

Manipulating Redox Kinetics of Sulfur Species Using Mott-Schottky Electrocatalysts for Advanced Lithium-Sulfur Batteries

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ABSTRACT: Lithium-sulfur (Li-S) batteries suffer from sluggish sulfur redox reactions under high-sulfur-loading and lean-electrolyte conditions. Herein, a typical Co@NC heterostructure composed of Co nanoparticles and semiconductive N-doped carbon matrix is designed as a model Mott-Schottky catalyst to exert the electrocatalytic effect on sulfur electrochemistry. Theoretical and experimental results reveal the

redistribution of charge and a built-in electric field at the Co@NC heterointerface, which are critical to lowering the energy barrier of polysulfide reduction and Li₂S oxidation in discharge and charge process, respectively. With Co@NC Mott-Schottky catalysts, the Li-S batteries display ultrahigh capacity retention of 92.1 % and a system-level gravimetric energy density of 307.8 Wh kg⁻¹ under high S loading (10.73 mg cm⁻²) and lean electrolyte (E/S = 5.9 μL mg_{sulfur}⁻¹) conditions. The proposed Mott-Schottky heterostructure in this work not only deepens the understanding of electrocatalytic effect in Li-S chemistry, but also inspires rational catalysts design for advanced high-energy-density batteries.

KEYWORDS: Mott-Schottky heterostructure, lithium-sulfur batteries, catalytic polysulfide conversion, lean electrolyte conditions

Rechargeable lithium-sulfur (Li-S) batteries have recently drawn tremendous academic and industrial interests due to the natural abundance of sulfur feedstock and ultrahigh intrinsic energy density ($\approx 2600 \text{ Wh kg}^{-1}$).^{1, 2} Such high specific energy derives from sulfur-related redox reactions that involve the reduction of solid sulfur molecules (S₈) to form solid lithium sulfide (Li₂S) with a series of dissolvable lithium polysulfides (LiPSs) intermediates on discharge and a reversible oxidation process on charge.^{3, 4} However, this multi-phase and multi-electron conversion chemistry faces some intrinsic challenges, including undesirable “shuttling” behavior of polysulfides and sluggish sulfur redox reactions.^{5, 6} Introducing electrocatalyst into sulfur cathode has

shown great potential in expediting polysulfides reduction in charge process^{7, 8} and/or Li₂S oxidation in discharge process^{9, 10}, thus efficiently suppressing polysulfide diffusion out of the cathode. Benefiting from this electrocatalysis effect, the lifespan and capacity of Li-S batteries have been updated to an acceptable level (>1000 cycles and > 1300 mAh g⁻¹) over the past few years.^{11, 12} These achievements, however, are usually obtained under a low sulfur loading (< 4 mg cm⁻²) and excess electrolyte (E/S > 15 μL mg_{sulfur}⁻¹), leading to low cell-level energy density (often < 200 Wh kg⁻¹).^{13, 14} Moreover, the exploration and discovery of catalysts mainly rely on trial and error, and the underlying catalytic mechanism in Li-S conversion chemistry still lacks deep mechanistic understanding.^{15, 16}

In Li-S electrochemistry, the kinetics of conversion reactions among sulfur species are strongly determined by surface adsorption and deposition as well as charge/mass transfer at the three-phase interface among catalysts, sulfur species, and electrolytes.¹⁷ These heterogeneous reactions intrinsically depend on the catalyst's surface electronic states. For most inorganic materials (such as N-doped C,¹⁸ CoP,^{15, 19} TiO₂,²⁰ MoS₂,²¹ and C₃N₄^{22, 23}), although they can immobilize polysulfides through a chemisorption mechanism, their semiconducting nature jeopardizes the electron transport to the surface-bonded polysulfide, thus slowing down the subsequent redox reactions of sulfur species (Figure 1a).²⁴ To this end, coupling these semiconductors with metallic phases as Mott-Schottky heterojunctions could be a proof-of-concept method to boost the catalytic activity by optimizing interfacial electronic interactions (Figure 1b). From basic semiconductor physics, the different energy structures between metal and

semiconductor can drive the flow of electrons across the interface until the heterojunctions reach the thermodynamic equilibrium state.^{25, 26} This manipulation of electron transfer in Mott-Schottky heterojunction can induce charge separation and an internal electric field at the interface, which not only propels the charge transfer and ion diffusion in lithium-ion batteries²⁷ but also modulates the energy barriers of photo/electrocatalytic reactions^{25, 28}. Despite these advances, catalytic understanding of Mott-Schottky heterostructure for Li-S conversion chemistry is rarely explored, yet critical to electrocatalysts design for multi-electron sulfur-based battery technology.

