

ELECTROCHEMISTRY

A long-life lithium-oxygen battery via a molecular quenching/mediating mechanism

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The advancement of lithium-oxygen (Li-O₂) batteries has been hindered by challenges including low discharge capacity, poor energy efficiency, severe parasitic reactions, etc. We report an Li-O₂ battery operated via a new quenching/mediating mechanism that relies on the direct chemical reactions between a versatile molecule and superoxide radical/Li₂O₂. The battery exhibits a 46-fold increase in discharge capacity, a low charge overpotential of 0.7 V, and an ultralong cycle life >1400 cycles. Featuring redox-active 2,2,6,6-tetramethyl-1-piperidinyloxy moieties bridged by a quenching-active perylene diimide backbone, the tailor-designed molecule acts as a redox mediator to catalyze discharge/charge reactions and serves as a reusable superoxide quencher to chemically react with superoxide species generated during battery operation. The all-in-one molecule can simultaneously tackle issues of parasitic reactions associated with superoxide radicals, singlet oxygen, high overpotentials, and lithium corrosion. The molecular design of multifunctional additives combining various capabilities opens a new avenue for developing high-performance Li-O₂ batteries.

INTRODUCTION

Lithium-oxygen (Li-O₂) batteries have the highest theoretical specific energy among all-known battery chemistries and are deemed a disruptive technology if a practical device could be realized (1–4). Typically, a nonaqueous Li-O₂ battery consists of a lithium metal anode separated from a porous oxygen cathode by an Li⁺ conducting electrolyte, and its operation relies on the stripping and plating of lithium on the negative electrode and the formation and decomposition of Li₂O₂ on the positive electrode (5). However, current Li-O₂ batteries suffer from challenges including low discharge capacity, poor round-trip efficiency, severe parasitic reactions, etc.

Toward solving these challenges and unlocking the energy capabilities of Li-O₂ electrochemistry, a great deal of research effort has been devoted to a fundamental understanding of the reactions and processes underpinning the operation of Li-O₂ batteries. It has been established that the discharge of an Li-O₂ battery proceeds with the oxygen reduction reaction (ORR) generating superoxide species (superoxide radicals O₂^{•−}/lithium superoxide LiO₂) that can either adsorb on the cathode surface followed by a surface mechanism or dissolve in the electrolyte solution followed by a solution mechanism (6, 7). The surface mechanism often produces Li₂O₂ with a thin layer morphology (a few nanometers in thickness) that rapidly passivates the cathode surface and leads to a low discharge capacity and a low

rate capability. On the contrary, the solution mechanism usually generates Li₂O₂ with a toroidal shape (up to a few micrometers in diameter) in the electrolyte solution and can support a high discharge capacity and a high rate capability. However, toroidal-shaped Li₂O₂ is not sufficiently close to the cathode surface, and a major part of the Li₂O₂ toroid is beyond the e-tunneling range during the charge process. Therefore, the oxidation of these Li₂O₂ proceeds through mechanisms beyond simply one-step electrochemical oxidation, including direct 2e[−] oxidation, solid-solution oxidation (via Li_{2-x}O₂), and stepwise oxidation (via LiO₂) (6, 8–10). Recent studies suggest that singlet oxygen (¹O₂) may also be involved in discharge and charge processes, during the disproportionation reaction of LiO₂ species, and in the 1e[−] oxidation of LiO₂ intermediates and 2e[−] oxidation of solid Li₂O₂ (11). Parasitic reactions occur inevitably during both discharge and charge processes. The parasitic reactions during discharge are mainly associated with the chemical attack of battery components (electrolytes and electrodes) by the reactive oxygen species (O₂^{•−}, LiO₂, Li₂O₂, and ¹O₂) (12, 13). However, the parasitic reactions during the charge process are more than chemical attack. Besides the chemical decomposition of battery components by reactive oxygen species generated during charging, the high voltage (>4 V) frequently applied during charging can also cause the electrochemical oxidation of the electrolyte and cathode materials, particularly in the presence of Li₂O₂ (14–16).

Many strategies have been devised to increase discharge capacity, lower charge overpotential, and alleviate parasitic reactions. These include using high donor number solvents and salts with the ability to decrease the reactivity (acidity) of Li⁺, certain additives that can stabilize superoxide radicals, and redox mediators that can relay electrons between cathode surface and dissolved oxygen (17–20). These approaches have been used to promote Li₂O₂ formation in the electrolyte solution rather than on the cathode surface and therefore increase the discharge capacity, among which the redox mediators demonstrate prominent performance. Nevertheless, the discharge product Li₂O₂ is difficult to be oxidized owing to its poor physical contact with the cathode surface and its intrinsically low electronic

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and ionic conductivity (21, 22). Oxygen evolution reaction (OER) redox mediators have also been proposed to efficiently oxidize Li_2O_2 . The oxidized form of redox mediators generated on the cathode surface can diffuse to the Li_2O_2 in the electrolyte solution and oxidize Li_2O_2 to O_2 with itself being regenerated (23–30). Because the OER redox mediators often have a redox potential that is much lower than the potential required to charge a Li-O₂ cell (usually >4.0 V), the energy efficiency of the Li-O₂ battery can be improved accordingly. Because of the low charging voltage, the decomposition of battery components during charging can be alleviated. Furthermore, to reduce the parasitic reactions caused by the attack of the reactive oxygen species, more stable electrolytes and cathode materials need to be used to construct Li-O₂ batteries (31–33). In addition, a few ORR redox mediators that can promote oxygen reduction to Li_2O_2 via a benign intermediate instead of the very reactive superoxide radicals have been used to mitigate the parasitic reactions (19, 34–40). Similarly, molecular $^1\text{O}_2$ quencher has also been used to deactivate the reactivity of $^1\text{O}_2$, which also contributes to the suppression of overall parasitic reactions (41). However, the combination of these functional molecules in one cell is rarely reported. The intrinsic reactivity of each component can easily conflict with another, leading to the weakening or even deactivating of the functionalities of these individual molecules.

All in all, to realize a high discharge capacity, a low charge overpotential, and less parasitic reactions for Li-O₂ batteries, it is desirable to have a multifunctional catalyst that simultaneously has the capabilities of ORR redox mediators, OER redox mediators, and superoxide radical quencher. However, the design and synthesis of such all-in-one molecules have not been reported. Here, we rationally designed and successfully synthesized a multifunctional superoxide radical quencher by grafting two redox mediator-active 2,2,6,6-tetramethyl-1-piperidinyloxy (TEMPO) functional groups to a quenching-active perylene diimide (PDI) backbone. The as-synthesized *N,N'*-di-(2,2,6,6-tetramethyl-1-oxyl-4-piperidinyl)-perylene-3,4,9,10-tetracarboxylic diimide (PDI-TEMPO) molecules can not only chemically quench superoxide species generated during discharge and charge processes to suppress the generation of $^1\text{O}_2$ but also act as a redox mediator to efficiently catalyze the formation and decomposition of Li_2O_2 in the electrolyte solution and lower the discharge and charge overpotentials of Li-O₂ batteries. The synergistic effect of TEMPO and PDI moieties can also induce more functionalities to minimize parasitic reactions such as preventing the corrosion of lithium anode. When adding PDI-TEMPO in the electrolyte, the Li-O₂ batteries proceed via an alternative chemical quenching mechanism, which achieves higher discharge capacity, lower charge overpotential, less parasitic reaction, and longer cycle life.

RESULTS

Synthesis and characterization of PDI-TEMPO molecules

The PDI-TEMPO reagent was synthesized by reacting 4-amino-2,2,6,6-tetramethylpiperidine with perylene-3,4,9,10-tetracarboxylic dianhydride, followed by an oxidation process (fig. S1A), yielding reddish-purple powders as the final product (42, 43). The PDI-TEMPO shows a good solubility in common electrolyte solvents used in Li-O₂ batteries, including diethylene glycol dimethyl ether (DEGDME) and dimethyl sulfoxide (DMSO) (fig. S1B). The molecular structures of PDI-TEMPO and intermediates were confirmed by several spectroscopic techniques including ultraviolet-visible (UV-Vis) spectroscopy,

mass spectrometry (MS), Fourier transform infrared (FTIR) spectroscopy, and electron paramagnetic resonance (EPR) spectroscopy (fig. S2). The strong absorption bands of amide at 1650 and 1680 cm^{-1} in the FTIR spectra indicate that the TEMPO functional groups have been covalently linked to the PDI backbone (fig. S2C). Fluorescence spectroscopy shows that the TEMPO radicals in PDI-TEMPO molecules quench the fluorescence of the PDI backbones, further verifying the successful synthesis of PDI-TEMPO molecules (fig. S3) (44).

