Electrolyte Stability and Discharge Products of an Ionic-Liquid-based Li-O₂ Battery Revealed by Soft X-Ray Emission Spectroscopy

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Abstract

We investigate the electrolyte stability and discharge products of an ionic-liquid-based Li-O$_2$ battery through soft x-ray emission spectroscopy (XES) experiments, which offer unique site specificity for detecting subtle changes in the local nitrogen and oxygen environment. We benchmark the valence electronic structures of the molecules composing the electrolyte, namely the solvent PP13(TFSI), the salt LiTFSI, and the PP13 cation. Then, the transformation of the electrolyte is shown using cathodes stopped at different discharge and charge stages. We provide experimental evidence that the nitrogen site of the salt is unstable during electrochemical operation. The chemical environment of the nitrogen atoms is gradually changed during the electrochemical cycle, indicating a breaking of S-N bonds. The ionic liquid solvent remains mostly as an ion pair, but some decomposition into PP13 cations and TFSI anions cannot be ruled out. Our results and detailed analysis also show that the discharge products in our cell consist of lithium peroxide with some amount of hydroxide; however, no carbonate is observed.
1. Introduction

Li-O$_2$ battery technology, with a potential energy density of 500 Wh/kg,$^1$ has been considered for the next generation of rechargeable batteries. However, low capacity, poor rate capability, low round-trip efficiency, and short cycle life are technical limitations$^{2,3}$ inherent to this battery caused by the special chemical environment and the complex chemistry of gaseous, liquid, and solid reaction products. The practical barriers and the complicated reactions require advanced characterizations, especially elemental- and site-sensitive (ideally \textit{in situ/operando}) spectroscopy, to clarify the complicated chemistry associated with critical challenges on both the electrode and the electrolyte.$^{4,5}$

The oxygen electrode is usually made of a three-dimensional network structure that features high surface area and excellent electronic conductivity. However, the electrode undergoes passivation during the discharge process because of the formation of poorly conducting Li$_2$O$_2$ on the carbon cathode.$^6$ Therefore, novel approaches are developed based on synthesizing advanced nanostructured carbon electrodes,$^{7,8,9}$ coating the existing carbon cathodes with non-carbon materials,$^{10,11,12}$ or designing a complete carbon-free electrode$^{13,14,15}$ to minimize the degradation of the oxygen electrode.

The electrolyte, based on a solvent and a lithium salt, is known to degrade gradually during the discharge and charge process.$^{16}$ At first, organic carbonate-based solvents were widely used in Li-O$_2$ cells. However, they were decomposing irreversibly during the electrochemical reaction. The
side-reaction products were found to be lithium carbonate and lithium alkyl carbonate.\textsuperscript{17} Thus, dimethyl sulfoxide, as well as ether-based solvents (i.e., dimethyl ether, diglyme, tetraglyme), were investigated, but low round-trip energy efficiency and long-term stability remain an issue with both solvents under the presence of Li\textsubscript{2}O\textsubscript{2}.\textsuperscript{18,19,20}

The most common salts used are LiPF\textsubscript{6}, LiClO\textsubscript{4}, LiN(SO\textsubscript{2}CF\textsubscript{3})\textsubscript{2}, LiNO\textsubscript{3}, and LiBr. The electrochemical performance was found to be affected by the salt concentration and the reactivity of the anion towards lithium peroxide formed during the discharge reaction.\textsuperscript{21,22,23} A practically viable electrolyte system is yet to be found for Li-O\textsubscript{2} batteries, and different pathways are currently explored, either by modifying the solvents with additives\textsuperscript{24} or by using ionic-liquid-based electrolytes\textsuperscript{25,26} or solid state ones.\textsuperscript{27}

