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In situ Fe K-edge X-Ray Absorption Spectroscopy study during cycling of Li$_2$FeSiO$_4$ and Li$_{2.2}$Fe$_{0.9}$SiO$_4$

Li ion battery materials

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In situ X-Ray Absorption Spectroscopy (XAS) results are presented for Li$_2$FeSiO$_4$ and Li$_{2.2}$Fe$_{0.9}$SiO$_4$, promising cathode materials for lithium-ion batteries. The aims are to establish the valence and local structure of Fe during charge and discharge to understand if the Fe$^{3+}$/Fe$^{4+}$ redox pair can be reached in the current battery design. It is found that the valence state changes between Fe$^{2+}$ and Fe$^{3+}$, with no evidence of Fe$^{4+}$ before the onset of electrolyte degradation. There is a reversible contraction and extension of the Fe-O bond lengths during cycling while the Fe-Si distance remains constant, which underlines the stability of the Li$_2$FeSiO$_4$ material. The same observations apply to Li$_{2.2}$Fe$_{0.9}$SiO$_4$ cathode material indicating that changing the stoichiometry does not provide any additional structural stability.

Introduction

Progress in the development of Li ion battery cathode materials is based upon the interplay of several defining features driven by the requirements of consumer markets. The materials should conform to criteria associated with energy density, cyclability, safety, and cost to ensure their future use in widespread energy storage applications.$^{1-3}$ The focus of electrochemical and structural characterization studies has been extended from robust layered intercalation oxides, such as LiCoO$_2$ and LiMn$_2$O$_4$ to polyoxoanion materials with tetrahedral structural units.$^4$

The quest for a suitable cathode material has included a focus on lithium iron silicates, which offer many benefits as prospective cathode materials while their complex polymorphism also makes them interesting from a fundamental perspective of solid state chemistry.$^5$ Using abundant elements, like iron and silicon, offer an economic alternative to the Co based system. Safety is a driving factor in battery development and the Si−O bond provides stability similar to the P−O bond in LiFePO$_4$ compounds.$^6$

Another advantage of the iron silicates, is the theoretical possibility to remove two Li ions from Li$_2$FeSiO$_4$, by utilizing the Fe$^{2+}$/Fe$^{3+}$ and the Fe$^{3+}$/Fe$^{4+}$ redox couples, thus, producing a higher capacity than 166 mAh$^{-1}$ for one Li ion. Although capacities greater than 200 mAh$^{-1}$ have been reported by several groups,$^6$ it has yet to be clarified whether it was due to Fe$^{4+}$ formation or electrolyte degradation.$^5$

Li$_2$FeSiO$_4$ has been intensively studied and the crystal structures have been deduced in the pristine (Li$_2$FeSiO$_4$), delithiated (LiFeSiO$_4$) and cycled (LiFeSiO$_4$) states, using X-ray diffraction (XRD) and neutron diffraction measurements.$^{5,9,11}$ Furthermore, the structural properties of various polymorphs have been investigated using computer modeling, including density functional theory (DFT) calculations to explore relationships between the macroscopic electrochemical properties and atomic structures.$^6,14,21$ In particular, the geometry of FeO$_4$ tetrahedra in terms of orientation, size and distortion is thought to influence the equilibrium potential measured in the different polymorphs. As a result the equilibrium potential in the first cycle of the material differs from subsequent cycles, corresponding to an irreversible change in the structure during the first cycle.$^{13,22}$

