The water catalysis at oxygen cathodes of lithium–oxygen cells

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Lithium–oxygen cells have attracted extensive interests due to their high theoretical energy densities. The main challenges are the low round-trip efficiency and cycling instability over long time. However, even in the state-of-the-art lithium–oxygen cells the charge potentials are as high as 3.5 V that are higher by 0.70 V than the discharge potentials. Here we report a reaction mechanism at an oxygen cathode, ruthenium and manganese dioxide nanoparticles supported on carbon black Super P by applying a trace amount of water in electrolytes to catalyse the cathode reactions of lithium–oxygen cells during discharge and charge. This can significantly reduce the charge overpotential to 0.21 V, and results in a small discharge/charge potential gap of 0.32 V and superior cycling stability of 200 cycles. The overall reaction scheme will alleviate side reactions involving carbon and electrolytes, and shed light on the construction of practical, rechargeable lithium–oxygen cells.
The Li–O₂ cells are operated with oxygen reduction to produce Li₂O₂ and its reverse oxidation to release Li⁺ ions and oxygen at cathodes (2Li⁺ + O₂ + 2e⁻ → Li₂O₂, $E^\circ = 2.96\text{ V vs Li}^{+/\text{Li}}$) in discharging and charging processes, respectively.¹⁻³ The intrinsic chemical and physical nature of Li₂O₂ and its intermediates can lead to instability of other cell components, such as carbon supports (> 3.50 V)⁴ and electrolytes, and formation of detrimental byproducts, typically Li₂CO₃ and lithium alkyl carbonates⁵⁻⁹. They can subsequently induce high overpotentials in both discharging and charging processes and eventually discharge/charge failure during cycles.¹⁰⁻¹³ A lot of efforts have been made to address these issues, but there are still many challenges for a practical Li–O₂ cell.¹⁴⁻¹⁶

The success of deploying Li–O₂ cells in the future will be crucially dependent on how small the overpotentials at the cathode side can be reduced and how many cycles they can work reversibly.¹⁵⁻¹⁷ Currently, two strategies have been widely applied for these issues. One is to use carbon-free or carbon-alternative cathodes, like nanoporous gold¹⁰, TiC¹¹ and conductive metal oxide supported Ru¹²⁻¹³, which can circumvent side reactions involving carbon. The charge overpotentials ensuring the decomposition of Li₂O₂ are reduced to ~0.54 V, corresponding to charge potentials of ~3.50 V (refs 10–13). This direct electrochemical oxidation of Li₂O₂ was revealed to involve multiple processes, in which the initial delithiation occurring at an overpotential of 0.44 V, possibly suggests the theoretical limit¹⁸⁻¹⁹. Alternatively, redox mediators have been introduced to chemically oxidize Li₂O₂ and the apparent charge potentials are typically determined by the mediators. For instance, tetrathiafulvalene and LiI reduce the charge potential to ~3.50 V, equal to a charge overpotential of 0.54 V (refs 20–23).

In principle, Li₂O₂ can be oxidized by mediators possessing higher redox potentials than its equilibrium potential; however, practical mediators are very limited. A more efficient and compatible one for further potential reduction is still lacking. By either the direct or indirect (mediators) Li₂O₂ oxidation strategy, it will be a great challenge to reduce the charge potentials to feasible values. Exploring new reaction mechanisms at the oxygen cathodes to significantly improve the round-trip efficiency and cycling stability is essential and urgently necessary for the development of Li–O₂ cells.

Although water in electrolytes has been found to affect the morphologies of discharge products and increase the discharge capacity,¹²⁻¹⁴ its presence in electrolytes or O₂ atmosphere resulted in rapid charge potential increase and hence cell death after several cycles.²⁶⁻²⁸ In addition to the main discharge product Li₂O₂, a byproduct LiOH was also detected.²⁷ We found that the decomposition of LiOH is strongly related to the applied catalysts, such as Ru nanoparticles supported on Super P (Ru/SP). To reduce the charge overpotentials of Li–O₂ cells, a trace amount of water in electrolytes and electrolytic manganese oxide (EMD, γ-MnO₂) nanoparticles are both utilized to favour the transformation of the discharge product from Li₂O₂ to LiOH and its following decomposition on charging. The MnO₂ incorporated in Ru/SP (Ru/MnO₂/SP) also allows for the water regeneration at the cathode during the discharge/charge cycles.

