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Iron speciation in minerals and glasses probed by M$_{2/3}$-edge x-ray Raman scattering spectroscopy

A. Nyrow · C. Sternemann · M. Wilke · R. A. Gordon · K. Mende · H. Yaş · L. Simonelli · N. Hiraoka · Ch. J. Sahle · S. Huotari · G. B. Andreozzi · A. B. Woodland · M. Tolan · J. S. Tse

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Abstract We present a spectroscopic study of the iron M$_{2/3}$-edge for several minerals and compounds to reveal information about the oxidation state and the local coordination of iron. We describe a novel approach to probe the iron M$_{2/3}$-edge bulk sensitively using x-ray Raman scattering. Significant changes in the onset and shape of the Fe M$_{2/3}$-edge were observed on ferrous and ferric model compounds with Fe in octahedral and tetrahedral coordination. Simulation of the spectra is possible using an atomic multiplet code, which potentially allows determination of e.g. crystal field parameters in a quantitative manner. A protocol is discussed that allows for determination of the Fe oxidation state in compounds by linear combination of spectra of ferric and ferrous end-members. The presented results demonstrate the capabilities of Fe M$_{2/3}$-edge spectroscopy by x-ray Raman scattering to extract information on the ratio of trivalent to total iron Fe$^{3+}$/∑Fe and local coordination. As x-ray Raman scattering is performed with hard x-rays, this approach is suitable for in-situ experiments at high pressure and temperature. It thus may provide indispensable information on oxidation state, electronic structure and local structure of materials that are important for physical and chemical processes of the deep Earth.

Keywords Iron speciation · minerals · x-ray scattering · x-ray absorption spectroscopy

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1 Introduction

Iron is the most abundant transition metal in the bulk Earth. It strongly influences the chemical and physical properties of iron-bearing minerals, glasses, and melts, which in turn affect many geological processes [Badro et al. (2003), Duffy (2008), Ono et al. (2007)]. At conditions of the deep Earth, the Fe$^{3+}$/∑Fe and the Fe coordination may vary dramatically, which is relevant for e.g. iron partitioning between Fe-Mg phases or behavior of iron during partial melting [Irifune and Isshiki (1998),...
In this paper we use non-resonant inelastic x-ray scattering, i.e. x-ray Raman scattering (XRS), in order to measure the iron $M_{\text{2,3}}$-edge, in a way similar to SXAS and EELS, but with high bulk sensitivity, which potentially allows for in-situ experiments at extreme conditions. The suitability of XRS for high pressure experiments has already been shown for various elements, e.g. carbon [Mao et al. (2003)], boron [Lee et al. (2005)], oxygen [Lee et al. (2008), Sahle et al. (2013)], silicon [Sternemann et al. (2013), Tse et al. (2013)], barium [Tse et al. (2011)], or vanadium [Ding et al. (2014)]. We study synthetic samples of FeO, Fe$_2$SiO$_4$, FeAl$_2$O$_4$, Fe$_2$O$_3$, and FePO$_4$, which serve as key examples for iron-bearing minerals and compounds with different iron oxidation state and coordination. They are used as a framework for evaluating the information that can be obtained by this technique for Fe-bearing mineral phases that are important throughout the deep Earth, ranging from Fe-Ni alloys of the Earth’s core [Antonangeli et al. (2010), Lin et al. (2010), Terasaki et al. (2011)] to Fe-Mg silicates in mantle [Fang and Ahuja (2008), Hofmeister (2006), Otsuka et al. (2010)] and crust, as well as extraterrestrial materials [Edwards et al. (2000), Xu and Lin (2000)].

First we give a short discussion on the theory of XRS spectroscopy followed by an overview on samples and experimental details. Then we demonstrate how XRS can be applied to measure dipole and multipole excitations and how the results can be compared to theory. Finally, we discuss in detail the sensitivity of XRS measurements to determine iron coordination and oxidation state and we use XRS to estimate the Fe$^{3+}/\sum$Fe ratio of glasses synthesized at different redox conditions.

2 Theory of x-ray Raman scattering

The inelastic x-ray scattering process occurs when an incoming x-ray photon is scattered by the electrons of the sample system, leading to a partial transfer of its energy and momentum. The energy and momentum transfer result in different excitations, such as valence band excitations, Compton scattering or phonon excitations [Bergmann et al. (2004), Hämäläinen and Manninen (2001), Krisch and Sette (2002), Schiöll (2007)]. Electronic excitations from a core state to a non-occupied state can occur if the transferred energy is in the vicinity of electron binding energies. Measuring those excitations allows for studies of absorption edges in both the dipole and the non-dipole limit. The technique that
exploits such phenomena is called non-resonant inelastic x-ray scattering or x-ray Raman scattering (XRS).