In this contribution, to investigate the Mott-Schottky effect on conversion chemistry in Li-S batteries, we put forward a paradigmatic Co@NC heterostructure consisting of Co nanoparticles encapsulated in a porous N-doped carbon matrix as a sulfur cathode host. In addition to the high electrical conductivity, polar reaction with LiPSs and smooth Li-ion surface diffusion, the metallic character of cobalt also can derive spontaneous rearrangement of the NC substrate's energy band, which facilitates charge transfer and induces a strong built-in electric field at the interface of the Co@NC heterostructure. These interfacial merits of Mott-Schottky electrocatalysts are beneficial to strengthening adsorption and conversion of polysulfides as well as lowering the energy barriers of sulfur reduction in the discharge process and Li₂S oxidation in the charge process. Consequently, the thus-derived Co@NC electrocatalyst renders Li-S batteries with remarkable cyclicality and gravimetric energy density under harsh test conditions, such as large current density of 4 C, ultrahigh sulfur loading of 10.73 mg cm⁻², low electrolyte/sulfur ratio of 5.9 μL mg⁻¹. The present work depicts the

intriguing potential of rationally designed Mott-Schottky electrocatalysts for advanced Li-S batteries.

RESULTS AND DISCUSSION

The Co@NC heterostructure was synthesized through a facile metal-organic framework-templated method (see details from Experiment section). Scanning electron microscopy (SEM) images in Figure S2 show that the obtained Co@NC composites possess a two-dimensional (2D) leaf-like morphology with a length of $\sim 5 \mu\text{m}$, a width of $\sim 3 \mu\text{m}$, and a thickness of $\sim 200 \text{ nm}$. From a transmission electron microscopy (TEM) observation (Figure 2a and S3), one can discern the uniform distribution of cobalt nanoparticles with an average diameter of $\sim 10 \text{ nm}$ on the porous and rough surface of the 2D products. Such a hierarchical structure has a large surface area ($410.7 \text{ m}^2 \text{ g}^{-1}$) and an average mesopore size ($\sim 4.5 \text{ nm}$) (Figure S4), which is beneficial for loading ample sulfur. The high-resolution TEM observation (Figure 2b) illustrates the encapsulation of Co nanoclusters by graphitic carbon nanocages, where the lattice fringes of 0.20 nm and 0.34 nm account for the (111) plane of metallic cobalt and (002) plane of graphitic carbon, respectively.²⁷ X-ray diffraction and Raman analysis (Figure S5 and S6) further confirm the coexistence of Co and graphitic carbon in the Co@NC composites. Additionally, the high-angle annular dark-field (HAADF) and the corresponding energy-dispersive X-ray spectroscopy (EDX) investigations (Figure 2c) imply the uniform distribution of Co nanoparticles inside the porous nitrogen-carbon framework. The weight percentage of Co nanoparticles is determined to be 11.6% by thermogravimetric analysis (TGA) (Figure S7).

The electronic environments of each constituent element in the thus-obtained Co@NC heterostructure were examined by X-ray photoelectron spectroscopy (XPS). As depicted in Figure S8b, the high-resolution Co 2p_{3/2} spectrum can be divided into two sub-bands at 778.5 eV and 780.3 eV, which correspond to metallic Co and Co-N species, respectively.²⁹ The N 1s spectrum in Figure 2d is resolved into pyridinic N (397.9 eV), pyrrolic N (400.4 eV), and graphitic N (401.1 eV).³⁰ The successful incorporation of N into the carbon matrix can also be confirmed by the C 1s spectrum (Figure S8d), in which a peak at 285.3 eV characteristic of C-N bonding is fitted. For comparison, pure Co metal and pure N-doped graphitic carbon (NC) counterparts (Figure S9 and S10) were also tested under identical conditions. We note that there is no characteristic Co-N peak for the pure Co sample (Figure S8b), which supports the interaction of Co and N species and accordingly positively charged Co species in Co@NC junction. Notably, the pyridinic N and C-N peaks of the NC sample are positively shifted to 398.1 and 285.6 eV in comparison with those of the Co@NC (Figure 2d and S8c).³¹ Namely, NC species in the Co@NC compound have a negative charge after metallic Co decoration. The above XPS results strongly reveal the electronic interactions between Co and NC, causing the injection of electrons from Co to NC at their coupling interfaces.

The surface electron transfer in Co@NC heterostructure is strongly affected by the work functions of the Co and NC, according to the band theory of solids.²⁶ Thus, the work function of the two materials was calculated using ultraviolet photoelectron spectroscopy (UPS) (Figure S11). Figure 2e shows that the work function value of

Co@NC (5.88 eV) is much smaller than that of NC (6.19 eV), evidencing the regulated electronic structure of NC substrates after the introduction of metallic Co. It is well known that N-doped C has a p-type semiconductor structure, since the N dopant can enlarge the bandgap of carbon support and thus give rise to lower valence-band maximum (E_v) and higher conduction-band bottom (E_c).²⁶ Hence, when a semiconducting NC is directly contacted with metallic Co with a smaller work function (5.00 eV), the electrons from Co will spontaneously move to NC until the same Fermi levels are reached on both sides. This self-driven charge density redistribution on the heterointerfaces is also known as the Mott-Schottky effect in solid-state physics.²⁸

The behavior of electron transfer between Co and NC was further theoretically elucidated by Bader charge analysis. The simulated Co@NC heterostructure model in Figure 2f and 2g explicitly illustrates the strong interfacial polarization between Co and NC and accordingly injection of electrons from surface Co atom to adjacent N atoms in NC, thus resulting in a built-in electric field pointing towards NC, which agree well with the experimental results (Figure 2h). In Li-S chemistry, this structural merit contributes to the chemical LiPSs-binding ability of Co@NC heterostructure, since the negatively charged NC side can improve adsorption of Li^+ while the positively charged Co side is prone to adsorb polysulfide anions.³² This conclusion is confirmed by a visual Li_2S_6 adsorption experiment (Figure S12) and additional XPS analyses of Co@NC/ Li_2S_6 composites (Figure S13). This strengthened the interaction between Co@NC and polysulfides by Mott-Schottky effect is important for inhibiting the polysulfides shuttling and accelerating their subsequent conversion in Li-S batteries.

