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Facet-engineered photoelectrochemical nanocatalysts toward fast kinetic lithium-air batteries†

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Despite lithium-air batteries having unprecedented theoretical energy supremacy, rambling Li₂O₂ growth typically impedes O2 diffusion and electron transfer, causing low capacity, poor charging kinetics and short cycle life. Herein, we propose synergistic catalysis in the cathode by engineering the crystal facet of Fe_2O_3 nanoplates and introducing a visible light-excited photochemical effect. The growth of Li₂O₂ is regulated with a uniform distribution, mitigating local product aggregation and thus reducing the barriers of O_2 diffusion and electron transfer. The improved reaction kinetics lead to \sim 420% capacity improvement, low charge potential, superior rate performance, and long-term stability (300 cycles) in actual air. We reveal a mechanism that photoelectrons around each catalyst node and the resultant homogeneous internal electric field increase charge transfer, decrease soluble intermediates, and enable high electrochemical reversibility of amorphous Li₂O₂ with an ordered process. This study provides inspiring insights for future research into controlling product ordering in Li-air batteries.

Broader context

Li-air batteries with ultra-high energy density promise the next-generation batteries for long-range electric vehicles. However, their actual energy density is far below the theoretical value because the diffusion transfer of the O2 reactant from air to the cathode interiors is largely hindered by the locally accumulated solid discharge products outside cathodes. Here, we demonstrate that introducing visible light energy is an effective strategy to enable batteries with largely enhanced discharge capability and cycling stability in actual air conditions. We reveal an intriguing mechanism that a photoelectron-induced internal microelectric field can change the product morphology and more importantly homogenize product growth within the cathodes, accelerating electron conduction and O2 diffusion. This simple but efficient approach shows great potential for advancing the practicality of Li-air batteries.

1. Introduction

Li-air batteries with an ultrahigh theoretical energy density $(\sim 3500 \text{ W h kg}^{-1})$ promise next-generation energy storage

solutions. Its energy superiority derives from the reactants that are O_2 gas in ambient air and the redox reaction of $O_2 \rightarrow$ Li₂O₂. During discharge, O₂ first transfers from ambient air to active sites in porous cathodes through diffusion.^{3,4} After that, O₂ receives one electron and is reduced to O₂ which can be adsorbed on the cathode surface followed by a further one-electron electrochemical reduction or dissolved in the electrolyte followed by a disproportionation reaction, leading to the formation of a Li₂O₂ film or particles determined by the current densities,^{5,6} electrolytes,^{7,8} cathode structures,^{9,10} and catalysts. 11-14 Nevertheless, premature failure happens when the discharge capacity is far from the high theoretical value of Li-air batteries, hence greatly plagueing their practicability. 15-17 The culprit of this problem is the uncontrolled deposition of Li₂O₂ that exhibits considerably low electronic conductivity (only 10⁻¹⁰-10⁻⁹ S cm⁻¹ at 100 °C)¹⁸ and insolubility $(4 \times 10^{-20} \text{ mol L}^{-1} \text{ in ether-based electrolytes}),^{19}$

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hindering the electron and oxygen transfer during the oxygen reduction reaction (ORR) and oxygen evolution reaction (OER) process. Undoubtfully, promoting fast mass (Li+, O2) and electron transfer processes is an indispensable route towards a high-energy-density Li-air battery with remarkable battery lifespan.20

To realize the high energy density of Li-air batteries, air cathodes need to achieve a high mass loading of active materials. and batteries are required to operate in ambient air rather than O₂ tanks.21 Compared with the large capacity of Li-O2 batteries (LOBs) in pure O₂ gas, Li-air batteries always show an apparent capacity decrease due to the slower O2 transfer in ambient air (~21% O₂ content). 22-24 Besides, more than 65% capacity loss was observed when the O₂ pressure decreased from 21 to 14 psi.²⁵ Notably, the O2 concentration and electronic conductivity are further subject to the limited diffusion space and high mass loading in a practical pouch cell configuration, leading to much more severe performance deterioration.²⁶ Such a vast gradient generated inside the thick electrode may correlate with the discharging voltage that reached the termination without fully utilizing the active sites of the air electrode. With the increase of the cell thickness, the energy density of the pouch cell decreased from 700 to <200 W h kg⁻¹. Moreover, the local nucleation and growth of Li2O2 particles together with byproducts (Li₂CO₃, etc.) and their inhomogeneous accumulation not only clog the oxygen-transport pores^{27,28} but also cause cathode

surface passivation.^{29,30} These local depositions mean that O₂ and e cannot transport deep inside the thick air cathodes, leaving plenty of unavailable "dead" active sites and leading to premature battery failure (Fig. 1b), which seriously compromises the primary superiority (high energy density) of Li-air batteries. In addition, the particle Li₂O₂ products are usually more difficult to decompose, leading to excessive charging polarization and further impairing the energy efficiency and long-term cycling stability of the batteries.³¹

In this work, we initiated a photoelectrochemical effect to guide Li₂O₂ nucleation, growth and reversible decomposition in air cathodes containing the Fe₂O₃ catalyst with high numbers of (110) facets (referred to as (110) Fe₂O₃) (Fig. 1a). The (110) facet of Fe₂O₃ possess superior visible light response and oxygen adsorption effect, enabling uniform distribution of photoelectrons, and promoting fast charge transport inside the electrode. In addition, the enhanced adsorption of oxygen species leads to a homogeneous nucleation and growth process of the Li₂O₂ products, avoiding local pore-clogging and premature battery failure. This accelerated O2 and charge transfer effect not only facilitates the enhancement of the discharge capacity but also promotes product decomposition during the charge process, especially in the air. (110) Fe₂O₃-based electrodes delivered a significantly increased capacity (~260% in O_2 and more promisingly $\sim 420\%$ in air) and extended cycle life (over 300 cycles) in the presence of visible light. The visible