Typically, the scattered photons are measured at fixed outgoing photon energy (6 - 20 keV) as the energy of the incident photons is scanned at energy-loss values in the vicinity of a certain absorption edge (e.g. 53 eV, 543 eV and 1071 eV for the iron M-, oxygen K- and sodium K-edge, respectively). In contrast to EELS or SXAS, XRS is highly bulk sensitive because the energy of the incoming and scattered photons is in the hard x-ray regime. The measured quantity in such an experiment is proportional to the dynamic structure factor, which is given by [Schülke (2007)]

\[
S(q, \omega) \propto \sum_f |\langle f| \sum_j e^{-i\mathbf{r}_j \cdot \mathbf{r}} | c \rangle|^2 \delta(E_c - E_f + \hbar \omega).
\]

For XRS the dynamic structure factor is proportional to the excitation probability from an initial core state |c\rangle with energy \(E_c\) to all non-occupied final states |f\rangle with energy \(E_f\). The momentum transfer vector is \(q\) and the sum is taken over the electrons of the system with position \(\mathbf{r}_j\). The delta-function ensures energy conservation with the energy transfer \(\hbar \omega\).

The dynamic structure factor contains element selective information about the local chemical and electronic structure as can be seen in expressed in terms of the transition matrix \(M_L(q, E)\) and the partial non-occupied electronic density of states \(\rho_L(E)\) with the angular momentum channel \(L\): [Soininen et al. (2006)]

\[
S(q, \omega) = \sum_L |M_L(q, E)|^2 \rho_L(E).
\]

According to equation (2) the weight of the matrix elements for different excitation channels (dipole and non-dipole) strongly changes with the magnitude of \(q\) giving rise to dominating dipole contribution (e.g. 3p to 3d) for \(q < \ll\) providing a signal similar to that obtained by SXAS and EELS. At higher \(q\) increasing contributions of non-dipole excitations are observed. In an experiment \(q\) can be easily tuned by the scattering angle \(2\theta\) due to \(q = \frac{2\pi}{\lambda} \sin(2\theta/2)\). Measurements performed at low and high \(q\) reveal a more complete picture of the electronic density of states [Soininen et al. (2006), Bradley et al. (2011)]. The special properties of XRS have made it an indispensable tool to accomplish experimental challenges if soft x-rays and electrons cannot be applied as a probe, see e.g. [Gordon et al. (2008), Inkinen et al. (2013), Lee et al. (2005), Lee et al. (2008), Mao et al. (2003), Mattila et al. (2005), Sahle et al. (2013), Soininen et al. (2005), Sternemann et al. (2005), Sternemann et al. (2013), Tse et al. (2011), Tse et al. (2013)].

In the case of transition metals atomic excitations dominate the XRS M- and L-edge spectra [Gordon et al. (2009)]. In fact, the 3p hole and the 3d hole wave functions overlap significantly and the final states are found by the vector coupling of these two wave functions. This so-called multiplet effect is well known in atomic physics and can be observed also in solids. Here, it has to be used rather than the density-of-states approach discussed above. We express the dynamic structure factor via

\[
S(q, \omega) = \sum_f \sum_{k=0}^{\infty} D_k |\langle f| j_k(qr) | c \rangle|^2 \delta(E_c - E_f + \hbar \omega) \times \delta(E_c - E_f + \hbar \omega)
\]

with the spherical Bessel function \(j_k(qr)\) of \(k\)-th order, which gives rise to the \(q\)-dependence, the radial wave functions \(f(r)\) and \(c(r)\), and coefficients \(D_k\) [Gordon et al. (2009)]. Since the initial state wave function (3p) is odd and the final d wave function (3d) even in presence of inversion symmetry, only transitions of odd parity and \(d - p = 1 \leq k \leq p + d = 3\) contribute in equation (3). Thus, only terms with \(k = 1\) (dipole transitions) and \(k = 3\) (octopole transitions) contribute to the XRS spectrum of the Fe M_{2/3}-edge [Haverkort et al. (2007)]. The relative weight of these two transitions is governed by the magnitude of the momentum transfer \(q\). This approach will be used later to discuss the Fe M_{2/3}-edge spectra.

