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TEM Studies of Carbon Coated LiFePO₄ after Charge Discharge Cycling

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Carbon coating has proven to be a successful approach to improve the rate capability of $LiFePO₄$ used in rechargeable $Li-ion$ batteries. Investigations of the microstructure of carbon coated $LiFePO₄$ after charge discharge cycling shows that the carbon surface layer remains intact over 100 cycles. We find micro cracks in the cycled material that extend parallel to low indexed lattice planes. Our observations differ from observations made by other authors. The differences between the orientations of crack surfaces in both studies can be reconciled considering the location of weak bonds in the unit cell and specimen geometry as well as elastic stress fields of dislocation.

Introduction

 $LiFePO₄$, first introduced by Padhi et al, is considered a promising low cost, environmentally benign cathode material for application in batteries for electric and hybrid vehicles (1). The removal of Li from LiFePO₄ proceeds via a two phase reaction in which heterosite (FePO₄) forms directly from the fully lithiated tryphilite LiFePO₄. LiFePO₄ and the delithiated FePO₄ belong to the group of phospho-olivines. The orthorhombic unit cell (SG 62, Pnma symmetry) consists of a distorted hexagonal close packed stacking of oxygen into which octahedrally coordinated Fe and Li ions and tetrahedrally coordinated P-ions are embedded. The migration of Li-ion takes place parallel to the b direction in tunnels between $FeO₆$ octahedra and $PO₄$ tetrahedra (2). The unit cell dimensions are a=10.33Å, b= 6.01\AA , c= 4.69\AA for LiFePO₄ and a=9.76Å, $b=5.75\text{\AA}$, $c=4.76\text{\AA}$ for FePO₄. The change in unit cell volume when going from tryphilite to the heterosite amounts to approximately 6.8%, which induces a considerable amount of stress at the two phase interface (3). Despite its high theoretical capacity of 170 mAh/g a major obstacle to the commercialization of $LiFePO₄$ is its poor electronic conductivity that limits achievable discharge rates (1). Reduction in particle size (4), carbon coating (5-7) and addition of small metallic particles (8) are among the approaches used to enhance the electrochemical performance of $LiFePO₄$. We successfully employed carbon coating with pyromellitic acid and ferrocene to enhance the rate capability of electrodes based on LiFePO₄ (9). Studies of the virgin material by Energy Filtered Imaging (EFTEM) showed that a carbon layer thickness of about 10 nm is sufficient to improve the electrochemical behavior (10).

In the original paper Padhi introduced a simplified core shell model for the distribution of the two phases (1). The highly anisotropic unit cell of $LiFePO₄$ and the resulting preferred direction of Li diffusion however raise concerns about the validity of the core shell model. Recently Chen et al. investigated $LiFePO₄$ before and after chemical delithiation by transmission electron microscopy (11). In $Li_{0.5}FePO₄$ the authors found regions of alternating contrast that were sometimes separated by cracks with (100)

surface planes. A high resolution image taken from a thin region near a crack suggests that stripes of $LiFePO₄$ and $FePO₄$ extend parallel to (100) planes. The finding of alternating FePO₄ and LiFePO₄ zones was qualitatively confirmed by a high resolution electron energy loss study (12) reviving the debate about the location of the $LiFePO₄/FePO₄$ interface.

Here we report the results of investigations by Transmission Electron Microscopy on carbon coated $LiFePO₄$ particles retrieved from cycled electrodes. We found cracks similar to those observed after chemical delithiation. However a major difference exists in the orientation of the crack surfaces. We consider the geometry of crack formation in $LiFePO₄$ based on dislocation glide systems and compare the orientation of cracks found in our material to observations made on $LiFePO₄$ synthesized by a hydrothermal route (11).

Experimental

 $LiFePO₄$ was synthesized by a sol gel method using iron nitrate, lithium acetate, phosphoric acid and glycolic acid precursors as detailed in reference (9). After initial firing at 500 $^{\circ}$ C to decompose these precursors, 6wt% pyromellitic acid and 1wt% ferrocene was added and the mixture was subsequently fired to 600ºC for 10 hours under N₂ atmosphere. Laminated electrodes containing 80 wt $%$ active material, 8 wt % Kynar poly(vinylidene fluoride), 6 wt % SFG-6 synthetic flake graphite, and 6 wt % compressed acetylene black were prepared. Electrodes were punched out to 1.8 cm^2 size, with loadings of about 5-10 mg/cm² active material. 2032 size coin cells were assembled in a helium-filled glove box, using lithium metal as a counter electrode and 1 M LiPF₆ in 1:2 ethylene carbonate/dimethyl carbonate (EC/DMC) as the electrolytic solution. Cells were cycled galvanostatically between 2.0 and 3.9V using a MacPile II potentiostat/galvanostat. All cells were charged at C/25 rate, and discharged at either $C/25$ (cell 1) or $C/2$ (cell 2) rates. The cycling behavior of the two cells varied as shown in Fig.1. At the lower discharge rate of $C/25$ about 145 mAh/g was retrieved from the cathode per discharge with no apparent decline in performance over the range of the experiment. At the higher discharge rates used in cell 2, a smaller specific

