Introduction to Metal–Air Batteries: Theory and Basic Principles
Zhiwen Chang and Xin-bo Zhang

Nowadays, energy is the power engine that sustains the operation of our society. In the energy field, we are confronted with a daunting challenge caused by the gradual depletion of fossil fuel. To secure a safe and sustainable energy supply, renewable energies such as solar and wind have been developed. However, these energies are geographically limited and intermittent, thus calling for reliable electrical energy storage (EES) system for stable and efficient power delivery. Simultaneously, the growing number of transportation vehicles has made the development of reliable EES system a task of urgency. Among various EES systems, rechargeable batteries are the most promising to meet these needs thanks to their high energy density and high energy efficiency [1]. Among them, the lithium-ion battery (LIB), which is operated on the basis of intercalation mechanism, has played an important role in our society in the past two decades [2]. However, the low energy density of LIB has restricted its application as the energy supplier of next generation. Under this circumstance, the development of metal–air battery has provided a solution benefitting from its much higher energy theoretical energy density than that of LIB.

In contrast to the closed system of LIB, the metal–air battery are featured with an open cell structure, in which the cathode active material, oxygen, coming from ambient atmosphere. In general, the metal–air battery consists of metal anode, electrolyte, and porous cathode. Metals such as Li, Na, Fe, Zn, etc. can be used as anode materials in metal–air batteries. And the theory and battery electro-chemistry will be briefly discussed on the basis of metal–air battery with different metallic anodes in the following section, which will be discussed in detail in the following chapters.

1.1 Li–O₂ Battery

Of all rechargeable metal–air batteries, the Li–O₂ battery (usually the aprotic Li–O₂ battery) possesses an ultrahigh theoretical energy density and is a promising EES. The theoretical energy density of the Li–O₂ battery can be
Introduction to Metal–Air Batteries: Theory and Basic Principles

around 11 586 Wh kg$^{-1}$ based on the mass of Li metal alone. When the mass of Li and Li$_2$O$_2$ is involved, the theoretical energy density of the Li–O$_2$ battery is still as high as 3505 Wh kg$^{-1}$, which is much higher than that of LIB [3]. The exceptionally high energy density of Li–O$_2$ battery mainly originates from two aspects. First, oxygen, the cathode material, is sourced from outside environment rather than being stored within the battery, thus helping to reduce the weight of the assembled cell. Second, during discharge, the lithium anode can deliver an extremely high specific capacity and rather low electrochemical potential (−3.04 V vs standard hydrogen electrode (SHE)), ensuring a desirable discharge capacity and a high operation voltage, respectively [4]. The history regarding the development of Li–O$_2$ battery is introduced briefly as follows. The first prototype of Li–O$_2$ battery was reported by Semkow and Sammells [5]. In 1996, a Li–O$_2$ battery with polymer-based electrolyte was introduced by Abraham [6]. During the following couples of years, Read et al. have carried out relevant researches in the Li–O$_2$ field, and Bruce has demonstrated the rechargeability of the system [7–9]. Since then, numerous efforts have been devoted into the Li–O$_2$ field along with success of varying degrees. Currently, there are four types of Li–O$_2$ batteries under investigation, which can be categorized on the basis of the applied electrolyte species (aprotic, aqueous, hybrid, and all solid-state electrolytes) [10]. All the four types of lithium–air batteries use lithium metal and oxygen (air) as anode and cathode active materials, respectively. Their fundamental electrochemical reaction mechanisms are closely associated with the electrolytes used. Simultaneously, the schematic illustration of these four types of Li–O$_2$ batteries is provided in Figure 1.1, being able to provide the readers with an easy access to their configuration.

Compared with the Li–O$_2$ batteries with aqueous, hybrid, and solid-state electrolytes, the researches on the Li–O$_2$ battery has taken the dominant place. So in the following section, all the discussion is around the Li–O$_2$ battery with aprotic electrolyte. A typical Li–O$_2$ battery consists of a lithium-metal anode, a porous carbon cathode, and a separator saturated with aprotic electrolyte, which is shown in Figure 1.1. In principle, the Li–O$_2$ chemistry is based on the following conversion reaction: [11]

\[
2\text{Li} + \text{O}_2 \leftrightarrow \text{Li}_2\text{O}_2 \quad E^0 = 2.96 \text{V vs Li}^+/\text{Li} \quad (1.1)
\]