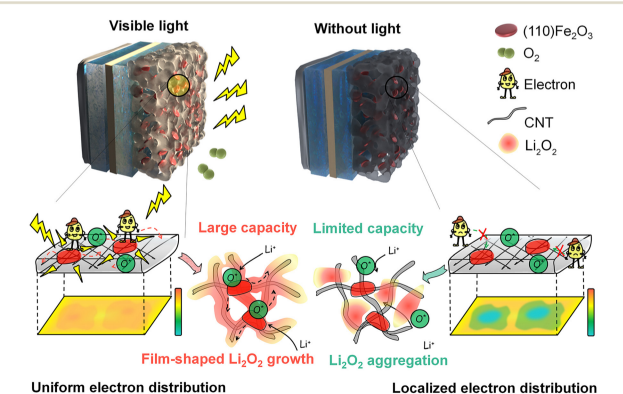


Fig. 1 Schematic of visible-light-induced photoelectron-rich microdomains and uniform Li₂O₂ deposition in the (110) Fe₂O₃ catalyst-based cathodes. (a) Instantaneously generated photoelectrons on each catalyst particle junction and improved catalysis of Fe₂O₃ with light modify Li₂O₂ formation, and the uniform deposition gives large discharge capacities. (b) The gradient distribution of electrons from electrode tabs to active sites and the unfavorable catalysis of Fe_2O_3 without light result in the local accumulation of large Li_2O_2 particles, pore-clogging, oxygen transfer obstruction and limited capacities.

light-regulated Li₂O₂ formation provides an innovative but effective way to improve the energy density, power density and cycling stability of Li-air batteries for practical use.

2. Results and discussion

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2.1. Material preparation and characterization

(110) Fe₂O₃ particles were synthesized by a cation-assisted template method, and the crystal phase was α-Fe₂O₃ based on the X-ray diffraction (XRD) patterns (Fig. 2a). The crystals are nanoplates with an average diameter of about 80 nm (Fig. 2b). In contrast, the shapes of referenced pristine Fe₂O₃ (pFe₂O₃) particles are irregular (Fig. S1, ESI†). The ratio of integrated XRD peak intensities $I_{(110)}/I_{(104)}$ increased from 0.77 for pFe₂O₃ to 1.24 for (110) Fe₂O₃. The transmission electron microscope (TEM) images in Fig. S2 (ESI†) show that the thickness of the nanoplates is less than 10 nm, and the high-resolution TEM (HRTEM, Fig. 2c) image of the atomic planes shows lattice

fringes with a spacing of 0.25 nm, corresponding to the (110) planes of Fe₂O₃. These results confirm the synthesis of (110) Fe₂O₃.

The effect of crystal facets on the photocatalytic activity of Fe₂O₃ was shown by ultraviolet-visible (UV-vis), X-ray photoelectron (XPS) and photoluminescence (PL) spectra. The UV-vis results (Fig. 2d) indicate that (110) Fe₂O₃ has a blue shift in the absorption edge compared to pFe₂O₃. The fitted Tauc plot (Fig. 2e) shows an increase in the bandgap from 2.00 eV of pFe₂O₃ to 2.08 eV of (110) Fe₂O₃. The valence band (VB) positions of (110) Fe₂O₃ and pFe₂O₃ analyzed by XPS valence band spectroscopy are at 1.96 eV and 2.80 eV, respectively (Fig. 2f). Accordingly, the relative band positions of (110) Fe₂O₃ and pFe₂O₃ were determined and shown in Fig. S3 (ESI†). The faceting lowers the conduction band (CB) position of Fe₂O₃ by 0.92 eV, making it below the redox potential of O₂/Li₂O₂. Thus the generated photoelectrons on (110) Fe₂O₃ can be transferred spontaneously and react with Li⁺ and O₂ in the discharge process.³² In addition, the PL spectra in Fig. 2g show

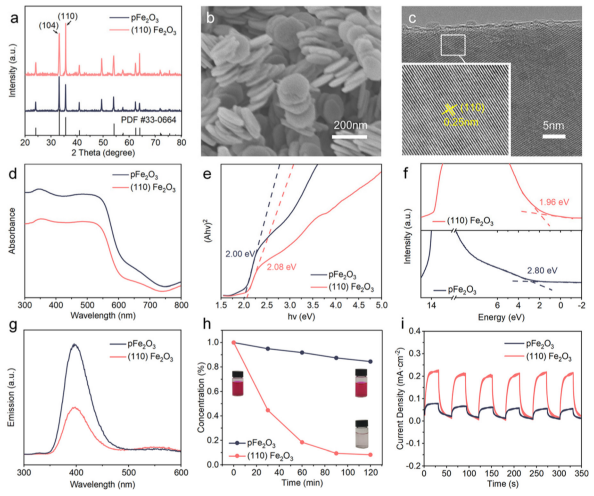


Fig. 2 Physicochemical and photoelectrochemical properties of faceted Fe₂O₃. (a) XRD patterns of (110) Fe₂O₃, pFe₂O₃, and PDF#33-0664 indexed to α -Fe₂O₃. (b) SEM image of the (110) Fe₂O₃ morphology. (c) HRTEM image showing the lattice fringes of the α -Fe₂O₃ particles assigned to the (110) facet. (d) UV-Vis absorption spectra of (110) Fe₂O₃ and pFe₂O₃. (e) Tauc plot based on the UV-vis absorbance data. The intercept of the tangent to the linear part of each curve and the X-axis represents the bandgap. (f) XPS valence band spectra of (110) Fe₂O₃. (g) PL spectra of (110) Fe₂O₃ and pFe₂O₅. (h) Photocatalytic degradation of RhB. (i) I-t curves at 0.898 V (vs. SHE).

that the luminescence intensity of (110) Fe₂O₃ is apparently lower than that of pFe₂O₃, indicating that the lifetime of photogenerated electrons is prolonged. This phenomenon is favorable to the photocatalytic performance, as shown by experimental observation of the photo-Fenton reaction degradation of rhodamine (RhB) on both materials (Fig. 2h). The degradation efficiency on (110) Fe₂O₃ (91.8%) is much higher than that on pFe₂O₃ (15.6%) after 2 h of light exposure. The photoelectrochemical properties of the samples were also investigated using amperometric I-t tests (Fig. 2i). The photocurrent density of (110) Fe₂O₃ is 0.22 mA cm⁻², much higher than that of pFe₂O₃ (0.08 mA cm⁻²). This enhanced photocatalytic performance confirms the effect of faceting on the improvement of the visible-light response.