The pristine material, γ−Li$_2$FeSiO$_4$, is prepared at temperatures between 600°C and 700°C and has a P2$_1$/n space-group. This structure is characterized by edge sharing LiO$_6$ and FeO$_6$ tetrahedra, as well as LiO$_2$ tetrahedra, of which half are pointing in opposite directions.$^{23}$ The delithiated material, inverse β−Li$_2$FeSiO$_4$, with space-group Pmn2$_1$, shows corner linked tetrahedra with all species pointing along the c axis. In this structure the SiO$_4$ tetrahedra are isolated from each other, sharing corners with LiO$_6$ and (disordered) (Li/Fe)O$_4$ tetrahedra. The cycled structure, β$\theta$−Li$_2$FeSiO$_4$, also
crystallise in the $Pmn2_1$ space-group. As for the delithiated material all tetrahedra are pointing in the same direction, sharing only corners with each other. Along the $a$ axis there are chains of ($Li$/$Fe$)O$_4$ parallel to chains of alternating (ordered) LiO$_4$ and SiO$_4$ tetrahedra.

X-ray Absorption Spectroscopy (XAS) is an experimental probe sensitive to changes in the valence and local structure of Fe sites, which can be used to address the questions regarding access to the Fe$^{3+}$/Fe$^{4+}$ redox couple, as well as the structural stability. There have only been two previous in-situ XAS studies of Li$_2$FeSiO$_4$ during cycling. The first study reported EXAFS analysis of only the first Fe-O shell. The second one only reported Fe XANES results.

For the purposes of this study, the focus will be placed on the $\gamma$ (pristine), inverse $\beta$ (delithiated) and the $\beta$ (cycled) polymorphs of Li$_2$FeSiO$_4$, probing changes in the local atomic structure with respect to crystalline order. Establishing how the local structure of Fe changes upon charge and discharge, especially during the first cycle, is essential for tailoring the next generation of cathode materials to the have desired electrochemical characteristics. This is achieved by using in situ EXAFS to monitor the length of the Fe–O bond and longer distance correlations during cycling. By using two samples with different Li and Fe contents in the series Li$_{2-x}$Fe$_x$SiO$_4$, i.e. Li$_2$FeSiO$_4$ and Li$_2$Fe$_{0.5}$SiO$_4$, we investigate the extent to which the structural changes during cycling can accommodate variations in stoichiometry.

In order to address the issue regarding the Fe$^{3+}$/Fe$^{4+}$ redox couple, in situ XANES has been undertaken to estimate the formal valence of the Fe atoms as Li is removed and inserted into the structure during battery cycling.

**Experimental**

**Sample preparation**

Stoichiometric amounts of Li acetate dehydrate (Acros), Fe (II) acetate and tetraethyl orthosilicate were mixed in ethanol with 2mL of acetic acid. After stirring, the suspension was transferred to a Teflon-lined autoclave and heated to 130°C for 12 hours. The resulting gel was dried under vacuum at 80°C, mixed with sucrose and ball milled under acetone for 30 minutes. Following acetone evaporation, the sample was annealed under flowing nitrogen at 600°C for 10 hours. The materials were then transferred to an Argon filled glove box (oxygen and water levels < 1ppm).

**Sample characterisation**

Power X-ray diffraction data were collected on a STOE STADI/P diffractometer operating in transmission mode with Fe K$_\alpha$ radiation ($\lambda = 1.936$ Å) to eliminate Fe fluorescence. X-ray diffraction shows the samples to be highly pure (Figure 1). The Li$_2$FeSiO$_4$ sample is judged to contain approximately 2% impurities probably in the form of Li$_2$SiO$_3$ and Fe$_{1-x}$O. The Li$_2$SiO$_3$ is expected to be inactive during lithiation/delithiation. The Fe$_{1-x}$O is only expected to be active at low voltages (conversion reactions) and this was not seen in our electrochemical measurements. The Li$_2$Fe$_{0.5}$SiO$_4$ sample is judged to be highly pure with negligible impurities. Note however that the diffraction patterns are not highly sensitive to the Li/Fe ratio.

**Figure 2. Electrochemical measurements of a) Li$_2$FeSiO$_4$ and b) Li$_2$Fe$_{0.5}$SiO$_4$.** The black curves refer to galvanostatic charging and discharging of 1½ cycle, whereas the colored curves show the data of the cells used for the in-situ XAS experiments. The labels mentioned here are used for referring to the XAS data shown in the other figures.

