and extend the available database on the activity of iron oxide components in silicate melts, we present below new a_{FeO}^{Iiq} and a_{FeO3}^{Iiq} data for SiO₂-rich melts, in presence of a significant proportion of ferric iron and for both anhydrous and hydrous conditions.

3. EXPERIMENTAL STRATEGY

3.1. Principle

In this study, we have retained the principle of measuring a_{FeO}^{Iiq} and a_{FeO3}^{Iiq} under relatively oxidizing fO₂ conditions by equilibrating a Fe-bearing alloy with the silicate melt. Noble

metal alloys (Fe-Ir) were preferred to the Fe-Ni alloys used in previous studies for several practical reasons. (1) Fe-Ir alloys are stable over a wide range of fO2, being limited only by the Ir-IrO₂ equilibrium (log fO₂ \sim 9 log units above NNO). In comparison, $a_{FeO}^{}^{liq}$ and $a_{Fe2O3}^{}^{liq}$ cannot be measured at fO₂ >NNO if Fe-Ni alloys are used (Fig. 1). (2) The Fe-Ir alloy (ygFe, Ir phase) forms a continuous solid solution from 0 to 100 at % Fe in the temperature range appropriate for equilibration with silicate melts, i.e., between ~900°C and up to \sim 1400°C (Massalski, 1992). (3) Ir has very low solubilities in silicate melts, <100 ppb (Amossé and Alibert, 1993; O'Neill et al., 1995) and the chemical properties of melts at equilibrium with metallic Ir are not expected to be significantly modified by the presence of dissolved Ir. (4) The mixing properties of the Fe-Ir alloy solid solution are known (Schwartzendruber, 1984) so that a_{Fe} can be calculated directly from X_{Fe}^{Fe-Ir} . Woodland and O'Neill (1997) confirmed the accuracy of the use of Fe-Ir alloys to calculate aFe at high-pressure high-temperature conditions by crosschecking equilibria involving Fe \pm Ir, olivine and quartz. Despite these advantages, the Fe-Ir alloy-silicate equilibration method cannot be used above fO_2 imposed by the limit of detection of iron in the alloy (see Fig. 1). For the analytical conditions of this study (limit of detection of Fe of about 500 ppm, see below), this maximum fO_2 was found to be near NNO+2. Above this fO_2 value, $a_{FeO}^{\ \ liq}$ and $a_{Fe2O3}^{\ \ liq}$ were determined through the hematite-silicate instead of the Fe-Ir alloy-silicate equilibrium.

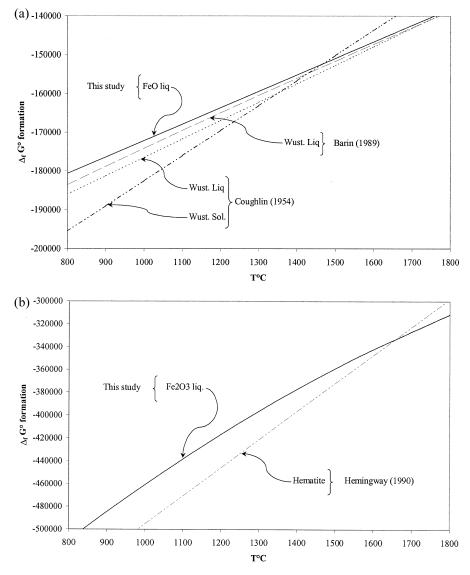


Fig. 2. (A) Gibbs free energy of formation from the elements at P and T of liquid FeO, liquid wustite and solid wustite. (P = 1 bar). For details of source data see Table 1. (B) Gibbs free energy of formation from the elements at P and T of liquid Fe₃O₃ and solid Hematite. (P = 1 bar). For details of source data see Table 1.

3.2. Standard States and Calculation of Activities

Previous determinations of activities of iron oxides in silicate melts have used various stoichiometries, standard states and thermodynamic data for the iron oxide melt components (Table 1). For the ferrous oxide component, a standard state of molten wustite (Fe_{0.947}O liquid) has been commonly chosen despite the fact that molten wustite contains ≥ 13 wt.% Fe₂O₃ at fO₂ conditions corresponding to IW (see Bowen and Schairer, 1934; Schumann and Ensio, 1951). The difference in stoichiometry between FeO and molten wustite was accounted for by various means (for details see Doyle and Naldrett, 1986; Doyle, 1988; 1989; Holzheid et al., 1997; O'Neill and Eggins, 2002). Snyder and Carmichael (1991) adopted standard states of solid wustite and hematite, respectively, for the ferrous and ferric oxide components. To eliminate inconsistencies due to differences in standard states and/or thermodynamic data, and to allow comparison between studies, the same liquid standard states and thermodynamic data have been adopted, as detailed below, and activities from previous studies have been recalculated.

In this study, stoichiometric FeO and Fe₂O₃ liquids were chosen as melt components. Their standard states were calculated from the elements at temperature and pressure. Because thermodynamic data are available for neither component, their standard state properties needed to be estimated. The properties of liquid FeO were estimated from a thermodynamic analysis of the pseudobinary system SiO₂-FeO \pm Fe₂O₃ at 1 atm (Bowen and Schairer, 1934; Schumann and Ensio, 1951; Bodsworth, 1959). The approach and method of numerical optimization are detailed in appendix of this paper. Results (Table 1; Fig. 2a) yield an expression for the Gibbs free energy of formation for liquid FeO at 1 bar, which is near that adopted in early and

Table 2. Starting compositions

							-								
#	1	2	3	4	5	6	7	8	9	10	11	12	Ca1	Ca2	Ca3
SiO ₂	75.84	78.33	75.77	77.17	75.57	77.08	75.66	76.77	76.2	73.44	70.45	69.29	76.32	74.17	76.33
Al_2O_3	11.93	11.77	12.31	12.07	12.64	11.79	11.92	11.45	11.50	11.11	12.26	11.70	11.96	13.29	11.92
FeO*	0.56	0.83	1.08	1.55	1.85	2.58	2.86	3.38	4.25	6.07	7.32	9.28	2.08	2.39	1.97
CaO	0.01	0.03	0.00	0.01	0.01	0.02	0.03	0.04	0.03	0.03	0.00	0.02	0.86	1.87	2.51
Na ₂ O	4.07	4.05	4.24	4.02	3.84	3.74	3.50	3.71	3.67	3.00	3.45	3.16	3.22	2.99	2.40
K ₂ Õ	5.46	4.97	5.34	4.72	4.97	4.76	4.84	4.67	4.36	4.05	4.60	4.20	4.27	3.73	2.81
Total**	97.88	99.90	98.75	99.85	98.88	99.90	98.79	99.98	100.06	97.71	98.08	97.65	98.72	98.45	97.95
A1/CNK***	0.95	0.98	0.97	1.06	1.08	1.04	1.08	1.03	1.07	1.19	1.15	1.20	1.04	1.08	1.03

* FeO total analyzed on EMPA is reported.

** Original total on EMPA.

*** $Al_2O_3/(CaO + Na_2O + K_2O)$ in mole.

recent studies (Schumann and Ensio, 1951; Bodsworth, 1959; Matsuzaki et al., 1998). The activity of liquid FeO is then calculated from equilibrium (1) and Eqn. 2 at known P, T, fO₂ and a_{Fe} (from X_{Fe}^{metal}). The molar volumes of FeO liquid and Fe metal were taken respectively from Lange and Carmichael (1990) and Woodland and O'Neill (1997). This yields the following expression for a_{FeO}^{liq} :

$$a_{FeO}^{Iiq} = a_{Fe} (fO_2)^{1/2} \exp ((-226244 + 42.49T + P. (0.483 + 8.633. 10^{-5}T - 3.784.10^{-6}P))/RT)$$
 (3)

For liquid Fe₂O₃, we have taken the entropy of melting of hematite estimated by Ghiorso and Sack (1995). From the melting point and the standard state thermodynamic properties of hematite (Ghiorso and Sack, 1991), together with Cp_{Fe2O3}^{liq} (Lange and Navrotsky, 1992), the thermodynamic properties of liquid Fe₂O₃ were calculated at any temperature (Fig. 2b; Table 1). The activity of Fe_2O_3 is then calculated from the equilibrium:

$$2\text{Fe} + 3/2\text{O}_2 \Leftrightarrow \text{Fe}_2\text{O}_3$$
 (4)

metal gas melt where:

$$a_{Fe2O3}^{liq} = a_{Fe}^{2} (fO_{2})^{1.5} \exp(\Delta G_{(5)}^{\circ}/RT)$$
(5)

at known P, T, fO_2 and a_{Fe} (from X_{Fe}^{metal}). $\Delta G^{\circ}(5)$ is taken from Table 1. The molar volumes of liquid Fe₂O₃ and Fe metal are taken respectively from Lange and Carmichael (1990) and Woodland and O'Neill (1997). The following expression for a_{Fe2O3}^{liq} is then obtained:

$$a_{Fe203}^{liq} = a_{Fe}^{2} (fO_{2})^{1.5} \exp((-0.0562 \text{ T}^{2} + 374.59 \text{ T} - 846564 + P (1.334 + 0.8.10^{-4} \text{ T} - 0.24510^{-4} \text{ P}))/\text{RT})$$
(6)

We ignore any volume dependence of Fe₂O₃ on the melt composition or structural positions of Fe₂O₃ because no data exists and, in the pressure range of this study, the volume effect on the calculated activity should negligibly affect the activities.

