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A 98.2% energy efficiency Li-O₂ battery using a LaNi₀.₅Co₀.₅O₃ perovskite cathode with extremely fast oxygen reduction and evolution kinetics

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ABSTRACT

Rechargeable lithium-oxygen (Li-O₂) batteries have been regarded as a promising energy storage device, but its practical use is impeded by its low energy efficiency. Herein, a bi-functional catalytic perovskite LaNi₀.₅Co₀.₅O₃ (LNCO) is employed as the cathode of an efficient Li-O₂ battery with a molten nitrate salt electrolyte at 160 °C. It displays a stable low charge–discharge overpotential 50 mV with a high energy efficiency (EE) 98.2 % at 0.1 mA cm⁻² for over 100 cycles. The excellent performance is attributed to the extremely fast oxygen reduction and evolution kinetics on the surface of LNCO. The discharge product is Li₂O with a porous and fluffy morphology which facilitates the transfer of oxygen and other intermediate species. It is noted that Li₂O as a discharge product enables a theoretical specific energy density of 5200 Wh kg⁻¹, which is superior to the Li₂O₂ as product giving 3500 Wh kg⁻¹ for those ambient temperature Li-O₂ batteries.

1. Introduction

Rechargeable lithium-oxygen (Li-O₂) batteries are highly promising energy storage devices with a theoretical specific energy up to 3500 Wh kg⁻¹, with Li₂O₂ as the discharge product of the mostly reported ambient temperature testing cells, which far exceeds that of the state-of-the-art lithium-ion batteries (LIBs) of around 380 Wh kg⁻¹ [1–3]. The mostly explored Li-O₂ battery consists of a lithium anode, a carbon-based cathode, an electrolyte composed of lithium salt and organic solvent, and works at close to ambient temperature. The cathode reaction is given in Eq. 1, where Li₂O₂ is often formed during discharge and decomposed during the charge step [2,4],

\[ 2\text{Li}^+ + 2e^- + \text{O}_2 \rightarrow \text{Li}_2\text{O}_2 \]  (1)

Since the first proof-of-concept Li-O₂ battery, great effort has been devoted to optimizing the battery components, including the electrolyte, cathode substrate and catalyst [5–85–7]. However, most reported Li-O₂ batteries presents a high charge–discharge overpotential and low energy efficiency (EE) [9–11]. The underneath reason is the sluggish reaction kinetics at the cathode side, where oxygen reduction reaction (ORR) and oxygen evolution reaction (OER) take place during the discharge and charge reactions, respectively. The lack of efficient catalysts hinders the cathode reaction and thus leads to undesired electrochemical performance [12,13]. In recent years, perovskite materials with a general formula of ABO₃ have been explored as ORR and OER catalysts due to their high electronic conductivity, oxygen affinity and catalytic activity towards oxygen species [14–16]. Kim et al [17] incorporated LaCo₀.₉Fe₀.₁O₃ (LCFO) nanowires with reduced graphene oxide (rGO) sheets and utilized the composite as the cathode of a Li-O₂ battery. The battery with LCFO@rGO cathode displayed a charge–discharge overpotential 0.98 V. Hou et al [18] prepared a LaFe₀.₉LaFe₀.₁O₃ (LFCO) composite with carbon paper and used it as the Li-O₂ battery cathode. They reported a charge–discharge overpotential of 1.29 V. Nevertheless, Li-O₂ battery has not achieved a comparable EE, still often lower than 70 %, to the state-of-the-art LIBs, around 98 % for commercial LIBs. In addition, the highly reactive intermediate species, such as singlet oxygen, superoxide anion (O₂⁻), lithium superoxide (LiO₂) and lithium peroxide (Li₂O₂) formed during discharge/charge reactions attack the carbon...
materials and the organic molecules in the cell causing irreversible changes and leading to a low cyclability [9,19,20].