Figure 1 : Comparison of the specific capacities observed at discharge rates C/25 and $C/2$.

capacity of 125mAh/g is observed that begins to deteriorate after about 50 chargedischarge cycles. The cells were stopped after approximately 100 charge discharge cycles and disassembled. The graphite/binder/powder mixture was scraped off the Al collectors and washed repeatedly in NMP to dissolve the binder followed by a wash in acetone. The particle/acetone solution was dispersed on a holey carbon grid to prepare TEM specimens. For comparison, $Li_{0.5}FePO_4$ obtained by chemical delithition of LiFePO₄ produced by a hydrothermal method (11) was also studied. TEM investigations were performed on a JEOL 3010 at the National Center for Electron Microscopy in Berkeley and the JEOL 2010 at the University of New Orleans. Both microscopes were operated at 200kV. Experimental electron diffraction patterns were compared to electron diffraction patterns simulated with the software packet Desktop Microscopist using unit cells published in literature (3).

Results

No qualitative microstructural differences were observed between particles retrieved from cell 1 and cell 2. The most prominent feature observed in the cycled particles is the formation of large cracks parallel to low indexed lattice directions. The cracks start at the specimen surface and extend into the interior of the particle. An example from a particle retrieved from cell 1 is shown in Fig. 2 together with the corresponding diffraction pattern. The orientation of the crack surface is parallel to (010) lattice planes. Close examination of the particle surface in specimens from both electrodes showed that the carbon surface layer has remained intact over more than 100 charge discharge cycles. Fig. 3 shows an example of a particle in [011] orientation demonstrating homogeneous layer thickness over a wide range of particle surface orientations. A close look at the selected area diffraction patterns in Fig. 2b and Fig 3b shows that spot splitting can be observed on diffraction spots further away from the incident beam (see arrow). The small separation of the spots is typical for the small difference in lattice parameters between

Figure 2 : Image showing cracks parallel to (010) planes and corresponding diffraction pattern taken from a $LiFePO₄$ particle retrieved from cycled cell. Cracks are observed repeatedly in particles retrieved from cycled cell 1 and 2.

Figure 3 : Carbon surface layers on cycled particles show no damage or orientation dependence of layer thickness (cycled cell 1).

Figure. 4 : Image (a.) and diffraction pattern (d.) taken from $Li_{0.5}FePO_4$ synthesized from hydrothermally produced LiFePO₄. The crack surface is oriented parallel to (100) planes. The enlarged section in b. shows that the crack surface extends over several lattice planes (see arrow). Further enlargement in c. shows that twinning or stacking faults are present in the material.

closely related phases like $LiFePO₄$ and $FePO₄$. The analysis of single crystal electron diffraction patterns taken from the cycled and from the as synthesized material showed that no preferred particle orientation could be found in the prepared TEM specimens.

Fig. 4a shows a micrograph from chemically delithiated $Li_{0.5}FePO_4$. Cracks similar to the ones observed after charge discharge cycling were observed. However in these specimens the orientation of the crack surface is parallel to (100) planes as can be seen from a comparison to the diffraction pattern shown in Fig. 4d. A magnified view of the crack surface in Fig.4b shows that the materials separation is not clear cut between two lattice planes. Instead the crack surface extends over several lattice planes as indicated by an arrow in Fig. 4b. Further enlargement of the area near the crack surface in Fig. 4c shows a change of the image contrast that indicates presence of twins or stacking faults in this area. Imaging of a delithiated particle in 2 beam condition shows a concentration of strain along lines that are parallel to [101] directions in the plane of the particle, see Fig. 5. The particles synthesized by the hydrothermal method have a plate like shape with the [010] direction parallel to the surface normal of the particle.

Figure 5 : $Li_{0.5}FePO_4$ imaged with diffraction intensity originating from (020) planes. The dark lines lying diagonally in the particle plane indicate presence of stress in the surrounding regions. The line directions are approximately parallel to [101] directions.

Discussion

The finding that the carbon surface layer withstands charge discharge cycling over many cycles without any visible damage or alterations is encouraging for future practical implementation of the coating method used in this study. The presence of micro cracks in particles retrieved from both cathodes suggests that these features are not responsible for the observed difference in electrochemical behavior. This is in contrast to a finding published in (13) where observed capacity loss is attributed to crack formation. In order to fully eliminate crack formation as a factor affecting electrochemical behavior a more quantitative investigation of the number of cracks observed in particles from either cathode is needed.

The different orientations of crack surfaces observed in cycled powders produced by sol gel method and in chemically delithiated powders produced hydrothermally is a disturbing result that asks for an explanation. We looked at the building blocks of the

microstructure to understand how the breaking of bonds between the building blocks of this structure can explain both observations. In a second approach we estimate the order of elastic stresses in the material that needs to be overcome to form cracks by dislocation glide.

