Title
Electrolyte Engineering to Improve Capacity and Rechargeability in the Lithium-Oxygen Battery

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Electrolyte Engineering to Improve Capacity and Rechargeability in the Lithium-Oxygen Battery

By

Colin M. Burke

A dissertation submitted in partial satisfaction of the requirements for the degree of Doctor of Philosophy in Chemical Engineering in the Graduate Division of the University of California, Berkeley

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in the Lithium-Oxygen Battery

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by

Colin M. Burke
Abstract

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Doctor of Philosophy in Chemical Engineering

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Professor Bryan D. McCloskey, Chair

A primary goal in rechargeable battery research is developing batteries with higher specific energies, with motivations including increasing electric vehicle range and enabling new deep space technologies. One such option, the nonaqueous lithium-oxygen (Li-O₂) battery, consists of a lithium negative electrode, a lithium salt and ether-based electrolyte, an electrolyte-soaked porous carbon positive electrode, and a gaseous oxygen headspace, and operates via the electrochemical formation (discharge) and decomposition (charge) of lithium peroxide (Li₂O₂). With an estimated theoretical specific energy of 3330 Wh/kg active material (Li₂O₂), more than four times that of current lithium-ion positive electrode materials, and a relatively low cost of battery components, the nonaqueous lithium-oxygen (Li-O₂) battery has garnered significant research attention over the past decade.

Unfortunately, critical challenges have been identified that prevent the realization of a high-capacity, rechargeable Li-O₂ battery. The ultimate discharge product, Li₂O₂, is insoluble in the most stable nonaqueous electrolytes and is a wide-band gap insulator, so during discharge it forms as a solid on the cathode’s carbon support and electronically passivates it, preventing further discharge after only a small amount of Li₂O₂ has formed. Li₂O₂ and its electrochemical intermediates also undergo irreversible side reactions with the nonaqueous electrolytes and carbon positive electrodes studied to-date, causing poor battery rechargeability.

In this work, the nonaqueous electrolyte of the Li-O₂ was engineered toward addressing these challenges and achieving a high-capacity, rechargeable Li-O₂ battery. Toward increasing achievable discharge capacity, the ability of electrolytes to induce solubility of the intermediate to Li₂O₂ formation, lithium superoxide (LiO₂), was studied, as this enables a solution mechanism of growth whereby Li₂O₂ grows in large, aggregated structures, allowing more Li₂O₂ to form before cathode passivation. First, the effect of the lithium salt anion on LiO₂ solubility was studied. To do so, a typical lithium battery salt,
lithium bis(trifluoromethane) sulfonimide (LiTFSI), was partially exchanged for the more strongly electron-donating lithium nitrate (LiNO_3) in Li-O_2 battery electrolytes. During galvanostatic conditions, a correlation between LiNO_3 concentration and discharge capacity was observed. Titrations and scanning electron microscopy of cathodes extracted from discharged batteries confirmed Li_2O_2 formation in aggregated structures in cells that partially employed LiNO_3 as an electrolyte, indicative of an increase in the solution mechanism with the addition of LiNO_3. The increase in LiO_2 solubility was attributed via \(^{7}\)Li NMR to a lower free energy of Li\(^+\) in the electrolyte as a result of the addition of the strongly electron donating NO\(^-\) in the lithium solvation shell. Differential electrochemical mass spectrometry (DEMS) showed similar oxygen evolution on charge with and without LiNO\(_3\), indicating no deleterious effect on cell rechargeability with the addition of LiNO\(_3\).

Second, as anion selection induces the solution mechanism by lowering the free energy of Li\(^+\) in solution, non-Li alkali metal cations and alcohols were studied as methods of inducing the solution mechanism by lowering the free energy of the superoxide anion (O\(_2^-\)) in solution. Galvanostatic cycling of Li-O_2 batteries containing non-Li alkali metal salts showed a small increase in the achievable discharge capacity, attributed to the softer acidity of non-Li alkali metal cations more favorably solvating the soft base O\(_2^-\). However, gas analysis of a sodium-oxygen battery with a small amount of Li\(^+\) salt added to the battery electrolyte showed Li\(^+\) quickly scavenges any non-Li alkali metal cation-associated O\(_2^-\), and the resultant LiO_2 quickly disproportionates into the insoluble Li_2O_2. It is therefore anticipated that an increase in Li-O_2 battery capacity upon the addition of non-Li alkali metal cations is only expected at high currents, when oxygen reduction rates are sufficiently high to allow some O\(_2^-\) to temporarily avoid Li\(^+\) in solution. Ppm quantities of methanol, ethanol, and 1-propanol were added to ether-based Li-O_2 battery electrolytes as analogues to water, which has been previously shown to induce the solution mechanism due to its strong Lewis acidity lowering the free energy of O\(_2^-\) in solution. The additives induced a two-fold increase in battery capacity, though with little trend in the capacity as a function of the additive’s Acceptor Number. These results highlight the complexity of interactions between the constituent species in an electrolyte in terms of their Lewis basicities, Lewis acidities, and other physicochemical properties.

While the formation of Li_2O_2 in large aggregated structures increases the achievable discharge capacity, an electrolyte-soluble redox mediator is required to oxidize aggregated Li_2O_2 on charge and shuttle electrons back to the electrode surface. The rechargeability of Li-O_2 batteries containing redox mediators in the presence of water impurities, which are likely difficult to eliminate in practical lithium-air batteries, was studied. Specifically, the effect of water contamination in the electrolyte on the promising redox mediator lithium iodide (LiI) was studied. DEMS and titrations of cathodes extracted from discharged batteries confirmed recent reports that lithium hydroxide (LiOH) formed as the dominant discharge product via a 4 e^{-}/O_2 process. However, isotopic labeling and DEMS were used to show LiOH is not reversibly oxidized back to
its reactants (Li$^+$, O$_2$, H$_2$O). Rather, titrations, scanning electron microscopy (SEM), energy-dispersive X-ray spectroscopy (EDX), and galvanostatic cycling of batteries under an argon atmosphere showed charge current in batteries containing both LiI and H$_2$O is a complex mixture of side reactions and redox shuttling.

With LiOH identified as an undesirable discharge product, the mechanism for Li$_2$O$_2$ degradation to LiOH in the presence of LiI and H$_2$O was studied. Galvanostatic cycling of lab-scale Li-O$_2$ batteries containing LiI and H$_2$O in DME and dimethyl sulfoxide (DMSO) showed that DMSO prevents Li$_2$O$_2$ degradation to LiOH. Cyclic voltammetry of these electrolytes showed DMSO exhibits a higher potential for iodide oxidation than DME, indicating iodide-mediated H$_2$O$_2$ reduction is more difficult in DMSO than DME. The ability of an additive to reduce H$_2$O$_2$ is therefore identified as a key consideration in Li$_2$O$_2$’s stability in the presence of water impurities. A tangential important finding during this study was the difficulty in selection of an appropriate reference electrode for studying redox mediators in Li-O$_2$ batteries, as electrodes with too high of a lithium intercalation voltage will chemically oxidize the redox mediator, while electrodes with too low of a lithium intercalation voltage exhibit spontaneous oxygen consumption in a Li-O$_2$ battery.
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Chapter 1: Introduction to the lithium-oxygen battery

Portions of this chapter are adapted from Ref. 1 with permission from The Royal Society of Chemistry.\(^1\)

1.1 Motivation

The state-of-the-art rechargeable battery for high specific energy and high energy density applications is the lithium-ion battery. Shown in Figure 1.1a, the conventional lithium-ion battery consists of a graphite negative electrode and a lithium transition metal oxide positive electrode separated by an electrolyte of lithium salt in carbonate solvents, and operates via reversible lithium intercalation between the two solid electrodes. As such, the lithium-ion battery’s specific energy – energy stored per mass – and energy density – energy stored per volume – are given by the lithium storage capacity of its electrodes. Commercially available lithium-ion batteries have more than three times the specific energy and energy density of previous rechargeable battery chemistries, such as nickel-metal hydride and silver-zinc, and as a result have revolutionized personal electronics and fully electric vehicles.\(^2\)

![Figure 1.1. a) Schematic of the conventional lithium-ion battery b) Schematic of the conventional lithium-oxygen battery](image)

A current focus in rechargeable battery research is identifying new materials or chemistries that could enable higher specific energy, higher energy density, and lower cost batteries than currently available lithium-ion batteries.\(^2\)\(^-\)\(^3\) These research directions are primarily driven by the understanding that, while still maintaining sufficient lifetime, safety, and power performance, a drastic reduction in battery cost and increase in battery energy density will be necessary to increase market penetration of electric vehicles.\(^4\) Improvements in specific energy, energy density, and cost would be welcomed for other technologies as well, including portable electronic devices, drones, and even space technology. As listed in NASA’s 2015 Space Power and Energy Storage Technology Roadmap, “the primary sub-goal for any space energy storage technology is to provide power at the highest possible specific energy with sufficient durability in the mission environment and, in the case of rechargeable storage, sufficient cycle life.” Of the
possible energy storage technologies, it lists that, “batteries that can safely store very large amounts of energy in small, low-mass packages enable the next generation of deep-space EVA suits that require advanced life support, communications, and computing equipment. All other missions are enhanced by having additional electrical power available without a mass penalty.”

As a result, multiple research directions are now being pursued in hopes of advancing beyond the limits imposed by conventional lithium-ion electrode materials. In the lithium-ion field, new high-energy electrode materials, such as silicon anodes and high voltage cathodes, are being developed. New battery chemistries, including lithium–oxygen, lithium–sulfur, and magnesium–ion, are being explored due to their active cathode materials’ high theoretical specific energies compared to lithium-ion cathode materials. Table 1.1 provides a comparison of the electrochemical performance of lithium sulfide ($\text{Li}_2\text{S}$), lithium peroxide ($\text{Li}_2\text{O}_2$), and various lithium-based metal oxides used as lithium-ion battery cathode materials.

Among the ‘beyond Li-ion’ battery cathode materials, $\text{Li}_2\text{O}_2$, the main discharge product of a nonaqueous lithium–oxygen ($\text{LiO}_2$) battery, has the highest theoretical specific energy. With two electrons extracted per $\text{Li}_2\text{O}_2$ and an approximate average discharge potential of 2.8 V vs. $\text{Li}^+/\text{Li}$, the approximate theoretical specific energy is 3330 Wh/kg active material. This is more than four times that of current lithium-ion positive electrode materials, e.g., 625 Wh/kg active material for $\text{LiCoO}_2$, approximating 0.6 electrons extracted per $\text{LiCoO}_2$ and an average discharge potential of 3.8 V vs. $\text{Li}^+/\text{Li}$. As a result, and in combination with the relatively low cost of the battery’s active materials, the $\text{LiO}_2$ battery has attracted significant research attention over the past decade.

It should be noted that although the values reported in Table 1 provide the impetus for Li–O$_2$ and lithium–sulfur (Li–S) research activity, more sophisticated treatments of realistic practical battery energy densities suggest only modest specific energy improvement, and no volumetric energy density improvement, may be achievable with fully developed Li–O$_2$ batteries over advanced lithium-ion batteries (e.g., those

<table>
<thead>
<tr>
<th>Active cathode material</th>
<th>Approximate reversible Li extraction/insertion (x)</th>
<th>Theoretical specific capacity ($\text{mA h g}^{-1}$)</th>
<th>Average discharge potential (V vs. Li/Li$^+$)</th>
<th>Typical voltage operating window (V vs. Li/Li$^+$)</th>
<th>Approximate theoretical specific energy ($\text{mW h g}^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{Li}_{0.8}\text{CoO}_2$</td>
<td>0.6</td>
<td>164</td>
<td>3.8</td>
<td>4.2-3.7</td>
<td>625</td>
</tr>
<tr>
<td>$\text{Li}<em>{1.0}\text{Mn}</em>{1.0}\text{Ni}_{2.0}\text{O}_4$ (high voltage spined)</td>
<td>1</td>
<td>147</td>
<td>4.7</td>
<td>5.0-3.6</td>
<td>690</td>
</tr>
<tr>
<td>$\text{Li}<em>{1.0}\text{Mn}</em>{1.6}\text{Sn}_{0.4}\text{O}_4$ (high voltage spined)</td>
<td>0.7</td>
<td>195</td>
<td>3.8</td>
<td>4.2-2.0</td>
<td>740</td>
</tr>
<tr>
<td>$\text{Li}<em>{1.0}\text{Mn}</em>{1.6}\text{Sn}_{0.4}\text{O}_4$ (LiNCM)</td>
<td>0.6</td>
<td>166</td>
<td>3.8</td>
<td>4.3-2.0</td>
<td>620</td>
</tr>
<tr>
<td>$\text{Li}<em>{1.2}\text{Mn}</em>{2.0}\text{Ni}<em>{0.5}\text{Co}</em>{2.0}\text{O}_4$ (NGM333)</td>
<td>0.85</td>
<td>267</td>
<td>3.65</td>
<td>4.7-2.0</td>
<td>970</td>
</tr>
<tr>
<td>$\text{Li}<em>{1.4}\text{Mn}</em>{2.0}\text{Ni}<em>{0.5}\text{Co}</em>{2.0}\text{O}_4$ (LiNCM)</td>
<td>2</td>
<td>1168</td>
<td>2.8</td>
<td>4.5-2.0</td>
<td>3330</td>
</tr>
<tr>
<td>$\text{Li}_{0.8}\text{CoO}_2$</td>
<td>2</td>
<td>1168</td>
<td>2.15</td>
<td>2.5-1.5</td>
<td>2510</td>
</tr>
</tbody>
</table>

Table 1.1
Reprinted from Ref. 1 with permission from The Royal Society of Chemistry.  

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using LMR-NMC and Li metal as the electrode materials).\textsuperscript{23} Tempering estimates are engineering considerations including the possible need for onboard storage of \(O_2\) (a closed-system, lithium-oxygen battery) rather than \(O_2\) directly from air (an open-system, lithium-air battery) to avoid water and carbon dioxide impurities that are detrimental to cell performance,\textsuperscript{24-26} as well as the use of a pure lithium metal negative electrode, which hinges on the same engineering of reversible, dendrite-free lithium metal stripping and plating that would enable lithium metal to be used in lithium-ion batteries. Of course, this conclusion is contingent on the successful development of LMR-NMC and other high-energy Li-ion electrode materials, which are currently limited by fundamental challenges that result in voltage and capacity fade during extended cycling.\textsuperscript{16,18} Additionally, it is difficult to estimate some critical parameters for a practical Li-O\(_2\) battery, such as separator thickness, electrolyte volume to positive electrode ratio, and carbon electrode mass and porosity, as we still work to study and address fundamental electrochemical challenges. Thus, while tempered from initial estimates, a modest increase in specific energy via low-cost active materials motivates the study of the lithium-oxygen battery.

1.2 The lithium-oxygen battery

Originally discovered by Abraham et al. and later more fully developed by Bruce et al. and McCloskey et al.,\textsuperscript{27-30} the current conception of the nonaqueous lithium-oxygen battery is depicted in Figure 1.1b. The battery consists of a lithium negative electrode, a lithium salt and ether-based electrolyte, an electrolyte-soaked porous carbon positive electrode, and a gaseous oxygen headspace, and operates via the electrochemical formation and decomposition of lithium peroxide, \(\text{Li}_2\text{O}_2\). Discharging the battery results in lithium oxidation at the lithium anode (reaction 1.1) coupled with oxygen reduction at the positive electrode to form \(\text{Li}_2\text{O}_2\) on the carbon surface (reaction 1.2). Applying a charging current to the battery results in \(\text{Li}_2\text{O}_2\) oxidation to gaseous oxygen at the carbon positive electrode and lithium plating at the lithium negative electrode.

\[
\text{Li}(s) \leftrightarrow \text{Li}^+ + e^- \quad U_0 = 0 \text{ V vs. Li}^+/\text{Li} \quad (1.1)
\]

\[
2\text{Li}^+ + 2e^- + O_2(g) \leftrightarrow \text{Li}_2\text{O}_2(s) \quad U_0 = 2.96 \text{ V vs. Li}^+/\text{Li} \quad (1.2)
\]

The ideal overall reversible cell reaction is therefore:

\[
2\text{Li}(s) + O_2(g) \leftrightarrow \text{Li}_2\text{O}_2(s) \quad U_0 = 2.96 \text{ V vs. Li}^+/\text{Li} \quad (1.3)
\]

\(U_0\) is the half-cell standard equilibrium potential of the reaction. The replacement of a conventional lithium-ion’s transition metal oxide cathode with gaseous oxygen significantly decreases active material cost and gives the lithium-oxygen battery its high theoretical specific energy, motivating its study.

1.3 Challenges

While researchers have found various organic solvents and cathode materials that exhibit the desired \(\text{Li}_2\text{O}_2\) electrochemistry as the major process on both discharge and
charge, even the most stable of these combinations currently result in low attainable discharge capacities, or small specific energies, and poor rechargeability at the cell level. Thus, even before addressing the engineering challenges of O\textsubscript{2} sourcing and lithium metal reversibility as described in the motivation, three more fundamental electrochemical challenges, involving lithium peroxide at the carbon positive electrode, need to be addressed, as shown in Figure 1.2, in order to achieve high-practical specific energy, rechargeable lithium-oxygen batteries.

![Figure 1.2](image)

**Figure 1.2.** Three critical challenges at the positive electrode of the lithium-oxygen battery. Adapated from Ref. 1 with permission from The Royal Society of Chemistry.

First, no organic solvent-based electrolyte studied to-date has shown sufficient stability for a commercially viable battery. Researchers quickly discovered carbonate-based electrolytes like those employed in lithium-ion batteries promote irreversible lithium alkyl carbonate (LiRCO\textsubscript{3}, with R being either Li or a small organic moiety, such as –CH\textsubscript{3} or –C\textsubscript{2}H\textsubscript{5}) formation, but even the most stable solvents, ethers\textsuperscript{29} and amides\textsuperscript{33}, participate in irreversible side reactions in the presence of Li\textsubscript{2}O\textsubscript{2} and/or its electrochemical intermediates.\textsuperscript{1} This decomposition has been shown to form solid, insoluble, and oxidatively strong side products, such as Li\textsubscript{2}CO\textsubscript{3}, at the positive electrode, leading to increasing charge overpotentials and subsequently exacerbated decomposition.\textsuperscript{34-35}

Second, all porous carbons studied to-date, while cheap and high surface area, show decomposition in the presence of Li\textsubscript{2}O\textsubscript{2} in operating cells. McCloskey et al. showed that Li\textsubscript{2}O\textsubscript{2}-induced reactions with a \textsuperscript{13}C carbon cathode led to Li\textsubscript{2}CO\textsubscript{3} formation during...
charge, leading to rising charge overpotentials and exacerbated decomposition. While noble metals and typical oxygen catalysts are more inert to Li$_2$O$_2$, they have been shown to catalyze side reactions, with carbon’s poor dissociation of O$_2$ actually desired to reduce the formation of parasitic side processes.

Third, Li$_2$O$_2$ is a wide-band gap insulator and is insoluble in the most stable solvents, like ethers. Therefore, in these electrolytes, Li$_2$O$_2$ forms as a conformal coating on the porous carbon scaffolding, and, after only a small amount of Li$_2$O$_2$ has formed, i.e., a small discharge capacity has been achieved, the Li$_2$O$_2$ electronically passivates the cathode, preventing further discharge. This results in small practical capacities, and thus a small energy per mass of cell components.

These fundamental electrochemical challenges at the oxygen cathode limit the technology readiness level of the lithium-oxygen battery, as they prevent the realization of even lab-scale cells that have high capacities and long cycle lives, which are necessary before addressing engineering challenges like oxygen purification and storage for scale-up to low-mass battery packs. Thus, there is a need for electrolyte and cathode selection and engineering to achieve a high-capacity, rechargeable lithium-oxygen battery.

1.4 Outline of dissertation

This dissertation details my experimental studies in electrolyte engineering to address these fundamental electrochemical challenges at the positive electrode in the lithium-oxygen battery.

Chapter 2 outlines our experimental setups, with emphasis on considerations necessary in scaling performance behavior from lab-scale batteries.

Chapters 3 and 4 discuss methods for increasing discharge capacity by inducing solubility of lithium superoxide, LiO$_2$, the intermediate to lithium peroxide formation, on discharge. This enables a solution mechanism of growth whereby Li$_2$O$_2$ grows in large, aggregated structures, allowing more Li$_2$O$_2$ to form before cell death. Chapter 3 discusses how the appropriate selection of a lithium salt anion can induce LiO$_2$ solubility and increase discharge capacity without adversely affecting rechargeability. Chapter 4 discusses efforts to broaden this understanding of interactions to other electrolyte additives.

One difficulty with making lithium peroxide in large, aggregated structures on discharge is that Li$_2$O$_2$ – which is the final discharge product, as lithium superoxide, LiO$_2$, is an unstable intermediate – is still an insulator and is insoluble in the electrolytes. As such, Chapters 5 and 6 will discuss work related to redox mediation in the lithium-oxygen battery. Chapter 5 discusses the implications of combining the otherwise promising redox mediator lithium iodide with ppm quantities of water in the otherwise nonaqueous electrolyte, as it causes lithium hydroxide, LiOH, formation. Chapter 6 extends this water contamination work toward an understanding of the interactions leading to LiOH formation, and discusses the importance of selecting an appropriate reference electrode when studying redox mediators.

Chapter 7 contains a summary and is followed by the references for all chapters.
Chapter 2: Characterizing performance of Li-O₂ batteries


2.1 Considerations when characterizing lab-scale lithium-oxygen cells

As discussed in Chapter 1, new electrolytes and electrode materials are needed to solve the lithium-oxygen battery’s fundamental electrochemical challenges. These challenges include electronic passivation of the cathode as well as electrochemical and chemical instabilities of both the electrolyte and carbon electrode in the presence of lithium peroxide and its intermediates.

If a lithium-oxygen battery were built to scale, at an acceptable weight and volume and with acceptable materials, the four main criteria, after safety, to assess whether a new electrolyte or electrode material improves the battery performance would be power, specific energy/energy density, electrical efficiency, and cycling stability. These could all be gleaned from the voltage response of a cell discharging and charging at a constant current, as shown in Figure 2.1.

![Figure 2.1](image)

*Figure 2.1.* Voltage response of a lithium-oxygen cell galvanostatically discharging (bottom line left to right) to a cutoff of 2 V and then charging (top line right to left) to an identical capacity. The cell contains an XC72 carbon positive electrode and 1 M LiTFSI in dimethoxyethane (DME) as the electrolyte. This cell is presented with gas analysis in Figure 3.9.
For a cell discharging at constant current, the power, \( P \), of a lithium-oxygen cell is given by:

\[
P = IV_d \quad (2.1)
\]

where \( V_d \) is the voltage at any given time during discharge, as seen in Figure 2.1, and \( I \) is the constant current being drawn.

The achievable energy, \( E \), from a lithium-oxygen cell is given by:

\[
E = Q_d V_d \quad (2.2)
\]

where \( Q_d \) is the achievable discharge capacity before the voltage drops precipitously, calculated via:

\[
Q_d = It_d \quad (2.3)
\]

where \( I \) is the constant current being drawn for \( t \) time until the voltage decrease. In our work, \( Q_d \) is measured as the capacity at which the cell voltage reaches a 2 V cutoff, as seen in Figure 2.1. In general, a precipitous fall in a battery’s voltage corresponds to the inability of the battery’s desired chemical reaction to support the applied constant current. This inability could be a result of charge transport issues or the full depletion of active materials, including active material transport limitations (e.g., oxygen diffusion). In the lithium-oxygen electrochemistry, the observed precipitous fall in voltage following the desired oxygen reduction reaction would continue until the cathode operating voltage falls out of the electrochemical stability window of the electrolyte (~1 V vs. Li/Li⁺), meaning the electrolyte would decompose reductively if the discharge were not stopped at 2 V. As introduced in Chapter 1, the sudden voltage decline, i.e., sudden exponential increase in discharge overpotential as observed in Figure 2.1, of the lithium-oxygen batteries studied herein is a result of \( \text{Li}_2\text{O}_2 \) forming as a thin film on the carbon electrode surface, which then electronically passivates the electrode and prevents further \( \text{Li}_2\text{O}_2 \) growth.\(^{39}\) The sudden exponential increase in overpotential is ascribed to a critical thickness through which electrons can no longer tunnel – as a resistor model would predict a linear increase in overpotential – and has been confirmed experimentally via measurements of charge transfer as a function of \( \text{Li}_2\text{O}_2 \) thickness via outer-sphere redox shuttles.\(^{32}\) Chapter 3 discusses methods for increasing the achievable discharge capacity in order to increase achievable energy.

The electrical efficiency of a lithium-oxygen cell is given by the difference between the voltages of discharge and charge plateaus. As the area under the discharge curve is the energy supplied by the cell, and the area under the charge curve is the energy required to charge the cell, operating voltages on both discharge and charge as close to the open circuit voltage (~2.85 V) as possible are desired to ensure an electrically efficient battery. As shown in Figure 2.1, the typical Li-O₂ battery does not exhibit a stable voltage plateau on charge, but rather a steadily increasing overpotential. As introduced in Chapter 1, this is a result of oxidatively stable decomposition products like \( \text{Li}_2\text{CO}_3 \) forming on the electrode surface, driving the overpotential up to > 4V in order to
oxidize them.\textsuperscript{34} Hence, there is a need for stable electrolytes and materials that do not degrade to these stable or any other side products.

Cycling stability is given by the quantity and quality (maintenance of power, energy, and electrical efficiency) of discharges and charges that can be achieved before the cell fails, i.e., can no longer sustain a discharge or charge current within the electrochemical stability window of the electrolyte. Figure 2.1 shows a cell’s first cycle, so the voltage response of additional cycles would be compared to Figure 2.1. This provides a measure of the lifetime of the cell.

Academic battery researchers aim to characterize the viability of new electrolytes or materials by testing these criteria in situ in small, unoptimized lab-scale cells and scaling up the findings to realistic, optimized cell configurations with various scaling factors, including electrode surface area, mass loading, and thickness. Unfortunately, these four criteria are not sufficient to characterize new electrolytes and electrode materials in lab-scale lithium-oxygen cells. Numerous undesired side reactions, both electrochemical and chemical, make it difficult to correctly measure power, energy, electrical efficiency and cyclability. For example, due to the reactive nature of the reduced oxygen species, voltage plateaus on discharge or charge exhibited by lithium-oxygen cells may correspond to undesirable reactions. This means that electrochemical signatures that appear to be related to the reversible, desired Li$_2$O$_2$ electrochemistry could actually be partially a result of unwanted side reactions. Additionally, if the discharging and/or charging is actually related to a modest amount of electrolyte and/or cathode degradation, the use of an unrealistically large volume of electrolyte in our lab-scale cells may still allow us to achieve many “cycles”, even though we are just slowly degrading our constituent components over each cycle. In other words, while the voltage response of a lithium-oxygen cell cycling at constant current such as in Figure 2.1 presents some of the challenges inherent the lithium-oxygen electrochemistry, as described above, measuring voltage response alone is not sufficient to fully understand the cell’s electrochemistry when comparing new materials and electrodes.