For experiments conducted above NNO+2, activities where obtained via the hematite-liquid equilibria (see above and Fig. 1):

$$\operatorname{Fe}_2\operatorname{O}_3 \Leftrightarrow 2\operatorname{FeO} + \frac{1}{2}\operatorname{O}_2$$
 (7)

hematite liquid gas and:

$$Fe_2O_3 \Leftrightarrow Fe_2O_3$$
 (8)

hematite liquid

and Lange and Carmichael (1990) for the melt components, and from Ghiorso and Sack (1991) for hematite (standard state properties, volume, activity-composition relations). This yields the following expressions:

$$a_{\text{FeO}}^{\text{liq}} = a_{\text{Fe2O3}}^{\text{Hm}} \text{fO}_2^{1/2} \exp((-342.43 \text{ T} + 998615 + P (-1.242 + 0.00046 \text{ T} - 7.537.10^{-6} \text{ P}))/\text{RT})$$
(9)

$$a_{Fe2O3}^{liq} = a_{Fe2O3}^{Hm} \exp((-0.0562 \text{ T}^2 + 128.59 \text{ T} - 38275 + P (-0.3027 + 0.00079 \text{ T} - 2.284 \text{ 10}^{-5} \text{ P})/\text{RT})$$
(10)

where a_{Fe2O3}^{Hm} is the activity of Fe_2O_3 in hematite. Uncertainties on a_{FeO}^{liq} and a_{Fe2O3}^{liq} were estimated by propagating the uncertainties in T ($\sim 10-15^{\circ}$ C), X_{Fe}^{metal} (5 to 15%), $\gamma_{\text{Fe}}^{\text{Ir-Fe}}$ (see Woodland and O'Neill, 1997), and fO₂ (0.05–0.3 log unit). For hydrothermal experiments, fO₂ uncertainties are due to uncertainty in the sensor method calibration and uncertainty in $f_{\rm H2O}$ in the melt (see hereafter). For 1 atm experiments, a crosschecking between fO₂ given by the zircon probe and fO2 estimated from the gas flow indicates maximum departures of ~ 0.05 log units. The mean uncertainties on a_{FeO}^{liq} and a_{Fe2O3}^{liq} are about 10% but can reach 18% for alloys with low X_{Fe}^{alloy} and 25% for the water-undersaturated experiments, which have the largest uncertainties on fO2.

4. EXPERIMENTAL AND ANALYTICAL TECHNIQUES

4.1. Starting Materials

Compositions of the 15 starting glasses are listed in Table 2. They have SiO_2 contents ranging between ~69 and ~78 wt.% and FeO_{tot} between \sim 0.5 and \sim 9 wt.%. The first series of glasses is calcium-free with several compositions (# 2, 4, 6, 8, 9) used previously to determine the effect of H₂O and fO₂ on the ferric-ferrous ratio of silicic melts (Gaillard et al., 2001). A second series of glasses contains different proportion of calcium. All glasses have nearly identical Si₄O₈ contents (between 56 and 63 mol.%, see below for a definition of the melt components). Within the series of calcium-free glasses, FeO_{tot} contents vary at constant KAlO₂/NaAlO₂. The calcic glasses have constant

	Starting composition		Run p	roducts (EMPA)	
	Glass (wt %)	Glass* (wt %)	Fe-Ir powders**	Fe-Pt wire***	Ternary Fe-Ir-Pt****
SiO ₂	49.58	51.1 (73)			
Al_2O_3	17.74	18.02 (32)			
FeO	8.04	4.80 (20)			
MgO	5.39	5.90 (21)			
CaO	10.61	11.90 (31)			
Na_2O	4.28	3.20 (15)			
TiÕ ₂	1.32	1.41 (23)			
Cr_2O_3	0.01	0.01 (2)			
K ₂ O	2.12	2.57 (12)			
Total	99.09	98.82 (79)			
Nb. points	14	18	23	8	1
X _{Fe}			0.063 (0.002)	0.185 (0.003)	0.12
X _{Ir}			0.935 (0.004)	0	0.55
X _{Pt}			Ô	0.815 (0.003)	0.33
a _{FeO}			0.081 (0.007)	0.077 (0.005)	?

Table 3. Test of Pt-Ir interaction: Runs performed at 1380°C, QFM, 1 atm, 10 days.

* Crystal-free glass analyzed after experiments. The two-sigma in brackets applies to the last number after the coma. The two-sigma value for FeO_{tot} equal +/-0.20 wt% demonstrates the homogeneity iron content in the melt and thus attainment of equilibrium of iron partitioning between liquid and metal.

** Pure Fe-Ir alloys analyzed: The composition of these grains represents about 90% of the accessible grains.

*** Pure Fe-Pt alloys analyzed: The composition of these alloys represents about 80% of the Pt-wire.

**** Example of an alloy Fe-Ir-Pt. The zone were these ternary alloys were founded was about 50µm around the Pt-wire.

 FeO_{tot} contents and $KAIO_2/NaAIO_2$, with CaO substituting for the alkali in equal molar proportions. Some glasses are slightly peraluminous and the others slightly peralkaline. Therefore, apart from the Ca = Na + K substitution and the small deviations from the metaluminous plane, the starting glasses can be viewed as pseudobinary mixtures between an aluminosilicate component of nearly constant composition and FeO_{tot}. All glasses were obtained from gels prepared with the method of Pichavant (1987), melted at ~1400 °C in air and then quenched. In most experiments, small amounts of iron were added (as Fe metal and/or Fe₂O₃) to compensate for the loss of iron to the sample

containers and Ir alloys (Table 3). Mixtures of silicate glass and Fe/Fe₂O₃ were homogenized and then ground in an agate mortar. The amount of iron added to the silicate base was estimated from Gaillard et al., (2001) for runs conducted in gold capsule, from Kawamoto and Hirose (1994) for runs with Au-Pd container and from Toplis and Carroll (1996) for Pt capsule.

For experiments at 1 atm, about 1 wt.% of pure iridium powder (grain size $\sim 10 \ \mu$ mm) was added to and homogeneously mixed with the silicate glass \pm Fe/Fe₂O₃ powder. Iridium-free mixtures were also prepared and run in parallel with the iridium-bearing mixtures, the

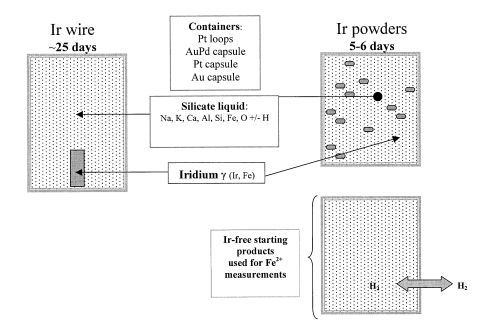


Fig. 3. Protocols used for Ir addition to the silicate base for 1 atm experiments and high-pressure runs.

# Run	# Sample	Starting products	Activity sensors	H ₂ O %wt	Δ NNO	$a_{\rm FeO}^{\rm liq}$	$a_{\rm Fe2O3}^{\rm liq}$
		D-1.28, 25 days					
1	1/1	$#3 + _{Fe2O3}$	Ir powders/Pt wires	0	-1.28	0.157/0.147	0.0009/0.0011
	1/2	$#3 + _{Fe2O3}$ #3 + Fe	Pt wire	"	"	0.159	0.0011
	1/3 1/4		Ir powders/Pt wires Pt wire	"	"	0.110/0.105 0.100	0.0004/0.0004 0.0013
	1/4	$#3 + _{Fe2O3}$ $#3 + Fe + _{Fe2O3}$	Ir powders/Pt wires	"	"	0.054/0.057	0.0001/0.0001
	1/6	$#3 + Fe^{2} + Fe^{2$	Pt wire	"	"	0.057	0.0004
	1/0	$#3 + Fe^{203}$ #3 + Fe	Ir powders/Pt wires	"	"	0.029/0.027	0.00004/0.00002
	1/8	$#3 + Fe + _{Fe2O3}$	Pt wire	"	"	0.028	0.0001
1160°C, 2	2 Kb, $fH_2^{b} = 1$	05 bar, water saturated, 4 d	ays				
2	2/1	$#5 + Fe + Fe_2O_3$	Ir powders	5.21	-1.25	0.106	0.0007
	2/2	$#5 + Fe + Fe_2O_3$	-	4.99	"	-	-
	2/3	$#5 + Fe + Fe_2O_3$	Ir powders	5.56	"	0.063	0.0003
	2/4	$#5 + Fe + Fe_2O_3$,	5.61	"	-	-
	2/5	$#5 + Fe + Fe_2O_3$	Ir powders	4.96	"	0.119	0.0009
	2/6	$#5 + Fe_2O_3$	— I	5.30	"	-	-
	2/7 2/8	$#5 + Fe_2O_3$	Ir powders/Pt capsule	5.03	"	0.028/0.032	0.00005/0.00006
1000°C 2		$#5 + Fe + Fe_2O_3$ 125 bar, 6 days	Pt capsule	5.00		0.051	0.0002
1000 C, 2 3	3/1 - 1	$#5 + Fe + _{Fe2O3}$	Ir Powders	3.03	-2.85	0.064	0.00007
5	3/2	$#5 + Fe + Fe^{2O3}$ $#5 + Fe$	"	5.55	-1.91	0.004	0.00001
	3/2	#5 + Fe	"	5.96	-1.91	0.107	0.0006
	3/4	$#7 + Fe + _{Fe2O3}$	"	5.40	-2.08	0.109	0.0005
	3/5	$#7 + Fe + \frac{Fe_{2}O_{3}}{Fe_{2}O_{3}}$	"	3.41	-2.66	0.076	0.0001
	3/6	$#3 + Fe + \frac{Fe_{2}O_{3}}{Fe_{2}O_{3}}$	"	6.28	-1.95	0.055	0.0001
	3/7	$#5 + _{Fe2O3}$	"	6.54	-1.91	0.042	0.00009
	3/8	$#3 + _{Fe^{2}O^{3}}$	"	5.15	-2.18	0.011	0.00000
	3/9	#3 + Fe	"	4.16	-2.39	0.104	0.0003
	3/10	$#5 + Fe + _{Fe2O3}$	"	6.65	-1.91	0.030	0.00005
	3/11	#5 + Fe	"	2.40	-3.16	0.028	0.00001
	3/12	$#5 + Fe + _{Fe2O3}$	"	6.00	-1.91	0.048	0.0001
100000	3/13	#5 + Fe + Fe		5.94	-1.91	0.050	0.0001
1000°C, 2		31 bar, water saturated, 16 c		5 10	0.8	0.011	0.00002
4	4/1 4/2	#2 #3	Ir Wire	5.12 5.95	$-0.8 \\ -0.8$	0.011 0.018	0.00002 0.00005
	4/2 4/3	#3 #4	"	5.24	-0.8	0.018	0.0001
	4/4	#5	"	5.32	"	0.023	0.0001
	4/5	#8	"	5.26	"	0.063	0.0007
	4/6	#10	"	5.07	"	0.085	0.0021
	4/7	#11	"	4.49	"	0.112	0.0035
	4/8	#12	"	4.50	"	0.120	0.0040
1000°C, 2	2 Kb, $fH2^{b} = 2$	22 bar, water saturated, 6 da	iys				
5	5/1	#11	Ir Wire	5.10	-0.5	0.159	0.0055
	5/2	#12	"	4.50	"	0.188	0.0087
,	,	.4 bar, water saturated, 6 da	J				
6	6/1	#1	Ir Powders	~ 6.68	+1.4	0.012	0.0002
	6/2	#2	"	~6.97	"	0.030	0.0015
	6/3	#5	"	~6.58	"	0.059	0.0056
	6/4 6/5	#6 #9	Ir powders/Magnetite	$\sim 6.97 \\ \sim 7.15$	"	0.076 0.092/0.087	0.0093 0.0136/0.01300
	6/6	#9 #10	Ir powders/Magnetite	~6.80	"	0.085/0.080	0.0118/0.0109
	6/7	#10 #1		~0.80 5.54	"	0.085/0.080	-
	6/8	#1 #2	_	5.43	"	_	_
	6/9	#5	_	5.26	"	_	_
	6/10	#6	_	5.23	"	_	_
	6/11	#9	Magnetite	~6.45	"	0.090	0.0132
	6/12	#10	Magnetite	~6.81	"	0.084	0.0110
1245°C, 0	0.001 Kb, air, 2	25 days	-				
7	7/1	$#Ca3 + Fe + Fe_2O_3$	Hm	0	+7.3	0.017	0.099
	7/2	$#Ca2 + Fe_2O_3$	"	"	"	0.017	0.102
	7/3	$\#Ca1 + Fe + Fe_2O_3$	"	"	"	0.017	0.102
	7/4	#3 + Fe	"	"	"	0.017	0.096