This enables the Li–O₂ cell to operate with a small discharge/charge potential gap and superior cycling stability. A reaction mechanism by converting Li₂O₂ to LiOH in the presence of both water and Ru/MnO₂/SP is proposed.

**Results**

**Low overpotentials at Ru/MnO₂/SP.** The Li–O₂ cells were constructed with Ru/MnO₂/SP pressed onto a carbon paper as cathodes, 0.5 M LiClO₄ in DMSO containing 120 p.p.m. of H₂O as electrolytes and LiFePO₄ in replacement of Li anodes. The LiFePO₄ is not a practical anode, but is able to avoid the reaction of Li metal and the trace amount of H₂O and any contamination from formation of the solid electrolyte interface layer on the Li anode. It has a stable potential of ~3.45 V vs Li⁺/Li regardless of the state of charge and favours the investigation on the discharge/charge behaviour in the presence of H₂O and the underlying reaction mechanism specifically at the oxygen cathode. The discharge/charge potentials at the oxygen cathode of the Li–O₂ cells are converted to against Li for discussion.

The carbon paper has been demonstrated to have negligible contribution to cell performance (Supplementary Fig. 1; Supplementary Note 1). The Li–O₂ cell with Ru/MnO₂/SP as cathode and the DMSO-based electrolyte containing p.p.m.-leveled H₂O is discharged and charged at 500 mA g⁻¹ (corresponding to 0.71 μA cm⁻² based on the Brunauer-Emmett-Teller, (BET) surface area of Ru/MnO₂/SP) as shown in Fig. 1a. A charge potential plateau as low as ~3.20 V, corresponding to an overpotential of ~0.24 V, can be clearly observed. Its corresponding dQ/dV curve in Fig. 1b shows sharp oxygen reduction reaction and oxygen evolution reaction peaks in the discharging and charging process, respectively, which are consistent with the flat discharge and charge plateaus. In addition, a small plateau at ~3.60 V is also obtained in the charge profile in Fig. 1a. It could be resulted from the affinity of DMSO with the

![Figure 1](https://www.nature.com/naturecommunications)
Ru/MnO₂/SP cathode that has a large contact angle of 62°, compared with the discharge and charge profiles of the Li–O₂ cell with a dimethoxyethane-(DME-) based electrolyte and the good wettability of DME on the cathode in Supplementary Fig. 2. This may enable the oxygen evolved in a charging process to accumulate in the voids between the residual discharge product and the cathode, as schematically described in Supplementary Fig. 3 and Supplementary Note 2. It would induce slow diffusion of electrolytes into the voids for the decomposition of the remaining discharge products and result in the relatively large overpotential at ~3.60 V, which is not observed in the affinitive DME-based electrolyte (Supplementary Fig. 2). Similar observations are also obtained in the other kinds of electrolytes, such as triglyme (G3)- and tetraglyme (G4)-based electrolytes (Supplementary Fig. 4). The trace amount of H₂O in electrolytes is demonstrated to play a crucial role on reducing charge overpotentials of Li–O₂ cells in all the widely used electrolytes, which was empirically considered as a negative factor in a Li-ion cell.