Consequently, the lithium-oxygen electrochemistry necessitates unique characterization tools to assess new electrolytes and electrode materials. These include a lab-scale battery design that uses a small ratio of electrolyte volume to positive electrode surface area and quantitative accounting for stability on the first few cycles of discharge and charge via pressure monitoring, differential electrochemical mass spectrometry (DEMS), $^1$H nuclear magnetic resonance spectroscopy (NMR), and chemical titrations of discharge products. Characterization techniques like X-ray diffraction (XRD), scanning electron microscopy (SEM), and other spectroscopic techniques can provide useful qualitative information, as well.

Also worth noting, Li$_2$O$_2$ forms on the carbon electrode surface and its thickness and growth morphology depend on numerous factors, including the chemical composition of the electrolyte, the chemical composition of the electrode, electrode porosity and tortuosity, oxygen diffusivity in the electrolyte, actual electrode surface area (as opposed to an electrode’s projected surface area), and current density. Therefore, it is difficult to compare intrinsic power, energy, electrical efficiency, and cyclability from lab to lab. In this work, capacities will simply be listed as mAh, rather than normalizing capacities to the carbon electrode weight or geometric area (mAh/mg or mAh/cm$^2$, respectively). Relevant carbon black composition, carbon black loading, cathode
geometrical surface areas, and current densities will be provided with the measurements described herein.

2.2 Lab-scale cell design

The lithium-oxygen cells used in our laboratory were developed by McCloskey et al. and were used extensively in their lithium-oxygen work. Shown in Figure 2.2, a typical cell consists of a 7/16” diameter lithium foil negative electrode, a 1/2” diameter glass fiber separator soaked with electrolyte, a 12mm diameter positive electrode of high surface area carbon on stainless steel mesh, and 80 µL of an electrolyte, typically a lithium salt or combination of lithium salts in a nonaqueous organic solvent (a common example being lithium bis(trifluoromethanesulfonyl) imide (LiTFSI) in 1,2-dimethoxyethane (DME)). A 1 mm tall, 1/2” OD 0.43” ID stainless steel ring is added to provide additional headspace. These cell parts are compressed between two stainless steel current collectors in a fused silica cylinder. A Swagelok body (not shown in Figure 1) encases and tightens the two current collectors together to a uniform headspace, and a perfluoroelastomer O-ring in each current collector is compressed against the silica housing to provide a hermetic seal. The top current collector is that for the positive electrode and has two soldered 1/16” OD stainless steel capillaries that extend to the bottom face in order to sample gases into and out of the cell. The cells are prepared in an argon glovebox with less than 0.1 ppm water to limit impurities, which have been found to have a large effect on the electrochemistry. The capillaries are capped prior to removal from the glovebox for attachment to the testing systems.

Figure 2.2. A schematic of the hermetically sealed Li-O₂ cell used in our lab. Adapted from Ref. 30.
The typical carbon positive electrodes used in this work were prepared as described previously.\textsuperscript{32} A mixture of 3:1 w:w ratio of Vulcan XC72 to PTFE binder in isopropanol (IPA) and water (4:1 water:IPA; and 15 mL total for 400 mg C) was sonicated for 30 seconds and homogenized for 6 minutes. A Badger model 250 air-sprayer was used to spray the slurry onto a \~4.5” x 4.5” piece of stainless steel mesh, which was rinsed with IPA and acetone and dried at 150°C for ten minutes prior to use. After letting the carbon air-dry, 12 mm diameter cathodes were punched out, rinsed with IPA and acetone, and dried at 150°C under vacuum for at least 12 hours. The cathodes were then transferred, while hot, into the glove box, and stored on a hot plate at 200°C immediately prior to use.

2.3 Gas analysis via cell pressure monitoring and differential electrochemical mass spectrometry (DEMS)

The testing systems used for our Li-O\textsubscript{2} cells are modeled after systems built at IBM.\textsuperscript{30} Two different systems were constructed at UC Berkeley by my colleagues and me (Figure 2.3): one system with pressure monitoring capability only, and one system with both pressure monitoring and differential electrochemical mass spectrometry (DEMS). The pressure monitoring system can test five cells at once, while the DEMS can test two.

![Figure 2.3](image)

Figure 2.3. (Left) Two-channel pressure monitoring and differential electrochemical mass spectrometry (DEMS) battery cell testing system. (Right) Five-channel pressure monitoring battery cell testing system.

For electrical measurements, each cell’s stainless steel current collectors are fastened into brass current collectors. These are connected to a Biologic VMP3 potentiostat allowing us to measure open circuit voltage, galvanostatically discharge and charge, perform cyclic voltammetry, perform electrochemical impedance spectroscopy, and more.

Simultaneously, for gas analysis, the cell capillaries from the cathode current collector are connected to 1/16” OD stainless steel capillaries on the system with Swagelok fittings. Figure 2.4 shows the process flow diagram of one channel, i.e., where one cell can be attached and cycled, as well as the connection of these channels to a vacuum pump, included in both systems, and an additional connection via a leak valve to a vacuum chamber and residual gas analyzer, included in the DEMS.
Opening the inlet valve allows gas to flow into the headspace. Upstream gas cylinders, tubing, and valves enable flowing pure oxygen, argon, isotopically enriched \(^{18,18}\)O\(_2\), carbon dioxide, nitrogen, hydrogen, and other gases as needed. The cell is prefilled with argon from assembly in the glovebox. Opening the purge valve, which is connected to vacuum via more stainless steel capillary tubing, a cross connector, and a typically open (see DEMS discussion below) solenoid valve, would evaporate highly volatile electrolytes, like dimethoxyethane (DME). Thus, to exchange the argon headspace with, for example, oxygen in order to discharge, the purge valve is left closed and the 6-way valve is toggled between position A and position B. In position A, the cell is isolated, and a 500 \(\mu\)L loop (high-precision sample loop for high-performance liquid chromatography) is connected to vacuum and thus evacuated. In position B, the cell is reconnected to the 500 \(\mu\)L loop, and consequently a portion of the cell headspace fills the loop. Toggling to A then evacuates this portion of the cell headspace, and, as the inlet valve remains open, more argon flows in, returning the cell to the line pressure. From calibration, eight cycles are necessary to fully exchange, within detection limits, the headspace of a typical cell with a gas of interest.

Closing both the inlet and purge valves with the 6-way valve in position B allows pressure monitoring of the cell via the in-line pressure transducer (P1 in Figure 2.4). This enables pressure decay during discharge, as oxygen is consumed, and pressure rise during charge, as \(\text{O}_2\), \(\text{CO}_2\), \(\text{H}_2\), and other gases are evolved.

In order to quantify the moles of gas consumed and evolved, the volumes of each testing channel and the volume of our cell were calibrated via an ideal gas approximation. First, we attached a known volume loop (e.g., another 500 \(\mu\)L high-precision sample loop for high-performance liquid chromatography) in place of the cell with the channel in position B. Second, we opened both the solenoid valve and the purge valve in order to evacuate both the cross and the channel. Third, we closed the purge valve and filled the channel with argon. After recording the pressure, we then closed the solenoid valve and then opened the purge valve to expand that gas into the full volume of the channel (in position B, everything between the inlet and purge valves, minus the cell) plus the known loop plus the cross (in position B everything between the purge valve and the solenoid valve). In the case of the DEMS, the leak valve was closed. Approximating the gas as ideal, the following equation then applies:

\[
P_{\text{initial}}(V_{\text{channel}} + V_{\text{loop}}) = P_{\text{final}}(V_{\text{channel}} + V_{\text{loop}} + V_{\text{cross}})
\]

where \(P_{\text{initial}}\) is the pressure measured before volume expansion, \(P_{\text{final}}\) is the pressure measured after volume expansion, and \(V_{\text{channel}}\), \(V_{\text{loop}}\), and \(V_{\text{cross}}\) are the volumes of those sections as described above. By measuring a set of loops of various known volumes (as well as a set of various initial pressures for statistical fidelity), we calculated both the channel volume and the cross volume. Knowing these values, we replaced the known volume loop with a dry cell, i.e., a cell as described above but with no electrolyte, and performed a similar volume expansion to calculate the volume of the cell. We then subtracted the electrolyte volume (e.g., 80 \(\mu\)L), and added the dry cell volume and channel volume to obtain the volume in position B between the inlet and purge for a typical cell. This is the volume then used to calculate the moles of gas consumed or evolved, approximating the gas as ideal, as we are typically operating at ambient
temperature and 0-1.5 atm pressure. Cells are discharged and charged with both the inlet and purge valves closed and the 6-way valve in position B. Knowing the cell and channel volumes and the ambient temperature via a nearby thermocouple, the pressure change measured by the in-line pressure transducer, P1 in Figure 2.4, can be converted to a change in moles of gas.

Electrolyte and cathode decomposition cause deviation from the ideal lithium-oxygen electrochemistry, as the constituent substances of lithium, oxygen, electrons, and/or lithium peroxide participate in unwanted side reactions with the electrolyte and cathode to form irreversible side products. As the ideal electrochemistry is two electrons per oxygen molecule (e⁻/O₂), any deviation from that value points toward electrolyte or cathode instabilities, i.e., poor rechargeability. Pressure monitoring allows a direct and real-time e⁻/gas measurement.

To directly measure e⁻/O₂ and to identify and quantify other gases evolved, we use the setup in Figure 2.4 in differential electrochemical mass spectrometry (DEMS) mode.

**Figure 2.4.** Process flow diagram of the pressure monitoring system and differential electrochemical mass spectrometer (DEMS) system. Not shown for both systems is the set of gas cylinders and selector valves to flow in a desired gas (argon, O₂, ^18^18O₂, CO₂, H₂, N₂, etc.). Shown is one cell’s channel; additional channel(s) are connected to the cross connector as labeled. The right arm of the cross on the DEMS system is connected to a leak valve to a vacuum chamber with a residual gas analyzer, whereas the left arm of the cross on the pressure monitoring system is connected to additional channels.
For this mode, a leak valve connects the right arm of the cross connector to a vacuum chamber containing a residual gas analyzer. Before charging, the cell headspace is exchanged for pure argon. Throughout charging, the 6-way valve is electronically actuated to pulse from A to B to A every couple minutes, sending gas from the cell headspace – predominantly argon, with small quantities of gases like oxygen, carbon dioxide, and hydrogen being evolved from the cell – via the 500 µL loop to the cross connector. Meanwhile, the solenoid valve (S in Figure 2.4) is electronically closed. The gas from the cell headspace therefore sits in the cross and leaks slowly into the chamber with the residual gas analyzer. This allows us to identify the gases being evolved from the cell.

To quantify the moles of gases evolved by the cell as identified by the residual gas analyzer, we prepared mixtures of gases of interest (for the Li-O₂ battery, typically oxygen, hydrogen, and carbon dioxide) at small concentrations in argon, approximating the gases as ideal and using pressure to calculate the concentrations. We then leaked these gases to the residual gas analyzer (RGA) and measured their ratios to the argon signal. We thus had a calibration for an actual ratio of oxygen to argon vs. the mass spectrometer’s measured ratio of oxygen to argon. (Of note, we first measured the 36:40 signal ratio of our pure argon, and from then on used the 36 peak to extrapolate an argon 40 peak, as the argon 40 signal, the dominant isotope of our dominant gas, would overwhelm our RGA.) Before charging a cell, while the cell is sitting at open circuit, we pulse the cell sufficiently – usually a minimum of eight times – to establish a reliable baseline for all mass signals of interest (typically 32 for O₂, 44 for CO₂, and 2 for H₂). The baseline was then subtracted from the values measured during charging. This corrected signal for each gas was then compared to the argon mass signal, and converted to an estimated actual ratio via the calibration. Knowing the volume of the 500 µL loop, the pressure of the gas in the 500 µL loop when it is briefly in position B via the pressure transducer P1, and the ambient temperature via a nearby thermocouple, we are able to calculate the total moles of gas in the 500 µL loop. Using ratios of the gases to argon via the mass spec and calibrations, we can then calculate the molar composition of the gas in the 500 µL loop. After charging has been completed, the cell is pulsed at least eight times to remove all of the evolved gases from the cell headspace, and usually eight times to reestablish and confirm the baseline. Throughout the DEMS process, the inlet valve is left open to a supply of argon gas to resupply the cell as gas is pulsed out. Of note, ¹⁸O₂ gives the same signal as ¹⁸Ar. As a result, the baseline 36 signal was used to extrapolate the argon 40, and then during charging, the 36 signal was used, minus the baseline. Additionally, each cell run on the DEMS is run alongside an identical cell operating under pressure rise, i.e., with both inlet and purge valves closed and the 6-way valve in position B. This is performed to ensure the total moles of gas evolved calculated from the DEMS matches the total moles of gas calculated via pressure rise.

By comparing the coulometry from the potentiostat with this regular gas quantification, e⁻/O₂ values can be determined throughout a battery’s discharge or charge. Additionally, the exact amount of oxygen evolved on charge (OER) can be compared with the amount consumed on discharge (ORR), giving another metric of rechargeability, as OER/ORR should approach unity. These values give important insight into the battery’s rechargeability. For example, in a discharging battery, an e⁻/O₂ value of 2.00 indicates the near perfect formation of Li₂O₂ on discharge (reaction 2.6). A subsequent
value of 2.50 e'/O$_2$ on charge indicates 20% of electrons are going into a non-oxygen evolving reaction. Therefore, DEMS allows us to quantify the rechargeability of each cell during cell cycling.

Based on error propagation of volume calibrations and DEMS gas calibrations, typical e'/O$_2$ values are within +/- 0.01 e'/O$_2$. Cells containing significant water quantities formed hydrogen at open circuit via reaction with the lithium anode, and correcting the pressure decay on discharge for this pressure rise at open circuit typically led to a reduction in precision to +/- 0.1 e'/O$_2$.

Of note, as described in Chapters 5 and 6 in further detail, e'/O$_2$ can be a misleading metric for Coulombic efficiency in some instances. Chapters 5 and 6 describe the 4 e'/O$_2$ formation of LiOH. This implies the four electron reduction reaction:

$$4\text{Li}^+ + 4\text{e}^- + \text{O}_2 + 2\text{H}_2\text{O} \rightarrow 4\text{LiOH} \quad (2.5)$$

However, as shown by Tulodziecki et al., LiOH is likely formed via the two electron formation of Li$_2$O$_2$, followed by the chemical reactions of Li$_2$O$_2$ with water to form LiOH and the consumption of H$_2$O via iodide, followed finally by the reduction of triiodide back to iodide:

$$2\text{Li}^+ + 2\text{e}^- + \text{O}_2 \rightarrow \text{Li}_2\text{O}_2 \quad (2.6)$$

$$\text{Li}_2\text{O}_2 + 2\text{H}_2\text{O} \rightarrow 2\text{LiOH} + \text{H}_2\text{O}_2 \quad (2.7)$$

$$\text{H}_2\text{O}_2 + 3\text{I}^- + 2\text{Li}^+ \rightarrow 2\text{LiOH} + \text{I}_3^- \quad (2.8)$$

$$\text{I}_3^- + 2\text{e}^- \rightarrow 3\text{I}^- \quad (2.9)$$

Overall, these electrochemical and chemical reactions sum to reaction 2.5, but the electrochemical step is still a 2 e'/O$_2$ process, rather than a typical 4 e'/O$_2$ reduction process such as typically observed on platinum electrodes. In other words, the presence of other compounds within a Li-O$_2$ battery can change the observed e'/O$_2$ values, as well as the ultimate products formed, even if the electrochemical step is still 2 e'/O$_2$.

### 2.4 Discharge product identification and quantification via chemical titrations

Other quantitative measures of rechargeability in the Li-O$_2$ battery are chemical titrations for the amount of Li$_2$O$_2$ on carbon cathodes extracted from cells at various states of discharge and charge. The titration protocol used here follows that developed previously.$^{32}$

To prepare cathodes for titration, immediately upon completion of the desired discharge and/or charge, cell headspaces are replaced with argon and the cells are capped and brought into an argon glovebox. The cathode and separator are extracted from each battery and placed in a 20 mL septum vial. Vacuum is pulled on this vial for roughly 3 minutes to remove residual electrolyte before tightening the septum and removing the vial from the glovebox for titration of the contents.
The titrations begin with the injection of 2 mL of ultrapure water (18.2 MΩ cm, Millipore) through the septum to react Li₂O₂ into LiOH and H₂O₂.

\[
\text{Li}_2\text{O}_2 + 2\text{H}_2\text{O} \rightarrow 2\text{LiOH} + \text{H}_2\text{O}_2 \quad (2.7)
\]

The protocol then proceeds with two chemical titrations. First, an acid/base titration is performed to quantify LiOH and any basic decomposition products present in the cell. Two drops of phenolphthalein are added as an indicator, and a standardized solution of 0.005 M hydrochloric acid is used as a titrant, with a pink to clear transition at the endpoint.

\[
\text{LiOH} + \text{HCl} \rightarrow \text{LiCl} + \text{H}_2\text{O} \quad (2.10)
\]

Second, an iodometric titration is immediately performed to quantify H₂O₂. An aqueous solution of 0.01 M potassium iodide and 3.5 M sulfuric acid are added in excess (typically 1mL each) along with a drop of molybdate-based catalyst solution to form I₂ in aqueous solution.

\[
\text{H}_2\text{O}_2 + 2\text{KI} + \text{H}_2\text{SO}_4 \rightarrow \text{I}_2 + \text{K}_2\text{SO}_4 + 2\text{H}_2\text{O} \quad (2.11)
\]

The molybdate-based catalyst solution is an aqueous solution of ammonium molybdate, ammonium hydroxide, and ammonium nitrate. Standardized 0.01 M sodium thiosulfate is then used as a titrant for I₂, with a yellow to clear transition at the endpoint.

\[
\text{I}_2 + 2\text{Na}_2\text{S}_2\text{O}_3 \rightarrow \text{Na}_2\text{S}_4\text{O}_6 + 2\text{NaI} \quad (2.12)
\]

An aqueous starch solution is typically added near the endpoint to complex with the remaining I₂ to create a more apparent dark blue to clear transition at the endpoint.

On cycled cathodes, the acid/base titration picks up basic decomposition products in addition to the LiOH formed from the reaction of water and Li₂O₂ in the septum vial, so the iodometric titration is typically taken for the Li₂O₂ yield. The Li₂O₂ percent yield is typically defined as the amount of Li₂O₂ formed during discharge, as quantified via an iodometric titration, to the amount of Li₂O₂ expected from coulometry, assuming an ideal 2e⁻/Li₂O₂ process. Controls of pure commercial Li₂O₂ powder were performed to verify the titrations’ accuracy.

A comparison of Li₂O₂ yields to e⁻/O₂ values for cells at various states of discharge and charge gives important insight into the formation, subsequent side reactions, and oxidation of lithium peroxide. For example, a near perfect 2.00 e⁻/O₂ value on discharge coupled with a less ideal 80% Li₂O₂ yield after discharge indicates the dominant formation of Li₂O₂ on discharge followed by chemical degradation of Li₂O₂ to other products, such as lithium formate (HCO₂Li) and lithium fluoride (LiF) via electrolyte solvent and salt decomposition.

2.5 Qualitative spectroscopic characterization techniques

Ex situ spectroscopic characterization techniques can provide useful information about lithium-oxygen cell performance as well. Techniques like X-ray diffraction
(XRD), Raman spectroscopy, X-ray photoelectron spectroscopy (XPS), and Fourier-transform infrared spectroscopy (FTIR) can be used to identify discharge products and decomposition products. Of note, however, these characterization techniques are all locally quantitative at best, i.e., they characterize random spots on a lab-scale electrode, rather than macroscopic performance of a battery material. They are useful in identifying possible reaction products and decomposition pathways, but gas analysis and chemical titrations should be used as the quantitative measurements for cell rechargeability. Intricacies can complicate their use as well, such as amorphous Li$_2$O$_2$ not visible via XRD, and decomposition of polyvinylidene fluoride (PVDF) binder in the carbon cathode showing similar peaks to lithium superoxide (LiO$_2$) in Raman spectroscopy.

Scanning electron microscopy (SEM) is useful to investigate changes in Li$_2$O$_2$ morphology. Energy-dispersive X-ray spectroscopy (SEM EDX) can also be used to identify discharge products.

$^1$H nuclear magnetic resonance spectroscopy (NMR) can be used to determine decomposition products in the electrolyte, as well as the pKa of ppm levels of water in different nonaqueous electrolytes. $^7$Li NMR can be used to determine the lithium environment in an electrolyte, as discussed in Chapter 3. NMR procedures for this work are detailed in Chapter 3, Chapter 4, and Chapter 6.
Chapter 3: Influence of electrolyte anion on Li-O_2 discharge capacity


Abstract
Among the ‘beyond Li-ion’ battery chemistries, nonaqueous Li-O_2 batteries have the highest theoretical specific energy and as a result have attracted significant research attention over the past decade. A critical scientific challenge facing nonaqueous Li-O_2 batteries is the electronically insulating nature of the primary discharge product, lithium peroxide, which passivates the battery cathode as it is formed, leading to low ultimate cell capacities. Recently, strategies to enhance solubility to circumvent this issue have been reported, but rely upon electrolyte formulations that further decrease the overall electrochemical stability of the system, thereby deleteriously affecting battery rechargeability. In this study, we report that a significant enhancement (greater than four-fold) in Li-O_2 cell capacity is possible by appropriately selecting the salt anion in the electrolyte solution. Using $^7$Li nuclear magnetic resonance, we confirm that this improvement is a result of enhanced Li$^+$ stability in solution, which in turn induces solubility of the intermediate to Li$_2$O$_2$ formation. Using this strategy, the challenging task of identifying an electrolyte solvent that possesses the anti-correlated properties of high intermediate solubility and solvent stability is alleviated, potentially providing a pathway to develop an electrolyte that affords both high capacity and rechargeability. To confirm the experimental results, an Ising model was developed with collaborators to predict Li$^+$ stability as a function of its solvation shell composition. We believe the model and strategy presented here will be generally useful to enhance Coulombic efficiency in many electrochemical systems (e.g., lithium-sulfur batteries) where improving intermediate stability in solution could induce desired mechanisms of product formation.

3.1 Introduction
The lithium-oxygen (Li-O_2) battery has garnered significant research interest in the past ten years due to its high theoretical specific energy (~3500 Wh/kg$_{Li_2O_2}$) compared to current state-of-the-art lithium-ion (Li-ion) batteries (~625 Wh/kg$_{LiCoO_2}$). Consisting of a lithium anode and an oxygen cathode, the nonaqueous Li-O_2 battery operates via the electrochemical formation and decomposition of lithium peroxide (Li$_2$O$_2$). The ideal overall reversible cell reaction is therefore:

$$2Li + O_2 \leftrightarrow Li_2O_2 \quad (2e^- \text{ process}) \quad U_0 = 2.96 \text{ V} \quad (3.1)$$

One challenge preventing the realization of the Li-O_2 battery’s high theoretical specific energy is that the discharge product, Li$_2$O$_2$, which is generally insoluble in aprotic organic electrolytes, is an insulator. Thus, on discharge, Li$_2$O$_2$ forms as a conformal coating on the cathode’s carbon support, as shown in Figure 3.1a. As such, it electronically passivates the cathode, resulting in practical capacities much smaller than
theoretically attainable.\textsuperscript{50} For a cell discharging at constant current, this is exhibited by a sudden exponential drop in voltage – or exponential increase in overpotential – as shown in Figure 2.1 and Figure 3.2, and is ascribed to a critical thickness of Li\textsubscript{2}O\textsubscript{2} through which electrons can no longer tunnel.\textsuperscript{39}

![Figure 3.1](image)

**Figure 3.1.** Possible discharge mechanisms in the nonaqueous lithium-oxygen battery. The thin film mechanism is dominant in typical ether-based electrolytes. The solution mechanism is induced by increasing solubility of the lithium superoxide intermediate in solution.

Recently, two reports have described the engineering of electrolytes to circumvent this passivation and improve Li–O\textsubscript{2} battery discharge capacity. Aetukuri et al. suggests that adding ppm quantities of water to a DME-based electrolyte increases the solubility of intermediates during Li\textsubscript{2}O\textsubscript{2} formation.\textsuperscript{25} This increased solubility allows a reduced oxygen species shuttling mechanism that promotes deposition of Li\textsubscript{2}O\textsubscript{2} aggregated toroid structures, as shown in Figure 3.1b. Of note, these structures are typically platelet-shaped, but are often referred in the literature as toroids, so we will use both terms interchangeably. The diffusion of the intermediates away from the electrode surface allows the surface to remain electronically accessible to Li\textsuperscript{+} and O\textsubscript{2}, promoting more Li\textsubscript{2}O\textsubscript{2} growth before the surface film meets a critical thickness, thereby leading to an increase in cell capacity. This increase in cell capacity with water content in the electrolyte is also consistent with reports by Gasteiger and coworkers.\textsuperscript{51-52} Aetukuri et al. reasons that this increase could be attributed to water’s significantly higher Gutmann Acceptor Number (AN) than DME, as the AN is a measure of a solvent’s Lewis acidity, and thus quantifies its ability to efficiently solubilize negatively charged species, such as the potential discharge product intermediate, superoxide (O\textsubscript{2}\textsuperscript{-}).\textsuperscript{53} Johnson et al. presents a related analysis, showing that an electrolyte solvent with a higher Gutmann Donor Number (DN), a measure of Lewis basicity,\textsuperscript{54} is more likely to induce toroid formation due to increased Li\textsuperscript{+} solvation efficiency, allowing solubility of O\textsubscript{2}\textsuperscript{-}.\textsuperscript{45} Johnson et al. further confirmed the presence of O\textsubscript{2}\textsuperscript{-} ions in Li\textsuperscript{+}-bearing high DN solvents using surface enhanced Raman spectroscopy, with others also confirming soluble O\textsubscript{2}\textsuperscript{-} formation.\textsuperscript{45, 55-57}

While water and certain organic solvents increase cell capacity via this solution mechanism, there is evidence that both decrease electrolyte stability. Water impurities in Li-ion electrolytes are known to enhance parasitic electrochemical side reactions, and Aetukuri et al. and Cho et al. showed that adding ppm quantities of water in Li–O\textsubscript{2} batteries leads to a decrease in electrolyte stability and irreversible reactions with the
Furthermore, using quantitative measures of battery rechargeability, high DN solvents, such as DMSO and N-methyl pyrrolidone, have been observed to be less stable than low DN solvents, such as acetonitrile and DME. Recently, Khetan et al. showed using thermodynamic analysis that an organic solvent’s ability to induce the solution mechanism is anti-correlated with its stability towards nucleophilic attack. Thus, Li-O₂ cells would benefit from an appropriately engineered electrolyte that both induces Li₂O₂ intermediate solubility and maintains or exceeds present electrolyte stability.