3 Sample description and experimental details

3.1 Samples

The sensitivity of the iron M_{2/3}-edge’s shape regarding the iron oxidation state and local coordination was probed for synthetic crystalline compounds with Fe^{3+} and Fe^{2+} in octahedral and tetrahedral configuration. The momentum-transfer dependence of XRS spectra was investigated using FeO (Fm3m) and α-Fe_{2}O_{3} (I\overline{R}3c, corresp. to hematite), whereas high-resolution XRS measurements were performed for synthetic FeAl_{2}O_{4} (Fd\overline{3}m, corresp. to hercynite [Andreozzi and Lucchesi (2002)]), Fe_{2}SiO_{4} (P\overline{b}n\overline{m}, corresp. to faujalite), FePO_{4} (P3\overline{2}12, iron(III) phosphate with berthilite-structure (AIPO_{4}), corresp. to rodolite) and α-Fe_{2}O_{3} (I\overline{R}3c). FeO and Fe_{2}SiO_{4} have Fe^{2+} in octahedral coordination, whereas FeAl_{2}O_{4} contains Fe^{3+} in tetrahedral coordination. α-Fe_{2}O_{3} and iron-(III)-phosphate have Fe^{3+} in octahedral and tetrahedral coordination, respectively. FeO and α-Fe_{2}O_{3} powders with 99.9% and 99.995% trace metals basis, respectively, were purchased from Sigma Aldrich.
The FeAl2O4 sample was synthesized and characterized by [Andreozzi and Lucchesi (2002)]. The chemical analysis by electron microprobe (EMP) shows 44.7±0.2 wt.% FeO, 55.5±0.4 wt.% Al2O3. Stoichiometric calculation from the EMP analysis and Mössbauer spectroscopy revealed an Fe2+/∑Fe ratio of 0.92 and 0.94, respectively, i.e. a small contribution of Fe3+. In FeAl2O4 85% Fe3+ occurs in tetrahedral coordination whereas 15% Fe2+ is octahedrally coordinated. FePO4 and Fe2SiO4 were analyzed by Fe K-edge XANES using the method of [Wilke et al. (2001)]. FePO4 (pre-edge centroid position at 7113.55±0.01 eV) did not show any contribution by Fe2+ and Fe3SiO4 (pre-edge centroid position at 7111.96±0.02 eV) no Fe3+. All used samples were characterized in addition by x-ray diffraction measurements at beamline BL9 [Krywka et al. (2007)] of the synchrotron radiation source DELTA (Dortmund, Germany) to verify sample structure and to exclude impurities. For XRS measurements, the powdered samples were pressed into pellets.

In addition to crystalline compounds, we measured three synthetic glasses to test potential for determining the Fe3+/∑Fe ratio. Two glasses were used with a composition of a transitional alkalic basalt from Iceland [Thy and Lofgren (1994)], RB0-1 and RB0-4, that were synthesized at 1350°C at two different oxygen fugacities and characterized by EMP, wet chemical analysis and Mössbauer spectroscopy, c.f. [Wilke et al. (2005)]. They have the following starting composition: 14.38 ± 0.21 wt% FeO, 47.88 ± 0.20 wt% SiO2, 13.22 ± 0.16 wt% Al2O3, 9.04 ± 0.08 wt% CaO, 4.68 ± 0.08 wt% Na2O, 4.04 ± 0.10 wt% TiO2 and 3.23 ± 0.06 wt% MgO. Their Fe3+/∑Fe were determined by wet chemistry analysis and Mössbauer spectroscopy to 0.83±0.04 (RB0-1) and 0.16±0.04 (RB0-4). In addition, we studied an Fe-doped haplogranitic glass synthesized in air at 1600°C (AOQ-2, 8.4 ± 0.2 wt% Fe2O3, 72.9 ± 0.5 wt% SiO2, 10.8 ± 0.2 wt% Al2O3, 4.5 ± 0.1 wt% K2O and 3.4 ± 0.2 wt% Na2O). A Fe3+/∑Fe of 0.63±0.04 was determined by Mössbauer spectroscopy (for experimental details see e.g. [Woodland and Jugo (2012)]) for low q, spectra obtained by 2 analyzers with the same momentum transfer could be summed up. At high q, spectra of all analyzers could be summed, which significantly enhances the statistical accuracy. This way of data processing was applied only for the experiments using the ID16 and P01 setup, whereas for data collected using the LERIX spectrometer each analyzer crystal was treated separately to determine the q-dependence of the Fe3+ signal.

3.2 Experiments

XRS spectra were acquired at 4 different experimental stations. In order to study the q-dependence of the Fe3+ signal, FeO (Fe2+) and α-Fe2O3 (Fe3+), both octahedrally coordinated, were measured at beamline PNC/XSD 20-ID of the Advanced Photon Source (APS) employing the LERIX spectrometer [Fister et al. (2006)]. Here, 19 Si analyzer crystals are arranged on a semicircle with a radius of 1 m for vertical scattering geometry covering the scattering angles from 9° to 171°, which corresponds to q between 0.8 Å−1 to 10.0 Å−1. The Si(555) reflection was used with an analyzer energy of 9.89 keV. The incident energy was monochromatized by a Si(111)-monochromator and an overall energy resolution of 1.5 eV was achieved. The M2,3-edges were measured for energy-loss values between 45 eV to 70 eV by variation of the incident energy. At beamline ID16 of the European Synchrotron Radiation Facility (ESRF), measurements of the Fe M2,3-edges of FeO, Fe2SiO4, FePO4, and FeAl2O4 were performed using the multiple element spectrometer for non-resonant inelastic x-ray spectroscopy of electronic excitations [Verbeni et al. (2009)], both for low and high q. The analyzer energy was set to 9.69 keV using the Si(660)-reflection and an overall energy resolution of 0.8 eV was obtained employing the Si(111) pre-monochromator together with a Si(220) channel-cut monochromator [Honkanen et al. (2013)]. Here, XRS spectra were collected utilizing a 9 analyzer array positioned at average scattering angles of 30° and 130° corresponding to q of 2.05 Å−1 and 9.1 Å−1, respectively. The iron M2,3-edges of iron bearing glasses were measured using the inelastic x-ray scattering spectrometer of beamline P01 at PETRA III (DESY), which is similar to that reported in [Verbeni et al. (2009)] but with vertical scattering plane and 5 Si(660) analyzer crystals. A total energy resolution of 0.6 eV was achieved utilizing a Si(311) monochromator. Here, the average scattering angle was 135° corresponding to a momentum transfer of 9.1 Å−1. Preliminary studies of the M2,3- and L-edges of FeO, Fe2SiO4 and α-Fe2O3 using XRS were conducted at beamline BL12XU of SPring8 [Cai et al. (2004)] to prove the feasibility of an XRS M2,3-edge study for the analysis of iron bearing minerals.

