Structure of NaFeSiO₄, NaFeSi₂O₆, and NaFeSi₃O₈
glasses and glass-ceramics

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ABSTRACT

The crystallization of iron-containing sodium silicate phases holds particular importance, both in the management high-level nuclear wastes and in geosciences. Here, we study three as-quenched glasses and their heat-treated chemical analogues, NaFeSiO$_4$, NaFeSi$_2$O$_6$, and NaFeSi$_3$O$_8$ (with nominal stoichiometries from feldspathoid, pyroxene, and feldspar mineral groups – i.e., Si/Fe = 1, 2, and 3 respectively) – using a variety of techniques. Phase analyses revealed that as-quenched NaFeSiO$_4$ cannot accommodate all Fe in the glass phase (some Fe crystallizes as Fe$_3$O$_4$), whereas as-quenched NaFeSi$_2$O$_6$ and NaFeSi$_3$O$_8$ form amorphous glasses upon quenching. NaFeSi$_2$O$_6$ glass is the only composition that crystallizes into its respective isochemical crystalline polymorph, i.e. aegirine, upon isothermal heat-treatment. As revealed by Mössbauer spectroscopy, iron is predominantly present as 4-coordinated Fe$^{3+}$ in all glasses, though it is present as 6-coordinated Fe$^{3+}$ in the aegirine crystals (NaFeSi$_2$O$_6$), as expected from crystallography. Thus, Fe can form the crystalline phases in which it is octahedrally coordinated, even though it is mostly tetrahedrally coordinated in the parent glasses. Thermal behavior, magnetic properties, iron redox state (including Fe K-edge X-ray absorption), and vibrational properties (Raman spectra) of the above compositions are discussed.

Keywords: Mössbauer, Fe redox, Raman, glass transition
INTRODUCTION

Crystallization of iron-containing sodium silicate phases is important, both in the management of high-level nuclear wastes and in geosciences (Ahmadzadeh et al., 2017; Bailey and Schairer, 1963; Bailey and Schairer, 1966; Cochain et al., 2012; Jantzen, 2011; Jantzen and Brown, 2007; Jeoung et al., 2001). The complex high-level nuclear wastes (HLW) stored in steel tanks at the Hanford site can contain more than 20 elements, among which iron (Fe) concentrations vary from about 5 to more than 30 wt% Fe$_2$O$_3$, for high-Al and high-Fe HLW, respectively (Kim et al., 2011; Kruger et al., 2013). Within the high-Fe wastes, Na$_2$O concentration varies from about 10 to 20 wt% depending on the cluster (Kim et al., 2011). Vitrification is the process used to immobilize radioactive HLW by converting it into a solid stable glass. SiO$_2$ is added in considerable amounts (with lower levels of other additives) to vitrify the radioactive HLW into a glass for immobilization. Consequently, HLW glasses have high concentrations of Si, Fe, and Na, and are thus potentially prone to crystallization of iron sodium silicate phases.

In particular, aegirine (NaFeSi$_2$O$_6$, also known as acmite), which is a clinopyroxene silicate phase, has been known to crystallize within some HLW glasses (Hrma et al., 1999; Jantzen and Edwards, 2015; Jantzen and Bickford, 1984; Jantzen et al., 1984; Vienna et al., 1996). Studying the crystallization of more than 100 simulant HLW glass compositions, Kim et al. (1994) reported that aegirine forms upon isothermal heat-treatment of glasses with high Na$_2$O (>10 wt%) and Fe$_2$O$_3$ (>7 wt%) contents, while aegirine is not observed in samples that have been canister-centerline-cooled (slow cooling profile recorded at the centerline of the Hanford HLW canisters). Surface crystallization of aegirine from simplified HLW glasses was confirmed by Plaisted et al. (2000), who showed that the aegirine contains other elements such as Cr and Ni.
and that its composition varies by temperature of the heat treatment. The formation of aegirine causes a small to moderate decrease in the chemical durability of final HLW glass waste form (Jantzen and Bickford, 1984; Jantzen et al., 1984; Jantzen et al., 2010). The spinel formation (i.e., magnetite $\text{Fe}_3\text{O}_4$ and related phases), however, has little or no effect on glass durability, whereas nepheline ($\text{NaAlSiO}_4$) and related aluminosilicate phases have the most detrimental impacts on aqueous chemical durability of crystallized HLW glass.

Iron and sodium are among the most common constituents of natural silicate melts, and both can have remarkable effects on their physical properties. Bailey and Schairer (1966) have extensively described how equilibrium crystalline and liquid phases in the system $\text{Na}_2\text{O}-\text{Al}_2\text{O}_3-\text{Fe}_2\text{O}_3-\text{SiO}_2$ are petrologically important for a wide range of alkaline igneous rocks. These crystalline phases include aegirine ($\text{NaFeSi}_2\text{O}_6$), 5.1.8 ($5\text{Na}_2\text{O} \cdot \text{Fe}_2\text{O}_3 \cdot 8\text{SiO}_2$ or $\text{Na}_5\text{FeSi}_4\text{O}_{12}$), nepheline ($\text{NaAlSiO}_4$, hexagonal), carnegieite ($\text{NaAlSiO}_4$, orthorhombic), albite ($\text{NaAlSi}_3\text{O}_8$), sodium metasilicate ($\text{Na}_2\text{SiO}_3$), hematite ($\text{Fe}_2\text{O}_3$), and different polymorphs of $\text{SiO}_2$ (quartz, tridymite, and cristobalite). In such a system, $\text{Fe}^{3+}$ can potentially behave similarly to $\text{Al}^{3+}$, though iron can play a special role; each valence state of iron plays a different structural role, and the redox ratio ($\text{Fe}^{3+}/\text{Fe}^{2+}$), can consequently influence the melting and crystallization behavior of magmas and properties of igneous rocks (Mysen and Richet, 2005). Hence, studying the speciation and structural role of Fe is crucial in understanding the structure of iron-containing silicate melts and glasses and their crystallization behavior (Komatsu and Soga, 1980).