The PDI-TEMPO molecule features two TEMPO moieties bridged by a conjugated PDI backbone (Fig. 1A). The multifunctional PDI-TEMPO molecule is specifically designed to enhance the electrochemical performance and reduce parasitic reactions of Li-O₂ batteries (fig. S4). The conjugated PDI backbone is an efficient electron acceptor to react with $\text{O}_2^{\bullet-}$ and can behave as a quencher by quenching the reactivity of $\text{O}_2^{\bullet-}$ (42). The TEMPO moieties can be electrochemically reduced and oxidized and therefore can be used as a soluble redox mediator for promoting both discharge and charge reactions in Li-O₂ batteries. The chemical grafting of these moieties via chemical bonds integrates these functionalities in one molecule and enables the intermolecular interactions, which may stabilize these moieties and present individual properties without interference. The chemical bondings between these two functional moieties can not only efficiently overcome the drawback of each group (insolubility for PDI-based materials and irreversible n-doping of TEMPO molecules) but also provide additional functionalities beyond the simple combinations. These synergistic properties can also efficiently suppress the generation of $^1\text{O}_2$ (41, 45, 46). Owing to the steric interactions between the conjugated PDI backbone and bulky TEMPO moieties, PDI-TEMPO molecules have a sterically twisted structure that the TEMPO moieties and PDI backbones are not on the same plane (Fig. 1B). Such unique molecular architecture can maintain optimal van Der Waals forces via π - π interaction between the conjugated PDI backbone and the basal plane of the graphitic carbon cathode. As a result, the molecules are drawn to the surface of the carbon cathode owing to the attraction between conjugated PDI backbone and carbon cathode, while they are prevented from depositing on the cathode because of the repulsion between the bulky TEMPO moieties and carbon cathode, which prohibited its diffusion to the lithium anode (47).

Redox behavior of PDI-TEMPO molecules

The redox behavior of PDI-TEMPO has been examined using a glassy carbon (GC) electrode in DEGDME electrolyte. As shown in the cyclic voltammetry (CV) results in Fig. 1C, PDI-TEMPO shows three pairs of redox peaks in the voltage range of 1.8 to 4.4 V. Two pairs of redox peaks (labeled by a, a' at ~2.8 V, and b, b' at ~3.8 V) are assigned to the reversible reduction (a and a') and oxidation (b and b') of TEMPO moieties, respectively, which is consistent with previous reports (37, 48, 49). The third pair of redox peaks started at ~2.3 V originate from the electrochemistry of the PDI backbone, where the corresponding oxidation peak merges with the oxidation peak of reduced TEMPO moieties, causing the merged peak position to downshift from 3.0 V (the oxidation potential of reduced TEMPO moieties) to 2.5 V (50). These redox pairs are demonstrated highly reversible, showing no visible decay after three cycles (fig. S5A). Even after 25 cycles in oxygen, the characteristic redox peaks remain in the following scan in the argon atmosphere, further verifying the high stability of these redox pairs (fig. S5B).

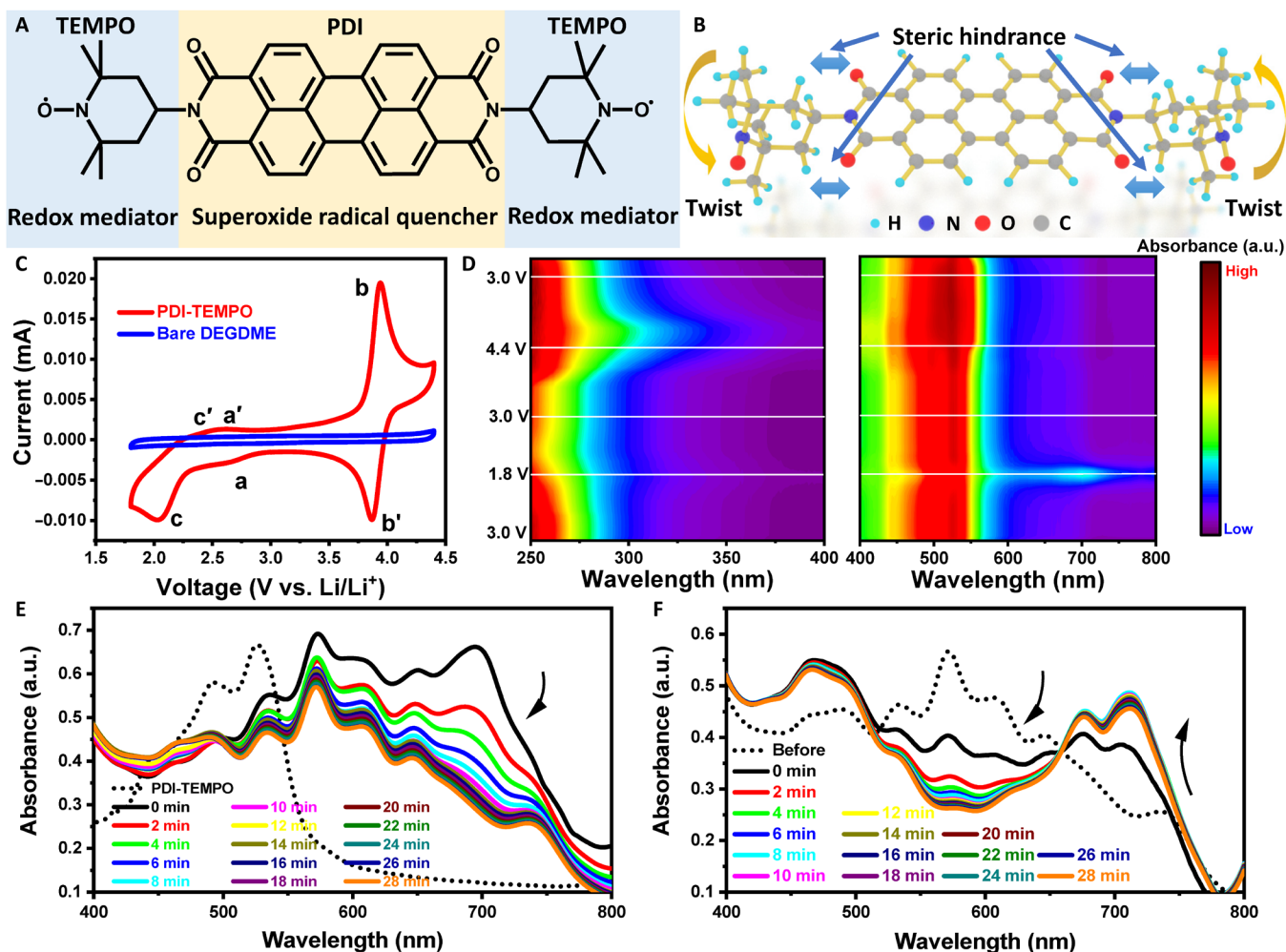


Fig. 1. Investigation of the redox capability of PDI-TEMPO in electrolytes. (A) The molecular structure and (B) stereostructure of PDI-TEMPO. a.u., arbitrary units. (C) CV curves of three-electrode cells with and without PDI-TEMPO in the DEGDM electrolyte in an argon atmosphere. The scan rate is 100 mV s^{-1} . The concentration of PDI-TEMPO is 5 mM. (D) In situ UV-Vis spectra of the electrolyte with PDI-TEMPO scanned in the range from 1.8 to 4.4 V. (E and F) UV-Vis spectra of the KO_2 -saturated DMSO solution with PDI-TEMPO (E) before and (F) after the addition of lithium salt. The marked PDI-TEMPO spectrum in (E) is the original PDI-TEMPO/DMSO solution.