**Electrochemical measurements**

Composite electrodes were prepared by ball-milling the active silicate material together with 8wt% of acetylene black and 10 wt% of EPDM in n-hexane. The obtained slurry was pressed on Al mesh and subsequently dried on Al discs having a typical mass of active material of 10–15mg/cm$^2$. Prior to use, the electrodes were further allowed to dry at 90°C in vacuum overnight. The in situ coffee bag-type cells consisted of the active electrode and a lithium metallic counter and reference electrode separated by a Celgard separator. The electrolyte used was a 0.4M solution of high quality LiTDL in EC:DEC (1:1 ratio by volume) the latter purchased from Aldrich. Both solvents and salt were used as received. The samples were charged and discharged at room temperature in galvanostatic mode between 2.0V and 4.8V (Li$_2$FeSiO$_4$) and 2.0V and 4.0V.
(Li$_2$FeSiO$_4$), with a ~13mAg$^{-1}$ current reflecting a C-rate of about C/25 based on complete lithium extraction/insertion using a MACCOR Series 2000 battery cycler (Figure 2).

We note that the polarisation of Li$_2$FeSiO$_4$ is higher than that of Li$_2$Fe$_{0.9}$SiO$_4$, but a clear explanation of that has not been found. One possibility is the formation of a solid-electrolyte interface (SEI), leading to an increased impedance of the cell and hence the polarisation. Nevertheless, the polarisation of Li$_2$Fe$_{0.9}$SiO$_4$ is similar to that observed by Deng et al.$^{30-32}$. They also reported a reduction in the polarisation upon doping with Zn$^{2+}$, Cu$^{2+}$ or Ni$^{2+}$. Clearly, in our case we can see Li$_2$Fe$_{0.9}$SiO$_4$ as a lithium doped Li$_2$FeSiO$_4$ as we replaced iron for lithium.

**In situ X-Ray Absorption Spectroscopy**

Data acquisition was carried out on BM26 (DUBBLE) EXAFS beamline at the European Synchrotron Radiation Facility (ESRF).$^{33}$ Spectra were collected at the Fe K-edge in transmission geometry at room temperature. The beam energy is tuned using a Si(111) double crystal monochromator, which gives an energy resolution of 1.2eV, and was calibrated using an Fe metal foil (setting the first inflection point equal to 7112 eV). The X-ray intensities are detected in ion chambers with 10% absorption in I$_0$ and 70% absorption in I$_1$. Data collection was made with 5eV steps through the pre-edge (7013-7093eV) regime which was shortened to 0.25eV steps while passing through the absorption edge (7094-7146eV) for the XANES measurements. The EXAFS were measured with a k-space interval of 0.05Å between 7146 and 7610eV, corresponding to $k_{\text{min}}$=3 and $k_{\text{max}}$=12Å$^{-1}$.

Figure 3. Calculated Li content for a) Li$_x$FeSiO$_4$ and b) Li$_2$Fe$_{0.9}$SiO$_4$. The black curves refer to galvanostatic charging and discharging of 1½ cycle, whereas the colored curves show the data of the cells used for the in-situ XAS experiments. The labels mentioned here are used for referring to the XAS data shown in the other figures.

The calculations of the amount of lithium extracted and inserted during battery cycling were based on the applied current and elapsed time of the charge and discharge and are plotted against the voltage. The extraction and insertion of lithium is governed by extraction and insertion electrons, thereby changing the valence of the transition metal ion. The latter being extracted from the XANES data below. Based on the model that lithium extraction above 4.2V is associated with electrolyte degradation (as opposed to further extraction of Li-ion from the material) the capacity for the electrolyte degradation has been subtracted, to predict not much lithium is extracted above 4.2V. The data of the calculated lithium contents in the samples are shown in Figure 3. The different states of charge/discharge processes are labeled 1-18 for the Li$_2$FeSiO$_4$ sample and 1-22 for the Li$_2$Fe$_{0.9}$SiO$_4$ sample. For both samples, the results of two cells will be shown here. More cells were analyzed, but they gave the same results (see Supplementary Information).