resulting glasses being used for the analysis of FeO and the determination Fe³⁺/Fe²⁺ of the glass. Starting products were suspended on Pt wires (sample/Pt mass ratio ~ 10) and subsequently placed in the hot spot of a 1 atm gas-mixing furnace.

For experiments at 2 kbar, wires of iridium metal (diameter 75 μ m, see Fig. 3) were initially used as the source of Ir. This method resulted in impracticably long experimental durations (i.e., ≥ 25 d), mainly because of the slow kinetics of diffusion of Fe in Ir metal. To decrease

		Table	4. commueu				
# Run	# Sample	Starting products	Activity sensors	H ₂ O %wt	Δ NNO	$a_{\rm FeO}^{\rm liq}$	a _{Fe2O3} liq
300°C, 0.001 Kb, air, 25 days							
3	8/1	#Ca1 + Fe	Hm	0	+6.7	0.028	0.135
	8/2	$#Ca2 + Fe + Fe_2O_3$	"	"	"	0.028	0.130
	8/3	$#3 + Fe_2O_3$	"	"	"	0.028	0.133
160° C, 2 Kb, fH2 ^b = 0.1 bar.	water satura	ted, 4 days					
)	9/1	$#3 + Fe_2O_3$	_	5.73	+4.9	_	_
	9/2	$#3 + Fe^{-1}$	Hm	6.06	"	0.026	0.070
	9/3	$#3 + Fe_2O_3$	Hm	5.79	"	0.026	0.069
	9/4	$#3 + Fe + Fe_2O_3$	"	5.77	"		
	9/5	$#Ca1 + Fe + Fe_2O_3$	"	5.55	"	0.026	0.070
	9/6	$\#Ca2 + Fe + Fe_2O_3$	"	5.58	"	0.024	0.069
	9/7	$\#Ca3 + Fe_2O_3$	"	5.84	"	0.023	0.068

Table 4. continued

the duration necessary for attainment of equilibrium, iridium was later introduced as in the 1 atm experiments, i.e., as powders of ~10 μ m grain size premixed with the silicate glass ± Fe and/or Fe₂O₃ (Fig. 3). As for 1 atm, iridium-free starting silicate glass ± Fe and/or Fe₂O₃ mixtures were run in parallel with the iridium-bearing mixtures. The starting products were loaded in noble metal capsules together with known quantities of distilled water. A majority of experiments were performed under H₂O-saturated conditions. The charges typically consisted of ~10 mg of H₂O for ~100 mg of the solid materials. For the H₂O-undersaturated experiments, H₂O in the charges ranged between 2 and 10 mg. Either Au, Au-Pd or Pt capsules (length 15–20 mm, water used depending on the experimental temperature (i.e., Au at 1000°C, runs # 3–6; Au-Pd and Pt at 1160°C, runs # 2, 9, Table 4).

4.2. Experimental Techniques

Experiments were conducted at both 1 atm and 2 kbar. For the 1 atm experiments, a vertical SiC resistance furnace was used. Temperature was monitored by a type S thermocouple and is known to within \pm 5 °C. CO₂-CO gas mixtures with proportions calculated from Deines et al. (1974) served to impose the experimental fO_2 which was measured directly by an yttria-stabilized zirconia probe with an uncertainty of 0.02-0.05 log unit (see above). A drop quench technique was used at the end of the experiments. The 2 kbar experiments were performed in an internally heated pressure vessel fitted with a rapid quench device modified after Roux and Lefèvre (1992). A two-windings furnace adapted from the device of Roux et al. (1994) was used. Temperature was measured by three type B thermocouples with estimated uncertainties (including thermal gradients) of $\pm 10^{\circ}$ C at 1000°C and $\pm 15^{\circ}$ C at the highest temperature investigated (1160°C). Pressure was imposed by Ar-H₂ mixtures, whose fH₂ served to control the redox conditions through the equilibrium:

$$H_2 + \frac{1}{2} O_2 \Leftrightarrow H_2 O \tag{11}$$

which shows that fO₂ is fixed if fH₂ is imposed to a H₂O-bearing system of known fH₂O (Scaillet et al., 1992; 1995). The fH₂ was not constantly monitored using a Shaw membrane but it was determined for each run using the Co-Ni-Pd redox sensors (Taylor et al., 1992; Pownceby and O'Neill, 1994). The fO₂ was subsequently calculated with fH₂O either taken from Burnham et al. (1969) for the watersaturated experiments, or computed from glass H₂O contents and the model of Silver et al. (1989) for the water-undersaturated experiments. Uncertainties in fO₂ were propagated through uncertainties in fH₂ calculated from redox sensor (1–5% depending on fH₂ value) and uncertainties in f_{H2O} of the melt (10%). At water saturation, fO₂ are shown to be accurate to within 0.1 log units and for water-undersaturated conditions, uncertainties rose to 0.3 log units.

4.3. Analytical Methods

Experimental products, including glasses, oxide minerals, and Fe-Ir alloys, were analyzed with a Cameca SX-50 electron microprobe of the

BRGM-CNRS laboratories at Orléans, France. Silicate glasses and oxide minerals were analyzed under the following operating conditions: acceleration voltage 15 kV, sample current 6 nA, counting time on peak 10 s. A beam size of 1–2 μ m was employed for oxide minerals whereas, for glass analysis, the beam was defocused to $\sim 10 \ \mu m$ to minimize the migration of alkali (Pichavant, 1987). The H₂O content of experimental glasses was estimated from the by-difference method (Devine et al., 1995), calibrated against a set of internal hydrous glass standards analyzed together with the unknowns during each microprobe session. Results were cross-checked with FTIR determinations on selected samples (see below and Gaillard et al., 2001). Fe-Ir alloys were analyzed under an acceleration voltage of 20 kV, a sample current of 20 nA, with counting times of 20 and 10 s on peak and background respectively and an electron beam focused to $1-2 \ \mu m$. Standards were pure metal Fe and Ir. Several other elements (S, Au, Pt, Pd, Si and Ni) were systematically analyzed together with Fe and Ir. Due to the small size ($\leq \sim 10 \ \mu m$) and porous texture of the Fe-Ir alloys, their analysis was difficult and the data indicated contamination with glass, more rarely magnetite. To overcome this problem, analyses were replicated until a minimum of 6-7 satisfactory analyses were obtained per sample, with the Si and Ir concentrations being taken as indicators of the extent of glass contamination. Under our analytical conditions, the limit of detection of Fe in the alloy is about 500 ppm.

A wet chemical technique, based on potassium dichromate titration, was used to measure the ferrous iron content (i.e., FeO) of glasses (Thornber et al., 1980; Gaillard et al., 2001). Details of the method and tests of reproducibility and accuracy were provided by Gaillard et al. (2001). The analytical uncertainty (expressed in wt.% FeO) increases with FeO, i.e., \pm 0.02 at 0.3 wt.% FeO and \pm 0.15 at 7 wt.% FeO. The Fe₂O₃ content of glasses was calculated by difference between total iron analyzed by electron microprobe (EMPA) and ferrous iron following:

$$Fe_2O_3 wt\% = (FeO_{tot} - FeO) 1.1113$$
 (12)

Only glasses from experiments performed with Ir-free starting mixtures were analyzed. The relationships between ferrous and total iron content were then established and fitted for Ir-free samples synthesized during each run (see results, 5.2.1, 5.2.2). These empirical relationships were used to calculate the FeO (and Fe₂O₃) contents of Ir-bearing glasses whose total iron contents were known from EMPA. Some glasses were not totally crystal-free, containing small amounts of either magnetite or hematite (see results). These Fe-oxides were shown to be in equilibrium with the melt and therefore, they should not be quenched phases (see results, 5.1.2). For these oxide-bearing samples, the FeO content of the glass was obtained by the empirical fit relating ferrous to total iron of the melt calibrated on crystal free samples. The ferric iron content in the glass was calculated from the measured total iron and calculated ferrous iron in the glass. This method was preferred to mass balance calculations that would consist in subtracting the contribution of oxides to the ferric-ferrous ratio of the sample to estimate the glass Fe^{3+}/Fe^{2-3} This mass balance method was shown to propagate significant uncertainties in some case (see Gaillard et al., 2001 for details).

