“Ship in a Bottle” Design of Highly Efficient Bifunctional Electrocatalysts for Long-Lasting Rechargeable Zn–Air Batteries

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Supporting Information

ABSTRACT: The poor durability of bifunctional oxygen electrocatalysts is one main bottleneck that suppresses the widespread application of rechargeable metal–air batteries. Herein, a “ship in a bottle” design is achieved by impregnating fine transition metal dichalcogenide nanoparticles into defective carbon pores that act as interconnected nanoreactors. The erected 3D porous conductive architecture provides a “highway” for expediting charge and mass transfer. This design not only delivers a high surface-to-volume ratio to increase numbers of exposed catalytic sites but also precludes nanoparticles from aggregation during cycling owing to the pore spatial confinement effect. Therefore, the long-term plague inherent to nanocatalyst stability can be solved. Moreover, the synergistic coupling effects between defect-rich interfaces and chemical bonding derived from heteroatom-doping boost the catalytic activity and prohibit the detachment of nanoparticles for better stability. Consequently, the developed catalyst presents superior bifunctional oxygen electrocatalytic activities and durability, out-performing the best-known noble-metal benchmarks. In a practical application to rechargeable Zn–air batteries, long-term cyclability for over 340 h is realized at a high current density of 25 mA cm⁻² in ambient air while retaining an intact structure. Such a universal “ship in a bottle” design offers an appealing and instructive model of nanomaterial engineering for implementation in various fields.

KEYWORDS: bifunctional catalysts, oxygen electrocatalysis, nanostructure design, defective carbon, Zn–air batteries, long-term cyclability

The recognition of global overconsumption of fossil fuels and associated environmental deterioration has spurred a strong imperative for the development of next-generation sustainable energy storage and conversion systems.1,2 The urgency is to develop inexpensive renewable energy storage technologies with high energy densities to reduce dependence on conventional fossil fuels. Rechargeable metal–air batteries are attracting renewed interest as a promising energy storage system because of the high energy density and enhanced safety as well as environmental compatibility.3 In particular, a Zn–air battery can theoretically store energy of as much as 1086 Wh kg⁻¹, up to 4 times higher than currently widely used lithium-ion batteries. The attractive energy density is derived from an open-cell configuration wherein a zinc anode is coupled with an air-breathing cathode that uses oxygen in atmospheric air as fuel for the electrochemical reactions.4,5 However, their widespread practical applications are largely hindered by the unsatisfactory rechargeability (i.e., lifetime) and low energy conversion efficiency, which mainly stem from the challenges encountered at the air cathode with sluggish kinetics and poor electrochemical stability for the repeated oxygen reduction (ORR) and evolution (OER) reactions.6,7 Therefore, the development of efficient and durable bifunctional electrocatalysts for both ORR and OER plays a pivotal role. Up to now, noble-metal electrocatalysts, such as platinum (Pt) and iridium (Ir), have been intensively investigated and recognized as the most highly active electrocatalysts for ORR or OER.8 Unfortunately, their
high cost, unsatisfactory catalytic bifunctionality, and stability inevitably preclude their widespread applications.9,10

Non-precious-metal based materials hybridized with carbon have become promising candidates for high-performance Zn−air batteries.11,12 Recently, numerous synthesis strategies have been reported, mainly including metal deposition onto carbon-based supports or carbon coating on metal particles.13−16 However, the former method cannot efficiently immobilize metal nanoparticles and control their growth during synthesis and even catalytic reactions, which may cause the loss or aggregation of nanoparticles and thus degrade their electrocatalytic durability and performance. Although the latter method can form a carbon-sealed confined space for metal-based particles, the absence of high and uniform porosity for sufficient mass transfer, as well as inefficient control of nanoparticle size and structural change due to poor consistency of the carbon-coating thickness, lead to limited electrocatalytic performance. Furthermore, numerous advantages of the carbon support should be exploited beyond its role as a nanomaterial carrier, which potentially include defect-induced modification and synergistic effects for enhanced electrocatalytic performance.17,18 Therefore, a rational structural design combined with defect engineering is critical to achieve a high bifunctional catalytic activity with enhanced durability.