Li-O2 battery with a eutectic mixed nitrate melt electrolyte operating at an elevated temperature (around 150 °C) has been investigated [21–24]. It emerges as a promising strategy for greatly improving the efficiency and stability. Xia et al. [24] incorporated a nickel nanoparticle-based cathode with molten nitrate as electrolyte and achieved a reversible Li-O2 battery giving a low overpotential 0.2 V at a current density 0.1 mA cm\(^{-2}\). They proposed a Li2O2-mediated ORR pathway with Li2O as the discharge product, as shown in Eq.2.

\[ 4\text{Li}^+ + 4\text{e}^- + \text{O}_2 \leftrightarrow 2\text{Li}_2\text{O} \] (2)

With such a premise, we are stimulated to develop a Li-O2 battery with a LaNi0.5Co0.5O 3 (LNCO) perovskite-based cathode without utilization of a carbon material. The LNCO serves as both a cathode matrix and bi-functional electrocatalyst, i.e., ORR and OER, in a molten nitrate electrolyte at 160 °C. The battery achieves a low charge–discharge overpotential 50 mV at a current density of 0.1 mA cm\(^{-2}\). A porous and fluffy Li2O is formed as the discharge product, facilitating the transfer of the reactants. In addition, density function theory (DFT) calculations demonstrate the electronic structure, band gap, total and partial density of states (DOS) of LNCO, supporting the excellent ORR and OER catalytic activity.

### 2. Experimental

#### 2.1. Preparation of the composite cathodes

LNCO was synthesized via a sol–gel method. The obtained powder was ground into fine powder and calcined at 1100 °C for 2 h. Subsequently, 1 g LNCO powder was mixed with 1 mL of LiNO3-KNO3 (both 99.999 %) water solution with a mole ratio of 42:58 and a total concentration of 0.25 g mL\(^{-1}\). 80 mg of the dried composite powder was sandwiched with two pieces of stainless steel (SS) meshes and was referred to as LNCO cathode. For the preparation of Super P carbon cathode, a sample from Hefei Kejing Materials Technology Co., ltd was used, 40 mg material was mixed with 40 μL LiNO3-KNO3 solution. The obtained composite powder was sandwiched with two pieces SS meshes and was used as SPC cathode. The procedures and methods are illustrated in Fig. 1.

#### 2.2. Cell assembling and galvanostatic cycling

The LNCO composite cathode and a lithium anode are sandwiched with a binary nitrate infiltrated glass fiber separator. In addition, a Li1.5Al0.5Ge1.5(PO4)3 (LAGP) is further inserted between the cathode and separator as previously reported [24]. The stacked components were sealed with a home-made Swagelok-type cell. Subsequently, pure oxygen (Research 5.0 grade, Linder AGA Co., ltd) was introduced into the cathode compartment and sealed at around 0.9–1.2 bar. The cell was then transferred to an oven and kept at 160 °C for 5 h at open circuit condition. The electrochemical performance was tested on a battery cycler (LANHE) at different current densities in a range of 0.1–2 mA cm\(^{-2}\) with a potential window of 2.6–3.5 V vs Li/Li\(^+\). For monitoring the battery pressure, a pressure transducer (Omega) was used.

The batteries after electrochemical measurement were transferred into the glove box again. The composite cathode was detached from the battery and rinsed with N-methylacetamide solvent (NMA, Sigma Aldrich) to remove the nitrate salts. The treated LNCO cathode was then dried at 60 °C under vacuum for 12 h to remove the solvent completely. The characterization technique and corresponding DFT calculations are described in Supporting information S1.