The electrochemistry of PDI-TEMPO has also been studied by in situ UV-Vis spectroscopy (Fig. 1D and fig. S6). During the potential scan from 3.0 to 1.8 V, the absorption band in the range of 250 to 350 nm associated with the TEMPO moieties gradually decreased, indicating that TEMPO moieties were reduced (fig. S6A). Simultaneously, the absorption band in the range of 450 to 550 nm associated with the pristine PDI backbone decreased, and the absorption at 700 nm from reduced PDI backbone increased (fig. S6B). The in situ UV-Vis spectroscopic study provides direct evidence that the PDI backbone has also been reduced following the reduction of the TEMPO moieties. Both reduction reactions are highly reversible because all absorption bands are recovered to their original states at the end of the reverse potential scan from 1.8 to 3.1 V (fig. S6, C and D). When the potential is further scanned to 3.6 V and beyond, the absorption band of the oxidized TEMPO moieties (250 to 350 nm) increases accordingly (fig. S6E) and recovers to their original intensities when the voltage is reversed to 3.0 V (fig. S6G). The peaks associated with the PDI backbone does not change because no redox reaction of PDI takes place in this potential range of 3.0 to 4.5 V

(fig. S6, F and H). These results unambiguously confirm that there are three highly reversible redox reactions of PDI-TEMPO in the operation potential range of Li- O_2 batteries.

The redox behavior of PDI-TEMPO molecules has been further elucidated through density functional theory (DFT) calculations. Figure S7 shows the visualized lowest unoccupied molecular orbital (LUMO), highest occupied molecular orbital (HOMO), and singly occupied molecular orbital (SOMO) of PDI-TEMPO molecules. The HOMO is fully occupied with electrons and thereby can only provide electrons. Its energy level corresponds to the oxidation potential. Similarly, the LUMO can only accept electrons, and its energy level relates to the reduction potential. The SOMO, on the other hand, is partially occupied by one electron and therefore can both provide and accept electrons, and its energy level represents both oxidation and reduction potentials. Furthermore, the positions of the molecular orbitals also represent the possible sites where the redox reactions are most likely to occur. As shown in fig. S7A (left), both LUMO and HOMO are distributed on TEMPO moieties of PDI-TEMPO, suggesting that the initial oxidation and reduction reactions

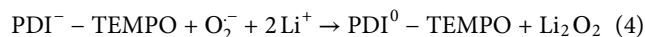
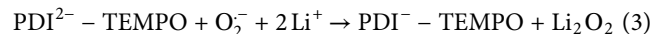
of PDI-TEMPO are based on reversible redox reactions of TEMPO moieties. Similarly, as presented in fig. S7A (right), the SOMO of the oxidized PDI-TEMPO (PDI-TEMPO⁺) is only distributed in TEMPO moieties, revealing that only TEMPO moieties are capable of being reduced (oxidized TEMPO moiety) and oxidized (neutral TEMPO moiety) at this stage. The SOMO of the reduced PDI-TEMPO (PDI-TEMPO⁻) in fig. S7A (middle) mainly locates on the PDI core and one TEMPO moiety. This designates that the further reduction of PDI-TEMPO would occur on the PDI backbone, while the oxidation reaction would take place on the reduced TEMPO moiety. The DFT calculation results are consistent with the electrochemical studies (Fig. 1, C and D).

The oxidative and reductive capability of PDI-TEMPO molecules shown in fig. S7B reveals that the SOMO of the reduced PDI-TEMPO (-1) is higher than the LUMO of O₂. Therefore, the reduced PDI-TEMPO can reduce O₂ during the discharge process. The SOMO of the oxidized PDI-TEMPO (+1) is notably lower than the HOMO of Li₂O₂, suggesting that it can decompose Li₂O₂ during the charge process. Thus, PDI-TEMPO molecules can boost both discharge and charge processes of Li-O₂ batteries owing to their unique redox properties.

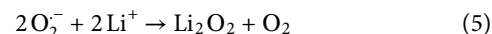
PDI-TEMPO molecule as a highly efficient O₂⁻ quencher

PDI-TEMPO with the unique molecular and electronic structure can not only facilitate the formation and decomposition of Li₂O₂ via the redox-active TEMPO functional moieties but also effectively chemically eliminate O₂⁻ through the conjugated PDI backbone as an O₂⁻ quencher. Thus, the parasitic reactions originated from O₂⁻ are expected to be substantially suppressed. We verified this hypothesis by dissolving potassium superoxide (KO₂) in the organic solvents, simulating the formation of O₂⁻ during the discharge process. UV-Vis spectra were collected to monitor reactions between O₂⁻ and PDI-TEMPO molecules (51). We used DMSO as the solvent because of the relatively high solubility of O₂⁻ species in DMSO. After adding PDI-TEMPO into the KO₂-saturated DMSO, the UV-Vis absorption band corresponding to the original PDI-TEMPO (PDI⁰-TEMPO, 450 to 550 nm) vanished immediately, and the band associated with the reduced PDI backbone (PDI⁻-TEMPO, 700 nm) appeared initially but decreased with time (Fig. 1E). Meanwhile, a series of absorption peaks in the range of 500 to 650 nm emerged and remained. These newly formed peaks correspond to the reduced PDI backbone with the acceptance of two electrons (PDI²⁻-TEMPO) (42). This reduction process is accompanied by a color change of the solution from red to brown and, lastly, blue (fig. S8). Therefore, the PDI backbone can be easily reduced by O₂⁻ (fig. S9A). After adding lithium salt [lithium bis(trifluoromethanesulfonyl)imide (LiTFSI)] in the aforementioned reduced PDI-TEMPO solution, the UV absorption associated with the PDI²⁻-TEMPO species gradually vanished, while the peaks corresponding to PDI⁰-TEMPO and PDI⁻-TEMPO reappeared (Fig. 1F). This phenomenon indicates that the reduced PDI backbone can be easily oxidized by O₂⁻ when lithium salt is present (fig. S9B). As there are still O₂⁻ presenting in the electrolyte, the PDI-TEMPO can still be reduced by O₂⁻ and then oxidized by lithium salt and O₂⁻ (fig. S9B). This is a dynamic process that lithium salt and superoxide radicals are continuously consumed to form Li₂O₂, facilitated by the chemical quenching reactions with PDI-TEMPO. This mechanism was further confirmed by EPR studies. The EPR peaks generated from the combination of 5,5-dimethyl-1-pyrroline *N*-oxide (DMPO) and O₂⁻ gradually

disappeared after adding PDI-TEMPO and lithium salt (fig. S10). The chemical reactions between PDI-TEMPO and O₂⁻ are proposed as below



Overall reaction



On the basis of the above experimental results, PDI-TEMPO molecules are confirmed to be a reusable soluble O₂⁻ quencher, which efficiently eliminates O₂⁻ by chemically reacting with O₂⁻ to form Li₂O₂. As a result, the O₂⁻ species will be immediately consumed once they are generated, and the parasitic reactions induced by O₂⁻ will be dramatically diminished. Furthermore, the multistep chemical reactions between O₂⁻ and PDI-TEMPO detour around the self-disproportion reactions of O₂⁻ (LiO₂), which significantly suppresses the generation of ¹O₂, eliminating parasitic reactions associated with ¹O₂ as well (fig. S4) (41, 45, 46). The deactivation of O₂⁻ by PDI-TEMPO has been further demonstrated by monitoring the reaction of KO₂ and DEGDME electrolyte for 1 week (fig. S11). The FTIR spectra of the DEGDME electrolyte containing PDI-TEMPO show no obvious change after a 1-week reaction, suggesting no detectable parasitic reactions between O₂⁻ and DEGDME solvent. However, the FTIR spectrum of the control sample (no PDI-TEMPO) shows an increase in peak at 1650 cm⁻¹, which signifies the formation of C=O, because of the attack by active oxygen species. Therefore, PDI-TEMPO in the electrolyte can simultaneously diminish parasitic reactions and promote the formation of Li₂O₂.

Discharge behavior of Li-O₂ batteries with PDI-TEMPO molecules

The electrochemistry of PDI-TEMPO was examined with CV using a GC electrode in an electrolytic cell. As shown in fig. S12, the ORR peak current in the PDI-TEMPO electrolyte is much higher than that in the bare DEGDME electrolyte. This clearly corroborates the PDI-TEMPO's capability of facilitating the discharge process. Similar results have also been obtained with an Li-O₂ cell containing a carbon paper positive electrode and a lithium foil negative electrode (figs. S13 and S14). Figure 2A shows the discharge-charge profiles of an Li-O₂ cell with a cutoff voltage of 2.0 and 4.6 V. The cell with PDI-TEMPO electrolyte delivers a specific capacity of 4.5 mAh cm⁻², which is more than 46 times higher than that of the cell with bare DEGDME electrolyte (less than 0.1 mAh cm⁻²; Fig. 2A and fig. S15). It is worth noting that the same configured cell with PDI-TEMPO electrolyte operated in an argon atmosphere only provides a specific capacity of 0.02 mAh cm⁻² (fig. S16), which is negligible compared with the capacity obtained in the oxygen atmosphere. Therefore, the high specific capacity should be entirely ascribed to the reversible formation and decomposition of Li₂O₂ (figs. S17 to S18), instead of the redox reaction of PDI-TEMPO dissolved in the electrolyte.