The ideal operation of a Li–O$_2$ battery is based on the electrochemical formation (discharge) and decomposition (charge) of lithium peroxide (Li$_2$O$_2$). The reduction proceeds through the following general steps:

\[
\text{O}_2 + \text{Li}^+ + e^- \rightarrow \text{LiO}_2 \quad (1.2)
\]

\[
\text{LiO}_2^* + \text{Li}^+ + e^- \rightarrow \text{Li}_2\text{O}_2 \quad (1.3)
\]

\[
2\text{LiO}_2 \rightarrow \text{Li}_2\text{O}_2 + \text{O}_2 \quad (1.4)
\]

It is worthy to note that the growth process of Li$_2$O$_2$ is very complicated. Currently, two models of O$_2$ reduction have been proposed, including surface-growth model and solution-growth model. During discharge, O$_2$ undergoes a one-electron reduction to generate O$_2^-$. In the surface-growth model,
the superoxide species (O$_2^-$ and/or LiO$_2^*$, where * indicates surface-adsorbed O$_2^-$ and LiO$_2$) adsorb on the cathode surface and undergo a second reduction, forming Li$_2$O$_2$ thin films (thickness <10 nm) on the cathode surface, severely passivating the cathode and limiting the maximum battery discharge capacity [12, 13]. In contrast, in the solution-growth pathway, the generated LiO$_2$ is dissolved in the electrolyte, which undergoes disproportionation, ultimately forming large Li$_2$O$_2$ toroidal crystals of variable sizes (typically <2 μm) [14–16]. Clearly, the reaction pathways can be affected by the solubility of LiO$_2$ in the electrolyte, which can be decided by the equilibrium between LiO$_2$ adsorbed on the electrode and LiO$_2$ dissolved in the electrolyte solution. Such an equilibrium is listed below: [17]

\[ \text{LiO}_2^* \leftrightarrow \text{Li}_{(\text{sol})}^{+} + \text{O}_2^{(\text{sol})}^- + \text{ion pairs} + \text{higher aggregates} \]  

(1.5)

According to Johnson et al. [18], at high discharge voltages (low discharge overpotentials) and in solvent with high donor number (DN), such as 1-methylimidazole (Me-Im, DN = 47), the Li$^+$ or Li$^{2+}$-containing species is solvated, and the equilibrium (1.5) is displaced to the right, yielding mainly soluble LiO$_2$. While in low DN solvents, such as CH$_3$CN (DN = 14), the solvation is weaker, and the equilibrium lies to the left, resulting in surface-adsorbed LiO$_2$ being dominant. In the latter case, the LiO$_2^*$ disproportionates or undergoes a second reduction to Li$_2$O$_2$ on the electrode surface, whereas in the former
Similarly, Aetukuri et al. have prompted a solution-based mechanism with trace amount of electrolyte additives, such as H₂O, which increases the solubility of LiO₂ with the O₂⁻ acting as a redox mediator [19]. In addition, the reaction pathways can be affected by other factors such as current density, the cathode adsorption ability, etc. [20, 21] As a typical example, in 2013, Xu et al. have promoted the solubility of superoxide into the electrolyte with Pd nanocrystal [21]. In his research, two kinds of cathodes including hollow spherical carbon (HSC) deposited onto carbon paper (CP) cathode (HSC deposited onto CP) and palladium (Pd)-modified HSC deposited onto CP cathode (P-HSC deposited onto CP) are prepared. In the P-HSC deposited onto CP cathode, the strong binding of superoxide (O₂⁻ or LiO₂⁻) with the carbon defect sites traps the superoxide on the cathode surface (a short diffusion path), leading to a poor superoxide availability on the prismatic crystal faces (a preferred nucleation site) and gives rise to the observed toroid aggregates for Li₂O₂. In contrast, in the P-HSC deposited onto CP cathode, the Pd nanocrystal surfaces are considered to be less sticky than carbon surfaces riddled with dangling bonds, which can weaken the binding of the generated superoxide to the substrate, thus enhancing the diffusion of the superoxide molecules away from the cathode surface, favoring the nucleation and crystallization of Li₂O₂ on the preferred prismatic crystal faces and leading to nanosheet-shaped structure growth.