### 3. Results

#### 3.1. The battery performance

To examine the electrochemical performance of LNCO as the bifunctional catalytic cathode, a Li-O2 battery with LNCO as cathode in a binary LiNO3-KNO3 eutectic nitrate melt was tested at 160 °C (Fig. S2). Fig. 2A plots the Galvanic charge and discharge profiles at 0.1 mA cm\(^{-2}\) within a voltage window between 2.6 and 3.5 V. The battery displays a stable discharge plateau at 2.80 V with a capacity of 7.96 mAh cm\(^{-2}\), while the charge step exhibits a plateau at 2.85 V, corresponding to a charge–discharge overpotential 50 mV and a remarkable EE value 98.2 %. The coulombic efficiency (CE) of the charge/discharge cycle is estimated as 97.2 %, demonstrating a high reversibility. This performance sharply contrasts to the SPC based cathode showing a CE only 47.1 %.
Fig. 2. Electrochemical performance of Li-O2 batteries with LNCO cathode. (A) Full discharge and charge curves of the cell with LNCO cathode (red curves) and with SPC cathode (blue curves). (B) Discharge and charge curves of different cycles with 0.1 mA cm\textsuperscript{−2}. (C) The corresponding EE and CE. (D) Corresponding pressure change during cycles. (E) Discharge and charge curves of different cycles with 0.5 mA cm\textsuperscript{−2}. (F) The corresponding EE and CE. (G) Voltage at different current (rate performance). (H) A plot of EE against current density with data in recent literature. Here only Li-O2 batteries cycled over 50 times are collected, and the values were calculated with the discharge/charge plateaus of the last cycle when the overall number of cycles was less than 100, otherwise the 100th cycle value if longer. The detailed information of the data and references are listed in Table S1. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
the cycling test, i.e., Fig. 2B, with a limited-capacity of 0.5 mAh cm\(^{-2}\) at 0.1 mA cm\(^{-2}\), two distinct plateaus are observed at 3.0 V and 2.78 V in the initial discharge step, where the higher plateau contributes to a relatively small capacity of less than 0.05 mAh cm\(^{-2}\) and the lower plateau at 2.78 V maintains much longer contributing to a major part of the capacity. It is noted that the full discharge profile shown in Fig. 2A also includes a tiny plateau comparable to a small discharge capacity at 3.0 V. Fig. 2C gives the corresponding EE and CE of the battery during cycling, where both experience an initial stabilizing period and quickly reach a stable value over 98 % during the following cycles. The corresponding real-time pressure in the cathode chamber during cycling was recorded and plotted in Fig. 2D. Linear pressure decreases and increases during discharge and recharge are observed, respectively. The pressure changes in Fig. 2D during cycles indicate highly reversible and efficient consumption and generation of oxygen ($\Delta P_{\text{charge}} / \Delta P_{\text{discharge}} \approx 97.62 \%$ after 3rd cycles), which is consistent with the charge/discharge profiles given in Fig. 2B. The cell was cycled again at a current density of 0.5 mA cm\(^{-2}\) for another 100 cycles and a stable charge–discharge overpotential 130 mV was measured corresponding to an EE of 94.7 % and CE of 99.82 %, respectively (Fig. 2E and 2F).

3.2. Material structure and surface states

The as-synthesized LNCO have a spherical morphology and an average particle size of 825 nm (Fig. 3A and Fig. S3). Fig. 3B shows the XRD patterns of the as synthesized LNCO, and the diffraction peaks are well consistent with JCPDS card No. 98–015-0874 (space group of R\(\overline{3}\) C, \(a = 5.6069 \, \text{Å}, \, b = 5.6069 \, \text{Å} \) and \(c = 13.4034 \, \text{Å}\)). The simulation with Material Studio confirms the perovskite structure of LNCO with the same symmetry. XPS spectrum indicates the presence of La, Ni, Co and O on the surface of LNCO (Fig. S5). Detailed signals have shown a partial overlap of the La 3d and Ni 2p spectra, where Ni 2P\(_{3/2}\) and Ni 2P\(_{1/2}\) can be assigned to the two peaks at 855.8 eV and 873.6 eV, respectively (Fig. 3C). These two peaks are further deconvoluted to show a bimodal feature, indicating the existence of Ni\(^{3+}/\)Ni\(^{2+}\) species. Similarly, the existence of Co\(^{3+}/\)Co\(^{2+}\) species are shown in Fig. 3D. The deconvoluted peaks correspond to the binding energies of Co 2P\(_{3/2}\) at 779.6 eV and 781.3 eV, and Co 2P\(_{1/2}\) at 794.9 eV and 796.8 eV. By using four-probe technique, the electric conductivity of LNCO is measured as 1.6 \times 10^3 S cm\(^{-1}\), which is consistent to reference [34].

