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
Hybrid Polymer Electrolyte for Lithium-Oxygen Battery Application

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FLORIDA INTERNATIONAL UNIVERSITY

Miami, Florida

HYBRID POLYMER ELECTROLYTE FOR LITHIUM-OXYGEN BATTERY
APPLICATION

A dissertation submitted in partial fulfillment of the

requirements for the degree of

DOCTOR OF PHILOSOPHY

in

MATERIALS SCIENCE AND ENGINEERING

by

Amir Chamaani

2017

To: Dean John Volakis
College of Engineering and Computing

This dissertation, written by Amir Chamaani, and entitled Hybrid Polymer Electrolyte for Lithium-Oxygen Battery Application, having been approved in respect to style and intellectual content, is referred to you for judgment.

We have read this dissertation and recommend that it be approved.

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Date of Defense: October 02, 2017

The dissertation of Amir Chamaani is approved.

Dean John Volakis
College of Engineering and Computing

Andrés G. Gil
Vice President for Research and Economic Development
and Dean of the University Graduate School

Florida International University, 2017

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DEDICATION

To my lovely wife, parents and parents-in-law,
without their love, patience and support, none of this would have been possible.

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ABSTRACT OF THE DISSERTATION
HYBRID POLYMER ELECTROLYTE FOR LITHIUM-OXYGEN BATTERY
APPLICATION

by

Amir Chamaani

Florida International University, 2017

Miami, Florida

Professor Bilal El-Zahab, Major Professor

The transition from fossil fuels to renewable resources has created more demand for energy storage devices. Lithium-oxygen (Li-O₂) batteries have attracted much attention due to their high theoretical energy densities. They, however, are still in their infancy and several fundamental challenges remain to be addressed. Advanced analytical techniques have revealed that all components of a Li-O₂ battery undergo undesirable degradation during discharge/charge cycling, contributing to reduced cyclability. Despite many attempts to minimize the anode and cathode degradation, the electrolyte remains as the leading cause for rapid capacity fading and poor cyclability in Li-O₂ batteries.

In this dissertation, composite gel polymer electrolytes (cGPEs) consisting of a UV-curable polymer, tetraglyme based electrolyte, and glass microfibers with a diameter of ~1 μm and an aspect ratio of >100 have been developed for their use in Li-O₂ battery application. The Li-O₂ batteries containing cGPEs showed superior charge/discharge cycling for 500 mAh.g⁻¹ cycle capacity with as high as 400% increase in cycles for cGPE over gel polymer electrolytes (GPEs). Results using *in-situ* electrochemical impedance spectroscopy (EIS), Raman spectroscopy, and scanning electron microscopy revealed that the source of the

improvement was the reduction of the rate of lithium carbonates formation on the surface of the cathode. This decrease in formation rate afforded by cGPE-containing batteries was possible due to the decrease of the rate of electrolyte decomposition. The increase in solvated to the paired Li^+ ratio at the cathode, afforded by increased lithium transference number, helped lessen the probability of superoxide radicals reacting with the tetraglyme solvent. This stabilization during cycling helped prolong the cycling life of the batteries. The effect of ion complexes on the stability of liquid glyme based electrolytes with various lithium salt concentrations has also been investigated for Li-O_2 batteries. Charge/discharge cycling with a cycle capacity of $500 \text{ mAh}\cdot\text{g}^{-1}$ showed an improvement as high as 300% for electrolytes containing higher lithium salt concentrations. Analysis of the Raman spectroscopy data of the electrolytes suggested that the increase in lithium salt concentration afforded the formation of cation-solvent complexes, which in turn, mitigated the tetraglyme degradation.

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1 INTRODUCTION

1.1 Current state-of-the-art Li-ion batteries

A battery is an electrochemical energy storage device which converts chemical energy into electrical energy. Chemical energy exists inside the electrodes of batteries and converts to the electrical energy via electrochemical reactions known as reduction/oxidation (redox) reactions. Batteries (cells) consist of three major components: a positive electrode (cathode), a negative electrode (anode), and an electrolyte. Figure 1-1 shows a typical representation of Li-ion batteries which consists of graphite as anode, lithium cobalt oxide as a cathode and a porous polymeric separator which electronically isolates the cathode and anode [1].

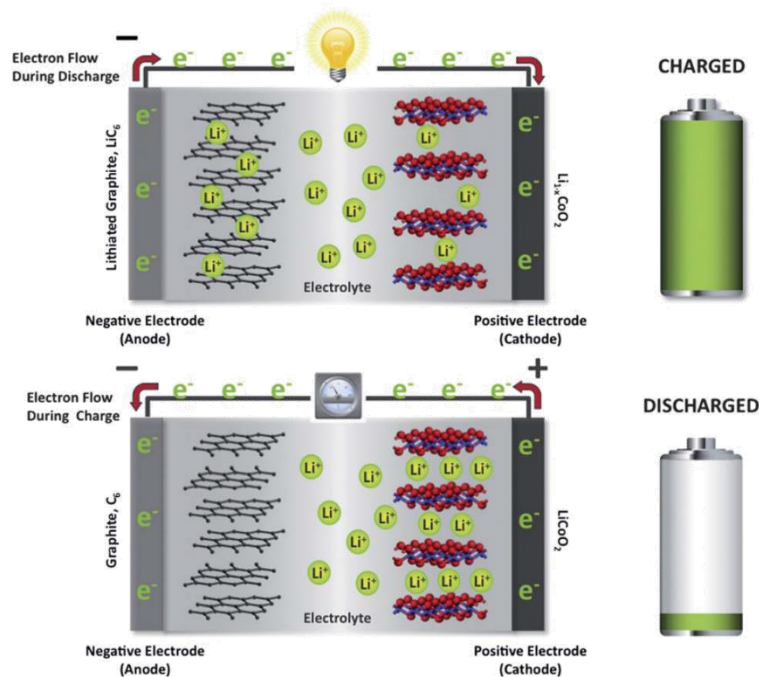


Figure 1-1: Schematic illustration of Li-ion battery consist of graphite as an anode and lithium cobalt oxide ($LiCoO_2$) as a cathode separated by a porous polymeric separator and filled with liquid electrolyte [1]

Each Li-ion cell is also filled with liquid electrolytes which convey the Li^+ between cathode and anode during cell operation. In Li-ion batteries, the typical electrolyte is a solution of LiPF_6 salt dissolved in an organic carbonate-based solvent mixture of ethylene carbonate (EC), dimethyl carbonate (DMC), and diethyl carbonate (DEC) [2]. During the charge process, Li^+ ions travel from the cathode to the anode and intercalate into a graphite anode crystal structure and electrons transfer from the cathode to anode through an external circuit. During discharge, the direction of Li^+ ions and electrons is reversed, and the electron moves from anode to cathode in the external circuit and delivers electricity. The reversible electrochemical reactions of electrodes (cathode and anode) happening during charge and discharge of a Li-ion battery involves the intercalation and deintercalation of Li^+ ions into and from the lattice sites of the anode and cathode active materials and can be expressed as follows [2]:

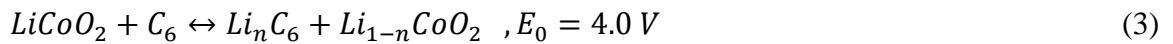
Cathode:



Anode:



Overall cell reaction:



Where, the value of n is about 0.6.

The standard free energy formation (ΔG_0) of the overall cell reaction can be related to the standard potential (E_0) of $\text{C}_6/\text{LiCoO}_2$ cell via the Nernst relation [2]:

$$\Delta G_0 = -nFE_0 \quad (4)$$

Where the F is the faradic constant, and n is the number of electrons involved the electrochemical reactions.

The specific energy of a Li-ion battery is usually expressed in Wh·kg⁻¹ as follows [2,3]:

$$\text{Specific Energy (Wh·kg}^{-1}\text{)} = \frac{\text{Cell Voltage (V)} \times 1000 \text{ (g)} \times 26.8 \text{ Ah}}{\text{kg} \times \text{equiv wt (g)}} \quad (5)$$

Where V is the cell voltage and 26.8 Ah is the Faraday constant. The equivalent weight of an electrochemical redox couple is obtained by dividing the weight of the electrode reactants (in grams) by the number of electrons transferred in the reaction. The energy density (Wh·L⁻¹) of Li-ion batteries could also be calculated from a formula (5) by using the equivalent volumes of the electrode reactants. Table 1-1 demonstrates the specific energy and energy density of common rechargeable batteries, including Li-ion batteries [3,4].

Table 1-1: Specific Energy and Energy Density of Commercial Rechargeable Batteries [3,4]

Battery Chemistry	Specific Energy, Wh·Kg ⁻¹	Energy Density, Wh·L ⁻¹
Pb-acid	30	80
Ni-Cd	40	90
Ni-MH	55	165
Ni-Zn	70	145
Ag-Zn	75	200
Li-ion	100-285	320-690

The specific energy and energy density of Li-ion batteries are limited by the reversible capacities of the cathode materials. The cathode materials used in Li-ion batteries are categorized into three major classes [5]: (1) layered lithiated transition metal oxides such as LiCoO_2 (LCO), $\text{LiNi}_{0.33}\text{Mn}_{0.33}\text{Co}_{0.33}\text{O}_2$ (NMC) and $\text{LiNi}_{0.8}\text{Co}_{0.15}\text{Al}_{0.05}\text{O}_2$ (NCA); (2) lithiated transition-metal spinel oxides such as LiMn_2O_4 and $\text{LiMn}_{1.5}\text{Ni}_{0.5}\text{O}_2$; (3) lithiated transition-metal phosphates like LiFePO_4 . Figure 1-2 depicts the crystal structure of these three cathode materials.

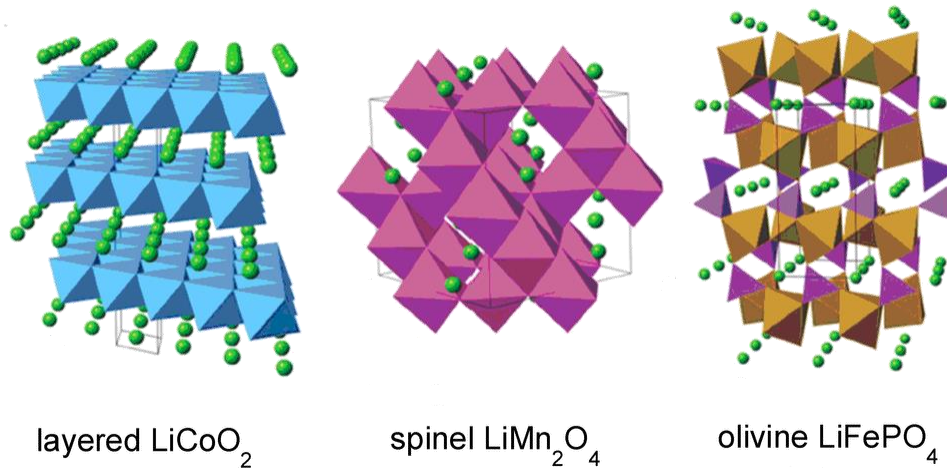


Figure 1-2: The crystal structures of common cathode materials in Li-ion batteries

Nanoarchitected cathodes with various morphologies, including zero dimension (0D; nanoparticles), one dimension (1D; nanotubes or nanowires), two dimension (2D; nanoplates or nanosheets), and three dimension (3D; hierarchical nanostructures), have also been developed to improve unique lithium storage properties and electrochemical performance (charge/discharge capacity, cycling stability, rate capability), and safety.

Figure 1-3 summarize some the recent nanostructured cathode materials developed for Li-ion batteries [6].

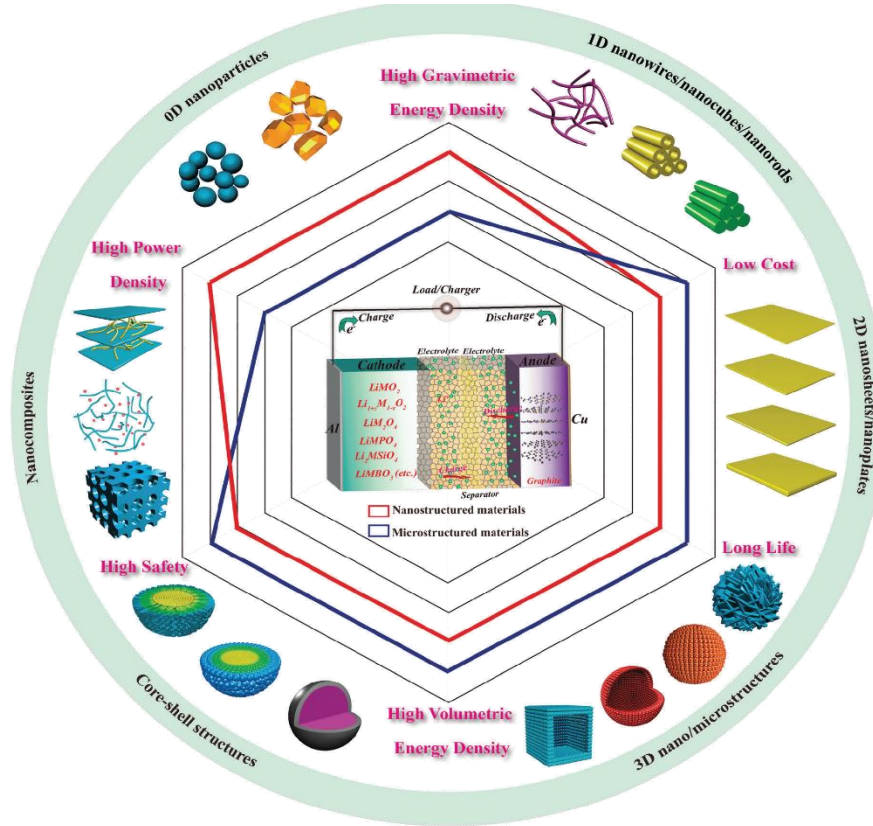


Figure 1-3: Illustration of the typical nanotechnologies applied in various cathodes for improving the Li-ion performance [6]

The graphite is a common material used in Li-ion battery anode with a theoretical specific capacity of $370 \text{ mAh} \cdot \text{g}^{-1}$ [2], which involves the transfer of one electron (or one equivalent of Li) per mole of C_6 . Graphite is used in different forms as active anode materials in Li-ion batteries including natural graphite, synthetic graphite, meso-carbon microbead (MCMB) graphite [2]. In Li-ion batteries, the power density is mostly limited by the graphite anode as the Li^+ ions diffusion rate into graphite is in the range of 10^{-9} - $10^{-7} \text{ cm}^2 \cdot \text{s}^{-1}$

¹ [7]. Therefore, many research studies have been devoted to developing new anode with higher energy and power density [8,9]. Figure 1-4 illustrates the active anode materials which have been used or under development for Li-ion batteries.

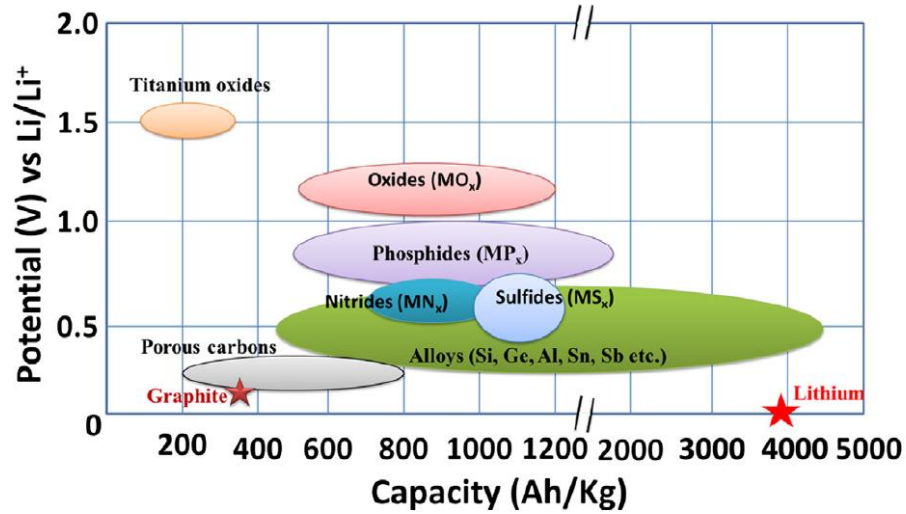


Figure 1-4: Schematic illustration of active anode materials for the next generation of Li-ion batteries [9]

1.2 Significance of Li-O₂ batteries

There is an urgency to minimize the consumption of fossil fuel and reduce the CO₂ emission due to the severe climate change. Therefore, the transition from gasoline-based vehicles to the partial electric vehicles like hybrid electric vehicles (HEVs) and plug-in hybrids (PHEVs) or fully electric vehicle (EVs) has begun [1,9,10]. Li-ion batteries have been in demand for electric vehicles for many years. Table 1-2 shows various Li-ion chemistries developed by different companies, which is currently being used in EVs ranked by present sales in the US [10].

Table 1-2: Batteries for Present Battery Electric Vehicles (EVs) sold in US [10]

Manufacturer	Model	Battery size (kWh)	Battery Chemistry	Battery Supplier	Vehicle range (mi)	Vehicle range (km)
Tesla	S	60–100	C/NCA	Panasonic/Tesla	208–315	334–508
Tesla	X	60–100	C/NCA	Panasonic/Tesla	208–315	334–508
BMW	i3	22,33	C/NMC	Samsung/Bosch	80,114	129,183
Nissan	Leaf	24,30	C/LMO (C/NMC)	AESC and LG Chem [†]	84,107	135,172
Volkswagen	e-Golf	24,35.8	C/NMC	Panasonic (Sanyo Div.)	83,124	135,200
Chevrolet	Spark	19	C/LFP	A123	82	132
Fiat	500e	24	C/NMC	Samsung/Bosch	87	140
Kia	Soul EV	27	C/NMC	SK Innovation	90	145
Smart	Fortwo EV	17.6	C/NMC	LG Chem	68	109
Ford	Focus EV	35.5	C/NMC	LG Chem	100	160
Mercedes	B-Class Electric	28	C/NCA, (C/NMC)	Panasonic/Tesla and SK Innovation [†]	85	137
Mitsubishi	i	16	LTO/LMO	Toshiba	62	100

While Tesla Motors and Chevrolet has announced that they will have a battery pack of 60 kWh and 85 kWh, their price is still far from the US Department of Energy ideal number (\$125 per kWh for a battery pack). Therefore, despite many developments in the Li-ion battery field, the current state-of-the-art of Li-ion batteries cannot meet many requirements for emerging applications such as future EVs. The energy density of 2 to 5 times greater of current Li-ion batteries are needed to meet the performance requirements of PHEVs with a 40–80 mile and EVs with a 300–400-mile driving range [1]. The next generation of advanced Li-ion batteries are approaching the performance needed for PHEVs; however, the recent development in energy density and price reduction of Li-ion batteries by Tesla Motors and Panasonic partnership is not sufficient for future EVs. Beyond Li-ion batteries (BLIs) such as Li-Sulfur and Metal-air batteries have been developed to replace the current Li-ion batteries. Figure 1-5 shows some the recent advances in Li-ion energy density and price pack along with the future battery chemistries [11].

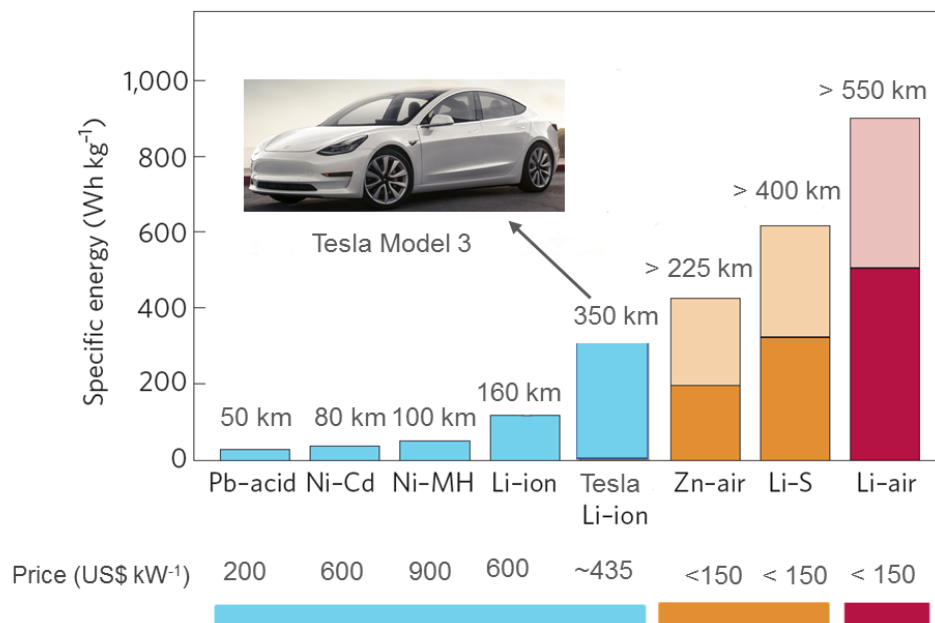


Figure 1-5: Practical specific energies for some rechargeable batteries, along with estimated driving distances and pack prices [11] (Note: some numbers in this Figure has been changed to reflect the recent advances in driving range and price pack of Li-ion batteries made by Tesla Motors.)

Among all BLI chemistries, the Li-air battery has the highest theoretical specific energy, and hence, it has attracted enormous research attention. Figure 1-6 shows the comparison of specific energy of various battery chemistries with gasoline [12]. The practical energy density of gasoline is $1700 \text{ Wh}\cdot\text{kg}^{-1}$ considering the average tank to wheel efficiency in the US is $\sim 13\%$, and theoretical energy density of gasoline is $13000 \text{ Wh}\cdot\text{kg}^{-1}$ ($13000 \text{ Wh}\cdot\text{kg}^{-1} \times 13\% = 1700 \text{ Wh}\cdot\text{kg}^{-1}$). The battery to wheel efficiency (battery systems) is about 90% , so the current energy density of Li-ion batteries ($100\text{-}265 \text{ Wh}\cdot\text{kg}^{-1}$) needs to be improved roughly 10-folds to make it competitive with gasoline. However, with the current state-of-the-art Li-ion batteries, reaching to the $1700 \text{ Wh}\cdot\text{kg}^{-1}$ is very optimistic. Since the energy density of lithium metal is $11680 \text{ Wh}\cdot\text{kg}^{-1}$, the practical energy density of gasoline (1700

$\text{Wh}\cdot\text{kg}^{-1}$) is only about 15% of the energy density of lithium metal. Hence, it is very conceivable that lithium-based batteries like Li-air could replace the gasoline easily.

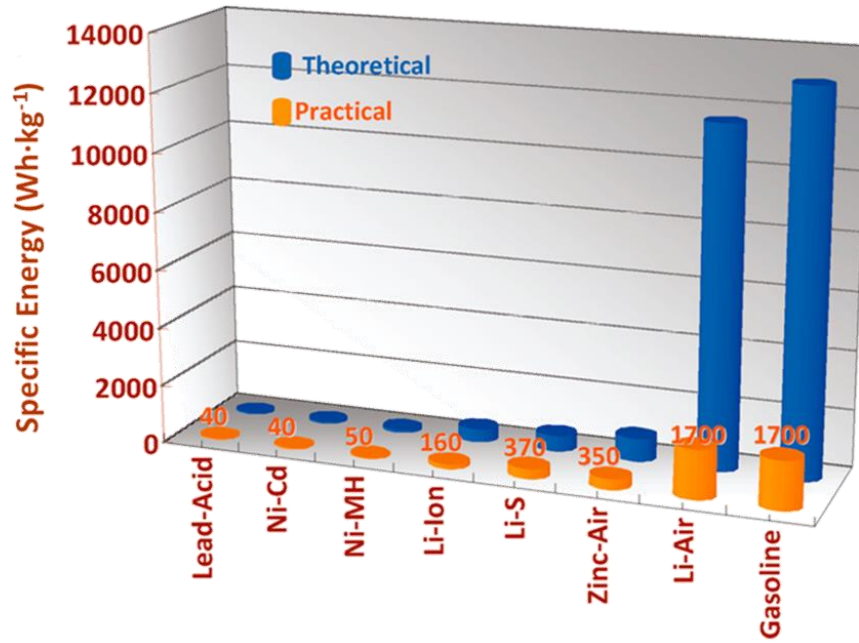


Figure 1-6: The specific energies ($\text{Wh}\cdot\text{kg}^{-1}$) for various types of rechargeable batteries compared to gasoline [12]

Although theoretical specific energy and energy density of Li-O₂ batteries are very high, the recent calculation has been shown that their practical system-level energy density and specific energy are not that much high considering the existing technologies. For example, Gallagher et. al [13] estimates system's practical specific energy and energy density for the Li-air batteries assuming two different methods for O₂ handling: (1) open Li-O₂ batteries with breathing systems with absorbers of humidity and removal of N₂ and CO₂ of the air and (2) close Li-O₂ batteries with battery packs inside the pressurized Oxygen vessel tank.

Figure 1-7 shows the comparison of their estimates for Li-O₂ and other possible batteries that are currently in research and development.

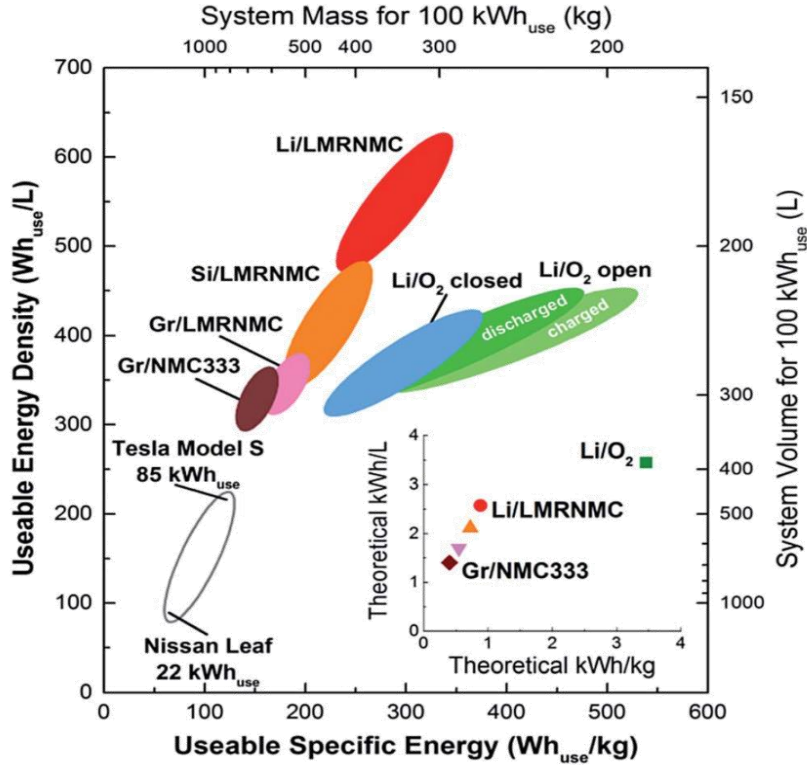


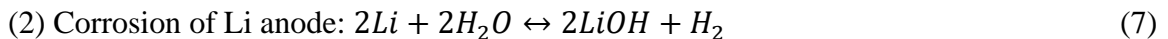
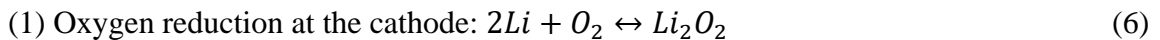
Figure 1-7: Calculated systems-level energy density and specific energy for 100 kWh of useable energy and 80 kW of net power at a nominal voltage of 360 V. (inset) Theoretical specific energy and energy density considering both anode and cathode active materials [13]

They argued that Li-O₂ batteries still have the highest projected useable specific energy, but are modest in comparison to this hypothetical advanced Li-ion battery using ANL Li-rich advanced cathode material (LMRNMC) in terms of energy density. Comparing their useable energy density and specific energy with the theoretical one (inset of Figure 1-7) of Li-O₂ batteries, it revealed that their estimated useable energy density and specific energy of Li-O₂ batteries are on order of magnitude smaller than a theoretical one, which is very

pessimistic. However, it is worth mentioning that their projections are based on the existing oxygen handling technology, and they did not consider any technological improvements in this aspect of the advanced batteries. Furthermore, other battery chemistries will undoubtedly compete with Li-air for the future EV since each of these battery chemistries now has their own technical challenges. For instance, advanced Li-ion batteries mentioned in Figure 1-7 using Li metal anode and Li-rich cathode material might give a practical, specific energy of $\sim 300 \text{ Wh}\cdot\text{kg}^{-1}$ and an energy density of $600\text{--}800 \text{ Wh}\cdot\text{L}^{-1}$ [14]. However, this also requires solving technical challenges of Li metal as an anode and the capacity fading in Li-rich cathodes. Thus, at this stage, there are a lot of uncertainties to pick a suitable chemistry among beyond Li-ion and advanced Li-ion batteries for future EVs.

1.3 Fundamental mechanisms of Li-O₂ batteries

The first primary lithium-air batteries were introduced by Littauer and Tsai in 1974 in which an aqueous alkaline solution was used as an electrolyte [15]. In their batteries, typical open circuit voltage was about 2.9–3.0 V, and a cell voltage of 2.0 V was achieved at current densities of approximately $200 \text{ mA}\cdot\text{cm}^{-2}$. The current efficiency of their cell was governed by the ratio of two competing reactions:



In their batteries, at OCV and low current density, self-discharge of the lithium anode was rapid, so the electrochemical efficiency of the cell was very low.

The first rechargeable Li-air battery was developed by Abraham and Jang [16] using a gel polymer electrolyte (GPE) containing a nonaqueous electrolyte. The cell consisted of a

lithium metal as an anode, a GPE and a carbon air electrode with a catalyst. Their GPEs consist of polyacrylonitrile (PAN) and a carbonate-based electrolyte containing LiPF₆ lithium salt. The observed OCV was around 3.0 V at room temperature, and the formation of lithium peroxide (Li₂O₂) on the surface of cathode after discharge was also confirmed. The capacity of the nonaqueous Li-air battery depends on the weight of the carbon cathode and its surface area as the discharge products Li₂O₂ is insoluble in nonaqueous electrolytes. In 2002, Read [17] developed a high capacity carbon air electrode using super P carbon black in a carbonate-based electrolyte (propylene carbonate (PC) and diethyl carbonate (DME)) containing LiPF₆ lithium salt. In 2006, Bruce and his coworkers reported the possibility of improved cycling of Li-air battery by using Super P carbon black with an electrolytic manganese dioxide. However, in 2010, Mizuno et al. [18] was reported that the lithium carbonate and lithium alkyl carbonate are the main discharge products, instead of Li₂O₂ in carbonate-based electrolytes. In the following years, other electrolytes like ether-based electrolyte [19,20] were developed for Li-air batteries. Currently, four types of Li-air (Li-O₂) are designed or are under development based on the electrolytes used: aprotic nonaqueous, aqueous, solid-state and hybrid (aqueous/nonaqueous) [21].

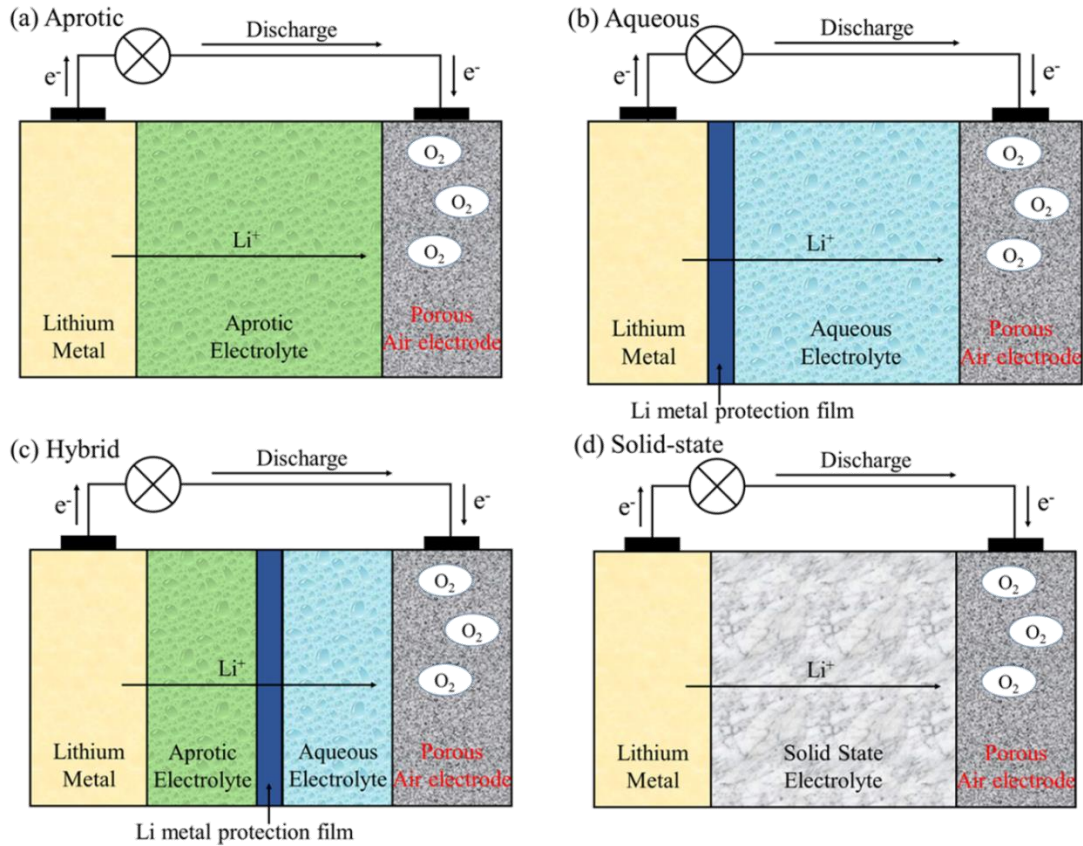


Figure 1-8: Different types of Li-O₂ batteries based on their electrolytes [21]

As can be seen from Figure 1-8, for all types of Li-O₂ batteries, an open system is necessary to get oxygen from the air and Li metal must also be used as the metal anode electrode to provide the lithium source for all the systems at the current stage. The configuration of Li-O₂ cells differs depending on the electrolyte employed. For instance, in the aqueous and hybrid aqueous/aprotic cells, a protective layer for Li metal is required to prevent the vigorous reaction of lithium with water. In aprotic Li-O₂ cells, porous carbon with sufficient surface area must be used as a cathode to serve as the reservoir for the insoluble discharge products. Of all Li-O₂ batteries, aprotic nonaqueous batteries have

gained much attention due to their similarities to the conventional Li-ion cells. In a typical Li-O₂ battery, the cell consists of Li metal as an anode and porous carbonaceous air cathode and Li⁺ containing aprotic electrolyte separating the cathode and anode. Figure 1-9 shows the discharge/charge mechanism of Li-O₂ battery in the nonaqueous electrolyte [12].

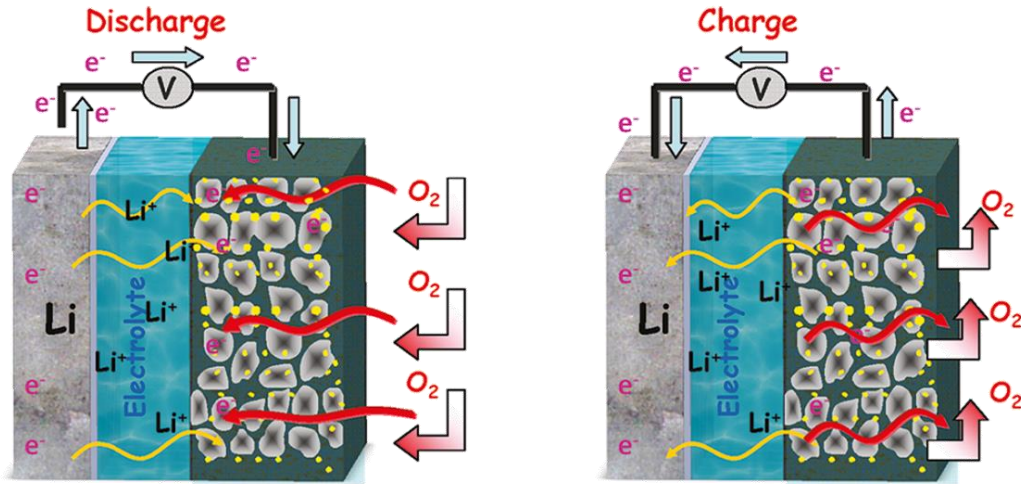
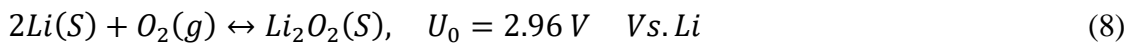


Figure 1-9: The discharge/charge mechanism of Li-O₂ battery in the liquid nonaqueous electrolyte [12]

During discharge, an oxidation reaction occurs at the anode ($\text{Li} \rightarrow \text{Li}^+ + \text{e}^-$) and electrons flow through an external circuit, and the Li⁺ ions generated from this reaction are transferred to the cathode through the electrolyte. At the cathode, the Li⁺ reduce oxygen to form Li₂O₂. Standard potential for the overall cell reaction, U_0 , can be calculated by the Nernst equation as follows:



In charge, the above reaction is reversed, and lithium metal is plated out on the anode, and O_2 is evolved at the cathode. The forward reaction happening in discharge is known as oxygen reduction reaction (ORR), and backward reaction is occurring in charge is known as oxygen evolution reaction (OER). Figure 1-10 shows a typical charge/discharge curve known as voltage profile of Li- O_2 battery [12]. As can be seen, the working voltage of this cell during discharge is approximately between 2.6 and 2.7 V, which is significantly less than the thermodynamical cell voltage, 2.96 V. This difference is called the discharge overpotential η_{dis} . During galvanostatic charging of the cell, the voltage increases to approximately 4.0 V. Hence the charge overpotential (η_{chg}) is significantly greater than the discharge overpotential (η_{dis}). The electrical energy efficiency for a charge/discharge cycle is only 65% ($2.6\text{ V}/4.0\text{ V}=65\%$) [12].