In this report, we describe the importance of the lithium salt anion in enhancing the solvation of electrochemically-formed intermediate species during Li-O₂ battery discharge, thereby enhancing discharge capacity. We present a study on two common Li-O₂ battery salts, lithium bis(trifluoromethane) sulfonimide (LiTFSI) and lithium nitrate (LiNO₃), dissolved in 1,2-dimethoxyethane (DME). These salts were selected because Schmeisser et al. found that TFSI⁻ and NO₃⁻ anions provided different DN in ionic liquids with common cations (NO₃⁻-containing ILs having higher DN than TFSI⁻-containing ILs). We also specifically selected NO₃⁻ because of its reported positive influence on Li-O₂ battery rechargeability compared to the more commonly used TFSI⁻. We found that electrolytes containing a high concentration of NO₃⁻ exhibited higher donicity, as verified using ⁷Li NMR, and provided an increase in battery capacity greater than four-fold compared to a battery employing a TFSI⁻ electrolyte, while not decreasing battery rechargeability, as measured using quantitative oxygen consumption and evolution. In order to theoretically quantify this enhancement, collaborators at Carnegie Mellon University and RWTH Aachen University developed an Ising model to describe the solvation shell of Li⁺. This analysis shows the origin of this enhanced solution process is due to the formation of ion pairs (Li⁺-NO₃⁻) in a DME solvent. The theoretical analysis further predicted that ion-pair formation and the associated enhancement in capacity would not be observed when DMSO is used as a solvent, which was then confirmed experimentally. We generalize this analysis to provide a rational basis for selection of electrolyte (solvent + salt) combinations for use in Li-O₂ batteries. We believe these results will have profound implications not only for Li-O₂ batteries, where a practical outcome of the solubility is an enhancement in battery capacity, but for other electrochemical systems (e.g., lithium-sulfur batteries) in which intermediate solvation may induce desired mechanisms of product formation.

3.2 Experimental methods

**Materials.** Lithium nitrate (BioUltra) and lithium bromide (ReagentPlus) were purchased from Sigma-Aldrich and were dried under vacuum in a heated glove box antechamber at 150⁰C for 24 hours before use. Lithium bis(trifluoromethane) sulfonimide (LiTFSI), 1,2-dimethoxyethane (DME), and dimethyl sulfoxide (DMSO) were purchased from BASF and used as received. Whatman QM-A glass fiber filters were purchased from VWR. Poly(tetrafluoroethylene) (PTFE, 60 wt% dispersion in H₂O) was purchased from Sigma-Aldrich. Vulcan XC72 was purchased from Fuel Cell Store and was filtered through a 60-mesh screen prior to use. T316 stainless steel 120 mesh, with wire diameter 0.0026”, was purchased from TWP Inc. Research-grade oxygen and argon were purchased from Praxair. 99% ¹⁸O₂ was purchased from Sigma-Aldrich. All electrolyte,
cell, and NMR sample preparation was completed in an argon-filled glove box with <0.1 ppm O₂ and <0.1 ppm H₂O. Lithium foil with a thickness of 0.01” was purchased from FMC Lithium. 99% Lithium chloride (BioUltra) was purchased from Sigma-Aldrich. 99.9% deuterium oxide (D₂O) was purchased from Cambridge Isotope Laboratories, Inc.

Cathode preparation. Cathodes of XC72 carbon on stainless steel mesh were prepared via the spray-coating method described in Chapter 2.

Carbon loading. Cell capacities depend on carbon loading. To keep carbon loading consistent, the cathodes used for any particular data set, such as the capacity measurements displayed in Figure 3.2 or the cathode morphology images in Figure 3.3, were all from the same batch of spray-coated cathodes. As a control, the capacity measurements presented in Figure 3.2 were repeated with a second batch of cathodes. All capacities changed proportionally (a slight increase for all cases), with the capacity of the cell using 0.5M LiNO₃/0.5M LiTFSI in DME maintaining just over a three-fold increase from the capacity of the cell employing 1M LiTFSI in DME. Cathodes contained on average 1.5-2 mg carbon/cm² electrode projected area.

Cell preparation. The LiO₂ cells used followed the same Swagelok design as described previously and described in detail in Chapter 2. All cells employed a 7/16” diameter lithium foil, a 1/2” diameter Whatman QM-A glass fiber separator, a 12mm diameter cathode of Vulcan XC72 carbon on stainless steel mesh, and a 1mm thick, 1/2” diameter stainless steel ring. The QM-A separators, like the cathodes, were rinsed with IPA and acetone and dried under vacuum at 150°C under vacuum for at least 12 hours before being transferred to the glove box and stored on a hot plate at 200°C. Each battery contained 80 µL of electrolyte.

Scanning electron microscopy. Discharged cathodes were characterized via scanning electron microscopy immediately after discharge. After replacing discharged cells’ headspaces with argon, the cells were transferred into the glove box, and the cathodes were removed. The cathodes were each rinsed with two 1 mL aliquots of DME and were subsequently dried under vacuum for at least five minutes in the glove box antechamber. The cathodes were then sealed in septa vials, removed from the glove box, and taken to the SEM. Immediately before imaging, the cathodes were removed from the argon-filled septa vials, placed on carbon tape on the SEM holder, and inserted into the SEM. From discharge completion to SEM insertion was typically one hour. From removing the cathodes from the septa vials to insertion into the SEM was typically less than 30 seconds. SEM was performed on a JEOL JSM-7500f.

Titration. The Li₂O₂ titration protocol used here followed that described previously and described in detail in Chapter 2. Of note, no titratable I₂ was observed from titrations on the separator alone, confirming the following two points: a) no NO₂⁻ that may have formed at the Li metal anode is present in the separator, and therefore NO₂⁻ does not result in a falsely higher Li₂O₂ yield as measured using the iodometric titration (NO₂⁻ also oxidizes I⁻); b) Li₂O₂ only forms on the cathode and not the separator.

Nuclear magnetic resonance spectroscopy. ⁷Li and ²³Na nuclear magnetic resonance spectroscopy measurements were completed on a Bruker AM-400 magnet with a 5mm Z-gradient broad band probe. Reference samples, those employing a chloride salt in D₂O, were prepared outside the glove box and were flame-sealed in melting point capillaries. Electrolyte samples were prepared inside the glove box and were placed, along with a reference capillary, in a Wilmad screw-cap NMR tube. All reference
samples were 3M of the chloride salt in D₂O. For the ²³Na NMR, 0.2M NaClO₄ was added. These molarities were taken from Schmeisser et al.⁶⁰ ²³Na NMR spectra are not reported here because of the poor solubility of NaNO₃ in DME. NMR spectra of only the reference and only the sample were taken to verify the identity of each peak.

3.3 Results and discussion

To characterize the effects of the electrolyte salt anion on discharge performance, Li-O₂ cells were prepared with electrolytes of varying concentrations of LiNO₃ and LiTFSI salts, totaling 1.0M Li⁺, in DME.

Figure 3.2a presents representative galvanostatic discharge profiles of these Li-O₂ cells as a function of the LiNO₃ salt concentration. The inset in Figure 3.2a shows the average cell capacity for each LiNO₃ concentration. Cell capacity increases more than four-fold over the LiNO₃ concentration range studied, clearly indicating the substantial effect of the Li⁺ counterion on cell capacity.

*Figure 3.2. a) Representative galvanostatic discharge profiles of Li-O₂ cells (450 μA/cm² under a 1.5 atm O₂ atmosphere to a 2V cutoff). Inset shows capacity dependence on LiNO₃ concentration. b) Li₂O₂ discharge yield as a function of LiNO₃ concentration. c) ⁷Li chemical shift of sample electrolyte solutions, versus a 3 M LiCl in D₂O standard, as a function of electrolyte LiNO₃ concentration. A less negative chemical shift represents a shift downfield. A 1.0M Li⁺ concentration was used for all electrolytes (DME used as the solvent), while the LiTFSI/LiNO₃ ratio was varied. The LiNO₃ concentration for each cell is provided in the figure. As an example, the cell labeled ‘0.1M LiNO₃’ contained 0.1M LiNO₃ and 0.9M LiTFSI. Error bars are one standard deviation of multiple experiments.*
Scanning electron microscopy (SEM) was performed on discharged cathodes to investigate changes in Li$_2$O$_2$ morphology, and hence changes in discharge mechanism, with increased LiNO$_3$ concentration. Figure 3.3 presents SEM images of a pristine cathode (Figure 3.3a) and images of cathodes from cells of identical electrolyte compositions as those studied in Figure 3.2, but discharged at 45 µA/cm$^2$ (Figure 3.3b-f). When comparing Figure 3.3a-c, the pristine, 0M LiNO$_3$, and 0.01M LiNO$_3$ cathodes appear indistinguishable. This implies a conformal coating of discharge product on the 0M LiNO$_3$ and 0.01M LiNO$_3$ cathodes, and is consistent with previous reports for 1M LiTFSI in DME.$^{25,37}$

Figure 3.3. a) Scanning electron microscope (SEM) image of pristine XC72 carbon cathode prior to discharge. b-f) Discharged cathodes in cells employing 1M total Li$^+$ concentration, with 0M LiNO$_3$ (1M LiTFSI), 0.01M LiNO$_3$, 0.1M LiNO$_3$, 0.5M LiNO$_3$, 0.7M LiNO$_3$, respectively. Cells were discharged at 45 µA/cm$^2$ to 0.9 mAh/cm$^2$ or a 2 V cutoff voltage, but all cells had at least 0.5 mAh/cm$^2$ capacity. Scale bars are 1µm.
A conformal coating of discharge product is indicative of a predominant thin film Li$_2$O$_2$ surface deposition mechanism. Originally outlined by Laoire et al., this mechanism is described by the following elementary steps $^{63-64}$:

$$\text{Li}^+ + e^- + O_2^* \leftrightarrow \text{LiO}_2^*$$  \hspace{1cm} (3.2)

$$\text{Li}^+ + e^- + \text{LiO}_2^* \leftrightarrow \text{Li}_2\text{O}_2^*$$  \hspace{1cm} (3.3a)

and/or

$$2\text{LiO}_2^* \leftrightarrow \text{Li}_2\text{O}_2^* + O_2$$  \hspace{1cm} (3.3b)

where ‘*’ denotes a species adsorbed to the cathode/Li$_2$O$_2$ surface. Importantly, in the DME/LiTFSI electrolyte, lithium superoxide (LiO$_2^*$) is insoluble and therefore remains adsorbed to the electrode surface, where a second charge transfer step (reaction 3.3a) or a disproportionation reaction (reaction 3.3b) results in the conformal Li$_2$O$_2$ coating observed in Figure 3.3b and c. $^9,37,63-66$

As the LiNO$_3$/LiTFSI ratio increases, the discharge morphology perceptibly changes. As seen in Figure 3.3d, when using 0.1M LiNO$_3$, nodular morphologies appear on the cathode surface. Increasing the LiNO$_3$ concentration to 0.5M and 0.7M finds these structures replaced with increasingly larger toroid structures, as seen in Figure 3.3e and f, respectively.

Of note, in preparing the electrolytes, we found the solubility limit of LiNO$_3$ in DME is approximately 1M. The capacity variability of the 0.7M LiNO$_3$/0.3M LiTFSI cell in Figure 3.2 is likely due to concentration polarization effects in the electrolyte, as 0.7M begins to approach the solubility limit of LiNO$_3$ in DME, such that LiNO$_3$ precipitation at the anode may occur. The LiNO$_3$ concentration range (<0.7M) we report was limited by this effect. Notably, we did not observe a maximum in capacity at any capacity less than saturation. Iliksu et al. recently reported a downturn in capacity above 0.7M LiNO$_3$, implying an optimal concentration of LiNO$_3$ below the solubility limit.$^{67}$ However, above 0.7M LiNO$_3$ we observed a large range of observed capacities that disallowed the assignment of any capacity trend. Some cells in this concentration range exhibited visible salt precipitation, which likely resulted in the observed capacity irreproducibility.

As described previously, the toroid morphology observed in Figure 3.3d-f is indicative of a solution mechanism of Li$_2$O$_2$ growth proceeding through solubility of the LiO$_2$ intermediate.$^{25,45,68}$ The solvation of LiO$_2$ into lithium cations and the redox active superoxide anion, O$_2^-$, follows the equilibrium reaction $^{25}$:

$$\text{LiO}_2^* \leftrightarrow \text{Li}^+\text{(sol)} + O_2^-(\text{sol})$$  \hspace{1cm} (3.4)

Solvated O$_2^-$ can then diffuse in solution to a growing Li$_2$O$_2$ toroid, where it combines with Li$^+$ to form adsorbed LiO$_2^*$ on the toroid surface. LiO$_2^*$ subsequently undergoes disproportionation according to reaction 3.3b, leading to the formation of Li$_2$O$_2$ on the toroid surface, as shown in Figure 3.1b.$^{25}$ The observed toroid formation on discharged cathodes from cells employing high LiNO$_3$/LiTFSI ratios supports the enhancement of this solution mechanism with increasing LiNO$_3$ concentration.

In further support of the solution mechanism, increasingly larger toroid structures were observed as current density decreased in cells employing 0.5M LiNO$_3$ (0.5M
LiTFSI) (Figure 3.4). This observation is consistent with previous reports where $\text{Li}_2\text{O}_2$ toroid formation was correlated to the enhancement of the solution $\text{Li}_2\text{O}_2$ formation mechanism at low currents.\textsuperscript{25,45,68}

Figure 3.4. Scanning electron microscope images of carbon cathodes from cells employing 0.5M LiNO$_3$/0.5M LiTFSI in DME discharged to 1 mAh capacity at currents of a) 10 $\mu$A, b) 25 $\mu$A, c) 50 $\mu$A, and d) 100 $\mu$A. Total electrode area was 1.1 cm$^2$ (12 mm diameter).

With the solubility mechanism confirmed, it was important to check that the increased solubility was indeed due to DN effects of NO$_3^-$ and not another experimental artifact, in particular water contamination. As a first control, the key experiments were repeated with new electrolyte solutions that used lithium nitrate powder that had been dried a second time under vacuum in a heated glove box antechamber at 150°C for 24 hours. All repeated experiments gave consistent results with their original counterparts. As a second control, water levels in the electrolytes were measured via Karl Fischer titration. All electrolytes had less than 70 ppm water, although the 1M LiTFSI (0M LiNO$_3$) had <10 ppm water content. Therefore, a battery employing an electrolyte of 1M LiTFSI in DME with 70 ppm of water was discharged at 450 mA/cm$^2$ to 2 V, analogous to the batteries in Figure 3.2. As observed in Figure 3.5, this battery saw an increase in capacity of 40% compared to a battery with 1M LiTFSI in DME, much smaller than the four-fold increase displayed in Figure 3.2 from the cell employing the 0.3M LiTFSI/0.7M LiNO$_3$ electrolyte.
Figure 3.5. Galvanostatic discharge of a cell employing nominally anhydrous (<10 ppm H₂O, purple curve) and 70 ppm H₂O added (red curve) to 1M LiTFSI in DME. A small water impurity in the LiNO₃ salt led to a linear increase in H₂O content with LiNO₃ concentration, with a maximum water content of 70 ppm H₂O in the 0.7M LiNO₃ cell. A 40% increase in capacity is observed when 70 ppm H₂O is added to a 1M LiTFSI cell. A greater than 400% increase was observed in the 0.7N LiNO₃ cell compared to the anhydrous 1M LiTFSI cell (Figure 3.2a in the main text).

Of note, we find that Li₂O₂ yield, as measured using an established peroxide titration technique, is generally unaffected by the electrolyte compositions studied here (Figure 3.2b), although a slightly higher Li₂O₂ yield may be observed at high LiNO₃ concentrations.

With a change in anion clearly inducing a solution Li₂O₂ growth mechanism, it can be reasoned that the NO₃⁻ anion is affecting LiO₂ solubility via enhanced Li⁺ solvation. The electrolyte anion can affect the overall electrolyte’s donicity (quantified by the Gutmann Donor number (DN), a measure of Lewis basicity), in turn affecting its ability to solubilize LiO₂ through enhanced solvation of Li⁺. We used ⁷Li NMR to probe the electron donicity felt by Li⁺ ions in our LiNO₃/LiTFSI in DME electrolytes as a proxy measurement of the relative effect of the anion on electrolyte DN.

Using NMR as a proxy for DN is a well-known technique, with Erlich et al. first proposing ²³Na NMR as an effective measurement for a solvent’s DN. Erlich et al. reasoned that a downfield ²³Na shift resulted from stronger interaction between the solvation shell molecules and the cation, thereby decreasing the cation’s shielding. The environment of Li⁺ in LiNO₃/LiTFSI in DME electrolytes cannot be determined via ²³Na NMR, though, as adding NaClO₄ to the electrolytes causes a white precipitate to crash out of solution (likely NaNO₃, as dissolving NaClO₄ in an anhydrous solvent containing LiNO₃ has been proposed as a method for making anhydrous LiClO₄).

We reason that ⁷Li NMR, in place of ²³Na NMR, can serve as a reasonable proxy of the relative donicity of Li⁺ electrolytes in a single solvent. A representative ⁷Li NMR
spectrum of a 0.5M LiTFSI/0.5M LiNO$_3$ in DME electrolyte versus our standard external standard of 3M LiCl in D$_2$O is shown in Figure 3.6.

![Representative $^7$Li NMR spectrum on 0.5M LiTFSI/0.5M LiNO$_3$ in DME. The chemical shift at 0 ppm corresponds to the Li shift of LiCl in D$_2$O, whereas the chemical shift at -0.39 ppm corresponds to the Li shift in the electrolyte.](image)

Figure 3.2c shows the $^7$Li chemical shift, referenced to an external standard of LiCl in D$_2$O, of each LiNO$_3$/LiTFSI in DME electrolyte. As LiNO$_3$ concentration increases, the $^7$Li peak shifts downfield, or becomes less shielded. Cahen et al. showed that the $^7$Li chemical shift of a lithium salt may display a concentration dependence, contingent, to a first approximation, on the DN of the solvent and the DN of the anion.$^{71}$ The DN of an electrolyte containing a low DN solvent and high DN anion, like Br$^-$ (DN=33.7) in acetonitrile (DN=14.1), exhibits an anion concentration dependence (DN values from Linert et al.$^{72}$). Conversely, electrolytes comprised of a high DN solvent with a relatively low DN anion, like ClO$_4^-$ (DN=8.44) in dimethyl sulfoxide (DN=29.8), do not exhibit a DN dependence on anion concentration. These trends agree with Linert et al., who found via solvatochromic dyes that the effective DN of an electrolyte depended on an interplay between the DN of the solvent, DN of the anion, and AN of the solvent.$^{72}$ For example, if the solvent’s DN was larger than the anion’s DN, then the electrolyte comprising the two had a DN similar to its solvent’s DN.

If LiNO$_3$ indeed has a higher DN than DME, then increasing the concentration of LiNO$_3$ will increase the number of NO$_3^-$ interacting with any particular Li$^+$, which, in turn, will lead to an increase in the electrolyte’s DN. Thus, we reason that the presence of a concentration dependence on $^7$Li chemical shift indicates NO$_3^-$ serving an active role in the electrolyte’s donicity, and the increasingly downfield shift of $^7$Li with increasing LiNO$_3$ concentration represents increasing donicity. In contrast, Figure 3.7 shows that indeed LiNO$_3$/LiTFSI salts in the high DN solvent dimethyl sulfoxide do not exhibit a
substantial change in $^7\text{Li}$ shift with increasing LiNO$_3$ concentration, and, as therefore expected, no statistically significant capacity increase is observed in DMSO-based electrolytes as the LiNO$_3$/TFSI ratio increases.

![Figure 3.7. a) $^7\text{Li}$ chemical shift of DMSO and DME-based electrolytes, versus a 3 M LiCl in D$_2$O reference, as a function of electrolyte LiNO$_3$ concentration. A less negative chemical shift represents a shift downfield. b) Discharge profiles (45 µA/cm$^2$, 2 V cutoff) as a function of LiNO$_3$ concentration for DMSO and DME-based electrolytes. A 1.0M Li$^+$ concentration was used for all cell electrolytes, while the LiTFSI/LiNO$_3$ ratio was varied. Labels correspond to discharges of cells using the following electrolytes: 1. 1 M LiTFSI in DME, 2. 0.5 M LiNO$_3$:0.5M LiTFSI in DME, 3. 1 M LiTFSI in DMSO, and 4. 0.5 M LiNO$_3$:0.5M LiTFSI in DMSO.]

To confirm the general correlation between enhanced Li$^+$ solvation and Li-O$_2$ battery capacity, another high DN anion, Br$^-$, was studied. As expected, similar trends in $^7\text{Li}$ NMR chemical shifts and Li-O$_2$ battery capacity are observed between 0.5M/0.5M LiBr/LiTFSI and similar LiNO$_3$/LiTFSI concentrations in DME and DMSO-based electrolytes (Figure 3.8). The LiBr-containing electrolytes had <10 ppm water content measured via Karl Fischer titration.

We note, however, that extreme care must be taken when using NMR techniques to compare and quantify solvent DN$s$, particularly between dislike solvent classes, such as protic and aprotic solvents, as was discussed by Gal and Laurence. Additionally, we recognize that $^7\text{Li}$ has comparable, and thus offsetting, diamagnetism and paramagnetism, meaning anisotropic factors can contribute significantly to the observed chemical shifts, unlike in the more commonly used $^{23}\text{Na}$ NMR for measuring solvent DN$s$. For the current study, our interest is only in the relative changes of the Li$^+$ chemical environment as a function of anion composition in a single aprotic solvent (both for DME and DMSO), such that $^7\text{Li}$ NMR provides useful qualitative, if not quantitative, values for comparison.
Figure 3.8. a) $^7$Li chemical shift of DMSO and DME-based electrolytes, versus a 3M LiCl in D$_2$O reference, as a function of LiBr concentration. A less negative chemical shift represents a shift downfield. Analogous to Figure 3.7a) in the main text, 0.5M Br$^-$ causes a noticeable downfield shift in the $^7$Li in DME, but not in DMSO. b) Discharge profiles (45 µA/cm$^2$, 1.5 atm O$_2$ atmosphere, 2 V cutoff), as a function of LiBr concentration for both DMSO and DME-based electrolytes. A 1.0 M Li$^+$ concentration was used for all cell electrolytes, with LiTFSI as the supporting salt. The LiBr concentration for each cell is provided. Analogous to NO$_3^-$ in Figure 3.7b) in the main text, 0.5M Br$^-$ provides over a three-fold increase in capacity in a DME-based electrolyte, but not in a DMSO-based electrolyte.

For a quantitative basis for the role played by the electrolyte anion, collaborators at Carnegie Mellon University and RWTH Aachen University developed a revised thermodynamic model for the solution electrochemical process. The solution mediated electrochemical growth of Li$_2$O$_2$ is triggered by the dissolution reaction given in reaction 3.4. The free energy change involved in this dissolution reaction is given by:

$$
\Delta G_{\text{sol}} = G_{Li^{+}_{\text{sol}}} + G_{O_2^{-}_{\text{sol}}} - G_{LiO_2^*} \quad (3.5)
$$

where $G_{Li^{+}_{\text{sol}}}$ is the free energy of the Li$^{+}$ ions in the electrolyte, $G_{O_2^{-}_{\text{sol}}}$ is the free energy of O$_2^-$ ions in the electrolyte and $G_{LiO_2^*}$ is the free energy of the adsorbed LiO$_2$ on the Li$_2$O$_2$ surface during discharge. To understand the role of salt anion on the equilibrium of the dissolution reaction, the stabilization of the solvated intermediates in the presence of the anion was explored by calculating the free energy of Li$^+$ ions. To a first approximation, the free energy of the Li$^+$ ions and thus the free energy of LiO$_2$ dissolution is largely dependent on the species that are present in Li$^+$ first solvation shell.$^{74-75}$ A modified Ising model$^{76}$ for the site occupancy in the solvation shell of Li$^{+}$ was developed by calculating interaction energies for the various ions with Li$^+$. The model uses inputs of the anion DN, solvent AN, solvent DN, and LiNO$_3$ concentration and is solved via the mean field approximation to calculate the occupancy of the Li$^+$ solvation shell for each of our electrolytes, the resultant average free energy of Li$^+$ (Figure 3.S1), and the resultant expected rate enhancement of the solution mechanism via

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The model shows that at 0.5M LiNO$_3$/0.5M LiTFSI, we would expect a ~4 fold enhancement in the rate of the solution process, in excellent agreement with experiment (Figure 3.S2). The model can be generalized to map out the entire electrolyte design space with a contour map of the Li$^+$ stabilization as a function of varying DN of solvent and anion. Please refer to Burke et al., PNAS 2015 for more information on the model.

With the ability of NO$_3^-$ to induce the solution mechanism confirmed experimentally and computationally, differential electrochemical mass spectrometry (DEMS) was used, as described previously, $^{30,32}$ to quantify the reversibility of the electrochemical reactions (Figure 3.9).

Figures 3.9a and 3.9d present galvanostatic discharge-charge profiles for cells employing 1M LiTFSI and 0.5M LiNO$_3$/0.5M LiTFSI. Discharge, or oxygen reduction reaction (ORR), overpotentials are similar between the two electrolytes. The cell employing LiNO$_3$ salt, however, exhibited a higher initial charge overpotential, which likely arises from an electronic transport limitation to Li$_2$O$_2$ toroid decomposition, although further studies are necessary to confirm this hypothesis. Figure 3.10 shows that for cells charged from the same discharge capacity, this initial charge overpotential does increase with increasing LiNO$_3$ concentration. As discussed in Chapters 5 and 6, this invites the use of an additive redox mediator to allow charge transfer during charge to occur through the electrolyte rather than the electronically insulating Li$_2$O$_2$ toroid.

Figures 3.9b and 3.9e show oxygen consumption as a function of capacity during discharge and charge for these two cells. On both discharge and charge, both cells show nearly identical $e^-$/O$_2$, indicating nearly equivalent degrees of reversibility. Figures 3.9c and 3.9f display gas evolution on charge from DEMS. Oxygen was clearly the dominant gas evolved for both cells, and in nearly identical quantities, with oxygen comprising 92% of the gases evolved from the 0M LiNO$_3$ (1M LiTFSI), and 93% of the gases from the 0.5M LiNO$_3$ (0.5M LiTFSI) cell. The ratio (OER/ORR) of the amount of oxygen evolved during charge (oxygen evolution reaction, OER) to the amount of oxygen consumed during discharge (ORR), an important metric of reversibility, is statistically equal for a cell employing 1M LiTFSI and a cell employing 0.5M LiNO$_3$/0.5M LiTFSI (OER/ORR ~0.82). Small quantities of carbon dioxide and hydrogen were evolved from each cell toward the end of charging, with the 0M LiNO$_3$ evolving slightly more carbon dioxide. The 0.5M LiNO$_3$ cell evolved slightly more hydrogen. Overall, DEMS shows that NO$_3^-$ induces the solution mechanism without deleteriously affecting the battery stability.
Figure 3.9. a, d) Galvanostatic discharge-charge curves for cells employing a-c) 1M LiTFSI in DME and d-f) 0.5M LiNO$_3$/0.5M LiTFSI in DME. b, d) Oxygen consumption during discharge and evolution during charge. c, f) Gas evolution rates for H$_2$, CO$_2$, and O$_2$ during cell charge. These were the only gases found to evolve during charge.
Figure 3.10. Representative discharge-charge profiles of cells of various LiNO₃ concentrations. All cells were discharged at 450 µA/cm² to 0.23 mAh/cm². A 1.0M Li⁺ concentration was used for all cell electrolytes, while the LiTFSI/LiNO₃ ratio was varied. The LiNO₃ concentration for each cell is provided in the figure legend.