3.3. Post-mortem analysis

The cathode after once discharge to 2.6 V gave rise to two extra XRD peaks at 33.65° and 56.41° with weak intensities (Fig. 4A), which are assigned to the (111) and (022) reflections of Li\(_2\)O (JCPDS No. 98–002-2402) and is in consistency with the Li 1 s XPS spectrum where the peak
of the binding energy at 53.6 eV demonstrates the existence of Li₂O (Fig. 4B). For a straightforward verification of the stoichiometries at the electrochemical steps, we utilized UV–vis spectroscopy and chemical titration for quantifying the oxygen containing species at different stages over the cycling (detailed methods are described in Supporting information S6) [24]. The cathode after a limited-capacity discharge to 2 mAh cm⁻² has 42.10 μmol Li₂O with no Li₂O₂, as shown in Fig. 4C and Fig. S6. The quantified Li₂O amount is nearly identical to the theoretically calculated value of 42.15 μmol, assuming the reaction happens as Eq. 3.

\[
2\text{Li}³⁺ + \frac{1}{2}\text{O}_₂ + 2\text{e}⁻ \rightarrow \text{Li}_₂\text{O}
\]  

(3)

For the recharged LNCO cathode, we find a small amount of Li₂O₂, 4.58 μmol, remains on the cathode surface, which is in consistency with the pressure monitoring result for the full discharge/charge cycle. In addition, the cathode after 100 cycles at recharged state contains a comparable amount of Li₂O 3.02 μmol. The SEM image in Fig. 4D shows the cathode after discharge, where porous and fluffy discharge product covers the LNCO particle surface. Elemental mapping confirms the existence of O element in the discharge product, which further demonstrates these domains corresponding to the Li₂O formed during the discharge process (Fig. 4E). Fig. 4F show the SEM images of the charged cathode after 100 cycles, where a complete removal of the porous and fluffy Li₂O is confirmed with showing an identical morphology as in the pristine LNCO.

3.4. Density functional theory verification

As shown in Fig. 5A, partial DOS profiles of 5d orbitals of La atoms, 3d orbitals of Ni and Co atoms, as well as 2p orbitals of O atoms contribute to the total DOS of LNCO, where a semi-metallic behavior is observed from the DOS distribution around the Fermi level (E₀) along both the spin-up and spin-down directions. Furthermore, the band energy profile shows a small band gap of 0.09 eV (Fig. 5B). In addition, the DOS of Ni, Co and O show an overlap in the energy range of −7 eV to 0 eV, suggesting a strong hybridization among the orbitals of O 2p and Ni and Co 3d [14,25,26].

The Gibbs free energy diagrams of both the ORR and OER steps are calculated to reveal the reaction pathway. The electrode reaction pathway is supposed as (1) Li + O₂ → Li₂O₂, (2) 2Li + Li₂O₂ → 2Li₂O. For simulation, we selected (110) and (100) facets of the LNCO surface, where they display adsorption strength towards the Li₂O₂ intermediate with Δ E_ads of −3.86 eV and −8.35 eV, respectively. As illustrated in Fig. 5C and 5D, at U = 0 V, the energy difference between every step relates to the binding energies between the intermediate species and the substrate facet. The calculated U₀ is 2.86 V, which is in consistency with the thermodynamic potential, 2.84 V at 160 °C. Here, the discharge and charge overpotentials are defined as η_{ORR} = U₀ − U_d and η_{OER} = U_c − U_d, respectively. The diagram presents the energy paths of the ORR/OER at different overpotentials over (1 1 0) facet of LNCO.

4. Discussion

4.1. Cathode material LNCO

The SEM and XRD results show that the LNCO phase was successfully synthesized with high purity and crystallinity (Fig. 3A and 3B). From the XPS spectrum, it is impossible to quantify the surface Ni species because of the overlap between Ni 2p and La 3d peaks in the same range. However, the XPS spectra indicates the simultaneous presence of Ni⁡/Ni²⁺ and Co³⁺/Co²⁺ redox couples on the surface (Fig. 3C and 3D). It has been reported that the simultaneous existence of Co³⁺ and Co²⁺ improves the Co-O bond strength and promote the dissociative
adsorption of oxygen on the LNCO surface, which greatly promotes the ORR catalytic activity. Furthermore, the co-existence of Ni\(^{3+}/\text{Ni}^{2+}\) and Co\(^{3+}/\text{Co}^{2+}\) redox couples on the LNCO surface exhibits a synergistic effect and promotes both the ORR and OER reactions [16].