Clearly, an optimization of the Li-O\textsubscript{2} system is necessary that requires additional understanding of the electrochemical and chemical reactions on the atomic/molecular level. Here, based on our previous results that soft x-ray emission spectroscopy (XES) can clearly differentiate the four key discharge products of non-aqueous Li-O\textsubscript{2} batteries, namely Li\textsubscript{2}O\textsubscript{2}, Li\textsubscript{2}O, Li\textsubscript{2}CO\textsubscript{3}, and LiOH,\textsuperscript{28} we employ the site specificity at the O and N K edges of XES to probe the decomposition mechanism of the electrolyte and the discharge products during the initial electrochemical cycle of the ionic-liquid-based Li-O\textsubscript{2} system. In this system, metallic lithium is used as anode, PP13(TFSI) [1-Methyl-1-Propylpiperidinium bis (trifluorosulfonylimide] as solvent, LiTFSI (lithium trifluorosulfonylimide) as salt (0.5 M), and a porous carbon electrode as cathode.
2. Materials and Methods

2.1. References and samples

As references, we investigated the gas diffusion layer (GDL) sheet (porous carbon cathode, as received), the PP13(TFSI) solvent, the Li(TFSI) salt (as received), and the electrolyte (PP13(TFSI) + 0.5M LiTFSI) (as prepared). The reference powders were pressed onto carbon tape and mounted on a sample holder. Liquid samples were impregnated on a piece of the GDL before being placed onto carbon tape.

Four samples, two during the discharge and two during the charge reaction, were prepared using the electrochemical cell described below. All parameters for the preparation of the four samples were kept constant, except for the potential at which the reaction was stopped. Potentials of 2.5 and 1.5 V were selected for the discharge reaction and labeled \( V_1 \) and \( V_2 \), respectively. For the charge process, the electrochemical reaction was stopped at \( V_3 = 3.5 \text{ V} \) and \( V_4 = 4.5 \text{ V} \). The potentiostat was programmed to charge or discharge the cell to the defined potential and hold this potential for 24 hours. Thereafter, the cell was disassembled inside an Ar-purged glove box, the cathode removed and mounted without further treatment on a sample holder using carbon tape. To avoid any air exposure, the sample holder was inserted into a transportable vacuum system and sealed under inert atmosphere inside the glove box. The transportable vacuum system was connected to the load lock of the synchrotron endstation and evacuated before transferring the sample holder into the analysis chamber.
2.2. Electrochemical set-up

For the current Li-O$_2$ electrochemical experiments, a lithium metal (Alfa Aesar, purity 99.9%) anode of 12 mm diameter was mounted on a titanium collector. The electrolyte consisted of LiTFSI salt (Alfa Aesar, 99.95% purity) dissolved in PP13(TFSI) ionic liquid solvent (IoLiTec, 99% purity and 48 ppm water). A 3 mm diameter piece of glass fiber separator (Whatman) was impregnated with 150 to 200 μl of the electrolyte and placed on top of the anode. A 7 mm diameter cathode was punched from a Gas Diffusion Layer (GDL) sheet (Freudenberg, H2315/C$_2$ – microporous layer, 90% porosity). The cathode and anode were dried under vacuum for 12 hours before introduction into an Ar-filled glove box (water and oxygen concentrations below 1 ppm).

The cell was first sealed in the Ar-filled glove box and connected outside to a high purity gas line (argon or oxygen) equipped with mass flow controllers for a controlled circulation of the gas through the cathode. The cell was flushed with argon, and then an oxygen flow of 10 sccm was applied for one hour at open circuit voltage (OCV) of 3.3 V. Thereafter, cyclic voltammetry tests were carried out with a SP-150 potentiostat (BioLogic) at room temperature and a scan rate of 10 mV/s. The cyclic voltammograms were acquired by starting at OCV, scanning first to 1.5 V, and subsequently to 4.5 V before returning to the OCV. All potentials in this paper are referenced to the Li$^+$/Li couple.

2.3. XES spectra acquisition
The experiments were performed at the undulator beamline 8.0.1.2 of the Advanced Light Source, Lawrence Berkeley National Laboratory, using the SALSA endstation\textsuperscript{29} equipped with a high-transmission soft x-ray spectrometer.\textsuperscript{30} The pressure during the experiments was $< 1 \times 10^{-8}$ mbar, and the estimated spot size $150 \ (h) \times 30 \ (v) \ \mu m^2$.