The charge process of Li–O₂ battery involves the electrochemical decomposition of Li₂O₂ formed on the air electrode, which is accompanied by several metastable states. In a similar case to the discharge process, the charge process of Li₂O₂ decomposition is also very complicated, involving many steps. In this aspect, several points such as surface delithiation and bulk oxidation of Li₂O₂ proposed by Gallant [22], the promoted one-electron charge reactions [23], etc. have been provided. Of note is that these achievements are insufficient to clarify the intrinsic mechanisms involving the discharge and charge, thus calling for more relevant efforts.

Although the study on the Li–O₂ batteries has been ongoing over the past decade, they have been hindered from commercialization due to several practical challenges. First is the sluggish reaction kinetics that lead to a high charge overpotential, thus inducing electrolyte decomposition. The second problem is the poor conductivity of Li₂O₂. Catalysts have been incorporated to promote the formation of amorphous Li₂O₂, whose conductivity is better than that of crystalline Li₂O₂. The third problem is the high oxidative ability of these species, degrading cathode and electrolyte to give parasitic products, which can passivate the active sites. In this aspect, various strategies have been proposed to enhance the chemical stability of carbonaceous cathode and fabricated noncarbonaceous cathodes. In addition, various kinds of catalyst have been applied to accelerate the reaction kinetics. Simultaneously, the selection of highly stable electrolyte is also required. Of note is that the oxygen and ionic conductivity of the electrolyte are also parameters that can affect the battery performances. After all, a Li–O₂
As mentioned in the above section, the Li–O\textsubscript{2} battery is viewed as a promising energy supplier of next generation. However, this battery suffers from terrible problems, including poor efficiency and low rate capability, which have restricted its practical application. Particularly, the serious charge overpotential over 1 V, the main reason for the low efficiency, also accelerates the degradation of the electrode and electrolyte. Under this circumstance, the Na–O\textsubscript{2} battery is viewed as a promising alternative of Li–O\textsubscript{2} battery, thanks to the merits of Na–O\textsubscript{2} battery compared with the Li–O\textsubscript{2} battery. In light of previous reports, the main advantage of a Na–O\textsubscript{2} battery is its much lower charge overpotential than that in typical Li–O\textsubscript{2} batteries. Simultaneously, in contrast to lithium, sodium resources are unlimited everywhere, and sodium is one of the most abundant elements in the Earth’s crust [17].

In 2011, the first Na–O\textsubscript{2} battery with molten sodium anode is reported by Peled et al. [24] However, this battery is operated at above 100 °C, thus limiting its practical application in our daily lives. Luckily, later in 2012, a Na–O\textsubscript{2} battery constructed at room temperature is reported by Janek’s group [25]. Since then, the researches in Na–O\textsubscript{2} battery conducted at room temperature have been surging. Currently, the Na–O\textsubscript{2} batteries can be split into two categories according to the used electrolyte, including nonaqueous and aqueous electrolyte [26]. It is worthy to note that the researches on the aprotic Na–O\textsubscript{2} battery have taken the dominant place, based on which all the discussions mentioned below are all around nonaqueous Na–O\textsubscript{2} battery. In a common configuration, the Na–O\textsubscript{2} battery comprises a sodium-containing anode (currently Na metal), a sodium-conducting organic electrolyte, and an air cathode. Unlike the cell chemistry in the Li–O\textsubscript{2} battery, where the targeted discharge product is only Li\textsubscript{2}O\textsubscript{2}, the discharge products in the Na–O\textsubscript{2} battery at least include Na\textsubscript{2}O\textsubscript{2} and NaO\textsubscript{2}. In detail, during discharge, the sodium metal anode is electrochemically oxidized to produce Na\textsuperscript{+} ions, which migrates across the electrolyte to reach the cathode. Simultaneously, the oxygen gas (O\textsubscript{2}) dissolved in the electrolyte is reduced to form superoxide species (O\textsubscript{2}–) or peroxide species (O\textsubscript{2}\textsuperscript{2–}), generating sodium superoxide (NaO\textsubscript{2}) or sodium peroxide (Na\textsubscript{2}O\textsubscript{2}), respectively. Upon charge, the generated NaO\textsubscript{2} or Na\textsubscript{2}O\textsubscript{2} is decomposed with the release of O\textsubscript{2} and Na\textsuperscript{+}.