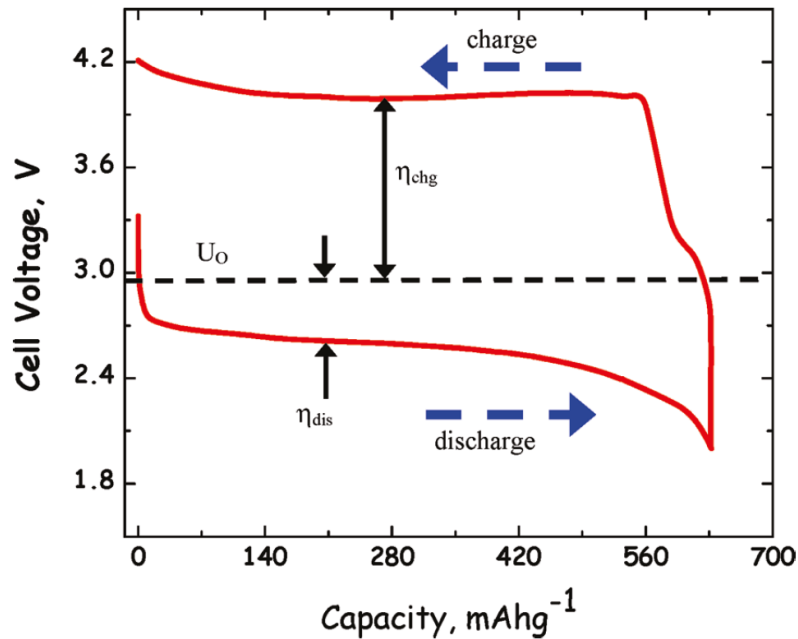


Figure 1-10: Typical voltage profile (charge/discharge curve) of Li- O_2 battery along with its overpotentials [12]

Many factors influence the charge or discharge overpotentials such as the deposition of side reaction products from the electrolyte and electrode degradation [22,23]. For example, the discharge processes depend on some competitive factors such as effective current density and voltage cut-off (overpotential) and whether the LiO_2 intermediate which is formed during discharge is dissolved in solution or adsorbed on the electrode surface. At high overpotentials and high current densities, O_2 is reduced to form Li_2O_2 , which grows as a film on the electrode surface [24,25]. However, at low current densities and overpotentials, Li_2O_2 can grow as surface films or large toroid-shape particles from a solution process, depending on the solvent or salt from which the electrolyte solution is formed or depending on additives in the electrolyte solution [26,27]. Figure 1-11 illustrates two different Li_2O_2 formation mechanism at low current density depending on the donor number of solvent used in electrolyte [23].

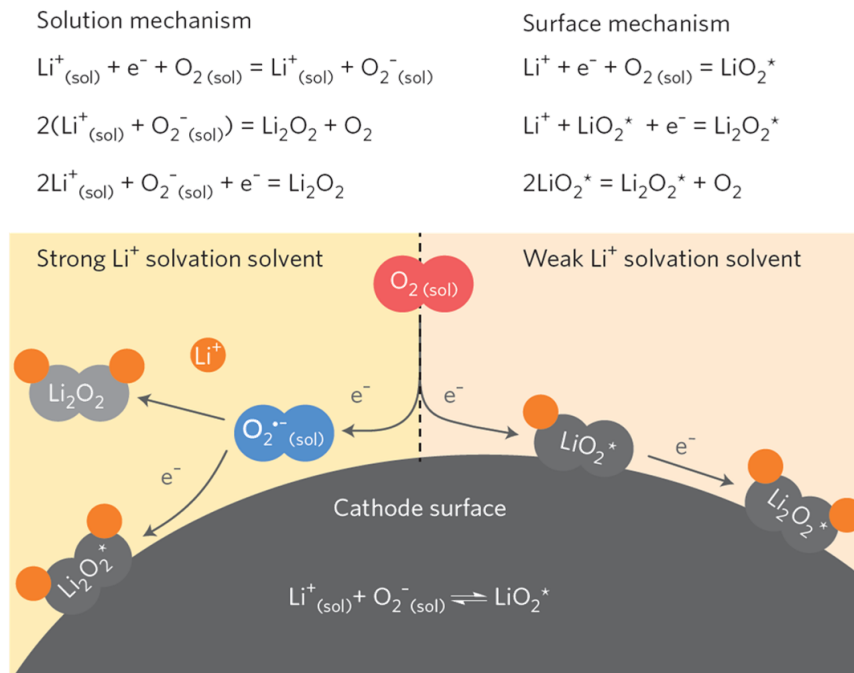


Figure 1-11: Reduction mechanisms in a Li- O_2 cell at low overpotentials depending on the donor number (DN) of solvent [23]

In a high donor number (DN) solvent, O_2^- is generated during discharge and dissolved in the electrolyte. Once the concentration of O_2^- reaches the solubility limit, it precipitates with Li^+ to produce LiO_2 on cathode surface and gets reduced or disproportionate to Li_2O_2 (solution-mediated formation). Large toroid-shaped Li_2O_2 can be formed via this mechanism, and accordingly, large discharge capacities can be obtained. On the other hand, in a low donor number solvent, LiO_2 is generated and deposited on cathode surface and further reduced via a disproportionation or electrochemical process to form a Li_2O_2 film on the cathode (surface-mediated formation). Figure 1-12 shows the scanning electron micrographs of toroid-shaped Li_2O_2 (solution-mediated formation) with the conformal film Li_2O_2 (surface-mediated formation) [26].

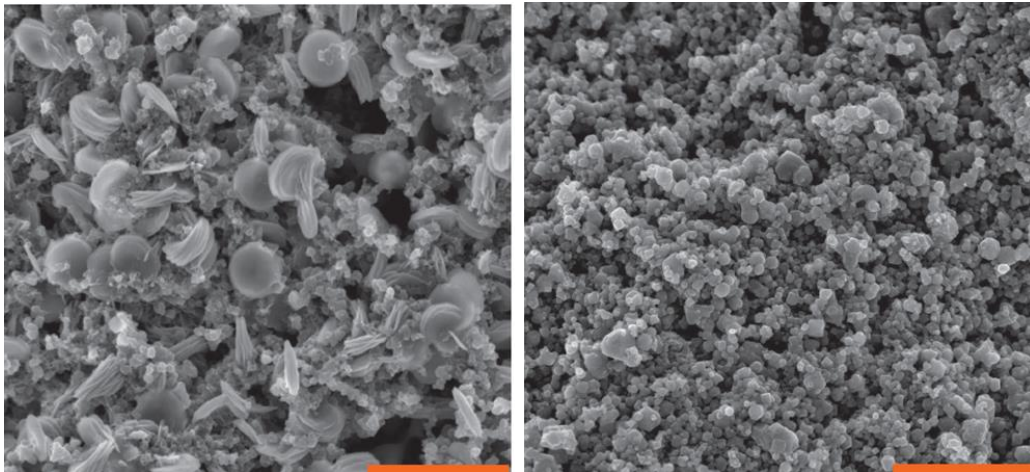


Figure 1-12: Scanning electron micrographs (right) of toroid-shape Li_2O_2 (solution-mediated formation) along with the (left) conformal film formation of Li_2O_2 (surface-mediated formation) on the Super P carbon black [26]

1.4 Challenges and degradation mechanisms in Li-O₂ batteries

Despite many research studies on the Li-O₂ batteries, they are still in their infancies, and many technical and fundamental challenges remains to be addressed before their commercialization [28]. Figure 1-13 depicts the summary of current challenges in Li-O₂ batteries.

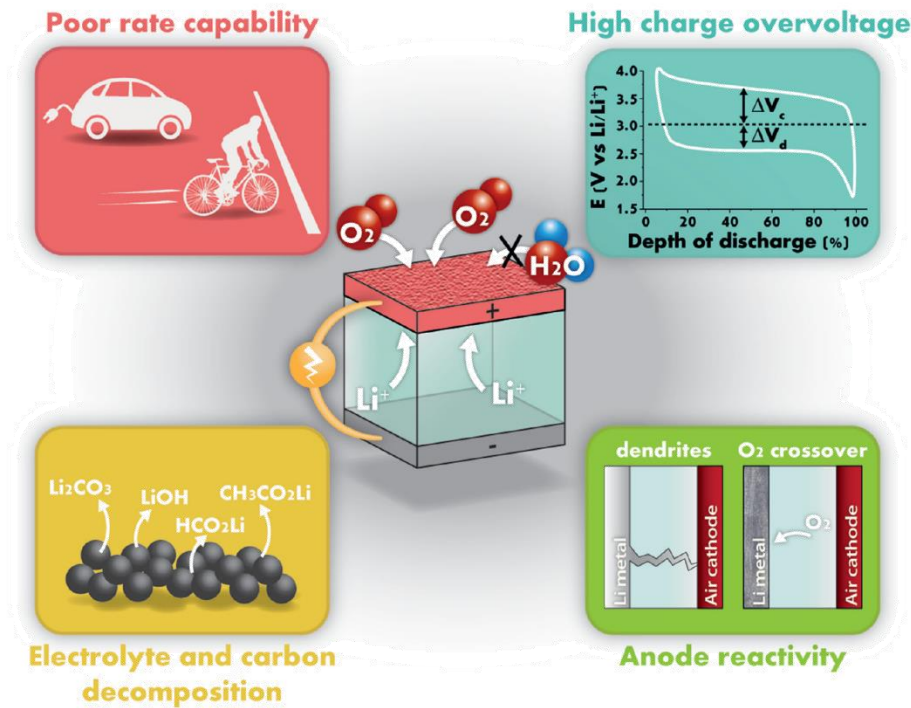


Figure 1-13: Summary of current challenges in Li-O₂ batteries [28]

Up to now, most of the research studies on Li-O₂ batteries have used only limited current densities (one or two orders of magnitude lower than those utilized in commercial Li-ion batteries), so the rate capability of Li-O₂ batteries must improve significantly to make them competitive with current Li-ion batteries. If the higher current densities cannot be achieved, the increase in O₂ flow can be considered as an alternative solution for transport

applications. Advanced chemical and electrochemical techniques have revealed that, to some degree, all components of Li-O₂ battery undergo undesirable chemical/electrochemical changes during discharge/charge cycling.

1.4.1 Lithium Anode Degradation

Metallic lithium is the main anode material used in Li-O₂ batteries due to its extremely low weight, the low negative potential (-3.04 vs. standard hydrogen electrode (SHE)) and high specific energy (11680 Wh·kg⁻¹). The usage of Li metal has its own safety concerns as the formation of lithium dendrites during repetitive lithium dissolution/deposition can result in poor cycling stability and internal short-circuiting. However, Li-O₂ battery failure owing to the dendrite growth has not been reported yet [29]. Recent studies have identified the reaction of Li metal with charge/discharge products and O₂ cross over from the cathode in Li-O₂ batteries [30–33]. Figure 1-14 illustrates the possible reaction is happening at the surface of the anode in Li-O₂ batteries.

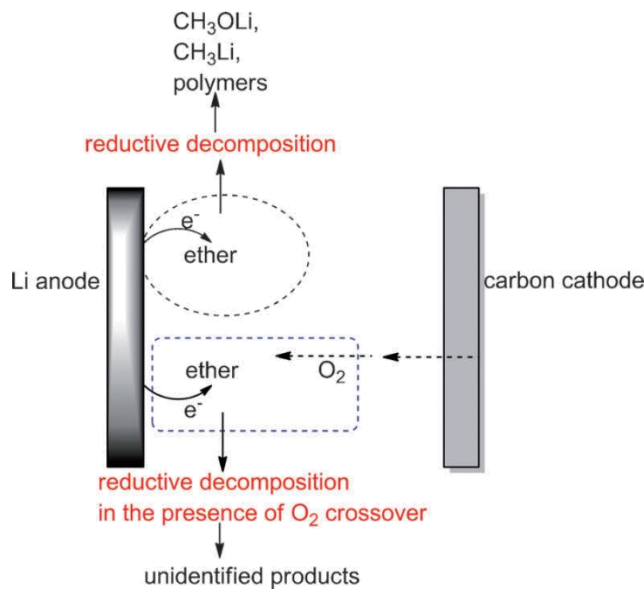


Figure 1-14: Possible Li metal reactions with electrolyte with/out O₂ [30]

Advanced approaches such as using oxygen and humidity impermeable separators [34] and artificial protective films [35] on the lithium have been proposed to minimize the Li anode degradation. Furthermore, replacement of lithium metal anode with lithiated carbon composites has also been proposed [36,37].

1.4.2 Cathode Degradation

As the main discharge products (Li_2O_2) of aprotic Li-O₂ batteries are insoluble in the electrolyte, they must be stored in a porous conductive matrix. Carbon in different allotropes have been used as cathode materials due to their high electronic conductivity, low cost, ease of fabrication and ability to catalyze the ORR/OER [38,39]. However, recent studies confirmed that carbon can react with discharge products and decompose during both discharge and charge in Li-O₂ batteries. It has been reported that carbon is chemically unstable above 3.5 V in the presence of Li_2O_2 and decompose to the lithium carbonates [22]. McCloskey et al. [22] reported that lithium carbonate (Li_2CO_3) and lithium alkyl carbonates (LiRCO_3) could be produced at the carbon- Li_2O_2 interface and Li_2O_2 -electrolyte interface, respectively due to the reaction of discharge product with Carbon cathode and electrolyte. Carbonate formation leads to an extra overpotential during charge, and subsequently, carbon reacts chemically with Li_2O_2 during charge to produce more lithium (Figure 1-15).

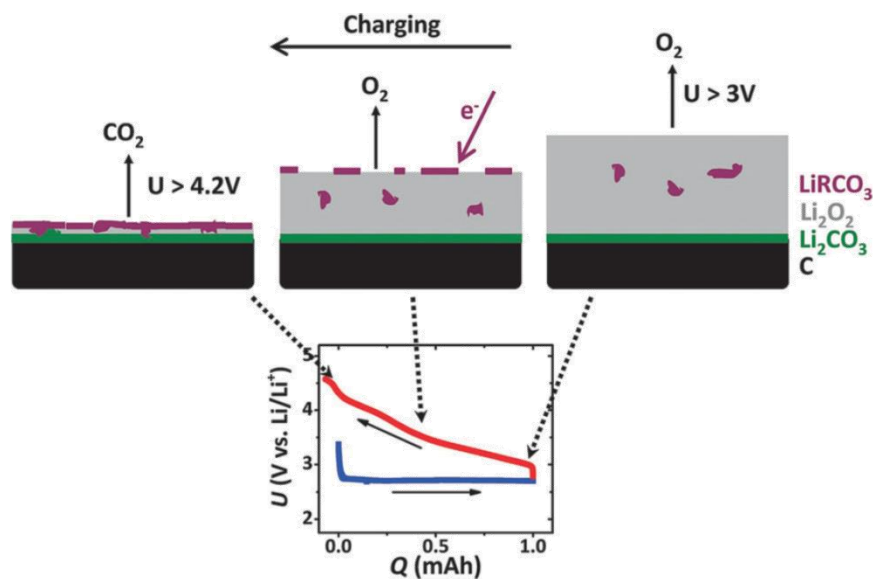


Figure 1-15: Proposed carbonate formation mechanism due to the reactivity of discharge products with carbon and electrolyte [22]

Due to the instability of carbon as cathode materials, many research has been devoted to replacing carbon. Bruce et al. [40] proposed nanoporous gold (NPG) cathode as a stable cathode for Li-O₂ batteries. Although NPG was stable and kinetics of Li₂O₂ oxidation was demonstrated to be faster than that of carbon cathodes, NPG cathodes are not suitable for cathode due to their high mass of gold, which in turn reduce the specific energy of Li-O₂ batteries significantly. Metal oxides [41] and metal carbides [42] have also been suggested for cathode materials. However, other side reactions were also reported for non-carbon cathodes [42].

1.4.3 Electrolyte Degradation

Despite many technological advanced in the development of stable cathode and anode for Li-O₂ batteries, electrolytes remain as a leading cause of rapid capacity fading and poor

cyclability [43]. The reactive oxygen species such as O_2^- , LiO_2 , Li_2O_2 and its derivatives are expected to coexist with molecular O_2 in the electrolyte owing to the ORR/OER and possible reaction between Li metal and dissolved O_2 [44,45]. The reaction between these reactive oxygen species with electrolytes has been considered as the main reason for electrolyte decomposition [46]. In general, the electrolyte decomposition pathways can be categorized into five groups as illustrated in Figure 1-16: (1) nucleophilic attacks, (2) auto-oxidation, (3) acid-base reactions, (4) proton-mediated reactions, and (5) reduction by Li [45].

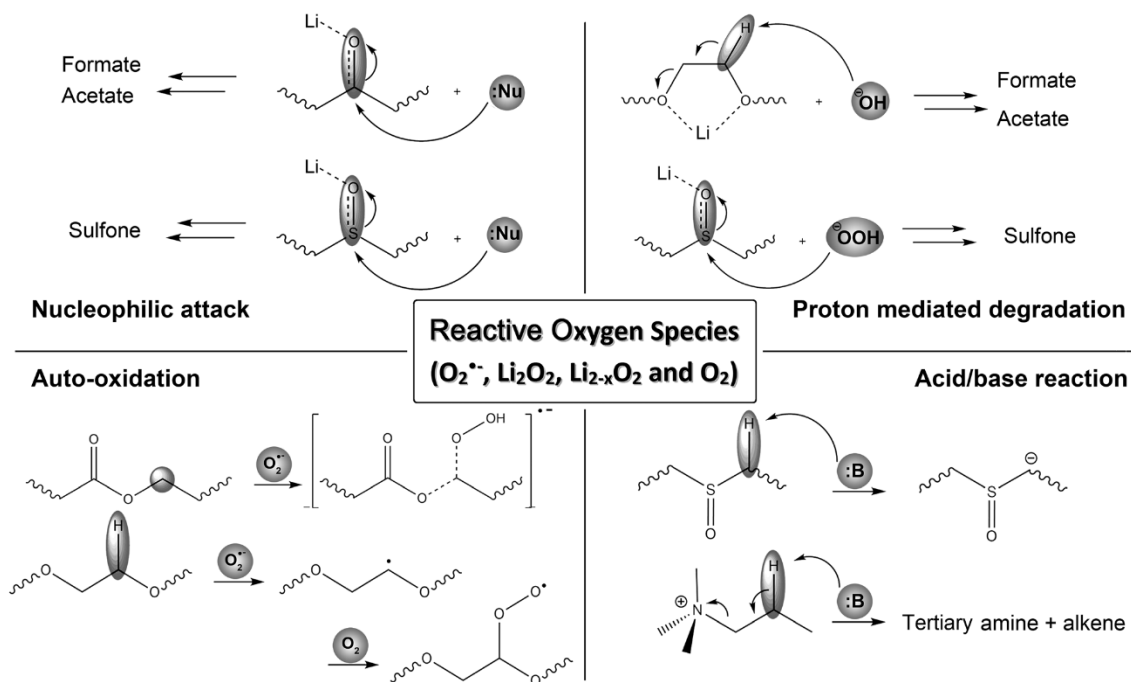


Figure 1-16: Schematic pathways of electrolyte decomposition by reactive oxygen species [45]

The decomposition mechanism is dependent on the chemistry of electrolyte used mostly solvent. In the following section, each of these decomposition pathways will be discussed briefly.

1.5 Electrolytes for Li-O₂ batteries

As mentioned earlier, the electrolyte has a profound influence on the reactions that occur at the anode and cathode and hence the overall cell operation of the nonaqueous Li-O₂ batteries. Electrolytes in Li-O₂ batteries need to have some certain requirements.

Table 1-3: Requirements on electrolytes for the nonaqueous Li-O₂ battery [15]

Conductivity	• Sufficiently high for the anticipated rate capability
Stability	• Within potential window used on discharge and charge • In contact with O ₂ and its reduced species on discharge • In contact with Li ₂ O ₂ and its intermediates on charge • In contact with the anode or a stable SEI formed on the anode
Low volatility	• To minimise evaporation at the porous O ₂ cathode
O ₂ solubility and diffusivity	• To ensure adequate rate of mass transport to the cathode
Able to wet the electrode surfaces	
Promotes some solubility of Li ₂ O ₂	Interaction with intermediates for high rate and packing density of Li ₂ O ₂
Safety, low cost and toxicity	

In the early stages of Li-O₂ battery research, organic carbonate-based electrolytes were widely used. However, it has been shown that the organic carbonates are unstable in Li-O₂ cells and a little or no evidence for Li₂O₂ formation on discharge was reported [47]. The nucleophilic attacks by O₂ to the C=O groups of carbonate-based electrolyte produce Li alkyl carbonates and Li₂CO₃[45,47]. Therefore, much attention shifted to other aprotic electrolytes for a Li-O₂ battery application.

1.5.1 Aprotic Liquid Electrolytes for Li-O₂ batteries

Ether solvents have been intensively studied for Li-O₂ batteries owing to their intermediate DN, which leads to the formation of both surface- and solution-mediated Li₂O₂ formation. They are also compatible with a lithium metal anode, stable to oxidation potentials up to 4

V versus Li/Li⁺, safe, low cost and low vapor pressure for higher molecular weights, such as tetraglyme (TEGDME) [15]. Various ether-based electrolytes in a linear form (glyme family) and a cyclic form such as 1,3-dioxolane have been explored. However, recent studies have confirmed that the decomposition of ether-based electrolytes also takes place gradually during cycling of Li-O₂ batteries to form lithium carbonates. The parasitic formation of these carbonates on the cathode eventually causes the battery failure [48]. It has been shown that the ether-based electrolyte undergoes decomposition by auto-oxidation mechanism in which superoxide radicals react with α -H in ethers [45].

Esters could also have been considered a good choice as electrolytes in the Li-O₂ batteries due to their high dielectric constant and low viscosity. However, most of the Ester solvents are prone to reaction with lithium metal [46]. Further, it has been shown that the Esters can be decomposed by the nucleophilic attacks [45].

Amides also are a major class of solvent that are known to be highly stable against nucleophilic attack and have been extensively studied for Li-O₂ batteries. Dimethylformamide (DMF) and N, N-dimethylacetamide (DMA) were investigated for the Li-O₂ battery applications. DMF could form Li₂O₂ upon discharge, and it decomposed upon charge, but some degree of side reactions upon cycling with the accumulation of Li₂CO₃, HCO₂Li, and CH₃CO₂Li were also reported [46,49]. In the case of DMA, high reactivity toward Li anode has been reported, which leads to the formation of unstable solid-electrolyte interphase (SEI). Unstable SEI formation on the anode could result in the formation of soluble decomposition products that are oxidized at the cathode surface upon charging [46]. Different approaches such as adding lithium salt additives (LiNO₃) to form

stable SEI on the anode was also proposed to make the amide-based electrolytes compatible with Li anode [50].

Sulfone-based electrolytes were also proposed as stable electrolytes against the superoxide attack (O_2^-) by theoretical calculation. However, many of sulfone-based electrolytes such as ethylmethylsulfone and tetramethylenesulfone have a low melting temperature around the room temperature ($\sim 27^\circ\text{C}$) which make them difficult to use in ambient temperature [46]. Further, some studies have shown sulfones are not stable against superoxide attacks and also they react with Li metal anode [15].

Dimethyl Sulfoxide (DMSO) electrolytes were also considered for Li- O_2 battery applications. Although the formation of Li_2O_2 was confirmed in cells using DMSO, other side products resulting from electrolyte decomposition such as $DMSO_2$, Li_2SO_3 , and Li_2SO_4 could be formed on the surface of cathode [51]. Li metal cycling efficiency of DMSO is low due to the reactivity of DMSO with Li metal [46].

Ionic liquids with different cations such as piperidinium, imidazolium, pyrrolidinium have also been studied for Li- O_2 battery applications due to their negligible vapor pressure, low flammability, high ionic conductivity and superior hydrophobicity and wide electrochemical window [52]. Some recent studies have proven the improved stability of ionic liquid over organic carbonate solvents for Li- O_2 batteries. However, some spectroscopic investigation on the discharge/charge products of cells using ionic liquids have also confirmed the existence of lithium carbonates. The limited cyclability of Li- O_2 cells using IL-based electrolytes and carbon electrodes suggests that side reactions are an issue that requires deeper investigation to confirm that these side reactions come from carbon decomposition or electrolyte decomposition [15].

1.5.2 Solid-State Electrolytes for Li-O₂ batteries

Solid-state electrolytes are considered competitive alternatives to liquid electrolytes for Li-O₂ batteries as they are safe, low cost, durable and flexible. They could also offer wider operational temperature range, and possibly longer cycle life owing to their ability to prevent lithium dendrite formation. The solid-state electrolytes are generally categorized into two general classes depending on their materials used: (1) Li⁺ ion conducting inorganic ceramics and (2) organic polymers. In the Li-O₂ cells, the advantage of solid-state electrolytes is that they act as a substantial barrier against diffusion of ambient gases and moisture toward the Li metal anode, and can also sustain a high operational temperature. These attractive features are the main driving force behind the development of Li-O₂ cells with solid electrolytes [46,53].

1.5.2.1 Ceramic Electrolytes

Ceramic electrolytes have also been explored for Li-O₂ batteries due to their relatively high Li⁺ conductivity, high thermal and chemical stability. Various types of ceramic electrolytes including sulfide, oxides, and phosphate were investigated in Li-ion batteries; however, only a few of them are employed in Li-O₂ batteries [54]. A family of ceramic electrolytes belong to Li-Al-Ge-PO₄ (LAGP) and Li-Al-Ti-PO₄ (LATP) systems, which possess relatively high ionic conductivity in the range of 10⁻⁴-10⁻⁵ S·cm⁻¹ are a good fit for Li-O₂ batteries. The first ceramic electrolyte in Li-O₂ batteries was reported by Kumar et al. [55] in which they investigated a LAGP ceramic electrolyte with a chemical composition of Li_{1+x}Al_xGe_{2-x}(PO₄)₃ (x = 0.5) (LAGP) as a solid-state electrolyte for Li-O₂ batteries. They suggested that LAGP-type ceramic electrolytes could also participate in Li₂O₂ formation [56]. Despite many attempts in the development of ceramic electrolytes with high ionic

conductivity, the high interfacial resistance between ceramic electrolyte/electrodes prevented the practical use of the solid-state ceramic at ambient temperature.

1.5.2.2 Solid polymer electrolytes

The first concept of solid polymer electrolytes (SPEs) was introduced by Prof. Peter Wright [57] in early 1973, and then their technological importance as a new class of electrolytes was discovered in the 1990s by Prof. M. Armand [58]. Since then, numerous polymer systems have been studied for lithium batteries, such as poly(ethylene) oxide (PEO). The PEO-based solid polymer electrolytes showed low ionic conductivity, poor mechanical properties, and narrow electrochemical window. Hence, many attempts have been made to explore solid new solid polymer electrolytes for lithium battery applications, such as composite polymer electrolytes, block copolymer electrolytes, and single-ion polymer electrolytes [53]. Solid polymer electrolytes (SPEs) have also been explored for Li-O₂ battery applications as they could offer a relatively high stability compared to the nonaqueous liquid electrolytes and a good protection for lithium anode to directly react with O₂ or H₂O. However, recent studies have shown that the chemical stability of polymers used in SPEs is questionable in the presence of discharge products [59]. Figure 1-17 illustrates the common polymer and their structures used in SPEs.

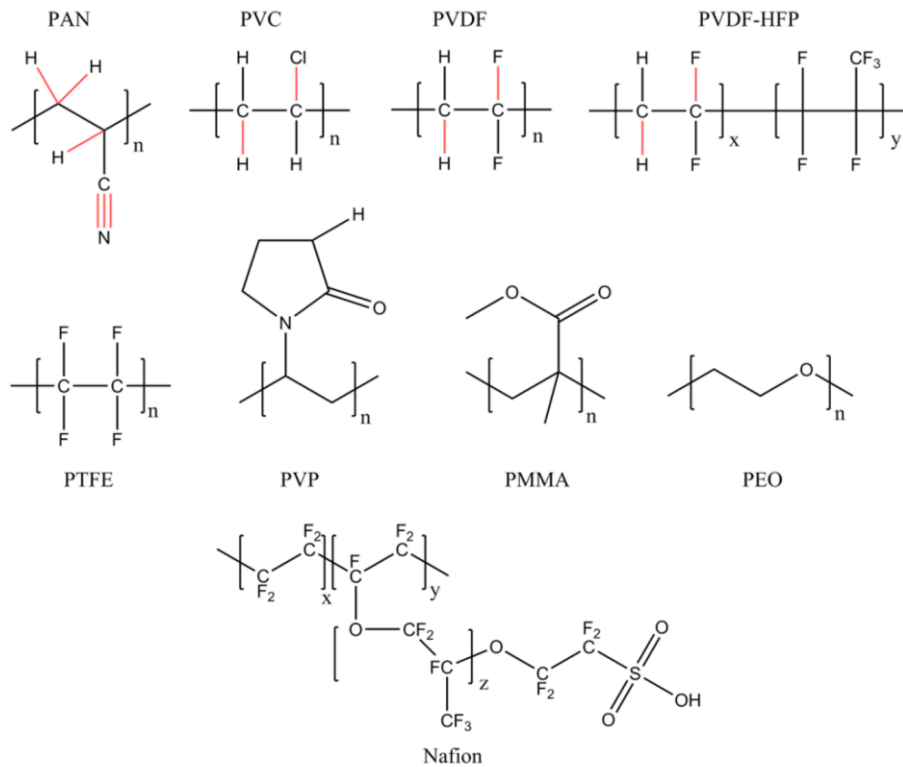


Figure 1-17: List of common polymers and their structures used in SPEs [59]

In SPE-based Li-O₂ batteries, the overall capacity is limited due to the absence of a liquid solution. Thus, most reported SPE-based Li-O₂ batteries were fabricated with thin and large-area carbon electrodes to obtain an acceptable capacity. Although impressive progress has been made on SPEs, the mechanism of ORR and OER in SPE-based Li-O₂ batteries is not yet identified. Moreover, the low ionic conductivity of SPEs shows poor reversible capacity [53].

1.5.2.3 Gel Polymer Electrolytes

Solid polymer electrolytes (SPEs) can be swollen with a liquid plasticizer to form gel polymer electrolytes (GPEs). These GPEs offer the ideal mechanical properties of SPEs

along with high ion conductivity of liquid counterparts. The first reported non-aqueous Li-O₂ battery was based on GPE containing polyacrylonitrile (PAN) polymer and carbonate electrolytes [16]. Other GPEs swollen by carbonates for Li-O₂ batteries was also reported [60]. However, using organic carbonates in their GPEs as plasticizers are questionable due to their tendency to decompose in the presence of oxygen radicals as mentioned earlier in this dissertation. Different GPEs using various polymer hosts and aprotic liquid electrolytes have been developed for Li-O₂ batteries [61].

Table 1-4: List of SPEs and GPEs used in Li-air (O₂) batteries [61]

Electrolyte type	Polymer matrix	Ionic conductivity ($\times 10^{-4}$ S cm ⁻¹)	Electrochemical window (V vs. Li/Li ⁺)	Operating atmosphere	Cycle life/cycle capacity
SPE	PAN	—	—	O ₂	3 cycles/ \approx 100 mAh g ⁻¹
SPE	PVA	2.85 (RT)	—	Ambient air	—
SPE	PEO	3.2 (55 °C)	Over 4.25 V	O ₂	—
SPE	PEO	—	—	O ₂	40 cycles/ \approx 210 mAh g ⁻¹
SPE	PEO	3 (RT)	Up to 4.6 V	O ₂	20 cycles/500 mAh g ⁻¹
GPE	Natural rubber	4.92 (RT)	—	O ₂	—
GPE	P(VDF-HFP)	—	—	O ₂	30 cycles/10 mAh
GPE	P(VDF-HFP)	10 (RT)	Up to 4.6 V	O ₂	50 cycles/500 mAh g ⁻¹
GPE	P(VDF-HFP)	5.49 (RT)	Up to 4.7 V	O ₂	40 cycles/500 mAh g ⁻¹
GPE	PMMA/PSt	1.27 (RT)	Up to 5.3 V	Ambient air, RH = 5%	100 cycles/500 mAh g ⁻¹
GPE	PVDF	—	Up to 4.7 V	O ₂	6 cycles/500 mAh g ⁻¹
GPE	PEO	—	—	O ₂	35 cycles/2.54 mAh
GPE	PVDF/pBQ	49.8 (RT)	—	O ₂	40 cycles/500 mAh g ⁻¹
GPE	P(VDF-HFP)	—	—	Ambient air, RH = 5%	180 cycles/500 mAh g ⁻¹
GPE	P(VDF-HFP)/ETPTA	2.843 (30 °C)	—	O ₂	100 cycles/500 mAh g ⁻¹
GPE	P(VDF-HFP)/ETPTA	2.843 (30 °C)	—	O ₂	90 cycles/500 mAh g ⁻¹
GPE	PFSA-Li	6.4 (RT)	—	O ₂	100 cycles/1000 mAh g ⁻¹

GPEs can improve the Li-O₂ battery performance in many ways. For example, it has been reported that the GPEs in Li-O₂ cell could minimize the electrolyte evaporation. GPEs could hinder the Li dendrite growth and form stable SEIs on the surface of Li metal anode [62]. They could also prevent Li metal corrosion by inhibiting the O₂ and humidity crossover [34,63,64]

1.6 Chemistry of Electrolyte Solution

Solvation is a process in which solute particles (molecules or ions) in a solution interact with the solvent molecules. The solvation of a solute has a significant influence on its dissolution and on the chemical reactions in which it participates [65]. The solvation energy can be defined as the standard chemical potential differences between solute in the solution state to their gaseous state as depicted in Figure 1-18.

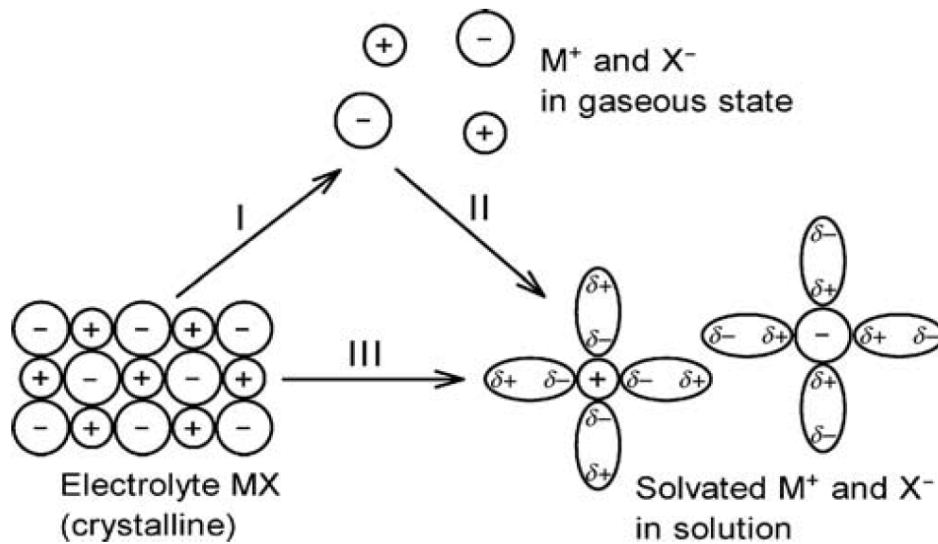


Figure 1-18: Schematic of dissolution process of solute MX in a solution [65]

As can be seen in process I, M^+ and X^- ions, which are strongly bound by electrostatic force in the crystal, are dissociated and converted to a gaseous state. In the process II, the M^+ and X^- ions in the gas phase dissolve into the solvent by solvation process. In Process III, the crystal of MX directly dissolves into the solvent and form solvated M^+ and X^- ions.

The Gibbs free energy of above-mentioned processes can be defined as follows:

$$\Delta G_{\text{III}}^0 = \Delta G_{\text{I}}^0 + \Delta G_{\text{II}}^0 \quad (9)$$

If the Gibbs energy of crystal lattice MX is denoted by ΔG_{lat} , ΔG_{I} is equal to $-\Delta G_{\text{lat}}$. Also, once MX is completely dissociated into free ions in the solution, ΔG_{II} could be obtained as the sum of the solvation energies of M^+ and X^- ions ΔG_{sol} . ΔG_{III} corresponds to the Gibbs energy of dissolution of MX, which is defined as ΔG_{s} , so the equation (9) can be rewritten as follows:

$$\Delta G_{\text{s}}^0 = \Delta G_{\text{sol}}^0 + \Delta G_{\text{lat}}^0 \quad (10)$$

Generally, ΔG_{sol}^0 and ΔG_{lat}^0 have large negative values, yet their magnitude close to each other in magnitude. Thus, ΔG_{s}^0 is relatively small. Moreover, the solute is easily soluble if the sum of the solvation energies of the ions constituting the solutes is larger than the lattice Gibbs energy (in absolute value) or very near to it.

Equation (10) could also be expressed by the solubility constant of MX ($K_{\text{sp}}(\text{MX})$):

$$\Delta G_{\text{s}}^0 = -RT \ln K_{\text{sp}}(\text{MX}) \quad (11)$$

In the electrolyte solutions, the interaction between ions (cations and anions) of salt and solvent molecules play a significant role in electrolyte properties such as ion transports and chemical and electrochemical behavior of electrolytes.

1.6.1 Ion-Solvent Interaction in Electrolytes

As described in the previous section, the solvation is an important parameter in the dissolution of solute (MX). It has been reported that ions of solute (MX) can interact with solvent molecules in different ways [65,66]. Hence the solvent properties can significantly affect the electrolyte solution properties. The most important solvent properties in considering solvent effects are the solvent permittivity and the solvent acidity and basicity. For instance, if the permittivity of one solvent is high ($\epsilon_r > 40$) and that of the other is low ($\epsilon_r < 10$), the difference in a chemical process in the two solvents is usually attributable to the influence of permittivity. On the other hand, in two high-permittivity solvents ($\epsilon_r > 40$) is often attributable to the influence of the acidity or basicity of the two solvents rather than the influence of permittivity. Table 1-5 shows ion-solvent interaction with their contribution percentage of total ionic solvation energy.

Table 1-5: Different Ion-Solvent interaction along with their contribution percentage [65]

1. Electrostatic interactions as expressed by the Born equation	($\geq 80\%$)
2. Electron (-pair) donor-acceptor interactions	($\leq 10\%$)
3. Interactions of anions with hydrogen bond donor solvents	($\leq 10\%$)
4. Interactions based on HSAB concept	($\leq 20\%$)
5. Interactions by back-donation from d^{10} -cation to solvent molecules	($\leq 10\%$)
6. Interactions related to the structure-making and breaking of solvents	($\leq 5\%$)

As it is clear the electrostatic interaction has the major contribution in ion-solvent interaction and can be defined as the difference between the electrostatic free energy of an ion in vacuum and that of the ion in a solution of relative permittivity. It has been shown that the difference between the electrostatic ion-solvent solvation energy in two high-

permittivity solvents is often less important than the difference in the solvation energies caused by other interactions [65,66].