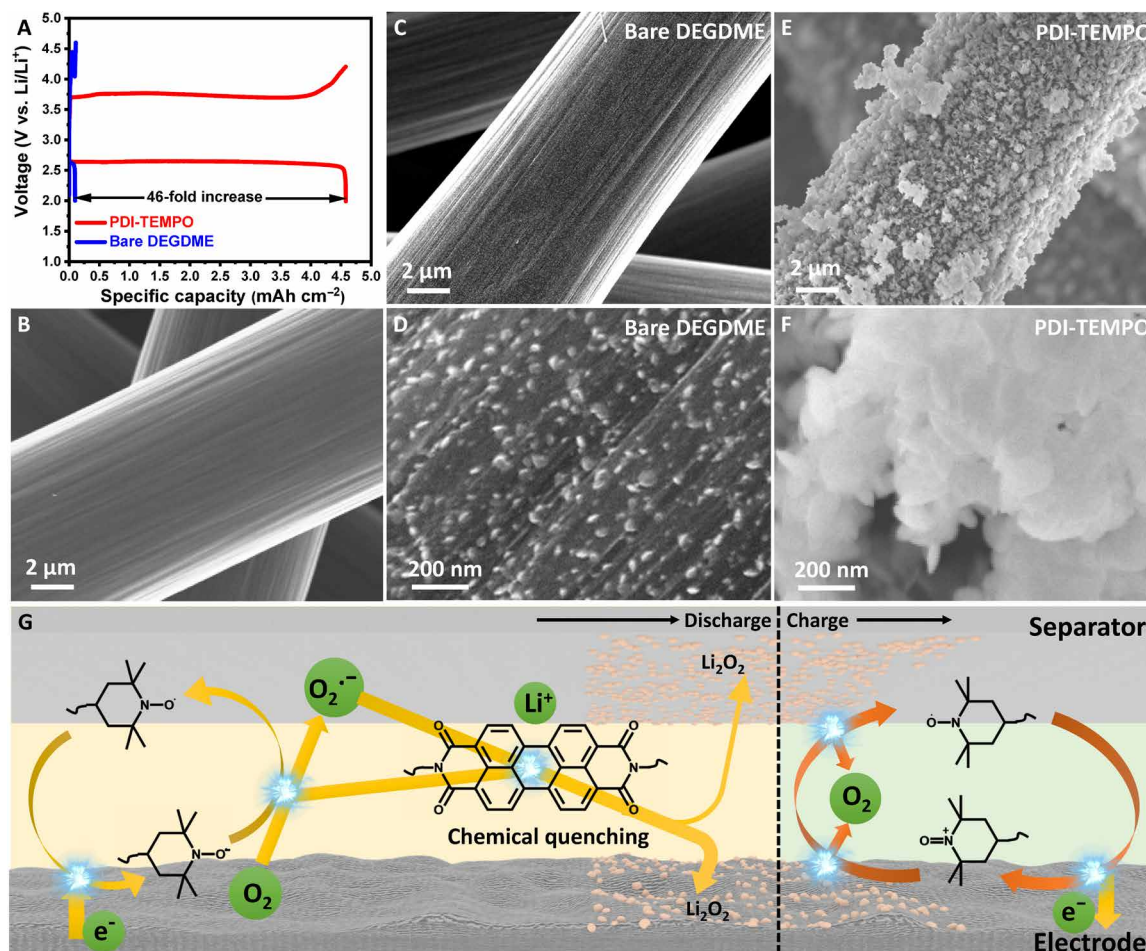


Fig. 2. Improvement of Li-O₂ batteries with the aid of PDI-TEMPO. (A) Full discharge-charge profiles of Li-O₂ batteries with PDI-TEMPO electrolyte and bare DEGDM electrolyte. Lithium foils are used as the anodes, and carbon papers as the cathodes. The current density is 0.1 mA cm⁻². The concentration of PDI-TEMPO is 5 mM. SEM images of the carbon paper electrodes (B) before and (C to F) after the first discharge. The electrolytes used are (C and D) bare DEGDM electrolyte and (E and F) PDI-TEMPO electrolyte. (G) Schematic illustration of the PDI-TEMPO facilitating the formation and decomposition of Li₂O₂ during the operation of Li-O₂ batteries.

The morphologies of discharged Li₂O₂ on the carbon paper electrode after the first discharge were characterized by scanning electron microscopy (SEM). Comparing to the pristine morphology in Fig. 2B, only a small amount of discharge products are observed on carbon fibers when discharged in the bare DEGDM electrolyte (Fig. 2, C and D). On the contrary, a large quantity of Li₂O₂ was generated on carbon fibers in the PDI-TEMPO electrolyte (Fig. 2, E and F). The cells with PDI-TEMPO electrolyte produced Li₂O₂ nanoparticles as the discharge product, which is different from the toroidal-shaped Li₂O₂ commonly obtained through a solution mechanism (fig. S19) (52, 53). This indicates that the discharge process may proceed via a different mechanism. We propose a chemical quenching process of O₂^{•-} by PDI-TEMPO molecules (54). As schematically shown in Fig. 2G, the discharge of the Li-O₂ cell is initiated by the reduction of TEMPO moieties, which reacts with oxygen molecules to catalyze the ORR process. The reduced oxygen species are then either directly released into the electrolyte or interacted with the reduced TEMPO moieties, which are then chemically quenched by the PDI backbone to generate Li₂O₂ nanoparticles. Therefore, the entire discharge process is mainly driven by the chemical quenching of O₂^{•-} in the electrolyte, which has also been verified by the rotating ring-disk

electrode (RRDE) measurements (figs. S20 and S21) (55). The above process is further enhanced when carbon nanotube (CNT) electrodes with higher specific surface area are used to accelerate the formation of O₂^{•-} (figs. S22 to S25). The results clearly corroborate that PDI-TEMPO molecules can efficiently relay electrons from the electrode surface to the dissolved O₂ to produce O₂^{•-}. Meanwhile, the O₂^{•-} can be chemically disproportionated to Li₂O₂ with the assistance of PDI backbones simultaneously (figs. S26 and S27 and movie S1).

Charge behavior of Li-O₂ batteries with PDI-TEMPO molecules

The PDI-TEMPO molecules can also function as a charging reaction redox mediator to efficiently improve the round-trip efficiency for Li-O₂ batteries (fig. S28). As shown in Fig. 2A and figs. S29 and S30, the charge voltage has been significantly decreased to 3.7 V (compared with 4.2 V in bare DEGDM electrolyte). This result is consistent with the CV curves (Fig. 1C), indicating that the redox reaction of PDI-TEMPO is responsible for the decrease in the charge overpotentials of the Li-O₂ cells.

To further analyze the discharge-charge process, galvanostatic intermittent titration technique (GITT) was conducted during the

discharge and charge processes. As shown in Fig. 3A, the discharge curves and the open-circuit potentials in the PDI-TEMPO electrolyte are more stable than that in the bare DEGDME electrolyte, owing to fast discharge kinetics in the PDI-TEMPO electrolyte. During charging, the equilibrium potentials of the cell with PDI-TEMPO electrolyte are much lower, ascribed to fast kinetics of Li_2O_2 decomposition in PDI-TEMPO electrolyte (Fig. 3B and fig. S31). Moreover, the enlarged GITT curves in the bare DEGDME electrolyte show three distinguishable charge stages (10, 56). The first stage (0 to 0.15 mAh cm^{-2}) displays a linear increase in voltage during the charge scan, which could be ascribed to the delithiation of the Li_2O_2 in the initial charge process. The second stage (0.15 to 0.35 mAh cm^{-2}) presents a temporary increase but followed by a drop of the charge voltage during each charge scan, indicating a typical Li_2O_2 decomposition through the nucleation process. The third stage (0.35 to 0.5 mAh cm^{-2}) exhibits a flat increase in the voltage, which may originate from the mixed reactions of Li_2O_2 decomposition and parasitic reactions at high overpotential. The GITT curves in PDI-TEMPO, however, show no divided charge stages. A flat plateau is observed, without temporary increase in charge voltage at each charge step, owing to the lack of typical Li_2O_2 decomposition through the nucleation process. Therefore, the charge process in the Li-O_2 cell with PDI-TEMPO electrolyte proceeds through a different mechanism from those in the cell with bare DEGDME electrolyte. The

charge process is triggered and accelerated by the chemical reaction between oxidized TEMPO moieties in PDI-TEMPO molecules and solid Li_2O_2 , as well as the quenching reaction between PDI-TEMPO and released superoxide species. Therefore, it proves that PDI-TEMPO can efficiently accelerate the kinetics of the OER.