Herein, we report a “ship in a bottle” nanocatalyst design based on fine pyrite-phase transition metal dichalcogenide nanoparticles rooted inside defective carbon pores. The pores function as individual nanoreactors that are interconnected with one another, in which oxygen catalysis efficiently proceeds under the steric pore environment. As a proof-of-concept, cobalt pyrite (CoS2) was selected as a model material because of its abundance and intrinsically conductive metallic nature.19 This design delivers multiple advantages: (i) the nucleation of CoS2 nanoparticles is controlled due to the spatial confinement effect of the carbon pores, which not only restricts nanoparticle overgrowth to guarantee a high surface-to-volume ratio and increase numbers of exposed catalytic sites but also prevents nanoparticles from aggregating during catalysis, solving the long-term issue of nanocatalyst stability; (ii) the interpenetrating and interconnecting porous carbon framework builds up a 3D conductive architecture as the nanoreactor to provide a “highway” for electron and mass transfer and facilitate fast catalytic kinetics; and (iii) the synergistic coupling effects between defect-rich interfaces and chemical bonding derived from sulfur doping boost the catalytic activity and prohibit the detachment of nanoparticles. The resultant catalyst presents a superior bifunctional electrocatalytic activity and durability toward ORR and OER and demonstrates excellent performance and long-term cyclability in the rechargeable Zn−air battery.

RESULTS AND DISCUSSION

A facile and universal impregnation strategy to achieve the recessed growth of CoS2 nanoparticles inside heteroatom-doped, defective carbon pores is illustrated in Scheme 1a. Ketjen Black EC600JD (KJ) carbon with adsorbed water vapor and ammonia functions as a porous built-in reaction host, whose microscopic morphology is shown in Figure S1a−c. After KJ carbon was soaked in cobalt(II) acetylacetonate organic solution, the cobalt precursor is hydrolyzed inside the
carbon pores under the alkaline environment derived from absorbed ammonia. Following this, the cobalt-hydrolyzed product undergoes thermal sulfurization, yielding the final product, in which CoS₂ is immobilized inside the S-doped, defective carbon pores (CoS₂/SKJ). Each individual pore acts as a nanoreactor, interconnecting with each other for efficient oxygen electrocatalysis, as illustrated in Scheme 1b.

Scanning electron microscopy (SEM) imaging (Figure 1a) reveals a uniform distribution and clean external surface for CoS₂/SKJ. No obvious morphology change is observed compared to KJ (Figure S1a), and no agglomeration forms on the external carbon surface, implying successful encapsulation of CoS₂ inside the carbon pores. As shown by the scanning transmission electron microscopy (STEM) imaging (Figure 1b), the bright nanodomains are the CoS₂ phase, whereas the dark domains are mainly the C phase from KJ carbon. The corresponding energy-dispersive X-ray spectroscopy (EDS) mapping confirms the presence of Co, S, and C elements in CoS₂/SKJ, and the homogeneous distribution of Co and S demonstrates that abundant CoS₂ active sites were exposed and dispersed uniformly throughout the porous carbon host. As shown in Figure 1c, the high-resolution transmission electron microscopy (HRTEM) image provides explicit evidence that CoS₂ nanoparticles with a particle size of ~8.0 nm are embedded in the carbon pores. Importantly, such small nanoparticles were maintained after a heat treatment at 450 °C because of the carbon pore confinement effect, which deters the CoS₂ nanoparticles from agglomerating even at high temperatures. The HRTEM image in Figure 1d reveals that the carbon framework intimately contacted with CoS₂ nanoparticles is highly graphitized, and the graphitization occurs not only around the CoS₂-loaded pores but also throughout the entire composite, building up an interconnected conductive network to offer fast electron transfer. KJ carbon exhibits a typical C (002) facet with a d-spacing around 0.337 nm (Figure S1d), while the graphitized carbon in CoS₂/SKJ presents slightly larger d-spacing (Figure 1d), which is attributed to the successful S-doping within the graphitized carbon skeleton. It is also noteworthy that the massive porous structures interpenetrate and interconnect with each other, creating a 3D nanoreactor architecture for ion transfer inside and through CoS₂/SKJ, which potentially provides significantly improved kinetics for electrocatalytic processes. The encapsulated CoS₂ demonstrates a high crystallinity of the primary particles, in which the lattice fringes with an interplanar distance of 0.276 nm correspond to the (200) crystal plane of CoS₂, which is evidenced by the fast Fourier transform (FFT) pattern (Figure 1e). The STEM image in Figure 1f shows one isolated CoS₂ nanoparticle with a diameter of around 8.0 nm immobilized within carbon pores, confirming the pore confinement effect, which is also reflected in the nanoparticle size distribution (Figure S2). As revealed in the corresponding electron energy loss spectroscopy (EELS) mapping (Figure 1g), the S element shows a homogeneous dispersion over the carbon skeleton in addition to the central CoS₂ phase, indicating the successful S-doping. The central-domain distribution of Co suggests the sole existence of intact cubic CoS₂. These results validate the encapsulation of immobilized CoS₂ into the interior pores of the S-doped carbon host. The morphology of commercial CoS₂ is shown in Figure S3. For comparison, a control composite was prepared by mechanically mixing KJ with commercial CoS₂ followed by the same thermal sulfurization process (denoted as CoS₂ plus SKJ). Without the pore confinement effect, CoS₂ plus SKJ

