Effects of the Electrode Wettability on the Deep Discharge Capacity of Li–O2 Batteries

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Supporting Information

ABSTRACT: The wettability of customized Li–O2 battery electrodes is altered by mixing acetylene black carbon particles with various binders. The wettability of the electrode can be characterized by the static contact angles between the electrode surface and nonaqueous electrolyte, which is 1 M bis(trifluoromethane)sulfonimide lithium salt (LiTFSI) dissolved in tetraethyl glycol dimethyl ether, and the double-layer capacitance measured by the cyclic voltammetry. Results show that electrodes containing poly(vinylidene difluoride) (PVDF) binder are lyophilic and increasing the fraction of poly(tetrafluoroethylene) (PTFE) increases the lyophobicity of electrodes. Li–O2 batteries are discharged at 0.1 mA/cm² with the cut-off voltage of 2.0 V. The discharge capacity of the electrode with 15% PVDF (36.5°) carbon coatings is 1665.8 mAh/g, whereas the customized electrode with 15% PTFE (128.4°) carbon coatings obtains the discharge capacity of 4160.8 mAh/g. However, the discharge capacity decreases to 3109.5 and 2822.9 mAh/g as the PTFE content further increases to 25% (135.5°) and 35% (138.5°), respectively. The electrode composed of two lyophobic carbon coatings on top and bottom and one lyophilic carbon coating in the middle has the static contact angle of 118.8° and acquires the highest specific discharge capacity of 5149.5 mAh/g.

1. INTRODUCTION

Greenhouse gas emissions from the production and consumption of fossil fuels have resulted in global climate change, acid rain, as well as other environmental consequences, which attract growing concerns from the world. Thus, renewable energy is emerging and thriving globally to replace the fossil fuels. However, the lower energy density and intermittency of renewable energy, such as solar energy and wind energy, limit their commercialization.1 Currently, the rechargeable battery is considered as one of the compromising technologies to efficiently store renewable energy. Among rechargeable batteries, the Li-ion battery has achieved a great success and has been widely applied in many devices and instruments, including mobile phones and electrical vehicles.2 The practical specific energy of the Li-ion battery (~200 Wh/kg), however, has already approached the theoretical specific energy (~600 Wh/kg).3

The Li–air battery using O₂ from the atmosphere displays a much higher theoretical energy density (~11 000 Wh/kg based on Li anode) than other rechargeable batteries.3 However, many obstacles, such as the low discharge–charge current rate, low charge discharge energy efficiency, and poor cycle life need to be overcome to commercialize the Li–air battery.4,5 Since CO₂ and H₂O in air would react with active components in batteries and deteriorate the performance, most laboratory experiments were conducted under pure O₂ environment.5,6 Experiments in this study have utilized pure O₂ so that the term Li–O₂ battery is presented throughout this paper. Generally, a rechargeable Li–O₂ battery using organic electrolyte consists of a lithium metal anode, a separator saturated with the organic electrolyte, and a porous cathode electrode (typically made from carbon or catalysts). The electrochemical reactions occurring on the anode and cathode as well as the overall reaction are shown below. The reversible cell voltage is 2.96 V.9

\[
\text{anode: } \text{Li} \xrightarrow{\text{discharge}} \text{Li}^+ + e^- \\
\text{cathode: } \text{O}_2 + 2e^- + 2\text{Li}^+ \xrightarrow{\text{discharge}} \text{Li}_2\text{O}_2(s) \\
\text{overall: } 2\text{Li} + \text{O}_2 \xrightarrow{\text{discharge}} \text{Li}_2\text{O}_2(s)
\] (1)

(2)

(3)

During the discharge operation, the Li metal is oxidized at the anode to generate lithium ion that transfers through the separator and reaches the porous cathode. The lithium ion reacts with oxygen from the ambient or gas channels and generates lithium peroxide (Li₂O₂) at the cathode. The oxygen reduction reaction (ORR) should occur both on triphase boundaries formed by the electrode material, the electrolyte and oxygen, as well as the interface of electrode material and electrolyte. The charging reactions at the anode and cathode...
are reversed from discharging reactions in which lithium peroxide is decomposed and oxygen is released at the cathode and lithium metal is deposited at the anode.