Another important contribution in ion-solvent interaction is electron pair donor (EPD) and electron pair acceptor (EPA) interactions. In ion solvation process, the solvent molecules approach a cation with their negative charge and approach an anion with their positive charge. Therefore, cation solvation is mainly related to the electron pair donor capacity (Lewis basicity) of the solvents and becomes stronger with the increase in donor number (DN) of solvent. The anion solvation, on the other hand, is closely associated with the electron pair acceptability (Lewis acidity) of the solvents and again becomes stronger with the increase in acceptor number (AN) [65,66].

The ion-solvent interactions can be studied by spectroscopic techniques like Infrared (IR), Raman and nuclear magnetic resonance (NMR) spectroscopy[67].

1.6.2 Ion-ion Interaction in Electrolytes

The coulombic force of attraction between two oppositely charged ions (M^+ and X^-) is inversely proportional to the relative permittivity of the solvent. Thus, solvents with high relative permittivity ($\epsilon_r > 40$) will be able to reduce the strong electrostatic attraction between oppositely charged ions and dissociate them into free solvated ions [66]. However, in relative low permittivity solvents, the complete dissociation becomes difficult, and part of the dissolved solute (MX) is not dissociated. The undissociated ions in low permittivity solvents contribute to chemical reactions and ion transport in the electrolyte solution. The ion association/dissociation of solute (MX) is strongly depended on the ion-association constant (K_A) which could be defined as follows:



$$K_A = \frac{[M^+X^-]}{[M^+][X^-]} \quad (13)$$

Upon solvation, based on the mutual geometric arrangement of the two ions and the solvent molecules different ion pairs can be formed. Figure 1-19 shows different ion pairs formed in the electrolyte solution.

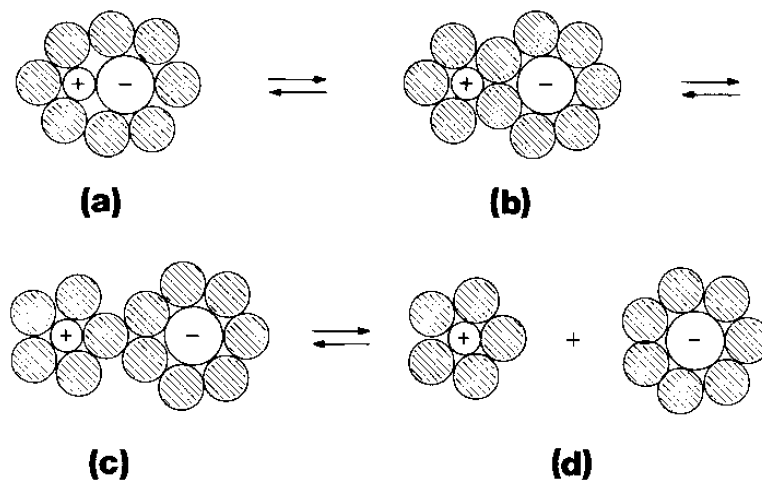


Figure 1-19: Schematic representation of different ion-pairs (a) Contact ion pair (b) Solvent shared ion pair (c) Solvent separated ion pairs (d) Free solvated ions (Shaded circles denote the solvent molecules) [66]

In contact ion pairs (CIPs), no solvent molecules intervene between the two ions that are in close contact. The ion pair separated by the thickness of only one solvent molecule is called a solvent-shared ion pair. In solvent-separated ion pairs (SSIPs) the primary solvation shells of the two ions are in contact so that some overlap of secondary and further solvation shells takes place. Further dissociation of the two ions leads to unpaired (free) solvated ions with independent primary and secondary solvation shells. In dilute solution

using low-permittivity solvents, the presence of ion-pairs even in dilute solutions were reported. With the increase of solute concentration, the formation of aggregate ions was also observed in high-permittivity aprotic solvents. For an alkali salts (LiX), ionic association strength can be affected by the negative charge delocalization, size, and steric effects of the anion X⁻. They could be categorized in three different classes [68]: (1) dissociated salts: LiN(SO₂CF₃)₂ (2) intermediate salts: LiClO₄ and LiBF₄ and (3) associated salts: LiCF₃SO₃, LiNO₃ and LiCF₃CO₂.

1.6.3 Salt-Inorganic Additive Interaction in Electrolytes

The first introduction of inorganic fillers in electrolyte solution for battery application was reported by Weston and Steele in 1982 [69], in which Al₂O₃ filler particles were added into PEO-based polymer electrolytes. They reported a significant improvement in the mechanical stability of a polymer electrolyte upon the addition of an inert filler, yet negligible reductions in ionic conductivity at low loadings. Their observation led to more research work to investigate the beneficial influence of inorganic fillers on an ion transport properties of electrolytes. Scrosati et. al [70] reported the increase in ionic conductivity of PEO-based solid polymer electrolyte (SPE) by incorporating nanosized TiO₂ and Al₂O₃ fillers. Since then many attempts have been made to improve the ionic transport properties of SPEs by ceramic fillers. Fillers can influence the ion transport mechanisms in SPEs in a variety of indirect or direct ways [71]. For example, one of the main concerns in SPEs is their low ionic transport properties at low temperatures due to lack of the amorphous phase in polymer structures. Ion transport in polymer electrolyte is due to segmental motion of polymer chains, which are significantly higher in the amorphous regions compared to crystalline counterparts [72]. Analogous to liquid plasticizers, small fillers may also add

free volume and speed up segmental dynamics and in turn improve the ion transports. Inorganic fillers could also directly participate in ion transport by increasing free Li^+ concentrations, Li^+ surface conduction, anion attraction, or as a Li^+ source [71].

The mechanism of filler participation in ion conduction can be expressed as follows: (1) fillers actively interact with the ion pairs. In this case, fillers with specific surface chemistries promote ion-pair dissociation level and increase the number of ions able to participate in conduction. It has been shown that acidic surface groups could attract anions, while basic surface groups attract cations. In either case, the corresponding counter-ion acts as mobile species. (2) The surface of the fillers provides an additional site for anion and/or cation migration, due to Lewis acid–base interactions between the salt and the particle surface. (3) The filler surface attracts either the anion or cation, which reduce the ion mobility. (4) Fillers can also behave as crosslinking sites for EO-segments and anions, changing the polymer chain structure at the interface and creating pathways for Li^+ transport independent of segmental motion.

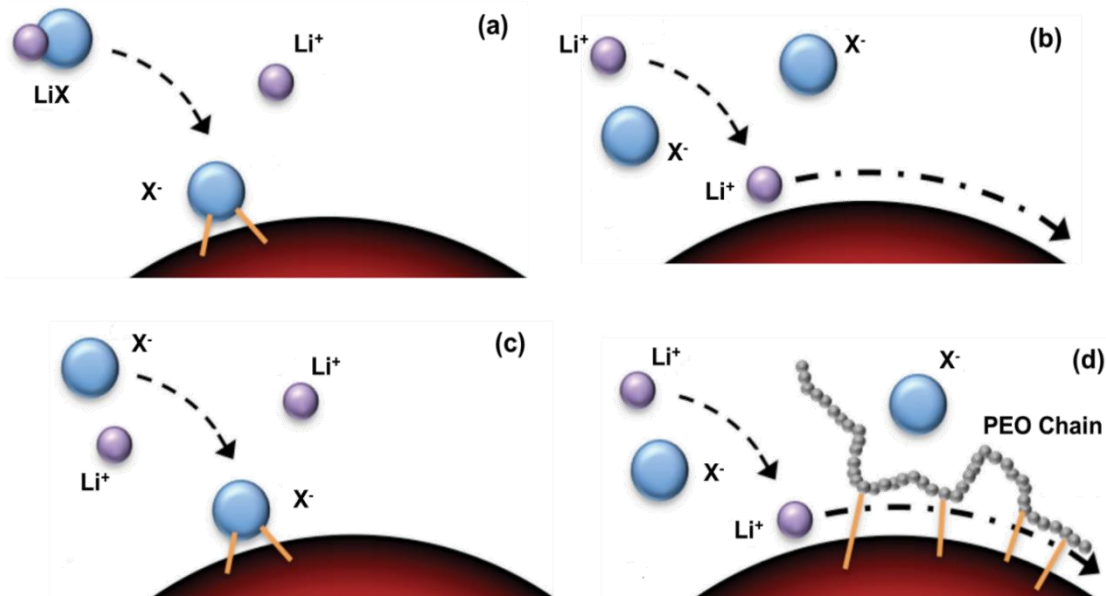


Figure 1-20: Schematic of the filler ion transport mechanisms in SPEs (a) Ion pair dissociation (b) surface transport (c) anion attraction (d) PEO chain promoted surface

Interaction between lithium salt and inorganic fillers in liquid electrolytes was also reported. Bhattacharyya and Maier in 2004 showed that active Li^+ surface conduction also occurs in liquid-based electrolytes with fillers known as soggy-sand electrolyte [73]. The liquid matrix allows for percolation of spherical particles at a lower loading, as in the absence of an adsorbed polymer layer the particles readily aggregate to form networks of complex fractal dimensionalities. Because of the low particle loading (<1–3 wt.%), soggy sand electrolytes with significant Li^+ surface transport display ionic conductivity above that of the pure liquid electrolyte. The mechanism of ion transport in soggy-sand electrolytes could be attraction of the ion pair at the surface of fillers and facilitates the ion pair dissociation. The counter-ion will then exist in the space charge region at the vicinity of particle liquid interface. At a threshold filler loading, ionic conductivity increases as percolation allows for long-range transport of the free ions in the space charge layer. Below

the threshold filler loading, the ionic transport properties will not change as percolation does not exist. Beyond threshold filler loading, the conductivity will continue to increase with increasing filler content until to a given filler loading (maximum loading). Beyond this maximum filler loading, conductivity decreases due to blocking of the percolative pathways and volume depletion effects [71]. Figure 1-21 shows the space charge at the vicinity of fillers and electrolyte interface with with/o filler percolation.

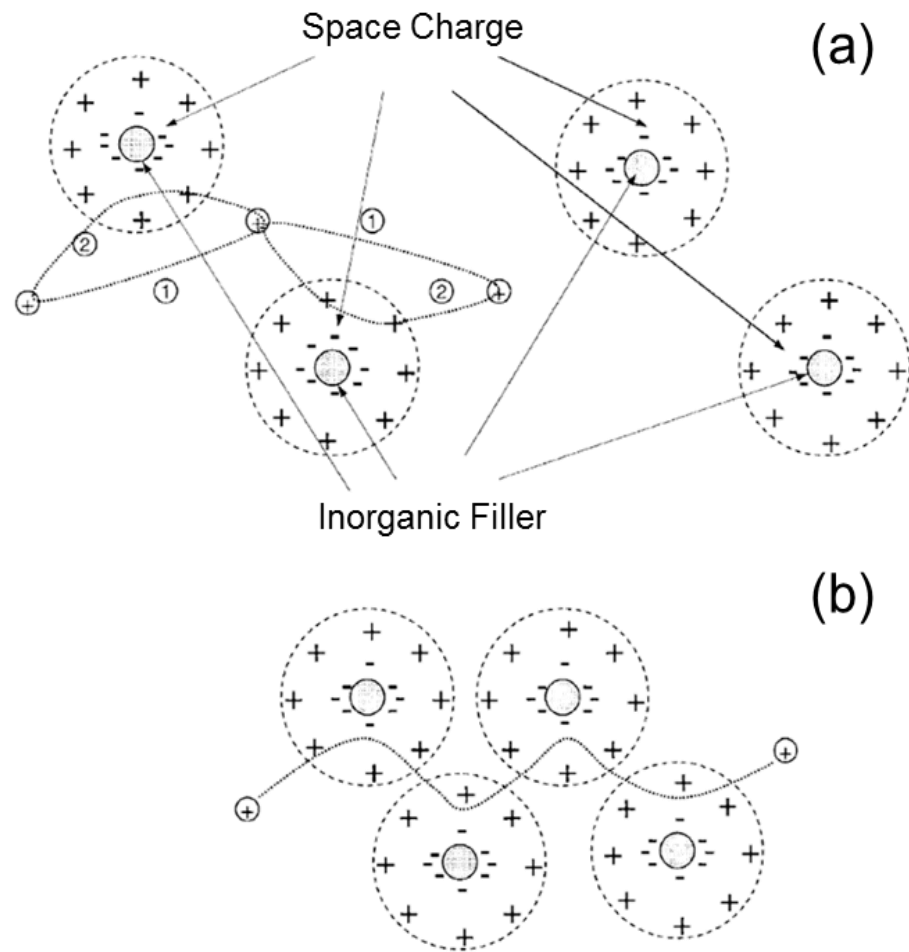


Figure 1-21: The inorganic filler in liquid electrolyte (a) below threshold filler loading (no percolation) (b) above or at threshold filler loading (percolation formed)

1.6.4 Ionic Transport Properties in Electrolytes

The value of conductance, L for a segment of solution immersed in an electric field is directly proportional to the cross-sectional area perpendicular to the field vector and is inversely proportional to the length of the segment along the field. The proportionality constant is the conductivity, σ , which is an intrinsic property of the solution [74]:

$$L = \sigma \frac{A}{l} \quad (14)$$

The conductance, L , is given in units of Siemens ($S = \Omega^{-1}$), and σ is expressed in $S \cdot \text{cm}^{-1}$.

Ionic conductivity, σ is the sum of contributions from all ionic species as the passage of current through the solution is accomplished by the independent movement of different species. Therefore, it is acceptable that each component of σ is proportional to the concentration of the ion, the magnitude of its charge $|Z_i|$, and the mobility, which is the limiting velocity of the ion in an electric field of unit strength. Once an electric field in the strength of ξ is applied to an ion, it accelerates under the force imposed by the field until the frictional drag force exactly counterbalances the electric force. Then, the ion continues its motion at that terminal velocity. The magnitude of the force applied by the field is equal to $|Z_i| e \xi$, where e is the electronic charge. The frictional drag force can also be approximated using Stokes law as $6\pi\eta r v$ where η is the viscosity of the medium, r is the radius of the ion, and v is the velocity. When the terminal velocity is reached, the ion mobility can be defined as [74]:

$$u_i = \frac{v}{\xi} = \frac{|Z_i| e}{6\pi\eta r} \quad (15)$$

The proportionality factor relating an individual ionic conductivity to charge, mobility,

and concentration turns out to be the Faraday constant so that the ionic conductivity can be defined as [65,74]:

$$\sigma = F \sum |Z_i| u_i C_i \quad (16)$$

The transference number for species i , which the fractions of the current carried by species I and J are called their transference numbers and is merely the contribution to conductivity made by that species divided by the total conductivity:

$$t_i = \frac{|z_i| u_i C_i}{\sum_j |z_j| u_j C_j} \quad (17)$$

1.7 Statement of Problems

Despite the high theoretical energy density of Li-air (O₂) batteries, they are far away from ideal energy storage systems for emerging applications such as electric vehicles (EVs). Li-O₂ batteries suffer from poor cyclability and quick capacity fading. Many fundamental research studies have been performed on the cathode and anode of Li-O₂ batteries to improve their performance; however, the electrolytes used in this field remains as one the leading causes of poor battery performance. Electrolytes with various solvents and lithium salts have been utilized for Li-O₂ batteries. Nonetheless, up to date, the choice of solvent and salt for chemical and electrochemical stable electrolytes remain a big challenge in this field. The aim of this dissertation was on the development of stable electrolytes for a Li-O₂ battery application. Recently, it has been shown that the stability of glyme-based electrolytes (common electrolytes used in the Li-O₂ battery) can be enhanced to some extent by changing the solvation of solvent and lithium salt. Some other research studies have also reported the improved performance of Li-O₂ batteries using gel polymer

electrolytes (GPEs) over liquid counterparts. The introduction of inorganic fillers in electrolytes (liquid and polymer electrolytes) have also been shown to enhance the transport properties of electrolytes by changing the ion pair association/dissociation. This research study aims to investigate the addition of inorganic filler materials to the electrolyte as a possible method to change the electrolyte properties for improved Li-O₂ battery performance. For this purpose, composite gel polymer electrolytes (cGPEs) and GPEs using common glyme-based liquid electrolyte and UV-curable polymer with and without one-dimensional (1D) borosilicate glass microfillers were developed, and their performance metrics were studied for Li-O₂ batteries.

In chapter 3, the performance of batteries using GPEs and cGPEs with different content of glass microfillers were investigated using different electrochemical characterization and spectroscopic techniques to obtain the optimum loading of glass microfillers. This work has been published:

Amir Chamaani, Neha Chawla, Meer Safa, Bilal El-Zahab, "One-Dimensional Glass Micro-Fillers in Gel Polymer Electrolytes for Li-O₂ Battery Applications", Electrochimica Acta 235 (2017) 56–63.

In chapter 4, the loading of glass microfillers was fixed to the optimum amount obtained in chapter 3 and lithium salt concentration were changed to investigate the sources and mechanism of improvement of cGPE-containing batteries. This chapter revealed that the glass microfillers reduce the rate of lithium carbonates formation originating from electrolyte decomposition. Using different spectroscopic and electrochemical techniques, a possible mechanism of improvement was also proposed.

Amir Chamaani, Meer Safa, Neha Chawla and Bilal El-Zahab, “Composite Gel Polymer Electrolytes for Improved Cyclability of Li-O₂ batteries” ACS Applied Materials and Interface 9 (2017) 33819–33826

In chapter 5, the performance of Li-O₂ batteries using glyme-based electrolytes with different salt concentrations was also investigated. The results confirmed that the increase in lithium salt concentrations would improve the battery performance by reducing the electrolyte decomposition. The spectroscopic results showed that increase in lithium salt concentration increases the formation of cationic complexes and in turn mitigate the electrolyte decomposition. This work was submitted to the Journal of Electroanalytical Chemistry, and it is still under review

Amir Chamaani, Meer Safa, Neha Chawla and Bilal El-Zahab, “Stabilizing Effect of Ion Complex Formation in Lithium–Oxygen Battery Electrolytes” (Journal of Electroanalytical Chemistry, Under review)

2 EXPERIMENTAL METHODS AND THEORY

2.1 Chemicals and Materials

The following is a list of all chemicals and materials used throughout this thesis:

Ethoxylated trimethylolpropane triacrylate (ETPTA, $M_w=428$, purity > 99.00%, Sigma Aldrich), 2-hydroxy-2-methyl-1-phenyl-1-propanon (HMPP, Photo-initiator, purity >97.00%, Sigma Aldrich), lithium bis(trifluoromethane) sulfonamide (LiTFSI, purity > 99.95%, Alfa Aesar), tetraethylene glycol dimethyl ether (TEGDME, purity > 99.00%, Alfa Aesar), N-Methylpyrrolidine (NMP, purity >97.00%, Sigma Aldrich), Polyvinylidene fluoride (PVDF, Alfa Aesar), multiwalled carbon nanotubes (CNTs, $D=5-20$ nm, $L=5$ μm , purity > 95.00% carbon basis, Sigma Aldrich) and Whatman glass microfiber filters (Binder free, Grade GF/B, Sigma Aldrich), Carbon cloth gas diffusion layer (CCGDL, thickness ~ 300 μm , FuelCell Earth), Polypropylene separator (Thickness ~ 25 μm , Celgard LLC.), Lithium foils (purity > 99.90%, MTI Crop.), Molecular Sieve beads (4Å, Sigma Aldrich). Polydimethylsiloxane (PDMS, SYLGARD 184 and curing agent, Dow Corning).

2.2 Li-O₂ cell assembly

2.2.1 Liquid Electrolyte Preparation

All liquid solvents used for liquid electrolytes and polymer electrolytes (gel polymer electrolytes (GPEs) and composite gel polymer electrolytes (cGPEs)) preparation were first dried over molecular sieve beads (4Å) for at least two weeks before use. The molecular sieves were always activated before use by heating at 250 °C under vacuum for 24 hours. Liquid electrolytes with different LiTFSI salt concentrations were prepared by dissolving an appropriate amount of LiTFSI salt (0.1, 1 and 3 mol·kg⁻¹) into TEGDME solvent in the

Ar-filled glove box with control O₂ and humidity content (Mbraun, <0.1 ppm O₂ and < 0.1 ppm H₂O).

2.2.2 Gel polymer and composite gel polymer electrolytes preparation

The glass microfibers used in this thesis as one-dimensional fillers for cGPEs preparation obtained by cutting Whatman glass microfiber discs (Binder-free, grade GF/B) and further fragmented by probe sonication in acetone for 3 hours. After fragmentation process, the glass microfibers were dried under high vacuum at 300°C for 48 hours and then stored in the Ar-filled glovebox for at least 2 weeks before use. GPEs with different LiTFSI salt concentrations were prepared by mixing of ETPTA monomer into liquid electrolytes (0.1, 1, and 3 mol·kg⁻¹) solution (80:20 wt.% of ETPTA/liquid electrolyte with 1:99 by weight HMPP: ETPTA monomer content as a curing agent). The cGPE preparation followed the same procedure with the addition of different fillers concentrations ranging from 0.5, 1, 2 and 5 wt.% of shredded glass microfibers. For cGPEs preparation, the appropriate amount of glass microfibers first was added to the liquid electrolytes in the Ar-filled glove box in sealed small vials and sonicated for 1 hours. The solution of ETPTA monomer and curing agent was then added to the sonicated mixture (glass fibers and liquid electrolytes) and further stirred inside the glove box for extra 1 hour to well-dispersed microfibers. The dispersed mixture solutions were transferred from vials by syringe and cast on 0.5" circular PDMS templates with the thickness of 150-200 μm layered on glass slides and exposed to UV-radiation (UVL-56 Lamp, λ=365 nm) in the Ar-filled glove box for 10 minutes which yielded free-standing and flexible films 150-200 μm in thickness.

2.2.3 Cathode Preparation and whole cell assembly

For cathode preparation, a carbon cloth gas diffusion layer (CCGDL) was first cut into 7/16" circular shape and dip coated by CNTs/PVDF (90:10 wt.% in NMP) slurry. Then all coated cathodes were dried at 120°C under vacuum overnight. The loading of CNTs was $0.5 \pm 0.03 \text{ mg}\cdot\text{cm}^{-2}$. For slurry preparation, 5mg of PVDF was first dissolved in 15 mL of NMP, and then 50 mg of CNTs added to PVDF/NMP solution. The CNTs/PVDF suspension in NMP was first mechanically mixed followed by bath sonication of 90 mins. The Li-O₂ batteries were prepared using modified Swagelok cells. For cell assembly, as-prepared 1/2" diameter GPEs/cGPEs or soaked Celgard polypropylene separators with liquid electrolyte was placed between 1/2" diameter lithium foil as an anode and 7/16" diameter CNT-CCGDL cathode. Before cell assembly, all cathodes were soaked with 20 μL liquid electrolyte. The amount of liquid electrolyte (20 μL) added to soak the cathodes were obtained based on trial and error. The whole cell was air-tight except for cathode side, which was fitted with stainless steel tube served as oxygen gas inlet. Ultra-high purity oxygen gas (Airgas, purity > 99.994%) was delivered into individual cell via gas manifolds containing humidity-resistant tubing. Throughout all tests under oxygen atmosphere, the flowing oxygen gas was kept at 5 psi gauge pressure via series of humidity-resistant valves/regulators. Figure 2-1 shows the modified Swagelok cell used in this study in assemble and disassemble mode. Figure 2-2 also showed the flowing-mode Li-O₂ battery testbed designed and built in the Lab.

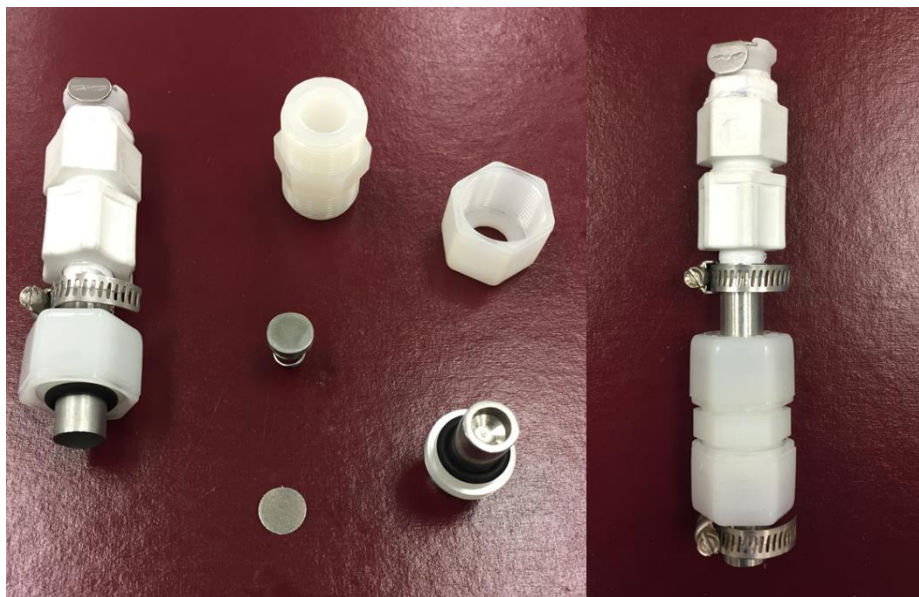


Figure 2-1: The modified Swagelok cell designed for Li-O₂ batteries in disassembled (left) and assemble (right) mode.

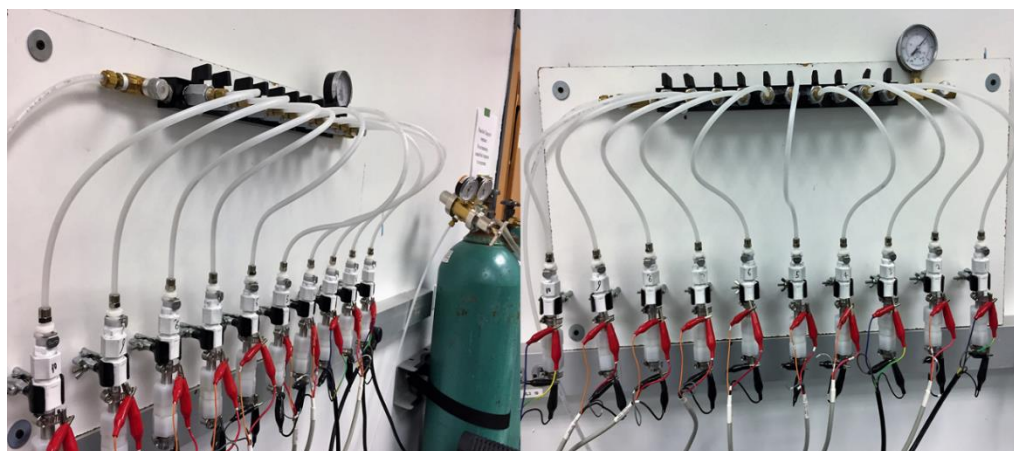


Figure 2-2: The flowing mode Li-O₂ battery testbed designed and built in this study

2.3 Electrochemical Characterizations

2.3.1 Galvanostatic charge/discharge

Galvanostatic charge/discharge is an electrochemical test to determine the capacity of the batteries. In a discharge process, a constant negative current is applied to the cell until a predefined cut-off voltage is reached and then during charging a current is reversed, and a positive current is applied to the cell. By this technique, the electrochemical performance of cell can be defined in a plot of cell voltage (volt) versus capacity (mAh). For the sake of consistency, the capacity of Li-O₂ batteries was normalized to the mass of active materials (CNTs) of the cathode and defined in mAh·g⁻¹. In this dissertation, the Li-O₂ batteries underwent the galvanostatic charge/discharge cycling at a current density of 250 mA·g⁻¹ to the cut-off voltages of 2.0-4.5 V or cut-off capacity of 500 mAh·g⁻¹. Figure 2-3 shows a single galvanostatic charge/discharge cycle of the Li-O₂ battery using CNTs-coated CCGDL in a voltage window of 2.0-4.5 V at a constant current density of 250 mA·g⁻¹ (250 mA·g⁻¹* ~0.5 mg of CNTs= ~125 μA).

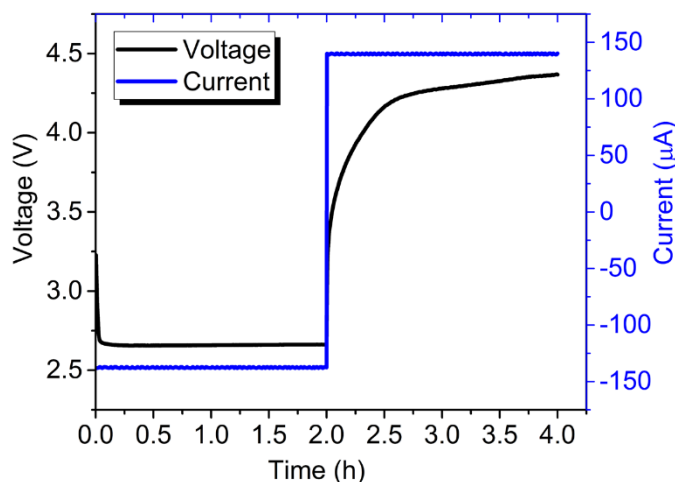


Figure 2-3: Example of typical galvanostatic charge/discharge cycling showing both the applied current and voltage response

2.3.2 Voltammetry tests

Voltammetry is the potentiodynamic technique where the potential is swept at a constant scan rate (V/s). A voltammogram is a plot of current versus voltage. Voltammetry technique is divided into two categories: Linear sweep voltammetry (LSV) and cyclic voltammetry (CV). In LSV, the voltage is only swept in one direction (cathodic or anodic) to investigate the one electrochemical reaction. However, in CV test, the voltage first is swept in one direction (forward direction) and then it is swept in a reversed direction (backward direction) to investigate the reversibility of electrochemical reactions. In this dissertation, LSV in forwarding reaction (anodic) was performed to study the electrochemical anodic stability of electrolytes when they are in contact with other components of the Li-O₂ cell. CV test was also performed on whole Li-O₂ cell to investigate the ORR and OER of cells using different electrolytes.

2.3.3 Chronopotentiometry

Chronopotentiometry is an electrochemical technique where the constant current is applied, and the voltage response is measured. In this dissertation, chronopotentiometry was also performed to investigate the oxidation stability of electrolytes. The Li-O₂ cells were assembled using actual cathodes (CNTs-coated CCGDL), Li anode and different electrolytes. Then the cells were charged without any predefined cut-off voltage before discharge process at the constant current density (250 mA·g⁻¹) used for galvanostatic charge/discharge cycling.

2.3.4 Li-ion transport measurements

2.3.4.1 Ionic Conductivity

The Ionic conductivity of electrolyte solutions is a measure of electrolyte ability to transport ions. The ionic conductivity measurement was performed by potentiostatic electrochemical impedance spectroscopy (EIS). For liquid electrolytes, the electrolytes were trapped between two stainless steel blocking electrodes by using 1 mm thick Teflon O-rings. For polymer electrolytes (GPEs and cGPEs), the electrolyte films were placed between two stainless steel electrodes. The ionic conductivity was determined using $\sigma = L/A \cdot R$, where L is the thickness of electrolyte films or the thickness of O-ring, A is the area of electrolyte films, or the area of O-ring and R is the bulk resistance obtained by high-frequency interception of EIS spectrum with abscissa. σ is the ionic conductivity in the unit of Siemens per meter (S/m).

2.3.4.2 Transference Number

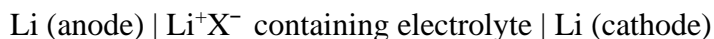
One of the most important parameters of battery electrolytes is the lithium transference number. Transference number is a dimensionless parameter which demonstrates the contribution of a particular charged species (Li^+) present in the electrolyte to the overall charge transport across the cell. In typical battery electrolytes, most of the ionic current is carried by anions, instead of cations (Li^+). In case of a simple binary electrolyte comprising of completely dissociated Li salt ($\text{Li}^+ \text{X}^-$), it can be expressed as follows:

$$t_{\text{Li}^+} = \left(\frac{I_{\text{Li}}}{I_{\text{Li}} + I_{\text{X}}} \right) = \left(\frac{I_{\text{Li}}}{I_{\text{total}}} \right) = 1 - t_{\text{X}^-} \quad (1)$$

Where t_{Li^+} is the lithium transference number, t_{X^-} is an anion transference number, I_{Li} is a current carried by Li^+ cations and I_{X} is a current carried by anions. Different methods have

been proposed to measure the transference number of electrolytes such as Hittorf method [75,76], galvanostatic polarization [77], potentiostatic polarization [78] and NMR method[77]. In this dissertation, DC polarization method modified by Bruce-Vincent has been used to determine the transference number. This method is used due to its compatibility with both liquid electrolytes and polymer electrolytes.

For the DC polarization method, the symmetrical Li/Li cell is assembled by placing the electrolyte films (GPEs or cGPEs) or presoaked Celgard separators with liquid electrolytes between two lithium electrodes as follows:



Lithium electrodes are used due to their abilities to reversibly exchange lithium ions, yet block the anions at the same time. If the symmetrical setup is polarized by applying a small DC voltage (usually 10 mV) across the cell, both the anions and cations start to move initially. Since the Li electrodes are only reversible for Li^+ ions, the moving anions are accumulated at the anode lowering the anions concentrations at the cathode, which generates a concentration gradient. Therefore, over time the initial current (I_0) starts to drop until a steady state current (I_{ss}) is reached which only originates from the non-blocking ionic (Li^+) species. This means that anions and cations at the beginning of polarization migrate together due to a DC voltage. The motion of the anions is reduced by the time during polarization and eventually comes to a complete stop. Thus, the lithium ions are the only moving species in electrolyte whereby the electric current is reduced. The first transference number measurement was made by the following formula:

$$t_{Li^+} = I_{ss}/I_0 \quad (2)$$

For the above equation, it is assumed that the current carried by lithium ions is the same at the beginning of the experiment ($t=0$) and in steady state ($t=\infty$) and furthermore are the only species that carry charge in the steady state. The above equation is for ideal cases when there is no reaction between existing species in electrolyte and Li electrodes. However, in the real cell, there is a reaction between the species in the electrolytes and Li electrodes, which leads to the formation of passivation layer known as a solid electrolyte interface (SEI) on the electrolytes. This SEI layer contributes to the ionic movement during cell polarization. Bruce and Vincent proposed a correction factor to the equation 2 to accounts this passivation film contribution [78]:

$$t_{Li^+} = \frac{I_{ss}(\Delta V - I_0 R_0)}{I_0(\Delta V - I_{ss} R_{ss})} \quad (3)$$

Where t_{Li^+} is a lithium transference number ΔV is an applied DC potential, R_0 is the initial resistance of the passivation layer, R_{ss} is a resistance of the passivation layer at steady state. So, the I_0 and I_{ss} are determined by DC polarization curve and R_0 and R_{ss} are measured by electrochemical impedance spectroscopy (EIS).

2.3.5 Electrochemical Impedance Spectroscopy (EIS)

2.3.5.1 Overview of EIS

Electrochemical impedance spectroscopy (EIS) has been extensively used in this dissertation to investigate the failure mechanism of Li-O₂ batteries using different electrolytes. The following describes some of important fundamentals of EIS, and a basic understanding of EIS is assumed.

Resistance (R) is an ability of a circuit element to resist the flow of electrical current. Based on the Ohm's law, the resistance could be defined as a ratio between voltage (V) and current (I), $R=V/I$. This relationship is for ideal resistance. An ideal resistance is defined based on a couple of assumptions: (1) it follows Ohm's Law at all current and voltage levels (2) its value is independent of frequency (3) AC current and voltage signal are in phase with each other through a resistor. Impedance is a more general circuit parameter like resistance. However, it is not limited by the simplifying properties as described for resistance. Electrochemical impedance is usually measured by applying an AC potential to an electrochemical cell and measuring the current through the cell. Electrochemical impedance is usually measured using a small excitation signal to make sure that cell's response is linear or pseudo-linear. In a linear or pseudo-linear system, the current response to a sinusoidal potential will be a sinusoid at the same frequency but shifted in phase.

If the excitation signal (potential) is expressed as a function of time [79]:

$$E_t = E_0 \sin(\omega t) \quad (4)$$

Where, ω is radial frequency. The relation between the radial frequency (ω) and frequency (f) is

$$\omega = \frac{2\pi}{f} \quad (5)$$

In a linear system, the response current is shifted in phase and can be expressed as:

$$I_t = I_0 \sin(\omega t + \varphi) \quad (6)$$

The impedance (Z) can be expressed as follows based on ohm's law:

$$Z = \frac{E_t}{I_t} = \frac{E_0 \sin(\omega t)}{I_0 \sin(\omega t + \varphi)} = Z_0 \frac{\sin(\omega t)}{\sin(\omega t + \varphi)} \quad (7)$$

So, the impedance is expressed in terms of a magnitude, Z_0 , and a phase shift, φ . Using the Euler's relationship, the impedance (Z) can also be defined as:

$$Z(\omega) = \frac{E}{I} = Z_0 \exp(j\varphi) = Z_0(\cos \varphi + j \sin \varphi) \quad (8)$$

As can be seen in equation (7), the expression for $Z(\omega)$ is composed of a real and an imaginary part. If the real part of $Z(\omega)$ is plotted on the X-axis and the imaginary part of $Z(\omega)$ is plotted on the Y-axis of a chart, a Nyquist plot can be obtained. In Nyquist plot, the value of Y-axis is negative and each point on the Nyquist Plot is the impedance at one frequency. Figure 2-4 shows a typical Nyquist plot. Another popular presentation method of impedance is a Bode plot. The impedance is plotted with log frequency on the X-axis and both the absolute values of the impedance ($|Z|=Z_0$) and the phase-shift on the Y-axis (Figure 2-4)

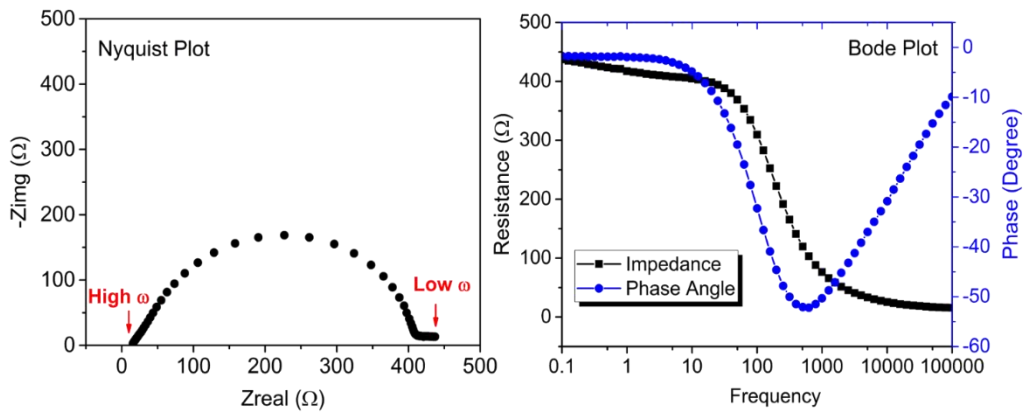


Figure 2-4: A typical Nyquist Plot (left) and Bode Plot (right)

It is worthy to mention that the EIS data are only valid if they are complied with the following criteria:

1- Linear: the system must obey Ohm's Law, $E = iZ$. The value of Z is independent of the magnitude of the perturbation.