For selected samples, infrared spectrometry was used to determine the glass OH and H_2O contents. Doubly polished glass wafers with

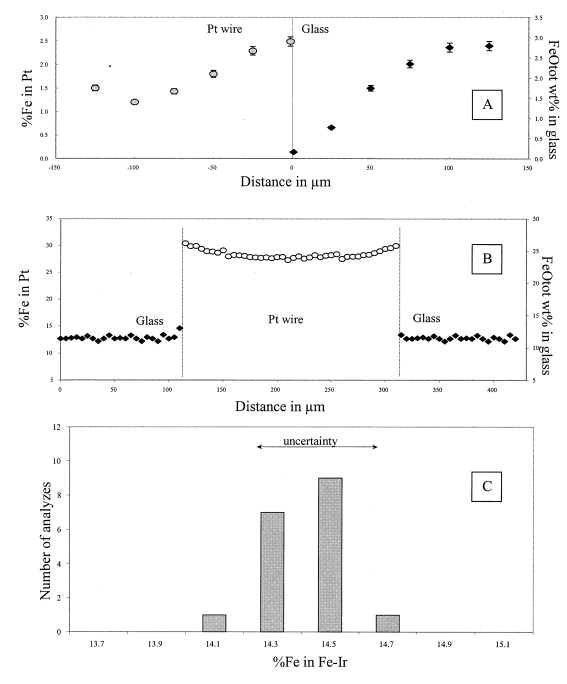


Fig. 4. Melt-Pt-Ir equilibria at one atm. (A): Non-equilibrated profile at the melt-Pt boundary (10 days, non tabulated); (B, C): Equilibrated, 25 days, (see Table 4, 5): (B) Profile in the Pt wire and at the melt-Pt boundary in sample #1/2. (C) Histogram of Fe content in 18 analysed Fe-Ir grains of the sample #1/3.

known thickness were placed under the beam of a Nicolet 760 Magna FTIR spectrometer. Analytical procedures were the same as described by Gaillard et al. (2001). Extinction coefficients from Newman et al. (1986) were used to calculate the OH and H_2O content (wt.%) from peak heights of the absorption bands at 4500 cm^{-1} and 5200 cm^{-1} respectively. Because fluid inclusions, Fe oxide minerals and/or metallic powders may be present locally, glass densities were not measured but calculated density agrees within less than 10% with recent density measurements for glasses of similar compositions (Behrens and Jantos, 2001).

5. RESULTS

5.1. Equilibrium and Internal Consistency

Determination of a_{FeO}^{liq} and a_{Fe2O3}^{liq} requires attainment of chemical equilibrium, either between Fe-Ir alloy and melt for $fO_2 < NNO+2$ (equilibria (1) and (5)) or between hematite and melt above NNO+2 (equilibria (8), (9)). Attainment of equilibrium was tested by performing time-series experiments. The

Table 5. Chemical composition of phases. (See Table 4 for activities calculated from alloys and/or oxides composition and

Methods					EMPA	4				FTIR	<u> </u>		Wet Chen	nistry
Phases		Glass	es or Ox	ides wł	ien speci	ified		Al	loys			Glas	ses	
# Sample	SiO_2	Al_2O_3	FeO	CaO	Na ₂ O	K ₂ O	H_2O	X Fe ^(Ir)	X Fe ^(Pt)	OH/H ₂ O	H_2O	FeO wt%	Fe ₂ O ₃	X Fe ₂ O ₃ /XFeO
1210°C, 0.	,		,	2										
1/1	69.08	11.75	10.37	0.00	3.11	4.23	-	0.17	0.30	n.d.	n.d.	9.55	0.91	0.04
1/2	68.14 71.06	11.50	11.85	0.07	3.08	4.28	-	0.14	0.31	n.d.	n.d.	10.37	1.65	0.07
1/3 1/4	71.96 71.65	11.75 11.82	7.35 7.89	$0.08 \\ 0.07$	3.35 3.29	4.55 4.49	-	0.14	0.26 0.28	n.d.	n.d. n.d.	6.99 7.60	0.40 0.32	0.03 0.02
1/4	74.21	11.82	4.26	0.07	3.54	4.49	_	0.11	0.28	n.d. n.d.	n.d.	4.13	0.32	0.02
1/6	74.04	12.01	4.67	0.00	3.59	4.65	_	0.11	0.25	n.d.	n.d.	4.50	0.19	0.02
1/7	75.68	12.05	2.10	0.04	3.80	4.80	_	0.07	0.18	n.d.	n.d.	2.06	0.05	0.01
1/8	75.91	12.10	2.29	0.06	3.71	4.86	_		0.22	n.d.	n.d.	2.20	0.10	0.02
1160°C, 2														
2/1	66.79	10.38	8.84	0.02	3.05	4.43	5.21	0.15		n.d.	n.d.	7.18	1.84	0.12
2/2	65.30	10.22	10.64	0.01	3.25	4.35	4.99	0.10		n.d.	n.d.	8.40	2.49	0.13
2/3	69.94	10.95	4.27	0.02	3.32	4.58	5.56	0.12		n.d.	n.d.	3.80	0.52	0.06
2/4 2/5	69.22 65.95	10.72	5.23	0.01	3.27 3.20	4.59	5.61	0.16		n.d.	n.d.	4.50	0.81 2.05	0.08
2/3	65.74	10.34 10.25	9.85 9.81	0.05 0.04	3.20 3.17	4.39 4.40	4.96 5.30	0.16		n.d. n.d.	n.d. n.d.	8.01 8.00	2.03	0.12 0.11
2/0 2/7	72.89	11.32	1.78	0.04	3.34	4.78	5.03	0.07	0.16	n.d.	n.d.	1.62	0.18	0.05
2/8	72.08	11.16	3.55	0.05	3.35	4.77	5.00	0.07	0.10	n.d.	n.d.	3.15	0.13	0.05
1000°C, 2					0.00		2.00		0110	indi	mai	0110	0	0.000
3/1	71.60	11.63	4.54	0.01	3.55	4.60	3.25	0.32		1.1/1.93	3.03			
3/2	71.67	11.76	0.82	0.01	3.52	4.47	6.71	0.11		1.38/4.17	5.55			
3/3	68.46	10.57	6.05	0.02	3.19	4.15	6.55	0.26		1.35/4.61	5.96			
3/4	68.00	10.64	7.15	0.01	3.13	4.15	5.85	0.28		1.28/4.12	5.40			
3/5	71.54	11.74	4.77	0.03	2.90	4.50	3.56	0.31		0.98/2.43	3.41			
3/6	70.38	10.62	3.27	0.02	3.56	4.21	6.59	0.21		1.37/4.91	6.28			
3/7	71.07	10.66	2.45	0.01	3.31	4.21	6.88	0.18		1.4/5.14	6.54			
3/8 3/9	72.88	11.72	0.70	0.03 0.01	3.69 3.29	4.44	6.02 4.99	0.11 0.31		1.37/3.78	5.15			
3/9	68.69 71.70	10.89 10.72	7.46 1.69	0.01	3.43	4.30 4.11	4.99 7.1	0.31		1.15/3.01 1.42/5.23	4.16 6.65			
3/11	75.40	11.34	1.81	0.02	3.70	4.46	2.66	0.10		0.85/1.55	2.40			
3/12	71.32	10.67	2.63	0.01	3.60	4.19	6.44	0.19		1.45/4.55	6.00			
3/13	71.44	10.74	2.41	0.02	3.67	4.23	6.77	0.16		1.39/4.55	5.94			
1000°C, 2	Kb, fH2	^b = 31	bar, wate	er satura	ated, 16									
4/1	73.70	11.47	0.61	0.01	3.70	4.57	6.25	0.04		1.35/3.77	5.12	0.52	0.11	0.09
4/2	73.20	11.50	0.98	0.00	3.80	4.53	5.95	0.055		n.d.	n.d.	0.79	0.21	0.12
4/3	72.37	11.48	1.22	0.02	3.78	4.60	6.35	0.07		1.34/3.91	5.24	0.97	0.28	0.13
4/4	72.82	10.98	1.81	0.03	3.40	4.63	6.09	0.095		1.34/3.97	5.32	1.39	0.47	0.15
4/5	72.10	10.45	3.12	0.02	3.57	4.29	6.01	0.12		1.26/4.00	5.26	2.40	0.80	0.15
4/6	71.16	11.10	4.65	0.00	3.18	3.79	5.01	0.14		1.23/3.84	5.07	3.68	1.08	0.13
4/7 4/8	69.21 68.29	11.64 11.82	6.37 7.18	0.03 0.00	3.32 3.47	4.14 4.06	4.36 4.22	0.16		1.06/3.43	4.49	4.90 5.59	1.64	0.15 0.14
4/8 1000°C, 2							4.22	0.165		1.04/3.46	4.50	5.59	1.77	0.14
5/1	68.38			0.02	3.65	4.45	5.10	0.16		n.d.	n.d.	4.00	1.59	0.18
5/2	67.16	11.28		0.02	3.51	3.97	4.50	0.17		n.d.	n.d.	6.15	1.97	0.14
1000°C, 2												0.10		
6/1	73.46	10.90	0.44	0.01	3.66	4.75	6.68	0.01		n.d.	n.d.	0.33	0.12	0.16
6/2	72.80	11.25	0.77	0.06	3.48	4.56	6.97	0.01		n.d.	n.d.	0.55	0.25	0.20
6/3	72.01	11.46	1.71	0.00	3.55	4.57	6.58	0.03		n.d.	n.d.	1.05	0.73	0.31
6/4	72.33	10.54	2.20	0.01	3.42	4.41	6.97	0.03		n.d.	n.d.	1.26	1.04	0.37
6/5	72.25	10.45	2.71	0.03	2.93	4.37	7.15	0.04		n.d.	n.d.	1.46	1.39	0.43
6/5-Mt	70.07	1.89	88.73	0.00	2.02	4.00	< 00	0.02			1	1.07	1 1 1	0.27
6/6	70.96	11.83	2.86	0.02	3.03	4.39	6.80	0.03		n.d.	n.d.	1.86	1.11	0.27
6/6-Mt	73 61	3.02	87.70	0.02	3 77	4.78	6.05			1.50/4.04	5.54	0.21	0.12	0.17
6/7 6/8	73.61 72.80	11.16 11.48	0.41 0.82	0.02 0.01	3.72 3.65	4.78 4.51	6.05 6.20			1.50/4.04	5.54 5.43	0.31 0.60	0.12 0.25	0.17
6/8 6/9	72.80	11.48	1.86	0.01	3.60	4.51	6.20 6.40			1.43/3.98	5.45 5.26	1.13	0.23	0.18
6/10	72.29	10.76	2.42	0.00	3.48	4.44	6.38			1.24/3.99	5.20	1.15	1.19	0.32
6/11	72.15	10.68	2.93	0.02	3.17	4.38	6.45			n.d.	n.d.	2.00/1.68	2.42/1.68	0.40
6/11-Mt		1.82	88.85											
6/12	71.18	11.43	2.97	0.02	3.18	4.25	6.81			n.d.	n.d.	3.09/1.93	4.35/1.16	0.27
6/12-Mt		2.94	87.93											
7														
7/1-gl	70.22	14.93	3.77	2.50	3.43	3.18	-					0.78	3.32	1.91
7/1-HM	72.26	6.24	82.09	1 70	2.04	2 50						0.00	2.02	1 07
7/2-gl 7/2-HM	73.26	13.08	3.23	1.78	2.96	3.59	-					0.68	2.83	1.87
1/2- H VI		5.16	82.28											