DEMS further showed only ¹⁸O₂ is evolved on charge after a discharge under ¹⁸O₂ of a cell employing 0.5M LiN¹⁶O₃/0.5M LiTFSI, confirming that NO₃⁻ does not participate in the electrochemical reaction other than to induce solubility of the intermediates, i.e., NO₃⁻ does not act as a oxygen transfer catalyst via a redox reaction involving a reduced form like NO₂⁻. This result agrees with a similar experiment using pure LiTFSI-based electrolytes,³⁷ implying that electroactive O₂ remains associated during both Li₂O₂ formation and oxidation. Note that this does not preclude NO₂⁻, formed via the reduction of NO₃⁻ at the lithium metal negative electrode, from acting as a redox mediator toward Li₂O₂ via the couple NO₂⁻/NO₂, as has been recently reported.⁷⁸

3.4 Conclusion

In conclusion, we have demonstrated Li⁺ counterion influence on promoting the solubility of electrochemical intermediates during a Li-O₂ battery discharge without further compromising electrolyte stability. Specifically, Li-O₂ batteries employing electrolytes of LiNO₃ and LiTFSI in DME display increasing capacity and increasing toroid formation with increasing LiNO₃ concentration. We ascribe intermediate solubility to enhanced stability of Li⁺ in solution by anions with higher effective donor numbers than that of the solvent, thereby also allowing increased stability of the electrochemically formed anion, O₂⁻, in solution.

The addition in LiNO₃ does not deleteriously affect cell rechargeability, evidenced via DEMS gas analysis, so this strategy can be easily combined with current efforts to identify novel, stable electrolytes. For example, collaborators at the private company Liox and the California Institute of Technology showed that the use of a
eutectic mixture of alkali metal nitrate salts as the electrolyte, without any organic solvent and operated at elevated temperature to melt the salt for ionic conductivity, showed superior reversibility to our typical 1M LiTFSI in DME electrolyte.\textsuperscript{46} The carbon cathode still degraded in the presence of lithium peroxide, but this represents a significant step forward toward a practical electrolyte that both induces the solution mechanism and is stable with lithium peroxide.

Further, a generalized Ising model was developed to predict Li\(^+\) solvation sphere occupation and the resulting stability of Li\(^+\) in electrolytic solutions. We envision this strategy for intermediate stabilization to be generally applicable to numerous nonaqueous systems in which stabilization of desired intermediates may lead to improved electrochemical efficiency. For example, in Li-S batteries, polysulfide intermediate speciation could potentially be controlled by simply tuning the Li\(^+\) anion, perhaps providing a route for increasing sulfur utilization.

3.5 Acknowledgments

The computational work described was performed by Vikram Pande and Professor Venkatasubramanian Viswanathan at Carnegie Mellon University and Dr. Abhishek Khetan at RWTH Aachen University, and is greatly appreciated. Special thanks to Alan Luntz, Dan Addison, Jeffrey Reimer, Hilda Buss, Jessica Nichols, and Christopher Dekmezian for helpful discussions and guidance on characterization of the materials studied here. The work at UC Berkeley was supported in part by previous work performed through the Laboratory Directed Research and Development Program of Lawrence Berkeley National Laboratory under U.S. Department of Energy Contract No. DE-AC02-05CH11231, and support for CMB was provided through the Vehicle Technologies Office under FOA DE-FOA-0000991 and start-up funds through the Department of Chemical and Biomolecular Engineering at UC Berkeley. The work at Carnegie Mellon University was supported through start-up funds through the Department of Mechanical Engineering at Carnegie Mellon University. A. K. thankfully acknowledges the funding for his doctoral studies by the Deustche Forschungsgemeinschaft (DFG).
3.6 Supporting information

Figure 3.S1 a) The occupation of the solvent (red line), TFSI⁻ (green line) and NO₃⁻ (blue line) in the Li⁺ solvation shell and b) the Li⁺ solvation energy (eV) as a function of the concentration of the NO₃⁻ anion. The rate enhancement of the solution process, \( r_S \sim \exp \left( \frac{-\Delta G_{\text{sol}}}{kT} \right) \), is marked on the right y-axis of b). The Li⁺ free energy is normalized relative to that of the case with 1M LiTFSI. There is a rate enhancement, \( r_S \), by a factor of \(~4\) as the concentration of NO₃⁻ is increased from 0.1M to 0.5M. Prepared by Vikram Pande and Professor Venkatasubramanian Viswanathan at Carnegie Mellon University and Dr. Abhishek Khetan at RWTH Aachen University.
Figure 3.S2. Plot showing the linear correlation between capacity (mAh) obtained from experiments (from Figure 3.2a inset) and rate enhancement of the solution process \( r_s \sim \exp\left(\frac{-\Delta G_{sol}}{kT}\right) \) as evaluated from the Ising model for various concentrations of LiNO\(_3\) in DME. The plot shows that the capacity varies exponentially with the Gibbs free energy of solvation of Li\(^+\) in the electrolyte. Prepared by Vikram Pande and Professor Venkatasubramanian Viswanathan at Carnegie Mellon University and Dr. Abhishek Khetan at RWTH Aachen University.
Chapter 4: Influence of electrolyte additives on Li-O₂ discharge capacity

Abstract

The achievable discharge capacity of the lithium-oxygen (Li-O₂) battery is limited by electrode passivation due to the deposition of the ultimate discharge product, lithium peroxide (Li₂O₂), which is an electronic insulator and insoluble in the most stable, ether-based electrolytes. Recent reports have shown electrolyte engineering can increase the achievable discharge capacity by inducing solubility of the electrochemical intermediate to Li₂O₂ formation, lithium superoxide (LiO₂). In this study, we investigate the effect of the Lewis acidity of electrolyte additives toward lowering the free energy of the superoxide anion (O₂⁻) in solution, increasing LiO₂ solubility and increasing achievable discharge capacity. The addition of non-Li alkali metal cations, which are expected to lower the free energy of O₂⁻ relative to lithium cations according to Pearson’s Hard-Soft Acid-Base theory, to ether-based electrolytes results a minimal increase in achievable discharge capacity. From the addition of lithium salt to a sodium-oxygen (Na-O₂) battery, it appears that in electrolytes containing both Li⁺ and non-Li alkali metal cations, Li⁺ quickly scavenges any formed O₂⁻. Therefore, the resultant capacity increase from the addition of non-Li alkali metal salts to a Li-O₂ battery is likely apparent only at high currents, where oxygen reduction occurs at sufficiently high rates to allow some O₂⁻ to temporarily avoid association with Li⁺ and remain in solution as non-Li alkali metal cation-associated O₂⁻, congruent with a recent report. As analogues to the strong Lewis acid water, the addition of ppm quantities of various alcohols to ether-based electrolytes show a two-fold increase in discharge capacity, but with little trend in the Lewis acidity of the alcohols. This highlights the complexity of interactions involving Lewis basicity, Lewis acidity, and other physicochemical properties of each constituent species in an electrolyte and their effect on Li-O₂ battery performance.

4.1 Introduction

A critical issue preventing the realization of a high-capacity, rechargeable lithium-oxygen (Li-O₂) battery is electrode passivation by the ultimate discharge product, lithium peroxide (Li₂O₂). Li₂O₂ is insoluble in the most stable nonaqueous electrolytes, like ethers, and is a wide-band gap insulator. In these electrolytes, Li₂O₂ is therefore conformally deposited on the cathode’s carbon support during discharge (Figure 3.1a), and it electronically passivates the cathode, preventing further discharge after only a fraction of the lithium and oxygen have reacted.¹⁹

As discussed in Chapter 3, the electrolyte can be engineered to circumvent this cathode passivation by enabling a solution-mediated growth mechanism of Li₂O₂ (Figure 3.1b). This mechanism is predicated on the solubility of the intermediate to lithium peroxide formation, lithium superoxide (LiO₂), to its constituent ions, Li⁺ and O₂⁻. The diffusion of these soluble ions away from the electrode surface enables the formation of thermodynamically favored aggregated Li₂O₂ structures and simultaneously leaves the electrode surface accessible for further reaction. Two methods were originally reported to enhance LiO₂ solubility and trigger this solubility-driven growth mechanism. Johnson et
al. presented that the use of a solvent with a high Gutmann Donor Number – a measure of Lewis basicity – like dimethyl sulfoxide (DMSO), increases LiO$_2$ solubility by stabilizing (or lowering the free energy of) Li$^+$ in solution.\textsuperscript{45} Similarly, Aetukuri et al. presented the use of ppm quantities of water (H$_2$O), which has a high Gutmann Acceptor Number – a measure of Lewis acidity – to stabilize (or lower the free energy of) O$_2^-$ in solution.\textsuperscript{25} While both methods showed increased battery capacity via the formation of aggregated Li$_2$O$_2$ structures as confirmed via SEM, neither method proved promising, as each has been shown to decrease battery rechargeability. Strongly electron donating solvents have been shown to be more susceptible to nucleophilic attack by lithium peroxide and thus are more prone to decomposition,\textsuperscript{59} and water impurities have been shown to cause irreversible reactions with the lithium anode.\textsuperscript{24,25}

As discussed in Chapter 3, an appropriately selected lithium salt anion can also induce the solution mechanism of Li$_2$O$_2$ growth. As electron donors, anions have been found to play a role in stabilizing cations in solution.\textsuperscript{72} The most common lithium-oxygen electrolytes use weakly electron donating anions, like bis(trifluoromethyl)sulfonamide (TFSI). The addition of a lithium salt with a strongly electron donating anion, like the nitrate anion (NO$_3^-$), induces LiO$_2$ intermediate solubility without the deleterious effects of water or a strongly electron donating solvent. At increasingly higher concentrations of the strongly electron donating anion, more anions enter the solvation shell of Li$^+$, and thus LiO$_2$ solubility is further enhanced and battery capacity further increased.\textsuperscript{40}

While an appropriately selected anion can increase capacity by four times, there are inherent limitations to the method. Higher concentrations of the anion in the lithium cation solvation shell equate to salt association, so salt solubility and possibly electrolyte conductivity limit the effectiveness of anion selection. Further, anion selection works to increase solution stability of Li$^+$, but Li$^+$ is already very stable in a 1 M solution compared to the superoxide anion, O$_2^-$.\textsuperscript{63}

In this chapter, we investigated the effect of additives to a typical lithium salt/organic solvent electrolyte toward increasing achievable discharge capacity via preferential solvation of O$_2^-$. Non-Li alkali metal cations, like sodium, potassium and cesium, are expected to be stronger electron acceptors from the superoxide anion, O$_2^-$, than lithium cations, as Pearson’s Hard Soft Acid Base theory suggests that increased cation size better suits electron acceptance from the soft base O$_2^-$.\textsuperscript{63} This is supported by the increased solubility of sodium superoxide, NaO$_2$ – the major discharge product in sodium-oxygen batteries – over LiO$_2$ in DME.\textsuperscript{79} Thus, adding these cations to the electrolyte is expected to increase O$_2^-$ solubility, and as a result increase the achievable discharge capacity of lithium-oxygen batteries. Lab-scale Li-O$_2$ batteries containing non-Li alkali metal salts in addition to lithium salt show a minimal increase in achievable discharge capacity. From studying the addition of lithium salt to a sodium-oxygen battery, it appears that in electrolytes containing both Li$^+$ and non-Li alkali metal cations, Li$^+$ quickly scavenges any non-Li alkali metal cation-associated O$_2^-$. This indicates the resultant capacity increase in a lithium-oxygen cell is minimal and only apparent at high currents, as recently reported,\textsuperscript{80} where oxygen reduction occurs quickly enough that some O$_2^-$ temporarily avoids Li$^+$ and remains in solution as non-Li alkali metal cation-associated O$_2^-$.\textsuperscript{63}

In addition to non-Li alkali metal cations, we investigated the effect of ppm quantities of methanol, ethanol, and 1-propanol on the discharge capacity of ether-based
Li-O₂ toward better understanding water’s role in increasing discharge capacity. Galvanostatic discharges show these alcohols provide a two-fold increase in discharge capacity, though with little trend in their Gutmann Acceptor Numbers. This work highlights the complexity of interactions between Lewis basicity, Lewis acidity, and other physicochemical properties of each constituent species.

4.2 Experimental methods

**Materials.** Lithium bis(trifluoromethane) sulfonimide (LiTFSI), 1,2-dimethoxyethane (DME), dimethyl sulfoxide (DMSO), Whatman QM-A glass fiber filters, poly(tetrafluoroethylene) (PTFE, 60 wt% dispersion in H₂O), Vulcan XC72, stainless steel mesh, research-grade oxygen and argon, lithium metal foil, lithium chloride (LiCl), and 99.9% deuterium oxide (D₂O) were all sourced as described in section 3.2. Lithium triflate (LiCF₃SO₃) was purchased from BASF and used as received. Potassium triflate (KCF₃SO₃, 98%) and cesium perchlorate (CsClO₄, 99.995%) were purchased from Sigma-Aldrich and used as received. Sodium triflate was purchased from Sigma-Aldrich and dried at 80 °C under vacuum for at least 12 hours prior to use. Sodium metal was purchased from GalliumSource. AvCarb P50 carbon paper was purchased from Fuel Cell Earth. Water used to contaminate the electrolytes was ultrapure (18.2 MΩ cm) via a Millipore filtration system. Methanol (anhydrous, 99.8%) and ethanol (anhydrous, 99.5%) were purchased from Sigma-Aldrich. 1-propanol (anhydrous, 99.7%, Sigma-Aldrich) was obtained from the UC Berkeley College of Chemistry Reuse Facility.

**Electrolyte preparation.** All electrolyte and cell preparation was carried out in an argon glove box with <0.1 ppm O₂ and <0.1 ppm H₂O. Water, methanol, ethanol, and 1-propanol were quickly and carefully added to electrolytes in the glovebox via micropipette. Water was added at a concentration of 2000 ppm by volume. Methanol, ethanol, and 1-propanol were subsequently added at the equivalent molarity as water, such that by volume, they were added closer to 4500 ppm, 6500 ppm, and 8300 ppm, respectively.

**Cathode preparation.** Cathodes of Vulcan XC72 carbon on stainless steel mesh were prepared via the spray-coating method described in Chapter 2. For sodium-oxygen batteries, P50 carbon paper was punched out in 11 mm diameter discs, which were subsequently rinsed with IPA and acetone prior to drying at 150 °C under vacuum for at least 12 hours. The cathodes were then transferred, while hot, into the glovebox, and stored on a hot plate at 200 °C immediately prior to use.

**Cell preparation.** Lithium-oxygen batteries following the Swagelok design detailed in Chapter 2 were prepared with lithium metal as the negative electrode, Whatman QM-A as a separator, Vulcan XC72 carbon on stainless steel mesh as the positive electrode, and 80 µL of electrolyte as detailed in section 3.2. Sodium-oxygen batteries were prepared similarly, with an 11 mm diameter sodium metal negative electrode, 1/2” Whatman QM-A separator, 11 mm P50 carbon paper positive electrode, and 80 µL of electrolyte.

**Cell characterization.** Achievable discharge capacity was measured as the capacity at which a typical cell discharging at a constant current reached a 2 V cutoff. Pressure monitoring and titrations were performed as described in Chapter 2.
Nuclear magnetic resonance spectroscopy. As described in Chapter 3, $^7$Li nuclear magnetic resonance spectroscopy measurements were completed on a Bruker AM-400 magnet with a 5mm Z-gradient broad brand probe. Each electrolyte was placed in an NMR tube along with a sealed melting point capillary containing 3 M LiCl in D$_2$O as a reliable lithium reference shift.

4.3 Results and discussion
4.3.1 Non-Li alkali metal cations

To characterize the effect of non-Li alkali metal cations on achievable discharge capacity, lab-scale Li-O$_2$ batteries containing non-Li alkali metal cations were discharged at constant current. Figure 4.1 shows representative discharge curves for Li-O$_2$ batteries containing a typical electrolyte of 1 M lithium bis(trifluoromethane) sulfonimide (LiTFSI) in dimethoxyethane (DME) and 0.005 M cesium perchlorate (CsClO$_4$) or potassium triflate (KCF$_3$SO$_3$) salts discharged at a constant current of 0.2 mA/cm$^2$. The addition of non-Li alkali metal salts shows little effect on achievable capacity as compared to the Li$^+$-only electrolyte.

![Figure 4.1](image)

Figure 4.1 Galvanostatic discharge profiles for cells containing 1M LiTFSI in DME (brown), 1M LiTFSI in DME with 0.005M CsClO$_4$ (green), and 1M LiTFSI in DME with 0.005M KCF$_3$SO$_3$ (blue) at 0.2 mA/cm$^2$ to a 2 V cutoff under a constant O$_2$ pressure of ~1.5 atm.

The low concentrations of potassium and cesium were selected based on recent work on the addition of non-Li alkali metal cations to lithium-lithium symmetrical cells to inhibit lithium dendrite growth during electrochemical cycling. As lithium has the lowest standard reduction potential of the alkali metals, a potassium salt electrolyte in contact with a piece of lithium metal would be expected to plate potassium on the lithium electrode while lithium dissolves into the electrolyte.
\[
\begin{align*}
K^+ + e^- & \leftrightarrow K \quad U_0 = -2.931 \text{ V vs. SHE (4.1)} \\
Cs^+ + e^- & \leftrightarrow Cs \quad U_0 = -3.026 \text{ V vs. SHE (4.2)} \\
Li^+ + e^- & \leftrightarrow Li \quad U_0 = -3.040 \text{ V vs. SHE (4.3)}
\end{align*}
\]

However, the standard reduction potentials of the larger alkali metals are sufficiently close to lithium’s that, according to the Nernst equation, the non-Li salts are thermodynamically stable against lithium metal at low concentrations.\(^8\)\(^2\) For a one-electron process at 25 °C, a modified Nernst equation is given by:

\[
U = U_0 - 0.05916 \ln \left( \frac{a_{\text{red}}}{a_{\text{ox}}} \right) \quad (4.4)
\]

where \(U\) is the effective reduction potential, \(U_0\) is the standard reduction potential, \(a_{\text{red}}\) is the activity of the reductant, and \(a_{\text{ox}}\) is the activity of the oxidant. In this case, \(a_{\text{red}}\) is taken as unity for the solid alkali metal, and \(a_{\text{ox}}\) is approximated as the concentration of the cation that is dilute in solution. This equation can be used to estimate the concentration of a non-Li alkali metal cation at which the effective reduction potential equates the standard reduction potential of lithium, i.e., the maximum concentration of a non-Li alkali metal cation before plating onto lithium metal is thermodynamically favored. 0.005 M is a sufficiently low concentration for both K\(^+\) and Cs\(^+\) to expect no non-Li alkali metal plating at the lithium electrode.

Interestingly, recent reports point to the ability of larger concentrations of potassium to significantly increase the achievable discharge capacity of nonaqueous lithium-oxygen batteries. Landa-Medrano et al. showed that the addition of 0.1 M potassium triflate (KCF\(_3\)SO\(_3\)) to a 1 M lithium triflate (LiCF\(_3\)SO\(_3\)) in tetraglyme (TEGDME) electrolyte resulted in a three-fold increase in attainable Li-O\(_2\) battery capacity.\(^8\)\(^0\) Matsuda et al. similarly showed the addition of 50 mM KCF\(_3\)SO\(_3\) increased the achievable discharge capacity two-fold.\(^8\)\(^3\) While the Nernst equation predicts only 0.014 M potassium is stable in thermal equilibrium with lithium metal, these reports imply a kinetic barrier prevents significant potassium plating on the lithium metal negative electrode.

Furthermore, these recent reports showed that a significant capacity increase with non-Li alkali metal cations only occurs at high current densities. This is interesting, as all previous methods of inducing the solution mechanism have shown an increase in the solution mechanism with a decrease in current density, as decreasing the rate of LiO\(_2\) production increases the likelihood of LiO\(_2\) dissolution over immediate reduction or disproportionation to form surface Li\(_2\)O\(_2\).\(^2\)\(^5\),\(^4\)\(^0\),\(^4\)\(^5\) In the case of a non-Li alkali metal salt additive, an increase in the solution mechanism hinges on the electrochemical formation of the non-Li alkali metal superoxide, rather than lithium superoxide, in order to take advantage of the increased solubility of the non-Li alkali metal superoxide over LiO\(_2\). Landa-Medrano et al. propose that high current densities enable KO\(_2\) formation and dissolution with temporary avoidance of O\(_2^-\) with Li\(^+\) ions.

Upon contact in solution, Li\(^+\) immediately reacts with O\(_2^-\) to form LiO\(_2\) and consequently Li\(_2\)O\(_2\) upon additional charge transfer or disproportionation.\(^8\)\(^0\),\(^8\)\(^4\) This mechanism is supported by the addition of lithium salt to a sodium-oxygen battery.
Sodium-oxygen batteries exhibit the 1 e⁻/O₂ formation of sodium superoxide, NaO₂, as their dominant discharge process. Shown in Figure 4.2, the addition of 0.1 M LiCF₃SO₃ to an electrolyte of 1 M NaCF₃SO₃ in DME in a sodium-oxygen battery exhibits a 1.95 e⁻/O₂ process on discharge until it nears the capacity associated with lithium exhaustion, whereby the discharge process switches to an expected 1.08 e⁻/O₂ process. This confirms that Li⁺ quickly scavenges any non-Li alkali metal cation-associated O₂⁻ in metal-oxygen batteries containing both Li⁺ and non-Li alkali metal cations.

Figure 4.2 Potential and pressure profiles for a sodium-oxygen cell containing a sodium anode, P50 carbon paper cathode, and 80 µL of 1M NaCF₃SO₃ and 0.1M LiCF₃SO₃ in DME. Cell was rested at open circuit for 0.5 hr, discharged at 0.2 mA/cm² to 1 mAh capacity, rested at open circuit for 10 min, charged at 0.2 mA/cm² to 3.5 V (reached at 0.82 mAh charge capacity), and rested at open circuit for 10 min. There is sufficient lithium for 0.21 mAh (94 min) of an ideal 2 e⁻/O₂ process, noted by the dotted line.

We tested similar lithium-oxygen cells in our lab to those recent reports mentioned above, particularly focusing on high current discharges where a substantive difference in capacities was observed between purely Li⁺ containing electrolytes and those containing additional small K⁺ or Cs⁺ inclusions. Figure 4.3 shows a small increase in capacity with 0.1 M KCF₃SO₃ at higher current, though a smaller capacity increase than prior reports.⁷⁰, ⁸³ Of note, all cells exhibited a 2.0 e⁻/O₂ process on discharge, indicating little to no formation of KO₂ – a 1 e⁻/O₂ product – as a final product. Toward studying even higher current densities, we performed chronoamperometry in an attempt to maximize formation of non-Li alkali metal-associated O₂⁻ before Li⁺ scavenging. Figure 4.4 shows the resultant current and the corresponding O₂ consumption rates from monitoring cell pressure of cells discharged at 2.3 V for 10 hours. The cell without the potassium additive, displayed in Figure 4.4a, actually exhibited a larger integrated capacity of 1.20 mAh compared with 1.01 mAh for the cell with K⁺, displayed in Figure
4.4b. The axes in Figure 4.4 are scaled for $2 \, e^-/O_2$, showing that each cell also maintained a $2 \, e^-/O_2$ process throughout discharge.

![Galvanostatic discharge profiles](image)

**Figure 4.3** Galvanostatic discharge profiles from cells containing 1M LiCF$_3$SO$_3$ in DME (brown) and 1M LiCF$_3$SO$_3$ in DME with 0.1M KCF$_3$SO$_3$ (blue) at 2 mA/cm$^2$ to a 2 V cutoff under a closed headspace of O$_2$.

While these results are seemingly at odds with the recent literature, various experimental differences could play a role. A different lithium metal surface layer in our lab-scale cells could cause more potassium to plate out, decreasing the amount of potassium active at the cathode. Geometric factors, such as electrode surface area and the ratio of electrolyte volume to electrode surface area, could also differ. It is likely the addition of non-Li alkali metal salts modestly increases the achievable discharge capacity of Li-O$_2$ batteries at high current densities, although not as substantially as the inclusion of ppm quantities of water or high Lewis basicity anions (e.g., NO$_3^-$).
Figure 4.4 Chronoamperometry at 2.3 V under a closed O₂ headspace of cells containing a) 1M LiCF₃SO₃ (LiOTf) in DME and b) 1M LiCF₃SO₃ in DME with 0.1M KCF₃SO₃ (KOTf). Axes for current (brown line) and O₂ consumption rate (gold points) are scaled for a 2 e⁻/O₂ process.
4.3.2 Alcohols

As described in the introduction, another additive considered to increase cell capacity by lowering the free energy of $O_2^-$ in solution is water. Aetukuri et al. showed that the addition of 2000 ppm of $H_2O$ resulted in a six-fold increase in achievable discharge capacity of a Li-O$_2$ battery containing an ether-based electrolyte. Aetukuri et al. argued that water’s high Gutmann Acceptor Number, a measure of Lewis acidity, results in preferential solvation of $O_2^-$ and a concomitant lowering of the free energy of $O_2^-$ in solution.\textsuperscript{25}

Toward investigating this mechanism, we studied the effect on Li-O$_2$ discharge capacity of adding ppm quantities of anhydrous methanol, ethanol, and 1-propanol to an ether-based electrolyte. Considering the chemical similarity of water and alcohols (R-OH, with R=H for water and R=alkyl substituents for alcohols), increasing the alkyl chain length correlates to decreasing Lewis acidity, or decreasing Acceptor Number, as shown in Table 4.1. Electrolytes consisting of our typical, most-stable electrolyte of 1 M lithium bis(trifluoromethane)sulfonimide (LiTFSI) in dimethoxyethane (DME) were prepared with 2000 ppm H$_2$O by volume or an identical molarity of anhydrous methanol, ethanol, or 1-propanol, labeled as “2000 ppm equivalent” in Table 4.1.

Table 4.1 presents the average achievable discharge capacity of lab-scale Li-O$_2$ batteries containing these electrolytes from galvanostatic discharges at 0.5 mA/cm$^2$, normalized by the milligrams of carbon in the electrode, which is a proxy for the available surface area. The alcohols induce a two-fold increase in battery capacity relative to no additives. However, water increases the capacity more than five-fold, and there is little trend in the capacity as a function of the additive’s Acceptor Number. Of note, each electrolyte composition corresponded to a near 2 e$^-$/O$_2$ process, and titrations confirmed the formation of Li$_2$O$_2$ as the dominant discharge product.