XES spectra were acquired with non-resonant excitation at the N and O K edge, using excitation energies of 433.7 and 547.4 eV, respectively. To minimize beam damage, the XES spectra were recorded while constantly scanning the sample with a speed of 50 $\mu m/s$ across an area of $1.2 \times 1.2 \ \text{mm}^2$ (corresponding to an exposure time of $\sim 0.6$ seconds for any given spot on the sample). The reference spectra of the different components involved were normalized to the maximum of the signal. To follow the evolution of the reaction at different potentials, the spectra were normalized to the integrated area of the spectrum.

The excitation energy scale was calibrated at the N and O K edge by measuring an XAS spectrum of N$_2$ gas\textsuperscript{31} and TiO$_2$ (anatase),\textsuperscript{32} respectively. The energy scale of the emission energy was calibrated using the elastically scattered Rayleigh lines.

3. Results and discussion

3.1. Cyclic voltammetry

Figure 1 displays the cyclic voltammetry measurements under argon flow (green, one cycle) and two consecutive cycles under oxygen flow (red). The first scan was run under an inert atmosphere of argon to provide a
background voltammogram. As can be seen, no appreciable current is observed over the full potential range of the oxygen reduction reaction (ORR) and oxygen evolution reaction (OER). The first cycle under oxygen indicates that the ORR starts at a potential of 2.5 V and reaches a maximum current of 80 μA at 1.5 V. On the reverse scan, the OER starts around 3.2 V and is smeared out over a large potential range (up to, at least, 4.5 V). The second oxygen cycle shows a similar redox process, with a decrease of the discharge capacity compared to the first cycle, indicating a partial passivation of the electrode surface or presence of side reaction products with the electrolyte.

3.2. Valence electronic structure of solvent and salt, as seen at the N K edge

We first determine the signature of the electrolyte at the N K edge by analyzing the spectra of the different molecules involved, namely PP13(TFSI), LiTFSI, and PP13. Figure 2 displays the measured nitrogen K XES spectra of the electrolyte, Li(TFSI), and PP13(TFSI). The N K-edge spectrum of LiTFSI represents the local partial density of states (PDOS) of the TFSI anion. The spectrum of the salt is described by a prominent peak at 393.5 eV, two weak features at ~386.4 eV and ~381.7 eV, a low-energy shoulder at ~391.5 eV, and a high-energy shoulder at ~395.6 eV. This is in agreement with the derived N K XES spectrum of TFSI published by Kanai et al.\textsuperscript{33}

The spectrum of the solvent consists of emission from both the PP13 cation and the TFSI anion. It is characterized by a main peak at 393.5 eV
accompanied by a high-energy shoulder at $\sim 395.2$ eV, a low-energy shoulder at $\sim 390.2$ eV, and some weak features at $\sim 383.7$ and $\sim 381.7$ eV. The spectral contribution of the N atom in the PP13 cation can now be obtained by computing the difference between the spectra of PP13(TFSI) and LiTFSI (the latter weighted by a factor of 0.5). As can be seen, the spectral contribution of the PP13 cation is defined by a peak at $\sim 395$ eV [giving rise to the PP13(TFSI) high-energy shoulder at $\sim 395.2$ eV], by a broad peak on the low-energy side centered at $\sim 391.5$ eV, and a weak feature at $\sim 383.7$ eV.

The spectrum of the electrolyte is expected to be mostly a combination of the solvent and salt spectra. Indeed, it mainly consists of a peak related to the TFSI molecule at 393.5 eV and high- and low-energy shoulders related to the PP13 cation (at $\sim 395$ and $\sim 391.5$ eV, respectively). Interestingly, small spectral lineshape enhancements are observed between 381 and 385 eV, indicating that there may be internal molecular interactions in the electrolyte composition.

3.3. Stability of nitrogen site in the solvent and salt molecules

In the previous section, we have shown that the two molecules in the electrolyte feature a specific valence electronic structure at the nitrogen site, which can be probed and distinguished with N K XES. In the following, we use this site specificity to follow the evolution of nitrogen atoms from the salt and the solvent at different stages of the reaction during the first cycle of the cyclic voltammogram. Figure 3 shows the N K XES spectra of the electrolyte, as prepared and as stopped at potentials $V_1$, $V_2$, $V_3$, and $V_4$. 