Methods					EMI	PA				FTIR	FTIR Wet Chemistry			nistry
Phases		Glasses	or Ox	ides v	vhen sp	ecified	l	Alloy	ys			Glasse	es	
# Sample	SiO ₂	Al_2O_3	FeO	CaO	Na ₂ O	K ₂ O	H_2O	X Fe ^(Ir) X	K Fe ^(Pt)	OH/H ₂ O	H_2O	FeO wt%	Fe ₂ O ₃	X Fe ₂ O ₃ /XFeO
7/3-gl	75.29	11.86	2.73	0.85	3.14	4.15	-					0.54	2.43	2.02
7/3-HM		3.81	84.01											
7/4-gl	75.18	12.19	2.45	0.02	3.48	4.63	-					0.52	2.14	1.85
7/4-HM		3.95	83.94											
1300°C, 0.001 Kł	, air, 25	days												
8/1-gl	75.14	11.56	3.85	0.84	3.03	4.08	-					0.77	3.42	2.00
8/1-HM		3.56	84.75											
8/2-gl	72.91	12.78		1.63	2.93	3.62	-					1.07	3.57	1.50
8/2-HM		4.78	82.77											
8/3-gl	75.03	11.93		0.01	3.40	4.65	-					0.73	2.69	1.66
8/3-HM			83.39											
1160°C, 2 Kb, fH														
9/1-gl		10.96								n.d.	n.d.	0.54	1.62	1.35
9/2-gl	70.83	10.89		0.05	3.26	4.48	6.06			n.d.	n.d.	0.70	2.55	1.64
9/2-HM			85.32											
9/3-gl	70.82	10.80		0.02	3.15	4.56	5.79			n.d.	n.d.	0.72	2.49	1.56
9/3-HM			86.45											
9/4-gl	70.83	10.80		0.00	3.19	4.50	5.77			n.d.	n.d.	0.73	2.51	1.55
9/4-HM			85.24											
9/5-gl	70.71	10.97		0.79	2.92	3.89	5.55			n.d.	n.d.	0.84	2.26	1.21
9/5-HM			86.38											
9/6-gl	69.75	11.81		1.66	2.80	3.44	5.58			n.d.	n.d.	0.85	2.03	1.08
9/6-HM			86.40											
9/7-gl	70.96	10.89		0.03	3.47	4.58	5.84			n.d.	n.d.	0.68	1.91	1.26
9/7-HM		1.60	85.32											

Table 5. Continued

activity determinations were cross-checked with independent equilibria, involving either noble metals others than Ir or magnetite. Strictly speaking, no reversals were performed (i.e., compositions of alloys were not reversed by duplicating experiments starting with pure Ir with others performed for example with $Fe_{50}Ir_{50}$ alloys). However, Fe metal was present in most

starting mixtures (Table 4) and alloy compositions were thus approached from both sides (i.e., pure Fe and pure Ir). Chemical equilibrium for the metallic iron and iron oxide components was demonstrated by using the following criteria: (1) attainment of homogeneous chemical compositions for glasses, for Fe-Ir alloys and for noble metal suspension wires/contain-

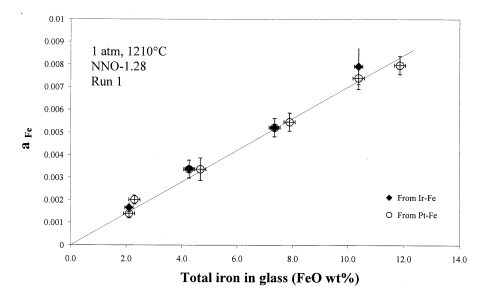


Fig. 5. Evidence for the consistency between Fe-metal activities calculated from Pt-Fe and, when available, Ir-Fe alloys. The total iron content of the melts is plotted against the calculated Fe metal activities that define, within error bars, a single trend. # Run 1, see Table 4 and 5 for data.

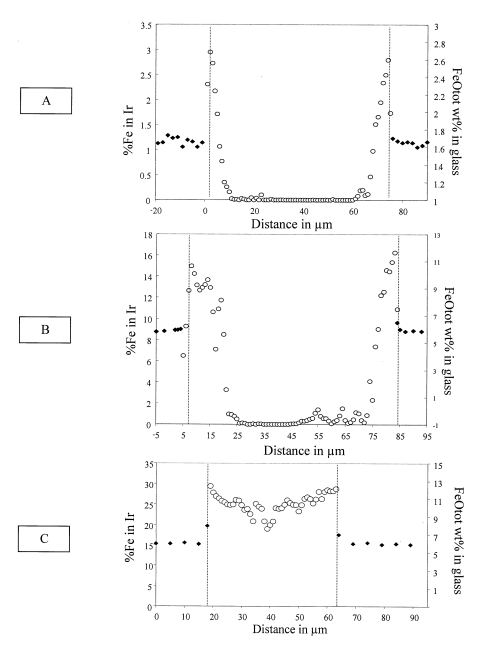


Fig. 6. Melt Fe-Ir interactions at 2 kb, 1000°C and water saturation illustrated by chemical profiles at the melt-Ir boundary. Non equilibrated after 4 days (A, run non-tabulated) and 10 days (B, sample #5/9 see Table 4-5):; (C): Nearly equilibrated after 25 days (run non-tabulated because the redox sensor was polluted by sulfide and subsequently unusable).

ers, (2) internal consistency of a_{Fe} , a_{FeO}^{liq} and a_{Fe2O3}^{liq} calculated from independent equilibria, which is the basic definition of thermodynamic equilibrium (equality of chemical potentials). These different lines of evidence are detailed below.

5.1.1. 1 atm experiments

At ~1200°C, several weeks were needed for attainment of equilibrium. The main problem encountered is the loss of iron to the Pt suspension wire. This causes iron depletion in the melt near the wire and produces heterogeneity in FeO_{tot} of glasses. At 1210°C, experiments of 25 d (run 1, Table 4) were required to obtain glasses with homogeneous FeO_{tot} (Fig. 4a, b). Once

the melt iron content is homogeneous, the analyzed Fe-Ir alloys are also homogeneous (Fig. 4c), suggesting that the kinetics of equilibration between Fe-Ir alloy and melt is faster than the kinetics of Fe equilibration with Pt loops by diffusion in the anhydrous melt (see also Woodland and O'Neill, 1997). However, to ensure that equilibrium was reached, each 1-atm run was performed for durations of 25 d. At constant fO₂, the Fe³⁺/Fe²⁺ of glasses synthesized from starting products with different proportions of metal Fe and Fe₂O₃ are close to each other (see Tables 4 and 5, runs # 1, 7, 8), suggesting that the iron redox equilibrium is attained or closely approached. In run # 1 (Table 4), compositions of the Pt-Fe suspension wires (Fig. 4b)

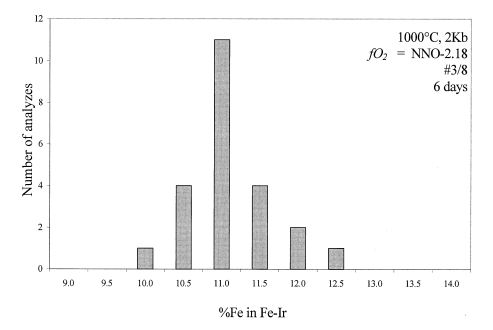


Fig. 7. Histogram of Fe content in 23 analysed Fe-Ir grains of the charge #3/8. The experiment lasts 6 days (See Table 4-5).

served to crosscheck the activity calculation from Ir-Fe (Heald, 1967; Schwartzendruber, 1984). Results show that a_{Fe}^{metal} (and hence μ_{Fe}^{metal}) are identical within error in both alloys (Fig. 5), the two activity sensors yielding nearly identical a_{FeO}^{Iiq} and a_{Fe2O3}^{Iiq} (Table 4). Because the Pt wire and the Ir powders are not in direct contact but are separated by the melt, the identity in μ_{Fe}^{metal} implies that each alloy is in equilibrium with the silicate melt (equilibria (1), (5)). For runs # 7 and 8, performed in air, hematite-liquid equilibria because the Fe concentration in the alloys is below detection under these fO₂ conditions. However, a_{FeO}^{Iiq} and a_{Fe2O3}^{Iiq} are

similar in runs performed with starting materials having different proportions of metal Fe and Fe_2O_3 (Table 4). This is interpreted as evidence for attainment of equilibrium in these two runs.