Figure 4. XANES calibration curve indicating position of the absorption edge at half height $E_{1/2}$ for standards; Fe foil, LiFePO$_4$, FePO$_4$, Fe$_3$O$_4$ and Fe$_2$O$_3$. The standards used for calibration are represented by solid squares while the open squares (Fe foil) are included for comparison.

**XAFS Data analysis**

To determine the valence state of the materials the position of the absorption edge at half height $E_{1/2}$ were measured for a number of standards, i.e. LiFePO$_4$, Fe$_3$O$_4$, FePO$_4$ and Fe$_2$O$_3$ (Figure 4) with varying valence state on the Fe-ions. For comparison the position of the Fe-foil is included in the graph but not included in the calibration curve regarding the valence states.

For EXAFS the absorbance ($\mu(t) = \ln(I_0/I)$) as a function of X-ray energy (E) was analyzed using VIPER$^{34}$ by fitting the pre- and post-edge backgrounds to obtain $\chi(k)$ where $k$ is the modulus of the wave vector of the photoelectron.$^{34}$ The EXAFS spectra give information about bond lengths and coordination numbers. Structural parameters were refined using a least squares fitting routine in EXCURVE9.27$^{35}$ whereby the single scattering fast curved wave theory was used to model the data. The key parameters...
in the model are R, N, and 2σ², which correspond to the radial distance, the number of coordinating atoms and the Debye-Waller (DW) factor (due to static and thermal disorder) for the ith shell of the neighboring atoms, respectively. The additional parameters in the model were calculated ab initio and are: f(k,R), the backscattering amplitude of the scatterer; 2δ(k), the phase shift due to the absorbing atom potential; Φ(k,R), the phase shift due to the scatterer; λ(k), the inelastic mean free path of the photoelectron. Lastly, the parameter S(ω)(k), the amplitude reduction factor due to many electron processes, was estimated to be equal to 0.64 by fitting to reference samples of Fe₂O₃ and γ-Fe₂O₃. A refinement was conducted on k¹ weighted data until R₀, the goodness of fit factor was minimized. This resulted in good fits for the entire k range with R factors less than 30% for the majority of data. The expected errors in R are ±0.02 and in DW factor ±20%.

For the EXAFS measurements, both Li₂FeSiO₄ and Li₂FeO₂SiO₄ have been fitted with a structural model consisting of four oxygen atoms around the central Fe cation at an initial distance of 1.96 Å. This is followed by a secondary shell of four silicon at a starting distance of 3.15 Å, and a further oxygen shell at 3.49 Å. This model is based on analyses of the crystal structures reported for pristine, delithiated, and cycled Li₂FeSiO₄ (Table 1). In fitting this model to the experimental EXAFS, the neighboring Li atoms are disregarded due to their negligible effect on the photoelectron scattering. Refinement of the data was performed by allowing several parameters to vary: E₀ (the Fermi energy, relative to E_l edge position), R, DW, while the coordination number (N) and amplitude reduction factor (S(ω)(k)) were held fixed.

<table>
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</tr>
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<td>4</td>
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<td>7</td>
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<tr>
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<td>4</td>
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<tr>
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<td>7</td>
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</tr>
<tr>
<td></td>
<td>O</td>
<td>3.47</td>
<td>7</td>
</tr>
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</table>

Table 1 A summary of structural parameters for Fe sites in the pristine, delithiated and cycled Li₂FeSiO₄ from crystal structures reported in the literature. Debye Waller refers to static disorder in this instance.