4.2. Electrochemical performance

The LNCO cathode-based Li-O\(_2\) battery operating at 160 °C exhibits excellent electrochemical performance. The battery displays with an ultra-low charge–discharge overpotential 50 mV, the corresponding remarkable EE and CE also indicate a high reversibility of the electrode reactions (Fig. 2A). As shown in Fig. 2B, the higher plateau at 3.0 V gradually diminished and the lower plateau at 2.78 V further extended while retaining the voltage during cycling. For the charge step, the initial few charge curves were also accompanied with a two-plateau feature, with a higher plateau at 3.1 V that gradually diminished after 10 cycles. Meanwhile, the lower plateau positioned at 2.83 V, contributing to the major charge capacity and accordingly prolonged as the battery cycled. The charge–discharge overpotential is calculated based on the major charge–discharge plateaus. This Li-O\(_2\) battery also demonstrates a remarkable stability with showing an unchanged charge–discharge overpotential 50 mV over 100 cycles. According to the corresponding real-time pressure in the cell as plotted in Fig. 2D, the consumed oxygen is determined as 5.46 × 10\(^{-6}\) mol per discharge cycle. Assuming 4e\(^{-}\) transfer per mol of oxygen, the total amount of electron transfer of one discharge is 1.31 × 10\(^{19}\). The discharge capacity was measured as 0.565 mAh per discharge, corresponding to about 2.03C of electricity and electron transfer of 1.17 × 10\(^{19}\). The eminent electrochemical performance of the LNCO cathode-based Li-O\(_2\) battery at elevated temperature indicates the extremely fast ORR and OER kinetics on the LNCO surface.

\[
\Delta \rho_{\text{discharge}} = nRT
\]

Compared with other Li-O\(_2\) batteries applied with other cathode materials, the Li-O\(_2\) cell with LNCO cathode shows relatively low overpotential and thus exhibits high EE, as shown in Fig. 2H. It is attributed to the following reasons: (i) The elevated temperature boosts the reaction kinetics. (ii) The high ionic conductivity of the molten nitrate allows fast Li\(^{+}\) ionic transfer. (iii) LNCO exhibits a high electrical conductivity. It guarantees sufficient electric transfer on the surface of cathode during discharge and charge, resulting in good rate performance [27]. (iv) LNCO cathode enhance the catalytic performance of ORR and OER, thus reduce the overpotential, especially on charge.

4.3. Discharge product analysis

In both XPS spectrum and XRD pattern (Fig. 4A and 4B) of LNCO cathode after discharge, there are no peaks of other lithium-species, such as Li\(_2\)O\(_2\), Li\(_2\)CO\(_3\), etc., suggesting that only Li\(_2\)O was formed as the discharge product. In addition, the peaks of Li\(_2\)O in XRD disappear completely after recharge and the pattern of LNCO remains the same, indicating the total decomposition of discharge product Li\(_2\)O and the stability of the LNCO. For the quantification analysis (Fig. 4C), we
believe that the remaining Li2O detected after charging step is due to the partial dissolution of Li2O into the nitrate electrolyte, and the saturation is achieved during the 1st cycle [28]. The subsequent cycles arise minimal effect of this dissolved amount, which is in line with the CE change in the initial few cycles. The above results suggest a highly reversible formation and decomposition of Li2O during discharge and charge steps. It is noted that such a Li2O with porous and fluffy feature is for the first time discovered in the Li2O batteries with molten nitrate system, which differs from previous reports where Li2O with well-defined octahedral shapes deposited on the metal nanoparticles under similar operation conditions (Fig. 4D) [24,28]. The unique structure and morphology of the Li2O in this work favors the high catalytic activity of the LNCO surface. Previous works on sputtered Li2O batteries reported that thin film like Li2O discharge product with inferior crystallinity tend to decompose at a lower charge potential than the highly crystallized Li2O [29]. Herein, the porous and fluffy Li2O likely provides abundant pathways for breathing O2 in and out during the discharge and charge steps and facilitates the transfer of other intermediates [30]. These effects enabled the fast reaction kinetics of this battery with much higher EE and cyclability than those reported in [21,24,31,32]. Fig. 4F shows the SEM image of the cathode after recharge and a complete removal of the porous and fluffy Li2O showing an identical morphology as in the pristine LNCO. It has been known that Li2O as the discharge product results in a 5200 Wh kg$^{-1}$ specific energy density, much higher than the cell with Li2O2 as the discharge product, i.e., 3500 Wh kg$^{-1}$ [24].