2.2. Discharge capability in Li-O₂ batteries and actual Li-air batteries

To demonstrate the effect of light and catalysts towards the Li-oxygen-based reactions, LOBs were assembled with different cathodes, namely (110) Fe₂O₃-loaded CNTs, pFe₂O₃-loaded CNTs or unloaded CNT-based cathodes. The LiTFSI/tetraglyme based electrolytes and Li metal anodes were coupled with these cathodes. Cyclic voltammetry (CV) was performed without (Fig. 3a) and with visible light (Fig. 3b) to clarify the reactions during discharge and charge. The electrocatalytic activities of the ORR and OER are enhanced. In the discharging process, compared to pFe₂O₃ which shows a reduction peak at 2.15 V, the onset potential of the reduction reaction in the (110) Fe₂O₃ battery was increased to 2.77 V and two reduction peaks at 2.52 V and 2.34 V could be distinctly found, which may indicate the two-step electron ORR process. In the charging process, two oxidation peaks at 3.20 and 3.39 V appeared in the presence of (110) Fe₂O₃, while pFe₂O₃-based batteries showed no obvious oxidation occurring under 4.0 V. With the presence of light (Fig. 3b), due to the lowered band position of (110) Fe₂O₃, the photoelectrons could be involved in the Li-O2 reactions to facilitate their kinetics, especially for the reduction process. The peak current density reaches 0.46 mA cm⁻² in (110) Fe₂O₃based batteries, while it is only 0.20 mA cm⁻² in pFe₂O₃-based ones. The Tafel slopes for the (110) Fe₂O₃ and pFe₂O₃ cases under light (Fig. S5, ESI†) were 146 mV dec⁻¹ and 194 mV dec⁻¹, respectively, which indicates that faceted (110) Fe₂O₃ has a better electrocatalytic performance as well as a significant photogenerated current that promotes the ORR process and enables an improved discharge ability and rate performance, especially with light illumination.

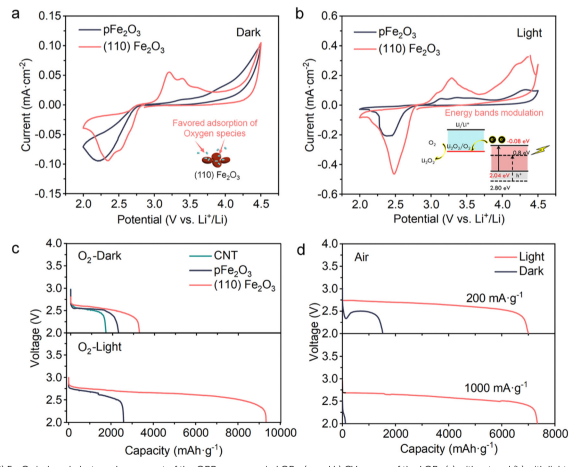


Fig. 3 (110) Fe₂O₃ induced photo-enhancement of the ORR processes in LOBs. (a and b) CV curves of the LOBs (a) without and (b) with light at 0.2 mV s⁻¹; the inset schematics illustrate the electrocatalytic and photocatalytic effects for Li-O2-based reactions. (c) Galvanostatic discharge profiles of (110) Fe₂O₃, pFe₂O₃, and CNT samples at 200 mA g^{-1} in O₂. (d) Galvanostatic discharge profiles of (110) Fe₂O₃ samples at 200 mA g^{-1} and 1000 mA g^{-1} in air.

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In a pure oxygen environment, the (110) Fe₂O₃-based cathodes exhibited an increased specific capacity of 3276 mA h g⁻¹ without light (Fig. 3c). In comparison, the unloaded CNT and pFe₂O₃ cases were 1678 mA h g⁻¹ and 2275 mA h g⁻¹, respectively. An improvement is further observed in the presence of light. After the addition of light, the discharge specific capacity of the (110) Fe₂O₃-based cathodes was unexpectedly increased to 9329 mA h g⁻¹, in sharp comparison to only 2596 mA h g⁻¹ of the pFe₂O₃ case. Considering that both materials have similar specific surface areas, 183.84 m² g⁻¹ for (110) Fe_2O_3/CNT and 183.42 m² g⁻¹ for pFe₂O₃/CNT (Fig. S6 and S7, ESI†), this apparent increase is attributed to the capacity provided by (110) Fe₂O₃ under light. Notably, the difference in capacity was more significant when the batteries were operated in the ambient air, as illustrated in Fig. 3d. Compared to the small capacity (1670 mA h g⁻¹ at 200 mA g⁻¹) in the dark conditions, the light-assisted discharge process exhibits reduced polarization (~0.25 V) and much increased capacity (over 7000 mA h g⁻¹), which retained no degradation of capacity even under high current densities of 1000 mA g⁻¹. Overall, the (110) Fe₂O₃-based electrode produced a $\sim 420\%$ capacity increase even in the air, indicating the excellence of the photochemical effect on (110) Fe₂O₃-based cathodes.

2.3. Light-induced surface deposition of discharge products

Changes in the composition and morphology of the discharge products on the (110) Fe₂O₃-based cathodes in dark/light conditions were investigated by XRD, SEM and XPS. The XRD pattern in Fig. 4a shows a characteristic peak at 34.8° corresponding to the (101) plane of crystalline Li₂O₂ generated by discharge in dark conditions, while the products are amorphous when the discharge process is performed in the presence of light, which is indicated by the absence of characteristics peaks of Li₂O₂. These amorphous products were verified to be mainly Li₂O₂ by Raman (Fig. S8, ESI†) and XPS spectra (Fig. 4f and g). The SEM images (Fig. 4b and c) indicate the differences in Li₂O₂ morphology. In dark conditions, micron-sized Li₂O₂ spherical particles are randomly deposited on the cathode, while film-like Li₂O₂ is formed consistently on the surface of CNTs and Fe₂O₃ nanorods. SEM images captured at different discharge stages were applied to further clarify the photo-organized Li₂O₂ formation process (Fig. 4d). At the discharge depths of 500, 1500, 3000 and 5000 mA h g⁻¹ and full discharge at 2.0 V, Li₂O₂ first nucleates on the nanoplate-shaped (110) Fe₂O₃ surface, then grows along the electrode surface, and finally covers the electrode surface uniformly.