Results and Discussion

Valence State during battery cycling

The galvanostatic curves as a function of time are plotted in Figure 2. In both samples a voltage plateau is observed at approximately 3.2V. In the case of Li₂FeSiO₄ a second plateau is distinguished at about 4.3V. The 4.3V plateau for the Li₂FeO₂SiO₄ sample is not reported here, but has been recorded in other cells. It has been considered in the literature that the high voltage plateau appearing beyond 4V is related either to the Fe⁴⁺/Fe⁴⁺ redox couple or to electrolyte degradation rather than lithium extraction from the host material as indicated in Figure 2 by the blue shaded area.

To determine if it is possible to access the Fe³⁺/Fe⁴⁺ redox pair with the current battery design XANES spectra for the two samples, Li₂FeSiO₄ and Li₂FeO₂SiO₄ during the first two charging cycles were recorded. The bottom right insets in Figure 5 clearly show the change in half-height position of the main absorption edge during the first cycle, which is due to the change in valence of Fe.

First Battery cycle

The oxidation states of Fe are shown in Figure 6 in relation to the degree of delithiation. In the case of pristine Li₂FeSiO₄, the estimated valence is Fe²⁺¹⁰. It may differ from exactly +2 due the small amount of Fe³⁺ impurities inherent in the synthesized material. However, it is evident that removal of lithium causes an increase in the valence of Fe from Fe²⁺¹⁰ to Fe²⁺₂⁺, after charging to 3.9V. Upon reinsertion of lithium the valence decreases until Fe reaches the original value of approximately +2.10. Further insertion of lithium reduces the valence of Fe to the value of +1.95. At this point more lithium has been inserted than removed. In real batteries this would be a purely hypothetical situation because the amount of lithium in the system is defined by the contents of the cathode, rather than a steady supply from a lithium anode. (It is also important to keep in mind the error bars of the XANES data).

Figure 5. In situ XANES data for samples at various points of charge and discharge for a) Li₂FeSiO₄ and b)Li₂FeO₂SiO₄. Top left: pre-edge. Central: normalized XANES. Bottom right: edge position at half height E₁/₂. Spectra of standard materials are included as reference.
The as prepared Li$_{2.2}$Fe$_{0.9}$SiO$_4$ material has an estimated valence state approaching Fe$^{2.3}$. This increases to Fe$^{2.85}$ upon charging to 4V and is further reduced to Fe$^{2.15}$ by inserting the same amount of lithium. Attempts to insert more lithium led to a further decrease in valence to Fe$^{1.85}$.

Second Battery cycle

The XANES measurements during charging to 4.8V do not show any evidence of Fe valence state higher than +3, and hence no evidence for the occurrence of Fe$^{4+}$. As a result, the observed plateau around 4.3V could be associated with electrolyte degradation. However, we note that several groups reported capacities higher than were theoretically predicted when charging beyond 4.2V. Some groups have inferred this to the oxidation of Fe$^{3+}$ to the Fe$^{4+}$ valence state, rather than electrolyte degradation and associated parasitic reactions. One difference in this battery design is the choice of salt, which will affect electrolyte reactions, and possible the lifetime of the Fe$^{4+}$ species, which might be very short lived. We recognise that our cells have been allowed to relax (to circa 4V) prior to XAS measurement, while the cells by Lv et al. were measured during cycling.

On the discharge, i.e. insertion of lithium, all the cells show that more lithium can be accepted than was extracted, irrespective if the cells were charged above 4V or not. This then would suggest a lower Fe valence than +2, but this is not supported by the XANES data (see section 3.2). Instead, it is suggested that the initial samples were slightly oxidized i.e. the Fe valence was originally higher than +2. A higher Fe valence would lead to less Li extraction on the charge. However, on the discharge, all available Fe$^{3+}$ is allowed to be reduced, and since there is more Fe$^{3+}$ than there was initial Fe$^{2+}$, more lithium can be inserted after the first charge cycle. This higher valence of the initial Fe in Li$_2$FeSiO$_4$ or Li$_{2.2}$Fe$_{0.9}$SiO$_4$ has been reported earlier by Lv et al. This Fe$^{3+}$ arises from the air oxidation of the nanoparticulate Li$_2$FeSiO$_4$ and Li$_{2.2}$Fe$_{0.9}$SiO$_4$ material. This air oxidation occurs both during transfer of the as-synthesised material to the glovebox and, in this instance, during the electrode preparation for the in situ measurements.