4.4. Catalytic activity analysis

As shown in Fig. 5A, the calculated total DOS are mainly formed by O 2p, Ni 3d and Co 3d orbitals from -6 eV to 4 eV while La 5d mainly contributes to the peak above 4 eV. The partial DOS of Ni, Co and O show an overlap in the energy range of ~7 eV to 0 eV, suggesting a strong hybridization among the orbitals of O 2p and Ni and Co 3d. In contrast, under the Fermi level, only a small amounts of hybridization exchange between La 5d and O 2p orbitals. Therefore, such a hybridization indicates the interaction force between Ni or Co and O is higher than that between La and O [33,34]. It is closely related to the structural stability of LNCO, indicates from the superior cyclability of the Li2O2 batteries with LNCO cathode shown as above. The results above suggest the key chemical properties of LNCO are affected by Ni and Co, instead of La. Meanwhile, it also indicates that the low spin state and high spin state co-exist for Ni ion and Co ion in LNCO from partial DOS. Ni3$^{+}$ with an electron configuration of $\text{t}_{2g}^{6}$$\text{e}_{g}^{2}$ at low spin and high spin states, respectively [35]. It owns more unpaired electrons in 3d orbital compared with Ni$^{2+}$ with an electron configuration of $\text{t}_{2g}^{5}$$\text{e}_{g}^{2}$. Similarly, Co$^{3+}$ owns more unpaired electrons in 3d orbital compared with Co$^{2+}$ [36]. The existence of the unpaired electron in 3d orbitals of Ni$^{2+}$ and Co$^{3+}$ positively affects the electric conductivity [35]. As a result, the Ni$^{2+}$/Ni$^{3+}$ and Co$^{2+}$/Co$^{3+}$ redox couples can also promote charge transfer between surface cation and adsorbates, which is consistent with the $e_{g}$ occupancy theory [37]. Furthermore, the orbital hybridization is accompanied with the d orbital splitting, where $\sigma$-orbital is primarily correlated to the ORR activity. Previous works reported that perovskite LaNiO3 with an $e_{g}$ occupancy of 1.2 had the highest ORR catalytic activity among the perovskites with different B metal species [31]. Since our LNCO is structurally comparable to LaNiO3, a favorable $e_{g}$ occupancy seems to have also been achieved in LNCO, thus leading to the excellent ORR activity. Furthermore, the existence of Co with a Co/Ni ratio of 1 in LNCO has realized a maximum covalency of B site metal–oxygen (B-O) bonds, which favors a kinetically fast OER [24,28-40]. Overall, LNCO with the above-mentioned electronic structure is suitable for an efficient bi-functional catalyst for both ORR and OER steps, leading to an exceptionally low overpotential and high EE for the Li2O2 battery.

The Gibbs free energy diagrams of both the ORR and OER steps are simulated, and we selected (1 1 0) and (1 0 0) facets of the LNCO surface. The weak binding energy between LNCO and Li2O2 on LNCO (1 1 0) facet favors the OER (Fig. 5G) [41]. Compared to the energy path profile of the (1 0 0) facet given in Fig. 5D, the (1 1 0) facet clearly shows a much smaller overpotential between the ORR and OER steps, suggesting a stronger likelihood of the occurrence of ORR and OER on the (1 1 0) facet than on (1 0 0). For the OER process, O2 first accepts electrons and reacts with Li$^{+}$ to generate LiO2 (Li$^{+}$ + O2 + 2e$^{-} = $LiO2). Due to the weak adsorption strength between LiO2 and (1 1 0) face on LNCO, the LiO2 further accept electrons and reacts with Li$^{+}$ to generate the discharge product Li2O2 (2Li$^{+}$ + LiO2 + 2e$^{-} = $2Li2O). During the charge process, the oxidation of Li2O2 happens via two steps: (1) the oxidation of Li2O to form LiO2 and Li$^{+}$; (2) the further oxidation of LiO2 to O2 and Li$^{+}$. DFT calculations further imply that during the charge, the rate limiting step is the oxidation of Li2O2 to form O2 and Li$^{+}$, and the weak adsorption interaction between Li2O and (1 1 0) facet of LNCO leads to a low charge overpotential.