Aegirine, a chain silicate from the clinopyroxene group, is the most well-studied sodium iron silicate phase. It melts incongruently, with separation of hematite at 990°C (Bowen et al., 1930) and has a monoclinic symmetry with space group $C2/c$ (Clark et al., 1969). Aegirine is the mineral name for the Na-Fe$^{3+}$ end-member of the class of rock-forming pyroxene minerals which...
in nature typically contains some impurities of Al, Cr, and Ca. The mineral aegirine is less common than other mixed metal Ca-Mg-Fe pyroxenes, such as augite, but is an important minor component (Bailey and Schairer, 1966; Bowen et al., 1930; Deer et al., 1992; Larsen, 1976). Another sodium iron silicate phase, known as 5.1.8 (Na$_5$FeSi$_4$O$_{12}$), has not been reported to occur in natural rocks. The “5.1.8” composition was first discovered by Bowen et al. (1930), and is extensively studied in our previous work (Ahmadzadeh et al., 2018).

In the sodium alumino-silicate groups, there are three important types of well-known minerals based on their SiO$_2$ content; nepheline or carnegieite (NaAlSiO$_4$, feldspathoid), jadeite (NaAlSi$_2$O$_6$, pyroxene), and albite (NaAlSi$_3$O$_8$, feldspar). Aegirine is the iron “equivalent” of jadeite, with the same crystal structure in which Fe$^{3+}$ substitutes for Al$^{3+}$ depending upon the pressure, though end-member jadeite is difficult to form except under extreme conditions (Nestola et al., 2007). NaFeSiO$_4$ (Fe-nepheline) and NaFeSi$_2$O$_8$ (Fe-albite) do not exist as known crystalline phases.

The present work describes crystallization studies of the three feldspathoid-, pyroxene-, and feldspar-type sodium iron silicate compositions. The structure of the parent glasses and the corresponding phases which form upon heat-treating the glasses are discussed. It should be noted that controlling the melting and/or heat-treatment atmosphere, and the subsequent Fe oxidation state, was not the aim of this work. The crystallization behavior of these compounds and the role of iron are presented through thermal analysis, X-ray diffraction, Raman, Mössbauer and X-ray absorption spectroscopies, and magnetic property measurements.

**MATERIALS AND METHODS**

Glasses nominally isochemical with the compositions of NaFeSiO$_4$ (‘Fe-nepheline’), NaFeSi$_2$O$_6$ (aegirine), and NaFeSi$_3$O$_8$ (‘Fe-albite’) were synthesized from Na$_2$CO$_3$ (Fisher
Scientific, >99%), Fe$_2$O$_3$ (Alfa Aesar, 98%), and SiO$_2$ (U.S. Silica, 99.7%) powders as starting materials. Each batch was melted twice for one hour in air in an electric furnace at 1500°C in platinum-10% rhodium crucibles, to make ~ 20 g of glass. Each melt was poured onto an Inconel plate to quench and form a glass. In the case of the Fe-nepheline composition, the melt was quenched more rapidly between two brass plates in order to attempt to avoid formation of Fe$_3$O$_4$ (magnetite), which was not completely achievable. The quenched glasses were then crushed into powders and isothermally heat-treated in the specific conditions selected to attempt to obtain maximum crystallization. The Fe-nepheline glass was heat-treated at 775°C for 7 h based on our previous study (Ahmadzadeh et al., 2017). The aegirine glass was heat treated at 900°C for 24 h based on the available phase diagrams in the literature (Bowen et al., 1930). The Fe-albite glass was heat treated at 1000°C for 24 h due to its higher SiO$_2$ content and known difficulty for nucleating its Al analogue, albite. In another test, this glass was also kept in the furnace for a longer time (72 h) at 900°C in an attempt to form the isochemical crystalline polymorph. All heat-treatments were followed by quenching samples in air. The compositions, melting temperatures, and heat-treatment conditions are summarized in Table 1.

For differential thermal analysis (DTA) and thermogravimetric analysis (TGA), the as-quenched glasses were crushed and sieved to a particle range of 63 to 125 µm. The data were obtained with a heating rate of 10°C min$^{-1}$ under constant N$_2$ flow (99.998% pure) with a SDT Q600–TA Instruments system.

An X’Pert Pro MPD (PANalytical, Netherlands) was employed to collect powder X-ray diffraction (XRD) patterns with Co Kα X-rays (λ = 0.1789 nm) at 40 kV and 40 mA, and data were analyzed using HighScore Plus software (PANalytical, Netherlands). For semi-quantitative
Rietveld refinement and quantification of the phase fractions, powders were mixed with 10 wt % CaF₂ as an internal standard.

Magnetic hysteresis loops were obtained using a vibrating sample magnetometer (VSM, PMC3900, Lakeshore Cryotronics, Westerville, OH) with maximum applied field of 1.8 T.

Room temperature $^{57}$Fe Mössbauer spectra were collected relative to $\alpha$-Fe over a velocity range of ± 12 or ± 6 mm s⁻¹ using a constant acceleration spectrometer with a 25 mCi source of $^{57}$Co in Rh. Attempts were made to fit the spectra using Lorentzian as well as Extended Voigt-Based Fitting (xVBF) paramagnetic doublets consistent with Fe$^{3+}$ and/or Fe$^{2+}$, using the Recoil analysis software package (Rancourt, 1998). The area ratio of the doublets was established, enabling estimation of the (Fe$^{3+}$/ΣFe) redox ratio based on fitted peak areas and assuming that the recoil-free fraction ratio $f(\text{Fe}^{3+})/f(\text{Fe}^{2+}) = 1.0$.