Analysis on the discharged/charged cathode. The discharged and charged Ru/MnO₂/SP cathodes in the DMSO-based electrolyte containing water have been characterized by X-ray diffraction (XRD). As shown in Fig. 2a, the discharge products are identified as a mixture of LiOH and Li₂O₂, referring to the standard powder diffraction files of 01-085-0777 and 00-009-0355, respectively. The diffraction peaks of LiOH become sharper and stronger at the cathode with a discharge capacity of 4,000 mAh g⁻¹ and addition of more electrolyte (Supplementary Fig. 5), suggesting more LiOH converted from Li₂O₂. To avoid the effect of MnO₂, quantification of LiOH and Li₂O₂ was conducted on the Ru/SP cathode with the same discharge capacity as in Fig. 1 via iodometric titration. The discharge product Li₂O₂ reacts with H₂O via Li₂O₂(s) + 2H₂O(l) → H₂O₂(g) + 2LiOH(aq) 30,31, where H₂O₂ further oxidizes iodide to iodine, the titrated target. The LiOH and Li₂O₂ in the discharged cathode were titrated via two steps (Supplementary Figs 6 and 7; titration processes in Supplementary Methods) and estimated to 16.02 and 1.48 μmol, respectively, which are in agreement with the XRD patterns of the discharged cathode in Fig. 2a. The majority of LiOH formed at a discharged cathode is revealed by the characteristic absorbance peak in the infrared (IR) spectra in Fig. 2b. Based on the electrons passing through the cathode and the oxygen derived from the discharge products of LiOH and Li₂O₂ by iodometric titration, the discharging process is a 1.97 e⁻/O₂ process, quite close to the theoretical value of 2.00 e⁻/O₂. It is confirmed that in other electrolytes, like DME-, G3- and G4-based electrolytes, the discharging process is also a ~2.00 e⁻/O₂ process, as listed in Supplementary Table 1. This suggests that LiOH is converted from Li₂O₂ via a chemical, not an electrochemical, process in the discharging processes.

After charge, the discharge products LiOH and Li₂O₂ were decomposed, as evidenced by the disappearance of their characteristic diffraction peaks in Fig. 2a. Further, the discharge products were directly observed by scanning electron microscope (SEM) in Fig. 2c. All the products are toroidal particles with obvious layering. These are quite similar to the toroidal aggregates in a Li₂O₂-only discharged cathode 32,33, which may suggest that the LiOH was derived from Li₂O₂ in the presence of H₂O in electrolytes by persisting the similar shape. The charged cathode becomes more porous after decomposition of the discharge products in Fig. 2d in comparison to the fresh cathode in Fig. 2e. These results indicate that the low charge overpotentials demonstrated in Fig. 1 and Supplementary Fig. 3 resulted from the electrochemical decomposition of LiOH.

Roles of Ru and MnO₂ during discharge and charge. In the Ru/MnO₂/SP cathode, all the components of Ru12,13, MnO₂ (ref. 34) and SP35 nanoparticles can act as oxygen reduction reaction catalysts. However, in the charging processes either SP or MnO₂ supported on SP (MnO₂/SP) did not show any plateaus at ~3.2 V but instead at >3.6 V (Supplementary Figs 8 and 9). Only the Ru/SP as revealed in Supplementary Fig. 10 presented low charge potentials. This suggests good catalytic activity of Ru nanoparticles on the decomposition of LiOH. Fig. 3 shows 50 cycles of discharge and charge of the Li–O₂ cell with the Ru/SP cathode. The charge potentials are sustained at ~3.20 V in the initial few cycles, and then quickly are increased to ~3.65 V in
the tenth cycle and beyond. It indicates that the majority of the discharge product Li₂O₂ after multiple cycles cannot be converted to LiOH, because the p.p.m.-leveled H₂O in the electrolyte was gradually consumed in the charging processes to produce H₂O₂. Hence, MnO₂ (EMD), the best disproportionation catalyst of H₂O₂ (ref. 36), was incorporated into Ru/SP to regenerate H₂O via 2H₂O₂(ℓ) → 2H₂O(ℓ) + O₂(ℓ) and make H₂O circulate during cycles and improve cycling stability of the Li–O₂ cell.

**Rate capability and cycling stability.** The Li–O₂ cell with Ru/MnO₂/SP and the DMSO-based electrolyte containing p.p.m.-leveled H₂O was examined at varied current densities, as shown in Fig. 4a. The polarization is obviously increased with the current density from 250 to 500 and 1,000 mA g⁻¹. The over-potential in the discharging and charging process at 250 mA g⁻¹ is 0.11 V and 0.21 V, respectively, leading to a small discharge/charge potential gap of 0.32 V. The Li–O₂ cell was continuously discharged and charged at 500 mA g⁻¹ for 200 cycles and ~800 h, and the selected runs of discharge and charge are shown in Fig. 4b. The discharge and charge curves during the first 150 cycles are almost overlapped except the first charge. A slight charge potential increase beyond the 150 cycles can also be observed, which could be related to the electrolyte instability during many cycles (Supplementary Fig. 11). This indicates superior cycling stability of the Li–O₂ cell, which is in sharp contrast to that without MnO₂ in Fig. 3. The discharge and charge capacities in the 200 cycles are almost constant, and the corresponding coulombic efficiency in each run is approaching 98% (Fig. 4c), indicative of good reversibility. This may be partially benefited from the conversion of chemically active Li₂O₂ to LiOH. The reversible formation and decomposition of LiOH and Li₂O₂ during the 200 cycles are further confirmed by ex situ XRD patterns in Supplementary Fig. 12 and SEM images in Supplementary Fig. 13 (Supplementary Note 3). The low charge potentials sustained for so many cycles, to the best of our knowledge, have never been achieved before. The small discharge/charge potential gap and good cycling stability of the Li–O₂ cell are rewarded by the ‘water catalysis’ at the Ru/MnO₂/SP cathode.