In all experiments several subsequent spectra were measured and summed up for each analyzer. The spectra were background corrected and finally normalized to the integral intensity between 49 eV and 68 eV energy loss. To increase the statistical accuracy, XRS spectra measured for similar q were added if the q-dependence was found to be weak, which will be discussed in more detail in the next section. For low q, spectra obtained by 2 analyzers with the same momentum transfer could be summed up. At high q, spectra of all analyzers could be summed, which significantly enhances the statistical accuracy. This way of data processing was applied only for the experiments using the ID16 and P01 setup, whereas for data collected using the LERIX spectrometer each analyzer crystal was treated separately to determine the
The procedures applied to separate the Fe M\textsubscript{2/3}-edge from the particle-hole excitation spectrum and Compton background for low and high \( q \), respectively, is similar to those reported by [Sternemann et al. (2008)]. For low \( q \), the Fe M\textsubscript{2/3}-edge can be found on the tail of the particle-hole excitation spectrum with its maximum at lower energy loss than the edge. Here, a Pearson function or likewise a Gaussian or a Lorentzian function can be used to approximate the underlying background (see figure 1 a)). With increasing \( q \) the particle-hole excitation spectrum transforms to the Compton spectrum. The maximum of the Compton peak moves to higher energy loss. This makes the subtraction of the Compton background difficult if the Compton maximum appears in the vicinity of the Fe M\textsubscript{2/3}-edge. Here, one can use either a Gaussian to approximate the Compton peak shape [Fister et al. (2009)] or the extraction procedure described in [Sternemann et al. (2008)]. For high \( q \) the Compton peak is located at much higher energy loss than the Fe M\textsubscript{2/3}-edge, so that it lies on a smoothly increasing background, which can be modeled as discussed for low \( q \) or, alternatively, by a linear fit (see figure 1 b)). As indicated in figure 1, different modeling of background does not significantly influence the shape of the Fe M\textsubscript{2/3}-edge as long as energy-loss values up to 15 eV above the edge onset are considered. For the data of the LERIX spectrometer, we used a Gaussian for background subtraction throughout the whole \( q \)-range, whereas for ESRF data a Lorentzian and a linear background was used for low and high \( q \), respectively.

4 Momentum transfer dependence of the Fe M\textsubscript{2/3}-edge

The \( q \)-dependence of XRS spectra of the M\textsubscript{2/3}-edge of octahedrally coordinated Fe in FeO (Fe\textsuperscript{2+}) and \( \alpha \)-Fe\textsubscript{2}O\textsubscript{3} (Fe\textsuperscript{3+}) is presented in figure 2 a)-c) for a large \( q \)-range between 0.8 to 10.0 \( \text{Å}^{-1} \). For \( q < 2 \text{ Å}^{-1} \) in the dipole limit (\( k=1 \), see equation 3), clear spectral differences between Fe\textsuperscript{2+} and Fe\textsuperscript{3+} are observed in the energy-loss region from 54 eV to 60 eV. Here, the spectra of Fe\textsubscript{2}O\textsubscript{3} show only one broad maximum at 58 eV, whereas the spectra of FeO show an additional shoulder at 55 eV. Both spectra exhibit a relatively high energy step for energy-loss values above 63 eV due to excitations into continuum states. With increasing \( q \) the spectral shape changes significantly due to the increasing contribution of octupole (\( k=3 \)) transitions to the excitation spectrum and only small contributions by continuum excitations. This is most prominently manifested by an intensity increase for energy-loss values below 55 eV. Finally, the spectral \( q \)-dependent changes are found to be weak for \( q > 8 \text{ Å}^{-1} \) and the spectra show an asymmetric shape with a maximum at 52 eV and a shoulder at higher energy loss in the case of Fe\textsuperscript{2+}. For Fe\textsuperscript{3+}, only a single maximum at 53.5 eV is observed.