Figure 1. (a) Overview SEM and (b) STEM images of CoS₂/SKJ as well as corresponding EDS mapping images. (c) TEM image of CoS₂/SKJ. (d) HRTEM image of graphitized carbon intimately surrounding CoS₂ nanoparticles. (e) HRTEM image and corresponding FFT diffraction pattern. (f) STEM image and (g) corresponding EELS elemental mapping and overlaid images of a single CoS₂ nanoparticle immobilized inside carbon pores.
displays much larger CoS2 particles with a diameter of 200 nm compared to CoS2/SKJ (Figure S4a,b). Besides, obvious agglomeration occurs in CoS2 plus SKJ, leading to a non-uniform distribution of CoS2 and SKJ carbon.

The crystalline nature of the as-prepared materials is revealed by the X-ray diffraction (XRD) (Figure 2a). All the peaks can be unequivocally attributed to cubic CoS2 with the space group $P\bar{4}3m$ (JCPDS 41-1471), confirming that the cobalt precursor was thermally sulfurized into cobalt sulfide. The crystal domain size of CoS2 along the (200) direction is calculated to be 8.3 nm via the Scherrer equation, which coincides with the results observed in HRTEM. Moreover, the peak at 23.8° corresponds to a typical carbon (002) plane, and its intensity in CoS2/SKJ slightly decreased compared with that in KJ. This change reflects the weakened graphitic crystallinity, and correspondingly numerous defects along the (002) direction that were likely formed as a result of S doping in the carbon skeleton.

Figure 2. (a) XRD patterns, (b) Raman spectra with a magnified inset, (c) $N_2$ adsorption–desorption isotherms, and (d) pore size distribution of KJ and CoS2/SKJ.

Figure 3. (a) TGA curves of CoS2/SKJ, KJ, and commercial pure CoS2 measured in air; high-resolution XPS spectra of (b) C 1s, (c) Co 2p, and (d) S 2p for CoS2/SKJ.
The Raman spectra of both KJ and CoS$_2$/SKJ show two scattering peaks located at around 1340 and 1590 cm$^{-1}$, which can be assigned to the disorder-induced D band with an A$_{2g}$ vibration mode of a defective C$_6$ ring, and G band with an E$_{2g}$ mode of sp$^2$-bonded C atoms, respectively.$^{26,27}$ The D/G intensity ratio ($I_D/I_G$) of CoS$_2$/SKJ is higher than that of KJ, thus further revealing its higher defective nature and structural distortion arising from the S doping. Besides, an obvious red shift of the G band is found in CoS$_2$/SKJ (1586 cm$^{-1}$) compared with KJ carbon (1592 cm$^{-1}$), which is a typical characteristic of n-type substitutional doping. Such a phenomenon can be attributed to the elongation of C–C bonds induced by the S-doping, which weakens the bond strength and thus decreases the vibrational frequency.$^{28,29}$ The XRD results coupled with Raman spectroscopy confirm that S atoms have been successfully incorporated into the carbon framework with the formation of numerous defects within CoS$_2$/SKJ, and that the S doping of graphitized carbon can be achieved by our approach.