**Reaction mechanism.** The equilibrium potential of LiOH in an aqueous solution is estimated to be ~3.42 V vs Li⁺/Li based on Nernst equation. However, in this studied electrolyte system the trace amount of H₂O and LiClO₄ are both the solutes and DMSO is the solvent. The equilibrium potential of LiOH is dependent on the concentration of H₂O, and it can be roughly estimated to be ~3.20 V, considering a concentration of 100 p.p.m. of H₂O in the electrolyte (see the estimation process in Supplementary Methods). This is in good agreement with the observed low charge potential plateaus in all the widely employed electrolytes DMSO-, DME-, G3- and G4-based electrolytes for Li–O₂ cells in Figs 1 and 4 and Supplementary Fig. 3.

Based on the above results, a mechanism for the discharging and charging process of the cell with Ru/MnO₂/SP and the electrolyte containing a trace amount of H₂O can be proposed and schematically described in Fig. 5a. On discharging, O₂ accepts electrons via the external circuit and is reduced to generate the primary discharge product Li₂O₂ (refs 40–42). At the same time, the Li₂O₂ reacts with H₂O from the electrolyte and is converted to LiOH via Steps (i and ii). Although Step (i) is an equilibrium,
it can be largely promoted to move forward by the conversion of one of the products H2O2 to H2O over MnO2 via Step (ii)36. The two steps (i and ii) occur sequentially, and quickly transform Li2O2 to LiOH as long as H2O remains in the electrolyte. This has been confirmed by the presence of substantial LiOH in discharged cathodes as revealed by both of the XRD patterns and the IR spectra in Fig. 2. This is consistent with the observations of Aetukuri et al.25, and Schwenke et al.26, where the H2O in electrolytes is possibly consumed by the employed Li anode or saturated by the product LiOH, and the lack of a promoter like MnO2 for Step (ii) made the equilibrium reaction in Step (i) to move backward and result in the major discharge product Li2O2 as detected.

In the following charging process, the resultant LiOH can be directly oxidized via Step (iii) to regenerate H2O at low charge potentials, by which the residual Li2O2 is then converted to LiOH via Steps (i) and (ii) and oxidized. To demonstrate the feasibility of Step (iii), commercial LiOH was ball-milled and then thoroughly mixed with Ru/SP with a ratio of 30:70 (wt/wt). The electrodes of Ru/SP with and without LiOH under Ar atmosphere were subjected to linear scanning voltammetry (LSV) at a low scan rate of 0.01 mV s−1 as depicted in Fig. 5b. The Ru/SP electrode incorporated with LiOH presents significant oxidation currents and an oxidation onset potential of ∼3.27 V as depicted in Fig. 5b, which is in sharp contrast with no oxidation response on the electrode without LiOH. At a carbon electrode pre-filled with LiOH, a high charge potential of >4.0 V was reported43. This suggests that over Ru/SP LiOH was oxidized at low potentials, though it may be dependent on its morphology and facet orientation and the catalyst. The gas evolved in a charging process can be identified as O2 by a gas chromatography (GC) mounted with a thermal conductivity detector, as evidenced by the sharp GC signal of O2 in Fig. 5c. As shown in the whole discharging and charging process.
process in Fig. 5a, H₂O is not consumed and can be circulated to behave like a catalyst. The total electrochemical reaction occurring at the oxygen cathode is 2Li⁺ + O₂ + 2e⁻ → Li₂O₂, consistent with the Li–O₂ cell chemistry.