The experimental results are compared with calculations employing an atomic multiplet code [de Groot (2005), de Groot (2008)] (see Fig. 2). The multiplet calculations were performed using the modified CTM4XAS package [Stavitski and de Groot (2010)] to obtain Fe absorption edges similar to the model used by van der Laan [van der Laan (1991)] for dipole and octupole transitions. Continuum excitations are not considered in the theoretical approach. To calculate the M\textsubscript{2/3}-edge, the values for the crystal-field splitting found in [de
Fig. 2 Upper panel: The effect of momentum transfer on the Fe M$\text{_{2/3}}$-edge for octahedrally coordinated Fe$^{2+}$ (FeO) and Fe$^{3+}$ ($\alpha$-Fe$_2$O$_3$) in the $q$ ranges 0.8-5.2 Å$^{-1}$ (a), 5.9-8.5 Å$^{-1}$ (b), and 8.9-10 Å$^{-1}$ (c). Lower panel: multiplet calculations for the dipole (d) and octupole (f) contribution as well as a 1:1 superposition of both to model the intermediate $q$-range (e).

Groot et al. (2005)] have been used. For a better agreement with the data, the Slater-Condon parameters were reduced to 64% of their atomic value and no spin-orbit splitting was applied. The calculated transition patterns were convoluted with a Lorentzian function (FWHM of 0.6 eV [Fuggle and Alvarado (1980)]) and with a Gaussian function (FWHM of 1.5 eV) to simulate the natural broadening of the states and the experimental resolution, respectively. In figure 2 d), calculations in the dipole limit (low $q$, $k=1$) are shown and are compared to results considering only octupole transitions ($k=3$, high $q$ limit), see figure 2 f). The modeled spectra resemble adequately both the spectral differences observed for the different oxidation states of iron and the $q$-dependence of the experiment although the weak maximum of the calculated Fe$^{2+}$ spectrum at high $q$ is not present in the measurement. Notably, for high $q$ the excitation spectra are dominated by octupole transitions with only small contribution by the continuum excitations. The spectrum of the dipole excitations appears broader compared to that of the octupole excitations. This may be explained by its higher energy position allowing its interaction with excitations into the continuum states [Sen Gupta et al. (2011), Sahle et al. (2014)]. The XRS measurements in the crossover regime, e.g. at $q = 5.89$ Å$^{-1}$, can be properly reproduced by a superposition of dipole and octupole multiplet calculations with a 1:1 ratio (figure 2 e)).

The observed $q$-dependence has two major implications for the study of iron speciation by XRS spectroscopy: i) Both, experiment and theory indicate very strong spectral changes due to iron oxidation state, which is observed particularly well at high momentum transfers; ii) The $q$-dependence of the spectra for $q > 8.11$ Å$^{-1}$ is negligible. Consequently, spectra measured with different analyzers at different $q$ in the high $q$-limit can be summed up, which significantly increases the statistical accuracy of the experimental results. Hence, measure-
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In this section we discuss how oxidation state (Fe$^{2+}$, Fe$^{3+}$) and/or coordination (octahedral, tetrahedral) of iron change the shape of the Fe M$_{2/3}$-edge. For this comparison, we will use the spectra obtained at ID16. The XRS spectra of the four reference compounds measured at low $q$ in the dipole limit are presented in figure 3 (left). The spectrum of $\alpha$-Fe$_2$O$_3$ (Fe$^{3+}$, octahedral) shows a strong excitation peak at 58.0 eV (B$^*$) and a weak pre-edge feature at 53.5 eV (A$^*$). These features are similar to those observed by Xiong et al. [Xiong et al. (2012)] using SXAS. Excitations into continuum states cause the high intensity in the post edge region. When compared to the XRS spectrum of FePO$_4$ (Fe$^{3+}$, tetrahedral), the pre-edge intensity A$^*$ is broader and less pronounced, whereas a sharper main peak (B$^*$) is observed. The edge onset for FePO$_4$ is slightly shifted to lower energy loss by 0.25 eV.

The XRS spectrum of Fe$_2$SiO$_4$ (Fe$^{2+}$, octahedral) is characterized by a two-peak structure with maxima at 54.5 eV and 57.3 eV (B$_1$ and B$_2$) together with a weak pre-edge feature at 52.8 eV (A). A comparison with the spectrum of FeAl$_2$O$_4$ (Fe$^{2+}$, 85% tetrahedral) yields a similar two peak spectral shape but the features B$_1$ and B$_2$ are broader and the edge onset is slightly shifted to higher energy by around 0.2 eV. Moreover, a less pronounced pre-peak intensity is observed in line with the findings for tetrahedral Fe$^{3+}$ in FePO$_4$ (berlinite structure). It must be recalled, that the FeAl$_2$O$_4$ sample contains 15% Fe$^{2+}$ in octahedral coordination [Andreozzi and Lucchesi (2002)], and this cation disorder might affects the spectral features slightly. Overall, small but