5.1.2. 2 kbar experiments

Results from the time-series experiments conducted with Ir wires are illustrated in Figure 6. At 1000°C, 2 kbar and for H₂O-saturated conditions, nearly flat Fe profiles in the Ir wire are obtained for durations of 25 d (Fig. 6c). For shorter durations, bulk equilibrium is clearly not attained. However, there is

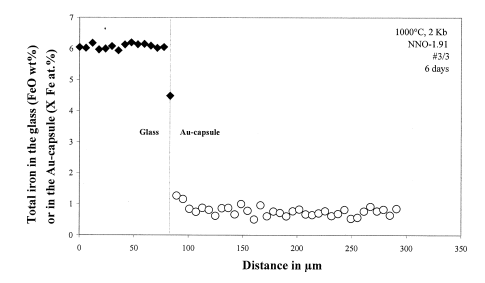


Fig. 8. Evidence of equilibrium attainment between melt and Au-capsule after 6 days-long experiments. Sample #3/3 (see Table 4-5).

no gradient in Fe concentration in the melt adjacent to the Ir wire. The Fe contents at the metal-melt interface are reproducible, suggesting local equilibrium between metal and melt (run # 4 and 5, Table 4, see also Fig. 6a, 6b). In comparison, experiments performed with Ir powders attained equilibrium much more rapidly. Under the same conditions as above (1000°C, 2 kbar, H₂O saturation), homogeneous Fe concentrations in Ir-Fe alloys were obtained after 6 d (run # 3, Table 4, Fig. 7). The establishment of ferric-ferrous equilibria in hydrous silicic melts requires only a few hours at 800-1000°C (Gaillard et al., 2002). In contrast, Fe loss to the capsule may require a significant amount of time because of a high container/sample mass ratio that makes the Fe mass transfer important. At fO₂ \ll NNO (run # 3, Table 4), Fe losses of about 50% were encountered for charges with the highest capsule/melt weight ratios. However, the Fe profile (Fig. 8) indicates that the Au capsule is homogeneous and there is no gradient in glass Fe concentrations at the interface (Fig. 8), suggesting equilibrium between Au capsule and glass. Therefore, alloying of Fe in the noble metal capsules does not delay significantly the attainment of equilibrium between Ir-Fe alloys and melt.

Activities of FeO^{liq} and Fe₂O₃^{liq} were also calculated from independent equilibria. In sample # 2/7 (run # 2, Table 4), results obtained from both activity sensors (Pt capsule and Ir powders) are nearly identical (difference of ~ 1400 Joules, Table 4) and suggest attainment of equilibrium between metal and melt because the Pt capsule and Ir powders are physically separated by the melt. Another test is provided by the magnetite-bearing charges of run # 6 (Table 4). Activities of FeO^{liq} and $Fe_2O_3^{liq}$ were calculated from both the metal-melt (Eqn. 1) and magnetite-melt equilibria (i.e., Fe_3O_4 243 \Leftrightarrow 3 FeO + $\frac{1}{2}$ O₂). For the latter, standard state properties were taken from Table 1 and Lange and Carmichael (1990) for FeO^{liq}, whereas for Fe₃O₄, standard state and mixing properties where calculated from Ghiorso and Sack (1991) and their composition was determined by EMPA. Results show that a_{FeO}^{liq} and a_{Fe2O3}^{liq} are within 10% from each other (samples #6/5, #6/6, Table 4), which is evidence for equilibration of the different iron components between magnetite, Ir-Fe alloy, and melt. The activity of the magnetite component in these crystals, calculated with Ghiorso and Sack (1991), are nearly identical to those calculated from magnetite-metal equilibrium (i.e., $Fe_3O_4 243 \Leftrightarrow 3Fe_3O_4 243$ $+ 2 O_2$). This conclusion demonstrates the internal consistency of the thermodynamic database used to evaluate $\mu_{\rm FeO}^{\rm liq}$, $\mu_{\rm Fe2O3}^{\rm liq}$, $\mu_{\rm Fe}^{\rm alloy}$ and $\mu_{\rm Fe3O4}^{\rm magnetite}$

5.1.3. Global vs. partial equilibrium

In the above section, we have presented evidence for the attainment of metal-melt, hematite-melt, magnetite-melt and magnetite-metal equilibria. It is worth stressing that this applies only to the different iron components (metallic iron in alloy, iron oxide components in melt, magnetite component in magnetite). Global equilibrium (defined as the equality of the chemical potential (μ_i^{ϕ}) of each component (i) in each phase (ϕ)) is not claimed. The absence of Ir in suspension wires or containers and, conversely, the absence of Au, Pt, or Pd in the Ir-Fe alloys is evidence for some disequilibrium in our experiments. Noble metals have very low solubilities in silicate melts (e.g., O'Neill et al., 1995), which would limit the flux of noble metals

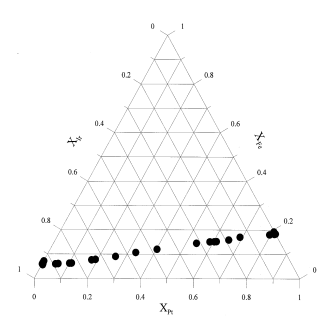


Fig. 9. Non-equilibrated Fe-Ir-Pt alloys and the effect of Pt/Ir on the Fe content of the alloys. The values of a_{FeO}^{liq} were calculated from the fO₂ and the composition of Ir-Fe binary (left, $a_{FeO}^{liq} = 0.081$) and Pt-Fe binary (right, $a_{FeO}^{liq} = 0.077$). See Table 3 and text (5.1.3).

and make their equilibration through the melt over long distances difficult. Although it is quite likely that equilibrium for the noble metals is attained locally between melt and either alloys or suspension wires or containers, equilibrium is clearly not attained between the various noble metal phases, and therefore global equilibrium is not attained. However, partial equilibrium is demonstrated for the different iron components and this is all what is needed for the measurement of their activities.

To illustrate this point, ternary Pt-Ir-Fe alloys were obtained in an experiment performed at high temperatures (1380°C), 1 atm and for 10 d at FMQ. The charge, made of a basaltic composition containing 8.04 wt.% FeO_{tot} plus Ir powders, was suspended to a Pt wire. A glass homogeneous in major elements was obtained (Table 3). Compositions of alloys were found to vary with their spatial position in the charge. Only Fe-Ir alloys were found in the main part of the glass. In a narrow zone (50 μ m) near the Pt wire, alloys with ternary compositions were found. The composition in this zone varies from Ir-free (rim of the Pt wire) to Pt-free (the farther away from the Pt wire, Fig. 9). The experiment is thus out of global equilibrium. Yet, a_{FeO}^{liq} calculated from each end-member binary composition (i.e., Ir-free and Pt-free) are nearly identical (respectively 0.077 and 0.081), suggesting that partial metalmelt equilibrium is attained for the Fe^{metal} and FeO^{liq} components. Differences in alloy Fe contents (Fig. 9) are not ascribed to variations of $\mu_{\rm FeO}^{\ \ liq}$ but rather result from changes in the mixing properties of the alloy controlled by the local Pt/Ir ratio. These ternary alloys were not found in other run products. This is probably due to difference in both run duration and temperature and in viscosity of melts that would affect the flux of noble metals and, therefore, the communication of their chemical potential. Nevertheless, we show that the absence of global equilibrium does not invalidate measurements of molten iron oxides activities.

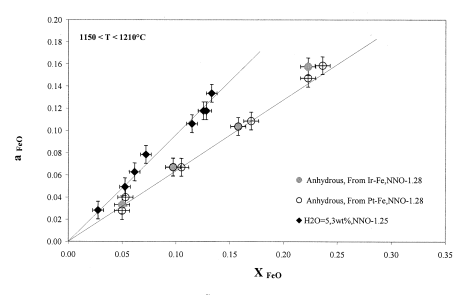


Fig. 10. Activity-compositions relationship for FeO^{liq} in anhydrous and hydrous melts under reducing conditions (Data extracted from # Runs 1-2, see Table 4-5). For the anhydrous compositions, a_{FeO}^{liq} are calculated from both Ir-Fe and Pt-Fe liquid equilibria. The straight line passing through the one atm data is an extrapolation of a_{FeO}^{liq} measured by Doyle et al. (1989).

5.2. Activity Results

The experimental conditions are presented together with the activities of FeO^{liq} and Fe₂O₃^{liq} (a_{FeO}^{liq}, a_{Fe2O3}^{liq}) in Table 4. The tabulated experiments satisfy to the conditions of equilibrium, except for runs # 4 and 5 (local metal-melt equilibrium, see above and Fig. 6). Neither non-equilibrated time-series experiments nor runs having experienced failure of the Co-Ni-Pd redox sensor are reported. The electron microprobe data for glasses, hematite, magnetite and alloys, together with the determinations of H₂O contents and the Fe³⁺/Fe²⁺ data are

given in Table 5. For the calculation of the mole fractions of FeO^{liq} and Fe₂O₃^{liq} (X_{FeO}^{liq} and X_{Fe2O3}^{liq}), the melt compositions were recast in terms of the following components: Si₄O₈, KAIO₂, NaAIO₂, CaAl₂O₄, Al₂O₃, CaO, H₂O, OH, FeO, Fe₂O₃. Glasses in experiments performed with the Ca-free starting compositions have various FeO_{tot} but nearly identical Si₄O₈, KAIO₂ and NaAIO₂ contents and all but 1 or 2 are peraluminous (Table 5). Therefore, a_{FeO}^{liq} and a_{Fe2O3}^{liq} are varied at constant composition of the aluminosilicate matrix. In experiments performed with the Ca-bearing starting composi-

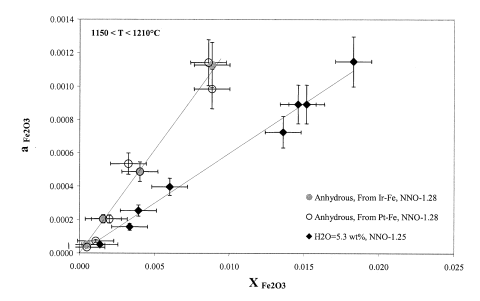


Fig. 11. Activity-compositions relationship for $Fe_2O_3^{liq}$ in anhydrous and hydrous melts under reducing conditions (Data extracted from # Runs 1-2, see Table 4-5). For the anhydrous compositions, a_{FeO}^{liq} are calculated from both Ir-Fe and Pt-Fe liquid equilibria.