For iron redox ratio analysis, in addition to Mössbauer spectroscopy, a solution-based spectrophotometric method (wet chemistry) was used. The reported average and standard deviation values are obtained from three replicate measurements of each sample. The procedure details are given in (Ahmadzadeh et al., 2018; Weaver et al., 2015) and references therein.

Fe K-edge X-ray absorption near edge structure (XANES) data were collected on the Beamline for Materials Measurement (bending magnet beamline BM-6) at the National Synchrotron Light Source II (NSLS II), Brookhaven National Laboratory, Upton, New York State, USA. The beamline is configured with a Rh-coated paraboloid collimating mirror, a Si (111) monochromator, and a flat harmonic rejection mirror. XANES data were acquired in transmission mode; an array of three ionization chambers, filled with N₂ and operated in a stable region of their current vs voltage curve, enabled concurrent measurement of incident and transmitted beam intensities from both a sample and a reference Fe foil. Measurements were
made in air at room temperature with the sample orientated at an incidence angle of 45° to the beam. Samples were prepared from finely ground powder specimens. These were homogenously dispersed in 60 mg of polyethylene glycol (PEG) and pressed in a 13 mm diameter die, to produce a thickness of one absorption length. Further details on data treatment of the XAS data, including the pre-edge, is described in the supplementary.

Raman spectra were collected on a Jobin Yvon-SPEX Horiba T64000 triple-spectrometer with confocal system through an Olympus BX40 microscope. Samples were excited using a 488 nm Coherent Genesis MX SLM laser running at 500 mW output power. The detector was a liquid nitrogen cooled CCD camera. The spectrometer was calibrated using the 520 cm⁻¹ line of single crystal silicon to within 1 cm⁻¹. Spectra were collected through a 200 x objective lens, from 25-1300 cm⁻¹, for count times of 200 s, with three subsequent spectra averaged.

RESULTS

Phase Analysis

X-ray diffraction patterns were obtained for the as-quenched (AQ) and heat-treated (HT) samples (See supplementary, Figure S1). Table 2 summarizes phase analysis of the three compositions, based on their XRD and Rietveld refinement results. As described in Table 2, depending on the ratio of Si to Fe, the as-quenched samples make either an amorphous phase (for Fe-albite and aegirine) or precipitate some iron as magnetite (for Fe-nepheline). Upon heat-treating the as-quenched samples, they may form hematite and/or aegirine, depending on the composition.

Thermal Analysis

The DTA and TGA results of the three as-quenched glasses are presented in supplementary, Figure S2. Increasing the SiO₂ content
leads to an increase in the glass transition temperature, $T_g$, of the amorphous phases, which is characterized by a subtle endothermic drop in the DTA thermographs, from 471°C in NaFeSiO$_4$, to 498°C in NaFeSi$_2$O$_6$, to 521°C in NaFeSi$_3$O$_8$ glasses. The $T_g$ decrease is probably related to overall decrease in average bond strength, since Fe-O bonds (both with Fe$^{2+}$ and Fe$^{3+}$) are considerably weaker than Si-O bonds (Luo and Kerr, 2006). Other observations about the thermal behavior are given in the supplementary.

**Magnetic Measurements**

Magnetization as a function of magnetic field measurements at room temperature (Figure 1) reveal that Fe-albite and aegirine as-quenched glasses contain no iron oxide (paramagnetic), whereas the as-quenched Fe-nepheline contains considerable amount of magnetite (ferrimagnetic). Further observations are provided in the supplementary.

**Mössbauer Spectroscopy**

Room temperature Mössbauer spectra of Fe-nepheline, aegirine, and Fe-albite as-quenched samples and crystallized aegirine sample were obtained (Figure 2). The fitted Mössbauer spectral parameters are given in the supplementary (Table S1). Aegirine and Fe-albite glasses and crystallized aegirine were fitted by two overlapping doublets, whereas the as-quenched Fe-nepheline was fitted by one doublet plus two sextets characteristic of magnetite (Fe$_3$O$_4$) (Oh et al., 1998). As-quenched aegirine and Fe-albite samples (Figure 2b and c) were fitted by xVBF functions, because at least three doublets were required in order to obtain robust fits using Lorentzian lines, and xVBF has also been successfully used to fit Mössbauer spectra for similar materials (Forder et al., 2013). This method provided robust fits with acceptable $\chi^2$ using only two doublets to fit the glass spectrum, one each representing Fe$^{2+}$ (lower intensity)
and Fe$^{3+}$ (higher intensity), according to their chemical shift (CS) and quadrupole splitting (QS) values obtained from the fits (Dyar, 1985; Dyar et al., 2006; Mysen and Richet, 2005).

Such $^{57}$Fe Mössbauer behavior, shown in Figure 2, is typical for iron-containing silicate glasses (Mysen and Richet, 2005). The fitted CS and QS values also reveal that the Fe$^{3+}$ and Fe$^{2+}$ are predominantly tetrahedrally coordinated in these glasses, though Fe$^{2+}$ can possibly have some five-fold coordination. In the spectrum for the Fe-nepheline as-quenched sample, there is an intense doublet assigned to four-coordinated Fe$^{3+}$ in the glass phase, in addition to the sextets attributed to the magnetite phase.