Fig. 3  High resolution measurements of the Fe M$_{2/3}$-edge on Fe$^{2+}$ and Fe$^{3+}$ (both octahedral and tetrahedral) for q = 2.05 Å$^{-1}$ (left) and q = 9.1 Å$^{-1}$ (right), respectively. See text for detailed discussion.
distinct differences due to local Fe coordination are
found for both oxidation states.
In contrast, the difference of the spectra for the two oxida-
tion states is striking (see also EELS measurements
by van Aken [van Aken et al. (1999)] and calculations
by van der Laan [van der Laan (1991)]). There is a large
energy shift of 2.5 eV between the edge onsets of Fe$^{2+}$
and Fe$^{3+}$ with a remarkable difference in line shape,
i.e. a double-peak structure of the Fe$^{3+}$ main edge in
contrast to a single peaked M$_{2/3}$-edge for Fe$^{3+}$. These
strong differences enable a separation between different
Fe oxidation states and should allow for precise deter-
mination of the Fe$^{3+}/\sum$Fe.
At high $q$, the line shape and energy onset of the Fe
M$_{2/3}$-edge changes completely. The XRS spectra of Fe$_2$O$_3$
and FePO$_4$ show a strong maximum (C') dominating
the M$_{2/3}$-edge spectra at the energy-loss position,
where the pre-peak is observed for low $q$. This differ-
ence is due to the dominating contribution of octupole
transitions at high $q$. In contrast to low $q$, there is a
slight energy shift of 0.25 eV for the edge onset to
lower energy-loss values for the octahedral compared
to the tetrahedral Fe$^{3+}$. The former also shows a less
pronounced feature D', which is most probably due to
small but still visible contributions from dipole excita-
tions.
Concerning the Fe$^{2+}$ compounds at high $q$, the spectral
shape of FeAl$_2$O$_4$ is very similar to that of Fe$_2$SiO$_4$. The
typical double-peak structure is conserved but shifted
as for Fe$^{3+}$ to lower energy loss and exhibits in con-
trast to low $q$ a more intense first peak at 52.3 eV (C$_1$)
and a less pronounced second peak at 54.0 eV (C$_2$).
We find the high-energy-loss feature (D) to be broader
and shifted to lower energy loss compared to Fe$^{3+}$ in
accordance to the broader line shape and energy shift
of Fe$^{3+}$ at low $q$.
Between tetrahedrally and octahedrally coordinated Fe$^{2+}$
at high $q$, only subtle differences are found, i.e. a shift
of the C$_2$ peak position to higher energy loss for octahe-
dral coordination and a small variation in the intensity
of peaks C$_1$ and C$_2$. This difference in intensity might
be due to small contribution of Fe$^{3+}$ and to a less ex-
tent due to octahedral Fe$^{2+}$ contained in the FeAl$_2$O$_4$
sample as discussed below. Apparently the spectral dif-
fences with respect to the different oxidation states
are at least as significant as observed for low $q$.
Our observations imply that the M$_{2/3}$-edge spectra en-
able to distinguish between tetrahedral and octahedral
coordination, particularly with spectra obtained at low
$q$. These relatively small differences between octahe-
drally and tetrahedrally coordinated Fe demand mea-
surements of the M$_{2/3}$-edge with high statistical accu-
curacy, particularly if the changes in pre-edge intensity are
to be employed. We would like to note, however, that
the use of pre-peak shape and intensity in the analy-
sis of local coordination might be not straightforward.
E.g. Xiong et al. [Xiong et al. (2012)] found significant
changes in the pre-peak for octahedrally coordinated
Fe$^{3+}$ in FePO$_4$.2H$_2$O compared to that in $\alpha$-Fe$_2$O$_3$.
These differences are related to the distortion of the
octahedra in FePO$_4$.2H$_2$O with respect to the regular
ones in Fe$_2$O$_3$.
To use these XRS spectra as references for character-
izing Fe$^{2+}/\sum$Fe of unknown samples requires references
with pure Fe$^{2+}$ and Fe$^{3+}$ oxidation states. As the FeAl$_2$O$_4$
 sample contains about 6% Fe$^{3+}$ at octahedral sites [An-
dreozzi and Lucchesi (2002)], it was corrected for the
Fe$^{3+}$ contribution by subtracting the XRS spectrum of
Fe$_2$O$_3$ weighted by 0.06. After subtraction, the resulting
spectrum was normalized according to the procedure
discussed above. The correction scheme is presented in
figure 4 for the high $q$ measurement. The corrected XRS
spectrum conserves the peak shift of structure C$_2$ but
the intensity ratio of peaks C$_1$ and C$_2$ changes slightly,
being now closer to the ratio found for Fe$_2$SiO$_4$. A cor-
rection of the FeAl$_2$O$_4$ spectrum for 15 % octahedrally
coordinated Fe$^{2+}$ can be disregarded due to the similar
shape of tetrahedral and octahedral Fe$^{2+}$ XRS spectra.

![Figure 4](https://example.com/figure4.png)

Overall the spectra measured at high $q$ show a much
better statistical accuracy as we were able to sum up
spectra acquired with several analyzer crystals, which
is advantageous in the analysis of oxidation state and coordination of an unknown sample.