The O2 diffusion in the porous cathode is a key factor that determines the discharge capacity of Li–O2 batteries. Therefore, many studies on Li–O2 batteries have paid considerable attention to the O2 diffusion in the porous cathode because poor O2 diffusion will reduce the capacity. The O2 diffusion in the porous cathode is related to carbon loading, pore clogging induced by the deposition of solid lithium peroxide, and the partial pressure of oxygen. In addition, the wettability between the electrolyte and electrode also significantly affects the performance of Li–O2 batteries because the O2 diffusion coefficient in air is several orders of magnitude higher than that in the liquid electrolyte. For instance, the O2 diffusion coefficient in nonaqueous electrolyte calculated by the Stokes–Einstein equation is on the order of 10−6 whereas the diffusion coefficient is on the order of 10−2 in air. The contact angle is often applied to characterize the wetting behavior between the solid and liquid surface. The electrolyte and carbon material in electrode possess different dielectric constants, which affect the wettability of the solid to the liquid. The contact angle of the electrolyte on the surface of carbon electrode increases when the difference of dielectric constant between the electrolyte and carbon electrode is larger. The electrolyte with a larger contact angle is less likely to wet the carbon electrode. Xu et al. showed that the wettability affected the O2 diffusion in the porous electrode and thus the discharge and charge capacities. The measured contact angles of 1 M bis(trifluoromethane)sulfonimide lithium salt (LiTFSI) in various nonaqueous solvents and their mixtures on the surface of both carbon and Teflon showed that the discharge capacity was improved when the contact angle increased. For example, the contact angle of 1 M LiTFSI in propylene carbonate (PC)/ethylene carbonate (EC) (1:1 wt) on the surface of carbon was 47°, whereas that of 1 M LiTFSI in PC/dimethoxymethane (DME) (1:1 wt) on the surface of carbon was 5°. The discharge capacity with the PC/EC solvent was 167.5 mAh/g, whereas the capacity was only 27.4 mAh/g with the PC/DME solvent. Binders such as poly(vinylidene difluoride) (PVDF) and poly(tetrafluoroethylene) (PTFE) are used to combine different types of small particles together and adjust the wettability of cathode electrode in Li–O2 batteries. Zhang et al. investigated the wettability of carbon electrodes using the electrolyte of 1 M LiPF6 in PC/EC with a weight ratio of 1:1. The electrode made from Super P carbon mixed with PTFE (8:2 wt) had the contact angle of 99°, whereas the electrode made from KB 600 carbon mixed with PTFE (8:2 wt) had the contact angle of 56°. The battery with lyophobic electrode obtained a higher discharge capacity (3175 mAh/g) than the battery with a lyophilic electrode (798 mAh/g).

On the other hand, strongly lyophobic electrodes may deteriorate the discharge–charge performance. The Li–O2 battery using lyophobic electrode was researched by Chen et al. in which the experimental data showed that the specific discharge capacity increased from 1365 to 2365 mAh/g when the PTFE weight ratio increased from 0 to 30%. However, the electrode composed of 40 wt % PTFE deteriorated the performance of Li–O2 battery such that the specific discharge capacity decreased to 2130 mAh/g. The PTFE weight ratio should be optimized because a higher PTFE content will reduce the amounts of triphase boundaries for the electrochemical reactions battery electrode. The contact angles of water on the surface of raw carbon paper and carbon paper with 30 wt % PTFE were measured to be 136.6 and 148.3°, respectively. It should be noted that the water and liquid electrolytes have very different dielectric constants: the dielectric constant of water is 78.3 at 25 °C, whereas that of tetraethylene glycol dimethyl ether (TEGDME) is 7.9. Therefore, the contact angle measured by water cannot represent the surface wettability between electrode and liquid electrolyte.