**Discussion**

The dependence of the charge potentials on the concentration of H₂O in electrolytes is shown in Fig. 6. In the dried DMSO-based electrolyte, one charge potential plateau can be obtained at ~3.65 V, which is attributed to the oxidation of Li₂O₂ and in good agreement with the literatures. When there is 120 p.p.m. of water in the electrolyte, the charge potential plateau is significantly reduced to ~3.2 V. This has been confirmed to the oxidation of LiOH that can be quickly converted from the primary discharge product Li₂O₂ via the reaction Li₂O₂ + 2H₂O → 2LiOH + O₂. The solution to the current challenges of Li–O₂ cells. With the potentials following the Nernst equation,

$$E = E^\circ + \frac{RT}{nF} \ln \left( \frac{P_{O_2}}{P_{O_2}^0} \right)$$

where $E$ is the electrode potential, $E^\circ$ is the standard electrode potential, $R$ is the gas constant, $T$ is the absolute temperature, $n$ is the number of electrons transferred, $F$ is the Faraday constant, $P_{O_2}$ is the partial pressure of oxygen, and $P_{O_2}^0$ is the standard pressure of oxygen. The resulting mixture was further stirred for 12 h. For the preparation of MnO₂/Ru/SP, the as-synthesized Ru/SP and MnO₂ nanoparticles with a weight ratio of Ru/MnO₂/SP = 7:5:7.5 were sonicated into an ethanol aqueous solution for 3 h and stirred overnight. Then, the product MnO₂/Ru/SP was collected and dried at ~80 °C under vacuum for 12 h.

**Cell assembly.** The electrode film composed of Ru/MnO₂/SP or Ru/SP and PTFE (A dispersion of 60 wt%, Du Pont-Mitsui Fluorochemicals Co. Ltd.) with a ratio of 85:15 wt% was rolled with a glass rod. The mass loading of Ru/MnO₂/SP or Ru/SP is ~0.5 mg cm⁻². The electrode film was pressed onto a hydrophobic carbon paper (SIGRACET Gas Diffusion Media, Type GDL, 33BA) to work as a cathode. The anode was consisted of LiFePO₄ (Sumitomo Osaka Cement, SP), and PTFE (70/20/10) pressed on an Al mesh. The amount of DMSO- and DME-based electrolytes in a coin cell was 100 μl and G3- and G4-based electrolyte was 50 μl, considering their different vapour pressures. The employed separator was a glass microfiber filter paper (GF/A, Whatman). The Li₂O₂/cell assembly in 2-M coin cell was conducted in an Ar-filled glove box that has a dew point of around ~90 °C (~0.1 p.p.m. of H₂O) and O₂ content below 5 p.p.m.. The cells stored in a glass chamber with a volume capacity of 650 ml were purged with O₂ (99.999%) before electrochemical tests.

**Characterization and measurements.** The contact angles of electrolytes on the cathodes were examined on AST Products with a model of Optima. The BET surface area of Ru/MnO₂/SP was measured to be 70.6 m² g⁻¹. Electrolytic MnO₂ (EMD, Bubendorf) was received from TOSOH, Japan and ball-milled to nanometre scale. Ru/SP was synthesized as follows: 85 mg of SP was stirred into a solution of ethylene glycol (EG, purity of >99.95%, Wako Chemicals) containing 7.5 mg of Ru in RuCl₃·xH₂O (purity of 36 ~ 42 wt% based on Ru, Wako Chemicals), and its pH was adjusted to 13 with 0.1 M of NaOH in EG. The suspension was then heated to 160 °C for 3 h with flowing N₂. After cooling to 80 °C, its pH was adjusted to 3 using 0.1 M HCl and the resulting mixture was further stirred for 12 h. The final product Ru/SP was centrifuged, washed with de-ionized water until the solution pH reached ~7 and dried at 80 °C in a vacuum oven for 12 h. For the preparation of MnO₂/Ru/SP, the as-synthesized Ru/SP and MnO₂ nanoparticles with a weight ratio of Ru/MnO₂/SP = 7:5:7.5 were sonicated into an ethanol aqueous solution for 3 h and stirred overnight. Then, the product MnO₂/Ru/SP was collected and dried at ~80 °C under vacuum for 12 h.

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