6 Determination of Fe\(^{3+}/\Sigma\)Fe

In this section we explore the possibility of using the Fe M\(_{2/3}\)-edge for determining the Fe oxidation state in minerals and compounds. In the light of the many techniques already available, we would like to point out that the M\(_{2/3}\)-edge measured by XRS may provide a substantially improved way of obtaining this information particularly in experiments performed in-situ at high pressure and temperature. Here, we aim at evaluating the technique for extracting the Fe\(^{3+}/\Sigma\)Fe in terms of procedure and precision. For doing so, we present measurements of the Fe M\(_{2/3}\)-edge on iron-containing glasses with Fe\(^{3+}/\Sigma\)Fe ratios ranging from about 0.1 to 0.9. We used spectra collected at high momentum transfer. The XRS spectra of the glasses were measured with slightly better energy resolution than the reference samples. To match the energy resolution of both experiments for the fitting procedure, the spectra of the glasses were artificially broadened by a convolution with a Gaussian of 0.6 eV FWHM.

The spectra of the glasses are presented in figure 5. The Fe M\(_{2/3}\)-edge of the RB0-4 glass (right) shows a shape similar to those of Fe\(_2\)SiO\(_4\) or FeAl\(_2\)O\(_4\). In contrast, the spectra of the AOQ-2 and RB0-1 glasses show shapes very similar to that of Fe\(^{3+}\)-oxide. It is noted that due to the strong multiplet excitations, contributions from other light elements contained in the samples do not influence the Fe M\(_{2/3}\)-edge spectrum significantly. This is evidenced by the Mg-bearing RB0-glasses for the Mg L-edge, which is located in the same energy range as the Fe M\(_{2/3}\)-edge.

For a quantitative analysis of the individual glasses, a superposition of Fe\(^{2+}\) and Fe\(^{3+}\) reference XRS spectra was fitted to the glass spectra. Here, three different sets of fits were performed. For the first fit, (I) spectra of octahedral Fe\(^{2+}\) and Fe\(^{3+}\) references (Fe\(_2\)SiO\(_4\) and Fe\(_2\)O\(_3\)) were used (blue line). For the second fit (II), FeAl\(_2\)O\(_4\) and FePO\(_4\) (both tetrahedral iron coordination) were used to investigate the influence of the coordination (red line). In silicate glasses, the Fe coordination shows site-to-site distribution with variable distortion or even mixed coordination. Hence, the use of spectra of crystalline samples with only one sort of coordination might not be appropriate. In order to analyze this, a weighted sum of all four reference spectra (Fe\(^{2+}\) and Fe\(^{3+}\), both octahedral and tetrahedral) was fitted to the glass spectra (III). For fitting procedures (II) and (III) we used the FeAl\(_2\)O\(_4\) XRS spectrum corrected for the Fe\(^{2+}\) contribution (c.f. figure 4).

The results are shown in figure 5. The general shape of the edge is reproduced well by all three ways of fitting with reference spectra. However, the fit revealed by superposition of all four references (green line) shows the best agreement, especially in the case of RB0-4 (5, right).

The quantitative results obtained by the fitting procedures can be found in table 1. The resulting Fe\(^{3+}/\Sigma\)Fe of the glasses extracted by a fit of Fe\(^{2+}\) and Fe\(^{3+}\) reference spectra for octahedral coordination tend to overestimate the Fe\(^{3+}/\Sigma\)Fe compared to the results revealed by Mössbauer spectroscopy. While the Fe\(^{3+}/\Sigma\)Fe of the RB0-glasses measured by XRS are close to the Mössbauer results, the deviation becomes very significant in the case of the AOQ-2 glass. Using FePO\(_4\) (tetrahedral coordination) as reference for Fe\(^{3+}\) significantly affects the results. Here, a better agreement is found for the AOQ-2 glass but the Fe\(^{3+}/\Sigma\)Fe of the RB0-glasses are underestimated. Thus, we conclude that the effect of coordination cannot be neglected in the fitting procedure although the spectra show only subtle differences as discussed before. The Fe\(^{3+}\) fraction of the RB0-1 glass seems to have predominantly octahedral coordination, while the tetrahedral coordination is preferred for the Fe\(^{3+}\) contribution in the AOQ-2 glass. To quantify this effect, the fitting procedure (III) was used with all four reference spectra to determine not only the Fe\(^{3+}/\Sigma\)Fe ratio but also the contribution of a certain coordination. This procedure reveals an improved agreement between the measured and fitted spectra of all glasses. Octahedral coordination is preferred for Fe\(^{2+}\) in all samples, while the fraction of tetrahedral Fe\(^{2+}\) can be neglected. In contrast, both octahedral and tetrahedral coordinations of Fe\(^{3+}\) contribute to the spectra for all samples, especially those with high Fe\(^{3+}/\Sigma\)Fe ratio. The RB0-1 glass seems to contain slightly more octahedral Fe\(^{3+}\) than tetrahedral Fe\(^{3+}\), while the AOQ-2 glass tends to have more tetrahedral Fe\(^{3+}\). Evidence for mixed coordination of Fe in silicate glasses has been already provided by various techniques (e.g. [Virgo and Mysen (1985), Drewitt et al. (2013), Farges et al. (2004), Wilke et al. (2005)]). In addition to mixtures of tetrahedral and octahedral coordination, also the presence of fivefold coordination in trigonal bipyramidal symmetry has been proposed [Rossano et al. (1999), Galois et al. (2001), Jackson et al. (2005)]. The difference observed here for the two compositions is consistent to the trends observed using the pre-edge of the Fe K-edge XANES [Farges et al. (2004), Wilke et al. (2005), Wilke et al. (2007)], that were interpreted as evidence that the coordination of Fe\(^{3+}\) is considerably influenced by the polymerization of the melt. Despite the similarity in the results, we would like to stress here that Fe M\(_{2/3}\)-
We showed that the q-dependence of the XRS spectra can be used to tune the sensitivity of the measurements. Because XRS is a bulk probe, Fe $M_{2,3}$-edge studies, which have been restricted to vacuum conditions so far, can now be conducted on samples that are incompatible with vacuum or even in experiments at high pressure and high temperature using e.g. a resistively or laser heated DAC. Thus, XRS provides an exciting tool to study the coordination, oxidation state and even the spin state [Nyrow et al., to be published] of iron in minerals, glasses and melts at conditions relevant for the Earth’s interior and gives access to unique complementary information on iron speciation at in-situ conditions for many geophysical and geochemical applications.