2- Stable: the system does not change with time and returns to its original state after the excitation is removed.

3- Causal: the response of the system is due only to the applied excitation.

The Kramers-Kronig (K-K) relations can be used to evaluate the linearity, stability, and causality of EIS data. The K-K relations demand that causal, complex plane spectral data shows dependence between magnitude and phase. The K-K relations will always be true for EIS data that is linear, causal, and stable. Therefore, if the real and imaginary part of spectral data does not comply with the K-K relations, the data must violate one of these conditions.

EIS spectrum obtained from electrochemical cells can be modeled as a network of electrical circuit elements known as an equivalent circuit model. The EIS response of an equivalent circuit can be measured and compared to the actual EIS response of the electrochemical cells. Based on the EIS response of electrochemical cells, different equivalent circuit model can be estimated. Each equivalent circuit model consists of some electrical elements. Table 2-1 shows some of the circuit elements commonly used to fit the equivalent circuit models [79].

Table 2-1: Common circuit elements used in EIS equivalent circuit models

Electrical Element	Impedance
R (Resistance)	R
C (Capacitance)	$1/j\omega C$
W (infinite Warburg)	$1/Y_0(j\omega)^{0.5}$
CPE (Constant phase element)	$1/Y_0(j\omega)^a$

2.3.5.2 Transmission Line Model (TLM) for porous systems

In the classical EIS measurement, a faradic electrochemical reaction at the planer electrode/electrolyte interface can be modeled by the Randles model as shown in Figure 2-5.

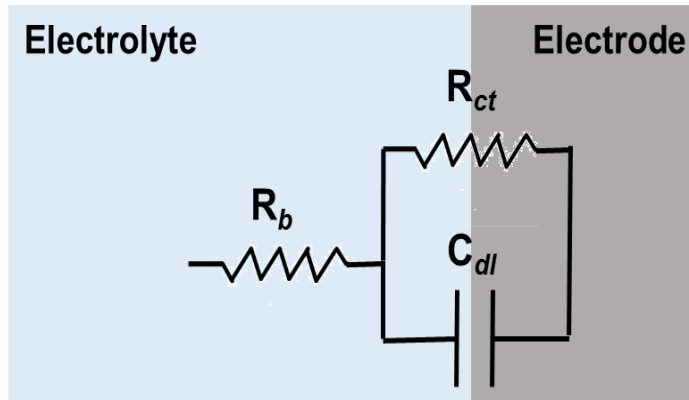


Figure 2-5: Typical circuit model (Randles model) describing the faradic process at planer electrode/electrolyte interface.

In the Randles model, the R_b represents the resistances from the electrolyte and R_{ct} represents all faradaic reactions that occur on the electrode's surface. C_{dl} describes nonfaradic capacitive charge storages [80]. The C_{dl} is often replaced by a CPE element for non-ideal capacitive behavior. Although, this model shows electrochemical interfaces of planar electrodes, it describes poorly the effect of porous electrodes that are used in most

electrochemical cells. In porous electrodes, mainly four electrochemical processes happen as shown in Figure 2-6: (1) mixed ions and electrons conduction as electric resistance (R_e), electrolyte bulk resistance (R_{sol}), and ionic resistance in pores (R_{ion}); (2) formation of an electric double layer at the electrode/electrolyte interface (C_{dl}); (3) charge-transfer reaction for faradic reactions as R_{ct} ; and (4) mass transfer to compensate for charge as diffusion [81].

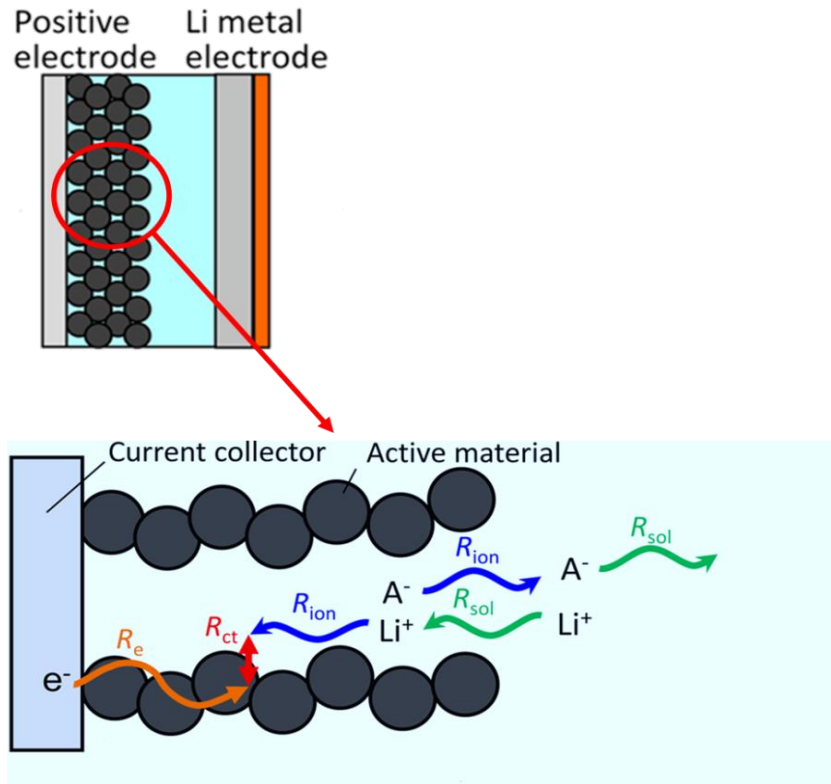


Figure 2-6: Schematic illustration of different resistances in porous electrodes [81]

To address these resistances in porous electrodes, the pores within porous electrode are modeled in a cylindrical shape with the following assumptions: Highly porous cathodes consist of the base electrode and the porous active electrode. The base electrode is an

inactive conductive material (metal foil or carbon cloth) where the porous active materials (electrodes) are deposited on it. Figure 2-7 shows a typical cylindrical pore of active porous materials on the base electrode flooded by the liquid electrolyte.

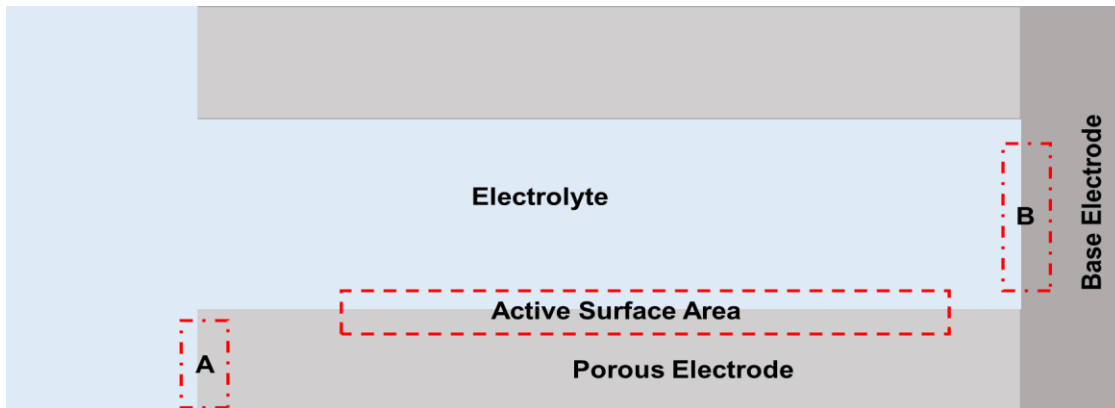


Figure 2-7: Different interfaces in porous electrode structure

Unlike the planar electrode where the electrochemical reaction happens on the surface of electrode, the rate of electrochemical reaction in porous cathodes is limited. This limitation arises from the fact that the accessibility of ions to the active interface is hindered by the small inner volume of the pores. Hence the diffusion rate of the ion in the pore becomes the dominating step in electrochemical reactions. The porous electrode could be categorized into three regions due to the restrictions of the electrochemical reactivity in porous electrodes. These interfaces are labeled as “A”, “B”, and “Active Surface Area” (see Figure 2-7) [82]. “A” represents the interface between the outer surface of the pores of porous electrode and the electrolyte. “B” describes interactions between base electrode and electrolyte. “Active surface area” describes the interactions between active materials of the porous electrode and electrolyte [80]. Transmission line model (TLM) is one the most promising circuit models to describe the electrochemical behaviors of those interfaces

existing in porous electrodes. Figure 2-8 shows a TLM model in a generic form. The model consists of several parallel and serially connected elements.

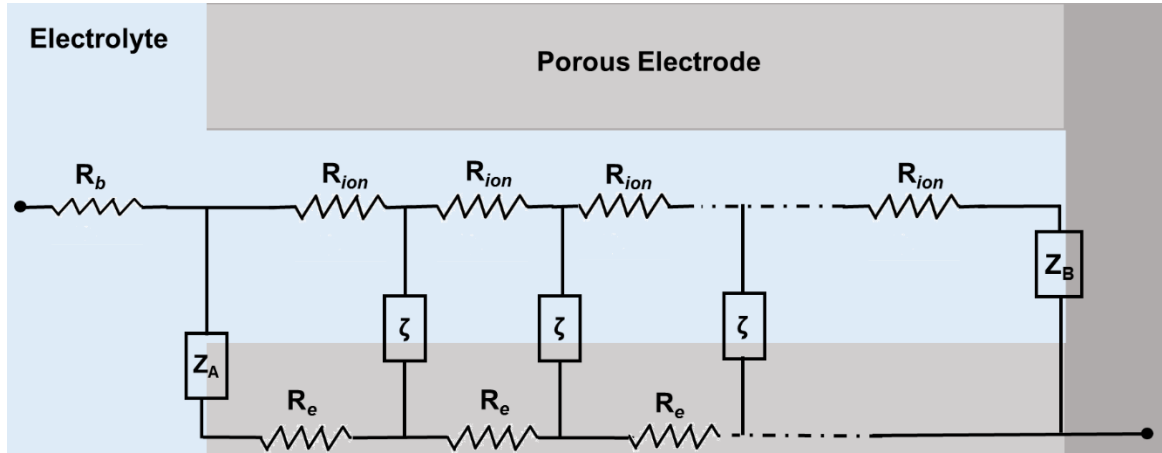


Figure 2-8 Scheme of a generic transmission line model

The interfaces “A” and “B” are represented by impedances Z_A on the outer surface of the pore and Z_B on the base electrode at the end of the pore, respectively. R_{ion} is the impedance of the electrolyte within the pore. It is important to mention that this impedance is different from bulk electrolyte resistance (R_b). R_e is the impedance of the porous electrode’s solid phase. ζ describes the impedance at the “Active surface area” as shown in Figure 2-8 [82]. The generic form of TLM is usually simplified due to the existing boundary conditions in real electrochemical cells. Bisquert [83,84] describes the simplified TLM for electrochemical energy storage systems. Based on their assumptions, the ionic resistance inside the pores are much higher than electronic resistance due to the electronic conduction in active materials used in batteries, so the resistive trail of R_e can be set as zero. Besides, it is assumed that the electrochemical reactions only happen in the active surface area and

no reaction occurs on the outer surface of electrode/electrolyte and base electrode/electrolyte interface. Hence both Z_A and Z_B is infinite (open-circuit). Applying the boundary condition, the generic form of TLM can be described as in Figure 2-9 for both faradic and nonfaradic reactions.

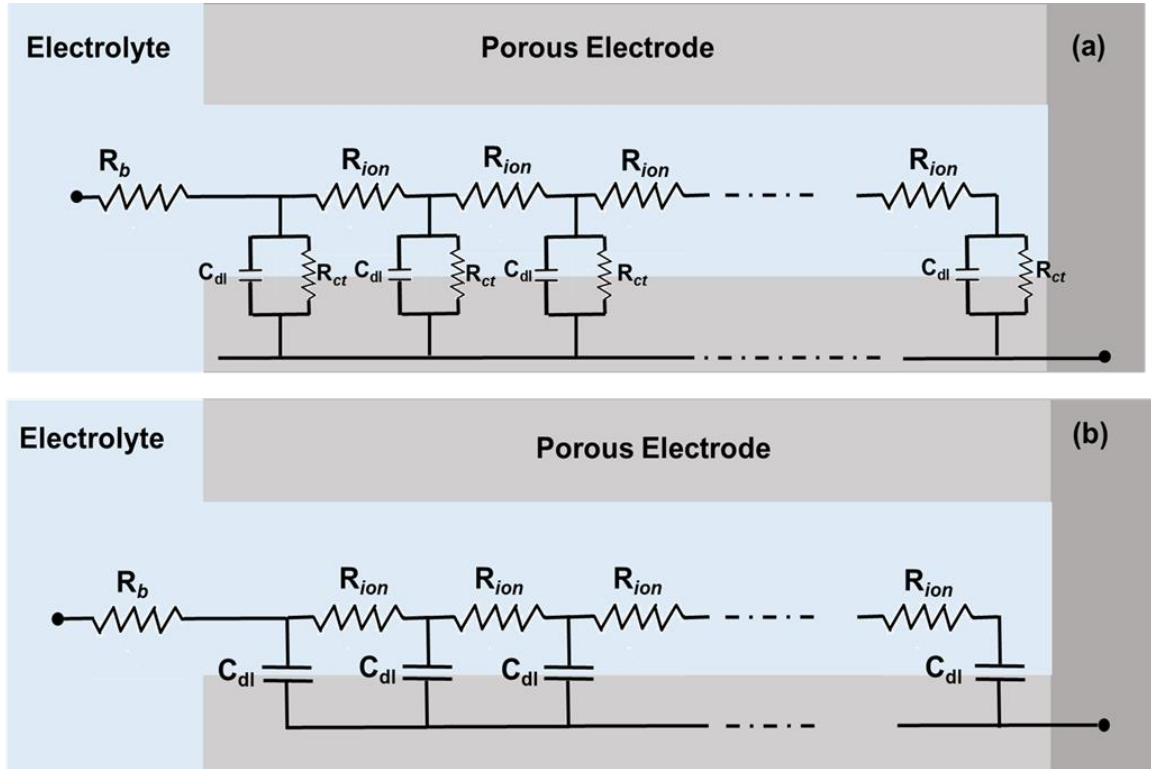


Figure 2-9: (a) Simplified TLM for porous cathodes describing faradic reactions at the active surface area (b) Simplified TLM for porous cathodes describing nonfaradic reactions at the active surface area

For faradic and non-faradic reactions at cylindrical porous electrodes, the overall impedance can be expressed as follows [80,81]:

$$Z_{\text{faradaic}} = \sqrt{\frac{R_{\text{ion},L} \cdot R_{\text{ct},A}}{(1 + j\omega R_{\text{ct},A} \cdot C_{\text{dl},A}) \cdot 2\pi r}} \coth \sqrt{\frac{R_{\text{ion},L} \cdot (1 + j\omega R_{\text{ct},A} \cdot C_{\text{dl},A}) \cdot 2\pi r}{R_{\text{ct},A}}} L \quad (9)$$

$$Z_{\text{nonfaradaic}} = \sqrt{\frac{R_{\text{ion},L}}{j\omega C_{\text{dl},A} \cdot 2\pi r}} \coth \sqrt{R_{\text{ion},L} \cdot j\omega C_{\text{dl},A} \cdot 2\pi r} L \quad (10)$$

Where, L and r are the length and radius of cylindrical pores, respectively.

For nonfaradic reactions, at the high frequency, the EIS spectrum shows a linear region with a 45-degree slope from the real axis followed by a vertical rise (Figure 2-10). The limiting value for Z_{real} and Z_{img} as $\omega \rightarrow 0$ is [81,85]:

$$Z_{\text{real}}(\omega \rightarrow 0) = \frac{R_{\text{ion}}}{3} \quad (11)$$

$$Z_{\text{img}}(\omega \rightarrow 0) = \frac{1}{\omega C_{\text{dl}}} \quad (12)$$

For faradic reaction, at high frequency, the EIS spectrum shows a linear region with a 45-degree slope from the real axis followed by the semi-circle at the low frequency (Figure 2-10). The limiting value for Z_{real} and Z_{img} as $\omega \rightarrow 0$ is [81,85]:

$$Z_{\text{real}}(\omega \rightarrow 0) = \frac{R_{\text{ion}}}{3} + R_{\text{ct}} \quad (13)$$

$$Z_{\text{img}}(\omega \rightarrow 0) = 0 \quad (14)$$

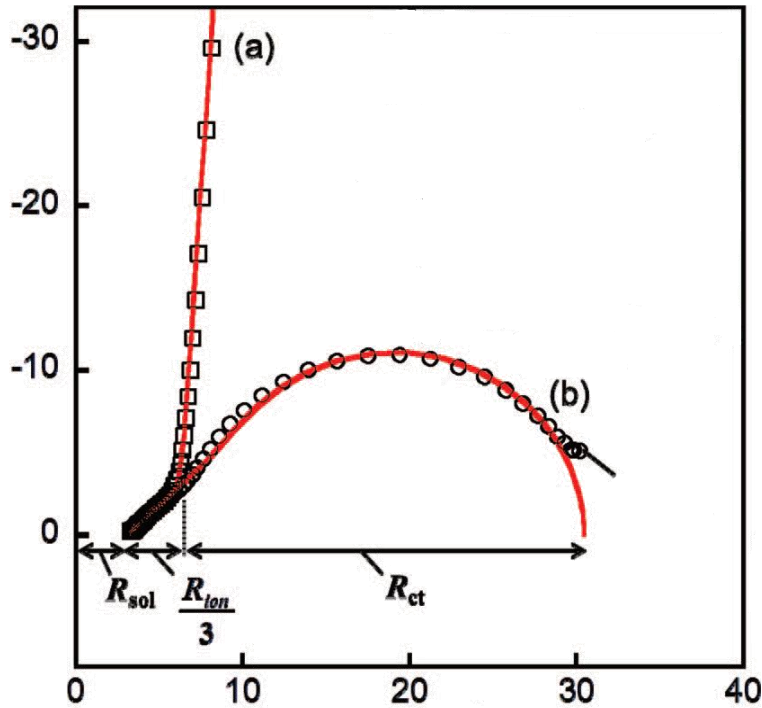


Figure 2-10: Typical Nyquist plots for cylindrical pores in (a) nonfaradic reactions and (b) faradic reactions [85]

2.4 Characterization techniques

2.4.1 Scanning Electron Microscopy (SEM)

Scanning Electron microscopy (SEM) is powerful imaging technique which could provide vast information about the morphology and topography of samples. In this technique, the sample is bombarded by high energy electrons and different electrons (secondary (SE) and backscattered electrons (BSE)) or characteristic X-rays is produced. These electrons and X-rays can be used to analyze the sample. In this thesis, SE electrons only used to investigate the morphology of glass microfibers and discharge products. To investigate the morphology of discharge products, the cathodes after discharge/charge process were harvested from Li-O₂ cells in the Ar-filled glovebox and rinse with TGDME solvent to

remove all residue of LiTFSI salt. Then samples were transferred to the vacuum chamber connected to the glovebox and dried at room temperature. Dried samples were also left in glovebox for a period to make sure that they are completely dried. To avoid any moisture or oxygen contamination, the SEM sample holder was first placed in glovebox and cathodes are fixed on the sample holder by double-sided copper tape. SEM sample holder with mounted cathode samples was then placed in Ar-filled bags and transferred to the microscopy room. The time from removing the sample from Ar-filled bags to the SEM was less than a minute to avoid any oxygen and humidity exposure. In this thesis, JEOL Multi-Beam FIB 4500 SEM at low vacuum was used to take micrographs of glass microfillers and cGPE. JEOL SEM 7000 was also used to investigate the cathodes. For the cGPE investigation, cGPE was first placed in acetone solvent for one day to remove all TEGDME liquid electrolytes inside cGPE. Then cGPE without TEGDME solvent was placed in a vacuum chamber to remove all acetone solvent.

2.4.2 X-Ray Diffraction (XRD)

X-ray diffraction (XRD) is very powerful and popular technique to investigate the crystalline structures of materials. In this dissertation, XRD was used to investigate the microstructures of glass microfibers and discharge products. XRD (Siemens 5000D X-ray Diffractometer) was used to obtain the microstructures of glass microfillers. For discharge products investigation, Bruker GADDS/D8 (XRD) with MacSci rotating Molybdenum anode ($\lambda = 0.71073 \text{ \AA}$) operated at 50 kV generator, and 20 mA current was also used to collect the diffraction pattern of discharge products. A parallel X-ray beam in size of 100 μm diameter was directed on to the cathode samples, and diffraction intensities were recorded on large 2D image plate during the exposure time.

2.4.3 Raman Spectroscopy

Raman spectroscopy is a spectroscopic technique commonly used to obtain the fingerprint of different chemical structures. This technique depends on inelastic scattering of monochromatic light, usually a laser. A laser light interacts with molecular vibrations and its energy being shifted. The shift in energy of the laser gives information about the vibrational mode of the chemical species. For experimentation, the sample is illuminated by the laser beam and reflected radiation from that illuminated spot is collected by a lens and transferred to the monochromator. Elastically scattered radiation at the wavelength corresponding to the laser line is filtered out by either a filter, while the rest of the collected light is dispersed onto a detector. In this dissertation, the Raman spectroscopy (BaySpec's NomadicTM, an excitation wavelength of 532 nm) was used to investigate the discharge products on the cathode samples. Raman was also performed on the liquid electrolytes, GPEs, and cGPEs to investigate the interaction of ETPTA polymer and glass microfillers with LiTFSI salt and interaction of LiTFSI salt with TEGDME molecules at different salt concentration.

2.4.4 Brunauer–Emmett–Teller (BET)

Brunauer-Emmett-Teller (BET) surface analysis was also used in this dissertation to investigate the surface area of the glass microfillers. The isotherm absorption-desorption of N₂ at 77K was performed using Tri-Star II Micromeritics.

2.4.5 Thermal Analyses

Thermogravimetric Analysis (TGA) and Differential scanning calorimetry (DSC) was performed in this dissertation using SDT Q600. TGA was used to examine the purity of

glass microfillers used in this study. SDT was also carried out on the ETPTA and GPEs to determine the glass transition of the polymer.

3 COMPOSITE GEL POLYMER ELECTROLYTE: EFFECT OF INORGANIC FILLERS CONTENT

3.1 Background

As mentioned earlier in this dissertation, the Li-O₂ batteries suffer from poor cyclability due to the reactivity of lithium anode with oxygen crossover, cathode decomposition, and electrolyte evaporation and decomposition [11,22,30,46,86–88]. Electrolyte decomposition has been previously reported to yield the formation of solid by-products covering gradually the surface of electrodes and causing rapid capacity fading in Li-O₂ batteries [31,48]. It has been suggested that some electrolyte properties such as ionic conductivity, lithium transference number, and electrolyte-electrodes interface could affect the electrochemical performance of Li-O₂ batteries [46,89,90]. Despite many attempts aiming at developing stable electrolytes for Li-O₂ battery application, the choice of solvent and salt of electrolyte remains one of the biggest challenges to develop reliable Li-O₂ batteries. Ceramic electrolytes and solid polymer electrolytes (SPEs) have been used in Li-O₂ batteries to overcome the liquid electrolytes' drawbacks. However, their high interfacial resistance and low ionic conductivity limit their practical applications at ambient temperatures [53]. Gel polymer electrolytes (GPEs) composed of liquid electrolytes in polymer matrices have been successfully used in Li-ion battery applications due to their high ionic conductivity and low interfacial resistances [91–94]. GPEs of different polymer-solvent pairs have been explored in Li-O₂ batteries and were shown to efficiently protect

the lithium anode from oxygen crossover, and limit electrolyte evaporation [63,64,95–100]. The incorporation of inorganic fillers to both liquid and polymer electrolytes (SPEs and GPEs) has been shown to improve the Li^+ transport properties such as ionic conductivity and lithium transference number through the interaction of fillers with the polymer, solvent, or salt [101–108]. Although the incorporation of some inorganic fillers into different GPEs for Li-O₂ battery application have also been initiated [109–111], most research studies have emphasized on zero-dimensional ceramic particles (e.g., nano and microparticles), and little attention has been given to one-dimensional fillers. In the present study, we investigate the influence of one-dimensional glass micro-fillers in gel polymer electrolytes using ethoxylated trimethylolpropane triacrylate polymer and tetraglyme-based solvent for the Li-O₂ battery application. Discharge/charge cycling of Li-O₂ batteries using gel polymer electrolyte with different glass micro-filler contents along with different electrochemical and microstructural characterization techniques have been used to evaluate the performance metrics of composite GPEs.

3.2 Experimental details

The GPEs and cGPEs were prepared as described in chapter 2. Figure 3-1 shows the schematic of GPE and cGPEs (1 mol·g⁻¹ LiTFSI salt concentration) preparation with their photographs after UV curing process. The glass microfibers content in cGPEs were changed from 0.5, 1, 2, 5 wt.% (hereinafter cGPEs-0.5%, cGPE-1%, cGPE-2% and cGPE-5%).

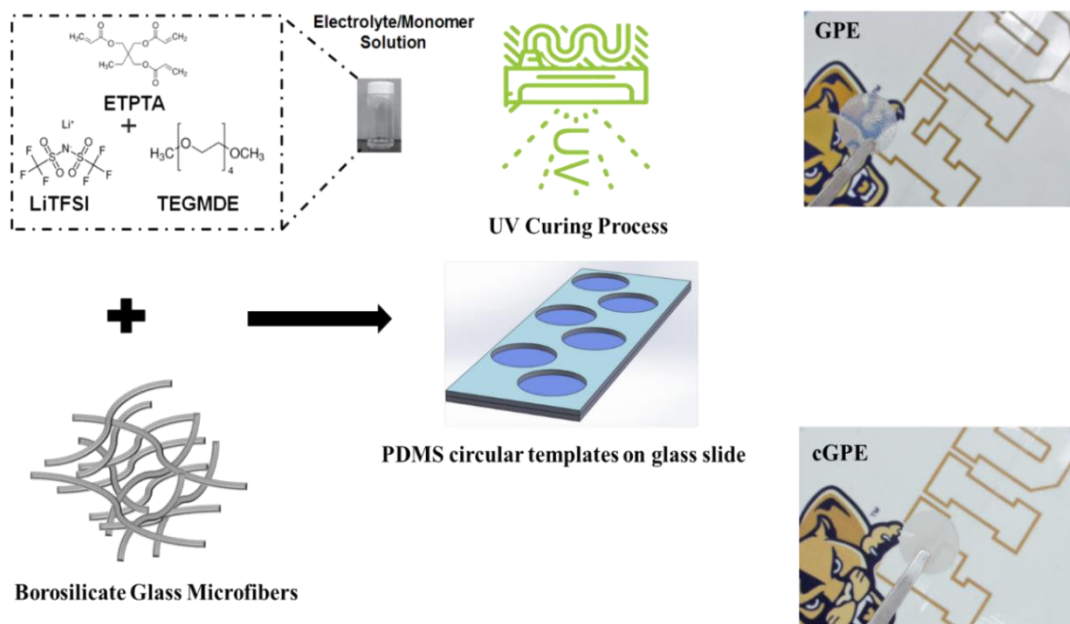


Figure 3-1: Schematic representation illustrating the GPE and cGPEs preparation along with photographs depicting the physical appearance of GPE and cGPE-1%

The ionic conductivity was measured by trapping the GPE and cGPEs with different content of glass microfibers between two blocking stainless steel as described in chapter 2. The content of ETPTA monomer to TEGDME electrolyte in all GPEs in this dissertation was optimized based on their ionic conductivity and mechanical properties in such way to have the highest ionic conductivity and free-standing film. Table 3-1 depicts the ionic conductivity of GPEs with their corresponding electrolyte content. As it is clear the ionic conductivity of GPEs increase with the increase in electrolyte content. 85 wt.% of TEGDME was the highest content of electrolyte in GPEs which gives freestanding films; however, their mechanical properties were very poor, and films were easily torn (Figure 3-2).

Table 3-1: Ionic conductivity of GPEs versus their electrolyte content

Electrolyte content in GPEs (wt.%)	Ionic conductivity ($mS \cdot cm^{-1}$)
100 (pure electrolyte)	2.56
15	$6.35 \cdot 10^{-9}$
20	$3.4 \cdot 10^{-4}$
50	0.016
75	0.5
80	1.0
85	1.1

Hence, the GPEs with the ratio of 20:80 (ETPTA: Electrolyte) by weight were prepared and investigated in the entire dissertation.



Figure 3-2: GPEs with 85 wt.% electrolyte content showing poor mechanical properties

Li^+ transference number was also determined by the Bruce-Vincent method as described in chapter 2. Two-stage DC polarization [112] was utilized in this dissertation to accurately determine the I_0 . The first stage ran with the fast sampling rate (intervals between measurements at 10 ms) lasting for 80 seconds. The external potential step is applied with

a delay of about 20 seconds in respect to starting of this stage to properly determine initial current. The second stage executed immediately after the first stage with the slower sampling rate (intervals between measurements of 1 s). This step ran until a steady-state current (I_{ss}) is reached. CNTs coated carbon cloth gas diffusion layer (CCGDL) cathodes were also prepared as described in experimental section. Figure 3-3 illustrates the optical and SEM images of pristine and CNTs-coated CCGDL.

Galvanostatic charge/discharge tests were performed using MTI battery tester at the constant current density of $250 \text{ mA}\cdot\text{g}^{-1}$ within the voltage range of 2.0–4.5 V. In-situ EIS measurements were also conducted using a Gamry Reference 600 in the frequency range of 100 kHz to 100 mHz using 10% of DC discharge current during cycling test. All charge/discharge and EIS studies were carried out at 25°C. Cathodes after charge/discharge were also extracted from Li-O₂ cells and characterized by Raman spectroscopy (BaySpec's Nomadic™, an excitation wavelength of 532 nm) and XRD (Bruker GADDS/D8 with MacSci rotating Molybdenum anode ($\lambda = 0.71073 \text{ \AA}$)).

3.3 Results and Discussion

The SEM micrograph of the glass microfibers in Figure 3-4 confirms the one-dimensional morphology of the glass microfillers with an average diameter of approximately 1 μm and an aspect ratio exceeding 100. Figure 3-4 also shows the microstructural characterization of glass microfillers using XRD, BET, and TGA in addition to depicting the cross-section of cGPE-1%. XRD pattern of the microfibers exhibited a broadened peak, demonstrating that the glass fillers used in this work have an amorphous phase. The surface area of microfibers was measured to be around $0.5 \text{ m}^2 \cdot \text{g}^{-1}$ by N_2 adsorption–desorption isotherm at 77 K, indicating that the microfibers were non-porous. The purity of the microfibers was confirmed using TGA analysis up to 1000°C with no detectable humidity and impurities. A cross-sectional SEM image of the cGPE-1% (Figure 3-4 (e-f)) shows that the microfibers are uniformly distributed with no apparent agglomeration in any direction.

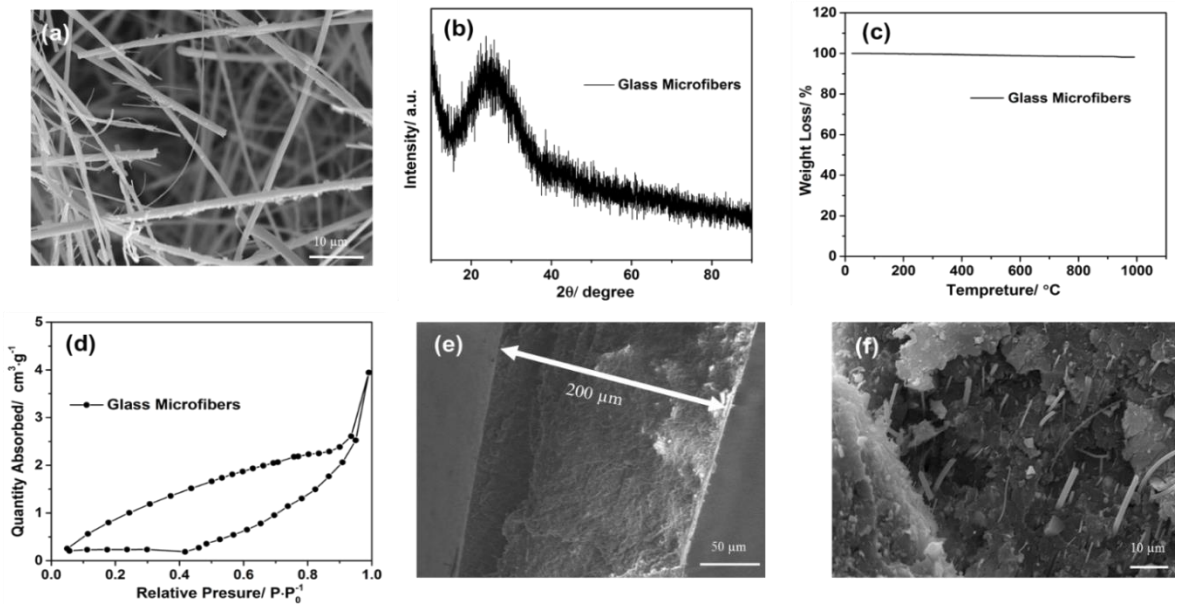


Figure 3-4: (a) SEM micrograph (b) XRD pattern (c) TGA analysis (d) BET surface area measurement of glass microfibers (e-f) the SEM micrographs of cross-sectional cGPE-1%

The polymer ETPTA was selected in this dissertation due to its following advantages. (1) The simplicity of the preparation of GPEs [113] (2) electrochemical stability in the voltage range of 2.0-4.5 V [95] (3) high mechanical properties which afforded high electrolyte contents (80%) in the GPEs [113] and (4) its demonstrated compatibility with Li-O₂ battery components [97]. Thermal properties of ETPTA polymer was also measured. Figure 3-5 shows the thermal stability and glass transition (T_g) of ETPTA polymer used in this dissertation. As it is clear, the ETPTA polymer is stable up to ~350°C suggesting that the ETPTA is thermally stable and can protect the battery from thermal runaway. The glass transition of ETPTA is also measured to be 73°C [113]. The high glass transition of ETPTA demonstrates a little contribution of ETPTA into ion conduction in GPEs and cGPEs.

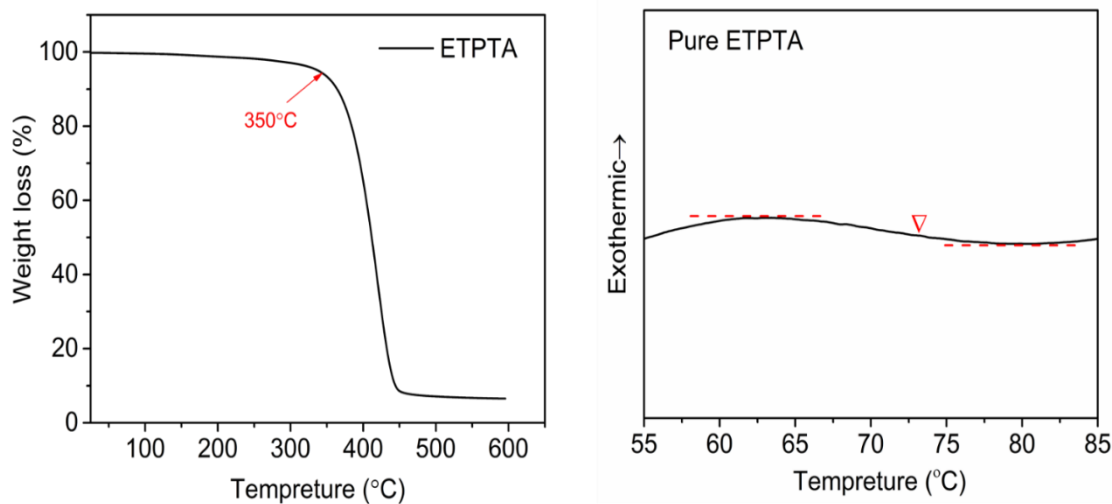


Figure 3-5: Thermal properties of ETPTA polymer (a) TGA analysis showing the thermal stability of polymer (b) DSC analysis showing the glass transition (T_g) of ETPTA polymer

The ionic conductivity (σ) and lithium transference number (t_{Li^+}) of the GPE and cGPEs were measured at room temperature (25°C), and the results are summarized in Table 3-2.

Figure 3-6 shows typical Chronoamperometric curves of symmetrical Li/Li cells using GPE and cGPE-1% after 10 mV of DC polarization along with the corresponding impedance spectra before and after polarization.