It is well-known that a high porosity and large surface area are crucial attributes of catalysts with good activity, and thus, the pore texture of the catalytic materials is investigated by N$_2$ adsorption and desorption measurement. As displayed in Figure 2c, a typical type-IV isotherm with a characteristic H3 hysteresis loop in the medium pressure range reveals the mesoporous nature. The specific surface area of KJ is 1433 m$^2$ g$^{-1}$, and its pore volume is 2.6 cm$^3$ g$^{-1}$. Correspondingly, KJ exhibits a narrow pore size distribution (PSD) of around 4.0 nm (Figure 2d), which is slightly smaller than the size of CoS$_2$ nanoparticles confined inside pores. This suitable pore size can effectively immobilize CoS$_2$ nanoparticles and prohibit detachment from the carbon pores, thus presenting a "ship in a bottle" design. After the thermal sulfurization with the formation of CoS$_2$ inside the pores, the specific surface area drops to 993 m$^2$ g$^{-1}$, while the pore volume decreases to 2.0 cm$^3$ g$^{-1}$. This high porosity of CoS$_2$/SKJ is favorable for exposing a high density of active sites and supplying large diffusion space for reactants, which are beneficial for its electrocatalytic performance. Compared to KJ, CoS$_2$/SKJ maintains similar shapes of the isotherm curve and PSD plots except with decreased pore volume, and there is no shift of the PSD peak position between CoS$_2$/SKJ and KJ. This strongly suggests that CoS$_2$ nanoparticles were immobilized within the pores. The small CoS$_2$ crystal size is derived from the adsorbed water vapor confined within the carbon pores, where its limited amount further determines the size of the cobalt-hydrolyzed product, followed by the sulfurization process.

Figure 4. (a) ORR curves of various catalysts in O$_2$-saturated 0.1 M KOH solution at 1600 rpm. (b) LSV curves of CoS$_2$/SKJ at different rotating speeds (inset: K–L plots obtained under various potentials). (c) OER curves and (d) Tafel plots of various catalysts at 1600 rpm in N$_2$-saturated 0.1 M KOH electrolyte. (e) differences between the ORR $E_{1/2}$ and OER $E_{p-10}$ of different catalysts. (f) ORR and (g) OER activities of CoS$_2$/SKJ and CoS$_2$ plus SKJ before and after 3500 cycles, respectively. (h) ORR and (i) OER chronoamperometric response of CoS$_2$/SKJ and Pt/C, CoS$_2$/SKJ, and Ir/C at a constant potential of 0.7 and 1.6 V, respectively.
process, in which CoS2 is formed without particle agglomeration and crystal growth owing to the pore confinement effect.

The loading amount of CoS2 within the carbon pores is evaluated by thermogravimetric analysis (TGA), as shown in Figure 3a. Commercially pure CoS2 is also studied by TGA for determining the loaded CoS2 content in CoS2/SKJ, which is calculated to be 15.5 wt %, while the carbon content is 84.5 wt % (see calculation details in the Supporting Information). The burned products of CoS2/SKJ collected after thermal treatment at 900 °C in air were investigated using XRD, as shown in Figure S5. To further investigate the elemental compositions and bonding configurations in CoS2/SKJ, X-ray photoelectron spectroscopy (XPS) is conducted (Figure 3b–d). The high-resolution C 1s spectrum (Figure 3b) not only shows peaks corresponding to C−C at 284.8 eV and C=O at 289.0 eV but also exhibits an apparent C−S peak located at 285.6 eV, implying that S was successfully doped into the carbon lattice network by forming the C−S covalent bond.32,33 In the high-resolution Co 2p spectrum (Figure 3c), a pair of characteristic peaks located at 780.5 and 796.5 eV are assigned to Co 2p3/2 and Co 2p1/2 with two satellite peaks at 785.5 and 802.2 eV, respectively, confirming the presence of Co2+ in the CoS2 phase.32,33 Figure S6 compares the Co 2p spectra of as-prepared CoS2/SKJ and pure CoS2. It is clearly seen that Co peaks in CoS2/SKJ slightly shift to the lower energy region compared to those of pure CoS2. This is because the electron cloud is expected to show a bias between CoS2 and S-doping carbon framework, forming a strong electronic coupling between these two components. Figure 3d presents the S 2p spectrum, in which the peaks at 162.6 and 163.7 eV are attributed to S 2p3/2 and S 2p1/2, respectively, corresponding to the disulfide S2−.32,34 A pair of characteristic peaks appearing at 164.4 and 165.3 eV belong to C−S−C and C=S bonds, respectively, validating that our designed thermal sulfurization process leads to S incorporation into the carbon skeleton resulting in C−S bonded groups.32,35 The existence of the C−S−C bond, which has been recognized as an effective active site, contributes to considerable catalytic activity.36 The peak centered at 168.4 eV arises from a SO3− species, such as sulfonate or sulfate, which is probably formed by the surface oxidation of sulfur due to the exposure in air.37,38 Therefore, the S-doping and carbon defective nature of CoS2/SKJ have been unveiled by XRD, Raman, and XPS results, which plays a key role for enhancement of the electrocatalytic activity.

