

Dependence of Li₂FeSiO₄ Electrochemistry on Structure

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Supporting Information

ABSTRACT: Small differences in the FeO₄ arrangements (orientation, size, and distortion) do influence the equilibrium potential measured during the first oxidation of Fe²⁺ to Fe³⁺ in all polymorphs of Li₂FeSiO₄.

Lithium transition metal silicates, Li_2MSiO_4 (M = Fe, Mn, Co),¹ Lihave attracted a lot of interest recently as positive electrodes for Li-ion batteries due to their potentially high theoretical capacities and to their rich crystal chemistry as they undergo complex polymorphism. Polymorphs of these "tetrahedral" structures can be classified into low- and high-temperature forms, which differ in the ordering/distribution of cations within tetrahedral sites of a hexagonal close-packed-based arrangement of oxygen.² Recent progress in the structure determination of Li_2MSiO_4 polymorphs² paves the way for a detailed study of the mechanisms of Li extraction/insertion from/into these silicates as a function of local and/or long-range order of cations. The achievements in the synthesis of electrochemically active Li_2FeSiO_4 (small, carboncoated particles) prompted us to study in detail their electrochemical properties as a function of the crystal structure adopted.

We carefully prepared and used three polymorphs of Li_2FeSiO_4 with very high purity. A pristine powder had been obtained first under hydrothermal conditions³ at 200 °C, labeled as LFS@200 and subsequently annealed at either 700 °C (LFS@700) or 900 °C (LFS@900) for 6 h and quenched to room temperature. Hydrothermal synthesis, experimental techniques used (X-ray diffraction, microscopy, electrochemical characterizations, Mössbauer spectroscopy, electrode preparations), and C-coating procedure are given in the Supporting Information.

The differences in the local environments of Fe^{2+} for between the three polymorphs are shown in Scheme 1. They differ in the interconnectivity of LiO_4 and SiO_4 tetrahedra and in their respective orientations along a given crystallographic direction. For all three polymorphs, only corner sharing occurs between FeO₄ and SiO₄ due to their strong discrepancy in O–O edge lengths. Each corner of a given FeO₄ tetrahedron, for all three polymorphs, is connected to two Li⁺ and one Si⁴⁺ cation.

- In LFS@900 (*Pmnb* S.G.),^{2d,e} a given FeO₄ tetrahedron shares two edges with LiO₄ tetrahedra
- In LFS@700 $(P2_1/n \text{ S.G.})$,^{2a,d} only one edge is shared between FeO₄ and LiO₄
- In LFS@200 (*Pmn*2₁ S.G.),^{1b} only corner sharing occurs

Scheme 1. Local Environments around FeO₄ Tetrahedra (In Green) in the Three Polymorphs of Li_2FeSiO_4 (LiO₄ in Gray, SiO₄ in Blue)



Table 1. Average Distances of Fe–O, Li–O, and Si–O Bonds and FeO₄ Distortion

	LFS@200	LFS@700	LFS@900
d_{avg} (Fe–O)/Å	2.076(3)	2.032(2)	2.026(1)
d_{avg} (Li–O)/Å	1.968(2)	1.970(1)	1.964(2)
d_{avg} (Si–O)/Å	1.636(2)	1.634(2)	1.656(4)
FeO ₄ distortion ⁵ /Å ³	2.3×10^{-4}	9.9×10^{-4}	12.8 × 10 ⁻⁴

As seen in Table 1, the differences in local environments around a given FeO₄ tetrahedron are immediately translated into varying degrees of distortion and average Fe-O bond lengths. Compared with usual distances in tetrahedral environments of oxides,⁴ LFS@900 and LFS@700 show contraction of the average Fe-O distances so as to accommodate edge sharing with the slightly smaller LiO₄ tetrahedra. Importantly, shorter average Fe-O interatomic distances imply stronger (more covalent) Fe-O bonds, which should have a direct effect onto the redox potential of the Fe^{2+}/Fe^{3+} couple. Indeed, seminal work by Goodenough's group on polyanion structures (sulfates, tungstates, molybdates, phosphates)⁶ demonstrated that a more ionic Fe-O bond (hence longer) resulted in lowering the energy of the antibonding states and thus increasing the voltage vs Li⁺/Li⁰ at which oxidation of Fe^{2+} into Fe^{3+} would take place. This is particularly true for LiFePO₄, which shows edge sharing between FeO₆ and PO₄ and displays^{6b,c} the highest value of voltage vs Li for iron in phosphates: $3.42 \text{ V vs Li}^+/\text{Li}^0$.

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Figure 1. (a) Equilibrium potentials measured from GITT. (b) Derivative plots obtained from PITT in the first oxidation of three polymorphs.