It is clear from these results that the two samples behave similarly after the first charge cycle, provided we have a lithium source at the negative electrode. Hence, we do not have any indication of stabilisation in the case of the Li$_{2.2}$Fe$_{0.9}$SiO$_4$ sample. It is further stressed that once the material is being used in a cell without a Li source, such as with carbonaceous negative electrodes, these Li$_{2.2}$Fe$_{1.4}$SiO$_4$ materials need to be made impurity free.

Local structure

Coordination numbers

In Figure 5 the top left insets show the details of the pre-edge peaks of the XANES spectra. The pristine Li$_2$FeSiO$_4$ sample displays two distinct peaks in the pre-edge region of the spectra. The peaks correspond to Fe$^{+2}$ in a tetrahedral environment. As lithium is removed from the cathode Fe$^{2+}$ is oxidised to Fe$^{3+}$ giving rise to a singular pre-edge feature, corresponding to Fe$^{3+}$ in a tetrahedral environment. Upon re-insertion of Li the two peaks, characteristic of Fe$^{3+}$ in the tetrahedral environment, re-appear again in the pre-edge spectra.

Figure 6: Amount of lithium (from electrochemistry) vs valence (from XANES data). The dashed red and blue lines indicate a theoretical value of Fe valence, during lithiation/delithiation for Li$_2$FeSiO$_4$ and Li$_{2.2}$Fe$_{0.9}$SiO$_4$ respectively.

Figure 7: Radial distribution functions by Fourier Transform of k$^2$ weighted Fe EXAFS spectra (calculated in the k range of 3.0 to 12 Å$^{-1}$) for pristine, delithiated and cycled structures. a) Li$_2$FeSiO$_4$ b) Li$_{2.2}$Fe$_{0.9}$SiO$_4$. The best fit parameters for the EXAFS are given in Table 2.

Bond Lengths

Figure 7 shows the Fourier transform (FT) of the EXAFS spectrum for the two samples at different stages during the first charging cycle. There are no reports in the literature for crystal
structures of Li$_2$Fe$_{0.9}$SiO$_4$ so the same approach to EXAFS analysis has been used for both Li$_2$FeSiO$_4$ and Li$_2$Fe$_{0.9}$SiO$_4$ samples.

Considering the two as prepared samples there is a prominent peak in both spectra, corresponding to the first shell of Fe-O bonds at approximately 2Å. A second prominent peak is observed at approximately 3.5Å associated with the Fe-Si neighbours and a second Fe-O shell. Table 2 shows the fitted results for the two samples.

The radial distribution functions shown in Figure 7 for both samples show reversible differences in both amplitude and bond length of the first shell (Fe-O) as the sample is charged and then discharged. The changes in bond length of the first shell (Fe-O) are shown in Figure 8. Combining the estimated valence from the XANES measurements with the fitted value of Fe-O bond lengths presents a linear, inverse relationship. As expected when Fe is located in a tetrahedral environment with a variable valence state of Fe$^{+2}$ to Fe$^{+3}$.

The pristine Li$_2$FeSiO$_4$ sample has an estimated valence of $+2.10$, in a tetrahedral environment with a bond length of 1.96Å in fair comparison with the literature, which reports results for Fe$^{+2}$ in a tetrahedral environment with a typical bond length of 2.00Å. This compares with Sirisopanaporn et al who reported the $\gamma$ structure to have a Fe-O bond length of 2.03Å while a similar theoretical value of 2.05Å was reported by Eames and coworkers.