Hitherto, the above results and analysis have demonstrated that our designed small CoS2 nanoparticles encapsulated inside pores of S-doped and highly defective carbon nanomaterials have been successfully prepared, thereby realizing a “ship in a bottle” design. All the exhibited features endow the product (CoS2/SKJ) with great potential as an exceptional electrocatalyst for Zn-air batteries.

The bifunctional electrocatalytic performance of CoS2/SKJ was evaluated via the linear sweep voltammetry (LSV) measurements. As a control, S-doped carbon was prepared by direct thermal sulfurization of KJ (denoted as SKJ). LSV measurements of CoS2/SKJ, SKJ, commercial CoS2 nanoparticles and KJ, as well as noble-metal catalyst (i.e., Pt/C and Ir/C) were conducted for comparison. The polarization curves of various prepared catalysts measured at a rotation speed of 1600 rpm are shown in Figure 4a. Compared to KJ, SKJ shows a higher onset potential of 0.80 V (versus the reversible hydrogen electrode, RHE) and diffusion-limited current density of 4.8 mA cm−2, which confirms the significant role of S-doping in enhancing ORR catalytic activity. CoS2 plus SKJ shows a low onset potential of 0.82 V and diffusion-limited current density of 4.8 mA cm−2, which is similar to the commercial CoS2 nanoparticles (0.81 V and 4.1 mA cm−2). By comparison, CoS2/SKJ exhibits a prominent improvement on ORR performance, with a higher onset potential of 0.92 V and larger diffusion-limited current density of 5.2 mA cm−2. Moreover, CoS2/SKJ shows a half-wave potential (E1/2) of 0.84 V, which is only about 10 mV more negative than that of commercial Pt/C catalyst but much higher than that of CoS2+SKJ (0.76 V), SKJ (0.69 V), commercial CoS2 nanoparticles (0.74 V), and KJ (0.66 V), highlighting the superior electrocatalytic activity of the CoS2/SKJ catalyst. To further investigate the electron-transfer kinetics of the CoS2/SKJ catalyst, polarization curves were measured at different rotation speeds. As shown in Figure 4b, CoS2/SKJ shows a well-defined plateau of diffusion-limited currents at all rotation speeds, where, as expected, the current densities increase with the rise of rotation speeds, which is owing to the expedited mass transfer of oxygen molecules from the electrolyte to the electrode surface. More importantly, the R–L plots of CoS2/SKJ present good linearity and near parallelism at different potentials, revealing the first-order reaction kinetics toward the concentration of dissolved oxygen.39 The electron-transfer number (n) for CoS2/SKJ obtained from the R−L slope is 4.0, signifying the fast kinetics with a four-electron-transfer reaction pathway to reduce oxygen directly to OH−. The superior reaction kinetics are attributed to the fast electron transfer benefiting from an interconnected conductive carbon framework, which not only surrounds the CoS2-impregnated pores but also is present within the whole composite. These results confirm the superior ORR catalytic activity of CoS2/SKJ.