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### Table 1

<table>
<thead>
<tr>
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<th>RB0-1</th>
<th>AOQ-2</th>
<th>RB0-4</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Fe^{3+}/\sum Fe$ XRS (I)</td>
<td>$0.89\pm0.01$ ($R^2=0.9890$)</td>
<td>$0.73\pm0.04$ ($R^2=0.9780$)</td>
<td>$0.18\pm0.02$ ($R^2=0.9274$)</td>
</tr>
<tr>
<td>$Fe^{3+}/\sum Fe$ XRS (II)</td>
<td>$0.74\pm0.05$ ($R^2=0.9884$)</td>
<td>$0.61\pm0.04$ ($R^2=0.9880$)</td>
<td>$0.14\pm0.04$ ($R^2=0.8561$)</td>
</tr>
<tr>
<td>$Fe^{3+}/\sum Fe$ XRS (III)</td>
<td>$0.82\pm0.03$ ($R^2=0.9964$)</td>
<td>$0.69\pm0.05$ ($R^2=0.9911$)</td>
<td>$0.18\pm0.01$ ($R^2=0.9291$)</td>
</tr>
</tbody>
</table>

### 7 Conclusion

In this paper we demonstrated the capabilities of XRS spectroscopy to study the $M_{2,3}$-edge of iron in different iron bearing minerals and compounds and showed how XRS spectra can be used to reveal information about the oxidation state and the local coordination of iron. The determination of the $Fe^{3+}/\sum Fe$ can be performed with high sensitivity due to the strong spectral changes observed at the Fe $M_{2,3}$-edge. Our data provide strong evidence that estimation of iron coordination of $Fe^{2+}$ and $Fe^{3+}$ in glasses or even melt is feasible. We showed that the q-dependence of the XRS edges with improved statistics are required to conduct a reliable analysis of the average coordination number of iron in glasses. Still, this preliminary result is particularly exciting, as this method would not only allow determining the iron oxidation state at in-situ conditions but also detecting coordination changes as a function of pressure and/or temperature, hardly feasible by any other method.

### Fig. 5

Fe $M_{2,3}$-edge spectra of glasses with different $Fe^{3+}/\sum Fe$ shown together with the results of the fits employing different reference spectra (see text for details).

#### RB0–1 glass

- $Fe^{2+}$oct + $Fe^{3+}$oct (I)
- $Fe^{2+}$tetr + $Fe^{3+}$tetr (II)
- All references (III)

#### AOQ–2 glass

- $Fe^{2+}$oct + $Fe^{3+}$oct (I)
- $Fe^{2+}$tetr + $Fe^{3+}$tetr (II)
- All references (III)

#### RB0–4 glass

- $Fe^{2+}$oct + $Fe^{3+}$oct (I)
- $Fe^{2+}$tetr + $Fe^{3+}$tetr (II)
- All references (III)
References


[Drewitt et al. (2013)] Drewitt JE, Sanloup C, Bythchkov A, Brassamin S, Hennet L: Structure of (Fe,Ca)$_{1-x}$O$_{y}$(SiO$_2$)$_{1-y}$ liquids and glasses from high-energy x-ray diffraction: Implications for the structure of natural basaltic magmas. Physical Review B 87, 224201 (2013)


Iron speciation in minerals and glasses probed by M$_{2,3}$-edge x-ray Raman scattering


[Sobolev et al. (1999)] Sobolev VN, McCammon CA, Taylor LA, Snyder CA, Sobolev NV: Precise Moessbauer milliprobe determination of ferric iron in rock-forming minerals and limitations of electron microprobe analysis. Amer.