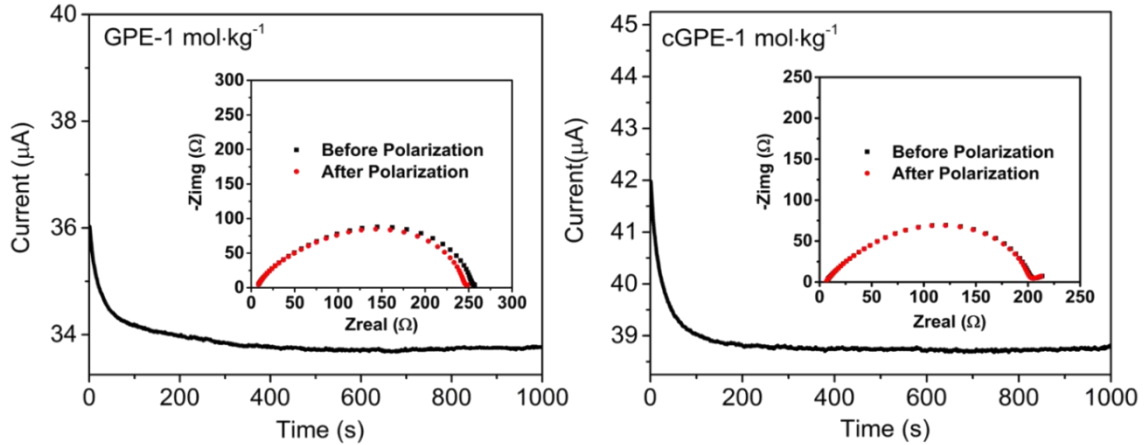


Figure 3-6: Chronoamperometric curves of Li/GPE and cGPE-1% /Li cell after 10 mV of DC polarization at 25°C where the electrolyte films contain 1 mol·kg⁻¹ LiTFSI salt. Insets: electrochemical impedance spectra before and after polarization.

As can be seen from Table 3-2, the GPE shows a good ionic conductivity of 1.02 mS·cm⁻¹ and a t_{Li^+} of 0.53. The transference number of GPE is in agreement with previously reported values for glyme-based GPEs [114,115]. The ion conduction contribution of ETPTA in GPEs was negligible. The LiTFSI salt was added to ETPTA polymer to make solid polymer electrolytes (SPEs), and their ion conductivities were also measured. In the saturated LiTFSI salt concentration, the ionic conductivity of SPE at 25°C was measured to be around 0.005 mS·cm⁻¹. Thus, comparing the ionic conductivity of GPE and SPE at 25°C confirms that ETPTA does not significantly contribute to ion conduction in GPEs. Upon addition of the micro-fillers, both σ and t_{Li^+} increased with the increase in fillers content until 1%, then started to drop upon increasing the fillers content. This increase is attributed to the interaction between the filler materials and the ions in the cGPE, namely the

adsorption of the TFSI counter-ions on the surface of glass microfibers and the formation of ion-ceramic complexes. This interaction increases the ion pair dissociation as previously reported [73,101,108,116]. This dissociation increases the free Li^+ , affording them to be transported easier through the percolating pathways of the microfibers. This means that the Li^+ transport enhancement is dependent on the formation of continuous filler network [117,118]. Hence filler loading more than the percolation threshold is needed. Conversely, excessive filler loading led to their aggregation and thus blockage of the conducting pathways of Li^+ ions [108,117,118]. This trade-off in effect of the addition of fillers suggests the existence of an optimum loading. The optimum loading amount was determined to be 1% in this work.

Table 3-2: Summary of ionic conductivity and lithium ion transference numbers of GPE and cGPEs with different glass micro-filler contents.

Gel Polymer Electrolytes	Transference Number, t_{Li^+}	Ionic Conductivity, σ (mS/cm)	Li^+ Conductivity, σ_{Li^+} (mS/cm)
GPE	0.53	1.02±0.05	0.54
cGPE-0.5%	0.58	1.12±0.02	0.65
cGPE-1%	0.66	1.40±0.02	0.92
cGPE-2%	0.52	0.95±0.05	0.50
cGPE-5%	0.48	0.75±0.02	0.36

The oxidation stability limit of GPE and cGPE-1% was also determined using a chronopotentiometric stability test and linear sweep voltammetry (LSV) under the oxygen atmosphere (Figure 3-7). For the chronopotentiometric test, Li-O₂ batteries using GPE and cGPE-1% were assembled and charged with no cutoff voltage at the current density of 250 mA·g⁻¹ without prior discharging for 10 h. For LSV tests, GPE and cGPE-1% were placed

between lithium foil as a reference and counter electrode, and a standard cathode (MWCNTs-CCGDL) as a working electrode. Voltage was scanned at a rate of $1 \text{ mV}\cdot\text{s}^{-1}$.

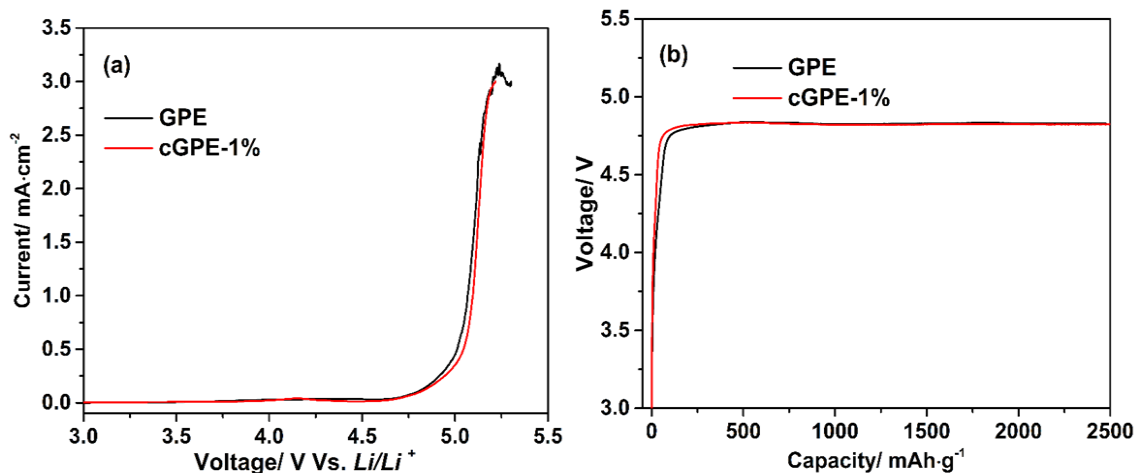


Figure 3-7: (a) Linear sweep voltammograms for Li-O₂ batteries using GPE and cGPE-1% with the scanning rate of $1 \text{ mV}\cdot\text{s}^{-1}$ under oxygen. (b) Chronopotentiometric test of Li-O₂ batteries using GPE and cGPE-1% at $250 \text{ mA}\cdot\text{g}^{-1}$ for 10 hours.

From the voltammograms and chronopotentiometric tests, the microfibers do not change the oxidation stability of GPE [119], as GPE and cGPE-1% show anodic stabilities up to 4.75 V. Furthermore, anodic stability at around 4.75 V for GPE and cGPE-1% suggests that the polymer used in this study is electrochemically stable under oxygen environment since the anodic stability of 1M LiTFSI in TEGMDE has been also reported at 4.75 under oxygen (Figure 3-8) [48].

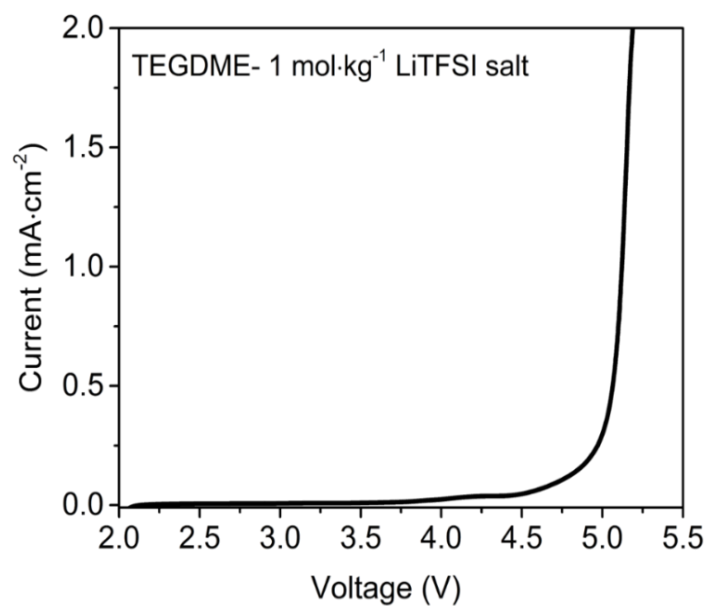


Figure 3-8: Linear sweep voltammogram of Li-O₂ battery using liquid electrolyte with 1 mol·kg⁻¹ LiTFSI

To provide more details on the contribution of microfibers in GPE on Li-O₂ battery performance, galvanostatic discharge/charge cycling studies were performed at current densities of 250 mA·g⁻¹ (0.125 mA·cm⁻²) and fixed cycle capacities of 500 mAh·g⁻¹ in the voltage window 2.0-4.5. To initiate the charge/discharge experiment, the Li-O₂ cells were rested under pressurized extra pure oxygen gas (5 psi gauge pressure), and open circuit voltage (OCV) of cells was monitored. In our experiment, it turned out that the rate of OCV change after 5 hours is less than 5 mV·h⁻¹ (Figure 3-9).

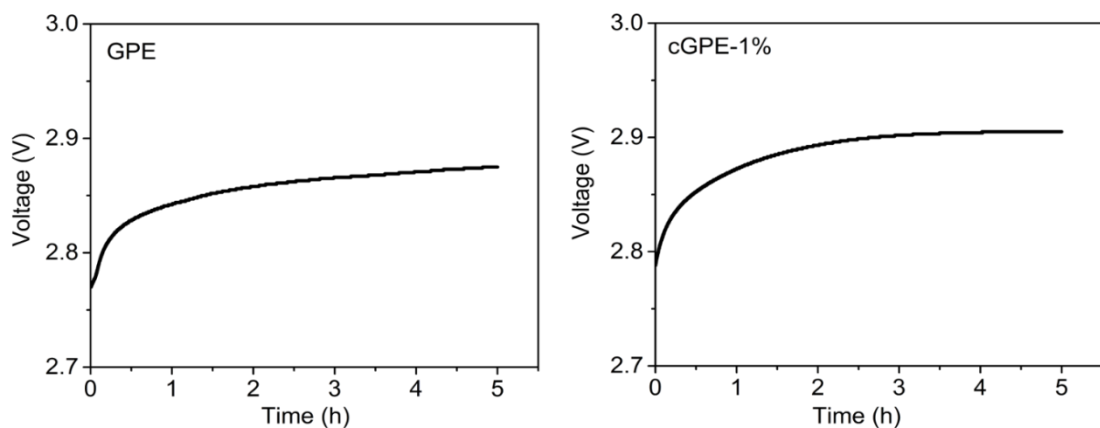


Figure 3-9: Open circuit voltage (OCV) of Li-O₂ cells using GPE and cGPE-1% under oxygen versus time

Hence, 5 hours of resting was considered as the relaxation time for all Li-O₂ cells before charge/discharge cycling test. From Figure 3-10, cells using cGPE-1% show the highest discharge cycling performance with a median performance of 54 consecutive discharge cycles comparing to cGPE-0.5% of 38, cGPE-2% of 30 cycles, cGPE-5% of 27 cycles, and the no-filler GPE of 29 cycles. Similar to the ionic conductivity and transference number experiments, cGPE-1% had the best performance, indicating the impact of the improvement of the ionic conductivity and transference number of the cGPE on the full cell. This observation is in agreement with previous studies showing that electrolytic properties in Li-O₂ batteries such as ionic conductivity, lithium transference number, and stability of interfacial resistances play a significant role in Li-O₂ battery behavior [46,89,90].

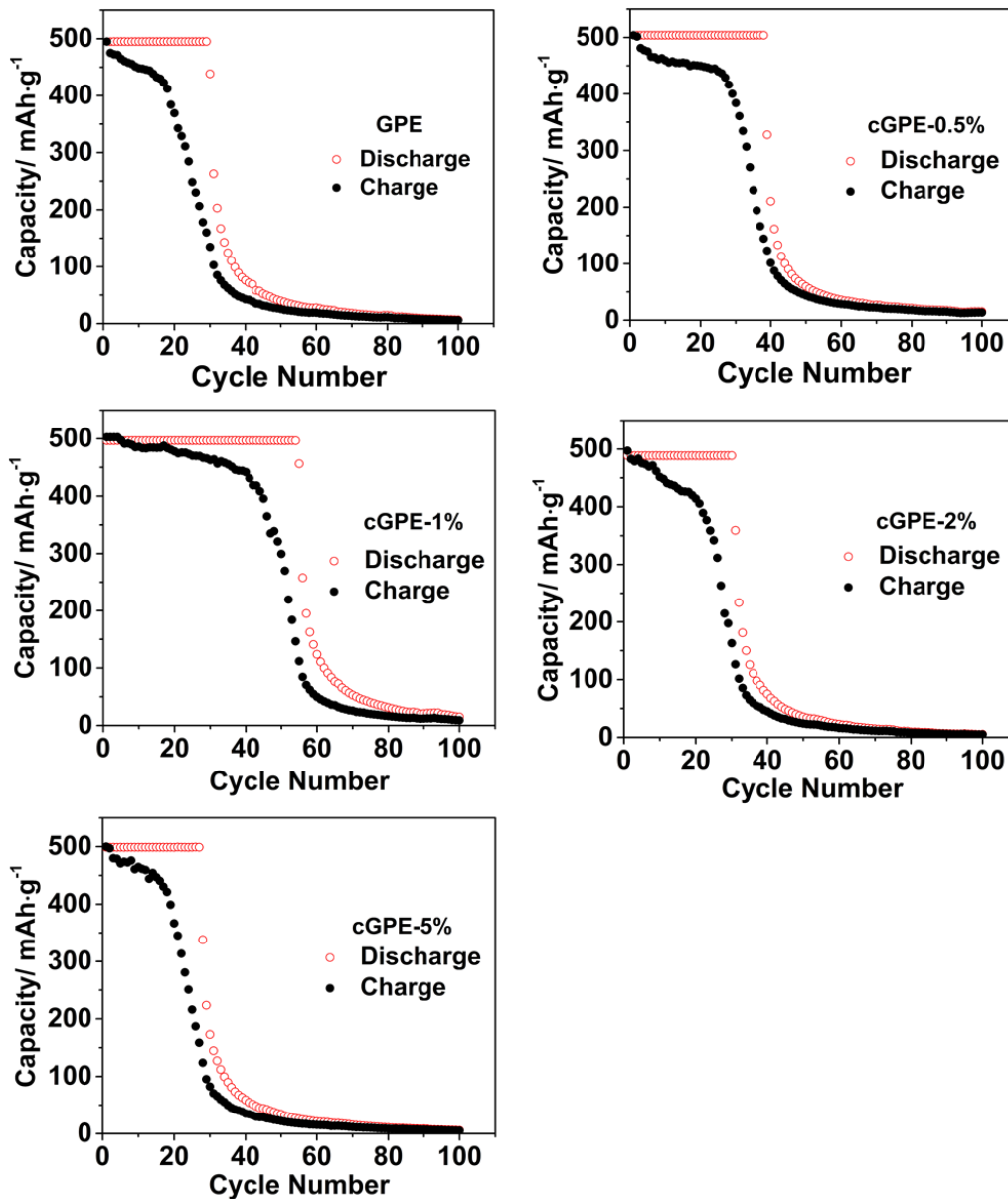


Figure 3-10: Cyclability of the Li-O₂ batteries for fixed charge and discharge cycle capacities of 500 mAh·g⁻¹ at a current density of 250 mA·g⁻¹ with voltage cutoffs of 2.0–4.5 V for GPE and cGPEs with various glass microfibers contents.

Figure 3-11 also shows the voltage profile of Li-O₂ batteries using GPE and cGPEs with different glass microfibers at current density of 250 mA·g⁻¹ (0.125 mA·cm⁻²) and at fixed cycle capacities of 500 mAh·g⁻¹ in the voltage window 2.0-4.5. As can be seen, the charge

and discharge voltage plateau of cells using GPE and cGPEs does not change significantly confirming that the glass microfibers do not interfere with OER and ORR process.

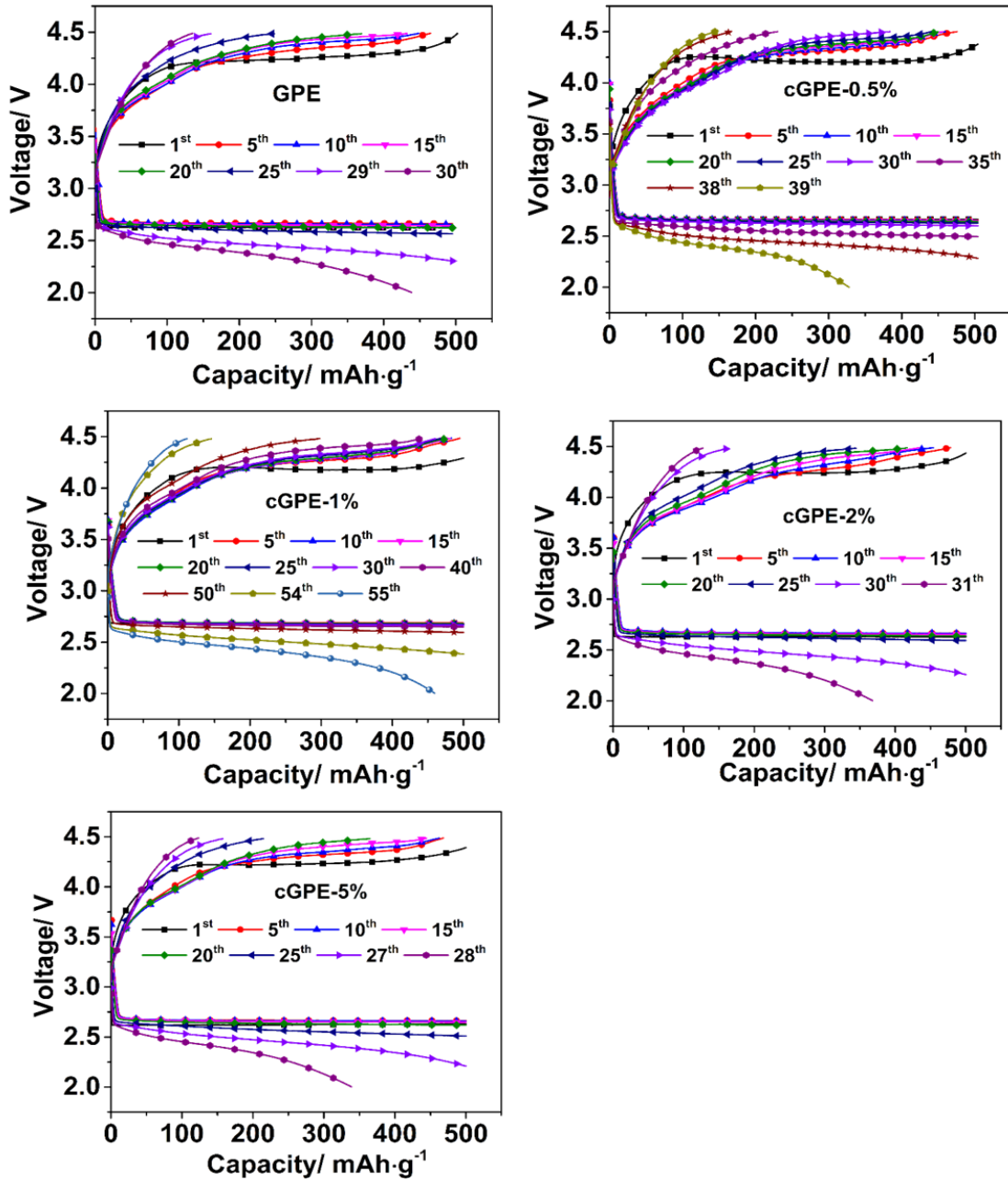


Figure 3-11: The voltage profile of Li-O₂ batteries using GPE and cGPE with different glass microfiber content

To further confirm the contribution of glass microfibers in ORR and OER, the cyclic voltammetry (CV) was performed on the Li-O₂ batteries using the GPE and cGPE-1% in a voltage windows of 2.0-4.5 V at a scan rate of 1 mV·s⁻¹ under oxygen (Figure 3-12).

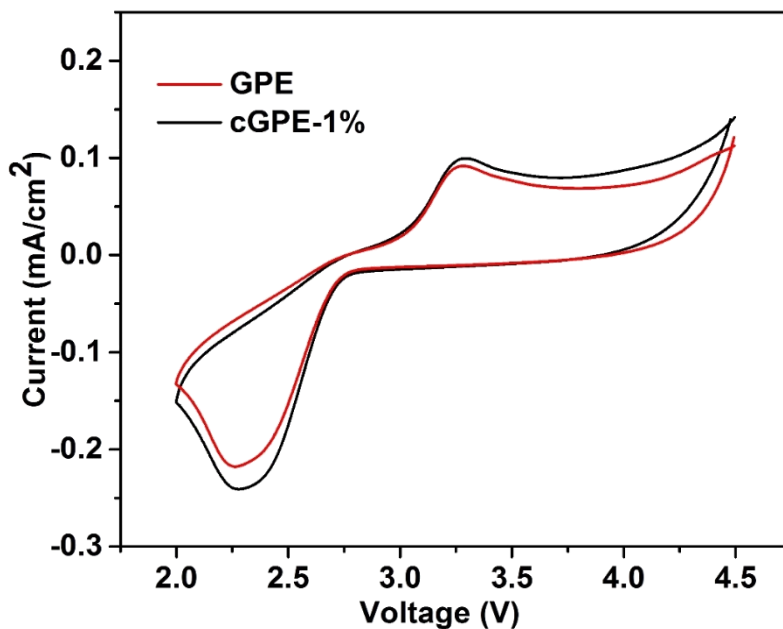


Figure 3-12: Cyclic voltammograms of Li-O₂ batteries using GPE and cGPE-1% in voltage windows of 2.0-4.5 V and scan rate of 1mV·s⁻¹

As can be seen, the onset of ORR peak occurs at 2.8 V for both GPE and cGPE-1%. The peak at 3.3 V could also be attributed to the OER. The similarity between the ORR and OER of cells using GPE and cGPE-1% proves that the glass microfibers do not alter the Li₂O₂ formation/decomposition. The slight higher cathodic and anodic current in the cell using cGPE-1% could be due to the higher ionic conductivity of cGPE-1% over GPE.

To confirm the formation of Li₂O₂ on the cathode, oxygen cathodes were investigated by XRD and Raman spectroscopy. Figure 3-13 shows the XRD patterns and Raman spectra of oxygen cathodes using both GPE and cGPE-1%. The diffraction peaks of Li₂O₂ can be observed after the discharge process suggesting that Li₂O₂ is the main product in the

discharge process with either electrolyte. Raman spectra of discharged cathodes similarly show Raman shifts at 790 cm^{-1} which corresponds to Li_2O_2 formation [120]. Raman spectra also show the two Raman shifts at 1350 cm^{-1} and 1580 cm^{-1} corresponding to the D and G bands of carbon nanotubes. The formation of Li_2O_2 on the surface of cathodes using GPE and cGPEs suggest that discharge capacities mainly result from the formation of Li_2O_2 .

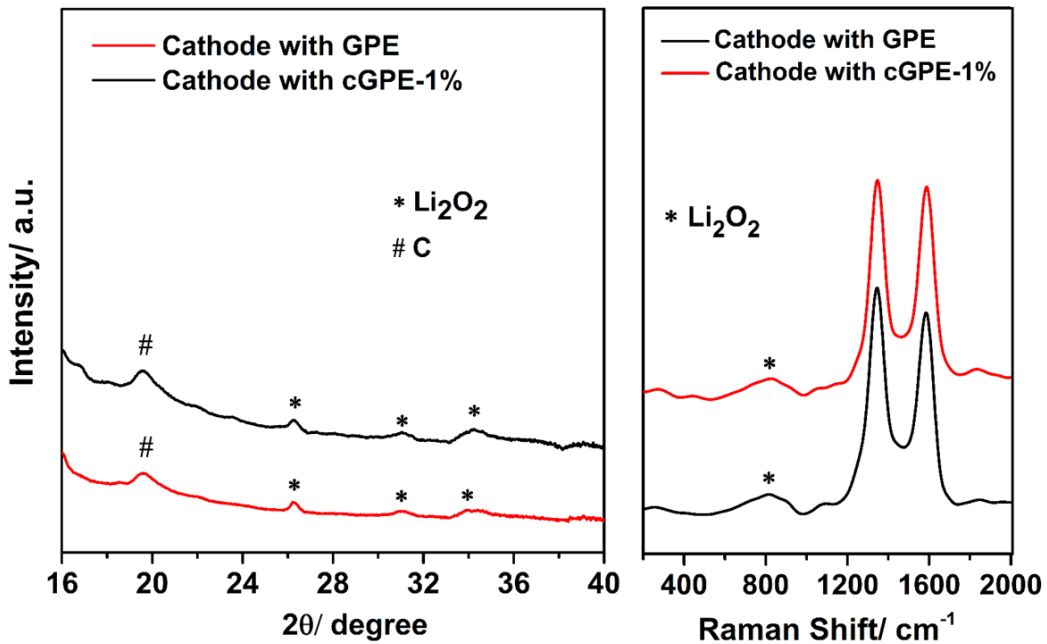


Figure 3-13: XRD patterns (left) and Raman spectra (right) of oxygen cathodes after a $500\text{ mAh}\cdot\text{g}^{-1}$ discharge of $\text{Li}-\text{O}_2$ batteries using GPE and cGPE-1%.

In order to measure the contribution of background capacity of CCGDL to total capacity of $\text{Li}-\text{O}_2$ batteries per cycle [121], the $\text{Li}-\text{O}_2$ cell was assembled using CNTs-free CCGDL with presoaked Celgard separator with $1\text{ mol}\cdot\text{kg}^{-1}$ LiTFSI salt electrolyte and charged/discharged in a voltage windows of 2.0-4.5 V at $125\text{ }\mu\text{A}$ (equivalent to the $250\text{ mAh}\cdot\text{g}_{\text{CNT}}^{-1}$ current density of $\text{Li}-\text{O}_2$ cells with CNT loading of 0.5 mg, which was the typical loading used in this work). Figure 3-14 shows the voltage profile of cell using CNT-

free GDL in full discharge mode along with cell using CNT-coated GDL using GPE-1 mol·kg⁻¹ LiTFSI with partial 500 mAh · g_{CNTs}⁻¹ at 125 μA current in a voltage window of 2.0-4.5V. The calculated capacity of cell using GDL up to 2.60 V was only 1 μAh. Moreover, the actual partial capacity of cell using CNT-coated GDL up to the 2.60 V was 250 μAh (500 mAh · g_{CNTs}⁻¹ * 0.5 mg of CNTs), so the capacity background contribution of GDL to the total capacity per cycle was only 0.4 %.

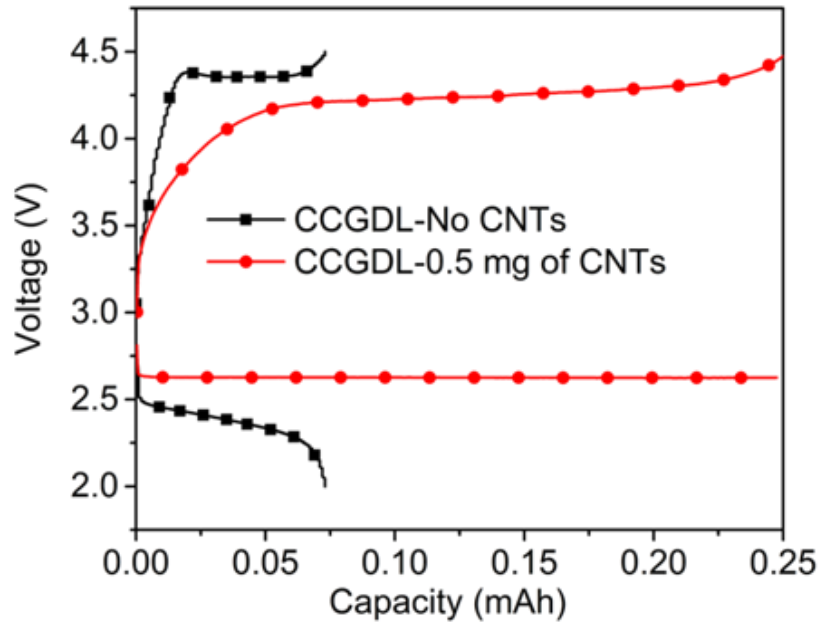


Figure 3-14: The voltage profile of Li-O₂ cell using CNT-free CCGDL in full discharge/charge mode along with cell using CCGDL-0.5 mg of CNTs in partial 500 mAh·g⁻¹ charge/discharge capacity mode at 125 μA current and voltage windows of 2.0-4.5 V

These results confirm that the CCGDL contribution to the total capacity of cells in partial charge/discharge of 500 mAh·g⁻¹ is very negligible. In situ EIS studies have been performed between 100 kHz to 100 mHz on Li-O₂ batteries using GPE and cGPEs after discharge during cycling and the Nyquist plots of cells using GPE and cGPE-1% are shown

in Figure 3-15 (a-b). The Nyquist plots consist of a semicircle corresponding to the total interfacial resistance between the electrolyte and the electrodes with a tail in the low-frequency region representing a semi-infinite Warburg resistance due to a diffusion controlled process of lithium ions and oxygen in the cathode [122,123]. Figure 3-15 (c) shows the change of interfacial resistance for all Li-O₂ batteries using GPE and cGPEs with different microfibers contents versus discharge cycle numbers up to their failure cycle (last discharge cycle with the capacity of at least 500 mAh.g⁻¹).

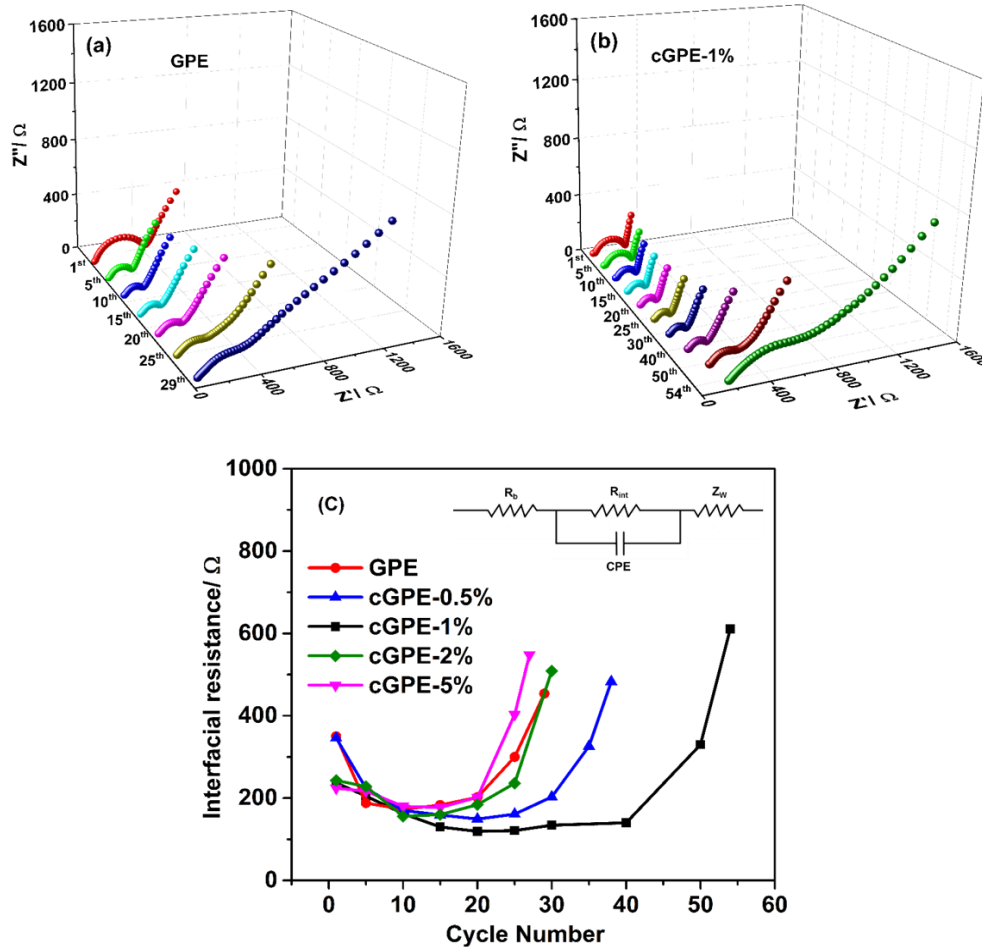


Figure 3-15: AC impedance Nyquist plots of Li-O₂ batteries using GPE (a) and cGPE-1% (b) during cycling after discharge and (c) Evolution of interfacial resistance of all Li-O₂ batteries using GPE and cGPEs against select cycle numbers.

As can be seen, adding the glass microfibers into GPE reduces the interfacial resistance of Li-O₂ cell even before cycling (Figure 3-16).

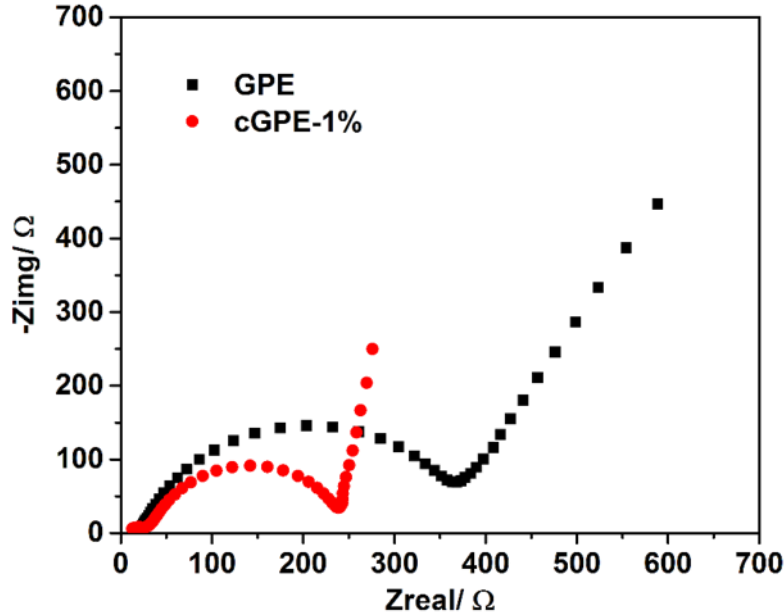


Figure 3-16: The EIS spectra of Li-O₂ cells using GPE and cGPE-1% before charging under Ar atmosphere

This could be due to the fact that the glass microfibers lead to hold more liquid electrolytes in cGPEs and in turn reduce the contact of liquid electrolyte and Li metals, which react with Li metal. Further, the interfacial resistance for cells started to decrease during the initial cycles and then steadily grew until their failure cycles. A decrease in interfacial resistance upon cycling in metal-O₂ and Li-ion batteries has been previously observed due to dendrite growth [124] or partial dissolution of the passivation film on the anode-electrolyte interface [125,126]. In later cycles, the observed increase in the interfacial resistance can be explained by the accumulation of the charge/discharge products/byproducts on the anode-electrolyte [30–32] and cathode-electrolyte interfaces

[123,127,128]. From Figure 3-15 (c), the interfacial resistance of Li-O₂ battery using cGPE-1% is more stable upon cycling comparing to cells using GPE and other cGPEs leading to 54 stable discharge cycles. This stabilization of interfacial resistance can be attributed to the Li⁺ transport enhancement in cGPE-1% which did not reduce the contact between the electrolyte and electrodes and maintained the active sites for specific resistance during cycling by providing sufficient Li⁺ ions at interfaces [125]. Excess of fillers in cGPEs (> 1%) could not stabilize the interfaces due to insufficient Li⁺ ion transport properties. In addition, the formation of poor electronic conductive Li₂O₂ and charge/discharge byproducts increases the cathode resistance and cathode-electrolyte interfacial resistance, resulting in high charge overpotentials [109,128]. Therefore, stabilizing the cathode-electrolyte interfacial resistance contributed to improving the charge capacity of cGPE-1% over GPE and cGPE containing cells, as can be observed in voltage profiles of Figure 3-11. Moreover, it is noteworthy that the poor rechargeability of all Li-O₂ batteries using GPE and cGPEs (Figure 3-10) was mainly due to the low OER activity of MWCNT cathodes as previously observed [88,129]. Various catalysts and redox mediators have been proposed to improve the OER and rechargeability of Li-O₂ batteries [130,131].

Lithium plating/stripping tests were also performed using symmetric Li/GPE/Li and Li/cGPE-1%/Li cells at the same current density used for cycling and plating and stripping cycle times of 2 hours each. From Figure 3-17, the voltage profile of cells using both GPE and cGPE-1% decreases during the initial cycles and then starts to grow. The decrease in overpotential at initial cycles was due to a heterogeneous Li plating/stripping process [132].

The continuous increase in voltage profile of cells is indicative of the growth of passivating layers produced by electrolyte and electrolyte decomposition products' reactivity with lithium [132].

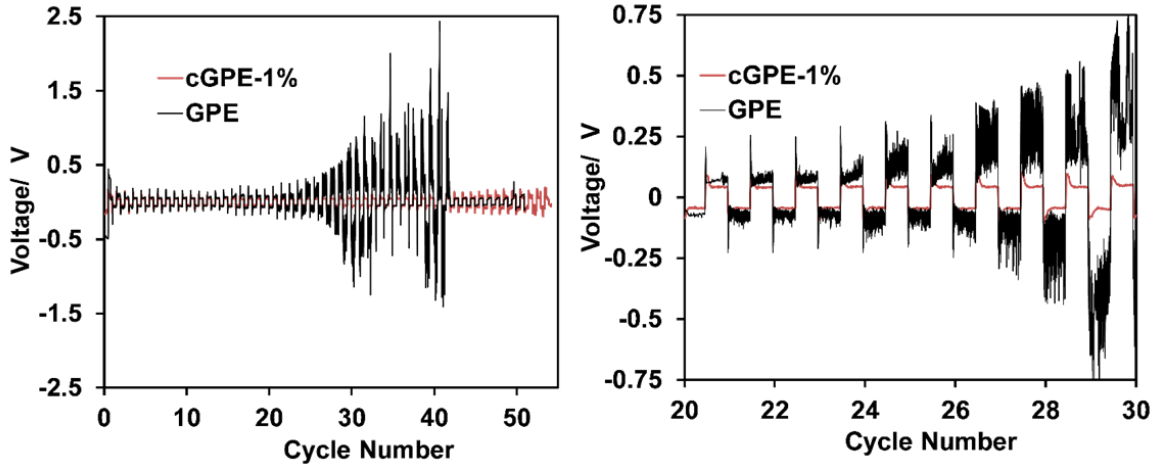


Figure 3-17: Voltage profile of the Li plating/stripping measurements performed using the symmetrical Li/cGPE-1%/Li cell and Li/GPE /Li cell at a current density of $0.125 \text{ mA}\cdot\text{cm}^{-2}$ and a plating and stripping times of 2 hours each.