The charging reaction has been further studied by a quantitative in situ differential electrochemical mass spectrometry (DEMS) using less volatile tetraethylene glycol dimethyl ether (TEGDME) electrolyte, during which the O_2 evolved and the charge passed were monitored simultaneously. The oxygen evolution during the charge process with ether electrolyte has the same trend as the GITT results. A peak increase in the evolution at the initial stage is followed by a sharp decrease before the evolution increases again in the middle of the charge process (Fig. 3C). This corresponds to a typical Li_2O_2 decomposition through a nucleation process (10, 41, 56). The oxygen evolution in the PDI-TEMPO electrolyte shows a steady curve from the initial stage of charge. It proves that the chemical reaction between the oxidized PDI-TEMPO and Li_2O_2 dominates the charge process (Fig. 3D). The ratio of the charge passed and the O_2 evolved during charging of the Li-O_2 battery has been quantified to be $2.01 \text{ e}^-/\text{O}_2$, while for bare Li-O_2 cell, a $2.40 \text{ e}^-/\text{O}_2$ has been obtained. The enhanced charging reversibility confirms that PDI-TEMPO can also efficiently promote the charge process.

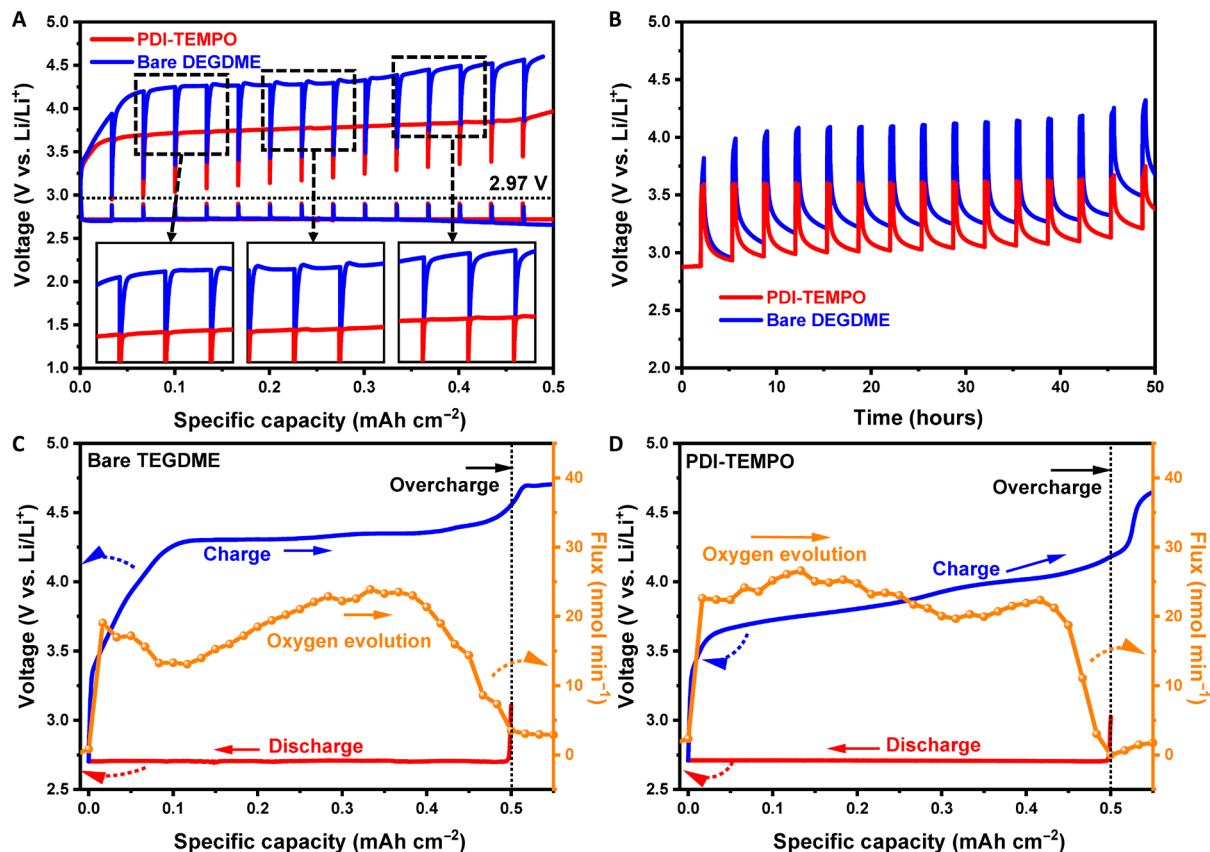


Fig. 3. Investigation of mechanisms within Li-O_2 batteries using PDI-TEMPO electrolyte. (A and B) GITT curves of the Li-O_2 batteries, which were acquired with a current density of 0.1 mA cm^{-2} for 0.5 mAh cm^{-2} and (A) a 60-min or (B) a 180-min time interval during the discharge and (B) charge processes. The insets are the enlarged images of selected areas. The concentration of PDI-TEMPO is 5 mM . (C and D) In situ DEMS analysis of the gas evolution during the charge process of Li-O_2 cell operation with (C) bare TEGDME electrolyte and (D) PDI-TEMPO electrolyte. The current density is 0.1 mA cm^{-2} .

PDI-TEMPO molecules to suppress parasitic reactions

The cycling performance of Li-O₂ cells containing PDI-TEMPO molecules has been markedly improved (Fig. 4, A and B, and figs. S32 and S33), because PDI-TEMPO molecules act as an O₂^{•-} quencher to eliminate active O₂^{•-} species and expedite both discharge and charge processes. Compared with only 39 cycles delivered by the Li-O₂ cell with bare DEGDME electrolyte, the cell with PDI-TEMPO electrolyte has shown a stable cycling performance over 300 cycles (Fig. 4B). As shown in Fig. 4A, the cell with PDI-TEMPO electrolyte displays stable charge potentials around 3.7 V during cycling, which is about 0.7 V lower than the cell with bare DEGDME electrolyte. The excellent electrochemical performance should be ascribed to the unique functions of PDI-TEMPO, which can efficiently diminish parasitic reactions and promote discharge and charge processes. To prove this point, we performed postmortem FTIR and x-ray diffraction (XRD) characterizations on the cathodes after cycling (Fig. 4, C to F). The discharge products were dominated by Li₂O₂ after the first discharge with PDI-TEMPO electrolyte, because strong FTIR peaks were found in the range of 400 to 700 cm⁻¹, and no peaks were spotted in the range of 1300 to 1700 cm⁻¹ in either electrolyte. These FTIR peaks disappeared after the first charge process, indicating highly reversible formation and decomposition of Li₂O₂ during discharge and charge processes. However, the FTIR peaks at 400 to 700 cm⁻¹ in bare DEGDME electrolyte remained in the charge process after five cycles, and the peaks associated with Li₂CO₃ in the range of 1300 to 1700 cm⁻¹ increased gradually with cycling (Fig. 4C). This confirms that a large quantity of Li₂CO₃ accumulated after several cycles in

the bare DEGDME electrolyte. On the contrary, the FTIR peaks in the range of 400 to 700 cm⁻¹ reversibly appeared and disappeared after discharge and charge processes in PDI-TEMPO electrolyte, corroborating that Li₂O₂ is the dominant discharge product (Fig. 4D). The XRD patterns of the corresponding cathodes also confirm the same conclusion. The XRD peaks of Li₂O₂ became weaker on cycling and eventually disappeared after 10 cycles when using bare DEGDME electrolyte (Fig. 4E). The XRD peaks of Li₂O₂ remained visible after 25 cycles in PDI-TEMPO electrolyte (Fig. 4F).