To determine the effectiveness of Mott-Schottky heterostructure in boosting polysulfide-involving reactions, cyclic voltammetry (CV) and electrochemical impedance spectroscopy (EIS) of symmetrical cells based on Co@NC and NC were analyzed in Li_2S_6 electrolyte. As exhibited in Figure 3a, four distinct redox peaks located at -0.095 (Peak A), -0.212 (peak B), 0.099 (peak C), and 0.198 V (peak D) can be detected from the CV of Co@NC cell. In contrast, the CV of NC cell only exhibits an inconspicuous pair of redox peaks with large voltage separation and low CV current. The profile discrepancy of CV results strongly validates the kinetics advantage of Co@NC for polysulfide redox conversion over NC. The EIS results (Figure S15) further reveal the high catalytic capability of Co@NC that can reduce the charge transfer resistance (R_{ct}) at the Co@NC/polysulfides interface. Subsequently, the kinetic evaluation of liquid-solid conversion from polysulfide to Li_2S was monitored by Li_2S precipitation in a potentiostatic discharge experiment.³³ As observed from the I-t curves in Figure 3b, the Li_2S nucleation peak possesses a higher current of 1.26 A g^{-1} after 310 s on the Co@NC electrode over that of NC electrode (0.41 A g^{-1} after 1010 s), indicating that the Co@NC electrocatalyst significantly improved the kinetics of Li_2S nucleation. The early nucleated Li_2S is of great importance to realize uniform distribution of Li_2S precipitation on Co@NC frameworks, which helps in maintaining high utilization of sulfur during cycling.¹⁷ The solid-liquid conversion from Li_2S to polysulfide was performed likewise by a potentiostatic charge method (Figure 3c).³³ Similarly, the Co@NC electrode exhibits shorter nucleation times (1850 s) and higher charging current responses (0.82 A g^{-1}) compared to NC counterpart (0.48 A g^{-1} after 2760 s),

pointing to propelled Li_2S dissolution kinetics on Co@NC catalysts. Primarily, the results signify that the Mott-Schottky effect on Co@NC heterojunction plays an essential role in electrocatalysis for all phase polysulfide-involving processes.

The superior electrocatalytic activity and hierarchical porous structure render the Co@NC Mott-Schottky electrocatalyst to be an auspicious cathode host for coin-type Li-S cells. In the resulting Co@NC/S composites, the sulfur content was controlled at 74.5 wt % and the close contact between S and Co@NC can promote the Mott-Schottky catalytic effect (Figure S16-S18). Figure 3d shows the CV profile of the Co@NC/S cathode at a scan rate of 0.1 mV s^{-1} . Two cathodic peaks appear at 2.308 and 2.025 V, which are related to the reduction of S_8 molecules to polysulfides (peak C1) and their further reduction to solid-state Li_2S (peak C2). The anodic peak (peak A) at 2.404 V is associated with the multistep reconversion of Li_2S eventually to S_8 .³⁴ The CV curves almost overlap during the first three cycles (Figure S19), demonstrating great reversibility of the sulfur redox reactions. With respect to the NC@S cathode, it exhibits the lower redox peak current and larger polarization between peak C1 and A as compared with the Co@NC/S cathode, suggesting a more rapid sulfur conversion kinetics in working Li-S batteries owing to the Mott-Schottky effect of Co@NC catalyst.

To quantify the catalytic activity of the Co@NC catalyst, the Tafel plots were calculated according to corresponding CV curves. As shown in Figure S20, the Co@NC/S cathodes are characterized by lower Tafel slopes for both Li_2S_n -to- Li_2S (69.2 mV dec^{-1}) and Li_2S -to- Li_2S_n (63.5 mV dec^{-1}) steps than those of the NC/S

cathodes (106.0 and 103.1 mV dec⁻¹, respectively), further demonstrating the significant kinetic promotion of sulfur species in both charging and discharging processes upon Mott-Schottky effect.³⁵ Activation energy (E_a) is also an important fundamental parameter for evaluating Li-S conversion chemistry.^{16, 36} Accordingly, we fitted the charge-transfer resistance measured at various temperatures into the Arrhenius equation and the E_a was derived to be 46.3 and 67.9 kJ mol⁻¹ for Co@NC/S NC/S cathode, respectively (Figure S21).⁷ Pertaining to the reduction of Li_2S_n into Li_2S , the rate-limiting step in the discharge process, the E_a difference between the Co@NC/S and NC/S cathodes is 23.9 kJ mol⁻¹ based on the Tafel analysis (see details in the Experiment section) (Figure 3e).³⁶ Likewise, the calculated E_a of the Li_2S oxidation process for the Co@NC/S cathode is 28.6 kJ mol⁻¹ smaller than that for the NC/S cathode. These dramatically reduced E_a values indicate that the reversible reactions between polysulfides and Li_2S are thermodynamically more favorable on the Co@NC Mott-Schottky junction.