Three well-defined (crystallographically) samples (LFS@-200, LFS@700, and LFS@900) were used for electrochemical characterization in order to investigate the influence of crystal structures on their electrochemical properties. Within the aim to homogenize the particle size and to improve electron contacts in the non-carbon-coated samples, we performed ball milling with Ketjen black under an argon atmosphere. Typical particle size used in our study was between 60 and 150 nm. Rietveld refinements of both pristine materials and electrode composites have been performed so as to confirm the crystal structures of the materials used (Figures S1 and S3 and Tables S1, S2, S4, and S5 in Supporting Information).

The precise positions of the Fe^{2+}/Fe^{3+} redox couple vs Li^+/Li^0 for each polymorph were carefully investigated using GITT and PITT techniques, which enable measurements close to the equilibrium potential (see Supporting Information). The use of GITT clarifies the difference in the first oxidation potential among the three polymorphs, as shown in Figure 1a. The lowest potential was observed in the case of LFS@900 (2.90 V vs Li^+/Li^0), while the highest was that of LFS@200 $(3.10 \text{ V vs Li}^+/\text{Li}^0)$. The equilibrium potential of LFS@700 was close to 3.00 V vs Li⁺/Li⁰. PITT measurements have been performed to substantiate the observed differences in the equilibrium potentials among the various polymorphs. The results are in agreement with those measured from GITT; the first oxidation potential (vs Li⁺/Li⁰) increases from LFS@900 to LFS@200, as shown in Figure 1b. One may note that the potentials measured from GITT are lower than those obtained from PITT. This could be explained by the activation energy needed for initial lithium extraction. Importantly, the coherence of the results obtained from both electrochemical techniques has confirmed the influence of the crystallographic structures on the position of the Fe^{2+}/Fe^{3+} redox potential in Li₂FeSiO₄. In other words, shorter average Fe-O bond lengths (i.e., higher covalency of the Fe-O bonds) and higher degree of distortion of the FeO_4 tetrahedra result in higher Fe^{2+}/Fe^{3+} redox energy (hence smaller voltage difference with Li⁺/Li⁰) through the inductive effect.

The use of PITT technique has pointed out another peculiarity of the rich polymorphism of Li₂FeSiO₄. The electrochemical measurements of the three polymorphs, besides showing the "classical" irreversible change of cell voltage from \sim 3.0 to \sim 2.8 V between the first and subsequent charges,⁷ highlight significant differences in the kinetics associated with these transformations.

As seen in Figure 2a-c, the high-temperature polymorphs (LFS@900 and LFS@700), with higher degree of "disorder" in terms of cation connectivity, are prone to transform into the thermodynamic favorable phase faster than the low-temperature



Figure 2. Derivative plots obtained from PITT measurements in the first, second, and fifth cycles for all three polymorphs.



Figure 3. (a) Galvanostatic curves in the first cycle. (b) Derivative plots of galvanostatic curves in the first cycle of LFS@700 polymorph.

(LFS@200) one. The first discharge of LFS@900 and LFS@700 occurs at a reduction potential of ~2.76 V vs Li⁺/Li⁰, while two reduction peaks at ~2.76 and ~3.04 V were observed in the case of LFS@200. The potential around 3 V is associated with the remaining initial Li₂FeSiO₄ phase, whereas the new Li-poor phase, originated upon first oxidation, shows the reduction potential at ~2.76 V. Moreover, an oxidation peak at 3.13 V could be observed in the second cycle of LFS@200, indicating a significant amount of initial LFS@200 phase (Figure 2b). Subsequently the irreversible transformation of LFS@200 is completed in the fifth cycle, while only first oxidation is needed for the structural rearrangement of LFS@700 and LFS@900.

Using conventional galvanostatic measurements, we observed that the applied current density is also important to the phase transformation kinetics. For instance, as seen in Figure 3, the residual of the initial LFS@700 phase could be observed only for the battery which cycled with high current density. Hence, not only the crystal structure but also the current density used in the experiment influence the kinetics of thermodynamic stabilization. Similar effects have been observed by varying the temperature at which the electrochemical tests were performed.

In this Communication, we used three very well-defined crystallographically pure Li₂FeSiO₄ samples crystallized in *Pmnb* (LFS@900), *P*2₁/*n* (LFS@700), and *Pmn*2₁ (LFS@200) space groups. Variations in the FeO₄ arrangements (orientation, size, and distortion) influence the equilibrium potential measured during the first oxidation of Fe²⁺ into Fe³⁺ in all polymorphs. The shorter (stronger) Fe–O bonds result in the higher splitting energy between bonding and antibonding states, and thus lowering the Fe²⁺/Fe³⁺ redox potential vs Li⁺/Li⁰. More surprisingly, phase transformation kinetics of Li₂FeSiO₄ upon cycling has structural, cycling rate, and temperature dependence.

ASSOCIATED CONTENT

Supporting Information. Experimental part, refined X-ray diffraction patterns, tables with crystallographic data, and SEM micrographs. This material is available free of charge via the Internet at http://pubs.acs.org.

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