Because of the significantly different O2 diffusivity in gas phase and electrolyte phase, electrodes with mixed wettability could create triphase boundaries that facilitate the oxygen transfer through gas phase (nonwetted pores). However, too many nonwetted pores will decrease the ionic conductivity of electrodes. Therefore, this study has conducted a systematic investigation on the effects of wettability of electrodes toward organic electrolyte (including both lyophilicity and lyophobicity) and its effects on the deep discharge capacity of Li–O2 battery. The wettability of electrode was adjusted by mixing carbon powders with different binders (PTFE and PVDF) at various weight ratios. The stability of binders has been systematically examined in the presence of commercial Li2O2, which showed that PVDF was unstable because of the presence of highly electron-withdrawing functional groups and an α or β hydrogen atom next to them. The PTFE binder was demonstrated to be stable in the presence of Li2O2. In addition, the PVDF binder was confirmed to degrade in Li–O2 batteries. The products formed in the PVDF degradation was measured by Raman spectroscopy and showed the Raman shifts of ~1123 and 1525 cm−1. However, the degradation can be prevented by drying the electrode under higher temperature, such as 200 °C. Thus, the stability of PTFE and PVDF binders can be guaranteed in this study if the electrodes are prepared according to the process in Experimental Section. The wettability of various customized electrodes was indicated by the contact angles of 1 M LiTFSI/TEGDME electrolyte on the customized electrode surface as well as the double-layer capacitance, which is proportional to the area of wetted pore surface in the electrode. After comparing the specific discharge capacities of electrodes with lyophilic and lyophobic carbon coating layers, the electrodes consisting of mixed coating layers were produced and tested to optimize the discharge–charge performance. The study shed light on the criteria to improve the O2 diffusion and increase the specific discharge capacity of Li–O2 batteries.

2. RESULTS AND DISCUSSION

2.1. Initial Deep Discharge. All these customized electrodes are tested by the deep discharge experiments at the current density of 0.1 mA/cm2. The discharge capacity of raw carbon cloth is negligible (<100 mAh/g) due to the low specific surface area. Therefore, the reported specific discharge capacity is calculated on the basis of the weight of carbon powders coated on the electrode without considering the weight of raw carbon cloth substrate and binders. The discharge curves with the highest discharge capacity among three repeated experiments of each electrode are shown in Figure 1, and the discharge capacities are compared in Figure 2. The lowest specific discharge capacity of 1665.8 mAh/g is achieved by the electrode with 15% PVDF carbon coatings. When the PTFE binder is used in the customized electrode, the electrode becomes more lyophobic. The specific discharge capacities of Li–O2 batteries increase from 3145.8 to 4160.8 mAh/g when the mass fraction of PTFE binder increases from 5 to 15% but...
decreases to 3109.5 and 2822.9 mAh/g when the mass fraction of PTFE binder further increases to 25 and 35%. The significant variation of specific discharge capacity can be attributed to the wettability change. To quantitatively correlate the discharge capacity with the electrode wettability, both the static contact angle and the double-layer capacitance of customized electrodes are measured and exhibited.

2.2. Contact Angle Measurement. The static contact angle between electrolyte and electrode surface is measured to indicate the surface wettability difference among customized electrodes with various contents of PTFE and PVDF binders. Instead of measuring contact angle between a water drop and electrode surface, the 1 M LiTFSI/TEGDME electrolyte (5 μL per drop) is dropped on the surface of customized electrodes to measure the static contact angle. The measured static contact angles (<90° for lyophilic surface and >90° for lyophobic surface) as well as images of electrolyte drops on electrode surfaces are shown in Figure 3.