Table 4.1 Discharge capacity for batteries containing 1M LiTFSI in DME and added H$_2$O, methanol, ethanol, or 1-propanol. The added H$_2$O is 2000 ppm by volume; the methanol, ethanol, and propanol were added such that equivalent moles were added as for the H$_2$O electrolyte (i.e., 4500 ppm, 6500 ppm, and 8300 ppm by volume, respectively). Gutmann Acceptor Numbers and Donor Numbers are from Knovel Solvents – A Properties Database.\textsuperscript{85}

<table>
<thead>
<tr>
<th>Additive to 1M LiTFSI in DME</th>
<th>Additive AN (kcal/mol)</th>
<th>Additive DN</th>
<th>Average capacity (mAh/mg)</th>
<th>Electrolyte $^{7}Li$ peak (ppm) referenced to 3M LiCl in D$_2$O</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>n/a</td>
<td>n/a</td>
<td>0.3</td>
<td>-1.18</td>
</tr>
<tr>
<td>2000ppm H$_2$O</td>
<td>54.8</td>
<td>18</td>
<td>1.7</td>
<td>-1.12</td>
</tr>
<tr>
<td>2000ppm equiv. methanol</td>
<td>41.5</td>
<td>30</td>
<td>0.6</td>
<td>-1.16</td>
</tr>
<tr>
<td>2000ppm equiv. ethanol</td>
<td>37.1</td>
<td>32</td>
<td>0.6</td>
<td>-1.16</td>
</tr>
<tr>
<td>2000ppm equiv. 1-propanol</td>
<td>33.7</td>
<td>30</td>
<td>0.7</td>
<td>-1.16</td>
</tr>
</tbody>
</table>

As seen in Table 4.1, the alcohols each have a higher Donor Number than water, in addition to their lower Acceptor Numbers. This could point to stronger interaction
between the alcohol and Li\(^+\) than water and Li\(^+\), at the expense of favorable solvation of O\(_2\). This hypothesis is supported by the effect of water on achievable discharge capacity when dimethyl sulfoxide (DMSO) is used as the electrolyte solvent. DMSO is a stronger Lewis base than DME, with a Gutmann Donor Number of 29.8 kcal/mol compared to DME’s 20 kcal/mol.\(^8\) As a strong Lewis base, DMSO has been shown to strongly solvate Li\(^+\), but it possibly also strongly solvates Lewis acid additives, like water, decreasing their interaction with O\(_2\). Table 4.2 shows the addition of 2000 ppm H\(_2\)O to a DMSO-based electrolyte results in the same ultimate capacity as 2000 ppm H\(_2\)O in DME. This corresponds to a three-fold increase in capacity over an anhydrous DMSO electrolyte, whereas 2000 ppm H\(_2\)O in DME corresponds to more than a five-fold increase in capacity over an anhydrous DME electrolyte. This shows the interaction of DMSO with Li\(^+\) and the interaction of H\(_2\)O with O\(_2\)\(^-\) are not mutually exclusive, as the ultimate capacity is less than the sum of the enhancement from just DMSO and from just 2000 ppm H\(_2\)O. Rather, the resultant capacity is at the least a complex mixture of the accepting and donating properties of each electrolyte constituent.

Table 4.2 Li-O\(_2\) battery capacity results for dimethyl sulfoxide (DMSO)-based electrolytes. Acceptor Number and Donor Number values are from Knovel Solvents – A Properties Database.\(^8\)

<table>
<thead>
<tr>
<th>Additive to 1M LiTFSI in DMSO</th>
<th>Additive AN</th>
<th>Additive DN (kcal/mol)</th>
<th>Average capacity (mAh/mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>None</td>
<td>n/a</td>
<td>n/a</td>
<td>0.6</td>
</tr>
<tr>
<td>2000 ppm H(_2)O</td>
<td>54.8</td>
<td>18</td>
<td>1.8</td>
</tr>
</tbody>
</table>

In an attempt to measure the effect of the alcohols’ higher Donor Numbers on Li\(^+\) solvation, \(^7\)Li NMR was performed, as discussed in Chapter 3. Figure 4.5 shows each electrolyte’s \(^7\)Li peak referenced to 3 M LiCl in D\(_2\)O, a reliable reference shift for lithium, contained in a sealed melting point capillary within the sample tube. The exact electrolyte peak shifts are listed in Table 4.1 as well. Only one \(^7\)Li peak was observed for each electrolyte composition, and is indicative of the average of all Li\(^+\) environments in the electrolyte. A downfield shift of this lithium peak relative to the reference indicates the lithium is on average interacting with stronger Lewis bases, indicating a higher apparent Donor Number of the electrolyte. There is a slight correlation between the \(^7\)Li peak and the capacity, but the magnitude of the change (e.g., 0.06 ppm from anhydrous to 2000 ppm H\(_2\)O) is small. To compare, an electrolyte of 0.5 M LiNO\(_3\)/0.5 M LiTFSI in DME, which induces a capacity increase similar to 1000 ppm H\(_2\)O, exhibits an electrolyte peak shifted over 0.5 ppm further downfield than the electrolyte peak of anhydrous 1 M LiTFSI in DME. Furthermore, \(^7\)Li peaks can be complicated by more factors than just electron density due to a relative balance in lithium’s paramagnetic and diamagnetic contributions.\(^7\) Thus, this perhaps simply highlights the difficulty in assessing Li\(^-\) solvation from \(^7\)Li NMR.
4.4 Conclusion

In conclusion, we studied the effect of adding non-Li alkali metal cations and ppm quantities of alcohols to ether-based Li-O\textsubscript{2} battery electrolytes on Li-O\textsubscript{2} discharge capacity. The addition of non-Li alkali metal salts induces a minimal increase in achievable discharge capacity in our lab-scale Li-O\textsubscript{2} cells. Gas analysis of the addition of lithium salt to a sodium-oxygen battery shows Li\textsuperscript{+} quickly scavenges any non-Li alkali metal cation-associated O\textsubscript{2}\textsuperscript{-} in electrolytes containing both Li\textsuperscript{+} and non-Li alkali metal cations. This corroborates that a capacity increase is only expected at high currents when oxygen reduction occurs quickly enough that some O\textsubscript{2}\textsuperscript{-} temporarily avoids Li\textsuperscript{+} and solubilizes as non-Li alkali metal cation-associated O\textsubscript{2}\textsuperscript{-}, as recently proposed.\textsuperscript{80}

The addition of ppm quantities of methanol, ethanol, or 1-propanol to ether-based Li-O\textsubscript{2} battery electrolytes induces a two-fold increase in battery capacity. However, there is little trend in the capacity as a function of the additive’s Acceptor Number, highlighting the complexity of interactions between Lewis basicity, Lewis acidity, and other physicochemical properties of each constituent species in an electrolyte.

Experimental confirmation of the ability of additive species to solvate and lower the free energy of the superoxide anion is difficult, as O\textsubscript{2}\textsuperscript{-} forms in situ and is a short-lived species in the presence of Li\textsuperscript{+}. Through the NASA Space Technology Research Fellowship (NSTRF) program, I have worked with collaborators at NASA’s Ames Research Center developing density functional theory and molecular dynamics modeling of additives’ influence on the solvation of O\textsubscript{2}\textsuperscript{-} in ether-based Li-O\textsubscript{2} electrolytes. Dr. John Lawson’s group has recently published robust molecular dynamics models of lithium electrolytes, both in the bulk and at electrochemical interfaces, which can determine
solvation shells and double layer behavior.\textsuperscript{86-87} Applying these models to ether-based electrolytes containing LiO\textsubscript{2} and either non-Li alkali cations or ppm quantities of water or alcohols could inform the important physicochemical properties of additives for preferential solvation of O\textsubscript{2}\textsuperscript{−}.

4.5 Acknowledgments

I gratefully acknowledge Chris Dekmezian and Howie Nguyen for working with me as undergraduate researchers on the effect of non-Li alkali metal cations on Li-O\textsubscript{2} discharge capacity. This work was supported by a NASA Space Technology Research Fellowship.
Chapter 5: Implications of 4 e⁻ oxygen reduction via iodide redox mediation in Li-O₂ batteries with trace water


Abstract

The nonaqueous lithium-oxygen (Li-O₂) electrochemistry has garnered significant attention due to its high theoretical specific energy compared to the state-of-the-art lithium-ion battery. The common active nonaqueous Li-O₂ battery cathode electrochemistry is the formation (discharge) and decomposition (charge) of lithium peroxide, Li₂O₂. In high-capacity cells, Li₂O₂ forms in large aggregated structures, but it is an insulator and insoluble in common Li-O₂ battery electrolytes and is therefore difficult to oxidize on charge. Recent research has focused on identifying small molecules that can serve as soluble redox mediators between Li₂O₂ and the electrode surface. One of the best candidates is lithium iodide, LiI, because the I⁻/I₃⁻ redox potential is only slightly higher (~3V vs. Li/Li⁺) than the equilibrium potential of Li₂O₂ oxidation to O₂ (~2.96V vs. Li/Li⁺). Interestingly, recent reports suggest that the introduction of lithium iodide to an ether-based electrolyte containing water at impurity levels induces a 4 e⁻ oxygen reduction reaction forming lithium hydroxide, LiOH, as the dominant discharge product. We provide quantitative analysis of the influence of LiI and H₂O on the electrochemistry in a common Li-O₂ battery employing an ether-based electrolyte and a carbon cathode. We confirm, through numerous quantitative techniques, that the addition of LiI and H₂O promotes efficient 4 e⁻ oxygen reduction to LiOH on discharge, which is unexpected given that only 2 e⁻ oxygen reduction is typically observed at undoped carbon electrodes. Unfortunately, LiOH is not reversibly oxidized to O₂ on charge, where instead a complicated mix of redox shuttling and side reactions is observed.

5.1 Introduction

Due to its high theoretical specific energy, the Li-air (or O₂) battery is currently being explored as a possible alternative to lithium-ion batteries. In typical nonaqueous Li-O₂ cell compositions, lithium peroxide (Li₂O₂) formation and oxidation has been identified as the reversible electrochemical reaction at the carbon cathode:

\[ 2\text{Li}^+ + 2e^- + \text{O}_2 \leftrightarrow \text{Li}_2\text{O}_2 \quad (2e^- \text{ process}) \quad U = 2.96 \text{ V vs. Li}^+/\text{Li} \] (5.1)

The forward reaction occurs on discharge and the reverse reaction occurs on charge. Unfortunately, many unsolved scientific challenges inhibit the achievement of reversible, efficient Li₂O₂ formation and oxidation, with two being particularly pressing. First, Li₂O₂ and its intermediates (O₂⁻ and LiO₂) are known to undergo parasitic side reactions with organic electrolytes and carbon cathodes during battery operation. When a realistically small electrolyte volume is used, these reactions limit Li-O₂ cell rechargeability to only a few cycles. Additionally, the products of these parasitic
reactions include solid carbonate/carboxylate species, which accumulate at the electrode surface, thereby blocking O\textsubscript{2} evolution from the Li\textsubscript{2}O\textsubscript{2}-electrolyte interface on charge.\textsuperscript{34} As a result, the potential on charge necessarily increases to values substantially higher (>1 V higher) than the equilibrium potential for O\textsubscript{2} evolution from Li\textsubscript{2}O\textsubscript{2}, creating an energy inefficient battery. With these parasitic reactions seemingly exacerbated at high potentials, this challenge perpetuates itself. Therefore, parasitic reactions are linked with energy inefficiency and are detrimental to rechargeability.

Second, Li\textsubscript{2}O\textsubscript{2} is a wide bandgap insulator and is insoluble in all studied organic electrolytes. Therefore, as it deposits on the cathode during discharge, it passivates the electrode surface, limiting Li-O\textsubscript{2} cell discharge capacity to a minor fraction of that necessary to compete with the specific energy of Li-ion batteries.\textsuperscript{38-39, 42, 48-50, 94} Recent reports have elucidated strategies to partially circumvent this passivation issue by triggering a solution-based mechanism for Li\textsubscript{2}O\textsubscript{2} formation.\textsuperscript{25, 40, 45-46} However, this mechanism results in Li\textsubscript{2}O\textsubscript{2} deposition in large toroids, as shown in Figure 5.1a, and perhaps Li\textsubscript{2}O\textsubscript{2} deposition away from the cathode surface.\textsuperscript{68} A fraction of Li\textsubscript{2}O\textsubscript{2} therefore becomes electronically isolated from the conductive cathode surface, likely contributing to the observed large charge overpotentials.

![Figure 5.1](image)

**Figure 5.1.** Schematics for charging lithium peroxide particles (pink) on carbon electrode (charcoal) in typical ether-based electrolytes (blue) **a)** without and **b)** with redox mediation.

Substantial research into redox mediators is being pursued to address these charge transport limitations. In the context of Li-O\textsubscript{2} batteries, redox mediators are small, soluble molecules with a reversible redox potential slightly above the equilibrium potential of the Li-O\textsubscript{2} battery (experimentally observed as ~2.85 V).\textsuperscript{37} During charge, as shown in Figure 5.1b, the mediator oxidizes at electronically accessible sites on the cathode, diffuses to electronically isolated Li\textsubscript{2}O\textsubscript{2}, and accepts charge via an outer sphere electron transfer from the Li\textsubscript{2}O\textsubscript{2}, allowing the Li\textsubscript{2}O\textsubscript{2} to be oxidized while regenerating the redox mediator.\textsuperscript{95} The battery therefore charges near the redox potential of the mediator, which, if appropriately selected, is only slightly above the open circuit potential of the battery, rather than reaching the high potentials required to presumably drive charge transport through Li\textsubscript{2}O\textsubscript{2}. The lower charge potential also provides a route to reduce parasitic processes observed at high potentials. Numerous redox mediators have now been studied in Li-O\textsubscript{2} batteries, including tetrathiafulvalene (TTF),\textsuperscript{96} (2,2,6,6-Tetramethylpiperidin-1-
yl)oxyl (TEMPO), tris[4-(diethylamino)phenyl]amine (TDPA), iron phthalocyanine (FePC), and lithium iodide (LiI).

LiI is a redox mediator of particular interest. Iodide, I, exhibits two redox couples in a common nonaqueous ether-based Li-O₂ electrolyte:

\[
3I^- \leftrightarrow I_3^- + 2e^- \quad U = \sim 3 \text{ V vs. Li}^+/\text{Li} \quad (5.2)
\]

\[
2I_3^- \leftrightarrow 3I_2 + 2e^- \quad U = \sim 3.5 \text{ V vs. Li}^+/\text{Li} \quad (5.3)
\]

The potentials listed are experimental oxidative onset potentials in dimethoxyethane, as shown in Figure 5.2 via cyclic voltammetry in lithium/porous carbon cells under an argon headspace. The first redox couple, ascribed to the I⁻/I₃⁻ redox reaction shown in equation 5.2, is just above the Li-O₂ equilibrium potential, and I⁻ is stable against reduction against lithium unlike many organic redox mediators.

![Figure 5.2. Cyclic voltammetry of cells under an argon headspace containing XC72 carbon electrodes and 0.25 M LiTFSI in DME (brown line), 0.25 M LiTFSI/0.2 M LiI in DME (green line), and 0.25 M LiTFSI/0.2 M LiI/500 ppm H₂O in DME (blue line). Each cell was reductively scanned to 2 V, anodically scanned to 4.25 V, and returned to OCV. Sweep rate was 10 mV/s. CVs were IR corrected with electrolyte resistance via PEIS impedance. PEIS parameters were 10 mV amplitude from 100 kHz to 50 mHz, 8 points per decade. CVs clearly show I⁻/I₃⁻ (oxidative onset at ~3.0 V) and I₃⁻/I₂ (oxidative onset at ~3.5 V) redox couples.](image)

Interestingly, recent reports indicated that in a typical nonaqueous Li-O₂ cell containing LiI and water impurities, Li₂O₂ was not the ultimate discharge product, but rather LiOH. This is interesting, as water by itself does not induce the formation of LiOH rather than Li₂O₂, and possibly provides a route to reducing parasitic reactions.
typically associated with Li$_2$O$_2$ formation and oxidation. Furthermore, an incredibly low voltage gap (~200 mV) between the discharge and charge voltage plateaus was observed, indicating that the cell achieved a remarkable energy efficiency.\textsuperscript{101} LiOH oxidation to O$_2$ and H$_2$O through the action of the I$^-$/I$_3^-$ redox couple (3 V vs. Li/Li$^+$) was reported as the active charge electrochemistry, although the I$^-$/I$_3^-$ redox potential lies below the standard potential for LiOH oxidation at standard conditions (3.45 V vs. Li/Li$^+$). Liu et al. indicate in their motivating work that the equilibrium of water, iodide, and oxygen is complex, and the role of polyanions (iodo-oxygen species) on the LiOH oxidation mechanism is poorly understood.\textsuperscript{101, 104-105} Therefore, while the reported electrochemistry of the I$^-$/I$_3^-$ couple for Li-O$_2$ batteries is intriguing, the exact activity of the I$^-$ and its oxidized states, I$_3^-$ and I$_2$, towards Li$_2$O$_2$/LiOH oxidation is unclear.

In this chapter, with the help of collaborators at the University of Waterloo and the private company Liox, we present quantitative insight into the influence of LiI and H$_2$O on the electrochemistry of a typical nonaqueous Li-O$_2$ cell employing a carbon cathode. LiOH is observed as the primary discharge product, along with minor parasitic side products, when LiI and H$_2$O are both present in the cell, with H$_2$O consumed in the oxygen reduction reaction. If H$_2$O (which is present in impurity quantities in our study) is fully consumed during discharge, the discharge electrochemistry reverts back to the more common 2 e$^-$ oxygen reduction reaction to form Li$_2$O$_2$. On charge, the concentrations of LiI and H$_2$O and the identities of the discharge products all have a dramatic influence on the observed electrochemistry and resultant voltage profiles. Isotopic labeling confirmed that O$_2$ evolution is only observed to occur from Li$_2$O$_2$, although reactions between Li$_2$O$_2$ and oxidized iodo species (I$_3^-$) can result in other product formation at low potentials (~3 V). No O$_2$ evolution was observed from electrochemically formed LiOH. Instead, LiOH reacts with I$_2$, which electrochemically forms at ~3.5V from I$_3^-$ oxidation, to form H$_2$O and LiIO$_3$, the latter of which can be solubilized in water-contaminated DME. Thus, while the 4 e$^-$ oxygen reduction activity on carbon electrodes in the presence of water impurities and I$^-$ is intriguing, the addition of LiI and water to a Li-O$_2$ battery does not promote a reversible oxygen electrochemistry in the cell compositions studied.

### 5.2 Experimental methods

Materials and methods described below are for my contributions to this study. Li-O$_2$ cells with slight variations in design and material were employed by our collaborators (supporting information figures) but showed consistent results, as discussed at the beginning of section 5.3.

**Materials.** Lithium iodide was purchased from Sigma-Aldrich and was dried under vacuum in a heated glove box antechamber at 150 °C for 24 hours before use. Lithium bis(trifluoromethane) sulfonamide (LiTFSI) and 1,2-dimethoxyethane (DME) were purchased from BASF and used as received. PTFE (both 60 wt% dispersion in H$_2$O and 1 µm particle-size powder) was purchased from Sigma-Aldrich. Vulcan XC72 was purchased from Fuel Cell Store and was filtered through a 60-mesh screen. Ketjenblack® (KB) was received from Toyota. T316 stainless steel 120 mesh, with wire diameter 0.0026”, was purchased from TWP Inc. Research-grade oxygen and argon were purchased from Praxair. 99% $^{18}$O$_2$ was purchased from Sigma-Aldrich. 90% H$_2$ $^{18}$O was purchased from Cambridge Isotopes. 99.9% deuterium oxide (D$_2$O) was purchased from Cambridge Isotope Laboratories, Inc. and used as received. Water used to contaminate
the electrolytes was ultrapure (18.2 MΩ cm, Millipore). All electrolyte and cell preparation was carried out in an argon glove box with <0.1 ppm O₂ and <0.1 ppm H₂O. Water was quickly and carefully added to electrolytes in the glovebox via micropipette. Lithium foil with a thickness of 0.01” was purchased from FMC Lithium.

Cathode preparation. The XC72 cathodes were prepared as described in Chapter 2. The Ketjenblack carbon cathodes were prepared via a slurry method. First, 20 mg of PTFE powder was added to 500 µL of IPA and sonicated for at least 1 hour. Second, a volume of this solution was mixed with four times its volume of ethanol and added to a vial containing lightly ground KB such that the final mixture was 85:15 w:w KB to PTFE. This mixture was sonicated and stirred for five minutes, three times each, until thick but flowing. If the mixture was too thick or started to dry out, more ethanol was added (usually about a couple of milliliters). The mixture was then spread over 12 mm diameter pieces of stainless steel mesh that had been cleaned with IPA and acetone and dried at 150°C. These were left to dry in a fume hood for 12 hours, and then were dried at 150 °C under vacuum for at least 12 hours and transferred hot to the glove box, similarly to the XC72 cathodes.

Cathode carbon loadings are identified in figure captions. Each set of data (high LiI DEMS, low LiI DEMS, titrations, etc.) used cathodes from the same batch with similar loading.

Cell assembly. Li-O₂ cells following the same hermetically sealed Swagelok design as described in Chapter 2 were built in an argon glove box with < 0.1 ppm H₂O and O₂. Cells consisted of 7/16” diameter pure lithium metal, 1/2” diameter Whatman QMA glass fiber as a separator, electrolyte, and a 12 mm diameter cathode of XC72 or KB carbon, with PTFE binder, on stainless steel mesh. Either 80 or 160 µL of electrolyte of varying concentrations of LiI, LiTFSI, and water in nominally anhydrous DME were added to each cell. Cells employing 80 µL contained one glass fiber separator; cells employing 160 µL contained two. The cells had a headspace of approximately 1.5 mL, which were discharged starting with a headspace of ~1.5 atm pure O₂, and charged starting with a headspace of ~1.2 atm Ar.

Gas quantification. To quantify oxygen consumption and gas evolution, such as shown in Figure 5.3, cells were attached to a differential electrochemical mass spectrometry (DEMS) setup with a pressure transducer within the closed cell headspace, as described in Chapter 2.

On discharge, oxygen consumption was quantified via pressure decay of a known cell headspace volume. Cells containing ≥2000 ppm of water showed a discernible pressure increase at open circuit potential due to hydrogen evolution from the water reacting with the lithium anode. Cells were left at open circuit potential for 30 minutes prior to discharge and 30 minutes after discharge to record the pressure rise. An average of these rises was subtracted from the pressure decay to obtain a corrected pressure change during discharge. Pressure rise was only ever witnessed before and after a 4 e⁻/O₂ process, never before or after a 2 e⁻/O₂ process, because in LiI-containing cells, a 2 e⁻/O₂ process only occurred if any intentionally added water was entirely consumed during LiOH formation, obviously disallowing parasitic water reduction to evolve hydrogen at the anode. Therefore, if a battery exhibited a switch from a 4 e⁻/O₂ process to a 2 e⁻/O₂ process in the middle of the discharge, it was approximated that the onset of the 2 e⁻/O₂ process was where pressure rise from hydrogen evolution at the anode ceased, which was
then taken into consideration during calculations of O$_2$ consumption in the pressure rise/decay measurements.

Of note, hydrogen evolution, and thus the reaction between water and the lithium metal electrode, is negligible for cells with low water contents (<2000 ppm). We do not see an increase in pressure over 30 minutes, with a resolution of +/- 0.2 torr. An increase in pressure of 0.2 torr over 30 minutes would correspond to a gas evolution rate of 0.001 µmol/min. The actual rate of H$_2$ evolution/LiOH formation at the anode will consistently decrease during discharge as water is consumed at both the anode and cathode, such that over the course of a 10 hour discharge (100 µA to 1 mAh), at most 0.6 µmol of LiOH could be formed at the lithium metal electrode.

For cells containing high water contents, the amount of LiOH formed at the lithium metal electrode is much smaller than that electrochemically formed at the positive electrode. Of the battery compositions studied, a battery containing 160 µL of electrolyte with 5000 ppm H$_2$O exhibited the most LiOH formation at the lithium metal electrode, as evidenced by hydrogen evolution at open circuit potential. This was confirmed via DEMS and then quantified via pressure measurements of a closed headspace. The initial rate of water consumption at the lithium metal electrode to form LiOH in these cells was 0.0088 µmol/min. Over the course of a 10-hour discharge (100 µA to 1 mAh), this corresponds to approximately 5.3 µmol of chemically-formed LiOH at the anode. We expect substantially less LiOH formation than this at the anode, as no H$_2$ evolution was observed after a discharge, indicating that the LiOH formation rate at the anode decreases to undetectable values during the discharge. At the cathode, these high water content batteries exhibit 37.3 µmol of LiOH formation. Thus, electrochemical formation of LiOH at the cathode is clearly the dominant source of LiOH in these high water content cells.

On charge, gas evolution was quantified by DEMS, as described in Chapter 2. Again, the cell headspace at the beginning of charge was pure argon. Whenever a cell was charged with the mass spec, an identical cell was charged with simple pressure rise measurements to verify that the gas evolution from the mass spectrometer (O$_2$, H$_2$, CO$_2$) matched the observed pressure rise.

*Titrations for LiOH and Li$_2$O$_2$ quantification.* Titrations of carbon positive electrodes for discharge product quantification were completed as described in Chapter 2.

Of note, there are three sources of base for the acid/base titration: LiOH formed as the ultimate discharge product, LiOH formed via the reaction of Li$_2$O$_2$ with water in the titration protocol, and basic decomposition products. As discussed more completely in section 5.3, on discharge, all three sources were uncoupled. The quantity of basic decomposition products was estimated as 24% of the quantity of Li$_2$O$_2$ formed, a consistent value for identical cells without lithium iodide. The quantity of LiOH formed as the discharge product was therefore the total amount of base titrated minus the quantity of basic decomposition products and the stoichiometric amount of LiOH formed from reacting the titrated value of Li$_2$O$_2$ with water. The expected values in Figure 5.4 were calculated via the observed discharge e'/O$_2$ value, i.e., a 4.0 e'/O$_2$ expects only LiOH formation, whereas a value of 3.0 e'/O$_2$ expects both Li$_2$O$_2$ and LiOH formed in a 1:2 ratio, and a 2.0 e'/O$_2$ expects only Li$_2$O$_2$ formation. On charge, the basic decomposition products and electrochemically formed LiOH cannot be decoupled, as the fate of the basic decomposition products formed during discharge is unknown on charge (i.e., they may or may not oxidize during charge).
Additionally, as the iodometric titration is based on quantitatively titrating I₂, I₂ formation from the I₃⁻/I₂ redox couple at high charge voltages is a false positive that makes the titration not applicable after high charge voltages, such as in Figure 5.4 at point iv.

Charing with no prior discharge. As shown in Figure 5.5 and Figure 5.6, typical cells constructed as detailed above were charged with a headspace of ~1.2 atm Ar with no prior discharge, i.e., no prior formation of Li₂O₂. The cells were charged at constant current to an arbitrary final capacity or a 4 V cutoff.