The crystallized aegirine sample shown in Figure 2d (fitted using Lorentzian peaks) exhibits a new strong doublet, in addition to the primary four-coordinated Fe$^{3+}$ doublet which is also observed in the glass sample. The new doublet, showing significantly lower QS and higher CS (0.40 and 0.31 mm/s, respectively), is due to octahedrally-coordinated Fe$^{3+}$ from crystalline aegirine, as Fe$^{3+}$ arranges 6-coordinated in this phase (Cameron et al., 1973). Considering the experimental error, these values are consistent with the QS and CS values reported in the literature (e.g., in (Baum et al., 1988), 0.39 and 0.33 mm/s, respectively) for pure aegirine crystals at room temperature. Overall, in the studied sodium iron silicate glasses, iron is shown to be predominantly present as four-fold coordinated Fe$^{3+}$ ($^{4}$Fe$^{3+}$), whereas crystal chemistry dictates the iron redox state and coordination in the crystalline phase(s).

Error! Reference source not found. Table S2 (see supplementary) presents the measured iron redox values of the two crystal-free samples, i.e., aegirine and Fe-albite glasses, obtained from the wet chemistry method, compared to those estimated from Mössbauer spectroscopy. Note that the iron redox values from Mössbauer spectroscopy are derived from the xVBF fits. The redox states from both
methods are consistent, and aegirine glass shows slightly higher Fe\textsuperscript{3+}/ΣFe ratio than Fe-albite glass.

**X-ray Absorption**

Figure 3 shows the results of the Fe K-edge data and analysis, including the XANES spectra (a), the extracted pre-edge features (b), and the structure field diagram of pre-edge centroid energy position and integrated intensity (c). The latter diagram gives an indication of Fe redox state and coordination number (CN). Qualitative comparison with the standard spectra for Fe\textsuperscript{3+}, CN=6 (\textsuperscript{6}Fe\textsuperscript{3+}, aegirine, natural mineral, NaFeSi\textsubscript{2}O\textsubscript{6}), Fe\textsuperscript{3+}, CN=4 (\textsuperscript{4}Fe\textsuperscript{3+}, FePO\textsubscript{4}, synthetic), Fe\textsuperscript{2+}, CN=6 (\textsuperscript{6}Fe\textsuperscript{2+}, FeCO\textsubscript{3}, synthetic), and Fe\textsuperscript{2+}, CN=4 (\textsuperscript{4}Fe\textsuperscript{2+}, staurolite, natural mineral, Fe\textsubscript{1.5}Mg\textsubscript{0.5}Al\textsubscript{0.1}Si\textsubscript{3.9}Al\textsubscript{0.1}O\textsubscript{22}(OH)\textsubscript{2}), indicates that the iron in the glasses mostly exists as \textsuperscript{4}Fe\textsuperscript{3+}. In general, the determined centroid energy and integrated intensity of the weak pre-edge features measured from our reference crystalline compounds are in good agreement with those reported by (Wilke et al., 2001), with the exception of our FePO\textsubscript{4} standard for which we determine a greater integrated intensity, most likely due to an overly thick sample resulting in a relatively suppressed edge step.

The weak pre-edge feature is associated with electronic transitions occurring between the 1s and 3d energy levels, and is a feature common to the K-edge spectra of many transition metals (Yamamoto, 2008). Lower oxidation states have fewer unfilled 3d levels, so transitions from the 1s levels become less probable and the intensity of this feature becomes less well-defined; these transitions are only weakly allowed for octahedral Fe since the initial and final states are centrosymmetric; whereas, for Fe in a non-centrosymmetric tetrahedral environment, the transition probability is enhanced due to admixture of unoccupied 4p orbitals in the final state. Previously, careful measurements of crystalline standards have been made (Wilke et al., 2001),
showing the relationship between Fe valence and coordination and pre-edge position and intensity, and further applied to glasses (Farges et al., 2004). It has also been shown that the extraction of Fe redox and coordination number from Fe XAS pre-edge features is complicated by monochromator resolution limitations (Cottrell et al., 2009; Galoisy et al., 2001; Wilke et al., 2005) and fitting method. Recent inter-synchrotron comparison of a set of reference glasses shows that with careful measurements and fitting procedures, an uncertainty in the centroid of ±0.1 eV is achievable (Fiege et al., 2017).

**Raman Spectroscopy**

The raw Raman spectra of the as-quenched glasses are shown in Figure 4. All of these spectra show a background decreasing in intensity from low to high Raman shifts. Di Muro et al. (2009) have studied a number of Fe-containing natural glasses and shown that the background slope of the raw Raman spectra correlates with the total iron content and its redox within the glasses. Figure 4 reveals a consistent trend for our samples; the slope of the background increases with increasing the Fe content and its redox state (Fe$^{3+}/\Sigma$Fe) from Fe-albite to Fe-nepheline.

The Raman spectra of the three as-quenched glasses (Figure 4) show intense low frequency envelopes with at least two bands; a main peak at ~440-460 cm$^{-1}$ and a shoulder at 530-550 cm$^{-1}$. The main peak at ~450 cm$^{-1}$, which was shown to be a polarized band by (Wang et al., 1993) for similar compositions, is assigned to the vibrations of bridging oxygens (BOs), i.e. (T–O–T), where T=Si,Fe, in tetrahedral rings. The high frequency (HF) band of the three spectra consists of a dominant peak at 900-960 cm$^{-1}$ and a shoulder at higher frequencies, i.e., ~1040-1100 cm$^{-1}$. Wang et al. (1993; 1995), have shown for similar compositions that the dominant peak is depolarized whereas the shoulder is polarized, suggesting that the dominant peak is for
the antisymmetric T–O–T stretching vibrations of BOs ($\nu_{as}(T–O–T)$), and the shoulder originates from the symmetric stretching vibrations of NBOs. This is a similar conclusion to that reached earlier by Mysen et al. (1980), who measured an aegirine glass spectrum similar to that shown here, with slightly greater intensity of the higher frequency band, though these authors attributed both bands to stretching of BOs, with one in a Fe-rich and the other in a Si-rich environment.