A stable voltage profile is observed for cGPE-1% containing cell indicates a uniform lithium deposition/stripping and stable SEI formation with no detectable failure for more than 50 cycles. GPE containing cell, however, shows non-uniform lithium deposition/stripping and unstable SEI formation that lead to the internal short-circuiting of the cell at cycle number 40. This improvement can be attributed to the fact that cGPE-1%, which has the highest ionic conductivity and transference number improvements, stabilizes the Li interface by promoting uniform Li plating/stripping and stable interfacial layers [103,133,134]. These observations are in good agreement with our cycling and EIS results.

In addition, the inclusion of glass micro-fillers can improve the mechanical properties of GPE and block the dendrite growth [135].

3.4 Conclusions

In this chapter, the introduction of 1D glass microfillers to GPE has shown to improve lithium transport properties, namely, ionic conductivity and transference number, under optimized content. Li-O₂ battery using cGPE-1% shows 86% more cycles than battery using GPE. This improvement is attributed to the stabilization of electrolyte-electrode interfacial resistances resulting from increased Li⁺ transport properties.

4 COPOSITE GEL POLYMER ELECTROLYTE: EFFECT OF LI SALT CONCENTRATION

4.1 Background

According to recent reports, the electrolyte remains as the leading cause for rapid capacity fading and poor cyclability in Li-O₂ batteries [136–138]. Commonly carbonate-based solvents used in Li-ion batteries have been shown to be unstable during the ORR process [47]. Consequently, ether-based electrolytes have been suggested for Li-O₂ battery applications [19,20]. Nonetheless, the stability of ether-based electrolytes remains a concern and recent studies reported on the reactivity between superoxide radical species formed during charge/discharge with the ether-based solvents [48,139,140]. The hydrogen abstraction from methylene groups in ether-based solvents by superoxide radical species and subsequent reactions can cause solvent decomposition and the formation of lithium carbonates. The progressive formation of these insulating decomposition products on the surface of the cathode yields high cell polarization, thus causing poor cyclability [39,48,138]. Functionalized ether-based solvents (e.g. 2,3-dimethyl-2,3-dimethoxybutane (DMDMB) and tri(ethylene glycol)-substituted trimethylsilane (1NM3)) have been proposed to eliminate the possibility of methylene hydrogen abstraction [138,141]. Increasing the concentration of lithium salt in electrolytes was also shown to improve the performance of Li-O₂ batteries [142–144]. The appropriate Li⁺ solvation with solvent molecules may increase the favorable accessibility of superoxides to Li⁺ by increasing the salt concentration, which in turn mitigates the electrolyte decomposition [142–144]. Gel polymer electrolytes (GPE) consisting of liquid electrolyte and polymer has been successfully used in lithium-ion batteries due to their high ionic conductivity and

mechanical stability [91,145,146]. Recently, GPE with different electrolytes and polymers have also been developed for Li-O₂ battery applications [61,64,97,98,147]. It has been reported that GPE improves the performance of Li-O₂ batteries by reducing the electrolyte evaporation, and preventing the lithium corrosion caused by oxygen crossover from air cathode [61,63,64]. Inorganic filler materials have been widely incorporated into electrolytes and were shown to yield an increase in ionic conductivity and Li⁺ transference number [103,104,108]. Croce et al. [101] showed that the Lewis acid groups on the surface of ceramic fillers in composite polymer electrolytes strongly adsorb the anions of the lithium salt and enhance its salt dissociation. Bhattacharyya and Maier [73] also observed the same anion adsorption behavior of ceramic fillers in non-aqueous liquid electrolytes. Incorporation of various ceramic fillers in different electrolytes have also been explored for Li-O₂ battery applications [64,95,99,109,125,128,148]. It has been shown that the inorganic fillers can improve the performance of Li-O₂ batteries by stabilizing the interfacial resistance and preventing lithium anode corrosion. In this chapter, we investigate the incorporation of one-dimensional glass microfillers into glyme-based GPEs on reducing the formation of parasitic electrolyte decomposition byproducts in Li-O₂ batteries. Charge/discharge cycling, EIS, SEM, and Raman spectroscopy have been used to assess the performance improvement of composite gel electrolyte (cGPE) containing batteries.

4.2 Experimental Details

For this chapter, liquid electrolyte solutions were first prepared by dissolving different concentrations of LiTFSI salt (0.1, 1.0 and 3.0 mol·kg⁻¹) in TEGDME. The GPE and cGPE with glass microfibers were prepared following the procedure detailed in our previous chapter. The glass microfibers content of cGPEs was chosen to be 1 wt.%, and only the lithium salt concentration was changed. The ionic conductivity and Li transference number were also measured as described in chapter 3. Figure 4-1 shows typical chronoamperometric curves of symmetrical Li/Li cells using GPEs and cGPEs with 0.1 and 3 mol·kg⁻¹ LiTFSI salt concentration with their corresponding impedance spectra before and after polarization.

Like chapter 3, galvanostatic charge/discharge tests were performed using MTI battery tester at the constant current density of 250 mA·g⁻¹ within the voltage range of 2.0–4.5 V. However, in-situ EIS measurements in this chapter were also conducted using a Gamry Reference 600, and the frequency range was extended to cover the 100 kHz to 10 mHz using 10% of DC discharge current during cycling test at OCV condition. All charge/discharge and EIS studies were carried out at 25°C. The interaction of ETPTA polymer, glass microfibers, and LiTFSI salt was also investigated by Raman spectroscopy.

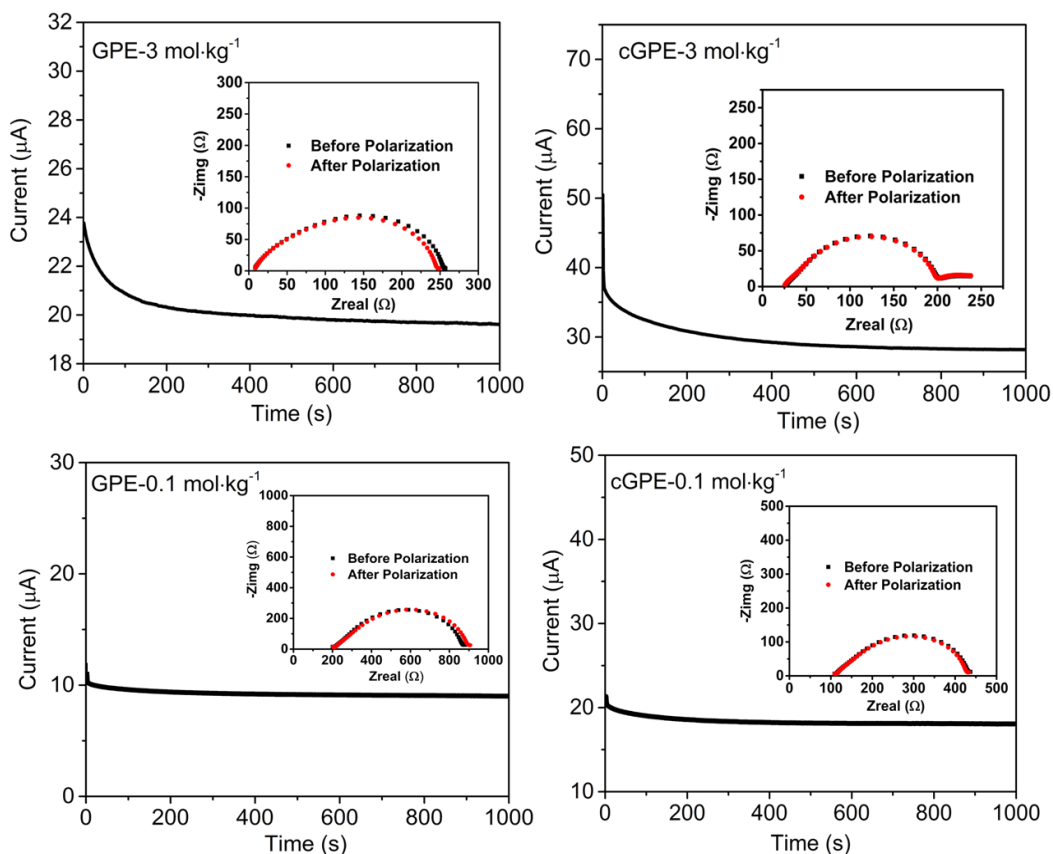


Figure 4-1: Chronoamperometric curves of Li/GPE and cGPE /Li cell after 10 mV of DC polarization at 25°C where the electrolyte films contain 0.1 and 3 mol·kg⁻¹ LiTFSI salt. Insets: electrochemical impedance spectra before and after polarization

All electrolyte samples including liquid electrolytes, GPEs, and cGPEs with various salt concentration were placed in airtight rectangular quartz cuvettes under argon inside the glovebox and then examined by Raman spectrometer (BaySpec's Nomadic™, 532 nm). Each spectrum was recorded with the exposure time of the 20s using the 10x objective lens at 25°C. Like chapter 3, the cathodes after different stage of charge/discharge were also characterized by the Raman spectroscopy, XRD, and SEM.

4.3 Results and Discussion

The ion transport properties of the GPE and cGPE prepared using various concentrations of lithium salt were evaluated using their corresponding methods described in the Materials and Methods. Table 4-1 summarizes the ionic conductivity (σ) and lithium transference number at room temperature (25°C).

Table 4-1: Summary of ionic conductivity and Li^+ transference number of GPE and cGPE at different salt concentrations

Electrolyte films	Transference Number t_{Li^+}	Ionic Conductivity σ ($\text{mS}\cdot\text{cm}^{-1}$)
GPE-0.1 mol·kg ⁻¹	0.40±0.02	0.13±0.02
GPE-1.0 mol·kg ⁻¹	0.50±0.03	1.02±0.05
GPE-3.0 mol·kg ⁻¹	0.54±0.02	0.68±0.02
cGPE-0.1 mol·kg ⁻¹	0.60±0.02	0.35±0.05
cGPE-1.0 mol·kg ⁻¹	0.64±0.02	1.40±0.02
cGPE-3.0 mol·kg ⁻¹	0.55±0.03	0.70±0.02

The ionic conductivity of GPE was shown to increase with increasing salt concentration from 0.1 mol·kg⁻¹ to 1.0 mol·kg⁻¹ due to the increase in the number of free ions. However, as the salt concentration increases to 3.0 mol·kg⁻¹, the ionic conductivity decreases as previously reported [149]. This decrease is mostly due to a drop in ion mobility caused by the higher viscosity of the liquid electrolyte soaking the polymer matrix [150,151]. The lithium transference number for GPE showed an increasing pattern with increasing salt concentration and followed the similar behavior of glyme-based liquid electrolytes and polyacrylate based GPE [150,151]. In Glyme-based electrolytes containing LiTFSI salt,

the increase in the salt concentration was shown to decrease the ionic association [152], and in turn, increases the Li^+ transference number.

The effect of the glass microfillers was demonstrated in increases in both σ and t_{Li^+} for the cGPE. This increase was owed to the immobilization of the TFSI^- , likely through interactions between Lewis acid groups on filler's surface and TFSI^- anions [108,110,116]. This ion-ceramic interaction yields an improvement in the ion pair dissociation and increases Li^+ dissociation [110]. At an optimized filler's content of 1 wt.% [148], a percolating network of glass microfibers is formed providing Li^+ transportation pathways. This increase was not apparent for the $3.0 \text{ mol}\cdot\text{kg}^{-1}$ cGPE, indicating insufficient surface groups for the immobilization of TFSI^- . This indicates that the effect of surface ion adsorption is reduced in environments with an abundance of free ions [153]. Therefore, with a fixed filler's loading, the effect of fillers was only apparent in cGPE with salt concentrations below a certain threshold, that once exceeded, the effect of fillers at that loading is reduced due to excessive amounts of ions in the solution.

Raman spectroscopy study was performed on liquid electrolytes, GPEs, and cGPEs to further investigate the interaction of ETPTA polymer and the microfillers with LiTFSI salt. Figure 4-2 shows that a Raman spectrum of TFSI^- anion which is assigned to the contraction and expansion mode of S-N coupled with CF_3 bending [154]. This Raman spectrum also represents the Li^+ and TFSI^- association [155]. The Raman shift between $736\text{-}742 \text{ cm}^{-1}$ is assigned to the unbounded TFSI^- (free anion or solvent-separated ion pairs (SSIPs)) [155,156] and Raman shift $\geq 744 \text{ cm}^{-1}$ show the bounded TFSI^- (contact ion pairs (CIPs) and ion aggregates) [155,156].

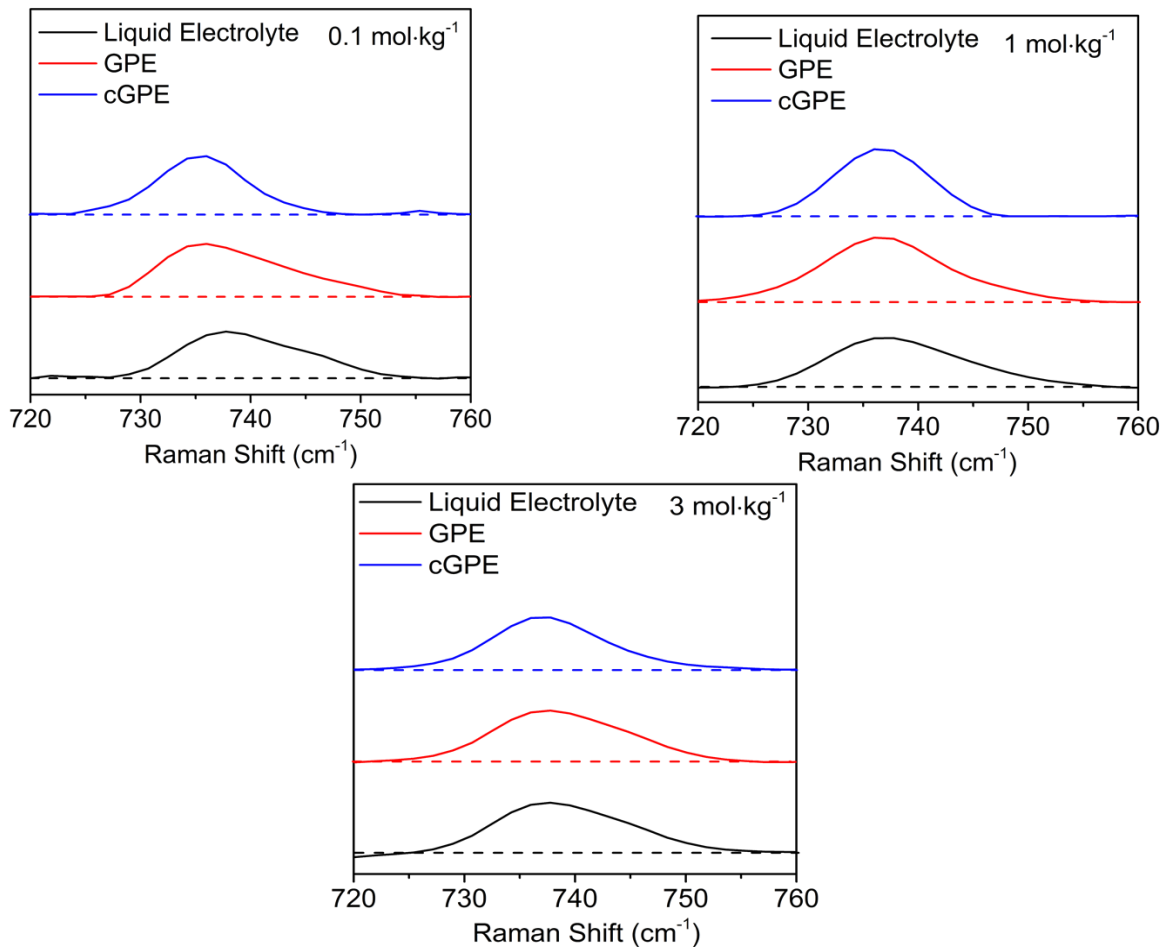


Figure 4-2: Raman spectra of Liquid electrolytes, GPEs and cGPEs with various LiTFSI salt concentrations showing Raman shifts of bounded and unbounded TFSI⁻ anions.

As shown in Figure 4-2, for all electrolyte samples, the addition of ETPTA polymer to liquid electrolytes with different salt concentrations does not change the shape of the TFSI⁻ Raman spectrum as they all exhibit two distinct Raman shifts for unbounded and bounded TFSI⁻ confirming that ETPTA polymer does not have any ion-trapping ability [145] and acts merely as an inert polymer. Furthermore, the addition of 1 wt.% glass microfillers to the GPEs at 0.1 and 1 mol·kg⁻¹ LiTFSI shows that the dissociation level of LiTFSI increases as the Raman shift of bounded TFSI⁻ (shoulder ≥ 744 cm⁻¹) was significantly reduced. However, cGPE at 3 mol·kg⁻¹ LiTFSI salt concentration indicates no significant changes

in Raman shift of bounded TFSI. These results are entirely in agreement with reported transference numbers. To further investigate the possible interaction of glass microfiber and liquid electrolyte with ETPTA polymer in GPEs, the glass transition of GPE was also obtained. The determination of glass transition (T_g) of the polymer matrix in GPEs with a small content of polymer (20 wt.%) is very challenging [157]. Hence to accurately determine the glass transition of ETPTA in GPE and cGPE-1%, the GPE and cGPE-1% with 50:50 wt.% ETPTA: liquid electrolyte ($1 \text{ mol}\cdot\text{kg}^{-1}$ salt concentration) was prepared. Figure 4-3 shows the DSC profile of GPE and cGPE-1%. As it is clear, the glass transition of ETPTA polymer inside GPE is very close to the pure ETPTA ($\sim 73^\circ\text{C}$) as obtained in the previous chapter. This similarity indicates that a little interaction between electrolyte and ETPTA in GPE. Furthermore, the addition of 1wt.% glass microfibers in GPE slightly reduce the T_g of ETPTA in cGPE-1%. As a result, reduction of polymer crystallinity is not a predominate reason for the ionic transport improvement of cGPE-1% over GPE [106].

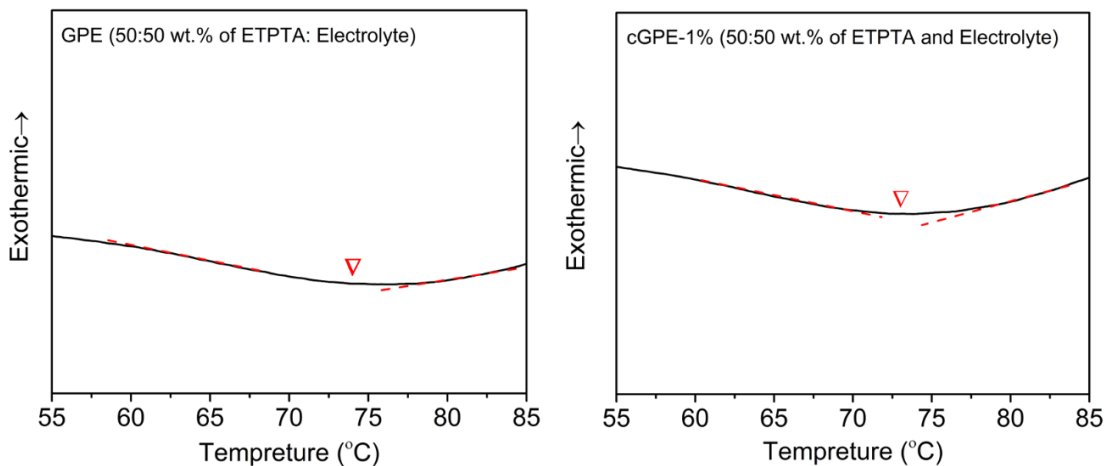


Figure 4-3: DSC profile indicating the glass transition (T_g) of (left) GPE containing 50:50 wt.% of ETPTA: Electrolyte with $1 \text{ mol}\cdot\text{kg}^{-1}$ salt concentration (right) cGPE-1% with the same ETPTA and polymer content

To examine the impact of glass microfillers on the performance of Li-O₂ batteries, charge/discharge cycling tests were performed at 250 mA·g⁻¹ for a cycle capacity of 500 mAh·g⁻¹ in a voltage window of 2.0-4.5 V. Figure 1 depicts the voltage profiles of GPE- and cGPE-containing Li-O₂ batteries using various salt concentrations up to the first cycle after their failure (first cycle with less than 500 mAh·g⁻¹). Cells using GPE-0.1 mol·kg⁻¹ ran for 12 discharge cycles compared to the GPE-1.0 mol·kg⁻¹ and GPE-3.0 mol·kg⁻¹ that ran for 29 and 40 cycles, respectively. This increase in cyclability with increased salt content has not been previously observed for GPE but was previously reported for liquid electrolytes [142,143]. Furthermore, cells using cGPE-0.1 mol·kg⁻¹ and cGPE-1.0 mol·kg⁻¹ showed 49 and 54 consecutive discharge cycles, respectively compared to cGPE-3.0 mol·kg⁻¹ with 43 discharge cycles. Using microfillers substantially improved the cyclability performance of Li-O₂ battery by 400% and 86% more cycles compared to their GPE counterparts for 0.1 and 1.0 mol·kg⁻¹, respectively. However, no significant improvement of cGPE over GPE was observed for 3.0 mol·kg⁻¹. This observation is consistent with the transference number improvements due to the inclusion of microfillers, proving the importance of Li⁺ transport properties of electrolytes on the Li-O₂ battery performance [89,148,158].

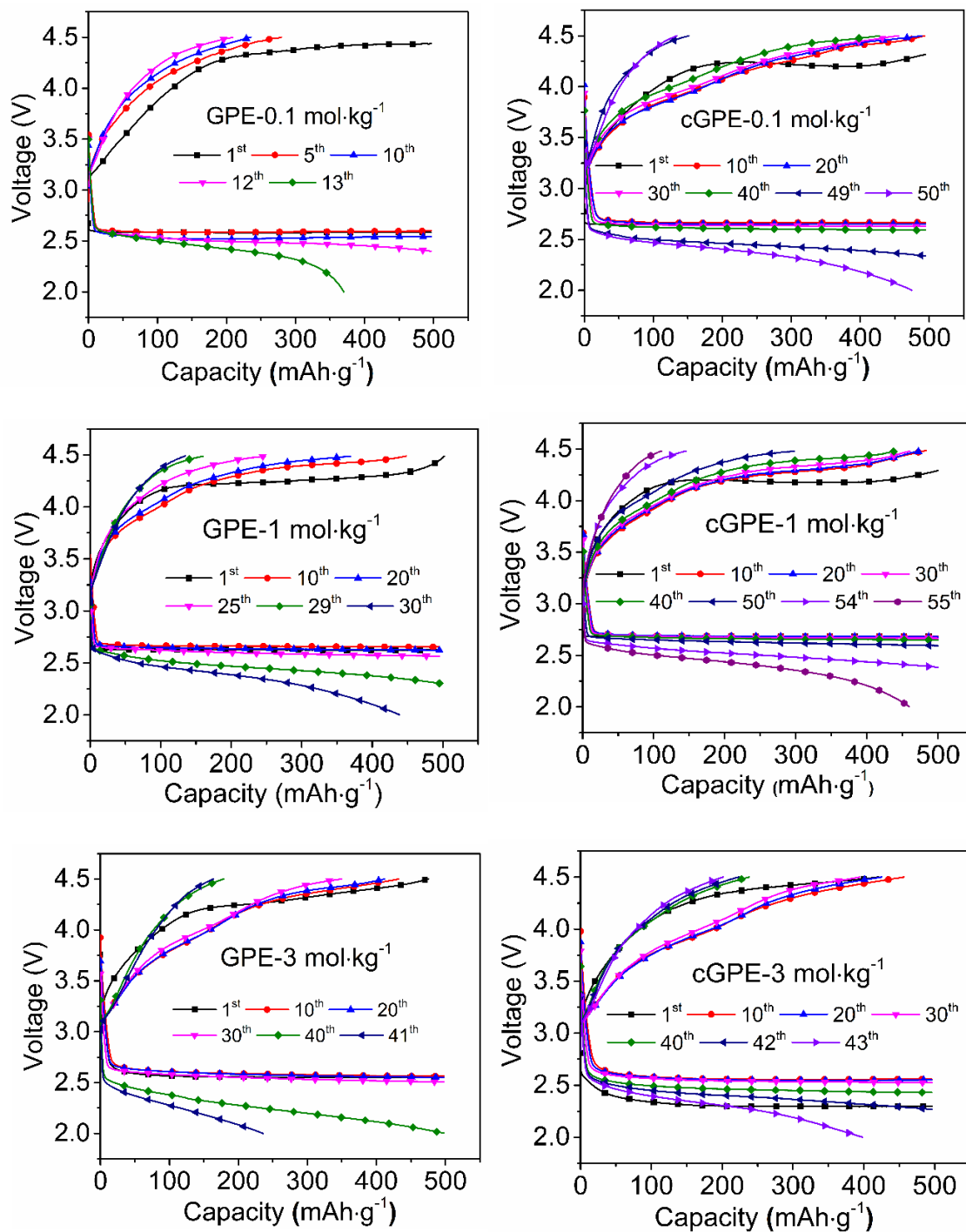


Figure 4-4: Voltage profile of Li-O₂ batteries using GPE and cGPE with different salt concentrations with a limited capacity of 500 mAh·g⁻¹ per cycle at current density of 250 mA·g⁻¹.

The formation of Li_2O_2 was confirmed by performing Raman spectroscopy on the surface of cathodes after one discharge cycle of $500 \text{ mAh}\cdot\text{g}^{-1}$. From Figure 4-5, all spectra show a Raman shift at $\sim 800 \text{ cm}^{-1}$ corresponding to the Li_2O_2 [120]. The presence of Li_2O_2 confirms that the discharge capacities of all Li- O_2 batteries are mainly due to ORR.

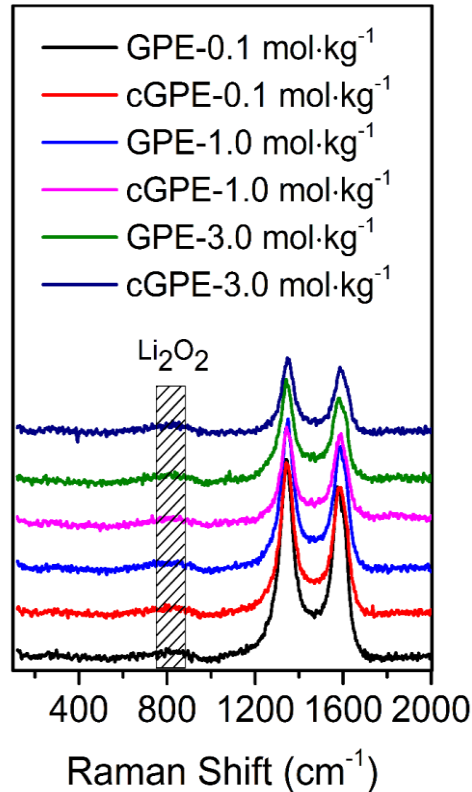


Figure 4-5: Raman spectra of $500 \text{ mAh}\cdot\text{g}^{-1}$ predischarged cathode using GPE and cGPE with different salt concentration. (Raman shifts at 1350 cm^{-1} and 1580 cm^{-1} correspond to the D and G bands of CNTs)

EIS has been recently used in metal- O_2 battery systems to conduct the in-situ determination of cell degradation mechanisms [159–162]. In this study, EIS was performed during cycling after discharge at OCV to evaluate the electrochemical behavior of GPE and cGPE.

Figure 4-6 (a) shows an example of a Nyquist plot of a Li-O₂ cell under oxygen using GPE-1.0 mol·kg⁻¹ after the discharge of the first cycle, 10th, 25th, and 29th (failure) cycles.

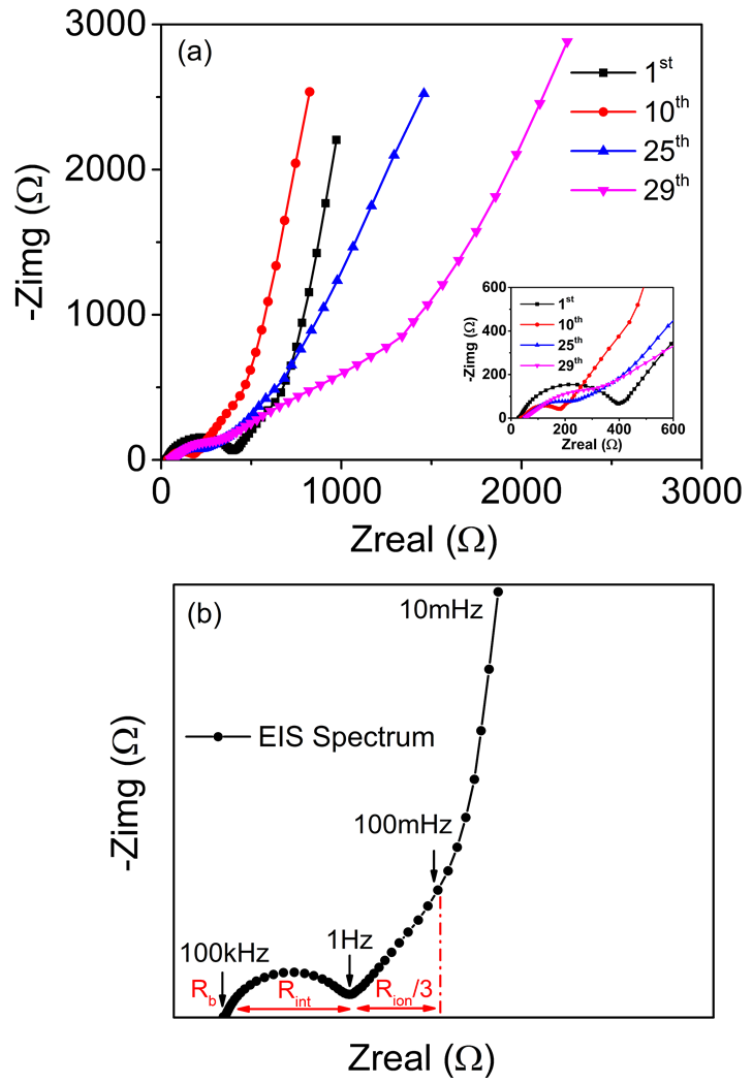


Figure 4-6: (a) Typical Nyquist plots of Li-O₂ batteries using GPE with 1.0 mol·kg⁻¹ salt concentration at OCV after first and failure cycle along with cycle 10th and 25th cycle (inset: shows close-up of Nyquist plots showing changes in interfacial resistances during cycling). (b) Transmission line model used to interpret the resistances of Nyquist plots.

The Nyquist plots consisted of a semicircle corresponding to the interfacial resistance (R_{int}) between the electrolyte (GPE or cGPE) and the electrodes, and a Warburg-like linear region followed by a rise, which is in agreement with other metal-O₂ EIS spectra at non-faradic conditions [159–161]. As described in chapter 2, transmission line model (TLM) has been used to determine the impedance behavior of the porous electrodes in Li-ion and metal-O₂ batteries [85,159,160]. In TLM, it is assumed that the Warburg-like linear region corresponds to the resistance of lithium-ion migration (R_{ion}) at the cathode [159,160]. Similar EIS behaviors were observed for GPE and cGPE at the other salt concentrations. The rise at low frequencies (less than 100 mHz) in the EIS spectra is consistent with the capacitive signal originated from the porous cathode [159,160]. However, it is worthy to mention that the EIS spectra deviate from the idealized porous cathode behavior as the rise at very low frequencies (<100 mHz) is tilted, and the slope of the Warburg-like line is not exactly 45°. It has been reported that this deviation is related to the frequency dispersion originating from a deficiency in the porous systems [163]. Bisquert [163,164] proposed a modified TLM circuit model to account this frequency dispersion by replacing the ideal capacitance in TLM model with constant phase element (CPE) to represent the capacitive behavior of porous cathode at non-faradic condition. Figure 4-7 (a) shows the modified TLM model used in this study. Based on this model, the ionic resistance (R_{ion}) could be estimated by the projection of Warburg-like line on the real axis of impedance as the limiting value of real impedance of porous cathode at low frequency ($\omega \rightarrow 0$) is equal to $R_{ion}/3$ [81,163–165]. The electrolyte resistance (R_b) and interfacial resistance (R_{int}) were also obtained from analyzing the real resistances of the semicircle of the EIS spectra at the high- and mid-frequency, respectively (Figure 4-7 (b)).

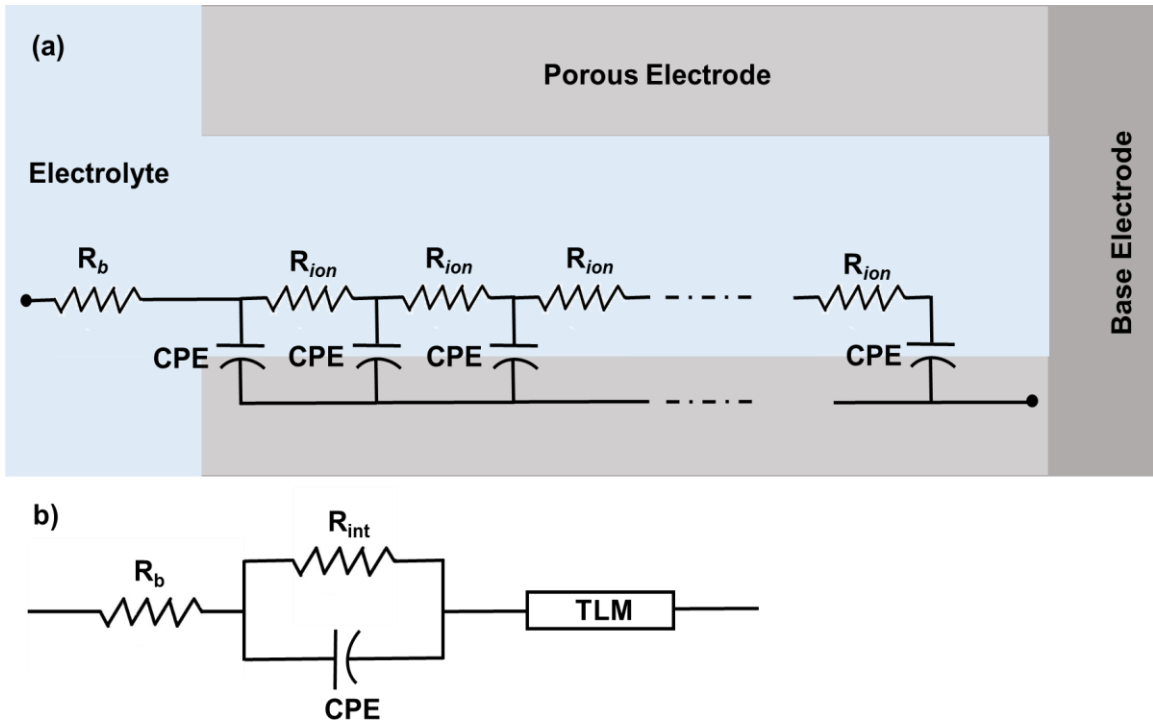


Figure 4-7: (a) Modified transmission line model (TLM) circuit model used to interpret the resistance of porous cathode. (b) The circuit model of the full cell Li-O₂ batteries.

As can be seen from Figure 4-6 (a), the Warburg-like line's slope steadily declines upon cycling. Furthermore, the diameter of the semicircle (R_{int}) is decreased in cycles 1 to 10, and then increased in cycles 10 and higher (Figure 4-6 (a) inset). Figure 4-8 shows the change of resistances R_{ion} , R_b and R_{int} versus discharge cycle numbers. R_{ion} was the dominant resistance in the cells which indicates that the increase in the resistance of cathode is the main cause of the failure. Shui et al. [32] and Knudsen et al. [160] also reported that the deactivation of the cathode was a major culprit in Li-O₂ battery failure. The increase in R_{ion} upon cycling indicates a physical pore is clogging within the porous cathodes caused by irreversible charge/discharge products deposition hindering Li⁺ transport inside the cathode [127].