As a rule, the reason for the Li₂O₂ deposition difference can be attributed to the existence of discharge intermediates, LiO₂,

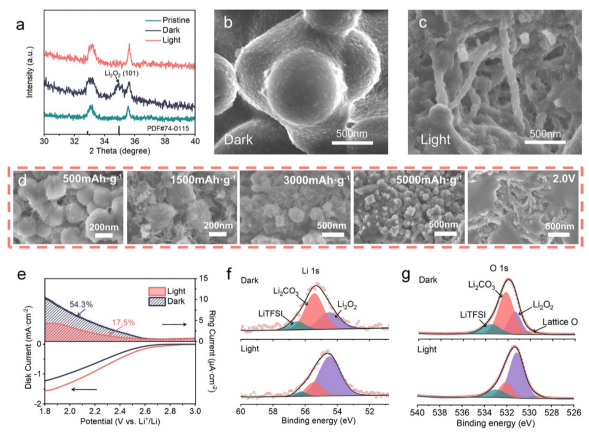


Fig. 4 Surface nucleation and growth process of products under visible light. (a) XRD pattern of the (110) Fe₂O₃ based cathodes after discharge. (b and c) Morphologies of the fully discharged (110) Fe₂O₃-based cathodes (b) without and (c) with light. (d) SEM images at different discharge depths at 200 mA g⁻¹. (e) RRDE measurements at 800 rpm. The disc potential increases from open circuit potential to 1.8 V vs. Li⁺/Li. E_{ring} was held at 3.5 V vs. Li+/Li. (f, g) XPS spectra of (f) Li 1s and (g) O 1s for the discharged cathode in dark/light conditions.

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whether formed in electrolyte (LiO_{2,sol}) or adsorbed on the cathode (LiO_{2,ads}).33-35 And the relative content of LiO_{2,sol} can be monitored by the rotating ring-disk electrode (RRDE) test^{34,36,37} (Fig. 4e and Fig. S9, ESI†). The disc electrode was loaded with (110) Fe₂O₃ and pFe₂O₃, respectively, and the ring electrode potential was set at 3.5 V (vs. Li⁺/Li) to detect the soluble intermediates. The disc and ring currents were monitored during a negative sweep from the equilibrium potential. The linear sweep voltammetry (LSV) curves obtained from the disk currents show an increase of the onset potential (2.85 V vs. 2.69 V) and a larger current density (1.56 mA cm⁻² vs. 0.98 mA cm⁻²) in the presence of light, showing that the illumination on (110) Fe₂O₃ greatly accelerates the ORR process. It is worth noting that the ring current decreased from 10.8 to 4.2 µA cm⁻² in the presence of light, and the weight of solution-phase reaction $(Q_{\rm ring}/Q_{\rm ORR})$ is revealed by this value. The $Q_{\rm ring}/Q_{\rm ORR}$ of the (110) Fe₂O₃/CNT cathode decreased from 54.3% in the dark to 17.5% in light conditions, indicating that the production of soluble intermediates was suppressed. This phenomenon was analyzed by Raman spectroscopy (Fig. S8, ESI†) which is generally conducted to detect soluble LiO₂ intermediates.³⁸ In dark

conditions, the discharged cathode showed obvious LiO₂ peaks at 1135 and 1480 cm⁻¹, while these peaks almost disappeared with light illumination. These experiment results suggest that the effect of light on (110) Fe₂O₃ decreases the amount of solvated LiO2 intermediates in the electrolyte and favors the ORR process on the electrode rather than in the electrolyte. This transformation has a positive impact toward the suppression of side products like Li₂CO₃, which may be attributed to less ¹O₂ generated during the disproportionation reactions, and they would further react with the carbon substrate at the cathode. 39-41 Changes in the compositions of the discharge products on (110) Fe₂O₃-based cathodes in dark/light conditions were investigated by XPS (Fig. 4f, g and Fig. S10, ESI†). Li₂O₂ and Li₂CO₃ species were observed on the (110) Fe₂O₃ based cathode, both in dark conditions and in light conditions, while the relative content of Li₂CO₃ decreased considerably after light discharge.

2.4. Boosted electron and oxygen transfer by the (110) Fe₂O₃ catalyst

The intrinsic reason for light-induced surface deposition of Li₂O₂ is revealed through the theoretical calculation and

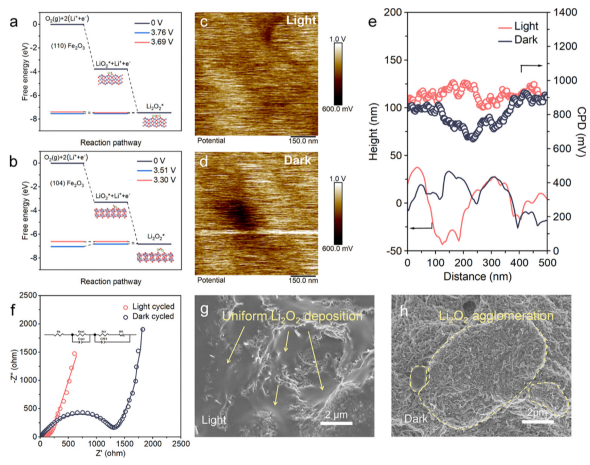


Fig. 5 Enhanced adsorption of oxygen intermediates and construction of a photoelectric field by (110) Fe₂O₃ to regulate Li₂O₂ deposition. (a and b) Calculated free energy diagrams for the ORR reactions on (a) (110) and (b) (104) facets. (c and d) CPD maps of the (110) Fe₂O₃ based cathode in (c) light and (d) dark conditions. (e) CPD and height curves. (f) Nyquist plots and equivalent circuit of the (110) Fe₂O₃ electrode after ten cycles with the current density at 300 mA g^{-1} and cycling specific capacity of 600 mA h g^{-1} . (g and h) Low magnification pictures of the morphologies of the fully discharged (110) Fe₂O₃-based cathodes (b) with and (c) without light.