It can also be seen in Figure 4 that increasing SiO$_2$ content systematically increases the frequency of HF envelope. This shift is related to decreased number of heavier Fe$^{3+}$ tetrahedra (as opposed to lighter Si$^{4+}$ tetrahedra) in the Fe-albite sample which lead to increased vibration frequencies (i.e., Raman shift). The HF envelope contains not only the silicate tetrahedral bands, $Q^0$, where n is the number of bridging oxygens, but also the $[4]$Fe$^{3+}$ related band, generally thought to be ~910-980 cm$^{-1}$ (Baert et al., 2011; Cochain et al., 2008; Di Genova et al., 2017; Magnien et al., 2006), near the lower frequency end of the band and hence weighting the envelope to lower frequencies when the tetrahedral Fe$^{3+}$ concentration is high. This band attributed to $[4]$Fe$^{3+}$ has been reported to shift to lower frequencies as Fe concentration is increased (Cochain et al., 2008), and does here as well (peak of Raman spectrum moves 965 cm$^{-1}$, 950 cm$^{-1}$, 920 cm$^{-1}$, for Fe-albite, aegirine, Fe-nepheline glasses, respectively), though there is some contribution of the silicate stretches. A Raman band attributed to Fe$^{2+}$ contributions has been reported at 1040-1050 cm$^{-1}$ (Cochain et al., 2008; Di Genova et al., 2017); however, the significant Raman scattering observed in this region in the Fe-nepheline glass is due rather to the Si-O stretching, since the peak of the Raman spectrum for a NaAlSiO$_4$ glass lies ~1015 cm$^{-1}$ when measured on the same instrument (not shown).
DISCUSSION

As-quenched glasses

As confirmed by XRD and magnetic measurements, NaFeSi$_2$O$_6$ and NaFeSi$_3$O$_8$ compositions have sufficient Si content to make pure glasses during quenching, while molten NaFeSiO$_4$ cannot readily quench into a glass at our quenching rate, without crystallization of excessive Fe as magnetite. Mössbauer spectroscopy revealed that iron in these glasses is predominantly present as tetrahedrally coordinated Fe$^{3+}$, while some Fe$^{2+}$ with higher coordination may also exist. Preparation of silicate glasses in which 100% of the iron is present as Fe$^{3+}$ can only be accessed under normal laboratory melting conditions if oxidizing agents (for example CeO$_2$) are added to the batch (Bingham et al., 2014), or if glasses with high basicity (i.e., high alkali contents) are used. Similar Mössbauer results to ours are reported by Fleet et al. (1984) and Henderson et al. (1984) for Fe-albite glass melted at 1450°C and 1 bar, though they used Lorentzian functions to fit their data, possibly resulting in lower reduced iron (~9% Fe$^{2+}$) than our measurements (~19% Fe$^{2+}$). Using neutron diffraction experiments and Empirical Potential Structure Refinement (EPSR) modeling, Weigel et al. (Weigel et al., 2008b; Weigel et al., 2006) have discussed that, in their aegirine glass melted at 1450°C for 30 min in air, the majority of Fe$^{3+}$ (~95%) is four-coordinated with oxygen, some Fe$^{3+}$ (~5%) is five-coordinated, and all Fe$^{2+}$ is five-coordinated. Their results also showed that $^{[4]}$Fe$^{3+}$ is randomly distributed in the glass network, acting as a network former. These temperatures are sufficiently close to our melting temperature (1500°C) that redox ratios obtained upon quenching should be similar (~12% Fe$^{2+}$ (Weigel et al., 2008a) vs. ~15% Fe$^{2+}$ our measurements), if redox equilibrium with the surrounding atmosphere was obtained or approached in both cases.
The XAS pre-edge results of our as-quenched glasses shown in Figure 3 indicate that, while iron primarily exists as Fe$^{3+}$ based on the centroid positions of the glasses, the centroids are slightly shifted towards Fe$^{2+}$ (lower energies) compared to the reference compounds. This is consistent with the presence of some Fe$^{2+}$ obtained from Mössbauer and wet chemistry colorimetric data, and consistent with the ‘mixing curves’ for $[^4]$Fe$^{3+}$ and $[^4]$Fe$^{2+}$ in silicate glasses (Jackson et al., 2005). However, no difference in the position of the centroids of the three studied glasses can be ascertained. This implies that the small differences between redox of these three materials are not distinguishable from the pre-edge results at this spectrometer resolution. Indeed, the shift from the Fe redox glass standards recently reported (Fiege et al., 2017) suggests that at most a 0.2 eV shift would be expected for glasses with Fe$^{3+}/∑$Fe of 80 to 100%. Moreover, the integrated intensities of the glasses, as shown in Figure 3-c, do not vary sufficiently to be able to make a meaningful interpretation of change, if any, of Fe coordination numbers. Our XAS data therefore demonstrate the speciation of Fe in the glasses to be predominantly $[^4]$Fe$^{3+}$ with some Fe$^{2+}$, probably $[^4]$Fe$^{2+}$, contribution. It should be noted, however, that some have suggested that Fe coordination in silicate glasses is between 4 and 5 for both Fe$^{3+}$ and Fe$^{2+}$ (Bychkov et al., 1993; Farges et al., 2005; Liu and Lange, 2006).