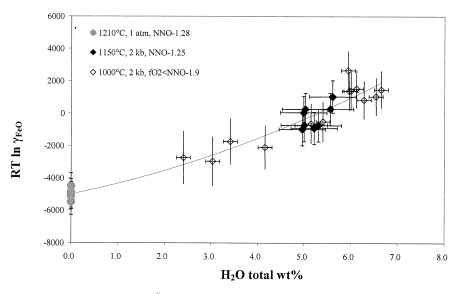


Fig. 12. Excess chemical potential of FeO^{liq} as a function of the water content under reducing conditions (Data extracted from # Runs 1-2-3, see Table 4-5). Water content is determined by difference on EMPA (black diamond-shaped) and FTIR (hollow diamond-shaped). The curve is a polynomial fit on the data.

tions, variations in a_{FeO}^{liq} and a_{Fe2O3}^{liq} are coupled with differences in the composition of the aluminosilicate matrix.

5.2.1. The effect of water on a_{FeO}^{liq} and a_{Fe2O3}^{liq}

For moderately reducing conditions (i.e., from NNO - 1 to NNO - 1.5) and under anhydrous conditions at 1210°C (run # 1), glass FeO_{tot} ranges between 2.10 and 11.85 wt.% (Table 5). Iron redox ratios (XFe₂O₃/XFeO) are very low (from 0.01 to 0.07) and positively correlated with FeO_{tot} (Table 5). The activity-composition relations for the anhydrous melts are shown in Figure 10. The agreement between activities determined from either the Ir-Fe or Pt-Fe (suspension wire) sensors is excellent. The anhydrous data show that a_{FeO}^{-liq} we measured is very close to the activities determined by Doyle (1988)

(straight line in Fig. 10) with $\gamma_{\text{FeO}}^{\text{liq}} = 0.7 (\gamma_{\text{FeO}}^{\text{liq}} = a_{\text{FeO}}^{\text{liq}}/X_{\text{FeO}}^{\text{liq}}$. The very good agreement between the Doyle's and our $a_{\text{FeO}}^{\text{liq}}$ emphasizes the consistency between our research and the previous studies devoted to measurement of thermodynamic properties of molten iron oxides in silicate melts. The effect of the addition of H₂O is apparent from run # 2 performed at 1160°C, 2 kbar, and under fO₂ conditions, NNO-1.25, very close to that of run # 1 (Table 4). The glasses (1.78 - 10.64 wt.% FeO_{tot}) have H₂O concentrations ranging from 4.96 to 5.61 wt.% (Table 5). The redox ratios of iron in these glasses (0.05 to 0.13) are significantly higher than in run # 1, although the fO₂ is nearly similar, and show a positive correlation with FeO_{tot} (see also Gaillard et al., 2001). Activities of FeO^{liq} follow Henry's law but $a_{\text{FeO}}^{\text{liq}}$ is higher in the

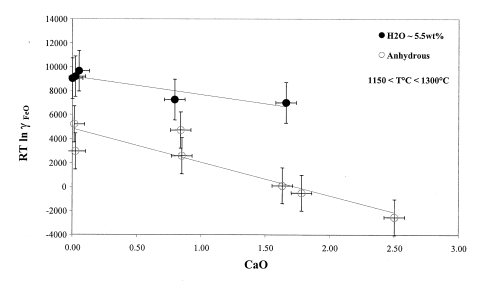


Fig. 13. Excess chemical potential of FeO^{liq} as a function of the calcium content under oxidizing conditions (Data extracted from # Runs 7-8-9, see Table 4-5).

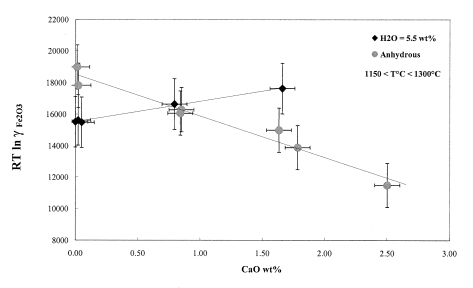


Fig. 14. Excess chemical potential of $Fe_2O_3^{liq}$ as a function of the calcium content under oxidizing conditions (Data extracted from # Runs 7-8-9, see Table 4-5).

hydrous than in the anhydrous samples at a given X_{FeO}^{liq} (with $\gamma_{FeO}^{liq} = 0.7$ for anhydrous samples against 1 for the hydrous series). In contrast, for $Fe_2O_3^{liq}$ (Fig. 11), addition of H_2O decreases a_{Fe2O3}^{liq} at a given X_{Fe2O3}^{liq} in comparison with a_{Fe2O3}^{liq} of anhydrous glasses under similar T-fO₂ conditions. The combined effect of water on a_{FeO}^{liq} and a_{Fe2O3}^{liq} implies that the ratio $\gamma_{Fe2O3}^{liq}/\gamma_{FeO}^{liq}$ is lower in hydrous than in anhydrous melts. This observation is consistent with the increase of the iron redox ratio (XFe₂O₃/XFeO) noted above for the hydrous glasses from run # 2 in comparison with the anhydrous glasses from run # 1 (see also Baker and Rutherford, 1996; Gaillard et al., 2001).

Constraints on the effect of H_2O on a_{FeO}^{Iiq} under isobaric conditions (2 kbar) are provided by run # 3 (Table 4). Glass

H₂O contents range between 2.40 to 6.65 wt.% (Table 4–5). Because of variations in H₂O concentration (and hence in fH₂O), fO₂ differs between samples, from NNO–1.91 to NNO–3.16 (Table 4). Under these conditions, very little Fe is present as Fe³⁺ and glass FeO_{tot} (0.70–7.46 wt.%) are equivalent to their FeO concentrations. For all samples from runs # 1–3, the excess chemical potential of FeO^{liq} ($\mu^{xs}_{FeO}^{Iiq} = RT \ln \gamma_{FeO}^{Iiq}$) is plotted against the water content (Fig. 12). Despite differences in T, P, fO₂ and FeO_{tot} between samples, the data consistently show that the addition of water progressively increases $\mu^{xs}_{FeO}^{Iiq}$. The deviation from ideality changes from negative at low and intermediate water contents to slightly positive above ~5 wt.% H₂O (Fig. 12).

Under strongly oxidizing conditions (NNO+4.9 < fO₂ <

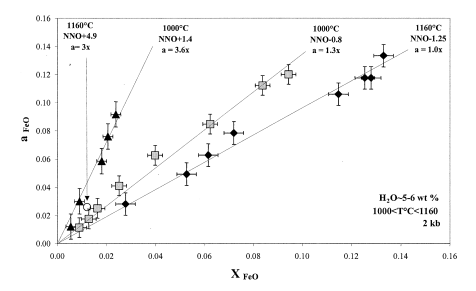


Fig. 15. The effect of fO_2 on activity-composition relationships for FeO^{liq} in H₂O-saturated melts at 2 kb. Data are extracted from # Runs 2-4-6-9 (See Table 4-5). The lines are best fit to the iso- fO_2 data and the resulting a-X relationships are given.

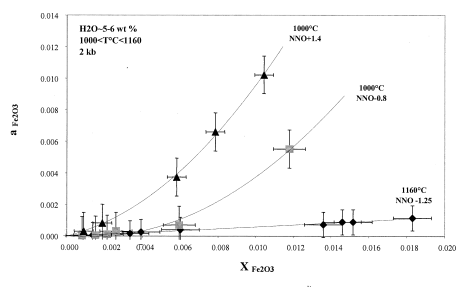


Fig. 16. The effect of fO₂ on activity-composition relationships for $Fe_2O_3^{liq}$ in H₂O-saturated melts at 2 kb. Data are extracted from # Runs 2-4-6 (See Table 4-5). Results from # Run 9 are not plotted here since it would have drastically changed the scale (see text).

NNO+7.3), the glasses are characterized by a narrow range of FeO_{tot} (2.45-3.15 wt.%) reflecting saturation with hematite (samples # 7/4, 8/3, 9/2 to 9/4, note that sample 9/1 is undersaturated in hematite, Table 4). In detail, the hydrous glasses saturated with respect to hematite at 1160°C have a higher FeO_{tot} than the anhydrous glasses at 1245°C, which shows that water (in the range 5.55 to 6.06 wt.% H₂O, Table 5) increases the solubility of hematite at high fO₂. For both series, glasses are strongly oxidized (XFe₂O₃/XFeO = 1.55-1.85, Table 5) and it is difficult to identify any effect of water on the ferricferrous ratio. The excess chemical potential of FeO^{liq} indicates positive deviation from ideality and seems slightly higher in the hydrous than in the anhydrous melts (Fig. 13). However, because glass FeO contents are low under such oxidizing conditions, errors in the determination of X_{FeO}^{liq} are important making estimation of the effect of water difficult. Excess chemical potentials of $Fe_2O_3^{liq}$ for hydrous and anhydrous melt overlap, so that the effect of water on $\gamma_{\text{Fe2O3}}^{\text{liq}}$ for strongly oxidizing fO_2 is not very marked (Fig. 14).

The FTIR results reveal that the water solubility decreases with increasing iron content. This is particularly obvious for runs 4 and 6 (Table 5) where the water solubility decreases from 5.2 to 4.5 wt.% as iron content varies from 0.6 to 7.2 wt.% FeO. However, because the effect of iron content on the extinction coefficient of OH and H_2O bands in glass is currently unknown, we consider that this observation, to be interpreted, needs confirmation by an independent method of water content measurement, which is beyond the scope of this study.