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experimental method. The adsorption of oxygen and intermediate species on the (110) and (104) facets of Fe₂O₃ were simulated by DFT calculations (Fig. S11, ESI†), and the decreases in free energy (ΔG) corresponding to the formation of LiO2 on the electrode surface and its later reduction were also calculated (Fig. 5a and b). The negative values of adsorption energy (E_{ads}) indicate that the thermodynamic process is favorable in both facets. The lower E_{ads} (-1.37 eV) on (110) than on (104) (-0.84 eV) indicates stronger oxygen adsorption on the former. $\Delta G_{(\text{LiO}_2)^*}$ and $\Delta G_{(\text{Li}_2\text{O}_2)^*}$ are respectively -3.76 eV and -3.69 eV for the (110) facets, whereas these values for (104) are -3.30 eV and -3.51 eV, respectively. Owing to the favored adsorption of oxygen species on the (110) facet, (110) Fe₂O₃ is more favorable for the electrode surface proceeded ORR process.

Except for oxygen-related reactants, the photo-induced morphology transformation is closely related to the charge distribution inside the porous cathode, which is determined by mapping the nanometer-resolution surface potential (Fig. 5c and d) using a Kelvin probe force microscope (KPFM). The values of linear contact potential difference (CPD) and height

in dark and light conditions are shown in Fig. 5e. In dark conditions, the CPD minimum is 655 mV and increases to 842 mV under light. The CPD values show larger variations in dark conditions than those in light conditions. In addition, the surface potential in light conditions is more evenly distributed than that in dark conditions (Fig. 5c and d). This difference suggests that light induces an internal-built electric field, which increases the kinetics for charge transfer, as shown by the electrochemical impedance spectra (EIS), both of fresh and cycled cathodes (Fig. 5f and Fig. S13, ESI†). The obtained charge transfer resistance (R_{ct}) and Warburg resistance (R_{w}) acquired from the equivalent circuits of EIS indicate the kinetics of electron transfer and oxygen transfer in the cathode, respectively. 42,43 The apparently lowered R_{ct} and R_{w} of the (110) Fe₂O₃ based cathode under light illustrates the unimpeded transfer process of oxygen and electrons, especially for the cycled conditions.

These results show that the light on (110) Fe₂O₃ plays a critical role in regulating the initial nucleation and subsequent growth of Li₂O₂ by anchoring intermediates on the cathode and homogenizing the charge distribution. As shown in Fig. 5g and h,

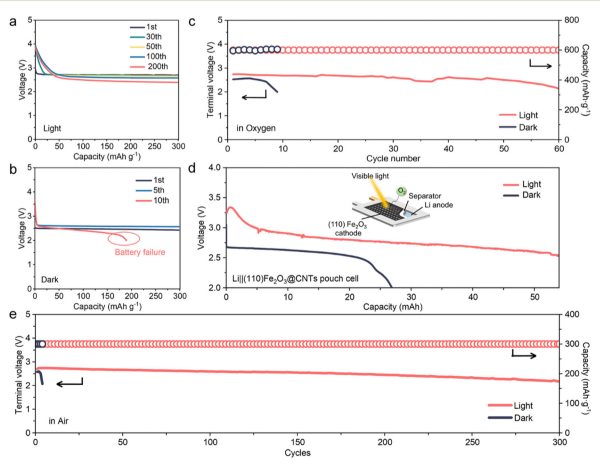


Fig. 6 Improved charge transfer kinetics and cycling stability induced by visible light on (110) Fe₂O₃ based cathodes. (a and b) Discharge profiles of the (110) Fe_2O_3 based cathode discharged in ambient air under (a) light and (b) dark conditions at 300 mA g^{-1} for different cycles. (c) The terminal discharge potential of each cycle for (110) Fe₂O₃ electrodes discharged in O₂ under dark and light conditions with the current density at 300 mA g⁻¹. (d) Discharge profiles for the (110) Fe_2O_3 based pouch cell at 0.05 mA cm⁻² in dark/light conditions. (e) The terminal discharge potential of each cycle for the (110) Fe₂O₃ electrodes discharged in ambient air under dark and light conditions with the current density at 300 mA g⁻¹

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Li₂O₂ produced under light is uniformly distributed on the whole surface, whereas the products produced without light accumulated on cathode domains (marked by the yellow dotted dash line). The local aggregation of products and their further obstruction of oxygen and electron-transport pathways leave plenty of unavailable "dead" active sites, leading to premature battery failure and limited capacity. This photo-assisted effect enables the smooth evolution of Li₂O₂ and the unimpeded transfer of reactants, and gives the cathodes the power to produce and store products associated with significantly improved capacities.

2.5. Cycling stability in actual Li-air batteries

The photo-assisted effect on (110) Fe₂O₃ significantly improves the cycling performance in O₂ and the actual air environment. In light conditions, the discharge plateau with reduced polarization is above 2.70 V and remains stable during cycling (Fig. 6a). This constant capacity has high reversibility and cycling stability accompanied by a very slight increase of R_{ct} (Fig. 5f and Fig. S13, ESI†). In sharp contrast, in dark conditions, the initial discharge plateau was lower (around 2.5 V), and during cycling, the polarization increased rapidly (Fig. S13, ESI†) and the battery failed after only 10 cycles accompanied by a significant increase of R_{ct} , which may result from byproduct accumulation (XPS spectra in Fig. 4f and g) caused by the large amounts of super-nucleophile LiO₂ intermediates (Fig. 4e).⁴⁴ An improvement in the pFe₂O₃ cathode was also observed under light (Fig. S14, ESI†), but it is less obvious than that on the faceted (110) Fe₂O₃ due to the lower photocatalytic activity (Fig. 2h, i and Fig. 3a). A Li-O₂ pouch cell was performed under dark and light conditions, respectively (Fig. 6d). Compared to the dark discharged profile, the light-assisted discharge plateau increased from 2.58 V to 2.80 V and the discharge capacity soared from 21 mAh to 53 mAh at a cut off voltage of 2.50 V. As for the cycling process conducted in the air environment, compared to only several cycles and fast failure of the battery performed without light, the discharge plateau of the lightassisted battery remained steady for over 300 cycles, which exhibits better reversibility and stability against the potential side-reactions present in ambient air.