The fact that sulfur redox reactions in Li-S batteries are always accompanied by Li-ion transfer encourages us to study the diffusivity of Li^+ at the Co@NC interface. Through CV testing at 0.1 to 0.5 mV s⁻¹, the lithium-ion diffusion coefficient (D_{Li^+}) can be calculated to be 6.17×10^{-8} , 7.55×10^{-8} , and 2.49×10^{-7} cm² s⁻¹ for the Co@NC/S cathode at peaks C1, C2, and A, respectively, which are an order of magnitude larger than those of NC/S cathode ($6.37 \times 10^{-9} \sim 6.28 \times 10^{-8}$ cm² s⁻¹), as shown in Figure 3f and S20. The enhanced Li^+ diffusion properties on the Co@NC/S surface may be attributed to four advantages of the Co@NC host: (1) hierarchical porous architecture that

shortens ion transport path; (2) in-built electric field in the Co@NC heterointerface that offers Li^+ transfer pathways; (3) synchronous regulation of polysulfide from immobilizing to conversion that alleviates the deposition of insulating layer on the electrode surface; (4) the barrier energy for Li^+ transfer on the Co (111) surface is theoretically estimated to be almost 10 times smaller than that on NC surface.³⁷

We also carried out density functional theory (DFT) calculations to simulate the oxidation and reduction reactions of sulfur electrochemistry on both NC and Co@NC substrates. The results in Figure 3g exhibit that the transformation of S_8 to Li_2S during discharge involves four intermediate LiPSs, namely, Li_2S_8 , Li_2S_6 , Li_2S_4 , and Li_2S_2 . Due to its largest positive Gibbs free (ΔG), the Li_2S_2 -to- Li_2S step is determined to be the rate-limiting step among all the steps, consistent with previous reports.³⁸ The ΔG for the formation of Li_2S on Co@NC is 3.53 eV, much smaller than that on NC (4.14 eV), clearly demonstrating that the reduction of S_8 is much easier on Co@NC surface. Figure 3h and i decipher the energy profiles of Li_2S decomposition during charge, which starts from an intact Li_2S molecule and ends with a LiS cluster and a single Li-ion and electron ($\text{Li}_2\text{S} \rightarrow \text{LiS} + \text{Li}^+ + \text{e}^-$).^{9, 38} The energy barriers for Li_2S decomposition on NC and Co@NC surfaces are calculated to be 0.67 eV and 2.01 eV, respectively. The lower energy barrier evidences the catalytic effects of Co@NC heterostructure for boosting Li_2S phase transformation. These theoretical simulations, together with above electrochemical analysis, evidently explain the Mott-Schottky effect on the Co@NC can simultaneously catalyze the S reduction and Li_2S oxidation, which is crucial to expedite electrochemical properties of Li-S energy storage systems.

After validating the superiority of the Co@NC Mott-Schottky electrocatalyst, the electrochemical properties of Li-S cells using Co@NC/S cathodes were then evaluated. The rate capability of Co@NC/S cathode with a sulfur mass loading of 1.5 mg cm^{-2} is presented in Figure 4a. The discharge specific capacities are as high as 1303, 1103, 971, 882, 815, 741, 634 mAh g^{-1} at 0.1, 0.2, 0.5, 1, 2, 3, and 4 C, respectively, suggesting excellent rate capability of the Co@NC/S electrode. When skipping back the current rate to 0.2 C, the discharge capacity can be recovered to 1064 mAh g^{-1} , indicative of good stability and structural robustness. All these discharge capacities are much higher than those of NC/S (4 C, 320 mAh g^{-1}) cathode. Figure 4b and S24 compare charge/discharge curves at different current rates, from which one can observe higher capacities of both high- and low-voltage plateau (Q_H and Q_L) as well as the larger ratio of Q_L/Q_H for the Co@NC/S cathode than those of NC/S cathode especially at faster rates, revealing that polysulfides species are effectively captured and converted by the Co@NC host. During the beginning of charge process, a voltage jump of NC/S cathode reflects a sluggish Li_2S activation process owing to the semiconducting properties of NC.³⁸ However, the addition of the Co@NC electrocatalyst significantly reduces the potential barrier from 2.29 to 2.20 V, further confirming the superior catalytic activity of the Co@NC heterostructure toward the kinetic oxidation process of Li_2S . Furthermore, it is worth noting that the voltage differences (ΔE) between the charge and the second discharge plateaus of the Co@NC/S electrode are significantly lower than those of the NC/S electrode at all current rates (Figure S24c). This decrease of cell polarization also contributes to the increase in energy efficiency (Figure 4c), which

further confirms that the Co@NC Mott-Schottky electrocatalyst effectively propels the kinetics of round-trip sulfur redox reactions ($S_8 \leftrightarrow S_6^{2-} \leftrightarrow S_4^{2-} \leftrightarrow Li_2S_2 \leftrightarrow Li_2S$) in a working cell.²³