The measurement of static contact angle is unsuccessful when the electrolyte drop is dripped on the surface of customized electrode containing 5% PTFE because the electrolyte droplet quickly spreads into the electrode. It indicates that the surface of 5% PTFE is lyophilic and more easily wetted by the electrolyte. When the PTFE content increases from 15 to 35%, the electrode becomes more lyophobic and the static contact angle increases from 128.4° to 138.5°. The mixed I electrode, which consists of one 15% PVDF coating sandwiched by two 15% PTFE coatings, displays a slightly lower static contact angle (118.8°) than the electrode with 15% PTFE. It may be caused by the fact that a small amount of PVDF binder is mixed on the top layer of carbon coating during the fabrication. In addition, the contact angles of customized electrodes with 15% PVDF carbon coatings as well as the mixed II electrode consisting of one 15% PTFE carbon coating sandwiched by two 15% PVDF carbon coatings are less than 90° and are unstable.

2.3. Double-Layer Capacitance Measurement. Since static contact angles represent the surface rather than internal wettability of customized electrodes, the double-layer capacitance of customized electrodes is also measured. This study employs an electrochemical approach to quantify the solid-liquid interfacial area and promote the understanding of wettability within customized electrodes. The electric double layer is created on the electrode surface that is wetted by the electrolyte because the customized electrode made of carbon powders is electrically conductive. The double-layer capacitance is related to the wetted interfacial area via the following equation

\[ C = \varepsilon_r \varepsilon_0 \frac{A_{sl}}{d} \]  

where \( C \) is the capacitance, \( \varepsilon_r \) is the relative permittivity of the electrolyte, \( \varepsilon_0 \) is the permittivity of the vacuum, \( d \) is the double-layer thickness, which can be approximated as Debye length, and \( A_{sl} \) represents the wetted pore surface. For an electrolyte with the given composition measured at the room temperature, parameters of \( \varepsilon_r, \varepsilon_0, \) and \( d \) can be considered as constants. Therefore, the change of the area of the wetted pore surface is proportional to the variation of the double-layer capacitance.

Figure 4 exhibits the relationship between the volume of added electrolyte and the measured double-layer capacitance. Considering the battery frame shown in Figure 8, the double-layer capacitance is related to the wetted pore surface in the electrode as well as the wetted area in the current collector. Thus, it is necessary to measure the capacitance when only the...
strongly lyophilic separator is inserted in the battery. Generally, as more electrolyte is added, more pore surface is wetted so that the double-layer capacitance increases. The customized electrode containing 15% PVDF has larger capacitances than that with 35% PTFE because the former electrode is more lyophilic and more pores are wetted by electrolyte. For example, the capacitance of the electrode with 15% PVDF was 12.9 μF more than that of the electrode with 35% PTFE at the volume of 20 μL. The capacitance of 35%-PTFE electrode combined with the separator is only slightly higher (<5 μF in all volumetric cases) than that of a single separator, as shown in Figure 4. It indicates that most of the electrolyte added into the lyophobic electrode is repelled into the lyophilic separator. The double-layer capacitance increases almost linearly with the amount of added electrolyte when the electrolyte is less than 60 μL. As the electrolyte is added, more electrolyte spreads into the separator. Therefore, the wetted surface in the current collector may account for a larger fraction of the double-layer capacitance. The sudden rise in the capacitance after 100 μL may be attributed to the fact that electrolyte is squeezed out from the battery frame and the extra double layer is created. Although the compression during battery assembly may partially compensate the capillary pressure within the electrode and cause measuring errors, especially when the electrolyte was more than 60 μL, the measurement of double-layer capacitance demonstrates the general trend that more pores are wetted by the electrolyte in the lyophilic electrodes.

2.4. Discussion. More lyophobic electrodes facilitate O2 diffusion by creating more gas paths. The improvement of O2 transfer results in a higher specific discharge capacity. However, the specific discharge capacity of the electrode decreases after further increasing the PTFE content to more than 15%. The influences of PTFE content on specific discharge capacity possesses good consistency with the published data.22 Further increasing the PTFE content generates more lyophobic pores that reduce the number of triphase boundaries for the ORR reaction and decreases the ionic conductivity. In Figure 5, the same trend as the static contact angle. In addition, the unstable discharge curves at the end of discharging process imply that lyophobic electrodes still experience O2 starvation on cathode, which increases the overpotential and causes the sudden death of the battery.