Fe redox ratio in melts is a function of temperature, pressure, composition, and oxygen fugacity (Mysen and Richet, 2005). We calculated the Fe redox ratio of the present glasses based on the equation provided by Jayasuriya et al. (2004), assuming the oxygen fugacity of 10$^{-0.7}$ for air. The Fe$^{3+}/∑$Fe values predicted are 66.3% (Fe-albite), 72.5% (aegirine), compared to measured values of 81.5% (Fe-albite, wet chemistry) and 83.3% (aegirine glass, wet chemistry) (Table S2). Other classic equations such as that of Sack et al. (1980) was also used, but the Fe$^{2+}$
computed is larger than those of Jayasuriya equation (Jayasuriya et al., 2004), and the latter is
closer to the measured values for the current compositions.

Our Fe-nepheline as-quenched sample only shows Fe\textsuperscript{3+} in the glass according to the
Mössbauer results, unlike the other two compositions, while the calculations for this composition
show Fe\textsuperscript{3+}/ΣFe of 81.9% for the melt at 1500°C. In the Fe-nepheline melt, the present Fe\textsuperscript{2+} in the
melt partitioned into magnetite (Fe\textsuperscript{2+}O · Fe\textsuperscript{3+}O\textsubscript{3}) while it was being quenched, and only Fe\textsuperscript{3+}
remained in the glass phase. If partial crystallization takes place during quench, the redox ratio of
the glass is different from that of its parent melt (Mysen and Richet, 2005).

Increasing Fe\textsuperscript{3+}/ΣFe with increasing iron content observed in these iron-bearing silicate
glasses is partially due to the following. Having less SiO\textsubscript{2} as the main glass former encourages
the iron (which is now at a larger fraction) to take a network forming role, and thus Fe\textsuperscript{3+}.
Systematic decrease in the vibrational frequency of the HF Raman band by decreasing the Si/Fe
ratio confirms increased number of heavier Fe\textsuperscript{3+} network formers (i.e., tetrahedra). The same
Raman trend has been reported by Wang et al. (1993) for similar compositions. They have
debated that introducing more Fe\textsuperscript{3+} at Si\textsuperscript{4+} sites in the glass network leads to a decrease in the
angle of T–O–T linkage which results in a bond length increase. As a result, in the Fe-albite
sample with a higher Si/Fe ratio, the frequency of the antisymmetric vibrations of BOs increases
(HF band shifts to higher frequencies).

The fraction of iron that exists as Fe\textsuperscript{2+} acts as network modifier (similar to Mg\textsuperscript{2+} and Ca\textsuperscript{2+}
in silicate glasses), resulting in network depolymerization. Moreover, even though Fe\textsuperscript{3+} is
normally a tetrahedral network former, it can also act as octahedral network modifier, even when
other cations can compensate the charge balance for its tetrahedral coordination (Mysen and
Richet, 2005). Therefore, while in some papers the three compositions studied here are assumed
to have “fully polymerized” network, a fraction of their oxygens is actually non-bridging (unlike the Al equivalent compositions), due to some Fe acting as network modifier. Even though some Fe crystallizes into magnetite during NaFeSiO$_4$ quench, the Si/Fe ratio in the glass still remains lower than that of NaFeSi$_2$O$_6$ and NaFeSi$_3$O$_8$, because the formed magnetite is only ~7.5 wt% (Table 2). The estimated residual glass composition for the Fe-nepheline as-quenched glass is ~NaFe$_{0.84}$Si$_x$.

Changes on heat treatment

Upon the heat-treatment of Fe-nepheline at 775°C for 7 h, the magnetite phase (spinel, Fe$_3$O$_4$) oxidizes to hematite (Fe$_2$O$_3$), and some Fe may be rejected from the glass into hematite, whereas a high fraction of the glass remains amorphous. With longer heat-treatment times at higher temperature, as discussed in (Ahmadzadeh et al., 2017), the amorphous phase can reject more Fe, creating additional hematite.

For the NaFeSi$_2$O$_6$ composition, on the other hand, a portion of the glass transforms to the aegirine phase during heat treatment at 900°C for 24 h. Trace amounts of hematite are also formed. It is interesting to note that even though the glass extensively forms aegirine crystals after the heat-treatment, no noticeable crystallization peak is observed in its DTA curve, while the aegirine melting peak is observed. Most likely, some aegirine crystals have formed in the DTA sample during non-isothermal heating, though not enough to show up as an exothermic peak. This is likely because the formation of aegirine is kinetically controlled by the relatively high heating rate of 10 °C min$^{-1}$ in the thermal analysis, as also suggested by other studies (Plaisted et al., 2000; Vienna et al., 1996).

The Mössbauer spectrum of the crystallized aegirine (Figure 2-d) confirms the presence of $^{[6]}$Fe$^{3+}$ in the crystal along with some $^{[4]}$Fe$^{3+}$ of the glass. The XAS pre-edge centroid of the
crystallized aegirine (NaFeSi$_2$O$_6$-900°C-24h) shown in Figure 3-c exhibits a shift to higher energies (+0.2 eV), combined with a shift to lower intensities, compared to those of the glasses. Comparing to the Mössbauer data (Figure 2), this observed pre-edge shift agrees with the conversion to ~100% Fe$^{3+}$ upon crystallization, considerable amounts of $[^6]$Fe$^{3+}$ in the aegirine crystal, and some remaining $[^4]$Fe$^{3+}$ in the residual glass. This change can be seen qualitatively in comparing the whole XANES spectra (Figure 3-a) of the natural aegirine standard, the NaFeSi$_2$O$_6$-900C-24h (crystallized) and the NaFeSi$_2$O$_6$-AQ (glass), with the intensity of the pre-edge features increasing visibly in this series, respectively.

In the case of NaFeSi$_3$O$_8$, due to high SiO$_2$ content and hence strong glass forming ability, the glass does not crystallize during heat-treatment at 1000°C for 24 h, even though some Fe leaves the glass as a small fraction of hematite. This composition can crystallize aegirine in addition to hematite upon prolonged isothermal heat-treatment times at lower temperatures (i.e., 900°C for 72 h).