The Co@NC Mott-Schottky electrocatalyst also improved the prolonged cycling stability of sulfur cathodes. As shown in Figure 4d, the NC/S cathode experiences continuous capacity fading over 500 cycles at 1 C and can only retain 28% (205.1 mAh g⁻¹) of its initial capacities. In sharp contrast, the Co@NC/S cathode delivers an initial discharge capacity of 909.4 mAh g⁻¹ and remains at 656.6 mAh g⁻¹ after 500 cycles with a capacity retention of 72 % and average coulombic efficiency of 99.8 %. Note that the capacity contribution from Co@NC heterostructure is negligible (Figure S25). Under a much higher rate of 4 C, the Li-S battery with Co@NC electrocatalyst also achieves unparalleled cycling stability with a capacity retention of 93.0 % after 800 cycles and stable Coulombic efficiency over 98% (Figure 4d). Such extraordinary cycling performance is prominent in comparison to other state-of-the-art Li-S studies on catalytic cathode hosts, based on sulfur loading, cycle numbers, and current densities, as summarized in Table S1.

The postmortem SEM investigation was conducted to analyze the structural integrity of cathodes and anodes that were disassembled from the cells after 100 cycles at 1 C. As exhibited in Figure 4e, large agglomerations are observed on the rough surface of the cycled NC/S cathode, demonstrating serious electrode passivation. Conversely, the Co@NC/S cathode after cycling almost retains its original morphology and exhibits a relatively smooth surface that only contains uniform solid precipitates

with no obvious large aggregations.³⁹ This morphologic evolution reveals that the Co@NC can effectively realize the homogeneous spatial distribution of electroactive sulfur species, which also helps to maintain low charge-transfer resistance during cycling, as shown in the Nyquist plots (Figure S28). As for the Li metal anode paired with the NC cathode (Figure 4f), it unveils a looser and rougher surface and many conspicuous cracks. In comparison, the cycled Li anode from the Co@NC/S cell still maintains a roughly compact surface with few pores/voids. Furthermore, the Li anode and separator from the Co@NC cell exhibit a much lighter color than that from the NC/S cell (Figure S29), further revealing that Co@NC helps to alleviate LiPSs shuttling and stabilize the S cathode and Li anode.

To gain further insights into the Mott-Schottky effect in Co@NC heterostructure, physically mixed Co metal and NC composite (denoted as Co/NC) was also utilized as a sulfur host and the rate and cycling tests of the corresponding sulfur cathode (denoted as Co/NC/S) were conducted under the same conditions. As shown in Figure S23, the Co/NC/S cathode displays a capacity of 474 mAh g⁻¹ at 4 C and retains a capacity of 547.5 mAh g⁻¹ after 500 cycles at 1 C. Both values are worse than those of Co@NC/S cathode, revealing that the extraordinary performance of Co@NC/S cathode originates from the Mott-Schottky catalytic effect in Co@NC heterostructure instead of simply synergistic effect between Co and NC.

To demonstrate the utilization of Mott-Schottky heterostructure in practical high-energy-density Li-S batteries, we also prepared Co@NC/S cathode with ultrahigh loading of 10.73 mg cm⁻² and cycled the cathode under starved electrolyte (E/S = 5.9

$\mu\text{L mg}^{-1}$). As displayed in Figure 4g, the cathode displays an initial capacity of 6.74 mAh cm^{-2} at 0.2 C, which increases to 38.17 mAh cm^{-2} after 17 cycles due to the activation process of the sulfur cathode as often observed in the literature.^{40, 41} A reversible capacity of 6.21 mAh cm^{-2} after 120 cycles remains with overall capacity retention as high as 92.1 %, which compares favorably with recently reported sulfur cathodes with raised loading ($> 5 \text{ mg cm}^{-2}$), as displayed in Figure 4h and Table S2. More crucially, at a current rate of 0.05 C, Co@NC/S cathode releases a high areal capacity of 12.87 mAh cm^{-2} (Figure S32). Accordingly, the gravimetric energy density is calculated to be 307.8 Wh kg^{-1} at a system level (see details in Table S3), which outperforms most of the previously reported Li-S batteries (usually $< 200 \text{ Wh kg}^{-1}$) (see details in Table S4).

High-temperature tolerance is another priority on the way to commercial Li-S batteries.⁴² At 55 °C (Figure S34), the Co@NC/S cathode exhibits a maximum areal capacity of 4.05 mAh cm^{-2} and maintains 80.3 % after 60 cycles under a sulfur loading of 4.29 mg cm^{-2} at 0.2 C. These capacities and cyclabilities are much better than those of NC/S cathode and highly competitive among the recently reported works on high-temperature Li-S batteries (see details from Table S5). These above-mentioned electrochemical performances of Co@NC/S cathode under ultrahigh sulfur loading, low electrolyte usage, and elevated temperature indicate that the Mott-Schottky heterojunction could be a new type of catalytic cathode host for advanced high-energy-density Li-S batteries.