To facilitate the oxygen diffusion without significantly sacrificing the ionic conductivity, electrodes with mixed wettability have been designed and tested. The mixed electrode structures shown in Figure 1c,d consist of three carbon coatings. In mixed I electrode, the top and bottom layers with 15% PTFE are lyophobic and the middle layer with 15% PVDF is lyophilic. The battery with mixed I electrode achieves the highest specific discharge capacity of 5149.5 mAh/g. Because of the lyophobicity of both the top and the bottom coatings, O2 can diffuse into the electrode without much diffusion resistance and reach the electrolyte that accumulates in the lyophilic coating in the middle. This design utilizes the porous structure inside the electrode more effectively. Also, adding a lyophilic layer in the middle of electrode may reduce the O2 diffusion resistance and reach the electrolyte that accumulates in the lyophilic coating in the middle. The mixed I customized electrode with stacking coatings (lyophobic coatings on top and bottom with a lyophilic coating in the middle) can be beneficial to promote the design of air electrodes and provide a valuable reference to fabricate high-capacity electrodes.

2.5. Discharge–Charge Cycle. Although this study is focusing on the deep discharge capacity of Li–O2 batteries with customized electrodes, the cycling performance is also investigated and analyzed. The first discharge–charge cycles of Li–O2 batteries with customized electrodes at 0.1 mA/cm2 are shown in Figure 6. The cut-off voltage of charge is set as 4.5 V. The mixed I 15% electrode as well as PTFE 15% electrode result in a discharge–charge cycle with the average coulombic efficiency close to 85%, whereas the battery with 15% PVDF

![Figure 5](image_url)  
**Figure 5.** Equivalent ionic resistance of the batteries with customized electrode of different wettabilities.

![Figure 6](image_url)  
**Figure 6.** First discharge–charge cycle of Li–O2 batteries with customized electrode.
achieves a slightly higher coulombic efficiency of 93%. The lower coulombic efficiencies of mixed I 15% electrode and PTFE 15% electrode can be contributed to the fact that more electrolyte in the lyophobic electrode evaporates because of much longer discharge and charge time, which may increase the ionic resistance during charge and thus deteriorate the coulombic efficiency.39 Besides, PVDF favors the thin Li2O2 film formed on the surface of electrode, whereas PTFE results in more Li2O2 toroids, which are more difficult to decompose during charging.30 The cycling performance of Li−O2 batteries with PTFE 15% electrodes is also studied with the cut-off capacity of 1000 mAh/g. Figure 7 shows that coulombic efficiency is more than 90% in the first nine cycles and the discharge−charge curves are shown in Figure S4. The cycling performance of Li−O2 batteries using customized electrodes is validated. Further study will focus on optimizing the design of customized electrode since the rational structure can increase the cycle life of Li−O2 batteries.31,32

3. CONCLUSIONS

The wettability of electrode was mainly altered by mixing acetylene black carbons with PTFE and/or PVDF binders at various weight ratios. The customized electrodes containing PVDF binder or less PTFE binder (<15%) were lyophilic, and their corresponding discharge capacities were lower than those of lyophobic electrodes. The electrode with 15% PTFE exhibited the deep discharge capacity of 4160.8 mAh/g because lyophobic electrode provided more gas path for O2 diffusion. This study also designed customized electrodes with stacked layers by combining the lyophobic layers with lyophilic layers. The mixed I electrode (15% PTFE carbon coatings on top and bottom, one 15% PVDF carbon coating in the middle) showed the highest deep discharge capacity of 5149.5 mAh/g among all of the experimental cases. The mixed wettability promoted the oxygen diffusion without significantly lowering the ionic conductivity or reducing the number of reaction sites. Therefore, configuring the wettability in the electrode design is an important direction to increase the discharge capacity of Li−O2 batteries. Further study may focus on improving the configuration of customized electrode wettability considering the spatial distribution of pore size31 and specific pore volume of various carbon powders.