Oxygen evolution activities are also investigated to confirm the suitability of CoS2/SKJ as bifunctional electrocatalysts for both ORR and OER. Figure 4c illustrates the LSV behavior of the as-prepared catalysts in N2-saturated 0.1 M KOH. As expected, CoS2/SKJ exhibits a OER potential of only 1.58 V at the current density of 10 mA cm−2, which is smaller than CoS2 plus SKJ (1.66 V), SKJ (1.73 V), commercial CoS2 (1.65 V), and KJ (1.76 mV), demonstrating the smallest over-potential of CoS2/SKJ among the obtained catalysts. In particular, the over-potential at 10 mA cm−2 (η10) of CoS2/SKJ (350 mV) is even smaller than that of commercial Ir/C catalyst (380 mV), revealing its excellent OER electrocatalytic activity. The fast OER kinetics of CoS2/SKJ is further revealed by its smaller Tafel slope (83 mV dec−1) relative to the other catalysts (Figure 4d). In addition, the activity improvement from KJ to SKJ reflects that S doping effectively contributes to OER catalytic activity enhancement.

To reveal the bifunctional activity of the CoS2/SKJ catalyst, the potential difference ∆E between the ORR (half-wave potential (E1/2) and OER potential at 10 mA cm−2 (E10)) is assessed (Figure S7), with a smallest value of ∆E (∆E = E10 − E1/2) indicating superior catalytic bifunctionality. The potential toward ORR and OER is reflected by comparing with the E° (OH−/O2) = 1.23 V. As illustrated in Figure 4e and Table S1, CoS2/SKJ exhibits a desirable ∆E of 0.74 V, smaller than Pt/C coupled with Ir/C, as well as the other as-prepared catalysts. More importantly, the low ∆E of CoS2/SKJ outperforms the majority of well-developed bifunctional catalysts reported recently (Table S2), unveiling the superior bifunctional electrocatalytic nature of CoS2/SKJ.
To further reveal the significant role of pore spatial confinement in catalyst stability, the electrochemical durability of CoS₂/SKJ and CoS₂ plus SKJ was investigated. As shown in Figure 4f and Table S3, CoS₂/SKJ exhibits much better ORR stability than CoS₂ plus SKJ, with a smaller $E_{1/2}$ loss (1.2%) than CoS₂ plus SKJ (6.6%) after 3500 cycles. For the OER stability evaluation after 3500 cycles (Figure 4g and Table S3), CoS₂ plus SKJ shows a substantial increase in $\eta_{j=10}$ (500 mV), whereas CoS₂/SKJ mostly maintains its OER activity with a $\eta_{j=10}$ of 370 mV. The inferior stability of CoS₂+SKJ is precisely owing to the absence of the pore confinement effect, which results in a severe agglomeration after a long-term cycling (Figure S8), causing inevitable loss of catalytic activity. Furthermore, CoS₂/SKJ and noble-metal benchmark catalysts were also evaluated via a chronoamperometric (CA) measurement at 0.7 V (Figure 4h). CoS₂/SKJ retains 97.5% of the initial ORR current, far superior to the 57.0% retention of Pt/C. Likewise, the excellent OER stability of CoS₂/SKJ is also revealed by the CA test at 1.6 V, with only 9.8% current decay after 30 000 s, whereas Ir/C presents a much larger current decay of 42.0% (Figure 4i).

Building upon the promising ORR/OER electrocatalytic activity and durability presented above, CoS₂/SKJ was used as a bifunctional catalyst within the air cathode to assemble rechargeable Zn–air batteries. Figure 5a exhibits a schematic of the Zn–air battery system. Commercial Pt/C plus Ir/C were also assembled in the Zn–air battery under same technique condition as the reference. The galvanodynamic charge and discharge curves are illustrated in Figure 5b. Both Zn–air batteries deliver a similar open-circuit voltage of ca. 1.4 V, but CoS₂/SKJ exhibits more-desirable charging and discharging curves. A narrower voltage gap between charge and discharge polarization voltages is found for CoS₂/SKJ, which reveals its better charge and discharge performance than the commercial Pt/C plus Ir/C mixture catalyst. Besides, CoS₂/SKJ delivers a higher power density (over 104 mW cm$^{-2}$) than Pt/C–Ir/C hybrid catalyst (84 mW cm$^{-2}$) (Figure 5b). This is because increased exposure of accessible active CoS₂ nanoparticles assures a higher power generation per unit area of electrode. At
a certain current density, the battery energy efficiency is based on the ratio of the discharge potential to the charge potential.\textsuperscript{40}