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The spectra were normalized to the integrated area of the spectrum. As shown in Figure S1, we find that the spectral shapes change slightly, and that the intensity of the prominent TFSI-related peak at 393.5 eV decreases during the course of the first cycle from OCV to $V_4$, indicating a change in the local chemical environment of (some of) the nitrogen atoms. In order to clarify the mechanism, the experimental data at different stages of the reaction were fitted by a linear combination of the two components PP13(TFSI) and LiTFSI. The composition evolution of these components during the discharge and the following charge process is shown in Figure 4. During discharge, the percentage of LiTFSI decreases already at potential $V_1$, from 60% to 18%. During charge, it further decreases, reaching 3% at the end of the charge. This clearly indicates that the nitrogen site of the salt is unstable when applying a voltage different from the OCV. Further nitrogen atoms are gradually removed from the TFSI anion during the electrochemical cycle, indicating, most probably, a breaking of the S-N bond in the anion. The observed decomposition of the anion is very similar to what has been observed with N-methyl-N-propyl-pyrrolidinium TFSA (P13TFSA)$^{34}$ and calculated for N, N-propyl-ethyl-pyrrolidinium (P13TFSI)$^{35}$, where the TFSA and TFSI anions were shown to be less stable against reduction than the P13 cation. The decomposition of the solvent is partially in agreement with the quantum molecular dynamics simulation by Nishino et al.$^{36}$, who conclude that the PP13(TFSI) molecule behaves as a complex without spontaneous dissociation of the PP13(TFSI) ion pair. We cannot currently make conclusions about the remaining fragments, but our results and analysis
unambiguously show that the nitrogen site of the TFSI anion of the salt and solvent is highly unstable under the reduction of oxygen and decomposition of Li$_2$O$_2$.

3.4. Discharge products

During the discharge, it is proposed that only lithium peroxide is formed.$^{2,18}$ However, depending on the electrolyte, it has been shown that other side-products like lithium carbonate, lithium hydroxide, and/or lithium fluoride can also be present.$^{16,19}$ These side-products could be due to decomposition of the electrolyte or a chemical reaction with the carbon cathode. Here, we characterized the discharge products by analyzing the O K XES spectra of the cathode obtained at potential V$_2$. As the cathode was measured ex situ without further treatment, the carbon cathode, the electrolyte, and the discharge products contribute to the oxygen signal at this voltage. Figure 5a displays the O K XES spectra of the as-received carbon cathode (GDL), the as-prepared electrolyte, and the cathode (V$_2$) after being kept at V$_2$ = 1.5 V for 24 hours. As can be seen, the oxygen contribution from the carbon cathode is very small (approx. 1/70) compared to the electrolyte. The O K XES spectrum of the electrolyte is dominated by two main peaks, at ~523 and ~527 eV. The spectrum reflects the local PDOS of the four oxygen atoms of the TFSI molecule. The spectrum of the cathode at potential V$_2$ is very similar to the one of the electrolyte, as displayed in Figure S2, suggesting the formation of very small amounts of O-containing discharge products.
To quantify the different oxygen contributions, the spectrum is fitted with three components, as described in the following. First, we performed a three-component fit, using the spectra of Electrolyte, GDL, and lithium peroxide. The result of the fit and the corresponding magnified residual (x10, labelled V₂-fit residual) is shown in Figure 5b, top. The fit gives a good description of the spectrum with the following spectral weights: 1.5% GDL, 95.0% electrolyte, and 3.5% lithium peroxide. A three-component analysis with GDL, electrolyte, and lithium carbonate or lithium hydroxide alone fails to describe properly the rising shoulder and the highest energy side peak of the O K spectrum at the potential V₂. The respective spectra of Li₂O₂, Li₂CO₃, and LiOH are shown at the bottom of Fig. 5b for comparison. Therefore, we conclude that the dominant discharge product is Li₂O₂ and the spectral weight is similar to the amount expected for the time of reaction from the analysis of the first reduction peak.