4. EXPERIMENTAL SECTION

4.1. Electrolyte Preparation. TEGDME (99%) purchased from Sigma-Aldrich was used as received to prepare the electrolyte. The electrolyte was prepared in a glovebox with <1 ppm water and <5 ppm oxygen. The electrolyte salt, LiTFSI (99.95%), from Sigma-Aldrich, was dissolved in TEGDME and the concentration was 1 mol/L.

4.2. Air Cathode Preparation. AvCarb 1071 HCB plain carbon cloth fabric purchased from Fuel Cell Store was used as the substrate of the customized cathode electrode on cathode. Conductive acetylene black (purchased from MTI Corporation) mixed with various weight ratios of PTFE was dissolved in the ethanol solution to form the carbon slurry. The substrate carbon cloth was dipped into the carbon slurry and dried in the atmosphere for 24 h. Then, the electrode was heat-treated at 350 °C for 30 min. The electrodes with customized coatings of 5, 15, 25, and 35% (in weight) of PTFE were obtained. In addition, electrodes with 15% PVDF carbon coatings and electrodes with mixed carbon coatings (both 15% PTFE and 15% PVDF) were prepared following the same procedure aforementioned. The schematic views on the structure of the customized electrodes are displayed in Figure 8. The targeted carbon loadings of all customized electrodes were 2 mg/cm2. The measured carbon loadings of the fabricated electrodes were 2.1 ± 0.1 mg/cm2.

4.3. Battery Assembly. The battery was constructed by sandwiching the oxygen diffuser, customized cathode electrode, separator, and lithium chip in the battery frame shown in Figure 9. The battery frame was customer-designed and made from highly corrosion-resistant grade 2 titanium. The open ratio of oxygen diffuser was 50% and the separator was a Whatman GF/L of electrolyte in the separator and in the cathode filter from Fisher Scientific with the diameter of 2.1 cm. The lithium chip with the diameter of 1.56 cm was purchased from MTI Corporation. The PTFE gasket and O-ring purchased from McMaster-Carr were inserted to prevent oxygen leaking. All batteries were assembled in the glovebox by adding 60 μL of electrolyte in the separator and in the cathode electrode. The substrate layer faced the O2 side and the 1 M LiTFSI/TEGDME electrolyte was added on the coating layer.

4.4. Wettability Measurements. The contact angles between 1 M LiTFSI/TEGDME and surfaces of customized
electrodes were measured using ramé-hart Model 190 contact angle goniometer at the room temperature. The static contact angle, which can be maintained for 120 s during measurement, was used to characterize the surface wettability. Each measurement was conducted at three different locations on the surface of customized electrodes using 5 μL of electrolyte drops. Double-layer capacitance measurement was carried out by the cyclic voltammetry (CV) tests with SP 150 potentiostat from BioLogic Science Instrument. The experimental setup was the same as the battery assembly in the discharge experiment. The 1 M LiTFSI/TEGDME electrolyte was added on the surface of the electrode by a pipette. All CV measurements were performed from −0.2 to 0.2 V at the scanning rate of 0.2 V/s and repeated at least three times. The EIS data were analyzed by the EC Lab software (V11.02).

4.5. Deep Discharge Measurements. Assembled batteries were connected to a four-channel Arbin MSTAT4 battery tester. The pure oxygen was supplied at a flow rate of 0.3 sccm, and the pressure of the pure oxygen was maintained at 10 kPa gauge pressure. After resting for 1 h, deep discharge tests were conducted at room temperature and repeated at least three times. First discharge—charge cycles of Li−O2 batteries with Mixed I, 15% PTFE and 15% PVDF carbon coatings, were also tested and analyzed.

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