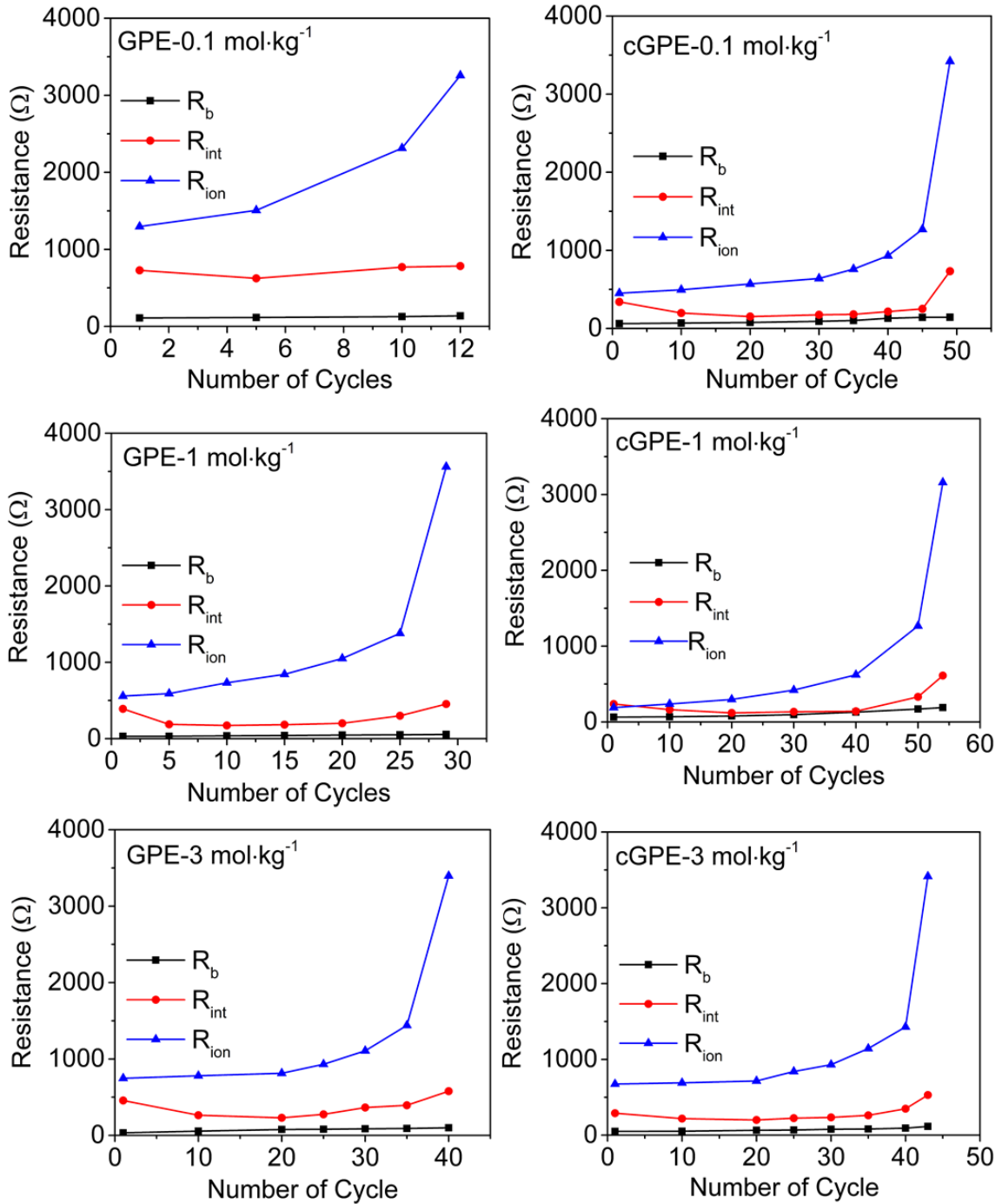


Figure 4-8: The change of resistances of Li-O₂ batteries using GPE and cGPE with different salt concentrations during cycling up to their failure

Tracking R_{ion} of cells using GPE- and cGPE-containing batteries at various salt concentrations (Figure 4-8) shows that for 0.1 and 1.0 mol·kg⁻¹ cGPE-containing batteries

had lower R_{ion} after the first discharge cycle and remained consistently lower than in their GPE-containing counterparts. In addition, the rate of growth of the R_{ion} also increased at a slower rate than for GPE-containing batteries, indicating that pore-clogging depositions mainly in the cathode were less problematic in batteries containing the cGPE. However, for the GPE and cGPE with $3.0 \text{ mol}\cdot\text{kg}^{-1}$ no significant reduction in R_{ion} growth was observed, which again suggests that the effect of the microfillers at this concentration was ineffective. This observation is consistent with the transference number and cycling comparisons previously discussed. To investigate the composition of the pore-clogging deposits, Raman spectroscopy was performed on the surface of cycled cathodes. Figure 4-9 shows the Raman spectra of cathodes using the GPE and cGPE at 0.1 and $1.0 \text{ mol}\cdot\text{kg}^{-1}$ recovered from Li-O₂ batteries after the 12th and the 29th discharge cycles (the failure cycles of GPE-containing batteries), respectively. Cathodes of GPE-containing batteries show pronounced Raman shifts at 1082 cm^{-1} , which correspond to the formation of lithium carbonates [120,166]. Conversely, the cathodes recovered from cGPE-containing batteries cycled under the same conditions and for the same number of cycles show less pronounced Raman shifts at 1082 cm^{-1} , indicating the lesser formation of lithium carbonates. These results confirm both the location (cathode) and nature (lithium carbonates) of the depositions leading to the pore-clogging and consequently the growth of R_{ion} . The presence of lithium peroxides and lithium carbonates were also confirmed by XRD (Figure 4-10).

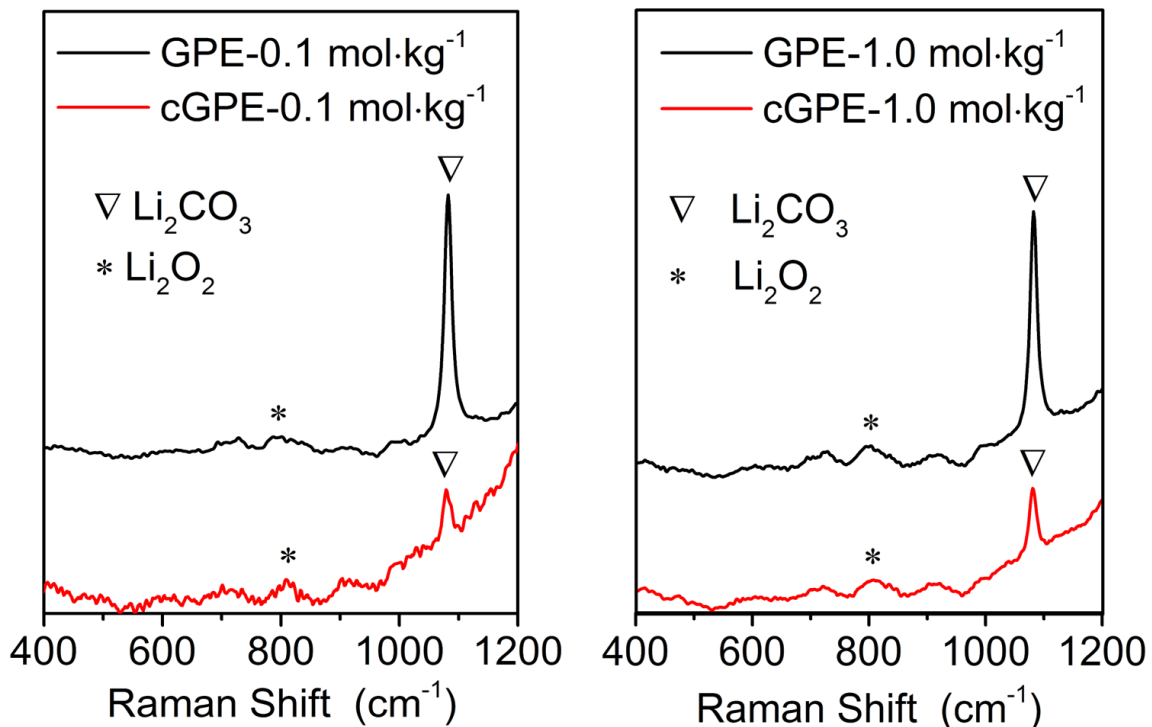


Figure 4-9: Raman spectra of cycled cathodes after discharge using GPEs and cGPEs with 0.1 and 1 mol·kg⁻¹ salt concentration at cycle number where the batteries using GPEs fails (cycle 12th for 0.1 mol·kg⁻¹ and cycle 29th for 1 mol·kg⁻¹).

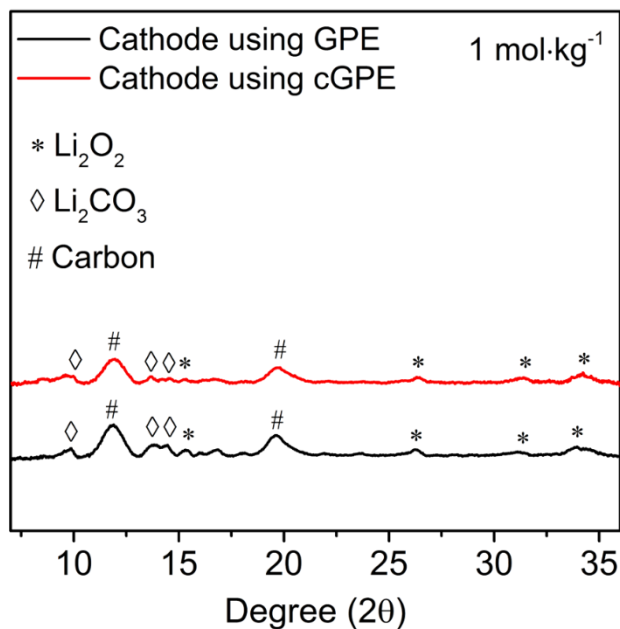


Figure 4-10: XRD patterns of cathodes using GPE and cGPE (1 mol·kg⁻¹ LiTFSI salt concentration) showing the presence of lithium peroxide and lithium carbonates.

Lithium carbonates can be formed by decomposition of carbon cathodes and electrolytes [22,48,140]. However, the formation of carbonate species during cycling of Li-O₂ batteries using tetraglyme-based electrolyte is suspected to be dominated by electrolyte decomposition [39]. This suggests that the addition of glass microfibers to the GPEs at 0.1 and 1.0 mol·kg⁻¹ reduces the electrolyte decomposition.

The cathodes of cells after the failure discharge cycle (the failure cycles of GPE-containing batteries) were recovered and visually inspected using SEM.

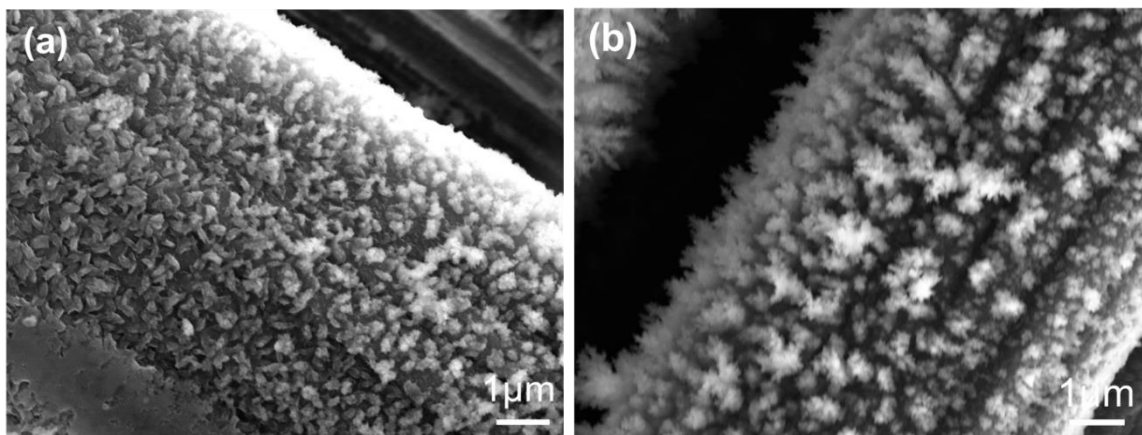


Figure 4-11: SEM micrograph of cycled cathodes in Li-O₂ batteries using cGPE (a) and GPE (b) with 1.0 mol·kg⁻¹ salt concentration after the failure discharge cycle.

In Figure 4-11, the cathode of batteries containing cGPE-1.0 mol·kg⁻¹ shows toroid-shaped discharge products (mostly Li₂O₂) covered by some fuzzy needle-like structures, previously reported as lithium carbonates [167]. In contrast, the cathode of batteries using GPE-1.0 mol·kg⁻¹ are mostly covered by the fuzzy needle-like structures and almost completely burying the toroid-shaped discharge products. This confirms the formation of the passivating lithium carbonates layer on the cathode. The continuous formation of these insulating lithium carbonates yielded large voltage hysteresis with an increase in cycling

(Figure 4-4). The impact of cycling on the interfacial resistance (R_{int}) in the Li-O₂ batteries is shown in Figure 4-12.

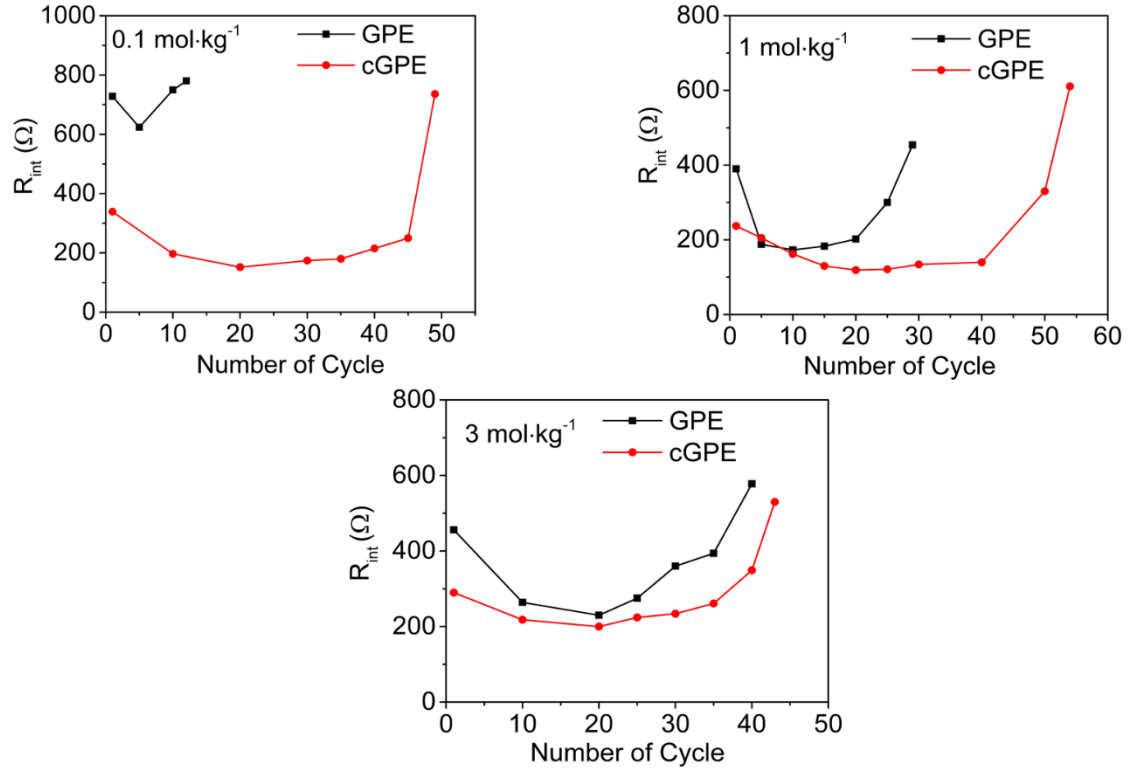


Figure 4-12: The change of interfacial resistance (R_{int}) of cells using GPE and cGPE with different salt concentrations during cycling up to their failures.

The R_{int} decreases in the initial cycles before it starts to grow in later cycles until the failure cycle. A similar phenomenon was also observed in our previous chapter [148]. As previously observed, the interfacial resistance of Li-O₂ batteries is initially governed by anode/electrolyte interface [127,160]. Hence the decrease in the R_{int} at initial cycles is mainly related to the dissolution of the passivation film on the anode/electrolyte films interface [125]. However, the increase in R_{int} in later cycles could be related to the accumulation of irreversible charge/discharge by-products on the anode/electrolyte [31]

and cathode/electrolyte interfaces [127]. To further distinguish the contribution of cathode and anode in our EIS spectra, two sets of symmetrical cells using Li/electrolyte film/Li, and cathode/electrolyte film/cathode were assembled. Potentiostatic EIS with AC amplitude of 5mV in the frequency range of 100k Hz to 100 mHz were performed on both Li/Li and cathode/cathode symmetrical cells. In the case of cathode/cathode symmetrical cells, both precycled and fresh cathodes were used. For precycled cathode/cathode symmetrical cells, both cathodes were initially cycled 50 times for limited charge/discharge capacity of 500 mAh·g⁻¹ at the same current density of 250 mA·g⁻¹ in separate cells before they were assembled in a new symmetrical cell. The cathodes after cycling were recovered from cells and rinsed with acetonitrile to remove the electrolyte salt before they were placed in symmetrical cells under oxygen environment. Figure 4-13 illustrates the EIS spectra of symmetrical cathode/cathode and Li/Li symmetrical cells along with EIS spectrum of the full Li-O₂ cell. As can be seen, Li/Li cell exhibits a semicircle corresponding to the interfacial resistance of the surface of Li and gel film electrolyte followed by charge transfer resistance. Fresh cathode/cathode symmetrical cell also shows a typical behavior of porous systems consisting of a Warburg diffusion resistance followed by a capacitive rise. Comparing the EIS spectrum of the full Li-O₂ cell with fresh cathode/cathode and Li/Li symmetrical cells confirms that the interfacial resistance of the full Li-O₂ battery is mostly controlled by the interfacial resistance of Li/electrolyte film. On the other hand, the precycled cathode/cathode symmetrical cell shows a semicircle followed by the Warburg-like resistance. This indicates that the irreversible charge/discharge products are accumulated on the cathode and contributes into the interfacial resistance upon cycling. Hence, at the beginning of cycling test, the interfacial resistance of full Li-O₂ cell is mostly

governed by the anode/electrolyte film interface. Then in later cycles, the cathode/electrolyte film interface was started to contribute to interfacial resistance.

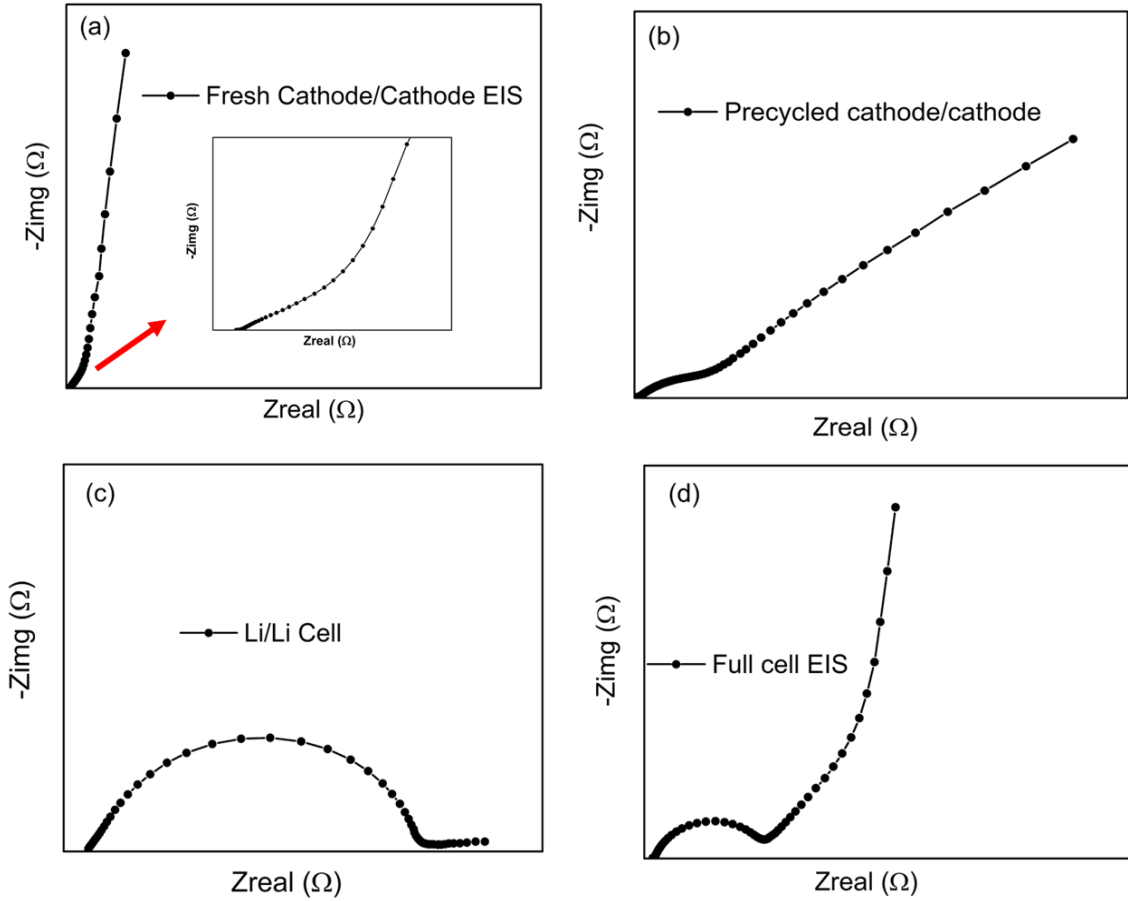


Figure 4-13: (a) EIS spectrum of (a) fresh cathode/cathode symmetrical cell (b) precycled cathode/cathode symmetrical cell (c) Li/Li symmetrical cell (d) full Li-O₂ battery

Yi et al. [109,128] also showed that the formation of lithium carbonates by-products on the cathode/electrolyte interface could increase the interfacial resistance in Li-O₂ batteries during cycling. The addition of microfillers (especially in 0.1 and 1.0 mol·kg⁻¹ salt concentration) maintained a lower R_{int} throughout the cycling, and can be credited to stabilizing the electrolyte and reducing its decomposition rate.

To further investigate the contribution of each cell component in the failure of Li-O₂ batteries, the Li-O₂ cells using GPE and cGPE-1% with 1 mol·kg⁻¹ LiTFSI salt concentration were assembled and cycled with the limited capacity of 500 mAh·g⁻¹ at a current density of 250 mA·g⁻¹. EIS was performed before cycling and after their failure cycles. The cells after failures were opened inside Ar-filled glovebox, and extra 20 μL electrolytes were added to soak the cathode. EIS was again performed on the cells. The cells with added electrolyte rested under oxygen for 5 hours and charged/discharged at the same condition. As can be seen from Figure 4-14, the ionic resistance (R_{ion}) of cells after failure increases as explained earlier. After soaking failed cathodes with extra electrolyte, the ionic resistance (R_{ion}) of cell reduced (Figure 4-14). The failed cells after soaking with extra electrolyte only ran for one cycle, and again they failed. EIS spectra of cells after their second failure was very close to their first failure EIS spectra. These results confirmed that the batteries were not failed due to the drying of cathodes. Secondly, although the ionic resistance (R_{ion}) of both cells reduced temporarily after addition of extra electrolyte, it did not prevent the failure of cells as they only ran for one cycle. The Li-O₂ cells using GPE and cGPE-1% were also cycled up to their failure and then their Li anodes were replaced with fresh one inside the glovebox. The cells with new Li anode and old cathode were not able to recover and they failed immediately. These results were confirmed that the Li anode is not responsible for Li-O₂ battery failure.

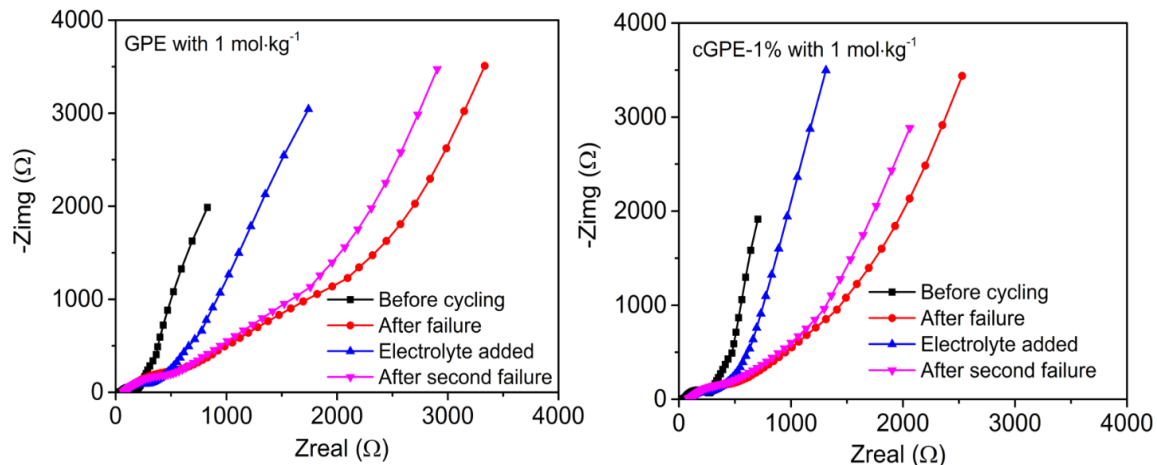


Figure 4-14: EIS spectra of Li-O₂ cells using GPE and cGPE-1% in different conditions

From the results obtained using the charge/discharge cycling, EIS, and Raman spectroscopy, the improvement in Li⁺ transport properties, especially lithium transference number correlated strongly with the improvement in cyclability of Li-O₂ batteries, mainly due to the stabilization of the electrolyte. At the cathode, the main source of instability of the electrolyte during discharge is the reaction of superoxide radicals with the glyme-based solvent molecules. Thotiyl et al. confirmed the prevalence of this reaction during discharge by tracking the ¹³C isotopic species in the decomposition products [39]. These superoxide radicals under ideal conditions would be reacting with the Li⁺ to yield Li₂O₂. However, in electrolytes with inefficient Li⁺ transport properties, the ratio of solvated Li⁺ to loosely bound ion pairs composed of solvated Li⁺ and TFSI⁻ is reduced [151,152], which in turn, increases the probability of the reaction of superoxide radical with the solvent [143,144] and promotes the electrolyte decomposition. Therefore, since an increase in transference properties of the electrolyte is associated with the higher ratio of solvated Li⁺, GPE and cGPE with improved transference numbers help reduce electrolyte degradation. The

degradation and eventual formation of lithium carbonates can, therefore, be reduced in Li-O₂ batteries using higher transference number electrolytes such as the cGPE presented in this work. Figure 4-15 demonstrates the possible mechanism of electrolyte stabilization in cGPEs.

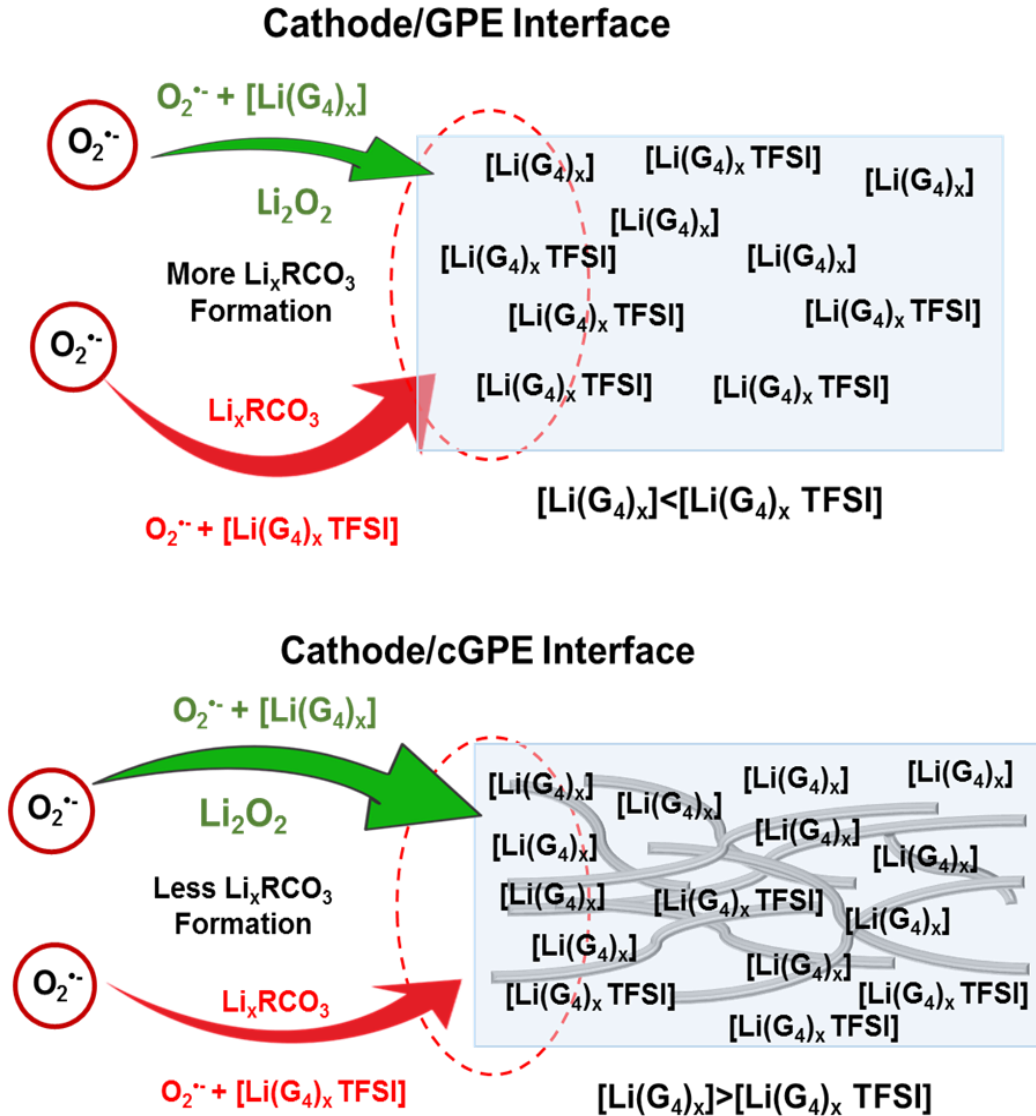


Figure 4-15: Schematic representation of proposed mechanism of electrolyte stabilization in Li-O₂ batteries in cGPEs over GPEs

4.4 Conclusions

In this chapter, gel polymer electrolytes (GPE) and glass microfillers-containing composite GPE (cGPE) were developed for their use in Li-O₂ batteries. Using various concentrations of lithium salt, it was observed that the resulting cGPE had improved lithium transport properties. Li-O₂ batteries containing cGPE cycled up to 5-fold more cycles of 500 mAh·g⁻¹ capacity per cycle compared to batteries containing GPE. This improvement was determined to be due to the reduction of the growth of ionic resistances in the cathode and its electrolyte interface. This resistance was traced back to the formation of lithium carbonates on the cathode due to the degradation of the tetraglyme-based solvent in the electrolyte. We theorize that the improved lithium transference of the cGPE increased the ratio of solvated to coupled Li⁺ which reduced the probability of superoxide radicals reacting with the tetraglyme-based solvent during cycling.

5 EFFECT OF ION PAIRS FORMATION ON THE STABILITY OF GLYME ELECTROLYTE

5.1 Background

Conventional carbonate-based solvents used in Li-ion batteries were shown to be severely decomposed by reactive oxygen species ($O_2^{\cdot-}$) during ORR process [47]. Therefore, new aprotic solvents such as Dimethyl sulfoxide (DMSO) [168], Dimethylformamide (DMF) [49] and Dimethylacetamide (DMA) [169] have been developed for Li-O₂ batteries. Recent studies, however, proved their reactivity toward Li anode and superoxide species [49,170,171,51]. Ether-based electrolytes which are relatively stable toward Li anodes are now the common solvents used in Li-O₂ batteries [172,48]. Their instabilities against the reactive oxygen species were also reported.[48,140] It is generally accepted that radical superoxides ($O_2^{\cdot-}$) attack the allylene groups close to the ethylene oxide bonds and decompose them to the carbonates groups. The subsequent reaction between these carbonates and discharge products (Li_2O_2 and LiO_2) forms lithium carbonates [140]. The continuous formation of parasitic lithium carbonates at the cathode during cycling cause high cell polarization and poor cyclability [39,48]. Recently, few theoretical and experimental studies suggested that the increase in the lithium salt concentration of the electrolytes improves the performance of Li-O₂ batteries [142,143]. The improvement has been accredited to the increase in the number of coordinated solvent molecules with Li^+ , which have been shown to be more stable than uncoordinated solvent molecules towards the superoxide attack [142,143]. Li et al. also showed that the type of Li^+ solvation with solvent molecules could also affect the reactivity of superoxides with solvent molecules [143]. Despite these findings at concentrated electrolytes, the mechanism of electrolyte

stabilization at relatively low lithium salt concentration is not well understood due to the complex interaction of Li^+ and solvent molecules. This complexity is originating from the fact that the addition of alkyl metal salts such as (Li^+X^-) to aprotic solvents forms different ionic complexes such as contact ion pairs (CIPs), solvent-separated ion pairs (SSIPs), and ion aggregates [155,173]. The presence of these ion pair complexes is dependent on the salt and solvent properties, and salt concentration [155,174,152,175]. In this chapter, we investigate the effect of Li cation/solvent solvation on the stability of tetraglyme-based electrolytes containing various Lithium bis(trifluoromethane) sulfonamide (LiTFSI) salt concentration for the Li- O_2 battery application. Tetraglyme-based solvent and LiTFSI salt were chosen in this study due to their popularity in the Li- O_2 battery systems[176]. Charge/discharge cycling, Raman spectroscopy along with various electrochemical characterization have been used to evaluate the origin of the improved performance at different salt concentrations.

5.2 Experimental Details

TEGDME solvent was first dried over 4Å Molecular sieve in the Ar-filled glovebox. Liquid electrolytes were prepared by dissolving different mole fraction of LiTFSI salt ($X_{\text{LiTFSI}} = 0.02, 0.2, 0.4$) in TEGDME solvent. It has to be mentioned that the lithium salt concentrations used in this chapter are very similar to the previous chapter. The Ionic conductivity of liquid electrolytes was determined by AC impedance spectroscopy. The electrolytes were trapped between two stainless steel blocking electrodes by using 1 mm thick Teflon O-rings as described in chapter 3. Lithium transference number (t_{Li^+}) was also determined using the Bruce-Vincent method. Li/electrolyte/Li symmetrical cells were prepared by soaking the Celgard 2400 polypropylene before cell assembly. Figure 5-1

shows chronoamperometric curves of Li/electrolyte/Li cells along with their impedance spectra. Raman spectroscopy was also performed to analyze the interaction of TEGMDE molecules with LiTFSI salt. All liquid samples with different salt concentration were filled in airtight rectangular quartz cuvettes in an Ar-filled glovebox and then examined by BaySpec's Nomadic™ Raman spectrometer (excitation wavelength of 532 nm). Each spectrum was recorded with the exposure time of 50 s using the 50X objective lens at 25°C. All cathodes after discharge tests were recovered from the cells in an Ar-filled glovebox and discharge products/byproducts were further characterized by Raman Spectroscopy (BaySpec's Nomadic™). Galvanostatic discharge tests were performed at the constant current density of $250 \text{ mA} \cdot \text{g}^{-1}$ with the limited capacity of $500 \text{ mAh} \cdot \text{g}^{-1}$ using MTI battery analyzer.

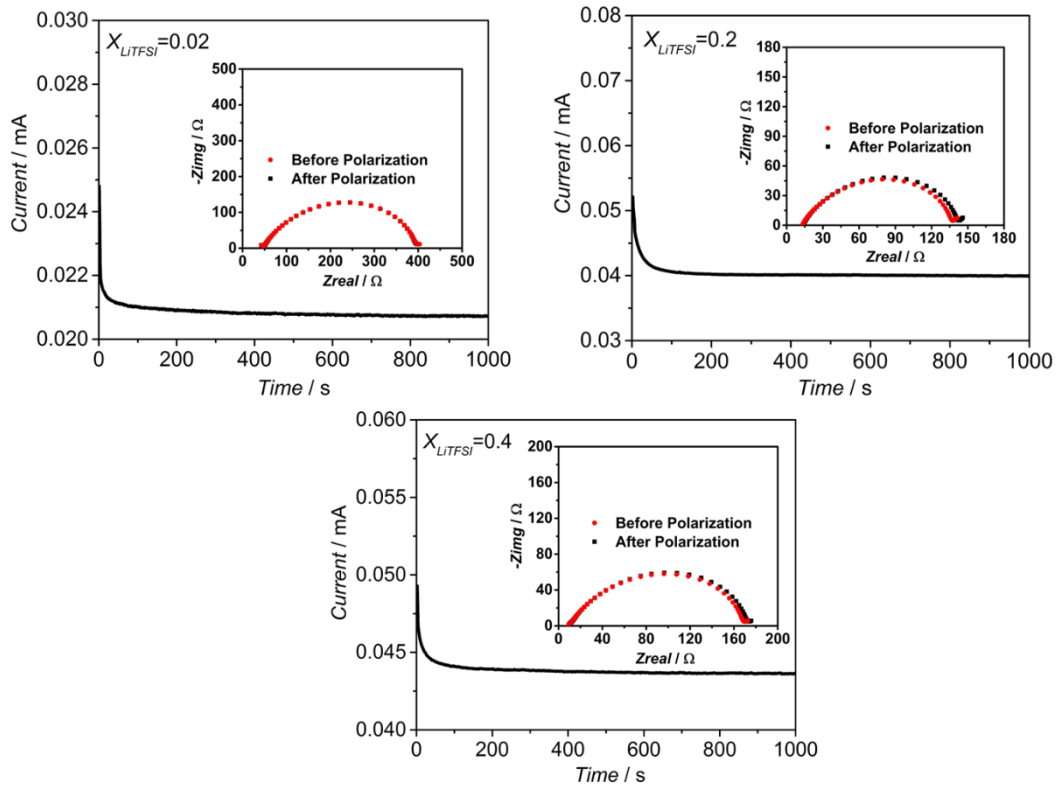


Figure 5-1: Chronoamperometric curves of Li/liquid electrolyte /Li cell after 10 mV of DC polarization at 22°C where the electrolytes contain different LiTFSI salt concentration. Insets: electrochemical impedance spectra before and after polarization

All cycling and electrochemical tests were carried out at 25°C. Galvanostatic electrochemical impedance spectroscopy (EIS) measurements were conducted in the frequency range of 100 kHz to 10 mHz using 10% of DC discharge current after discharges at open circuit voltage (OCV).

5.3 Results and Discussion

The ionic transport properties (ionic conductivity (σ) and Li transference number (t_{Li^+})) of electrolytes with different salt concentrations were measured at room temperature (25°C), and results are shown in Table 5-1. As it is cleared, the ionic conductivity of electrolytes increases with Li salt concentration from $X_{LiTFSI}=0.02$ to $X_{LiTFSI}=0.2$ and then decreases with further increase in Li salt concentration ($X_{LiTFSI}=0.4$). The increase in the ionic conductivity is associated with the increase in the number of charge carriers in the electrolyte [151]. However, the decrease in ionic conductivity is due to the increase in viscosity of the electrolyte which reduces the mobility of charge carriers [151]. The Li transference number of electrolytes, on the other hand, keeps increasing with Li salt concentration [151].

Table 5-1: The ionic conductivity (σ) and transference number (t_{Li^+}) of electrolytes with various salt concentration

Liquid Electrolyte	Transference Number, (t_{Li^+})	Ionic Conductivity, σ (mS·cm ⁻¹)
$X_{LiTFSI}=0.02$	0.39	0.31
$X_{LiTFSI}=0.2$	0.50	2.56
$X_{LiTFSI}=0.4$	0.56	1.13

Interestingly, the Li transference number of Liquid electrolytes are very close to those reported for GPEs indicating that the ETPTA polymer acts only as an inert polymer and EPTPA does not have any ion trapping ability. To investigate the effect of salt concentration of liquid electrolytes on the performance of Li-O₂ batteries, galvanostatic charge/discharge cycling was performed at the current density of 250 mA·g⁻¹ with the limited capacity of 500 mA·g⁻¹ in the voltage window of 2.0-4.5 V. Figure 5-2 shows the voltage profile of Li-O₂ cells using electrolyte with different salt concentrations up to their failures (cycle less than 500 mA·g⁻¹).