It was previously shown that NaFeSi$_2$O$_6$ and NaFeSi$_3$O$_8$ glasses do not form the respective silicate crystals upon slow cooling of the melts (5 °C min$^{-1}$ from 1500 to 600°C followed by rapid quench) (Marcial and McCloy, 2019). However, at least for aegirine, the phase has been observed to crystallize heterogeneously (surface crystallization) in non-isochemical compositions following a long induction for nucleation, confirming that formation of aegirine is controlled kinetically (Plaisted et al., 2000; Vienna et al., 1996). In the case of Fe-albite, the conditions for nucleation of the crystalline phase, if it exists, may mimic that of albite, where the nucleation time is extremely long and the temperature of maximum nucleation is well below $T_g$ in a region of high viscosity, making albite glass almost impossible to crystallize in a water-free, seed-free experiment (Zanotto and Cassar, 2017).
**Related mineral systems**

The $\text{Al}^{3+}$ equivalents of the compositions studied here, i.e., $\text{NaAlSiO}_4$ (nepheline or carnegieite), $\text{NaAlSi}_2\text{O}_6$ (jadeite), and $\text{NaAlSi}_3\text{O}_8$ (albite) can form distinct crystalline phases and are well-studied rock-forming minerals (Deer et al., 2004; Marcial et al., 2016; Marcial and McCloy, 2019; Palmer, 1994). $\text{Al}^{3+}$, for which we substituted $\text{Fe}^{3+}$, is tetrahedrally coordinated by oxygen atoms in nepheline and albite structures, while octahedrally coordinated in jadeite. Substitution of all $\text{Al}^{3+}$ by $\text{Fe}^{3+}$ did not form the same crystal structure for nepheline and albite, whereas $\text{NaFe}^{3+}$Si$_2$O$_6$ forms isomorphous crystals to $\text{NaAl}^{3+}$Si$_2$O$_6$. Nonetheless, it was shown in (Ahmadzadeh et al., 2017) that the nepheline structure can incorporate significant amounts of $\text{Fe}^{3+}$ (up to $x=0.37$ in $\text{NaAl}_{(1-x)}\text{Fe}_x\text{SiO}_4$) in $\text{Al}^{3+}$ sites. Moreover, Bailey and Schairer (Bailey and Schairer, 1963; Bailey and Schairer, 1966) have observed that albite can also accommodate a limited amount of iron in its structure, likely in Al sites, though most Fe found in feldspars is in impurities of magnetite and hematite (Montiel-Anaya and Franco, 2019).

Comparison with the other alkali oxide-$\text{Fe}_2\text{O}_3$-SiO$_2$ systems is warranted. In the $\text{K}_2\text{O}$-$\text{Fe}_2\text{O}_3$-SiO$_2$ system, the K analogues of all these three compounds (i.e. KFeSiO$_4$, KFeSi$_2$O$_6$, and KFeSi$_3$O$_8$) exist as crystalline phases (Faust, 1936; Lange et al., 1986). In fact, not only iron-leucite (KFeSi$_2$O$_6$) (Bell and Henderson, 1994), but also iron-kalsilite (KFeSiO$_4$) (Bentzen, 1983) and iron-orthoclase/ferri-sanidine (KFeSi$_3$O$_8$) (Lebedeva et al., 2003; Shchipalkina et al., 2019) are isomorphous to their Al equivalents, and $\text{Fe}^{3+}$ is tetrahedrally coordinated in all three compounds. The Li equivalent of aegirine (LiFeSi$_2$O$_6$), known as Li-aegirine, crystallizes in the same crystal structure as spodumene (LiAlSi$_2$O$_6$) and aegirine (NaFeSi$_2$O$_6$), i.e., monoclinic structure with space group $C2/c$, with $[^6]\text{Fe}^{3+}$ (Lottermoser et al., 1998; Zhang et al., 2002). However, this phase is not reported as a natural mineral. LiFeSiO$_4$ has been reported, but is not
known as a natural mineral (Honma et al., 2012; Nytén et al., 2006). To our knowledge, LiFeSi₃O₈ has only been calculated but not experimentally reported (Persson, 2014).

It is notable that the coordination of Fe³⁺ has been shown to be particularly strongly influenced by the nature of the alkali cation in some silicate glasses, wherein [CN]Fe³⁺ decreases (from 6 (Li) to 4 (K)) with increasing alkali ionic radius (Bingham et al., 2014). This is qualitatively consistent with the coordination behavior of Fe³⁺ in the crystalline systems described above. Even with the same coordination number, the ionic size may still influence the stability of a given crystal structure, such as for alkali-Fe³⁺-feldspar, where the distortion of the Na environment due to its smaller ionic size is less favorable for the feldspar structure than K (Shchipalkina et al., 2019).

**IMPLICATIONS**

Three sodium iron silicate compositions (NaFeSiO₄ – Fe-nepheline, NaFeSi₂O₆ – aegirine, NaFeSi₃O₈ – Fe-albite) have been studied. The high-SiO₂ compositions (aegirine and Fe-albite with SiO₂ > 50 wt%) can readily form glass upon quenching, whereas Fe-nepheline crystallizes magnetite even at relatively fast quench rates because of its lower Si/Fe ratio. In the glasses, although there is some Fe²⁺ (<20%, depending on the composition), most of the iron occurs as 4-coordinated Fe³⁺ cations. However, when Fe crystallizes as the sodium iron silicate mineral (i.e., aegirine) upon isothermal heat-treatment, it arranges as Fe³⁺ octahedra (6-coordinated). It was found that analysis of pre-edge XANES features is less sensitive than Mössbauer for the study of Fe redox and coordination in such compositions where Fe³⁺/∑Fe does not vary significantly. The as-quenched glasses show similar Raman spectra with intense low-frequency (200-600 cm⁻¹) and high-frequency (800-1200 cm⁻¹) envelopes, and systematic
changes of their frequency and relative intensity based on their changing Si/Fe ratio. Moreover, the thermal analysis of these glass reveals an increase in the glass transition temperature from $T_g=471^\circ$C in NaFeSiO$_4$ (Si/Fe=1) to $T_g\sim521^\circ$C in NaFeSi$_3$O$_8$ (Si/Fe=3), due to the stronger Si-O bonds substituting for Fe-O bonds.