5.2.2. The effect of fO_2 on a_{FeO}^{liq} and a_{Fe2O3}^{liq} of hydrous melts

Redox conditions dominate the activities of both FeO^{liq} and Fe₂O₃^{liq} in H₂O-saturated melts synthesized at 2 kbar. For FeO^{liq}, activity-composition relationships are shown in Figure 15 for 4 fO₂ ranging from NNO-1.25 to NNO+4.9, and at 2

temperatures (1000 and 1160°C). For all oxyisobars (O2 isobars), a_{FeO}^{liq} follows Henry's behavior. The Henry's law constant (constant activity coefficient, $\gamma_{\text{FeO}}^{\text{liq}}$) is positively correlated with fO₂, increasing from a value of ~ 1 at NNO-1.25 (see also Fig. 10) to \sim 4 at NNO+1.4. The point for NNO+4.9 (which corresponds to 3 nearly identical measurements, run # 9, Table 4) clearly shows that the positive dependence of $\gamma_{FeO}{}^{liq}$ on fO₂ does not apply for fO₂ above NNO+1.4. There should exist a maximum in $\gamma_{FeO}{}^{liq}$ for a fO₂ between NNO+1.4 and NNO+4.9. Activity-composition relationships for Fe₂O₃^{liq} are shown on Figure 16. As observed for FeO^{liq}, γ_{Fe2O3}^{liq} has positive fO₂ dependence. The a_{Fe2O3}^{liq} - X_{Fe2O3}^{liq} relationships, which are nearly ideal at NNO+1.4, show a strong negative deviation at NNO-1.25. At NNO-0.8 and NNO+1.4, a_{Fe2O3}^{liq} does not vary linearly with X_{Fe2O3}^{liq} and therefore $Fe_2O_3^{liq}$ does not follow Henry's law in the entire fO_2 range covered by Figure 16. The point at NNO+4.9 (run # 9, Table 4) plots well outside Figure 16 ($a_{Fe2O3}^{liq} = 0.069 - 0.070$ for $X_{Fe2O3}^{liq} = 0.02$). This demonstrates that the increase of $\gamma_{\rm Fe2O3}^{\rm liq}$ with fO₂ continues above NNO+1.4, in contrast to the behavior of FeO^{liq}.

5.2.3. The effect of Ca on a_{FeO}^{liq} and a_{Fe2O3}^{liq}

Comparison between the Ca-free and Ca-bearing runs at high fO2 (between NNO+4.9 and NNO+7.3) allows the effect of the Ca = K + Na substitution to be demonstrated (Fig. 13). Important differences appear between the 2 kbar hydrous and 1 atm anhydrous samples. FeOtot concentrations in hydrous glasses at equilibrium with hematite decrease with increasing CaO (run # 9, Table 4) whereas the reverse is observed for anhydrous glasses (runs # 7, 8, Table 4). The ferric-ferrous ratio decreases with increasing CaO for the hydrous glasses (samples 9/2 to 9/6, Table 4) but does not vary significantly for anhydrous glasses (samples 7/1 to 7/4 and 8/1 to 8/3, Table 4). Excess chemical potentials of FeO^{liq} decrease with CaO for

both hydrous and anhydrous compositions (Fig. 13), and the effect of water on μ_{FeO}^{xs} ^{liq} appears to be more marked as CaO increases. In contrast, the effects of CaO on RT ln γ_{Fe2O3}^{liq} for hydrous and anhydrous melts are barely distinguishable from each other except at the highest CaO concentrations (Fig. 14).

6. DISCUSSION AND CONCLUSIONS

6.1. Water and Fe³⁺/^Fe²⁺

Here above we have put in evidence different effects of water on $a_{\rm FeO}^{~~liq}$ and $a_{\rm Fe2O3}^{~~~liq}$ that are both fO_2 dependent. In terms of Fe₂O₃/FeO - fO₂ relationships, our results suggest that water addition increases ferric/ferrous under reducing conditions and has a less important or no effect under oxidizing conditions. This is in very good agreement with previous experimental studies (Baker and Rutherford, 1996; Gaillard et al., 2001) identifying similar effects of water. The recent results of Wilke et al. (2002) do not follow this trend probably due to the Na-poor and Ca-rich composition they used. Our results indeed suggest that substitution of Na by Ca should increase the activity coefficient of ferric iron in hydrous melts and therefore decreases the Fe^{3+}/Fe^{2+} of the melt. Such an effect of Ca on Fe^{3+}/Fe^{2+} of hydrous melt was also identified by Gaillard et al. (2001) on the Pinatubo glass matrix sample. The results of Moore et al. (1995) do not follow the trend we observed. Indeed, their conclusions suggested that water has no measurable effect on Fe³⁺/Fe²⁺ of silicate melts. However, their experiments were essentially conducted on peralkaline melts in contrast to our and Baker and Rutherford (1996)'s studies that concern metaluminous systems. We therefore anticipate that the conclusions we draw about the effect of water on the energetic of molten iron oxides in metaluminous melts cannot be extrapolated to peralkaline molten silicates.

6.2. Conclusive Remarks

The effect of water and fO_2 on a-X relationships in rhyolitic melts has been elucidated by equilibrating Fe-Ir, Fe-Pt alloys and hematite with molten silicates. We showed that under reducing conditions, water increases the activity coefficient of ferrous iron and has an opposite effect on ferric iron. Therefore, water incorporation in melts under reducing conditions result in a slight increase of Fe³⁺/Fe²⁺. Under oxidizing conditions, no significant effect of water can be measured. In contrast, a-X relationships for both ferrous and ferric iron components are strongly positively deviated as fO_2 increases. For FeO^{liq}, activity coefficient changes from ideality at QFM to ~4 at NNO+1.5. Combining these results with the previous works, a database covering compositions and conditions encountered in natural magmas is available.

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APPENDIX: DETERMINATION OF LIQUID FeO STANDARD PROPERTIES

Thermodynamic properties of pure liquid FeO (FeO^{liq}) were extracted from a numerical interpretation of the phase relations in the pseudobinary system SiO₂-FeO +/- Fe₂O₃. This system is constrained by the phase diagram of Bowen and Schairer (1934) and numerous measurements of FeO^{liq} activities in the liquid (61 metal-silicate equilibria compiling the works of Schumann and Ensio, 1951 and Bod-sworth, 1959). These activity measurements constrain the energetic of iron in the part of the system containing from 60 to 100 wt.% of molten iron oxides (mainly FeO with Fe₂O₃ increasing from ~0 to 13wt.% as total iron oxides increases from 60–100%). Precise extrapolation of the thermodynamic properties of pure *FeO*^{liq} is thus feasible. The excess Gibbs free energy of the system (G_{xs}) was fitted using Margules formalism with 3 interaction parameters between liquid SiO₂^{liq} and

Table A1. Optimization in the system $FeO-SiO_2\pm Fe_2O_3$. Upper Table: Gibbs free energy formula of the components used for the thermodynamic optimization of the molten system $SiO_2-FeO\pm Fe_2O_3$. Middle Table: Stoichiometry of equilibria used and corresponding evaluated component. Lower Table: Results of optimization.

Reference		Components		ΔG° formation at 1bar, $T^\circ K$ (J/				
Coughlin (1954)		Wustite solid		-265020 + 64.8 .(T)				
Coughlin (1954)		Wustite liquid		-237156 + 47	7.64 .(T)			
Robie et al (1978)		Si, O ₂						
Berman et Brown (1987)		Crystobalite		-898867 + 17	0.22 .(T)			
Berman et Brown (1987)		Tridymite		-902880 + 17	4.2 .(T)			
DeCapitani et Kirschen (1998)		SiO ₂ liquid	-891808 + 16	6.11 .(T)				
O'Neill (1987)		Fayalite $-1450800 + 30.26$.						
Equilibria				Evalua	ted Component			
$SiO_2^{cristobalite-tridymite} = SiO_2^{liq}$				SiO				
$Fe_2 \tilde{SiO}_4^{fayalite} = 2 FeO^{liq} + SiO_2$	liq			FeO	^{Ĩiq} & SiO ₂ ^{liq}			
$Fe_{0.947}^2 \vec{O}^{wustite} = 0.947 \text{ FeO}^{liq} + \vec{O}^{liq}$	$(1-0.947)/2 O_2^{gas}$			FeO				
$\operatorname{Fe}^{\operatorname{metal}} + \frac{1}{2} \operatorname{O}_2 = \operatorname{FeO}^{\operatorname{liq}}$, 2			FeO	liq			
Standard state for FeO ^{liq} *		Ν	Margules parameters for	or the excess free energy				
$\Delta H_{f}^{o}{}_{FeO}^{liq*}$ ΔS_{f}^{o}	e liq* FeO	W ₁₃	W ₁₂₃₄	W ₁₂₂₃	W ₁₁₁₂			
-226244 -42	49	-125350	57419	-16259	-10128			

* $\Delta G_{f_{FeO}}^{\circ}{}^{liq} = \Delta H_{f_{FeO}}^{\circ}{}^{liq} - T. \Delta S_{f_{FeO}}^{\circ}{}^{liq}$; Enthalpy and entropy of formation of FeO^{liq} applicable in the temperature range 910°C–1392°C. **Gxs = $X_{FeO}^{3}X_{SiO2}W_{1112} + X_{FeO}^{2}X_{SiO2}^{2}W_{1223} + X_{FeO}X_{SiO2}^{3}W_{1234} + X_{FeO}X_{FeO}X_{Fe2O3}W_{13}$

At saturation of Fe, wustite, cristobalite, tridymite, fayalite:

Average difference between measured-calculated chemical potentials ($\mu_{\text{FeO}}^{\text{liq}}$, $\mu_{\text{SiO2}}^{\text{liq}}$): 645 J/mol.

FeO^{liq} and a symmetric Margules formulation describing the interactions between FeO^{liq} and Fe₂O₃^{liq} (see Table hereafter) Interactions between SiO₂^{liq} and Fe₂O₃^{liq} were ignored given that the content of Fe₂O₃^{liq} at the liquidus of Si-bearing crystals are negligible and that ferric iron become an important species of the system when silica is present only at very low concentration (see Bowen and Schairer, 1934). The optimization was realized using a minimizing function constrained with inequalities allowing but restraining variations of liquidus temperatures of solids (see Berman and Brown, 1984, 1987 for LAP method). As initial values, the standard state molten wustite was adopted for liquid FeO (Coughlin, 1954 see Table 1). Throughout the minimization, both enthalpy and entropy of formation of liquid FeO were allowed to vary within 20%. The results including the thermodynamic properties of liquid FeO, the interaction parameters defining G_{xs} and the accuracy of the model are given in the following Table for liquid FeO and liquid SiO₂. The evaluated standard state for liquid FeO are very close to the standard state adopted by Schumann and Ensio (1951), Bodsworth (1959) and Matsuzaki et al. (1998) but significantly differ from the thermodynamic parameter of liquid wustite (see text and Table 1).