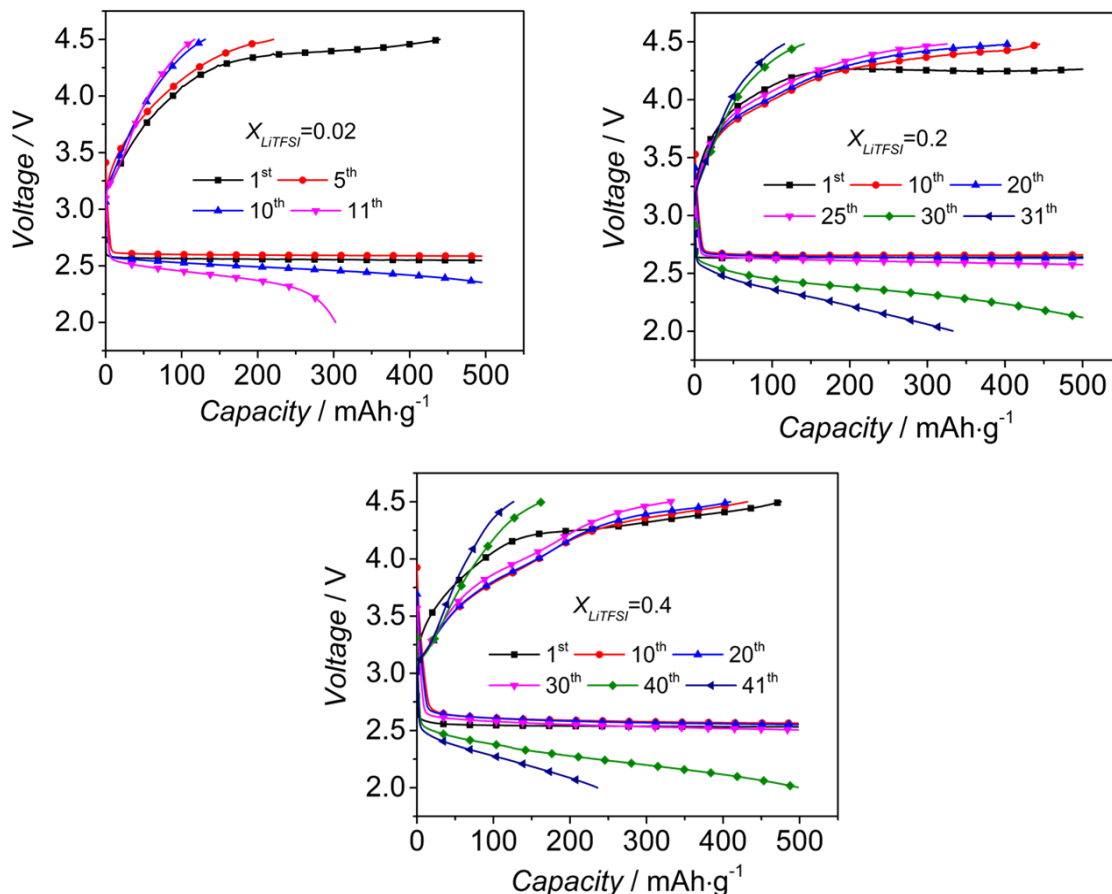


Figure 5-2: Voltage profile of Li-O₂ batteries using TEGDME electrolyte with various LiTFSI salt concentrations with a limited capacity of 500 mA·g⁻¹ per cycle at current density of 250 mA·g⁻¹ in a voltage range of 2.0-4.5 V.

As can be seen, Li-O₂ cells using an electrolyte with Li salt mole fraction of 0.02, 0.2 and 0.4 shows 10, 30 and 40 cycles, respectively. This shows that increase in salt concentration improves the cycleability of Li-O₂ batteries as previously was observed for ether-based electrolytes, as well [142,143]. Interestingly, the cyclability of cells using liquid electrolytes is similar to the cells using GPEs proving that the making the liquid electrolytes to the GPEs by adding ETPTA polymer does not improve the cell performance. Electrochemical impedance spectroscopy (EIS) was performed to investigate the failure mechanism of Li-O₂ batteries using electrolytes with different salt concentrations. *In-situ* EIS was carried out during cycling after discharge at open circuit voltage (OCV) under oxygen. Figure 5-3 shows the typical Nyquist plots of Li-O₂ batteries using an electrolyte with $X_{LiTFSI} = 0.2$ salt concentration after 1st, 5th, 25th and 30th (failure) discharge cycles.

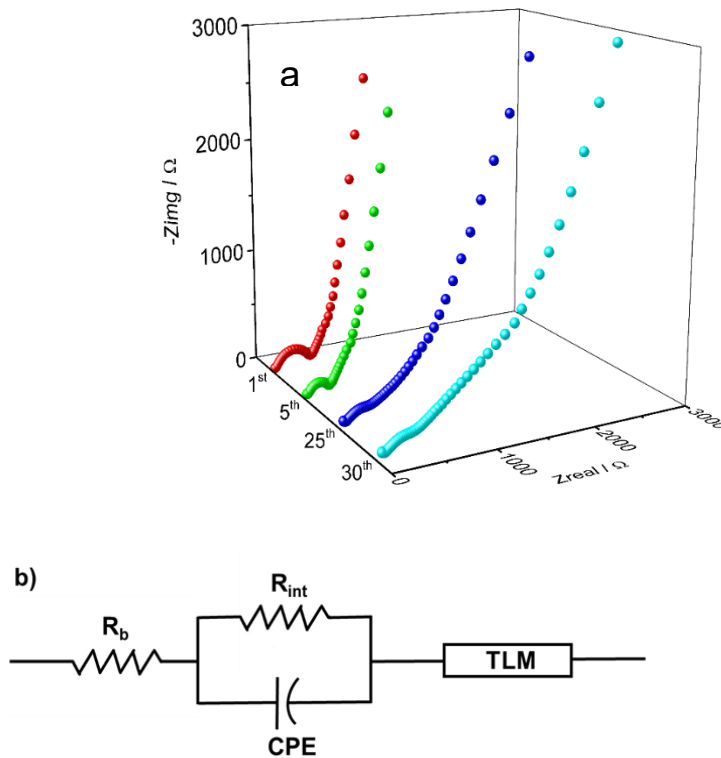


Figure 5-3: (a) Typical Nyquist plots of Li-O₂ batteries using TEGDME electrolyte with $X_{LiTFSI}=0.2$ salt concentration at OCV after first and failure cycle (30th) along with 5th and 25th cycle (b) Circuit model of the full cell as described in previous chapter

The Nyquist plots were similar to the Nyquist plots of cells using GPEs and cGPEs. They consisted of a semicircle corresponding to the interfacial resistance (R_{int}) between the electrolyte and the electrodes and a Warburg-like linear region followed by a rise at low frequencies. All impedance parameters (R_{ion} , R_b , and R_{int}) were obtained as described in the previous chapter.

Figure 5-4 shows the evolution of resistances R_{ion} , R_b and R_{int} of cells using different salt concentrations versus selected discharge cycle numbers up to their failure cycle. As can be seen, R_{ion} was the dominant resistance in all cells during cycling which indicates that the resistance within the cathode is the leading cause of the battery failure. Bardenhagen et al. [127] confirmed that the deposition of irreversible charge/discharge products on the cathode hinders the Li^+ transport within the pores of the porous cathode and in turn increases the R_{ion} . Tracking R_{ion} of cells shows that the R_{ion} increased at a slower rate with the increase in salt concentration indicating the pore clogging caused by irreversible charge/discharge products is less at high salt concentration.

Tracking the R_{int} upon cycling (Figure 5-5) also indicates that the R_{int} reduces at initial cycles and then starts to grow by cycling until the failure of cells as previously observed for GPEs and cGPEs. A decrease in interfacial resistance was reported for metal- O_2 cells using liquid electrolyte [159] quasi-solid state electrolyte [125] and gel polymer electrolyte as described in previous chapters [148,63]. Knudsen et al. [160] and Bardenhagen et al. [127] previously reported that the interfacial resistance (R_{int}) of Li- O_2 batteries is mostly controlled by anode/electrolyte interface at initial discharge process. Consequently, the decrease in the R_{int} at early cycles could be ascribed to the dissolution of the passivating film on the anode/electrolyte interface [125]. In the later cycles, however, the accumulation

of irreversible charge/discharge by-products on electrodes/electrolyte interface [127,31] causes the increase in R_{int} .

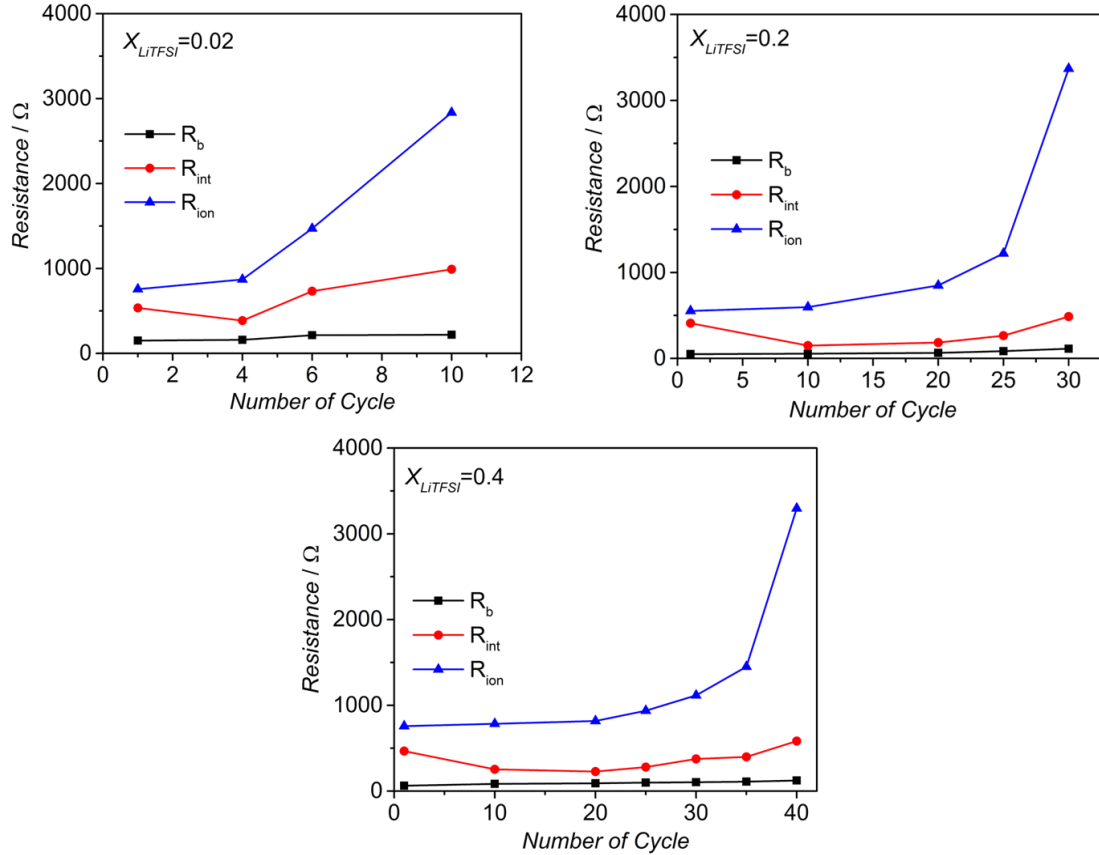


Figure 5-4: The change of resistances of Li-O₂ batteries using TEGDME electrolyte with different salt concentrations during cycling up to their failure (cycle 10th, 30th and 40th for $X_{LiTFSI}=0.02$ and 0.2 and 0.4, respectively).

Furthermore, the interfacial resistance (R_{int}) of Li-O₂ cell using $X_{LiTFSI}=0.2$ during cycling is less than that of Li-O₂ cells using $X_{LiTFSI}=0.02$ and $X_{LiTFSI}=0.4$ (Figure 5-5). Markus et al. also reported the same trend in interfacial resistance of Li-O₂ cells using different salt concentrations [177]. At low salt concentration ($X_{LiTFSI}=0.02$), the R_{int} is higher compared to other salt concentration due to diminished ionic conductivity [177]. The increase in

lithium salt concentration from $X_{LiTFSI}=0.2$ to $X_{LiTFSI}=0.4$ also reduces the R_{int} . The increase in Li salt concentration increases the viscosity of electrolyte, which causes the wettability between electrolyte and electrodes become poor [178].

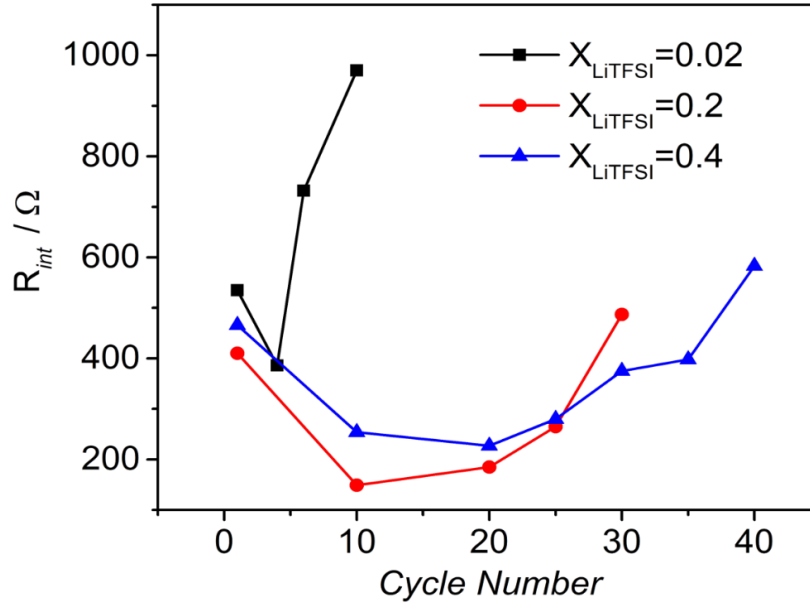


Figure 5-5: Interfacial resistance evolution of Li-O₂ cells using liquid electrolyte with various Li salt concentration versus selected cycles until their failure

Raman spectroscopy was also performed on the cycled cathodes after their failures to investigate the cause of the increase in R_{ion} and cell failure. Figure 5-6 illustrates the Raman spectra of failed cathodes harvested from Li-O₂ batteries with different salt concentrations. All spectra showed a Raman band at around 800 cm^{-1} confirming the formation of Li_2O_2 on the cathodes [120]. The presence of Li_2O_2 at all salt concentrations affirms that the discharge capacities of all batteries are due to oxygen reduction reaction (ORR). All cycled cathodes after their failures also showed a pronounced Raman band at 1082 cm^{-1} corresponding to the lithium carbonate species [120,166].

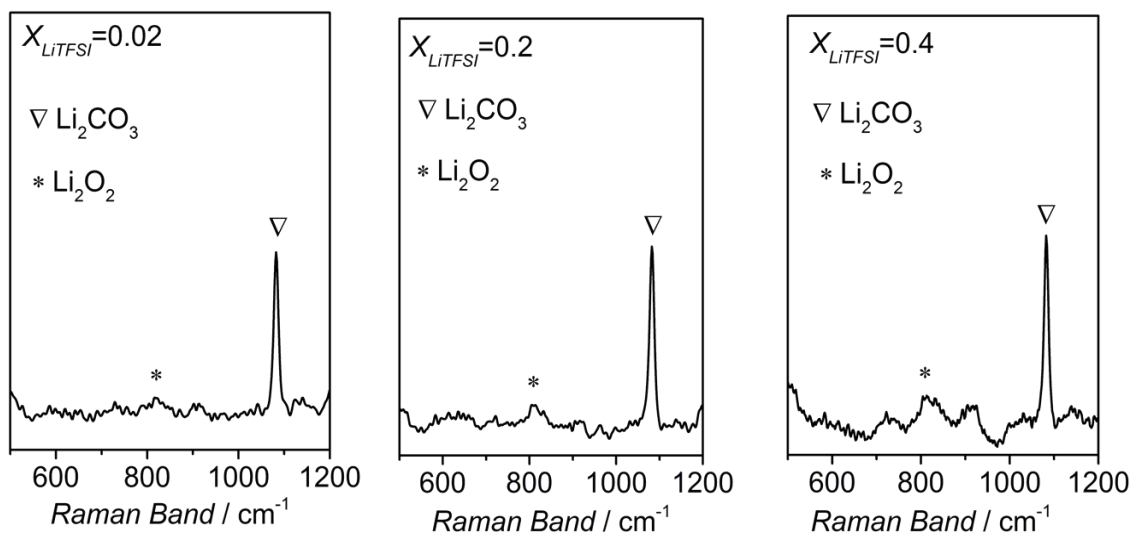


Figure 5-6: Raman spectra of cycled cathodes in Li-O₂ cells using TEGDME electrolyte with various salt concentration after their failure cycles (cycle 10th, cycle 30th and cycle 40th for cell using $X_{LiTFSI}=0.02$, 0.2 and 0.4, respectively).

As explained in previous chapter, the formation of carbonate species in Li-O₂ batteries has been associated with the decomposition of carbon cathodes and electrolytes [39,140,22]. However, Thotiyl et. al [39] and Xu et. al [140] proved that the majority of carbonate formation in Li-O₂ batteries using tetraglyme electrolyte is coming from electrolyte decomposition, not carbon cathode decomposition. Therefore, the results of cycling, EIS and Raman spectroscopy suggest that increase in Li salt concentration in the Tetraglyme electrolytes could improve the electrolyte stability in Li-O₂ batteries during cycling. It has been previously shown that reaction of radical superoxides ($O_2^{\cdot-}$ and LiO_2) with solvent molecules at the cathode could decompose solvent molecules and forms carbonate species [39,140]. To further investigate the effect of LiTFSI salt concentration on the stability of tetraglyme electrolyte, Raman spectroscopy was performed on the electrolyte solutions.

Figure 5-7 shows the Raman spectra of electrolyte solutions as a function of LiTFSI salt concentration. Deconvolution of Raman bands were performed by Gaussian-Lorentzian (Voigt) fitting function [179]. Pure TEGDME solvent exhibits the Raman bands at around 805, 820 and 850 cm^{-1} corresponding to the mixture mode for CH_2 rocking vibrations and C-O-C stretching vibration [175,180]. The addition of LiTFSI salt in the TEGMDE solvent introduced a new Raman band centered at around 740 cm^{-1} which could be assigned to the contraction and expansion mode of S-N coupled with CF_3 bending of TFSI^- anion [154]. Since the TFSI^- anion weakly interacts with TEGDME solvent molecules, the changes in the Raman band around 740 cm^{-1} are mostly correlated to the Li^+ cation and TFSI^- anion association [180]. Several types of solvates could be formed in TEGDME solution with various LiTFSI salt concentration due to the different Li^+ coordination with two conformers of the TFSI^- anion (cisoid (C_1) and transoid (C_2)) [155]. Hence it is difficult to conclusively assigned Raman bands to different solvates. However, it is generally accepted that the Raman bands between 736-742 cm^{-1} correspond to the unbounded TFSI^- (free and solvent separated ion pairs (SSIPs)) [155,156] and Raman bands $\geq 744 \text{ cm}^{-1}$ describe the bounded TFSI^- (contact ion pairs (CIPs) and ion aggregates (AGGs)) [155,156]. It is also known that upon addition of LiTFSI salt, a new Raman band at $\sim 865 \text{ cm}^{-1}$ known as a breathing mode were also appeared due to interaction between TEGDME and TFSI^- , which form cationic complexes ($[\text{Li}_x(\text{TGDME})_y]^+$) [175]. As can be seen from Figure 5-7, the increase in LiTFSI salt concentration increases the intensity of Raman band at $\sim 865 \text{ cm}^{-1}$ (cationic complexes) and reduces the intensity of Raman bands for free TEGDME molecules indicating the ratio of cationic complexes to free TEGDME increases.

Figure 5-7 also indicates that TFSI⁻ anions even at very low salt concentration ($X_{LiTFSI}=0.02$) have a strong ionic association tendency to Li⁺ as formation of CIPs and AGGs was confirmed in Raman spectrum (Raman band at ≥ 744 cm⁻¹). The similar results were also observed for TFSI⁻ in solvents with low Gutman donor numbers (DNs) [181,182]. The explanation for this high ionic association of TFSI⁻ could be due to the competitive solvation tendency of Li⁺ with both TEGMDE and TFSI⁻. The ratio of bounded to unbounded TFSI⁻ also increases with Li salt concentration confirming that the formation of anionic complexes (CIPs or AGGs, $([Li_n(TFSI^-)_m]^-)$) increases with Li salt concentration. Differentiating between CIPs (TFSI⁻ coordinated with only one cation) [183] and AGGs (TFSI⁻ coordinated with two or more cations) [183] using vibrational Raman spectroscopy is difficult [184]; however, Suo et. al [156] suggested that the Raman bands between 744-746 cm⁻¹ describes the formation of CIPs and Raman bands ≥ 747 cm⁻¹ corresponds to the AGGs. The formation of AGGs is unfavorable in electrolytes for battery applications as they significantly reduce the cationic transference number [183,185]. Furthermore, recent Raman spectroscopic investigation on the Mg(TFSI)₂ salt [183] showed that AGGs with bidentate ligands are formed at higher frequency (~ 752 cm⁻¹), yet the CIPs with monodentate ligands are formed at lower frequencies (~ 746 cm⁻¹). Although this distinct deconvolution of CIPs and AGGs with bidentate and monodentate ligands have not be reported for LiTFSI, Umebayashi et. al [186] confirmed the weak interaction of Li⁺ and TFSI⁻ at Raman band ~ 744 cm⁻¹ and strong solvation of TFSI⁻ and Li⁺ with bidentate ligands at Raman band ~ 750 cm⁻¹ [186]. The formation of AGGs with bidentate ligands significantly change the Li transference number as bidentate ligands provide strong interaction between Li⁺ and TFSI⁻ [187]. As can be seen from Figure 5-7,

the Raman band corresponding to the bounded TFSI⁻ appears at ~745 cm⁻¹ and increases with the Li salt concentration, which could suggest the formation of CIPs with monodentate ligands. Thus, at the studied salt concentrations, the increase in Li salt concentration will only increase the formation of CIPs with weak Li⁺ and TFSI⁻ interactions, which could liberate Li⁺ and participate in Li transference number. Besides, recent theoretical and experimental studies suggest that the cationic complexes are mostly contributed in the ionic transports in electrolytes rather than anionic complexes [188,189]. The Raman band corresponding to the cationic complexes at ~865 cm⁻¹ substantially increases with salt concentration, yet the Raman band for anionic complexes (~745 cm⁻¹) slightly increases. Hence, although both cationic and anionic complexes increase with Li salt concentration, the formation of cationic complexes are dominated in TEGDME solutions at the studied Li salt concentrations. The increase in Li⁺ transference number with Li salt concentration reported in Table 5-1 could be justified by the dominated formation of cationic complexes over anionic ones and the increase in the formation of CIPs (anionic complexes) with monodentate ligands.

Recent studies confirmed that uncoordinated glyme molecules are prone to the superoxide attacks [141], so based on the Raman spectroscopy, the formation of the cationic complexes could protect the TEGDME molecules from decomposition [142,190].

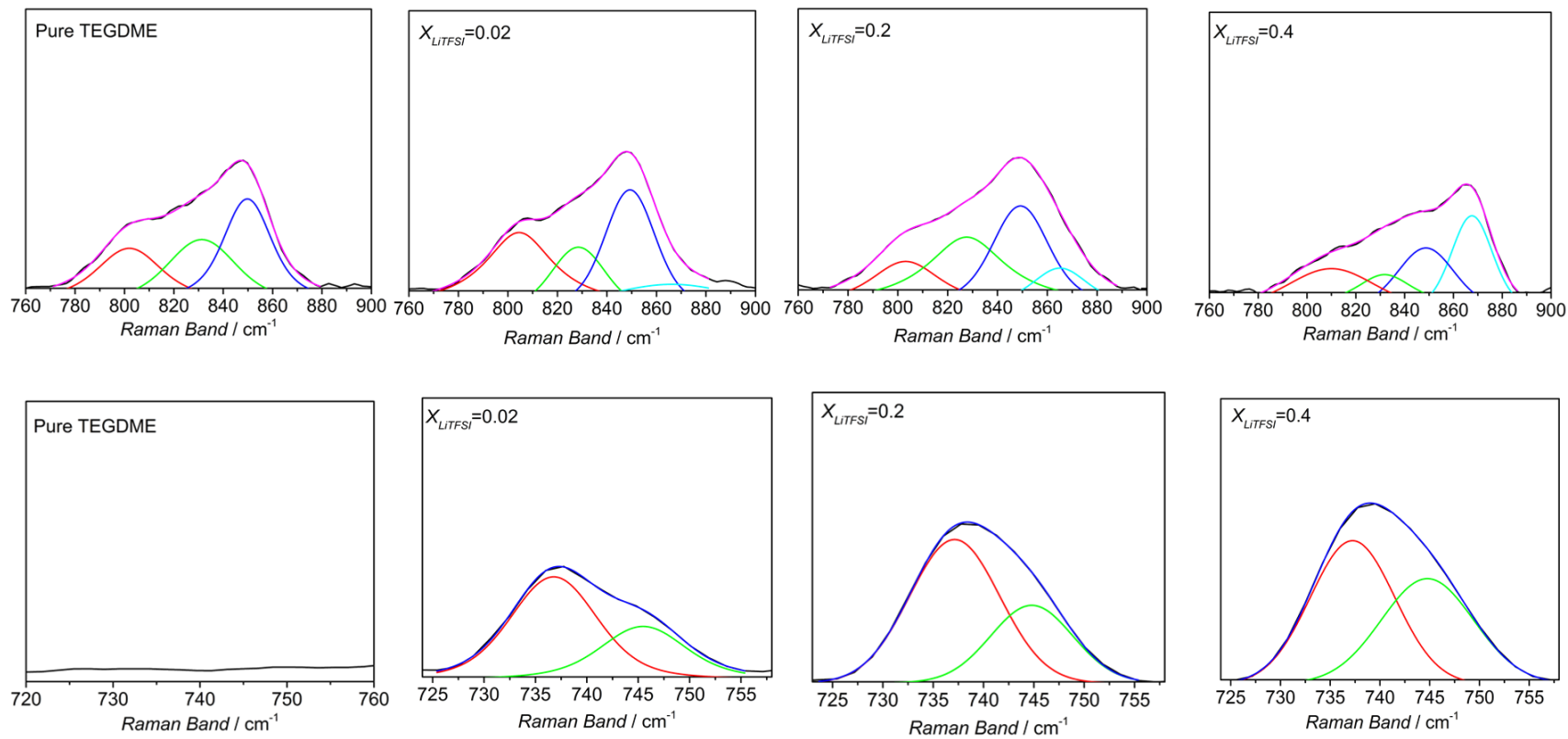


Figure 5-7: Raman spectra of electrolyte solution with various LiTFSI salt concentrations in the range of 800-900 cm^{-1} and 720-760 cm^{-1} at 22°C.

From the Raman spectroscopy results of electrolyte solutions, the protection mechanism of TEGDME with LiTFSI salt concentration could be explained by the fact that the ratio of solvated TEGDME with Li^+ to the unsolvated (free) TEGDME increases with salt concentration, which in turn protect the electrolyte from superoxide attack [142,143]. Furthermore, the Li salt concentration increases the Li transference number [151] in TEGDME solution and favors the reaction of superoxide with Li^+ instead of TEGDME molecules [143].

5.4 Conclusion

In this chapter, the influence of ionic complex formation in tetraglyme based electrolytes with various LiTFSI concentrations on the performance of Li-O₂ batteries have been investigated. Cycling results showed that the increase in LiTFSI concentration significantly improves the cyclability (as high as 300%). The *in-situ* electrochemical impedance spectroscopy (EIS) along with Raman spectroscopy revealed that the improvement was due to a slower growth rate of carbonate species originated from tetraglyme degradation within the porous cathodes. Raman spectroscopy analyses of the electrolytes at various LiTFSI concentrations suggest that the increase in the formation of cationic complexes likely lead to the stabilization of the tetraglyme molecules by protecting them from reactive superoxide attacks.

6 CONCLUDING REMARKS AND FUTURE WORKS

6.1 Concluding Remarks

Lithium-oxygen batteries have been considered as next-generation energy storage devices due to their high theoretical energy densities close to gasoline. However, many fundamental and technical challenges remain in this field. One of the most important challenges in Li-O₂ batteries is poor cyclability owing to the electrolyte decomposition. Recently, some attempts have been made to explore the performance of Li-O₂ batteries using polymer-based electrolytes (SPEs and GPEs) and their derivatives such as composite polymer electrolytes. Although some performance improvements have been reported in Li-O₂ batteries using polymer electrolytes and composite polymer electrolytes over liquid electrolytes, the exact source of improvement remains unknown and there are still a lot of room for improvement. The aim of this dissertation was to develop and characterize the hybrid polymer electrolytes based on UV-curable polymer, one-dimensional glass microfibers with diameter of approximately 1 μm and aspect ratio exceeding 100 and the conventional glyme-based solvent with various Li salt concentrations for lithium-oxygen battery application

In the first phase of this project, the content of one-dimensional glass microfibers in hybrid polymer electrolytes (composite gel polymer electrolyte, cGPEs) was optimized based on the performance of Li-O₂ batteries using GPE and cGPEs containing 1 mol·kg⁻¹ LiTFSI salt concentration with various glass microfibers content. It turned out that Li-O₂ batteries using cGPE with 1 wt.% glass microfibers showed the highest cyclability (54 cycles of 500 mAh·g⁻¹) compare to GPEs (29 cycles) and other cGPEs. The Li transport properties of GPEs and cGPEs have also revealed that at the optimum content of 1 wt.% glass

microfibers, the ionic conductivity and Li transference number improves as high as 37 % and 28%, respectively. Using in-situ electrochemical impedance spectroscopy (EIS), the battery performance improvement was traced to the stabilization of the interfacial resistances in the batteries owing to the improved Li⁺ transport properties.

In the second phase of this project, the GPEs and cGPEs containing 1 wt.% glass microfibers content with various salt concentrations ranging from 0.1, 1.0 and 3.0 mol·kg⁻¹ LiTFSI salt concentration were prepared to further investigate the exact source of improvement of glass microfibers in cGPEs, the sites (cathode or anode) at which the improvements were observed and the mechanism in which such improvement occurred. The cGPEs, with 1 wt.% of glass microfibers, demonstrated increased ionic conductivity and lithium transference number over GPE at various concentrations of lithium salt. Improvements as high as 50% and 28% in lithium transference number were observed for 0.1 and 1.0 mol·kg⁻¹ salt concentrations, respectively. However, at LiTFSI salt concentration of 3.0 mol·kg⁻¹, no significant improvement has been observed. Li-O₂ batteries containing cGPE similarly showed superior charge/discharge cycling for 500 mAh.g⁻¹ cycle capacity with as high as 86% and 400% increase in cycles for cGPE with 1.0 and 0.1 mol·kg⁻¹ over GPEs. Similar to the ionic properties of GPEs and cGPEs with 3.0 mol·kg⁻¹ LiTFSI salt concentration, a little improvement has been observed in cyclability performance of batteries using GPE and cGPE at this salt concentration. In situ EIS investigation using transmission line model (TLM) has been uncovered that the cGPE-containing batteries reduced the growth of ionic resistances (R_{ion}) in the cathode and its electrolyte interface. This resistance was traced back to the formation of lithium carbonates on the cathode due to the degradation of the tetraglyme-based solvent in the electrolyte.

We theorize that the improved lithium transference of the cGPE increased the ratio of solvated to coupled Li^+ which reduced the probability of superoxide radicals reacting with the tetraglyme-based solvent during cycling.

In the last phase of this project, the performance of Li-O₂ batteries using glyme-based (TGDME) liquid electrolytes with various salt concentration (0.1, 1 and 3 mol·kg⁻¹) was also investigated. It turned out that the increase in LiTFSI salt concentration from 0.1 to 3.0 mol·kg⁻¹ improves the cyclability of batteries as high as 200%. In-situ EIS and Raman spectroscopy investigation was shown that Li-O₂ battery mechanism failure was an increase in ionic resistance (R_{ion}) due to the accumulation of electrolyte decomposition products like lithium carbonate species on the cathode. Raman spectroscopy on the liquid electrolytes with various LiTFSI salt concentrations along with their ionic transport measurement revealed that the increase in LiTFSI salt concentration increases the ratio of solvated TEGDME with Li^+ . At the same time, increase in LiTFSI salt concentration improves the Li transference number, which in turn it could also increase the probability of superoxide reactions with Li^+ than TEGDME solvent molecules.

6.2 Future Works

The present dissertation has introduced the application of hybrid polymer electrolytes for Li-O₂ batteries. The results have shown that the introduction of one-dimensional ceramic glass microfillers in gel polymer electrolytes (GPEs) significantly improve the performance of Li-O₂ batteries by reducing the growth of electrolyte decomposition on the cathodes, which in turn causes the pore clogging of the porous cathode. Different spectroscopic investigations have revealed that the glass microfillers mitigate electrolyte decomposition by changing the solvation properties of Li salt and solvent. To the best of

our knowledge, this is the first report in the Li-O₂ battery field showing the possible mechanism of electrolyte stabilization by using the glass microfillers in hybrid polymer electrolytes. In the author viewpoint, this research study just opened a new horizon in the application of hybrid polymer electrolytes for electrolyte stabilization in Li-O₂ batteries. The following describes some of the unanswered questions, which deserve further investigation. The composition and surface functionality of borosilicate glass microfibers could affect the anion trapping ability of fillers, which in turn could significantly change the performance of Li-O₂ batteries. Further investigation on the composition of glass microfillers and tuning the content of boron element as one of the most efficient anion trapping element in glass deserve further investigation. Recently, it has been shown that the formation of singlet oxygen during charge/discharge cycling could attack the electrolyte and cause rapid electrolyte decomposition. Introducing different functional groups on the surface of glass microfillers could be one possible option to trapped these singlet oxygens to protect the electrolyte. To further investigate the proposed electrolyte stabilization mechanism, the isolation of glass microfillers in cGPEs on the anode/electrolyte interface or cathode/electrolyte interface will also reveal useful information.

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PUBLICATIONS

1. Amir Chamaani, Neha Chawla, Meer Safa and Bilal El-Zahab, “*One-Dimensional Glass Micro-Fillers in Gel Polymer Electrolytes for Li-O₂ Battery Applications*”, *Electrochimica Acta*, 235 (2017) 56–63
2. Neha Chawla*, Amir Chamaani*, Meer Safa and Bilal El-Zahab, “*Palladium-Filled Carbon Nanotubes Cathode for Improved Electrolyte Stability and Cyclability Performance of Li-O₂ Batteries*”, *Journal of Electrochemical Society*, 164 (2017) A6303-A6307, *Co-first Author
3. Meer Safa, Amir Chamaani, Neha Chawla and Bilal El-Zahab, “*Polymeric Ionic Liquid Gel Electrolyte for Room Temperature Lithium Battery Applications*”, *Electrochimica Acta*, 213 (2016) 587–593
4. Amir Chamaani, Meer Safa, Neha Chawla and Bilal El-Zahab, “*Composite Gel Polymer Electrolytes for Improved Cyclability in Li-Oxygen Batteries* (ACS Appl. Mater. Interfaces 9 (2017) 33819–33826
5. Amir Chamaani, Neha Chawla, Meer Safa and Bilal El-Zahaba, “*Stabilizing Effect of Ion Complex Formation in Lithium–Oxygen Battery Electrolytes*” (Journal of Electroanalytical Chemistry, under review)
6. Meer Safa, Yong Hao, Amir Chamaani, Ebenezer Adelowo, Neha Chawla, Chunlei Wang and Bilal El-Zahab, “*Capacity Fading Mechanism in Lithium-Sulfur Battery using Poly(Ionic Liquid) Gel Electrolyte*” (*Electrochimica Acta*, 10.1016/j.electacta.2017.11.185)

7. Amir Chamaani, Neha Chawla, Meer Safa, Bilal El-Zahab, “*The Effect of Li-Ion Dissociation Level in Gel Polymer Electrolytes on Electrochemical Performance of Li-O₂ Batteries*”, 231th ECS Meeting, New Orleans, LA, May 2017
8. Neha Chawla, Amir Chamaani, Meer Safa, Bilal El-Zahab, “*Investigation of Electrochemical Performance of Li-O₂ Batteries Using Catalyst Containing Carbon Nanotube Cathodes*”, 231th ECS Meeting, New Orleans, LA, May 2017
9. Amir Chamaani, Meer Safa, Neha Chawla, and Bilal El-Zahab, “*Hybrid Gel Polymer Electrolyte for Li-O₂ Batteries*” MRS Fall 2016 Meeting, Boston, MA, Nov 2016
10. Amir Chamaani, Meer Safa, Neha Chawla, and Bilal El-Zahab, “*Electrochemical Investigation of the Effect of Inorganic Fillers in Gel Polymer Electrolytes for Li-O₂ Batteries*”, 18th International Meeting on Lithium Batteries, Chicago, IL, Jun 2016
11. Neha Chawla, Amir Chamaani, Meer Safa and Bilal El-Zahab, “*Nanocatalysts for improved capacity retention in lithium-air batteries,*” MRS Spring Meeting 2016, March 2016
12. Neha Chawla, Amir Chamaani and Bilal El-Zahab, “*Multi-Walled Carbon Nanotubes as Cathodes for Lithium-Air Batteries: Effect of Catalysts and Additives*”, 228th ECS Meeting 2015, Phoenix, AZ, Oct 2015
13. Meer Safa, Amir Chamaani, and Bilal El-Zahab, “*Polymeric Ionic Liquid Electrolytes: Effect of Inorganic Fillers,*” 227th ECS Meeting 2015, Chicago, IL, May 2015
14. Amir Chamaani, Meer Safa, Neha Chawla and Bilal El-Zahab, “*Electrochemical Behavior of Flexible Gel Polymer Electrolyte with Fiber Fillers*”, 2nd Florida Statewide Graduate Student Research Symposium, Orlando, Florida, May 2015
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