3. Conclusion

In summary, we demonstrated an efficient photochemical strategy to regulate Li₂O₂ nucleation and growth behavior under visible light. An exceptional capacity increase and extended cycle life were realized, both in O_2 (~260%, over 60 cycles) and more promisingly in the actual air environment ($\sim 420\%$, over 300 cycles). This effect derives from two aspects. The (110) facet engineering of α-Fe₂O₃ improves its adsorption ability toward oxygen intermediates, suppressing the disproportionation reactions and byproduct generation. Additionally, the improved photocatalytic activities of (110) Fe₂O₃ cause rapid generation of photoelectrons and synchronous construction of a uniform photo-electric field throughout the entire electrode under visible light. The preferential nucleation of Li₂O₂ on (110) Fe₂O₃

surfaces and its subsequent consistent growth with the assistance of the photo-electric field induce the homogeneous distribution of solid Li₂O₂ depositions, preventing the local product accumulation and accelerating the O2 diffusion and electron transfer, thus prohibiting premature battery failure and promoting the reversible decomposition of Li₂O₂, achieving a long-term stable Li-air battery working in the actual air environment. This novel distributed electron transfer induced by visible light provides both material and conceptual breakthroughs for developing many other high-performance open battery systems that could promise the synergistic coordination of electric and light power.

4. Experimental section

4.1. Preparation of the cathode materials

All reagents in this work are analytically pure and used directly without further purification. Multiwalled Carbon Nanotubes (CNT, φ 10–20 nm) were purchased from Time Nano Co. Ltd, and used as a conducting matrix in Li-O2 cathodes. FeCl3·6H2O (Aladdin, 99%) and CH₃COONa·3H₂O (Aladdin, 99%) were used as precursors for hydrothermal synthesis of (110) Fe₂O₃. 2.76 g FeCl₃·6H₂O and 5.25 g CH₃COONa·3H₂O were dissolved in 35 mL anhydrous ethanol and stirred vigorously for 10 minutes. The solution was transferred into a 50 mL Teflon-lined stainlesssteel autoclave and heated in an oven at 200 °C for 24 h. After being cooled down, the resulting powder was vacuum-filtrated and then dried at 80 °C.

4.2. Characterization

X-Ray Diffraction (XRD, Rigaku MiniFlex-600) patterns were recorded to confirm the successful synthesis of (110) Fe₂O₃. Transmission electron microscopy (TEM, JEM 2100-F) was utilized to identify the microstructure and exposed facets of a single (110) Fe₂O₃ nanoparticle. Scanning Electron Microscopy (SEM, Regulus-8100) is used for morphology observation of Fe₂O₃ materials and the cathode. N₂ adsorption-desorption (Belsorp-mini) was conducted and the specific area of pFe₂O₃, (110) Fe₂O₃, pFe₂O₃/CNT and (110) Fe₂O₃/CNT could be calculated according to the Brunauer-Emmett-Teller (BET) mode. X-Ray photoelectron spectroscopy (XPS, Thermal Fisher Scientific Escalab-Xi +) was analyzed to identify the surface species via each detected element for the (110) Fe₂O₃ cathode being discharged. The UV-visible diffuse reflectance spectrum (DRS, Shimadzu UV-3600 Plus) was measured to characterize the light absorbance (a) of the samples in the range of 300 $< \lambda <$ 800 nm, and then the Tauc plot could be converted according to the Kubelka-Munk method. XPS valence band spectroscopy is conducted to determine the VB potential of the Fe₂O₃ sample by a VersaProbe III (Scanning ESCA Microprobe) SCA (Spherical Analyzer). Photoluminescence (PL) spectra were measured by a Steady state/Lifetime Spectrofluorometer (Edinburgh FLS1000) to characterize the lifetime of photoelectrons. Confocal Raman Microscopy (LabRAM HR Evolution) with an excitation

wavelength of 532 nm was used to measure and collect Raman spectra.

4.3. Electrochemical and photoelectrochemical performance

The photocatalytic activity of Fe₂O₃ was investigated by decomposing Rhodamine B (RhB) via irradiation with visible light. An amount of the Fe₂O₃ powders was dispersed in 100 mL of the RhB solution (3 \times 10⁻⁵ M). The prepared solution was stirred for 30 min in the dark to achieve an adsorptiondesorption equilibrium. Before the suspension was irradiated with 300 W Xeon lamps, 3 mL of H₂O₂ (30%) was added. After a given time interval, 4 mL of the mixture was collected, filtered and analyzed using the U-2910 single-beam spectrometer (Hitachi). Chronoamperometry measurement was tested at 0.87 V (vs. SHE) on an electrochemical workstation (Autolab) using a three-electrode system, in which the platinum electrode and Ag/AgCl electrode were used as a counter electrode and reference electrode, respectively. The contact potential difference (CPD) measurement was conducted with a Kelvin Probe Force Microscope (KPFM, Bruker Dimension icon) under an ambient atmosphere with Xeon arc light irradiation (400 nm < $\lambda < 800$ nm). The alternating current voltage was set to 0.5 V to avoid a high bias-induced band bending.