5.3 Results and discussion

We initially investigated the impact of various H₂O and LiI electrolyte concentrations on the discharge and charge voltage profiles of the Li-O₂ cell. The voltage response for a Ketjenblack positive electrode discharged and charged (100 μA/cm² to a capacity of 1 mAh/cm²) with different concentrations of LiI and H₂O in DME-based aprotic electrolyte is shown in Figure 5.S1. Only small changes are observed in the discharge voltage profile between cells with varying LiI and H₂O concentrations. On charge, the potential profile is clearly influenced by the LiI and H₂O concentrations. Generally, the higher LiI concentration cells show lower charging voltages compared to their lower LiI concentration counterparts. In fact, the 0.2M LiI, 500 ppm H₂O cell maintains a ~3 V plateau for the duration of charge, consistent with previous reports. As a result, the energy efficiency of this cell appears outstanding, as only a ~0.25 V gap between the discharge and charge voltages is observed. High water concentrations, in addition to low LiI concentrations, appear to result in higher final charging voltages, as well. Of note, other carbon-based electrodes exhibit only small differences in overpotentials compared to Ketjenblack electrodes. In particular, XC72 carbon black, a different type of high-surface area carbon, shows similar overpotentials to Ketjenblack (e.g. Figure 5.S1 and Figure 5.7 give comparison of Ketjenblack and XC72, respectively, with 0.05M LiI), and therefore results from cells comprised of either carbon electrode can be directly compared if needed. See figure captions for the specific carbon used for that data set. Figure 5.S2 shows that graphene-based electrodes exhibit similar overpotentials as well, and thus graphene-based electrodes were not studied further.

As discussed in Chapter 2, energy efficiency is not necessarily a measure of the reversibility of Li-O₂ cells, as the discharge and charge profiles could correspond to undesired reactions. The electrochemical reactions at various lithium iodide and water concentrations were quantified via a combination of differential electrochemical mass spectrometry (DEMS), pressure decay/rise measurements in a closed cell headspace, and acid/base and peroxide titrations, as discussed in the experimental section.

Figure 5.3 shows pressure decay (discharge) and DEMS (charge) results for batteries employing 0.2 M lithium iodide and water concentrations corresponding to nominally anhydrous (< 20 ppm), 500 ppm, and 5000 ppm. 0.2 M LiI was selected for these measurements because, at modest current densities and with the given overall cell geometry, this concentration, coupled with a low water concentration, enables the low ~3 V potential throughout the entire duration of charge, as observed in Figure 5.S1 and in previous reports. Pressure decay and DEMS allow for the direct correlation of observed oxygen consumption and evolution to the observed voltage profiles of our galvanostatic cycling.
Figure 5.3. High [LiI] DEMS results. Galvanostatic discharge and charge profiles (100 µA/cm²) with oxygen consumption and gas evolution for lithium-oxygen batteries employing a lithium anode, a cathode of ~1 mg Ketjenblack® with PTFE binder on stainless steel mesh (12mm diameter), and 160 µL of DME with 0.25 M LiTFSI and 0.2 M LiI as the electrolyte. Each column presents a battery with different added electrolyte water content: a-c) nominally anhydrous; d-f) 500 ppm D₂O, and g-i) 5000 ppm H₂O. The batteries were discharged under a headspace of ~1.5 atm oxygen and charged under a constant pressure headspace of ~1.2 atm argon. b), e) and h) show oxygen consumption on discharge and evolution on charge. The asterisk (*) denotes a value adjusted for a hydrogen evolution rate at open circuit potential, which arises from water reacting with the lithium anode. Gas evolution on charge was quantified via DEMS. c), f), and i) show quantitative gas evolution throughout the charge.

Focusing on the discharge processes of the cells presented in Figure 5.3, all three cells exhibit oxygen consumption with rates varying between a pure 2 e⁻/O₂ process and a pure 4 e⁻/O₂ process depending on the cell’s total water content. The nominally anhydrous battery exhibits a clear 2 e⁻/O₂ process for the duration of the discharge, indicative of lithium peroxide formation as is typically observed in nonaqueous Li-O₂
cells (Figure 5.3a-c). The battery containing 500 ppm water (Figure 5.3d-f) exhibits a 3.2 e⁻/O₂ process for the first 0.24 mAh of discharge, and a 2 e⁻/O₂ process for the remainder of the 1 mAh discharge. The initial 3.2 e⁻/O₂ process indicates a mix of lithium hydroxide formation (reaction 5.4) – a 4 e⁻/O₂ process – and lithium peroxide formation (reaction 5.1, from the introduction) – a 2 e⁻/O₂ process.

$$4\text{Li}^+ + 4e^- + O_2 + H_2O \leftrightarrow 4\text{LiOH} \quad (4e^- \text{ process}) \quad (5.4)$$

The switch from a 3.2 e⁻/O₂ process to a 2 e⁻/O₂ process likely occurs once H₂O is completely consumed, as 500 ppm of water in the electrolyte volume employed (160 mL) corresponds to a capacity of 0.24 mAh for LiOH formation (as calculated from the reaction 5.4 stoichiometry), and the discharge potential noticeably drops to a lower plateau at the onset of the 2 e⁻/O₂ Li₂O₂ formation process. The battery containing 5000 ppm water (Figure 5.3g-i) exhibits a near 4 e⁻/O₂ process for the duration of the 1 mAh discharge, indicating lithium hydroxide formation dominates at high water contents. Theoretically, as calculated from reaction 5.4, 5000 ppm water in the electrolyte volume employed (160 µL) corresponds to a capacity of 2.4 mAh for LiOH formation, so there is sufficient water to allow a 1 mAh discharge even after considering (the discernible) water consumption at the lithium anode. The presence of LiOH as a discharge product was further confirmed via X-ray diffraction (XRD) studies of the cathode, as shown in Figure 5.S3 for a cell with high LiI and H₂O content.

Quantification of both LiOH and Li₂O₂ was performed using established titrations on extracted cathodes from cells with identical composition as those studied in Figure 5.3 (Figure 5.4). For a battery containing 500 ppm H₂O and 0.2 M LiI (Figure 5.3d-f), 5.8 µmol LiOH and 15.8 µmol Li₂O₂ are expected to form during the 1 mAh discharge given the amount of O₂ consumed during the 4 e⁻/O₂ and 2 e⁻/O₂ processes; 5.1 µmol LiOH and 11.1 µmol Li₂O₂ were titrated (Figure 5.4a-b), giving yields of 87% LiOH and 70% Li₂O₂. The Li₂O₂ yield is consistent with previous reports for nominally anhydrous Li-O₂ cells employing Ketjenblack® cathodes. Therefore LiOH formation is more efficient than Li₂O₂ formation in these cells, although parasitic processes during LiOH formation still persist. As identified previously by both Kwak et al. and Liu et al., LiOH formation in a cell of this composition is an interesting finding, as the 4 e⁻/O₂ ORR reaction is uncommon in nonaqueous Li-O₂ batteries, and on carbon electrodes in general. However, Tulodziecki et al. recently showed this LiOH formation does not follow a typical 4 e⁻/O₂ reduction process such as typically observed on platinum electrodes. LiOH is likely formed via the 2 e⁻/O₂ formation of Li₂O₂, followed by the chemical reactions of Li₂O₂ with water to form LiOH and the consumption of H₂O via iodide, followed finally by the reduction of triiodide back to iodide. This work and mechanism is discussed further in section 6.1.

On charge, the three cells presented in Figure 5.3 exhibit very little oxygen evolution compared to the oxygen consumed during discharge. The total oxygen evolution from the nominally anhydrous battery, for example, is 2.4 µmol, or less than 13% of what would be expected assuming full oxidation of the discharge products. As the nominally anhydrous and 500 ppm water electrolytes exhibit Li₂O₂ as their dominant discharge product and the 5000 ppm water electrolyte exhibits LiOH as its dominant discharge product, the limited O₂ evolution during the cells’ ~3 V charge plateaus shows...
that the ~3 V charge plateau is predominantly related to electrochemical processes other than O₂ evolution, irrespective of the discharge product. Additionally, while the 5000 ppm battery exhibits a small amount of oxygen evolution during the step from the ~3 V plateau to the higher ~3.5 V plateau, DEMS shows negligible oxygen evolution otherwise. Therefore, LiOH is not reversibly oxidized to produce O₂ at these potentials. The cell chemistry on charge is extremely complex, being a function of the concentrations of I⁻ and H₂O, as well as the ratio of discharge capacity to total water content in the cell (and hence, the product distribution formed during discharge).

Numerous processes occur during charge at the ~3 V plateau. If substantial Li₂O₂ is formed, as is the case in the low water content cells (Figure 5.3a-g), it can disappear

**Figure 5.4.** Acid/base and iodometric titration results for cells identical to those studied in Figure 5.3 d-f) (a) and b) 500 ppm H₂O) and g-i) (c) and d) 5000 ppm H₂O). The acid/base titration quantifies lithium hydroxide and other decomposition products present; these are distinguished by using the expected value of basic decomposition products formed during lithium peroxide formation. On charge, the relative fates of the decomposition products and the electrochemically formed LiOH are unknown, so they cannot be decoupled. The iodometric titration gives the lithium peroxide results. *I₂ formation during the 3.5V plateau in c-d interfered with the iodometric titration’s ability to properly quantify Li₂O₂ at the end of charge (7.3 µmol of ‘Li₂O₂’ were titrated at the end of charge, even though only 0.3 µmol were produced during discharge).
during this plateau, as observed from iodometric titrations presented in Figure 5.4a-b. A small fraction of Li$_2$O$_2$ disappearance is ascribed to oxidation that evolves O$_2$ (Figure 5.3c, f) while the remainder appears to chemically react with I$_3^-$ to form a soluble product. I$_3^-$ was likely formed from I$^-$ oxidation during charge in Figure 5.4a, as the battery separator after charge was a faint yellow-red in color, indicative of I$_3^-$ in aprotic polar media. We note that I$_3^-$ formed during charge can result in a false positive in our iodometric titration, so that the difference in the Li$_2$O$_2$ titrated between points i and ii in Figure 5.4a (9.4 µmol difference, or 85% of the Li$_2$O$_2$ formed during discharge) represents a lower limit of the actual Li$_2$O$_2$ that disappeared on charge.

![Graph](image)

**Figure 5.5.** Charging cells under an argon headspace with no prior discharge (charge in this case goes left to right) containing XC72 carbon electrodes and 0.25 M LiTFSI in DME (brown line), 0.25 M LiTFSI/0.2 M LiI in DME (green line), and 0.25 M LiTFSI/0.2 M LiI/500 ppm H$_2$O in DME (blue line) at 100 µA/cm$^2$. If all I$^-$ were converted to I$_3^-$ in the cells containing LiI, only 0.29 mAh of charge capacity would be expected before voltage rise. The ability of the cells containing LiI to galvanostatically charge in perpetuity when no Li$_2$O$_2$ is ascribed to a redox shuttle, i.e., iodide being oxidized at the carbon positive electrode to triiodide, then diffusing to the lithium electrode, becoming electrochemically reduced back to iodide, and diffusing back to the positive electrode to repeat. Figure 5.5S shows the current dependence of redox shuttling in cells containing 0.05M LiI, as well as an inset schematic for redox shuttling.

The Li$_2$O$_2$ disappearance cannot account for the entirety of the electrochemical charging capacity, implying that the ~3 V plateau is related in part to I$^-$/I$_3^-$ redox shuttling between the anode and cathode. Figure 5.5 shows cells with an argon headspace charged at constant current with no prior discharge. A cell containing 0.25 M LiTFSI in DME exhibits immediate polarization, with the voltage rising to a cutoff of 4 V. Meanwhile, cells containing 0.2 M LiI, with and without 500 ppm H$_2$O, can be charged in perpetuity at a ~3 V plateau, with Figure 5.5 showing the first 1 mAh of charging. If all I$^-$ were converted to I$_3^-$ in this cell without any shuttling, a capacity of only 0.29 mAh would be
expected before voltage rise. This ability to “charge” without any prior formation of discharge product is ascribed to a redox shuttle mechanism, shown in the inset of Figure 5.S4, of iodide being oxidized at the carbon positive electrode, diffusing to the lithium electrode, becoming electrochemically reduced, and returning to the positive electrode to repeat. Figure 5.S4 shows similarly that a cell containing just 0.05 M LiI and 500 ppm H₂O that has yet to be discharged can be “charged” at low current densities at ~3 V in perpetuity, while Figure 5.S4b shows that an attempted charge of an identical cell without LiI exhibits an immediate sharp rise in voltage. Generally, the shuttling process is known to be strongly influenced by mediator concentration, diffusion coefficient, current density, and interelectrode distance. Figure 5.5 and Figure 5.S4 are evidence that an I⁻/I₃⁻ redox shuttle current can be maintained between the electrodes at a current density of roughly 100 µA/cm² which is the current density used in Figures 5.S1 and 5.3, in our typical cell geometries. Of note, analogous cells that employ higher concentrations of water, such as 5000 ppm, exhibit the two-plateau behavior as shown in Figure 5.3g. Figure 5.6 shows that a battery containing 4500 ppm H₂O charged with no prior discharge exhibits a step up to 3.5 V before subsequent perpetual charging. Complete conversion of I⁻ to I₃⁻ and I₃⁻ to I₂ in this cell corresponds to 0.107 mAh, near where the step occurs. This indicates water’s ability, perhaps through its reaction with and subsequent passivation of the lithium metal electrode, to inhibit the system’s ability to sustain the I⁻/I₃⁻ redox shuttling between electrodes.

![Graph showing potential vs. capacity for different electrode combinations](image)

**Figure 5.6** Charging cells under an argon headspace with no prior discharge (charge in this case goes left to right) containing XC72 carbon electrodes and 0.25 M LiTFSI, 0.05 M LiI, 4500 ppm H₂O in DME (blue line) or 0.25 M LiTFSI, 0.05 M LiI in DME (green line) at 50 µA/cm². Electrolyte volume is 80 µL. Complete conversion of I⁻ to I₃⁻ corresponds to 0.071 mAh; subsequent conversion of I₃⁻ to I₂ corresponds to 0.036 mAh.
When LiOH is the dominant discharge product, i.e., cells with high water content, it only disappears during the ~3.5 V plateau, and not the ~3 V plateau (e.g., Figure 5.3g), suggesting the I\(_3^-/I^-\) redox shuttle dominates at the ~3 V plateau. As shown in Figure 5.4a, acid/base titrations indicate that after a full discharge-charge cycle, cells with low water content – and thus those that exhibit the ~3 V plateau for the duration of charge – still contain all LiOH formed during discharge. This is consistent with SEM images (Figure 5.S5) showing that solid discharge products remain after charge, indicating that there is no activity for LiOH oxidation at this low charge potential. Cathodes discharged and charged in cells with high water content that reach the ~3.5 V plateau on charge, however, exhibit (after charge) only a minor fraction of the basic products formed during discharge. The ~3.5 V plateau is likely related to the I\(_3^-/I^-\) redox couple, which is active at these potentials, and I\(_2\) formation is clearly visible in the glass fiber separators of cells that reach this plateau. Recalling the minimal O\(_2\) evolution exhibited at even the ~3.5 V plateau (Figure 5.3g-i), this suggests that the dominant chemistry occurring at this plateau is a non-O\(_2\) evolving LiOH oxidation reaction involving the I\(_3^-/I^-\) redox couple, which will be discussed later.

Isotopic labeling of O\(_2\) and H\(_2\)O finds that no evolved O\(_2\) originates from LiOH formed during discharge. O\(_2\) evolution from LiOH during charge requires O-O bond reformation that occurs between statistically random oxygen atoms contained in the LiOH. LiOH forms during discharge through O\(_2\) bond splitting and reaction with H\(_2\)O. The oxygen evolution of a cell that was charged after being discharged under a pure \(^{18,18}\)O\(_2\) headspace, with H\(_2\)^\(^{16}\)O as the water impurity, is shown in Figure 5.3f (500 ppm H\(_2\)O cell). Pure \(^{18,18}\)O\(_2\) is evolved during charge. This result is also consistent with a cell discharged under \(^{16,16}\)O\(_2\) and containing a H\(_2\)^\(^{18}\)O impurity; Figure 5.S6 presents the oxygen evolution of a cell that was charged after being discharged to a capacity of 1 mAh in 0.2M LiI/5000 ppm H\(_2\)^\(^{18}\)O under \(^{16,16}\)O\(_2\). No \(^{18}\)O-enriched O\(_2\) is evolved at any point during the entire charge process. Both of these cell isotope configurations (Figure 5.3f and Figure 5.S6) would be expected to evolve significant \(^{16,18}\)O\(_2\) on charge (i.e., the evolved O\(_2\) would be enriched in the H\(_2\)O O isotope) if O\(_2\) evolution occurred from LiOH. In Figure 5.S6, the transition from I\(_3^-\) to I\(_2^-\) is accompanied by the evolution of a small volume of isotopically pure \(^{16}\)O\(_2\), indicating that the O-O bond in O\(_2\) remained intact throughout the discharge-charge process, as expected with Li\(_2\)O\(_2\) formation and oxidation. This confirms that the oxidation of LiOH to form O\(_2\) does not occur either chemically or electrochemically, and the small amount of O\(_2\) observed from cells containing LiOH as the primary discharge product can be ascribed to the oxidation of Li\(_2\)O\(_2\).

Interestingly, lowering the iodide concentration enables higher rates of oxygen evolution on charge when lithium peroxide is the primary discharge product. For example, Figure 5.7 presents pressure decay (discharge) and DEMS (charge) results for batteries discharged with 0.05 M LiI (instead of 0.2 M LiI, as was used in Figure 5.3) and 2000ppm of water in 80 µl of electrolyte (corresponding to 0.5 mAh of possible LiOH formation). When discharged beyond 0.5 mAh (Figure 5.7a-c), a switch from a 4 e^-/O\(_2\) process to a 2 e^-/O\(_2\) process is observed on discharge, and O\(_2\) evolution is observed on charge. The O\(_2\) evolution coincides with a rising charge voltage above ~3.2 V, and corresponds to 63% of the expected yield based on the O\(_2\) consumed during the 2 e^-/O\(_2\) discharge process. This OER/ORR value is similar to (though slightly lower than) the
yield in aprotic Li-O₂ cells in the absence of LiI. A battery discharged to a cutoff capacity of less than 0.5 mAh, where only a 4 e⁻/O₂ process is observed (Figure 5.7d-f), exhibits only minor oxygen evolution on charge. These results further confirm that only lithium peroxide, and not LiOH, can be oxidized to evolve O₂ in aprotic electrolytes on carbon cathodes. Batteries employing lower iodide concentrations are not able to maintain the desirable ∼3 V charge plateau, but do exhibit more oxygen evolution on charge than the higher LiI concentration cells, with the oxygen originating exclusively from lithium peroxide.

Figure 5.7. Low [LiI] DEMS results. a) and d) show galvanostatic discharge and charge profiles (200 µA/cm²) from lithium-oxygen batteries employing a lithium anode, a cathode of ∼2 mg XC72 carbon with PTFE binder on stainless steel mesh (12 mm diameter), and 80 µL of an electrolyte of 0.25 M LiTFSI, 0.05 M LiI, and roughly 2000 ppm H₂O in DME. The only difference between the batteries is the discharge capacity cutoff, as labeled. b) and e) show oxygen consumption on discharge and evolution on charge versus capacity. c) and f) show quantitative gas evolution throughout the charge. The asterisk (*) denotes a value adjusted for a hydrogen evolution rate at open circuit potential, which arises from water reacting with the lithium anode.
The DEMS and titration data clearly indicate that the addition of lithium iodide to a water-contaminated, nonaqueous lithium-oxygen battery enables a 4 e-/O₂ process involving the consumption of water to form lithium hydroxide on discharge. However, on charge, lithium hydroxide is not reversibly oxidized back to oxygen. Any lithium peroxide formed during discharge is only partially oxidized to evolve oxygen on charge, with higher oxygen evolution efficiencies observed at lower iodide concentrations. Consequently, the low 3 V plateau on charge obtained in Figure 5.S1 is not due to reversible oxygen electrochemistry, but rather to a complex iodo-oxygen electrochemistry that is highly dependent on the exact cell composition and testing conditions (e.g., LiI and H₂O concentration, current density, cathode composition, and cell geometry).

This still leaves open the question of the fate of LiOH at the high, ~3.5 V charge plateau (i.e., where it is observed to disappear). Contrasting reports exist that demonstrate the removal of LiOH from the electrode surface during charge of a Li-O₂/LiI cell. Liu et al. reports that I₃⁻ oxidizes LiOH to evolve O₂ during the ~3 V plateau. Our own results, as well as those from Kwak et al., demonstrate that LiOH does not react at the low potential plateau, but can be removed from the cathode surface at the higher (~3.5 V) voltage plateau, but without O₂ evolution. One possible explanation for this behavior is that LiOH oxidation results in LiIO₃ formation. In alkaline aqueous media, the following reaction occurs:

\[ 3I_2 + 6LiOH \leftrightarrow LiIO_3 + 5LiI + 3H_2O \quad (5.5) \]

Hence, it is possible that at high charge voltages, when the I₃⁻/I₂ redox couple is actively forming I₂, LiOH reacts to form small amounts of LiIO₃, given that a 6 LiOH:1 LiIO₃ stoichiometric ratio is expected from reaction 5.5.

To prove this hypothesis, our collaborators at the University of Waterloo developed a modified version of the iodometric titration to detect small amounts of LiIO₃ present on various positive electrodes charged to different capacities. A detailed description of this method is in the supporting information. The fraction of LiIO₃ was determined for electrodes stopped at various states of charge in either 0.05 M LiI or 0.2 M LiI/5000 ppm H₂O/DME electrolyte (Figure 5.S7). This water concentration was chosen to ensure LiOH was formed as the overwhelming dominant product on discharge. For low LiI concentrations (0.05 M), it is clear that a small quantity of LiIO₃ is produced (< 0.5 µmol) on cell charge. The small magnitude is understandable, as at 0.05 M LiI, complete conversion of I⁻ to I₂ equates to a maximum of 1.5 µmol LiIO₃ produced via reaction 5.5, assuming the discharge product is 100% LiOH. Nonetheless, the presence of LiIO₃ is confirmed on extracted positive electrodes with both EDX analysis, which shows LiIO₃’s characteristic 3:1 ratio of O:I (Figure 5.S8), and XRD analysis, which shows diffraction patterns characteristic of LiIO₃ (Figure 5.S9). A similar quantity of LiIO₃ is also observed for the cell discharged/charged in 0.2 M LiI at voltages below 3.4 V (< 0.5 µmol). However, at high voltages and significant capacity overcharge, and thus high concentrations of I₂, approximately 3.0 µmols of LiIO₃ are observed on the electrode surface, clearly indicating a higher reactivity of LiOH with I₂ than I₃⁻. Reaction between LiOH and I₂ (and the lack of a reaction between LiOH and I₃⁻) was confirmed with mock chemical experiments, as shown in Figure 5.S10. LiIO₃ is very soluble in H₂O (76 g/100
62 mL), and hence can easily dissolve in the 5000 ppm H$_2$O electrolyte. Dissolution is further increased with the production of H$_2$O accompanying LiIO$_3$ formation. This is a plausible explanation for the removal of LiOH from the electrode surface, but does not fully explain the removal of LiOH at low voltages by Liu et al.$^{101}$ We note that the total amount of water contained in our cells is small compared to the water included in cells studied by Liu et al. (e.g., 5000 ppm H$_2$O in 160 µL in our cells versus 45,000 ppm H$_2$O in 700 µL in Liu et al.). This distinction could give rise to the discrepancy observed at low voltages in each study, as both LiOH solubility and the chemical reaction rate between LiOH and I$_3^-$ increases with increasing water content.$^{104}$ Other factors that could play a role in this discrepancy are depth of discharge (i.e., amount of product formed) and the time delay between the end of discharge and the characterization of the discharge products.

5.4 Conclusion

In conclusion, the effect of LiI and H$_2$O on the Li-O$_2$ chemistry can be summarized as follows (Figure 5.8). On discharge, the presence of both LiI and H$_2$O promote a 4 e$^-$/O$_2$ process (LiOH production) instead of the normal 2 e$^-$/O$_2$ process (Li$_2$O$_2$ production) typical of an aprotic Li-O$_2$ cell employing an ether-based electrolyte and undoped carbon cathode. High H$_2$O concentrations result in almost stoichiometric 4 e$^-$/O$_2$ LiOH formation, whereas intermediate/low H$_2$O concentrations result in a mixture of Li$_2$O$_2$ and LiOH formation. On charge, a variety of processes occur which are dependent on the final discharge product composition and the concentrations of LiI and H$_2$O. I$^-$ oxidation, in the absence of any discharge product, features two redox couples, I$^-$/I$_3^-$ (E ~ 3.0 V) and I$_3^-$/$I_2$ (E ~ 3.5 V). The I$^-$/I$_3^-$ shuttle between the anode and cathode substantially contributes to the cell electrochemistry at low charge potentials (2.95 – 3.10 V). LiOH does not oxidize when this I$^-$/I$_3^-$ couple is active, although Li$_2$O$_2$ can react to either evolve O$_2$ or form unidentified iodo-oxygen species. The I$_3^-$/$I_2$ redox is observed after the initial I$^-$/I$_3^-$ redox in cells with high water content in the electrolyte. This transition results in O$_2$ evolution from Li$_2$O$_2$, presumably promoted by the formed I$_2$. The I$_3^-$/$I_2$ redox is active during a second charge plateau that takes place at ~3.5 V, and LiOH is observed to disappear during this process. LiOH reacts with I$_2$ to form a small amount of LiI$_2$O$_3$, which is soluble in the water-filled electrolyte via reaction 5.5, but does not evolve O$_2$. Based on our studies, while a low charge overpotential makes the use of I$^-$ seem appealing, the lack of O$_2$ evolution from Li$_2$O$_2$/LiOH oxidation in LiI-containing cells makes it a poor additive for efficient Li-O$_2$ cycling. Of the oxygen consumed on discharge, less than 13% is recovered on charge under the best-case ~3 V charge scenario. Hence, the low charge overpotential seen in LiI-containing Li-O$_2$ batteries is primarily due to iodo-oxygen electrochemistry, rather than reversible oxygen evolution. It should be noted, given our results, that other cell compositions (e.g., those with cathodes comprised of active materials other than carbon, higher water or LiI concentrations than studied here, other halide salts, etc.) may result in a substantially different electrochemistry, and more research is certainly necessary to fully understand the complex charge chemistry observed in this system.
Figure 5.8. Summary of processes in typical Li-O₂ batteries employing an ether-based electrolyte, carbon cathode, LiI, and water impurities.