The accumulation of by-products in the bare DEGDME electrolyte originates from parasitic reactions of active oxygen species (O₂^{•-} and ¹O₂) and the decomposition of electrolyte solvent. The continuous attack by O₂^{•-} and ¹O₂ to solvent molecules and cathode materials lead to the formation of a substantial amount of by-products during cycling. Moreover, high charging voltage is required to decompose both Li₂O₂ and by-products, resulting in the further decomposition of electrolyte solvent and generation of more ¹O₂. This vicious cycle continuously produces by-products and eventually terminates the Li-O₂ batteries. It is worth noting that adding TEMPO, a commonly used redox mediator, in the DEGDME electrolyte to lower the charge potential can also suppress parasitic reactions associated with large overpotentials to a certain extent during the charge process (fig. S34). It has recently been reported by Petit *et al.* and Liang *et al.* that the use of redox mediators could accelerate the relaxation of ¹O₂, making them a potential quencher for ¹O₂ (41, 57). However, we found that the cell with TEMPO redox mediator could only last for 60 cycles, and by-products still quickly accumulated during cycling (fig. S34).

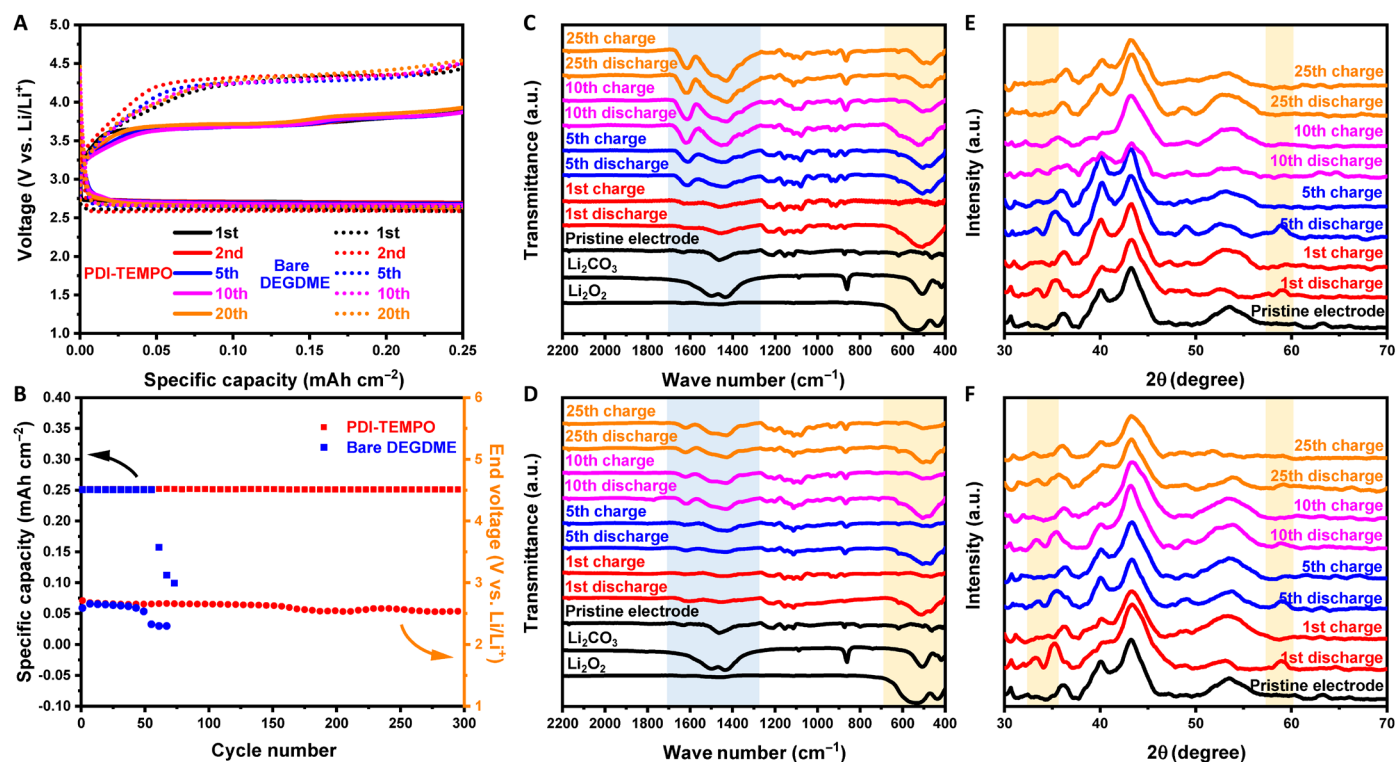


Fig. 4. Postmortem characterizations of electrodes after cycling. (A) The discharge-charge profiles and (B) cycling performance of Li-O₂ batteries with PDI-TEMPO electrolyte and DEGDME electrolyte. The reversible capacities are restricted to 0.25 mAh cm⁻². The current densities are 0.1 mA cm⁻², and the cutoff voltage is 2.3 V/4.6 V. The concentration of PDI-TEMPO is 5 mM. (C and D) Postmortem FTIR spectra of electrodes after cycling with (C) bare DEGDME electrolyte and (D) PDI-TEMPO electrolyte. (E and F) Postmortem XRD patterns of the electrodes after cycles with (E) bare DEGDME electrolyte and (F) PDI-TEMPO electrolyte. The highlighted blue areas are the places where the characteristic peaks for Li₂CO₃ appear, and the highlighted orange areas are the places where the characteristic peaks for Li₂O₂ appear.

This specifies that parasitic reactions associated with active oxygen species attack during the discharge and charge processes remained severe. The rational design of PDI-TEMPO molecules can efficiently lower the charge overpotentials and quench $O_2^{\bullet-}$ species, both of which are deemed as the origins of 1O_2 generation. Furthermore, TEMPO moieties are also reported to accelerate the relaxation of 1O_2 during the operation of Li- O_2 batteries, and two TEMPO moieties bridged by PDI backbone in one molecule probably enhance their overall capability in quenching 1O_2 (41, 57). Therefore, PDI-TEMPO molecules are able to suppress the generation of 1O_2 as well, leading to the comprehensive suppression of parasitic reactions. To verify this assertion, we conducted a demonstration experiment by adding 20 mM 9,10-dimethylantracene (DMA) in electrolytes as the trap agent for 1O_2 . The results in fig. S35 show that DMA is quickly consumed in DEGDME electrolyte over cycling, while TEMPO electrolyte displays a much lower consumption rate of DMA. The consumption rate of DMA in PDI-TEMPO electrolyte is the slowest, which almost equals the self-degradation rate of DMA in O_2 atmosphere. This result unambiguously verifies that PDI-TEMPO molecules can not only quench superoxide radicals and lower charge overpotentials but also suppress the generation of 1O_2 , leading to substantially reduced parasitic reactions.

Cycling performance of Li- O_2 batteries with PDI-TEMPO molecules

Unlike previously reported redox mediators that can easily shuttle to the negative electrode and react with the highly active lithium metal, PDI-TEMPO molecules can effectively overcome this issue in Li- O_2 batteries. Our rationally designed PDI-TEMPO molecules

contain a conjugated PDI backbone and two bulky TEMPO moieties. The steric hindrance between these two segments results in a twisted stereostructure of PDI-TEMPO molecule, in which TEMPO moieties are not on the same plane as the PDI backbone (Fig. 1B and fig. S36, A and B). This distinctive stereostructure enables PDI-TEMPO molecules to be restricted in the cathode region by van Der Waals forces via π - π interactions between the conjugated PDI backbone and the conjugated graphitic carbon in the cathodes (fig. S36C) (42, 47, 50). Meanwhile, such π - π interactions are weakened by the steric hindrance between the graphitic carbon and bulky TEMPO moieties, preventing PDI-TEMPO molecules from depositing on the cathode (Fig. 5A and fig. S36C). Consequently, PDI-TEMPO molecules are still in the electrolyte, but prohibited from migrating to lithium anode and reacting with lithium metal. Moreover, because PDI-TEMPO molecules quench the reduced oxygen species immediately after they are generated, these species are also restricted to the cathode region to form discharge products (fig. S26). Therefore, PDI-TEMPO molecules can also strongly inhibit parasitic reactions on lithium metal anodes from these small molecules. To verify this, lithium metal anodes were characterized by SEM after cycling. A comparison experiment was conducted by dissolving 10 mM TEMPO in DEGDME electrolyte. As shown in fig. S37, the lithium foil retrieved from TEMPO electrolyte is severely corroded, caused by the direct reactions between lithium metal and oxidized TEMPO molecules. The lithium foil in the bare DEGDME electrolyte also shows a corroded morphology (Fig. 5B), which originates from the parasitic reactions between lithium metal and the dissolved small molecules such as oxygen species (fig. S38). On the contrast, the lithium foil in PDI-TEMPO electrolyte shows no obvious corrosion after cycling