The prepared Fe₂O₃ powder, CNT and PVDF (poly(vinylidene fluoride)) were mixed with a mass ratio of 3:6:1, ground for 30 min to blend uniformly. After that, the slurry was spread on clean Carbon Paper (Avcarb P50T) with a diameter of 8 mm and dried in a vacuum oven at 100 °C for 24 h, and the Fe₂O₃/CNT cathode was obtained with the total average loading of Fe₂O₃ and CNT of 0.4-0.6 mg cm⁻². For a typical process for assembling a CR2032 coin Li-O2 battery, all the operations were conducted in an Argon atmosphere filled glove-box with both H₂O and O₂ contents below 0.5 ppm, in which the Fe₂O₃/CNT cathode was an oxygen electrode and photoelectrode, Li foil (12 mm in diameter and 0.5 mm in thickness) and glass fiber (Whatman, GF/A) were used as the counter electrode and the separator, respectively, and 80 µL of 1 M LiTFSI dissolved in TEGDME (DoDoChem, LK-001) was used as the electrolyte and dropped on the separator. The assembled Li-O2 battery was stored in a volume capacity of 250 mL sealed glass test device, which was filled with O2 and tested on a LAND CT2001A multichannel battery system. The Li-O2 pouch cell assembled by a (110) Fe₂O₃/CNT cathode (2 cm \times 5 cm), lithium foil (3 cm \times 6 cm) and glass fiber (Whatman, GF/A), Carbon paper and Cu-mesh were employed as a current collector for the cathode and anode, respectively. The pouch cell was packed by a plasticsealed Al-film with openings at the cathode side. The Li-air battery was assembled according to the above-mentioned process and operated under actual air. All the batteries were performed at room temperature (~25 degrees). An Autolab multichannel workstation was used to evaluate the cyclic voltammetry (CV) performance and electrochemical impedance spectra (EIS) in a frequency range from 10⁶ Hz to 10⁻² Hz of the coin cell. For photo-assisted discharge, the Xe lamp (CEAULIGHT-HXF300) (400 nm $< \lambda < 800$ nm) was opened

at the power of 130 W to irradiate the coin cell through the vessel and porous cathode shells.

The rotating ring-disk electrode (RRDE) test was performed with a similar procedure described by Sankarasubramanian et al.³⁷ The working electrode was a Φ 5 mm glassy carbon (GC) disk and Pt ring made by Phychemi (Hongkong) Co., Ltd. The slurry coated on the GC electrode was prepared with 2 mg Fe₂O₃ powder and carbon black blended with 15-20 µL Nafion dispersion (Alfa) was dispersed into isopropanol. The average loading is about 0.2 mg. The Pt electrode and partially delithiated LiFePO4 electrode were used as the counter electrode and reference electrode, respectively. The 0.2 M solution of lithium bis-trifluoromethanesulfonimidate (LiTFSI) (Sigma-Aldrich) in tetraethylene glycol dimethyl ether (DME) (Sigma-Aldrich) was prepared in an Argon filled glovebox after a further water removal process and used as the electrolyte. The water content was 36.3 ppm, measured by a Karl Fischer titration analyzer (Metrohm 899 Coulometer).

4.4. Computational section

All calculations in this work were carried out using the density functional theory (DFT) method as implemented in the VASP code. The electronic exchange–correlation energy was modeled using the Perdew–Burke–Ernzerhof (PBE) functional within the generalized gradient approximation (GGA). The projector augmented wave (PAW) method was used to describe the ionic cores. For the plane-wave expansion, a 450 eV kinetic energy cut-off was used after testing a series of different cut-off energies. A Monkhorst–Pack $3\times 3\times 1$ k-point grid was used to sample the Brillouin zone. The convergence criterion for the electronic structure iteration was set to be 10^{-4} eV, and that for geometry optimizations was set to be 0.01 eV Å⁻¹ on force. Gaussian smearing of 0.1 eV was applied during the geometry optimization and for the total energy computations.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 X.-D. Lin, Y. Gu, X.-R. Shen, W.-W. Wang, Y.-H. Hong, Q.-H. Wu, Z.-Y. Zhou, D.-Y. Wu, J.-K. Chang, M.-S. Zheng, B.-W. Mao and Q.-F. Dong, Energy Environ. Sci., 2021, 14, 1439-1448.
- 2 Y. Dou, Z. Xie, Y. Wei, Z. Peng and Z. Zhou, Natl. Sci. Rev., 2022, nwac040.
- 3 K. Chen, G. Huang and X.-B. Zhang, Chin. J. Chem., 2021, 39,
- 4 Y. Liu, J. Cai, J. Zhou, Y. Zang, X. Zheng, Z. Zhu, B. Liu, G. Wang and Y. Oian, eScience, 2022, 2, 389-398.
- 5 L. D. Griffith, A. E. S. Sleightholme, J. F. Mansfield, D. J. Siegel and C. W. Monroe, ACS Appl. Mater. Interfaces, 2015, 7, 7670-7678.
- 6 B. D. Adams, C. Radtke, R. Black, M. L. Trudeau, K. Zaghib and L. F. Nazar, Energy Environ. Sci., 2013, 6, 1772-1778.
- 7 D. Aurbach, B. D. McCloskey, L. F. Nazar and P. G. Bruce, Nat. Energy, 2016, 1, 1-11.
- 8 C.-L. Li, G. Huang, Y. Yu, Q. Xiong, J.-M. Yan and X. Zhang, J. Am. Chem. Soc., 2022, 144, 5827-5833.
- 9 Y.-C. Lu, B. M. Gallant, D. G. Kwabi, J. R. Harding, R. R. Mitchell, M. S. Whittingham and Y. Shao-Horn, Energy Environ. Sci., 2013, 6, 750-768.
- 10 J. Shui, F. Du, C. Xue, Q. Li and L. Dai, ACS Nano, 2014, 8,
- 11 A. Liu, X. Liang, X. Ren, W. Guan and T. Ma, Electrochem. Energy Rev., 2022, 5, 112-144.
- 12 G. Zhang, G. Li, J. Wang, H. Tong, J. Wang, Y. Du, S. Sun and F. Dang, Adv. Energy Mater., 2022, 12, 2103910.
- 13 Q. Zhang, C. Wang, Z. Xie and Z. Zhou, Energy Environ. Mater., 2022, 5, 1103-1116.
- 14 L. Guo, L. Tan, A. Xu, G. Li, G. Zhang, R. Liu, J. Wang, Y. Du and F. Dang, Energy Storage Mater., 2022, 50, 96-104.
- 15 X. Gao, Y. Chen, L. R. Johnson, Z. P. Jovanov and P. G. Bruce, Nat. Energy, 2017, 2, 1-7.
- 16 M. Balaish, X. Gao, P. G. Bruce and Y. Ein-Eli, Adv. Mater. Technol., 2019, 4, 1800645.
- 17 F. Z. Wang and X. L. Li, ACS Appl. Mater. Interfaces, 2018, 10, 26222-26232.
- 18 V. Viswanathan, K. S. Thygesen, J. S. Hummelshøj, J. K. Nørskov, G. Girishkumar, B. D. McCloskey and A. C. Luntz, J. Chem. Phys., 2011, 135, 214704.
- 19 L. Cheng, P. Redfern, K. C. Lau, R. S. Assary, B. Narayanan and L. A. Curtiss, J. Electrochem. Soc., 2017, 164, E3696.
- 20 S. Zhang, M. Chen, X. Zhao, J. Cai, W. Yan, J. C. Yen, S. Chen, Y. Yu and J. Zhang, Electrochem. Energy Rev., 2021, 4, 336-381.
- 21 K. Chen, D.-Y. Yang, G. Huang and X.-B. Zhang, Acc. Chem. Res., 2021, 54, 632-641.
- 22 D. Geng, N. Ding, T. S. A. Hor, S. W. Chien, Z. Liu, D. Wuu, X. Sun and Y. Zong, Adv. Energy Mater., 2016, 6, 1502164.
- 23 E. J. Nemanick and R. P. Hickey, J. Power Sources, 2014, 252, 248-251.