CONCLUSION

In summary, we employed the Co@NC Mott-Schottky heterostructure as a cathode host to catalyze conversion chemistry in Li-S batteries. Combining results from theoretical calculations and spectroscopy analyses, we conclude that, in the Co@NC heterostructure, Co nanoparticles and porous N-doped C matrix are strongly coupled, which induces self-driven charge redistribution and a built-in electronic field at the interface. The interfacial effect, also known as the Mott-Schottky effect, in Co@NC catalyst helps to propel Li^+/e^- transport rates, chemisorb polysulfide species, reduce energy barrier for polysulfide reduction and Li_2S oxidation in the discharge/charge processes. Benefiting from these advantages, the Li-S batteries built with Co@NC catalysts exhibit impressive capacity retention (92.1 %) and gravimetric energy density (307.8 Wh kg^{-1}) under practical working applications, including ultrahigh sulfur loading (10.73 mg cm^{-2}) and a decreased electrolyte usage ($5.9 \mu\text{L mg}^{-1}$). The present work deepens the understanding of Mott-Schottky catalytic mechanism for high-efficiency Li-S conversion chemistry and broadens the horizons of electrocatalyst design for other multi-electron energy storage and conversion systems.

ASSOCIATED CONTENT

Supporting Information.

The Supporting Information is available free of charge at

Additional experimental details and characterizations

AUTHOR INFORMATION

Notes

The authors declare no competing financial interest.

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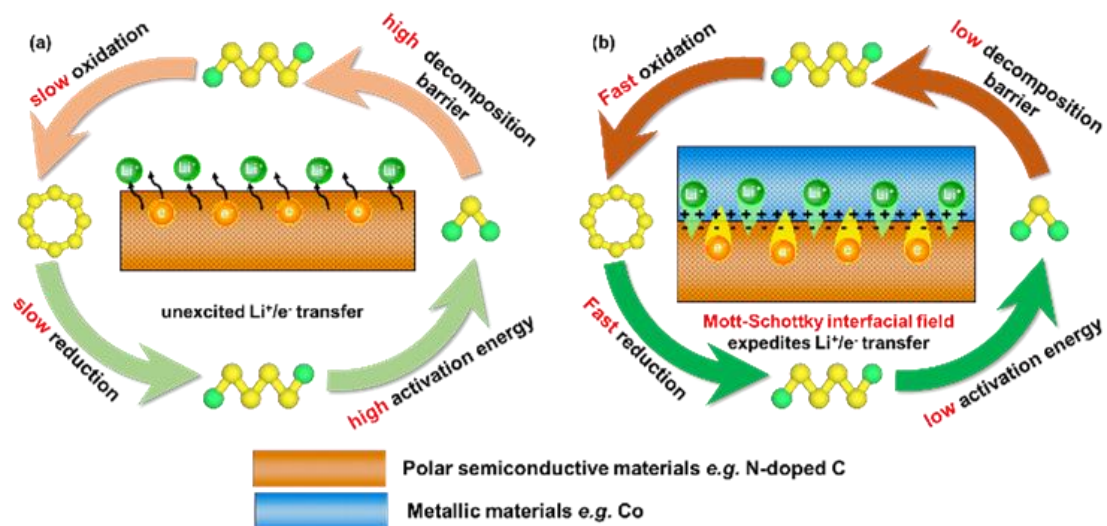


Figure 1. Schematic for Li-S batteries with different cathode hosts: (a) routine polar semiconductive materials; (b) Mott-Schottky heterostructures. The Mott-Schottky interfacial effect boosts Li⁺/e⁻ transfer and redox kinetics of sulfur species during charge/discharge processes.