Upon delithiation of the Li$_2$FeSiO$_4$ sample the estimated valence increases to $+2.95$ with an associated shortened bond length to 1.89Å, which is again consistent with the literature that reports a Fe-O distance of 1.88Å for Fe$^{+3}$ in a tetrahedral environment. Computations by Eames et al. on the inverse $\beta\gamma$ polymorph give a value of 1.92Å. It is evident that removing lithium from the sample causes an increase in valence and subsequent contraction of the Fe-O bond. As lithium is re-intercalated into the bulk of the cathode the estimated valence returns to $+2.10$ with an associated Fe-O bond length of 1.96Å. Then there is further re-insertion of Li which leads to a slightly lower valence of $+1.95$, seen as an additional lengthening of the Fe-O distance to 1.98Å. This is good agreement with Armstrong et al (2.03Å) and Eames et al (2.05Å) for the $\beta\gamma$ cycled structure.

The information obtained from fitting the first Fe-O shell includes relationship of the Debye Waller factor to valence state. It is evident from Figure 9 that the Debye Waller factor is high in the pristine material indicative of a distorted FeO$_4$ tetrahedral site. As the cathode undergoes charging, the DW factor increases slightly due to the mixed sites of Fe$^{+2}$ and Fe$^{+3}$ with different bond lengths. As the valence increases to approximately Fe$^{+3.6}$ the DW factor starts to decrease. This infers an increased proportion of Fe$^{+3}$ having a narrower range in Fe-O bond lengths in accordance with more regular FeO$_4$ tetrahedra. Eames et al. suggests that upon delithiation the distortion of the FeO$_4$ tetrahedra is reduced and the reduction in the Fe-O Debye Waller factor from the EXAFS analysis is consistent with this.

By analyzing the remaining shells it is possible to relate the local structural changes to the reported crystal structures for pristine, delithiated, and cycled materials. Figure 10 illustrates the variation in distances for the second and third shells. For both Li$_2$FeSiO$_4$ and Li$_2$Fe$_{0.9}$SiO$_4$ samples the second shell (Fe-Si) does not exhibit
discernible changes in amplitude or peak position while the third shell (Fe-O) shows increased amplitude and shifts to longer distance upon cycling. It indicates that the Fe-Si distance is relatively unchanged upon cycling, which is consistent with literature reports in Table 1. The Fe-Si distance in the pristine material is 3.15Å in good comparison with 3.12Å reported by Eames et al. Delithiation of the material does not change the Fe-Si distance of 3.15Å. This agrees with the predictions regarding the delithiated inverse βII structure where a Fe-Si distance of 3.11Å is reported. The strength of the Fe-O-Si connection provides the material with its thermal stability and therefore enhanced safety attributes. Note that the Fe-Si distance corresponds to Fe-O-Si connected tetrahedra, where Si-O is constant and Fe-O contracts during delithiation, but the relative orientation of the tetrahedra change to keep the Fe-Si distance approximately constant. This can be explained by a competitive tradeoff between the attractive forces of Fe-O and the cation–cation repulsion between Fe³⁺ and Si⁴⁺, which results in the Fe-O-Si bond angle increasing.