The study of these simplified compositions can help understand the potential phases that may crystallize within high-Fe nuclear waste glasses, which can affect their aqueous alteration behavior (Deshkar et al., 2019). It is shown that, in NaFeSi$_{x}$O$_{(2x+2)}$ type compositions, Fe crystallizes as iron spinel (i.e., magnetite) upon quench when $x=1$ (even at high quench rates); the melt can readily form a glass when $x\geq2$; and the glass can hardly crystallize when $x\geq3$. Increasing the Si/Fe ratio was found to increase the Fe$^{2+}$ fraction, consistent with known models, which has a network modifying role. The correlation between Si/Fe ratio and Fe oxidation state in such glasses, is important for geosciences and high-Fe nuclear waste management, as it can influence the melt polymerization, and as shown here, its crystallization behavior. Details of Fe chemistry are also important for nuclear waste vitrification processes, as the redox strongly influences foaming behavior and melt rate (Pokorny and Hrma, 2014). For a full understanding, future studies should be conducted with deliberate control of the Fe oxidation state of the starting glass, for these same compositions, in order to obtain a more complete understanding of the role of Fe conformation and Fe content on network topology and resulting thermal properties and viscosity.

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FIGURE CAPTIONS

Figure 1. Room temperature curves of magnetization – M, versus magnetic field – H, for a) as-quenched Fe-nepheline, b) as-quenched aegirine and Fe-albite, and c) heat-treated aegirine samples with maximum applied field of 1.8 T. The insets show the curves with maximum applied field of 0.2 T.

Figure 2. Fitted room temperature $^{57}\text{Fe}$ Mössbauer spectra of as-quenched (AQ) Fe-nepheline (a), Aegirine (b), and Fe-albite (c) samples, along with the crystallized aegirine heat-treated at 900°C for 24 h (d). The black points represent experimental data, blue line is total fit, and magenta lines are fits of individual Fe sites. See the online version of this article for color.

Figure 3. Fe K-edge X-ray absorption data. a) Measured XANES of glasses, crystallized glass, and standards, b) fits of the extracted pre-edge, and c) pre-edge integrated intensity versus centroid position (with respect to Fe foil at 7112 eV). Our measured samples are shown in red, whereas the measured standards are in blue. The same standards reported by Wilke et al (Wilke et al., 2001) are shown in filled black circles with their other reported values in open circles.

Figure 4. Raw (a) and treated (b) Raman spectra of NaFeSiO$_4$ (Fe-nepheline), NaFeSi$_2$O$_6$ (aegirine), and NaFeSi$_3$O$_8$ (Fe-albite) as-quenched (AQ) samples. The curves are offset along Y axis. The data shown in (b) is Long-corrected (Long, 1977), background-subtracted, and area-normalized.
Table 1. Studied sodium iron silicate samples

<table>
<thead>
<tr>
<th>Sample</th>
<th>Formula</th>
<th>wt% oxides</th>
<th>Melting temp (°C)</th>
<th>Isothermal heat treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe-nepheline</td>
<td>NaFeSiO₄</td>
<td>18.13 46.71 35.15</td>
<td>1500</td>
<td>775 7</td>
</tr>
<tr>
<td>Aegirine</td>
<td>NaFeSi₂O₆</td>
<td>13.42 34.56 52.02</td>
<td>1500</td>
<td>900 24</td>
</tr>
<tr>
<td>Fe-albite</td>
<td>NaFeSi₃O₈</td>
<td>10.65 27.43 61.92</td>
<td>1500</td>
<td>1000° 24</td>
</tr>
</tbody>
</table>

*The Fe-albite glass was also heat-treated for longer time (i.e., 72 h) at 900°C to try to crystallize the glass.

Table 2. Phase analysis of the as-quenched and heat-treated Fe-nepheline, aegirine, and Fe-albite samples, based on XRD and Rietveld refinement results

<table>
<thead>
<tr>
<th>Composition</th>
<th>Phase Analysis</th>
<th>As-quenched (AQ)</th>
<th>Heat-treated (HT)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaFeSiO₄</td>
<td>Phases (wt%)</td>
<td>Due to low silica content (Si/Fe=1), glass cannot accommodate all Fe easily, and excessive Fe crystallizes as Mgn.</td>
<td>79.2% Amrph + 20.8% Hmt</td>
</tr>
<tr>
<td>NaFeSi₂O₆</td>
<td>Amrph</td>
<td>Due to higher silica content (Si/Fe=2), glass accommodates all Fe.</td>
<td>7.4% Amrph + 91.4% Ae + 1.2% Hmt</td>
</tr>
<tr>
<td>NaFeSi₃O₈</td>
<td>Amrph</td>
<td>Due to high silica content (Si/Fe=3), glass accommodates all Fe.</td>
<td>95.5% Amrph + 4.5% Hmt (HT for 72 h at 900°C crystallizes 19% Ae too)</td>
</tr>
</tbody>
</table>

Amrph: amorphous; Mgn: magnetite; Hmt: hematite; Ae: aegirine.