Fig. 6 shows schematic drawings of the olivine structure projected into the (001) plane (Fig. 6a) and into the (100) plane (Fig. 6b). Fig. 6a shows how sheets of edge sharing FeO₆ octahedra (shaded brown) form (100) planes that are connected in [100] direction by PO_4 tetrahedra (shaded grey). The bond between two (100) planes is where the corners of PO_4 tetrahedra and FeO_6 octahedra meet- making this a weak bond which explains the formation of cracks between (100) planes. A plan view of a (100) plane in Fig. 6b shows that the sheets of edge sharing $FeO₆$ octahedra (shaded red) are better described by serrated chains of edge sharing $FeO₆$ octahedra parallel to the [100] direction. In [010] direction these chains are connected by PO_4 tetrahedra that extend above and below the plane of the drawing (shaded yellow and blue respectively). Hence the formation of cracks between (010) planes can proceed by breaking of the same type of bonds between FeO₆ and PO₄ corners as in the case of breaking between (100) planes. A more quantitative view is to consider actual stresses that need to be overcome in order to create a crack by a dislocation glide mechanism.

Figure 6 : Arrangement of PO_4 tetrahedra and FeO_6 octahedra in LiFePO₄ seen in the (001) plane (a.) and in the (100) plane (b.)

Dislocations are one dimensional lattice defects that can be described by a deviation of actual atom positions from their assigned lattice sites. The movement of such a defect through the volume of a specimen is a mechanism of plastic deformation. Another mode of deformation is twinning (not considered here). The magnitude and direction of the atom displacement caused by the presence of a dislocation is described by the Burgers vector ים.
ד *b* , for example Eispracement along unferent directions and in direction planes (gride planes) will cause
cracks in different directions. The combination of a Burgers vector and a glide plane is $\frac{5}{1}$ *b* =[100] describes a displacement in [100] direction by the length of one lattice constant in that direction $(a=10.33\text{\AA})$. For comparison the lengths of Burgers vectors in [010] and [001] direction are 6.01Å and 4.7Å respectively. Displacement along different directions and in different planes (glide planes) will cause called a slip system. Table I compares the slip systems that can lead to formation of cracks having (100) and (010) surface planes. Notice that a dislocation with the Burgers vector [001] is glissile in both the (100) and the (010) plane.

The elastic strain energy stored in a material due to presence of a dislocation is proportional to the product of Burgers vector principles calculations (14). Using the elastic constants published by Maxisch we $\frac{1}{i}$ *b* and bulk modulus G. For isotropic materials it follows that the strain energy is smallest for dislocations with the shortest Burgervectors- and those are the dislocations that will be formed. LiFePO₄ however is not isotropic as has been shown recently by Maxisch and Ceder by applying first estimate qualitative differences between elastic strain fields associated with dislocations of the three $\langle 100 \rangle$ type Burgers vectors. The estimated products G \vec{r} *b* for the dislocations considered here are listed together with the values used for the elastic moduli in the direction of the Burgers vectors in table II.

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It can be seen that a dislocation with a Burgers vector enable energy of a disfocution the formation of energy is ravorable in both observed
directions. A better judgment on the activated glide systems can be obtained by the r *b* =[001] that is glissile in both (100) and (010) planes has the lowest amount of elastic stress associated with it. For dislocation glide the elastic stress needs to be overcome. Therefore from the standpoint of elastic energy of a dislocation the formation of cracks is favorable in both observed superposition of misfit stresses between two phases with elastic stress of dislocation lines as has been shown in precipitation hardened alloys (15).

Finally looking at the particle geometry of the $LiFePO₄$ powders used in both studies (sol gel versus hydrothermal) one may speculate that in case of the plate like particles having [010] surface normals cracks between (010) planes cannot be seen in transmission. On the other hand particles synthesized by the sol gel method have an arbitrary orientation of surface planes, which makes it more likely to observe both: cracks between (010) and between (100) planes. A closer look into Fig. 2a shows that perpendicular to the cracks between (010) planes smaller crack are observed. From the indices in this zone axis these smaller cracks are roughly between (701) planes which are at an angle of 3.71º to (100) planes.

Summary

We investigated carbon coated $LiFePO₄$ subjected to charge discharge cycling by transmission electron microscopy. Our results show that the carbon surface layer produced by our method withstood the cycling and was fully intact after over 100 charge discharge cycles. Micro cracks have been observed independently of the applied discharge rate. The surface orientation of the cracks observed in this study differs from the orientation of cracks observed in other studies. However considerations of the

 ^{*} β Burgers vectors of $\langle 110 \rangle$ type are not considered here for a lack of data on the elastic constants in these directions.

microstructure and elastic strain energy due to dislocations within the particles show that both orientations are feasible. The particle geometry of the delithiated particles suggests that only cracks parallel to the incident electron beam are visible in transmission electron microscopy while those perpendicular to the beam would not be detected if they were present.

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