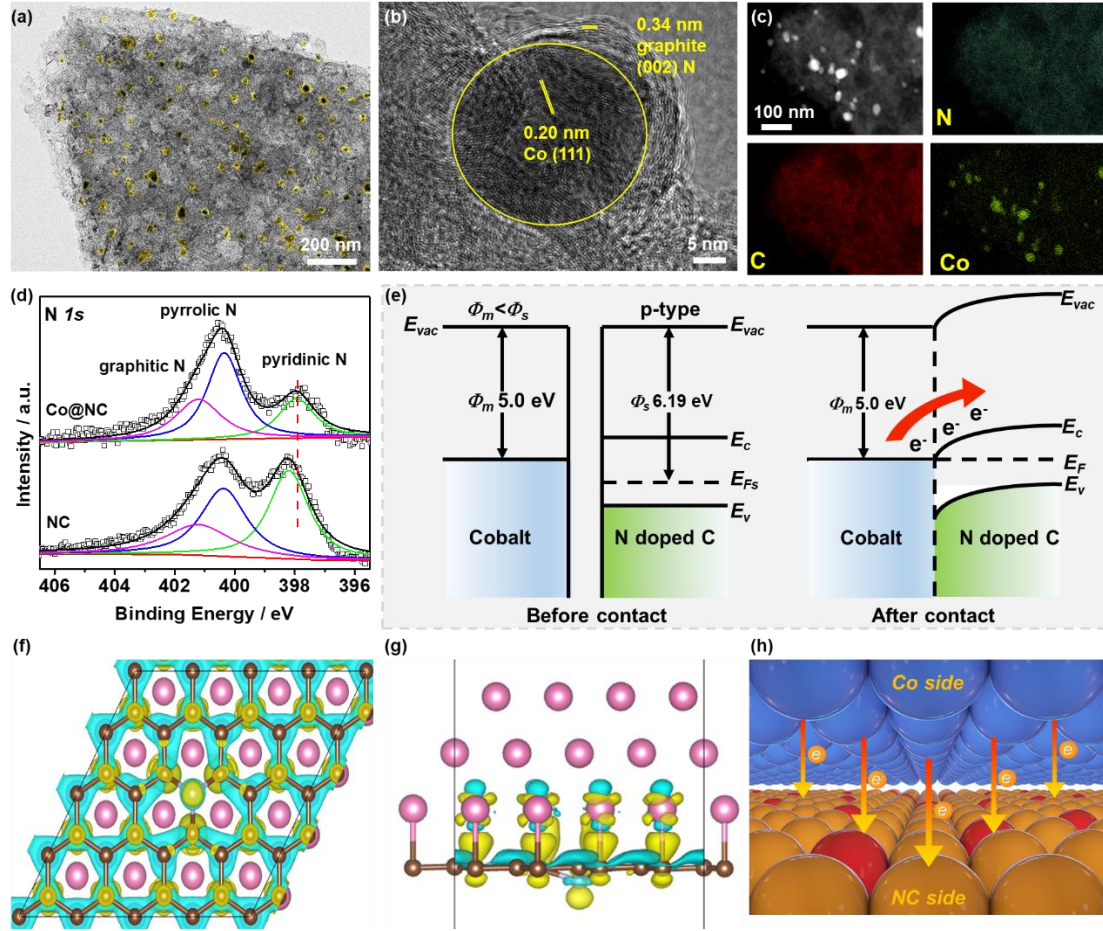


Figure 2. (a) TEM, (b) HRTEM, (c) HAADF images and the corresponding EDX mapping results of Co@NC heterostructure. (d) N 1s XPS spectra of NC and Co@NC. (e) Schematic illustration of the Mott-Schottky type contact of the Co@NC before and after contacting. (f, g) Differential charge density redistributions in Co@NC heterostructure. The yellow and blue isosurfaces represent charge accumulation and depletion in the space, respectively, and the isosurfaces level is 0.002 au. (h) Schematic diagram for the charge transfer between Co and NC.