Finally, in order to test whether the three-component description can still be significantly improved, the V₂ spectrum was analyzed by a four-component analysis, i.e., using GDL, electrolyte, Li₂O₂, and either Li₂CO₃ or LiOH as fit functions. The better agreement is obtained with LiOH (1.5% GDL, 93.4% electrolyte, 2.9% Li₂O₂, and 2.2% LiOH), as seen from the magnified residual labeled “adding LiOH (x10)”, compared to the case where Li₂CO₃ is added (magnified residual labeled “adding Li₂CO₃ (x10)”).

Therefore, we summarize that the main O-containing discharge product is Li₂O₂, with some lithium hydroxide also being present. However, based on our spectroscopic results, there is no significant amount of lithium carbonate formed during the discharge reaction. The presence of LiOH in
the ionic-liquid-based electrolyte would be in accordance with a recent microscopy study\textsuperscript{37} that shows the presence of LiOH crystallites, the amount of which depending on both the concentration of LiTFSI and the storage time of the discharged electrodes. Therefore, the presence of LiOH is mostly at the origin of the electrodes surface passivation of the electrodes and by that explains the decrease of the discharge capacity observed in the second reaction cycle.

Using XES, each of the compounds shows features of clearly distinguishable spectral shape that will allow \textit{in-situ} and \textit{operando} measurements in the future. As an outlook, Figure 6 presents a schematic view of the novel flow-through electrochemical cell that represents a first step towards \textit{in-situ} and \textit{operando} XES measurements; additional scanning capabilities will be required to avoid the beam-damage effects reported in the manuscript by Léon et al.\textsuperscript{28}

4. Conclusions

In this work, we used the site specificity of XES and a novel electrochemistry cell to probe the electrolyte and discharge products of an ionic-liquid-based Li-O\textsubscript{2} battery. We determined the valence electronic structure of PP13(TFSI), TFSI, and PP13 molecules at the N and O K edges. We observe that the spectral shapes change slightly and the chemical environment of some of the nitrogen atoms is significantly modified during the first discharge/charge cycle, suggesting a decomposition of the salt. The TFSI anion is found to be unstable at the nitrogen atom, while the ionic-liquid solvent appears to remain as an ion pair. Additionally, the O K
edge XES spectra show that lithium peroxide and a small amount of lithium hydroxide are formed as discharge products; however, the formation of lithium carbonate cannot be observed.

There are no conflicts to declare

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Figure captions

Figure 1. Cyclic voltammetry curve of Li/PP13(TFSI) + 0.5M LiTFSI/C (scan rate of 10 mV s⁻¹), using an argon flow (green curve, one cycle), followed by an oxygen flow (red, two cycles). For further ex situ analysis of the cathodes, the electrochemical reaction was stopped at potentials $V_1$ and $V_2$ during the discharge, and at potentials $V_3$ and $V_4$ during the charging of the Li-O₂ cell.

Figure 2. N K XES spectra of the electrolyte (red), the salt (LiTFSI, blue), and the solvent PP13(TFSI) (purple). The spectrum labeled “PP13” shows the weighted difference of the solvent and salt spectra, as calculated by the equation given below the molecular structures.

Figure 3. N K XES spectra (red circles) after stopping at the four potentials described in Figure 1, together with the corresponding two-component fits (solid black lines) using PP13(TFSI) (purple lines) and LiTFSI (blue lines), as well as the corresponding residuals.

Figure 4. Evolution of the relative spectral contribution of the PP13(TFSI) and LiTFSI components to the N K spectra of the electrolyte during the first cycle.

Figure 5: (a) O K XES spectra of the cathode electrode as received (“GDL”), the electrolyte as prepared (“Electrolyte”), and the cathode after discharge at potential $V_2$. (b) O K XES spectrum of the cathode after stopping at potential $V_2$ (red dots) and the corresponding fit (black) using three components (i.e., GDL, electrolyte, and lithium peroxide). Below the fit, three magnified (x10) residuals are shown: for the three-component fit
(“V₂-fit residual”) and for the four-component fits “adding LiOH” and “adding LiCO₃”, respectively. The spectra of the three reference compounds are also shown.

Figure 6. Schematic view of the novel flow-through electrochemical cell, designed for ex-situ, in-situ, and operando XES measurements with the SALSA endstation (SS: stainless steel, PEEK: polyether ether ketone).
Figure 1
Figure 2
Figure 4
Figure 5
TOC Graphic