5.5 Acknowledgments

This work was performed in collaboration with Dr. Robert Black, Ivan Kochetkov and Professor Linda Nazar at the University of Waterloo and Dan Addison and Dr. Vincent Giordani at the private company Liox. Helpful discussions with Prof. Clare Grey are gratefully acknowledged. CMB, BDM, VG and DA gratefully acknowledge support from the U.S. Department of Energy, Energy Efficiency and Renewable Energy Vehicle Technologies Office under award No. DE-0006869, and previous support that enabled this work through the Laboratory Directed Research and Development Program of Lawrence Berkeley National Laboratory under U.S. Department of Energy Contract No. DE-AC02-05CH11231. CMB gratefully acknowledges support from a NASA Space Technology Research Fellowship under award No. NNX16AM56H. LFN is grateful for funding for this work from NRCan through their ECO-EII program, and acknowledges NSERC for partial support through the Discovery and Canada Research Chair programs.
5.6 Supporting information

Collaborators’ experimental methods:

*Electrode X-ray diffraction.* XRD measurements were carried out using a Bruker D-8 Advance diffractometer employing Cu-Kα radiation (λ = 1.5406 Å). Samples were mounted on a silicon low-background holder using a moisture-protective barrier.

*Scanning electron microscopy.* Cathodes were removed from cells after cell completion. The cathodes were washed with THF, dried under vacuum and mounted onto SEM stubs with double sided carbon tape in a hermetically sealed argon-filled box. Transfer from the glovebox to the SEM was performed with minimal (i.e., 1 sec) exposure to the atmosphere upon placement into the SEM load-lock chamber. Analysis was performed with a Zeiss Ultraplus FESEM.

*LiIO₃ detection.* For quantitative determination of LiIO₃, a modified version of the iodometric titration for Li₂O₂ was used. Once electrodes + separators were removed from the cell, washed with THF, and dried, the entire contents were placed into ~ 5 mL of H₂O (exact volume known via mass balance). Coarse MnO₂ powder was added to the solution and stirred with the cell contents for approximately 2 hours. The addition of Li₂O₂ to H₂O produces H₂O₂, which is then oxidized on the surface of MnO₂ to give O₂ + H₂O. Hence, after 2 hours, the entirety of H₂O₂ was oxidized, and what remains is LiOH and LiIO₃ dissolved in a known volume of H₂O. The entire contents were then filtered 3 times through a 0.2 µm syringe filter to fully remove MnO₂. MnO₂ is a false positive for this measurement, and thus mock experiments were performed to ensure that all MnO₂ is removed after three-fold filtration. The final transparent liquid (2.5 mL) was used for titration. Approximately 50 mg KI and 0.5 mL of 5 M H₂SO₄ were added which turned the solution yellow. Excess I⁻ and H⁺ were added to ensure full conversion of the reaction LiIO₃ + 6H⁺ + 8I⁻ → 3I₃⁻ + 3 H₂O. In the presence of excess I⁻, I₂ converts to I₃⁻, which is then titrated using a thiosulfate solution in the same manner as described previously for quantification of Li₂O₂.
Figure 5.S1. Voltage profiles of Ketjenblack electrodes (4 mg carbon) galvanostatically discharged (lower curves, left to right) and charged (higher curves, left to right) in DME electrolyte with different LiI concentration (0.05 M and 0.2 M) and H₂O concentration (500 ppm and 5000 ppm) at 100 µA/cm². The total salt concentration (LiI + LiTFSI) was kept at 0.3 M. The total electrolyte volume used is 180 µL. At low LiI concentrations (0.05 M), regardless of H₂O concentration, the charge profile gradually slopes to higher voltage. At the higher LiI concentrations, an initial low voltage charge plateau (~2.95 V) is exhibited. Only at the end of charge is a rise in potential observed for 5000 ppm H₂O, while the low charge plateau continues for the low (500 ppm) H₂O concentration. Prepared by Dr. Robert Black, Ivan Kochetkov, and Professor Linda Nazar at the University of Waterloo.
Figure 5.S2. Comparison of reduced graphene oxide and C750 graphene positive electrodes with 0.2 M LiI and either 500 ppm or 5000 ppm H₂O. This demonstrates that regardless of the positive electrode source, the electrochemistry is relatively constant. All electrodes were loaded at 4 mg. In the case of RGO, to achieve loadings of ~4 mg, a relatively thick electrode had to be constructed that resulted in poor mechanical stability. We believe this contributes to the larger overpotential observed compared to that of G750 graphene and KB electrodes. Graphene oxide was prepared via the oxidation of graphite (Alfa Aesar), followed by reduction to rGO according to the procedure previously reported by Grey et al.¹⁰¹; material was centrifuged in the initial washing steps. Prepared by Dr. Robert Black, Ivan Kochetkov, and Professor Linda Nazar at the University of Waterloo.
Figure 5.S3. a) X-ray diffraction pattern of a positive electrode after discharge in 0.2 M LiI/5000 ppm H$_2$O/DME. The only crystalline product is LiOH; b) SEM micrograph of the same discharged positive electrode. Prepared by Dr. Robert Black, Ivan Kochetkov, and Professor Linda Nazar at the University of Waterloo.
Figure 5.S4. Li-O$_2$ batteries charged under 1 atm argon gas with no prior discharge as a function of current density. Electrolyte is 0.25 M LiTFSI and 500 ppm H$_2$O in DME with 0.05M LiI (a) and no added LiI (b). These cells were constructed with two Whatman GF/D glass fiber separators, 300 µL of electrolyte, and cathodes containing 2 mg of a 95:5 mass ratio Super P carbon to PTFE on 10 mm stainless steel mesh. If all I$^-$ were converted to I$_3^-$ in the cells containing LiI, only 0.27 mAh of charge capacity would be expected before a voltage rise. The interelectrode distance was ~500 µm. Prepared by Dr. Vincent Giordani and Dan Addison at Liox.
Figure 5.S5. SEM micrographs of the positive electrode surface at the end of charge. a) single discharge/charge profile of cells run in electrolyte with 500 ppm and 5000 ppm H₂O (0.2 M LiI/DME), and corresponding SEM micrograph at the end of charge for b) 500 ppm H₂O and c) 5000 ppm H₂O. Prepared by Dr. Robert Black, Ivan Kochetkov, and Professor Linda Nazar at the University of Waterloo.
Figure 5.S6. Voltage profile and corresponding oxygen evolution profile for a cell discharged under $^{16}\text{O}_2$ and charged in 0.2 M LiI/DME with 5000 ppm H$_2^{18}\text{O}$. On discharge, a reasonable fraction of total produced LiOH is $^{18}\text{O}$ labeled (some Li$_2\text{O}_2$ is also produced). On charge, only $^{16}\text{O}_2$ is observed from the oxidation of Li$_2\text{O}_2$. No signal from $^{18}\text{O}_2$ (LiOH oxidation) is observed at either low potential ($I_{1}/I_{3}$) or high potential ($I_{3}/I_{2}$). Prepared by Dr. Robert Black, Ivan Kochetkov, and Professor Linda Nazar at the University of Waterloo.
Figure 5.87. Electrochemical curves for batteries containing a) 0.05 M LiI/5000 ppm H₂O/DME and c) 0.2 M LiI/5000 ppm H₂O/DME, and corresponding LiIO₃ for b) 0.05 M and d) 0.2 M LiI at the points specified. At low LiI concentrations, little LiIO₃ is generated due to the small concentration of I₂. At high LiI concentrations, an abundance of LiIO₃ is observed at V > 3.5 due to the generation of large amounts of I₂, which reacts with the LiOH to form LiIO₃. Batteries were cycled at 100 µA/cm² and cathode carbon loading was 4 mg of KB. Prepared by Dr. Robert Black, Ivan Kochetkov, and Professor Linda Nazar at the University of Waterloo.
Figure 5.S8. Energy-dispersive X-ray spectroscopy spectrum of a Ketjenblack positive electrode extracted from cell containing 0.05 M LiI, 0.25 M LiTFSI, and 5000 ppm H₂O in DME after discharge at ~100 µA/cm² to 1 mAh and charge at ~100 µA/cm² to 1 mAh. As these electrolyte and cycling conditions lead to I₂ production on charge, LiOH and I₂ react to form LiIO₃ on the positive electrode, as exhibited by an O:I ratio of 3:1 on the positive electrode surface. Prepared by Dr. Robert Black, Ivan Kochetkov, and Professor Linda Nazar at the University of Waterloo.
Figure 5.S9. X-ray diffraction pattern of a Ketjenblack positive electrode extracted from cell containing 0.05 M LiI, 0.25 M LiTFSI, and 5000 ppm H$_2$O in DME after discharge at ~100 $\mu$A/cm$^2$ to 1 mAh and charge at ~100 $\mu$A/cm$^2$ to 1 mAh. As these electrolyte and cycling conditions lead to I$_2$ production on charge, LiOH and I$_2$ react to form LiIO$_3$ on the positive electrode. Dashed lines are the major reference markers from the JCPDS database, as shown in Figure 5.S10. Prepared by Dr. Robert Black, Ivan Kochetkov, and Professor Linda Nazar at the University of Waterloo.
Figure 5.S10. Powder X-ray diffraction patterns of a) LiOH, and resulting solids after mixing 2.0 mmol LiOH with b) 2.5 mmol LiI and 2mmol I₂ in DME, and c) 4 mmol I₂ in DME. The reaction period was ~ 8 hours. Reference markers for LiOH (blue, in a) and b)) and LiIO₃ (red, in c)) are from the JCPDS database. Prepared by Dr. Robert Black, Ivan Kochetkov, and Professor Linda Nazar at the Univeristy of Waterloo.
Discussion of Figure 5.S10:

Mock chemical reactions were performed to determine the degree of reactivity between I$_3$/I$_2$ with LiOH. The reflections corresponding to the starting LiOH product are shown in Figure 5.S10a above. A solution of I$_3^-$ was prepared with a mixture of 2.5 mmol LiI and 2 mmol I$_2$. In DME, the equilibrium I$^- + I_2 \rightarrow I_3^-$ (pKa ~ 6.5) is nearly shifted 100% towards the product, creating a solution of I$_3^-$ with excess I$^-$ (no I$_2$ present). The contents were left to stir for 12 hours. The reflections of the resulting solid product are the same as the starting LiOH, leading us to conclude that no chemical reaction occurs (Figure 5.S10b). The XRD reflections of the solid product obtained from the reaction of LiOH with a solution of pure I$_2$ (4 mmol in DME) indicate LiIO$_3$ is formed via the reaction: 3I$_2 + 6$ LiOH $\rightarrow$ LiIO$_3 + 5$ LiI + 3 H$_2$O. Other reactions involving I$_2$, reduced oxygen species (e.g., O$_2^-$) and the electrolyte solvent are also possible, as is shown in Figure 5.S11 below.

Of note, the lack of reaction between LiOH and I$_3^-$ observed here is in contrast to a similar study by Liu et al., who observed slow I$_3^-$ consumption (on the order of 10s of hours to complete consumption) when it was mixed with a DME solution containing 3-6 w% water and an excess of LiOH. The likely explanation for this discrepancy is the different water contents employed in each respective study (3-6 w% in Liu et al., <50 ppm here), as Liu et al. also suggest a fast reaction between LiOH and I$_3^-$ in aqueous solutions, implying that H$_2$O acts to solubilize LiOH or I$_3^-$, allowing each to react more rapidly.
Figure 5.S11. KO₂ reaction in the presence of LiI. To determine the effect of I⁻ in the presence of O₂⁻, a 5 mL solution of DME with 500 ppm H₂O, 0.2 M LiI and 0.1M LiTFSI was prepared, to which 1.5 mmol of KO₂ and 2.5 mmol of 18-crown-6 ether were then added to generate O₂⁻. The contents were left to stir for 2 hours, after which the solid contents were removed and washed. a) The ¹H NMR spectrum obtained from the solids shows the clear presence of decomposition products, as expected from O₂⁻ in the presence of DME. b) After the addition of KO₂ the supernatant, turned slightly orange, and then deepened to dark orange, indicating the presence of I₃⁻. Prepared by Dr. Robert Black, Ivan Kochetkov, and Professor Linda Nazar at the University of Waterloo.
Chapter 6: Considerations for redox mediators in the presence of water, including reference electrode selection

Abstract
In high-capacity lithium-oxygen (Li-O\textsubscript{2}) batteries, lithium peroxide (Li\textsubscript{2}O\textsubscript{2}) forms on discharge in aggregated structures away from the carbon electrode surface. As Li\textsubscript{2}O\textsubscript{2} is insoluble in the most stable, ether-based electrolyte and is an insulator, an electrolyte-soluble redox mediator is required to oxidize Li\textsubscript{2}O\textsubscript{2} on charge and shuttle electrons back to the electrode surface. One promising redox mediator, iodide (I\textsuperscript{-}), has recently been shown to promote Li\textsubscript{2}O\textsubscript{2} degradation to lithium hydroxide (LiOH) in the presence of water (H\textsubscript{2}O), which is problematic because LiOH is not reversibly oxidized to lithium, oxygen, and water on charge. It therefore is important to determine the critical considerations for predicting whether a redox mediator will induce Li\textsubscript{2}O\textsubscript{2} degradation to LiOH in the presence of water impurities, which are likely to be difficult to completely eliminate if lithium-air batteries are practically realized. In this study, we show via galvanostatic cycling of lab-scale Li-O\textsubscript{2} batteries containing LiI and H\textsubscript{2}O in DME and DMSO that DMSO prevents Li\textsubscript{2}O\textsubscript{2} degradation to LiOH. Cyclic voltammetry of these electrolytes shows DMSO exhibits a higher potential for iodide oxidation than DME, implying H\textsubscript{2}O\textsubscript{2} reduction is more difficult in DMSO. Thus, the most critical consideration in predicting Li\textsubscript{2}O\textsubscript{2} degradation to LiOH in the presence of water impurities is a redox mediator’s ability to oxidize and subsequently reduce H\textsubscript{2}O\textsubscript{2} in the electrolyte of choice. We also discuss reference electrode selection for studying redox mediators in Li-O\textsubscript{2} batteries, showing the study of redox mediators unstable to lithium metal – or of redox mediators in solvents unstable to lithium metal – requires the use of a lithium reference electrode with a lower lithium intercalation voltage than the redox mediator but a higher voltage than oxygen reduction.

6.1 Introduction
Redox mediators are likely necessary to enable charging of high capacity nonaqueous lithium-oxygen (Li-O\textsubscript{2}) batteries, as the majority of the dominant discharge product lithium peroxide (Li\textsubscript{2}O\textsubscript{2}) forms away from the electrode surface in aggregated structures. Li\textsubscript{2}O\textsubscript{2} is insoluble in common nonaqueous battery electrolytes and a wide band gap insulator,\textsuperscript{39} such that oxidizing aggregated Li\textsubscript{2}O\textsubscript{2} on charge requires electron shuttling through the electrolyte via a small molecule redox mediator.

As discussed in Chapter 5, iodide (I\textsuperscript{-}) has been studied as a possible redox mediator. However, the combination of lithium iodide and water impurities in the electrolyte leads to the formation of lithium hydroxide (LiOH) as the dominant discharge product, and LiOH is not reversibly oxidized back to its constituent components – lithium, oxygen, and water – on charge. Rather, charge current is carried via a combination of irreversible side reactions and redox shuttling between the electrodes of iodide and its oxidized forms.\textsuperscript{41} Thus, it is important to avoid LiOH formation on discharge.

Recently, Tulodziecki et al. elucidated the mechanism for LiOH formation on discharge in a Li-O\textsubscript{2} battery containing an ether-based electrolyte containing lithium
iodide (LiI) and ppm quantities of added water (H$_2$O). By adding potassium superoxide (KO$_2$), a stable superoxide at ambient temperature and pressure, to an electrolyte of 0.1 M LiTFSI and 0.2 M LiI in 1,2-dimethoxyethane (DME), Tulodziecki et al. showed that rather than the typical four-electron oxygen reduction associated with catalysts like platinum:

$$4\text{Li}^+ + 4e^- + \text{O}_2 + 2\text{H}_2\text{O} \rightarrow 4\text{LiOH} \quad (6.1)$$

the formation of LiOH in an ether-based electrolyte containing LiI and ppm H$_2$O follows the 2 e$^-$/O$_2$ formation of Li$_2$O$_2$, chemical degradation of Li$_2$O$_2$ to LiOH and hydrogen peroxide (H$_2$O$_2$), and H$_2$O$_2$ reduction via I$^-$ oxidation to triiodide (I$_3^-$):

$$2\text{Li}^+ + 2e^- + \text{O}_2 \rightarrow \text{Li}_2\text{O}_2 \quad (6.2)$$

$$\text{Li}_2\text{O}_2 + 2\text{H}_2\text{O} \rightarrow 2\text{LiOH} + \text{H}_2\text{O}_2 \quad (6.3)$$

$$\text{H}_2\text{O}_2 + 3\text{I}^- + 2\text{Li}^+ \rightarrow 2\text{LiOH} + \text{I}_3^- \quad (6.4)$$

$$\text{I}_3^- + 2e^- \rightarrow 3\text{I}^- \quad (6.5)$$

This is an elegant proposal, as it essentially corresponds to an iodometric titration (described in section 2.4) occurring within the cell – water reacting Li$_2$O$_2$ to LiOH and H$_2$O$_2$ followed by iodide reducing H$_2$O$_2$. To be clear, water impurities at these concentrations without iodide do not induce the reaction of Li$_2$O$_2$ to LiOH and H$_2$O$_2$ in DME. Tulodziecki et al. proposed two roles for iodide in causing this cascade of reactions. First, iodide lowers the pKa of dilute H$_2$O, evidenced by a downfield shift in the H$_2$O’s $^1$H NMR shift, increasing the water’s reactivity. Second, iodide reduces H$_2$O$_2$ and is consequently oxidized to triiodide, as the I$^-$/I$_3^-$ redox couple is at a potential only slightly above that of a discharging lithium-oxygen battery. This then enables the subsequent electrochemical reduction of I$_3^-$ back to I$^-$, resulting in an overall process that is 4 e$^-$/O$_2$ as exhibited via DEMS in Chapter 5.

It is important to determine if other redox mediators will cause this same Li$_2$O$_2$ degradation reaction to irreversible LiOH in the presence of water impurities. The lithium metal negative electrode currently used in Li-O$_2$ batteries will likely need to be protected from the electrolyte to prevent dendrite growth and/or to enable the use of organic redox mediator molecules that are unstable to lithium. With the lithium metal protected, some amount of water in the electrolyte may be tolerated in order to enable an air-breathing battery over a system including onboard oxygen storage. Thus, it is important to know the general interaction of H$_2$O and redox mediator molecules. This chapter discusses considerations for Li$_2$O$_2$ degradation to LiOH formation in the presence of water impurities in nonaqueous Li-O$_2$ batteries containing redox mediators. Our work confirms the proposed roles of iodide, with particular emphasis on the ability of iodide to reduce H$_2$O$_2$ leading to complete degradation of Li$_2$O$_2$ to LiOH. A tangential important finding during this study was the difficulty in selection of an appropriate reference electrode for studying redox mediators in Li-O$_2$ batteries. We discuss appropriate reference electrode selection and its importance in proper characterization of redox mediators in Li-O$_2$ cells.
6.2 Experimental methods

**Materials.** Lithium bis(trifluoromethane) sulfonimide (LiTFSI), 1,2-dimethoxyethane (DME), dimethyl sulfoxide (DMSO), Whatman QM-A glass fiber filters, poly(tetrafluoroethylene) (PTFE, 60 wt% dispersion in H$_2$O), Vulcan XC72, stainless steel mesh, research-grade oxygen and argon, lithium metal, lithium chloride (LiCl), deuterium oxide (D$_2$O), lithium nitrate (LiNO$_3$), and lithium bromide (LiBr) were all sourced as described in section 3.2. Acetonitrile (anhydrous, 99.8%), potassium dioxide, and lithium manganese (III,IV) oxide (electrochemical grade) were purchased from Sigma-Aldrich. Lithium iron phosphate-coated aluminum foil, lithium titanate spinel powder, and Super P carbon black were purchased from MTI Corp. Polyvinylidene fluoride (PVDF) was purchased from Solvay. N-methylpyrrolidone was purchased from Alfa Aesar. Water used to contaminate the electrolytes was ultrapure (18.2 MΩ cm) via a Millipore filtration system.

**Electrolyte preparation.** All electrolyte and cell preparation was carried out in an argon glove box with <0.1 ppm O$_2$ and <0.1 ppm H$_2$O. Water was quickly and carefully added to electrolytes in the glovebox via micropipette, with ppm quantities calculated by volume.

**Electrode preparation.** Cathodes of Vulcan XC72 carbon on stainless steel mesh were prepared via the spray-coating method described in Chapter 2. Electrodes of lithium iron phosphate (LiFePO$_4$) on aluminum foil were punched out as 7/16” diameter discs and used as is. Lithium titanate spinel (Li$_4$Ti$_5$O$_{12}$) and lithium manganese oxide (LiMn$_2$O$_4$) electrodes were prepared on stainless steel mesh via a slurry method. Either 11mm or 7/16” diameter stainless steel mesh discs were punched out, rinsed with IPA and acetone, and dried at 150 °C for at least ten minutes. Active material, Super P carbon, and polyvinylidene fluoride (PVDF) (via a 10 wt% solution in n-methylpyrrolidone (NMP)) were combined in a mass ratio of 90:5:5, typically totaling 1 g, with an additional 0.1 g NMP and mixed via a Thinky mixer for five minutes at 1500 rpm. Additional NMP was added with further mixing as needed to create a ketchup-like consistency. The slurry was then spread over the stainless steel mesh discs, which were subsequently dried at 110 °C under vacuum for ten minutes. The electrodes were then calendared to ~60% true density via an MTI rolling press, followed by further drying at 60 °C under vacuum for twelve hours. The electrodes were then transferred into the argon glovebox while hot and stored on a hot plate at 200 °C immediately prior to use. Loading was typically ~10mg for LTO and ~15-25mg for LMO.

**Cell preparation.** Lithium-oxygen batteries following the Swagelok design detailed in Chapter 2 were prepared with lithium metal as the negative electrode, Whatman QM-A as a separator, and Vulcan XC72 carbon on stainless steel mesh as the positive electrode, as detailed in section 3.2. Electrolyte volume was either 80 µL or 160 µL (for which two Whatman QM-A separators were used), as detailed in figure and table captions. Prior to use as the lithium source and reference electrode in Li-O$_2$ batteries, Li$_4$Ti$_5$O$_{12}$ and LiMn$_2$O$_4$ electrodes were prelithiated in cells containing a 7/16” lithium metal negative electrode, a 1/2” Whatman QM-A separator and 80 µL of 1 M LiTFSI in DME under an argon headspace.
Cell characterization. Electrochemical cycling, pressure monitoring, and titrations of discharged cathodes were performed as detailed in Chapter 2. The moles of Li$_2$O$_2$ formed on discharge are given via the iodometric titration. LiOH and basic decomposition products are both measured via the acid-base titration, and thus cannot be distinguished. The value for LiOH and basic decomposition products reported here is for that formed during discharge. LiOH formed upon reaction of Li$_2$O$_2$ with water during the titration protocol has been subtracted out as two times the moles of Li$_2$O$_2$ measured via the iodometric titration. As discussed in Chapter 5, in cells containing 1 M LiTFSI in DME as the electrolyte, which exhibits a 2.0 e$^-$/O$_2$ process on discharge, basic decomposition products are typically 24% of the quantity of Li$_2$O$_2$ formed. In cells with worse e$^-$/O$_2$, such as in 1 M LiTFSI in DMSO, this quantity is higher. Thus, even after subtracting a stoichiometric amount of LiOH formed from the reaction of Li$_2$O$_2$ with water in the titration protocol, a nonzero value for LiOH and basic decomposition products is still expected even when Li$_2$O$_2$ is formed as the dominant discharge product.

KO$_2$ disproportionation. Ex situ analysis of Li$_2$O$_2$ degradation in the presence of iodide and water impurities in various solvents was performed via potassium superoxide (KO$_2$) disproportionation experiments similar to Tulodziecki et al.$^{43}$ In an argon glovebox, electrolytes containing 0.25 M LiTFSI with and without iodide and water impurities were placed in vials. KO$_2$ powder was added to the electrolyte vials, resulting in immediate disproportionation upon contact with Li$^+$ according to:

$$\text{2KO}_2 + 2\text{Li}^+ \leftrightarrow \text{Li}_2\text{O}_2 + \text{O}_2(\text{g}) + 2\text{K}^+ \quad (6.6)$$

with the added KO$_2$ massed to be the limiting reagent. Tulodziecki et al. showed this in situ formation of Li$_2$O$_2$ enables fast degradation of Li$_2$O$_2$ to LiOH by the combination of iodide and ppm water in DME, according to reactions 6.3-6.5.

X-ray diffraction. Powder X-ray diffraction was performed on the powder formed via KO$_2$ disproportionation via Bruker D2 phaser PXRD with CuKα radiation (λ = 1.54178 Å). To prepare the powder for XRD, the liquid was decanted and the powder rinsed three times with pure DME before drying in the glovebox antechamber.

Nuclear magnetic resonance spectroscopy. $^1$H nuclear magnetic resonance spectroscopy measurements were performed on both a Bruker AM-400 magnet with a 5mm Z-gradient broad band probe and a Bruker Avance III 500 MHz magnet with a 5mm Z-gradient broad band probe. Each electrolyte was placed in an NMR tube along with a sealed melting point capillary containing 99.9% D$_2$O. The D$_2$O was used as a deuterium lock, and its residual proton peak was used as a reference for the electrolyte’s $^1$H spectrum.

6.3 Results and discussion
6.3.1 Considerations for Li$_2$O$_2$ degradation to LiOH

To probe the proposed roles of iodide in causing the degradation of Li$_2$O$_2$ to LiOH in the presence of water impurities in nonaqueous Li-O$_2$ batteries, 1,2-dimethoxyethane (DME) typically chosen for stability,$^{31}$ was replaced with dimethyl sulfoxide (DMSO), a solvent with a higher dielectric constant and higher Gutmann Donor Number and Acceptor Number.$^{85}$ Table 6.1 gives the e$^-$/O$_2$ and titration values for typical
lab-scale Li-O\textsubscript{2} batteries containing 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H\textsubscript{2}O in pure DME, pure DMSO, and mixtures of the two by volume discharged at a constant current of 0.2 mA/cm\textsuperscript{2}. As the amount of DMSO increases, the batteries trend from a 4 e\textsuperscript{−}/O\textsubscript{2} process to a 2 e\textsuperscript{−}/O\textsubscript{2} process on discharge, and the discharge product trends from predominant LiOH to predominant Li\textsubscript{2}O\textsubscript{2}, along with expected basic decomposition products. Of note, 6.8 \textmu mol Li\textsubscript{2}O\textsubscript{2} after a 0.5 mAh discharge corresponds to a 73% yield of an ideal 2.0 e\textsuperscript{−}/Li\textsubscript{2}O\textsubscript{2} process (9.3 \textmu mol Li\textsubscript{2}O\textsubscript{2} after 0.5 mAh), in agreement with Li\textsubscript{2}O\textsubscript{2} yields in 1 M LiTFSI in DMSO.\textsuperscript{32} These data indicate that DMSO disrupts iodide’s role in the degradation of Li\textsubscript{2}O\textsubscript{2} to LiOH in the presence of water.