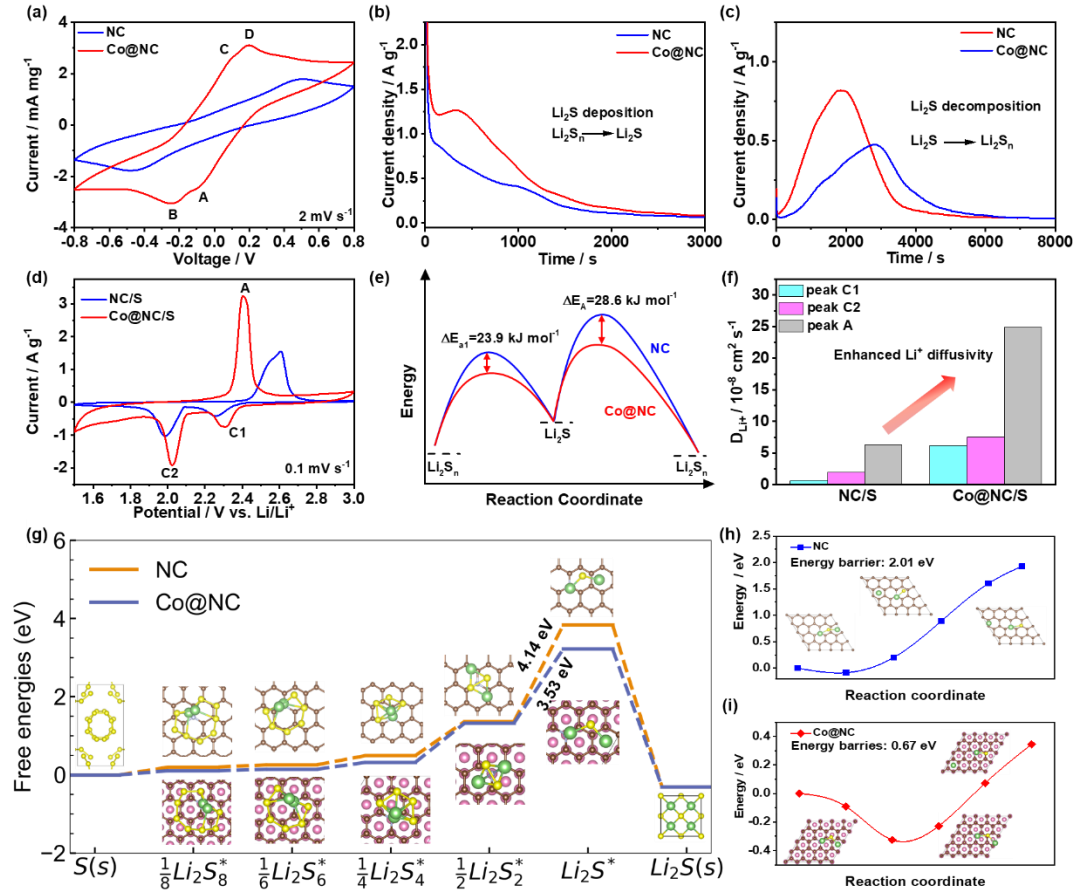


Figure 3. (a) CV profiles of Li_2S_6 symmetrical cells with NC and Co@NC electrodes. (b) Potentiostatic discharge profiles with a Li_2S_6 solution from the open circuit voltage to 2 V. (c) Potentiostatic charge profile at 2.40 V to evaluate dissolution kinetics of Li_2S . (d) CV profiles of Li-S batteries with NC/S and Co@NC/S cathodes. (e) The activation energies (E_a) of the formation and dissolution of Li_2S . (f) Li^+ diffusion coefficient calculated from the Randles-Sevcik equation. Free-energy evolution for (g) S reduction and (h, i) decomposition of a Li_2S cluster on NC and Co@NC supports, where the optimized adsorption configurations of polysulfides species and the initial, transition, and final structures of Li_2S are presented in the inset. The black, yellow, green, pink, and dark blue balls represent C, S, Li, N, and Co atoms, respectively.

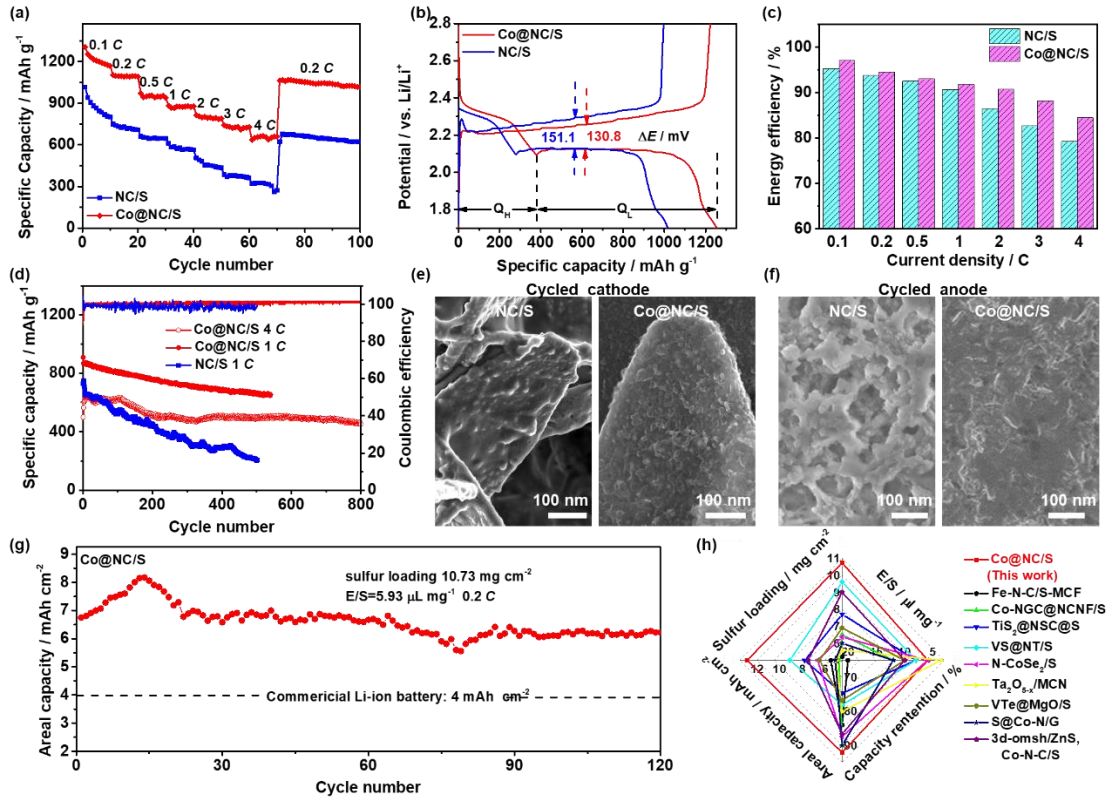


Figure 4. (a) Rate capabilities of NC/S and Co@NC/S cathodes. (b) Charge/discharge curves of different electrodes at 0.1 C. (c) Energy efficiency of different cathodes. (d) Long-term cycling performance of different cathodes at 1 and 4 C. SEM images of (e) sulfur cathode and (f) Li anode after 100 cycles at 1 C. (g) Cycling performance of Co@NC/S cathode with high sulfur mass loading of 10.73 mg cm⁻² at 0.2 C. (h) Electrochemical performance comparison between our Co@NC/S cathode and previously reported sulfur cathodes. The related data are collected from Table S2.

Table of Content (TOC) Graphic