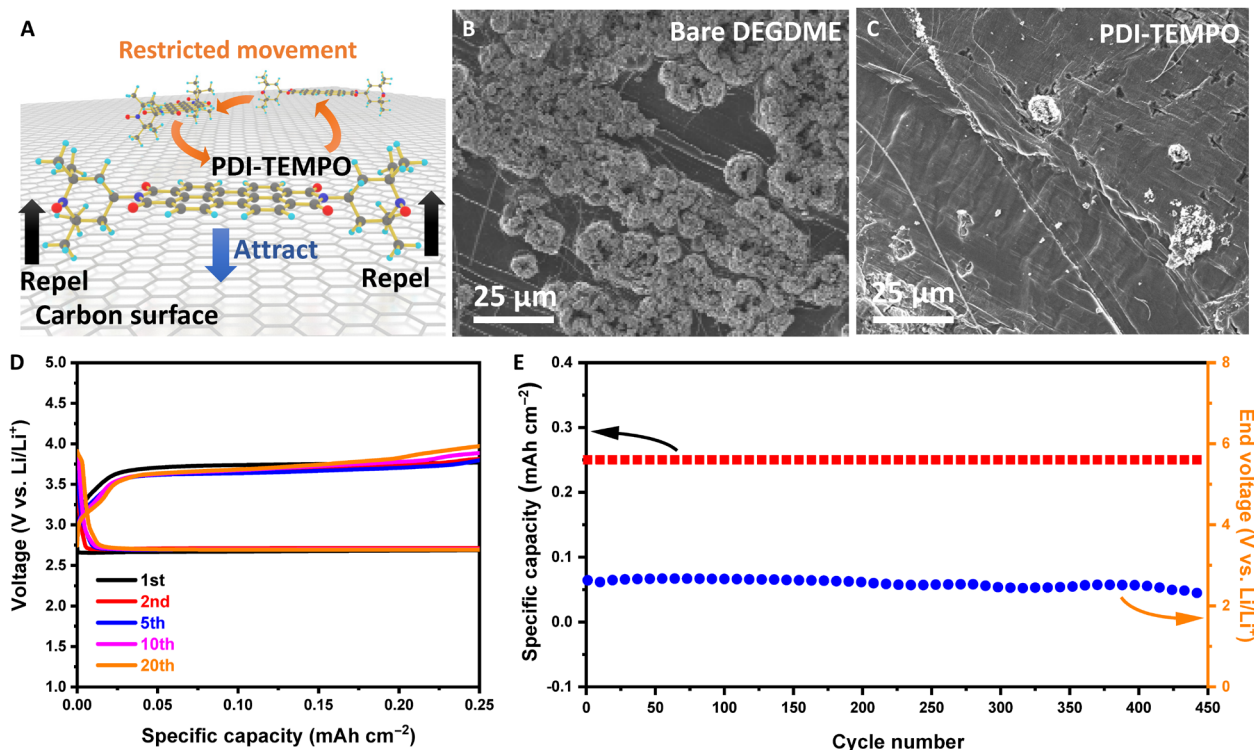


Fig. 5. Influences of PDI-TEMPO on lithium metal anode and cycling performance. (A) Schematic illustration of the interaction between PDI-TEMPO and graphitic carbon. (B and C) SEM images of the lithium metal anodes from (B) bare DEGDME electrolyte and (C) PDI-TEMPO electrolyte after 10 cycles. (D) Discharge-charge profiles and (E) cycling performance of the Li- O_2 batteries with 10 mM PDI-TEMPO electrolyte. The current density is 0.2 mA cm^{-2} .

(Fig. 5C), demonstrating that PDI-TEMPO molecules can suppress parasitic reactions caused by the dissolved oxygen species. Therefore, further increasing the concentration of PDI-TEMPO in the electrolyte until 10 mM should not deteriorate the electrochemical performance (concentration optimization in fig. S39). Instead, high concentration of PDI-TEMPO enhances its capability for quenching $O_2^{\cdot-}$ and suppressing 1O_2 , especially at elevated current densities (fig. S40). Hence, we subsequently doubled the concentration of PDI-TEMPO to 10 mM in electrolyte. As shown in Fig. 5D, the Li- O_2 cell with 10 mM PDI-TEMPO electrolyte presents low overpotentials and a significantly extended cycle life of more than 450 cycles (Fig. 5E and fig. S41). We further incorporated a Nafion membrane on lithium metal to prevent the dendrite growth and a porous graphene electrode to accommodate more Li_2O_2 (figs. S42 to S44). The Li- O_2 cell with PDI-TEMPO reagent has exhibited an ultralong cycle life exceeding 1400 cycles (fig. S45) (58).

DISCUSSION

In summary, a multifunctional PDI-TEMPO quencher has been rationally designed and synthesized for boosting the performance of Li- O_2 batteries. The PDI-TEMPO molecule contains a PDI backbone with the capability of quenching $O_2^{\cdot-}$, and redox mediator-active TEMPO moieties to catalyze both discharge and charge processes in the electrolyte solution. When added in the DEGDME electrolyte, the operation of the Li- O_2 cell undergoes a new chemical-quenching reaction mechanism, leading to the formation of Li_2O_2 nanoparticles. Furthermore, the unique twisted molecular structure confines PDI-TEMPO molecules in the cathode region, which can effectively prevent corrosion of lithium metal anode. Consequently, Li- O_2 cells containing PDI-TEMPO quencher demonstrated an increased discharge capacity, improved energy efficiency, diminished parasitic reactions, and long cycle life.

MATERIALS AND METHODS

Synthesis of PDI-TEMPO

The detailed synthesis process of *N,N'*-di-(2,2,6,6-tetramethyl-1-oxyl-4-piperidinyl)-perylene-3,4,9,10-tetracarboxylic diimide (PDI-TEMPO) is illustrated in fig. S1A (42, 43).

In brief, 0.51 ml (3 mmol) of 4-amino-2,2,6,6-tetramethylpiperidine (Sigma-Aldrich), 0.588 g (1.5 mmol) of perylene-3,4,9,10-tetracarboxylic dianhydride (Sigma-Aldrich), and 0.279 g (1.5 mmol) of zinc acetate (Sigma-Aldrich) were combined with 25 g of imidazole (Sigma-Aldrich) in a round bottle flask. The flask was evacuated and refilled with argon three times and then heated with a sand bath at 120°C for 3 hours, 160°C for 3 hours, 200°C for 3 hours, and, lastly, 240°C for 48 hours. After the mixture cools down to ~80°C, 200 ml of $CHCl_3$ was carefully added to the flask. The clear red solution was extracted by separatory funnel and washed five times with 50 ml of water. The obtained organic phase was then collected, dried over $MgSO_4$, and concentrated under reduced pressure to yield a purple solid (PDI-TEMPH).

The resulting compound PDI-TEMPH was further oxidized by dissolving in methanol at room temperature together with a catalytic amount of $Na_2WO_4 \cdot 2H_2O$ (Sigma-Aldrich) and 5 ml of 30% H_2O_2 (ChemSupply) (37, 48). The mixture was stirred at room temperature for 24 hours. After the reaction, the solvents were removed through Rotavapor (Buchi R-100), and isopropanol (Sigma-Aldrich) was

added to dissolve the crude product. After filtration, the filtrate was collected and dried under vacuum at room temperature. PDI-TEMPO was obtained as a purple-reddish solid. The solid was dried under vacuum for 24 hours before storing in the argon-filled glovebox. PDI-TEMPO can also be prepared by the direct reaction between 4-amino-2,2,6,6-tetramethylpiperidine-1-oxyl and perylene-3,4,9,10-tetracarboxylic dianhydride via a similar process, without the oxidation process.