- 24 A. K. Thapa, Y. Hidaka, H. Hagiwara, S. Ida and T. Ishihara, J. Electrochem. Soc., 2011, 158, A1483.
- 25 C. Tran, X.-Q. Yang and D. Qu, J. Power Sources, 2010, 7.
- 26 H. C. Lee, J. O. Park, M. Kim, H. J. Kwon, J.-H. Kim, K. H. Choi, K. Kim and D. Im, Joule, 2019, 3, 542-556.
- 27 C. O. Laoire, S. Mukerjee, K. M. Abraham, E. J. Plichta and M. A. Hendrickson, J. Phys. Chem. C, 2009, 113, 20127–20134.
- 28 Z. Lyu, Y. Zhou, W. Dai, X. Cui, M. Lai, L. Wang, F. Huo, W. Huang, Z. Hu and W. Chen, Chem. Soc. Rev., 2017, 46, 6046-6072.
- 29 J. Wang, Y. Zhang, L. Guo, E. Wang and Z. Peng, Angew. Chem., Int. Ed., 2016, 55, 5201-5205.
- 30 H. Gong, T. Wang, K. Chang, P. Li, L. Liu, X. Yu, B. Gao, H. Xue, R. Ma, J. He and J. Ye, Carbon Energy, 2022, 4, 1169-1181.
- 31 M. Wang, J. Chen, Z. Tian, W. Dai, B. Cui, X. Cui, D. Wang, Y. Xiao, X. Lian, C. Jiang, H. Yang, Y. Wang, Z. Sun, Y. Ding, Y.-Y. Sun, J. Zhang and W. Chen, Energy Environ. Sci., 2023, 16, 523-534.
- 32 D. Du, Z. Zhu, K.-Y. Chan, F. Li and J. Chen, Chem. Soc. Rev., 2022, 51, 1846-1860.
- 33 N. B. Aetukuri, B. D. McCloskey, J. M. Garcia, L. E. Krupp, V. Viswanathan and A. C. Luntz, Nat. Chem., 2015, 7, 50-56.
- 34 Y. Wang, N.-C. Lai, Y.-R. Lu, Y. Zhou, C.-L. Dong and Y.-C. Lu, Joule, 2018, 2, 2364-2380.
- 35 T. Liu, J. T. Frith, G. Kim, R. N. Kerber, N. Dubouis, Y. Shao, Z. Liu, P. C. M. M. Magusin, M. T. L. Casford, N. Garcia-Araez and C. P. Grey, J. Am. Chem. Soc., 2018, 140, 1428-1437.
- 36 L. Johnson, C. Li, Z. Liu, Y. Chen, S. A. Freunberger, P. C. Ashok, B. B. Praveen, K. Dholakia, J.-M. Tarascon and P. G. Bruce, Nat. Chem., 2014, 6, 1091-1099.
- 37 S. Sankarasubramanian, J. Seo, F. Mizuno, N. Singh and J. Prakash, J. Phys. Chem. C, 2017, 121, 4789-4798.
- 38 J. Wang, L. Ma, J. Xu, Y. Xu, K. Sun and Z. Peng, SusMat, 2021, 1, 345-358.
- 39 S. Dong, S. Yang, Y. Chen, C. Kuss, G. Cui, L. R. Johnson, X. Gao and P. G. Bruce, Joule, 2022, 6, 185-192.
- 40 N. Mahne, B. Schafzahl, C. Leypold, M. Leypold, S. Grumm, A. Leitgeb, G. A. Strohmeier, M. Wilkening, O. Fontaine, D. Kramer, C. Slugovc, S. M. Borisov and S. A. Freunberger, Nat. Energy, 2017, 2, 1-9.
- 41 Y. K. Petit, E. Mourad, C. Prehal, C. Leypold, A. Windischbacher, D. Mijailovic, C. Slugovc, S. M. Borisov, E. Zojer, S. Brutti, O. Fontaine and S. A. Freunberger, Nat. Chem., 2021, 1-7.
- 42 J. Huang, B. Tong, Z. Li, T. Zhou, J. Zhang and Z. Peng, J. Phys. Chem. Lett., 2018, 9, 3403-3408.
- 43 X. Luo, R. Amine, K. C. Lau, J. Lu, C. Zhan, L. A. Curtiss, S. Al Hallaj, B. P. Chaplin and K. Amine, Nano Res., 2017, 10, 4327-4336.
- 44 V. S. Bryantsev, V. Giordani, W. Walker, M. Blanco, S. Zecevic, K. Sasaki, J. Uddin, D. Addison and G. V. Chase, J. Phys. Chem. A, 2011, 115, 12399-12409.