**Table 6.1** Electron per oxygen values during discharge and titration values after discharge for cells containing 160 \textmu L of 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H\textsubscript{2}O in mixtures of DME and DMSO. The electrolyte volume and concentration of water correspond to sufficient water for 0.95 mAh of LiOH formation. Discharging was stopped at the labeled capacity.

<table>
<thead>
<tr>
<th>Solvent</th>
<th>e\textsuperscript{−}/O\textsubscript{2}</th>
<th>Capacity (mAh)</th>
<th>LiOH (and basic decomposition products) (\textmu mol)</th>
<th>Li\textsubscript{2}O\textsubscript{2} (\textmu mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DME</td>
<td>4.1</td>
<td>0.3</td>
<td>10.6</td>
<td>0.1</td>
</tr>
<tr>
<td>1:15 (v:v) DMSO:DME</td>
<td>3.4</td>
<td>0.5</td>
<td>14.9</td>
<td>1.4</td>
</tr>
<tr>
<td>50:50 (v:v) DMSO:DME</td>
<td>2.3</td>
<td>0.5</td>
<td>5.2</td>
<td>6.0</td>
</tr>
<tr>
<td>DMSO</td>
<td>2.2</td>
<td>0.5</td>
<td>3.8</td>
<td>6.8</td>
</tr>
</tbody>
</table>

Confirming the stability of Li\textsubscript{2}O\textsubscript{2} in the presence of LiI and H\textsubscript{2}O in DMSO, Figure 6.1 shows oxygen consumption and gas evolution data via pressure monitoring for Li-O\textsubscript{2} batteries containing 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H\textsubscript{2}O in the labeled mixtures of DME and DMSO, galvanostatically discharged and charged at 0.25 mA/cm\textsuperscript{2}. The trend toward a 2 e\textsuperscript{−}/O\textsubscript{2} process on discharge coincides with increased rechargeability on charge, as observed by the larger amount of gas evolved during charge (noting that prior reports have shown that most gas evolved in a DMSO-based Li-O\textsubscript{2} cell is oxygen). As an additional screening, ex situ product formation analysis similar to that in Tulodziecki et al. was conducted in an argon glovebox. KO\textsubscript{2} was added to lithium-containing electrolytes. Figure 6.2 shows the X-ray diffraction spectra of the dried powders that precipitated from this KO\textsubscript{2}/electrolyte mixture. The powder formed in 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H\textsubscript{2}O in DMSO shows Li\textsubscript{2}O\textsubscript{2} signatures, similar to that formed in 0.25 M LiTFSI in DME. The powder from 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H\textsubscript{2}O in DME shows a distinctly different XRD spectrum, with unassigned signatures similar to, though not exactly, LiOH and LiIO\textsubscript{3}.\textsuperscript{41, 109} Additionally, the electrolyte in the vial containing 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H\textsubscript{2}O in DME turned yellowish orange, indicating the formation of triiodide via reaction 6.4, as described in Tulodziecki et al.,\textsuperscript{43} while the solution in the vial containing 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H\textsubscript{2}O in DMSO did not exhibit a color change.
Figure 6.1 a) and c) Galvanostatic cycling of lithium-oxygen cells containing 160 µL of 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H₂O in mixtures of DMSO and DME in ratios by volume of 1:15 and 50:50, respectively. b) and d) Oxygen consumption and gas evolution from pressure decay and pressure rise measurements.

Figure 6.2 X-ray diffraction of powders formed by adding potassium superoxide as the limiting reagent to vials of lithium-containing electrolytes. Electrolytes were i) 0.25 M LiTFSI in DME, ii) 0.25 M LiTFSI 0.2 M LiI 2000 ppm H₂O in DME and iii) 0.25 M LiTFSI 0.2 M LiI 2000 ppm H₂O in DMSO. * denotes peaks expected for Li₂O₂, as seen in recent reports.⁴³,¹⁰⁹
DMSO’s effect on the iodide-induced degradation of Li$_2$O$_2$ was studied through the two roles of iodide presented by Tulodziecki et al.: 1) lowering the pKa of H$_2$O and 2) chemical equilibrium shift due to the continuous consumption of H$_2$O$_2$.\textsuperscript{43}

The relative pKa of H$_2$O was observed via $^1$H NMR. Electrolytes of DME, DMSO, and acetonitrile (MeCN) – a high dielectric constant, low Gutmann Donor Number and low Gutmann Acceptor Number solvent – with and without 0.2 M LiI and 2000 ppm H$_2$O were placed inside NMR tubes along with sealed melting point capillaries containing 99.9% D$_2$O. Figure 6.3 shows the $^1$H NMR spectra, referenced to the residual proton peak (assigned 4.7 ppm in each spectra) in the D$_2$O capillary.

![Figure 6.3](image)

**Figure 6.3** $^1$H NMR of labeled electrolytes. Peak at 4.7 ppm is from residual protons in 99.9% D$_2$O in a sealed melting point capillary. Arrows point to peaks assigned to ppm H$_2$O in the electrolytes. For spectrum (i), these two peaks align, as labeled.

A large H$_2$O downfield shift, i.e., a strong deshielding of the proton or lowering of H$_2$O pKa, was observed with the addition of LiI in DME (comparing spectrum i vs. spectrum ii). In both acetonitrile and DMSO, however, the addition of LiI also moves the H$_2$O shift further downfield (iv vs. v and vii vs. viii respectively). Though the shifts are not as far downfield as in DME (i vs. ii), both DMSO and MeCN exhibit an iodide-induced lower pKa of H$_2$O, and thus predict increased LiOH formation. However, as the pressure rise and powder XRD results above show, Li$_2$O$_2$ is the predominant discharge product in 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H$_2$O in DMSO. Additionally, in the absence of LiI, the proton shift of 2000 ppm H$_2$O is similar in the three solvents, with the shift in DME (spectrum ii) slightly further downfield than that in DMSO (viii) and MeCN (v). This
might indicate more LiOH formation in the DME electrolyte, but previous literature suggests water impurities in DMSO and acetonitrile induce more LiOH and/or lithium oxyhydroxide (LiOOH) formation than water impurities in DME, according to spectroscopic characterization of discharged cathodes.\textsuperscript{110-112} Thus, the work herein indicates an iodide-induced lower pKa of H$_2$O is not the most important consideration for iodide-induced Li$_2$O$_2$ degradation.

Turning to the ability of iodide to continuously reduce H$_2$O$_2$, it is well known in the literature that iodide oxidation, i.e., triiodide and iodine (I$_2$) formation, is highly dependent on solvent.\textsuperscript{113-115} To probe this solvent effect on iodide oxidation experimentally, cyclic voltammetry, beginning with an oxidative scan, was performed on batteries under an argon atmosphere containing 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H$_2$O in DME and DMSO. Shown in Figure 6.4, iodide oxidation in DMSO occurs about 0.5 V vs. Li/Li$^+$ higher than in DME.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure6.4.png}
\caption{Cyclic voltammetry of batteries under an argon atmosphere containing 160 µL of 0.25 M LiTFSI, 0.2 M LiI, and 2000 ppm H$_2$O in DME and DMSO as labeled. Sweep rate was 20 mV/s and CVs began with oxidative scan.}
\end{figure}

Thus, iodide oxidation is more difficult in DMSO than in DME. While this is an electrochemical measurement, it points to a less favorable driving force for triiodide formation in DMSO, implying iodide-induced H$_2$O$_2$ reduction is more difficult in DMSO than in DME. This serves as an explanation for the permanence of Li$_2$O$_2$ as the predominant discharge product in LiI- and H$_2$O-containing electrolytes with DMSO as the solvent. Of note, this also correlates with the previously described observation that KO$_2$ disproportionation in LiI- and H$_2$O-containing DME is coupled with the electrolyte turning yellowish orange, evident of iodide oxidation, whereas KO$_2$ disproportionation in LiI- and H$_2$O-containing DMSO exhibits no color change. The higher oxidation potential of iodide in DMSO is likely linked to stronger solvation of iodide due to its higher Gutmann Acceptor Number than DME.\textsuperscript{85}
Table 6.2 Electron per oxygen values during discharge and titration values after discharge for cells containing 160 µL of 0.25 M LiTFSI and 0.2 M LiBr with and without 2000 ppm H₂O in DME. The electrolyte volume and concentration of water correspond to sufficient water for 0.95 mAh of LiOH formation. Discharging was stopped at the labeled capacity.

<table>
<thead>
<tr>
<th>Electrolyte</th>
<th>e/O₂</th>
<th>Capacity (mAh)</th>
<th>LiOH (and basic decomposition products) (µmol)</th>
<th>Li₂O₂ (µmol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25M LiTFSI, 0.2M LiBr in DME</td>
<td>2.2</td>
<td>0.3</td>
<td>2.8</td>
<td>4.1</td>
</tr>
<tr>
<td>0.25M LiTFSI, 0.2M LiBr, 2000ppm H₂O in DME</td>
<td>2.3</td>
<td>0.3</td>
<td>3.1</td>
<td>4.1</td>
</tr>
</tbody>
</table>

In further support that DMSO precludes H₂O₂ reduction via more difficult iodide oxidation, electrolytes of lithium bromide (LiBr) in DME were similarly studied. Bromide oxidation is well known to occur at higher potentials than iodide oxidation in common nonaqueous electrolytes. (Cyclic voltammetry was not performed in our laboratory to avoid bromine formation.) As expected given the higher oxidation potential of Br⁻ than I⁻, Table 6.2 shows that LiBr- and H₂O-containing DME-based electrolytes exhibit near 2 e⁻/O₂ processes on discharge. Titrations of discharged cathodes also point to Li₂O₂ as the dominant discharge product, along with basic decomposition products as expected, as 4.1 µmol Li₂O₂ after a 0.3 mAh discharge corresponds to a 73% yield of an ideal 2.0 e⁻/Li₂O₂ process (5.6 µmol Li₂O₂ after 0.3 mAh).

Figure 6.5 H NMR of labeled electrolytes. Peak at 4.7 ppm is from residual protons in 99.9% D₂O in a sealed melting point capillary. Arrows point to peaks assigned to ppm H₂O in the electrolytes.
Meanwhile, $^1$H NMR spectra for electrolytes with 1500 ppm H$_2$O with and without LiBr in DME, presented in Figure 6.5, show that the addition of LiBr to DME causes a nearly identical downfield shift in the ppm H$_2$O peak as with the addition of LiI. For reference, lithium nitrate (LiNO$_3$) does not induce the same downfield shift in the ppm H$_2$O peak, as shown in Figure 6.5. Thus, it seems the most critical consideration in predicting Li$_2$O$_2$ degradation to LiOH in the presence of water impurities is an additive’s ability to oxidize and subsequently reduce H$_2$O$_2$ in the electrolyte of choice, rather than the additive’s effect on the pKa of the dilute water.

$6.3.2$ Implications of redox mediators on reference electrode selection

A tangential important finding during this study was the difficulty in selection of an appropriate reference electrode for studying redox mediators in lithium-oxygen batteries.

In studying the effect of other solvents and other redox mediators on the degradation of Li$_2$O$_2$ to LiOH in lab-scale batteries, the use of a pure lithium metal anode limits the options. Organic redox mediators degrade quickly against lithium metal,$^{95,102}$ and some interesting solvents do not form a stable interface on lithium, such as acetonitrile.$^{31}$

Lithium iron phosphate (LiFePO$_4$, referred to as LFP), is commonly used as an alternative lithium source and reference electrode in lithium-oxygen studies, especially with acetonitrile.$^{31,45}$ LFP exhibits reversible lithium intercalation near 3.5 V vs. Li/Li$^+$ with a stable voltage profile due to its two-phase behavior:

$$\text{Li}^+ + e^- + \text{FePO}_4 \leftrightarrow \text{LiFePO}_4 \quad U \sim 3.5 \text{ V vs. Li}^+/\text{Li} \quad (6.7)$$

Unfortunately, LFP’s standard reduction potential for lithium intercalation is too high for use with iodide.

![Figure 6.6 Vials containing 0.25 M LiTFSI and 0.2 M LiI in DME under an argon atmosphere and (left to right) no electrode, a lithium iron phosphate electrode, a lithiated lithium manganese oxide electrode, and a lithiated lithium titanate spinel electrode. The LiFePO$_4$ was delithiated 50% in a cell containing 1 M LiTFSI in DME and an argon headspace prior to insertion in the vial. The LMO and LTO were 50% lithiated in cells containing 1 M LiTFSI in DME and argon headspaces prior to insertion in the vial.](image-url)
As iodide oxidation occurs near 3 V vs. Li/Li$^+$ in DME, contact between an electrolyte of LiI in DME and an LFP electrode causes immediate chemical lithiation of the LFP and oxidation of the iodide. Figure 6.6 shows vials containing 0.25 M LiTFSI and 0.2 M LiI in DME under an argon atmosphere, with the second vial from the left containing an LFP electrode that has been previously delithiated 50% in a battery containing 1 M LiTFSI in DME under argon. The electrolyte immediately has turned orange, indicating chemical oxidation of iodide, likely to iodine. This ongoing chemical reaction at the reference electrode complicates the use of LFP to study iodide redox mediation at the carbon/oxygen working electrode in a Li-O$_2$ battery. LFP should be avoided as a reference electrode in Li-O$_2$ batteries containing redox mediators with redox potentials below that of lithium intercalation in LFP, as has been recently reported.$^{117}$

While chemical oxidation of the redox mediator provides a ceiling for possible lithium intercalation materials as reference electrodes in Li-O$_2$ batteries, spontaneous oxygen reduction provides a floor. As exhibited by the desired discharge reaction in the Li-O$_2$ battery, oxygen reduction occurs ~2.8 V vs. Li/Li$^+$ in DME. Lithium metal reacts spontaneously with oxygen, but forms a semi-stable passivating layer, allowing a lithium-oxygen battery employing a typical 1 M LiTFSI in DME electrolyte to rest stably under oxygen for hours. Some lower-voltage lithium intercalation materials, while avoiding chemical oxidation of the redox mediator, react spontaneously under oxygen. For example, lithium titanate spinel (Li$_4$Ti$_5$O$_{12}$, referred to as LTO), exhibits reversible lithium intercalation near 1.55 V vs. Li/Li$^+$ in DME with a stable voltage profile, forming lithiated lithium titanate spinel (Li$_7$Ti$_5$O$_{12}$, referred to as LLTO):\[
3\text{Li}^+ + 3e^- + \text{Li}_4\text{Ti}_5\text{O}_{12} \leftrightarrow \text{Li}_7\text{Ti}_5\text{O}_{12} \quad U \sim 1.55 \text{ V vs. Li}^+$/Li $$(6.8)$$

Partially delithiated LLTO is stable against iodide oxidation, as shown in the rightmost vial of Figure 6.6, where there is no color change of a 0.25 M LiTFSI and 0.2 M LiI in DME electrolyte upon insertion of the partially delithiated LLTO. However, LLTO reacts spontaneously under oxygen, losing nearly 80% state-of-charge as it undergoes the reaction:\[
\text{Li}_7\text{Ti}_5\text{O}_{12} + \text{O}_2 \leftrightarrow \text{Li}_2\text{O}_2 + \text{Li}_5\text{Ti}_5\text{O}_{12} \quad (6.9)
\]
perhaps through a lithium superoxide (LiO$_2$) intermediate, as shown by Piana et al.$^{118}$ Figure 6.7a shows a battery containing LLTO versus lithium metal with an O$_2$ headspace resting at open circuit for 10 hours before constant current charging (delithiating), discharging (lithiating), and charging again between 2 V vs. Li/Li$^+$ (charge) and 1.1 V (discharge) cutoffs. Pressure monitoring shows O$_2$ is spontaneously consumed throughout the 10 hour rest period, and only ~0.43 mAh – a fraction of the second charge – indicated with an arrow in Figure 6.7a, occurs with both a stable voltage profile and a stable pressure. Interestingly, electrochemical formation of Li$_2$O$_2$ on the surface of the lithium titanate spinel electrode seems to extend the usable capacity. Figure 6.7b shows a similar battery charged to a 4.5 V cutoff before a subsequent discharge and charge. The usable charge capacity with stable voltage and stable pressure, as indicated by the arrow, is ~0.83 mAh. Stability in both voltage and pressure when charging the electrode versus
lithium under oxygen is necessary to use the electrode as a lithium source and reference electrode in a lithium-oxygen battery.

**Figure 6.7** Galvanostatic cycling with pressure monitoring of lithiated lithium titanate spinel versus lithium in cells containing 1 M LiTFSI in DME and oxygen headspaces. Lithium titanate spinel electrodes were prelithiated versus lithium in cells containing 1 M LiTFSI in DME and argon headspaces before use in the exhibited cells. Labeled sections on the graphs correspond to i) open circuit for 10 hr, ii) charging at ~0.2 mA/cm$^2$ to a) 2 V and b) 4.5 V, iii) open circuit for 0.5 hr, iv) discharging at ~0.2 mA/cm$^2$ to a) 2 V and b) 4.5 V, v) open circuit for 0.5 hr, vi) charging at ~0.2 mA/cm$^2$ to a) 2 V and b) 3.6 V (arbitrary), and vii) open circuit for 0.5 hr. The cell in b) was quickly refilled with oxygen at ~42 hr, resulting in the sudden increase in pressure. The arrows point to the usable capacity as a lithium reference electrode with stable voltage and stable pressure: a) ~0.43 mAh and b) ~0.84 mAh.

Between the two challenges of chemical oxidation of the redox mediator and spontaneous oxygen reduction, a possible mid-voltage option for a lithium reference electrode is lithium manganese oxide (LiMn$_2$O$_4$, referred to as LMO). LMO has been
shown to withstand overlithiation of ~0.7 Li$^+$ to Li$_{1.7}$Mn$_2$O$_4$ at ~3 V vs. Li/Li$^+$ in carbonates, presenting a lithium intercalation voltage slightly above oxygen reduction.

$$0.7\text{Li}^+ + 0.7\text{e}^- + \text{LiMn}_2\text{O}_4 \leftrightarrow \text{Li}_{1.7}\text{Mn}_2\text{O}_4 \quad U \sim 3 \text{ V vs. Li}^+$/Li$$

Figure 6.8a shows the galvanostatic discharge profile and pressure data for a battery containing LLMO versus lithium metal with an O$_2$ headspace resting at open circuit for 10 hours before charging (delithiating), discharging (lithiating), and charging again at constant current between 3.6 V vs. Li/Li$^+$ (charge) and 2.7 V (discharge) cutoffs.

**Figure 6.8** a) Galvanostatic cycling with pressure monitoring of lithiated lithium manganese oxide versus lithium in cells containing 1 M LiTFSI in DME and oxygen headspaces. Lithium manganese oxide electrodes were prelithiated versus lithium in cells containing 1 M LiTFSI in DME and argon headspaces before use in the exhibited cells. Labeled sections on the graphs correspond to i) open circuit for 10 hr, ii) charging at ~0.076 mA/cm$^2$ to 3.6 V vs. Li/Li$^+$, iii) open circuit for 0.25 hr, iv) discharging at ~0.076 mA/cm$^2$ to 2.7 V, v) open circuit for 0.25 hr, vi) charging at ~0.076 mA/cm$^2$ to 3.6 V (arbitrary), and vii) open circuit for 0.25 hr. b) Comparison of the pressure of this cell with the ambient temperature, measured via a thermocouple near the cell exterior.
This electrode shows 1.7 mAh of delithiation capacity with a stable voltage plateau before a sudden voltage rise on the first charge, as well as repeatability on the next cycle. Pressure monitoring of the cell shows a pressure decay of 0.5 torr/hr, with fluctuations due to temperature as shown in Figure 6.8b. While this is not ideal for use as a lithium reference electrode, its steady nature means it could be accounted for, and a 2.0 e-/O₂ process and a 4.0 e-/O₂ process in this cell would consume 13.4 torr/hr and 6.7 torr/hr, respectively, or more than ten times this consumption. Unfortunately, additionally, an electrolyte of 0.25 M LiTFSI and 0.2 M LiI in DME shows a color change to faint yellow with the addition of 50% delithiated LLMO electrode, shown in the second vial from the right in Figure 6.6. The partially delithiated electrode reaches an open circuit potential just above 3 V vs. Li/Li⁺, and iodide to triiodide onset, as shown in Figure 5.2, occurs at ~2.95 V vs. Li/Li⁺. Thus, LLMO might work best as a lithium source and reference electrode for redox mediators with higher redox potentials than iodide, such as tetrathiafulvalene (TTF), which exhibits a redox couple at ~3.5 V vs. Li/Li⁺.

6.4 Conclusion

In conclusion, the most critical consideration in predicting Li₂O₂ degradation to LiOH in the presence of water impurities is an additive’s ability to oxidize and subsequently reduce H₂O₂ in the electrolyte of choice. Galvanostatic cycling of lab-scale Li-O₂ batteries containing LiI and H₂O in DME and DMSO shows that DMSO prevents Li₂O₂ degradation to LiOH, and cyclic voltammetry of these electrolytes show DMSO exhibits a higher potential for iodide oxidation than DME, implying H₂O₂ reduction is more difficult in DMSO. As the low pKa of dilute water in DMSO via ¹H NMR would predict LiOH formation, an additive’s effect on the pKa of the dilute water likely plays a minor role in final product formation.

The use of a lithium metal negative electrode limits the options of solvents and redox mediators for studying the degradation of Li₂O₂ to LiOH, so we investigated other possible lithium reference electrodes for use in lab-scale Li-O₂ batteries. High voltage lithium intercalation materials like lithium iron phosphate are unsuitable with most redox mediators, as their lithium intercalation voltages are higher than the redox potentials of most Li-O₂ redox mediators, leading to chemical oxidation of the redox mediator upon contact with the intercalation material. Low voltage lithium intercalation materials like lithiated lithium titanate spinel are unstable to oxygen, though after significant exposure to O₂, LLTO can provide some usable capacity – a stable voltage profile and stable O₂ headspace pressure – for use in a lab-scale Li-O₂ battery. Overlithiated lithium manganese oxide is a possible mid-voltage option that is stable to O₂, though it exhibits slow oxygen consumption, and its intercalation voltage (~3 V vs. Li⁺/Li) is slightly too high for use with iodide.

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Chapter 7: Summary

In this work, the nonaqueous electrolyte of the lithium-oxygen (Li-O_2) battery was engineered toward achieving a high-capacity, rechargeable battery.

Toward inducing a high-capacity solution mechanism of lithium peroxide (Li_2O_2) growth, the typical lithium battery salt lithium bis(trifluoromethane) sulfonimide (LiTFSI) was partially exchanged for lithium nitrate (LiNO_3), as NO_3^- is a stronger electron donor than TFSI^- and therefore provides enhanced stability of Li^+, and hence the superoxide intermediate, in the electrolyte. Li-O_2 batteries employing electrolytes of both LiNO_3 and LiTFSI in 1,2-dimethoxyethane (DME), totaling 1.0 M lithium cation (Li^+) concentration, displayed increasing capacity with increasing LiNO_3 concentration. Titrations of cathodes extracted from discharged batteries confirmed Li_2O_2 formation as the dominant discharge process, and scanning electron microscopy (SEM) of discharged cathodes showed increasing toroid formation with increasing LiNO_3 concentration, confirming the solution mechanism. ^7Li nuclear magnetic resonance (NMR) spectroscopy was used to attribute this increase in the solution mechanism to enhanced Li^+ stability in solution, which in turn induces solubility of lithium superoxide (LiO_2), the intermediate to Li_2O_2 formation. In addition, an Ising model was developed with collaborators to predict Li^+ stability as a function of its solvation shell composition. Differential electrochemical mass spectrometry (DEMS), via a custom-built instrument, was used to show the addition of LiNO_3 did not deleteriously affect battery rechargeability.

As anion selection induces the solution mechanism by lowering the free energy of Li^+ in solution, non-Li alkali metal cations and alcohols were studied as methods of inducing the solution mechanism by lowering the free energy of the superoxide anion (O_2^-) in solution. The addition of non-Li alkali metal salts induced a minimal increase in the achievable discharge capacity of Li-O_2 batteries. Gas analysis of the addition of lithium salt to a sodium-oxygen battery showed Li^+ quickly scavenges any non-Li alkali metal cation-associated O_2^- in electrolytes containing both Li^+ and non-Li alkali metal cations. This corroborates a recent report that a capacity increase is only expected at high currents when oxygen reduction occurs quickly enough that some O_2^- temporarily avoids Li^+ and solubilizes as non-Li alkali metal cation-associated O_2^- The addition of ppm quantities of methanol, ethanol, or 1-propanol to ether-based Li-O_2 battery electrolytes induced a two-fold increase in battery capacity with little trend in the capacity as a function of the additive’s Acceptor Number, highlighting the complexity of interactions between the constituent species in an electrolyte in terms of their Lewis basicities, Lewis acidities, and other physicochemical properties.

As the high-capacity solution mechanism of Li_2O_2 causes some Li_2O_2 to form in aggregated structures away from the electrode surface, the addition of small molecule redox mediators to the electrolyte was studied as a means of charging Li_2O_2. The effect of water (H_2O) contamination in the electrolyte on the promising redox mediator lithium iodide (LiI) was studied. DEMS and titrations of cathodes extracted from discharged batteries confirmed recent reports that lithium hydroxide (LiOH) formed as the dominant discharge product via a 4 e^-/O_2 process. However, isotopic labeling and DEMS were used to show LiOH is not reversibly oxidized back to its reactants (Li^+, O_2, H_2O). With the help of collaborators, titrations, SEM energy-dispersive X-ray spectroscopy (SEM EDX), and galvanostatic cycling of batteries under an argon atmosphere were used to show...
charge current in batteries containing both LiI and H₂O is a complex mixture of side reactions and redox shuttling.

With LiOH identified as an undesirable side product, the mechanism for Li₂O₂ degradation to LiOH in the presence of LiI and H₂O was studied. Galvanostatic cycling of lab-scale Li-O₂ batteries containing LiI and H₂O in DME and dimethyl sulfoxide (DMSO) showed that DMSO prevents Li₂O₂ degradation to LiOH. Cyclic voltammetry of these electrolytes showed DMSO exhibits a higher potential for iodide oxidation than DME, implying iodide-mediated H₂O₂ reduction is more difficult in DMSO. This points to the ability of an additive to reduce H₂O₂ as a key consideration in Li₂O₂’s stability in the presence of an additive and water impurities.

As some redox mediators and solvents are unstable with a lithium metal negative electrode, other possible lithium-containing electrodes were investigated for use in lab-scale Li-O₂ batteries. Placement of a lithium iron phosphate (LFP) electrode in a vial of an iodide-containing electrolyte showed a color change attributed to chemical oxidation of iodide to triiodide, indicating LFP’s lithium intercalation voltage is too high for stable use in a LiI-containing Li-O₂ battery. Meanwhile, DEMS of lithiated lithium titanate spinel (LLTO) showed spontaneous oxygen consumption, indicating its lithium intercalation voltage is too low for use in Li-O₂ batteries. A possible mid-voltage option was shown to be lithiated lithium manganese oxide, though it exhibits slow oxygen consumption, and its intercalation voltage (~3 V vs. Li⁺/Li) is better suited for redox mediators with slightly higher first oxidation potentials than iodide.
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