Characterizations

The 1H nuclear magnetic resonance spectra were recorded on an Agilent 500 Spectrometer at 25°C in *d*-chloroform. The infrared spectroscopy measurements were conducted on a Nicolet Magna 6700 FT-IR spectrometer. All spectra were obtained using 4- cm^{-1} resolution and 64 scans at room temperature. The spectra of ATR-FTIR were obtained using 4- cm^{-1} resolution and 16 scans at room temperature with argon protection. A field emission scanning electron microscope (Zeiss Supra 55 VP) was used to observe the electrode morphologies. Nitrogen-sorption measurements were carried out at 77 K with a Micromeritics 3Flex surface characterization analyzer. The specific surface area was calculated by the Brunauer-Emmett-Teller (BET) method. UV-Vis spectroscopy was conducted through an Agilent Cary 60 UV-Vis spectrophotometer, scanning from 800 to 200 nm. All the electrospray ionization-MS spectra were recorded on Waters liquid chromatography-MS system with the condition setting as follows: capillary voltage, 3.0 kV; extractor voltage, 2.0 V; RF lens, 0.5 V; source temperature, 120°C; desolvation temperature, 200°C; desolvation gas flow, 500 liter hour $^{-1}$; pump flow, 20 μl min $^{-1}$. Sample concentration was 1 to 5 mg μl^{-1} in MeOH, and the cone voltage was varied from 15 to 80 V depending on the samples. EPR was collected using a Bruker EMX X-Band ESR Spectrometer. EPR spectra were obtained under 120 K and then normalized on the basis of the weight of the samples used for measurement. XRD measurement was performed on a Bruker D8 X-ray diffractometer using $Cu K\alpha$ radiation. For the XRD analysis, the electrodes from the disassembled cells were washed and dried first and then sealed with "Parafilm" to exclude moisture and carbon dioxide from the discharge products, which are very sensitive to normal atmospheric air components.

Electrochemical characterizations

All the electrochemical characterizations were conducted on a CH Instruments 660E electrochemical workstation and BioLogic VMP3 potentiostat. The CV for two-electrode configuration was operated by using a lithium metal foil as the anode, glass fibers as separator soaking with electrolyte, and a porous electrode as the cathode. Carbon paper electrodes were prepared by stacking three layers of pre-cut carbon papers (10-mm diameter). CNT cathodes were prepared by casting the slurry of CNT and polytetrafluoroethylene (PTFE) (8:2) in iso-propylene/water on the pre-cut carbon paper and dried before use. Porous graphene electrodes were prepared via the same method (59). The electrolyte was prepared by dissolving 5 mM PDI-TEMPO in DEGDME with 0.5 M LiTFSI. The scan rate was 0.5 mV s $^{-1}$. The CV for three-electrode configuration was operated with a Pt wire as the counter electrode, an Ag wire as the reference electrode, and a GC electrode as the working electrode. The scan rate was 100 mV s $^{-1}$.

The discharge/charge performances were evaluated by a Neware Battery Testing System. The discharge-charge performances were

evaluated by assembling Li-O₂ batteries. A two-electrode Swagelok-type cell with an air hole (0.785 cm²) on the cathode side was used to test the electrochemical performances. A two-electrode coin cell (CR2033, Neware) with several open holes on the cathode side was used for long-term cycling. The cells were assembled in an argon-filled glove box with water and oxygen level less than 0.1 parts per million. Lithium foils were used as anodes, and polyvinylidene difluoride (PVDF)/glass fibers were used as separators. The lithium foils were immersed in propylene carbonate solution containing 5% fluoroethylene carbonate for 5 days before use. A Nafion membrane was used between the lithium anode and separator for long-term cycling. The electrolytes were prepared by dissolving 0.5 M LiTFSI in DEGDME with 5 mM PDI-TEMPO. The typical amount of electrolyte used in a single cell was 150 μl. The assembled cell was gas-tight except for the cathode side window, which is exposed to the oxygen tank. All measurements of Li-O₂ batteries were conducted in a gas tank of 1-atm dry oxygen atmosphere. The capacities of the batteries were calculated on the basis of the total area of the electrodes. GITT was performed using an Li-O₂ cell at a current density of 0.1 mA cm⁻². A 1-hour/3-hour rest step was given after 10 min of each discharge and charge step. The cutoff capacity was set to 0.5 mAh cm⁻².

A modified Swagelok-type Li-O₂ cell was linked to a commercial magnetic sector mass spectrometer (Thermo Fisher Scientific) by a specially designed gas-purging system for in situ DEMS measurement. The flow rate of purge gas was set at 1 ml min⁻¹. During the charge process, high-purity Ar was used as the carrier gas. The DEMS cells were assembled in the argon-filled glove box for the electrochemical testing.

The RRDE measurements were performed using an RRDE with a GC disk and a Pt ring (Pine Research Instrumentation, USA) (55). A homemade three-electrode configuration was used with Pt wire as the counter electrode, Ag wire as the reference electrode, and RRDE as the working electrode. The electrolytes used for the measurements were the DEGDME electrolyte and PDI-TEMPO electrolyte. To prevent potential mask effect by the redox current of PDI-TEMPO, the PDI-TEMPO electrolyte was prepared by dissolving 0.5 mM PDI-TEMPO in the DEGDME electrolyte. The rotation speed of the RRDE was held at 1600 rpm for the tests. To ensure the generation of currents for OER and ORR from both electrolytes during the tests, an oxidation potential was held constantly at 4.1 V versus Li/Li⁺, and a reduction potential was held constantly at 2.3 V versus Li/Li⁺. As the currents generated for OER come from the consumption of Li₂O₂, which was generated during the ORR process, the tests of the OER efficiency were conducted straight after tests of the ORR efficiency. The tests of the ORR efficiency were carried out in the oxygen atmosphere, and the ones of the OER efficiency were in the argon atmosphere. To further eliminate the currents from side reactions and self-redox reactions, a control experiment was carried out using the same setting and only operated all the measurements in the argon atmosphere. The calculation of the efficiency could be corrected by the equation below

$$\text{Efficiency} = (I_{\text{ring}} - I_{\text{ring-corrected}}) / (I_{\text{disk}} - I_{\text{disk-corrected}}) \quad (6)$$

The I_{ring} and I_{disk} are the currents obtained on the ring and disk electrodes during the ORR and OER processes. The $I_{\text{ring-corrected}}$ and $I_{\text{disk-corrected}}$ are the currents obtained on the ring and disk electrodes

during the control experiment. More details will be found in the illustration in figs. S20 and S21.

Theoretical calculations

The HOMO and LUMO in this work were performed with the Gaussian 09 program, through the conventional DFT calculations using the hybrid exchange-correlation functional B3LYP on the molecules. The Basis Set was triple-zeta valence with polarization (TZVP) (60).

Demonstration experiments

The demonstration experiments of the scattering of Li₂O₂ during the discharge process were performed by assembling the Li-O₂ cell through a special configuration. A carbon paper was added between the lithium anode and the glass fiber separator. CNT electrodes were used as cathodes in the cell. The cell was directly discharged to 2.0 V at a current density of 0.1 mA cm⁻². The inserted carbon paper was taken out for SEM, and the glass fiber was taken out for phenolphthalein tests. The phenolphthalein tests were performed by first soaking the glass fiber into 3 ml of water. Then, two drops of freshly made phenolphthalein/ethanol solution were added to the aqueous solution dropwise. The color-changing process was recorded by video and photoshoots.

The demonstration of the capacity loss during the charge process was performed by changing the electrolyte/separator after the discharge process. The Li-O₂ cells with different electrolyte configuration were first discharged to 0.25 mAh cm⁻² at a current density of 0.1 mA cm⁻². The cells were then taken to an argon-filled glove box and disassembled. The cathodes were washed with clean DEGDME solvent three times and dried under vacuum. The dried cathodes were then assembled into Li-O₂ cells with fresh lithium metal anodes, a glass fiber separator soaked with DEGDME electrolyte. The as-assembled cells were then directly charged to 4.6 V at a current density of 0.1 mA cm⁻².

The demonstration of the suppression of ¹O₂ during cycling was performed by adding 20 mM DMA in all the electrolytes. To avoid possible reactions between lithium metal and DMA, an LiFePO₄/C electrode was used instead of the lithium metal foil as the anode. The Li-O₂ cells were cycled with a capacity of 0.25 mAh cm⁻² at a current density of 0.1 mA cm⁻². The cells were then taken to an argon-filled glove box and disassembled. All the battery components were soaked in 9 ml of DEGDME solvent. UV spectra were performed on the extracted solution. A reference was prepared by resting the bare DEGDME electrolyte with 20 mM DMA in oxygen for the same time of each cycling period to monitor the self-degradation of DMA in O₂.

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at <https://science.org/doi/10.1126/sciadv.abm1899>

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