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<tbody>
<tr>
<td>Point 1</td>
<td>O</td>
<td>1.97 (1)</td>
<td>4</td>
<td>0.015 (1)</td>
<td>30</td>
</tr>
<tr>
<td>P₂/n (γ₁)</td>
<td>Si O</td>
<td>3.13 (1)</td>
<td>4</td>
<td>0.018 (3)</td>
<td></td>
</tr>
<tr>
<td>Pristine</td>
<td>O</td>
<td>3.48 (4)</td>
<td>7</td>
<td>0.042 (9)</td>
<td></td>
</tr>
<tr>
<td>Point 8</td>
<td>O</td>
<td>1.89 (1)</td>
<td>4</td>
<td>0.007 (1)</td>
<td>15</td>
</tr>
<tr>
<td>Pmn₂, (i-β₀)</td>
<td>Si O</td>
<td>3.12 (1)</td>
<td>4</td>
<td>0.039 (9)</td>
<td></td>
</tr>
<tr>
<td>Delithiated</td>
<td>O</td>
<td>3.64 (2)</td>
<td>7</td>
<td>0.033 (8)</td>
<td></td>
</tr>
<tr>
<td>Point 17</td>
<td>O</td>
<td>1.98 (1)</td>
<td>4</td>
<td>0.013 (1)</td>
<td>31</td>
</tr>
<tr>
<td>Pmn₂, (β₀)</td>
<td>Si O</td>
<td>3.14 (2)</td>
<td>4</td>
<td>0.021 (4)</td>
<td></td>
</tr>
<tr>
<td>Cycled</td>
<td>O</td>
<td>3.43 (6)</td>
<td>7</td>
<td>0.059 (9)</td>
<td></td>
</tr>
</tbody>
</table>

Table 2. Structural parameters for Li₂FeSiO₄ and Li₁₂Fe₀.₅SiO₄ at states of charge associated with the pristine, delithiated and cycled materials – corresponding to points in Figure 2. The coordination number (N) remained fixed while the distance (R) and Debye Waller (2$\sigma$) were free to vary until the Rf was minimized. Uncertainty of the last digit is given in parentheses.

Figure 10 shows that the third shell Fe-O distance gets longer approximately linearly with change in valence. The distances obtained from EXAFS fitting increase in length from 3.40Å to 3.65Å from pristine to delithiated material. The same trend of lengthening the Fe-O distance in the 3rd shell is seen in the reported crystal structures in Table 1, from Armstrong (pristine) and Eames et al (delithiated and cycled). The increase in the Fe-O distance, corresponding to the third shell, arises from Fe-O-Si-O linkages and increases due to the increase in the Fe-O-Si bond angle as discussed previously.

Conclusions

Detailed in situ Fe K-edge XANES and EXAFS results have been presented for highly pure samples of Li₂FeSiO₄ and Li₁₂Fe₀.₅SiO₄ cathode materials.

- Analysis of the in situ XANES data shows that the Fe valence remains between Fe²⁺ and Fe³⁺, while Fe⁴⁺ was not evident in this voltage regime. Instead, we attribute any additional capacity to electrolyte degradation at voltages above 4.2V in this study.
- Air oxidation of the sample during electrode preparation accounts for the substantial difference between charge and discharge capacities during the first electrochemical cycle. Therefore, more lithium can be inserted upon discharging, than can be removed during charging.

In situ Fe K-edge EXAFS data have been analysed to include details of the first three neighbouring shells, Fe-O, Fe-Si and Fe-O. These data obtained during the first cycle with charging to 4.8V provide an important experimental comparison for the structural models of pristine, delithiated and cycled Li₂FeSiO₄ which have been reported in the literature.

- The quantitative Fe valence estimates from XANES, and the Fe-O bond length from EXAFS are both consistent with the cycling of valence state from Fe²⁺ to Fe³⁺ in the reported structures. The first shell Fe-O DW factor, the second shell Fe-Si, and the third shell Fe-O distances also change according to previously reported structural models.
- The constant Fe-Si distance, despite changes to the environment of the FeO₄ tetrahedra, underlines the stability of the Li₂FeSiO₄ material, and hence its desirability in battery applications from a safety point.

This study has also presented the first results for the closely related Li₁₂Fe₀.₅SiO₄ cathode material. All of the above observations also apply to this material suggesting that changing the stoichiometry in this manner does not provide any additional structural stability.

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Notes and references

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The valence and local structures of Fe during battery cycling of Li$_3$FeSiO$_4$ and Li$_{2.2}$Fe$_{0.9}$SiO$_4$ are studied by in-situ (XAS) measurements.