4.5. The mechanism

Based on both the experimental and simulation results, we propose an ORR mechanism on the LNCO cathode surface as in Fig. 6. This initiates as oxygen molecule (O$^{2−}$ad) diffuses and adsorbs onto LNCO surface (Eq. 4), where it is reduced to Li2O2 ad with the incorporation of Li$^{+}$ from the electrolyte (Eq. 5). This Li2O2ad is further reduced with a consecutive electron acceptance from LNCO, forming the surface adsorbed LiO2ad (Eq. 6). Due to the solubility of Li2O in molten nitrates, the LiO2ad tends to be dissolved, as described in Eq. 7, and indicated in Fig. 6 with dotted arrows. This pathway is less energy-intensive and accounts for the upper-plateau in the discharge curves (Fig. 2A, 2B and 2E). With the gradual saturation of the O$^{2−}$ species close to the LNCO surface, the deposition of solid Li2O onto LNCO is triggered, which corresponds to the lower discharge plateau. The in-situ formed Li2O on the LNCO surface is supposed to moderately bonded onto the LNCO, so that a free growth of Li2O is prohibited. As a result, porous and fluffy Li2O rather than the previously reported highly crystalline Li2O octahedra crystallites are formed during ORR [24]. The dissolved Li2O gradually saturates in the entire molten salt phase during the cycling, which eventually leads to the diminish of the upper discharge plateau, as shown in Fig. 2B and 2F. However, with the gradual diffusion and distribution of the O$^{2−}$ species in the molten nitrate, a part of dissolved Li2O is hard to be recharged. It is the reason why the CE is less than 100 % in Fig. 2C and 2F, especially at the first three cycles.

\[
\begin{align*}
O_2 + e^- & \rightarrow O^{2-}_{ad} & (4) \\
2O^{2-}_{ad} + 2Li^{+} + 2e^- & \rightarrow LiO_2^{ad} & (5) \\
2LiO_2^{ad} + 2Li^{+} + 2e^- & \rightarrow 2Li_2O_{ad} & (6) \\
Li_2O_{ad} & \rightarrow 2Li^{+} + O_2^{−} & (7)
\end{align*}
\]

5. Conclusions

In summary, we adopted a perovskite LaNiO3$\delta$,CoO3$\delta$ (LNCO) as a bi-functional catalyst for ORR/OER and as the cathode substrate for a Li2O2 battery with a molten nitrate electrolyte at an elevated temperature 160 °C. The battery showed an ultra-low charge–discharge overpotential 50 mV with a record-high energy efficiency (EE) 98.2 % and remained a remarkable cyclability over 100 cycles with unchanged overpotential and EE values at a current density 0.1 mA cm$^{-2}$. Quantitative analyses confirm the stoichiometric formation of Li2O2 as the only discharge product via a 4e$^{-}$ transfer pathway. Computational simulations indicate that the superior performance of LNCO is related to its semi-metallic characteristics, as well as the co-existence of surface Ni$^{2+}$/Ni$^{3+}$ and Co$^{2+}$/Co$^{3+}$ redox couples that efficiently catalyze both ORR and OER steps. As a result, we have for the first time discovered the formation of a
porous and fluffy Li$_2$O phase as the discharge product on the LNCO perovskite oxide cathode, which enhances the kinetics of the ORR and OER. A surface 4e– discharge reaction mechanism with two different follow-up steps is proposed. Our work indicates a great potential of the perovskite materials as cathode of the future rechargeable Li–O$_2$ battery. We envision a prosperous future of such energy storage systems operating at moderate temperatures in various applications.

CRediT authorship contribution statement

Qiyan Qiu: Conceptualization, Methodology, Investigation, Writing – original draft. Zheng-Ze Pan: Conceptualization, Funding acquisition, Methodology, Writing – review & editing. Penghui Yao: Formal analysis. Jiashu Yuan: Methodology. Chun Xia: Validation. Yicheng Zhao: Validation. Yongdan Li: Supervision, Funding acquisition, Writing – review & editing.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/jcej.2022.139608.

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