In terms of cell chemistry, the free enthalpy of formation of \( \Delta_f G(\text{Na}_2\text{O}_2) = -437.5 \text{ kJ mol}^{-1} \) [27]. The standard open cell potential for the formation of Na\textsubscript{2}O\textsubscript{2} is estimated to be \( E(\text{Na}_2\text{O}_2) = 2.33 \text{ V} \). And the theoretical energy density of a Na–O\textsubscript{2} battery based on Na\textsubscript{2}O\textsubscript{2} can be \( \sim 1600 \text{ Wh kg}^{-1} \). In 2013, Li et al. reported the application of nitrogen-doped graphene nanosheets (N-GNSs) as cathode catalyst in the Na–air battery, which has promoted the formation and
decomposition of $\text{Na}_2\text{O}_2$ [28]. In their study, N-GNSs displayed a discharge capacity two times greater than their pristine counterpart, as well as superior electrocatalytic activity as a cathode material for sodium–air batteries. The enhanced performance of N-GNSs is attributed to the active sites introduced by nitrogen doping. In another case, Liu et al. reported the formation of $\text{Na}_2\text{O}_2$ nanosheets, which grow on the NiCo$_2$O$_4$ nanosheets fabricated on Ni foam [29]. In this research, the Na–air battery based on the NiCo$_2$O$_4$ nanosheets exhibits an initial discharge capacity of 1762 mAh g$^{-1}$ with a low polarization of 0.96 V at 20 mA g$^{-1}$.

In the pursuit of a Na–O$_2$ battery with excellent performances, the $\text{Na}_2\text{O}_2$ is not the desirable discharge product. Generally, the battery with $\text{Na}_2\text{O}_2$ shows poor reversibility and high overpotentials, similar to the problems in recharging Li–O$_2$ batteries [24, 30, 31]. In response, the generation of NaO$_2$ rather than $\text{Na}_2\text{O}_2$ should be promoted, given the fact that the charge and discharge overpotentials can be reduced to less than 200 mV when NaO$_2$ forms [32, 33]. From the thermodynamic point of view, NaO$_2$ is less stable than $\text{Na}_2\text{O}_2$. With a free enthalpy of formation of $\Delta_f G(\text{NaO}_2) = -437.5 \text{kJ mol}^{-1}$, the formation of NaO$_2$ is energetically rather similar to that of $\text{Na}_2\text{O}_2$, with a small energy difference of only 12 kJ mol$^{-1}$ [27]. This small difference is also expressed by the quite close standard open cell potentials, $E(\text{NaO}_2) = 2.27 \text{V}$ and $E(\text{Na}_2\text{O}_2) = 2.33 \text{V}$. Theoretical calculations indicate that the relative thermodynamic stability of NaO$_2$ and $\text{Na}_2\text{O}_2$ depends on the crystallite size; nanosized NaO$_2$ appears to be more stable than $\text{Na}_2\text{O}_2$ of equal size, and nucleation of NaO$_2$ might thus indeed be favored [34, 35]. So far, various researches regarding the formation and decomposition of NaO$_2$ are reported. These products have been shown to have different morphologies: NaO$_2$ in micron-scale cubic shapes, nanorods, and conformal films.

Just like the case in Li–O$_2$ battery, it is critical to gain insights into the oxygen reduction kinetics in the presence of Na ions and the nucleation and growth mechanisms of discharge products. In 2015, Vitoriano et al. reported the rate-dependent effects on the size and distribution of the discharge product [35]. In addition, the type of solvent can affect the discharge reactions as well. In the report by Luntz et al. [36], strong solvent–solute interactions in long-chain ethers shift the formation of NaO$_2$ toward a surface process, resulting in submicrometric crystallites. In contrast, short chains, which facilitate desolvation and solution precipitation, promote the formation of large cubic crystals (c. 10 μm). In addition, the current rates are also reported to affect the growth of NaO$_2$ [35]. At low rates, the entire thickness of the carbon nanotube (CNT) carpet was more homogeneously covered with faceted nanoparticles. While at high rates, micron-sized cubes were formed on the top and bottom interfaces between the CNT carpet and electrolyte. In fact, nucleation and growth under electrochemical conditions are complex processes [37], and future mechanistic studies are required to obtain a more coherent picture. For these considerations, relevant discussions are only provided here.

Apart from the mentioned controversies, there is a general agreement in literature that Na–O$_2$ batteries can be a good alternative to overcome some of the drawbacks of Li–O$_2$ batteries. Thus, their promising electrochemical performance could open new opportunities beyond LIB. However, the development of
sodium–air battery is still at its infancy, and more efforts are necessary to realize its practical application.

References