As shown in Figure 5c, at the current density of 120 mA cm\textsuperscript{−2}, the energy efficiency of CoS\textsubscript{2}/SKJ is 1.5 times higher than that of the noble-metal benchmark, which is attributed to its provision of unimpeded access for reactants reaching toward the active sites (i.e., the high-efficiency catalyst utilization). Moreover, the more salient electrochemical performance of the CoS\textsubscript{2}/SKJ air electrode is also due to the interconnected conductive carbon network and interpenetrating porous structure, which guarantee fast charge transfer, as well as enhanced accessibility of oxygen species and electrolyte to the electrocatalytic sites.

To further investigate the efficiency and cycle stability of CoS\textsubscript{2}/SKJ for the Zn-air battery, a galvanostatic charge−discharge test is performed at a high current density of 25 mA cm\textsuperscript{−2}, which cycles with 40 min charging and 40 min discharging. As illustrated in Figure 5d, the initial charge−discharge voltage gap of CoS\textsubscript{2}/SKJ is 0.92 V, smaller than that of the noble-metal benchmark Pt/C plus Ir/C catalyst (1.01 V). More importantly, the Zn−air battery assembled with CoS\textsubscript{2}/SKJ exhibits a superior durability without noticeable voltage fading for both charge and discharge processes over 340 h of continuous operation, whereas the battery using Pt/C plus Ir/C catalyst delivers a limited cyclability with operation time of less than 30 h. The poor stability of Pt/C plus Ir/C catalyst is because of the dissolution and redeposition, migration and aggregation, and sintering of the noble-metal particles as well as the formation of the insulating platinum oxides under operational conditions.\textsuperscript{39} These could lead to an increase in the particles' size and the particle detachment, causing the irreversible loss of active sites and degradation of electrocatalytic performance.\textsuperscript{34} The battery energy efficiency of Pt/C−Ir/C hybrid catalyst reduces far faster than that of CoS\textsubscript{2}/SKJ, leading to significant charge and discharge performance losses. Particularly, the battery with CoS\textsubscript{2}/SKJ can also withstand a very high current density (25 mA cm\textsuperscript{−2}), presenting an extremely low decaying rate of 0.039 millivolts per cycle and a small voltage gap of 0.93 V after 255 cycles, much better than the noble-metal benchmark with a high voltage decaying rate (32.44 millivolts per cycle) and voltage gap (1.74 V) (Figure 5e). Such long-lasting cyclability over this time scale in this work is evidently superior to other recently reported results, as shown in Figure 5f and associated detailed information in Table S4. Undoubtedly, the impressive stability and long-term rechargeability of CoS\textsubscript{2}/SKJ can be attributed to the pore confinement design. Specifically, the carbon pores provide spatial confinement to restrict the overgrowth of active CoS\textsubscript{2} nanoparticles and prevent them from aggregating during the catalysis and cycle process. At the same time, S doping within the carbon framework can modify the electronic and steric pore environment with the formation of the C−S bond, which may afford strong affinity toward CoS\textsubscript{2}, which can, in turn, immobilize CoS\textsubscript{2} nanoparticles inside the pores. To affirm the structural evolution upon cycling as the evidence for the above analysis, SEM, TEM, and STEM observations were carried out on the CoS\textsubscript{2}/SKJ electrode after cycling over 340 h. Figure S9 demonstrates that the intact 3D architecture was retained during the battery cycling. CoS\textsubscript{2} nanoparticles are found still immobilized well inside the carbon pores with an original size of around 8.0 nm without obvious agglomeration or detachment (Figure 5g and h). This manifests the robust carbon framework structure and effective pore spatial confine-

ment effect, which contributes to the excellent durability. Finally, the application prospect of the device is demonstrated in Figure 5i. A total of three CoS\textsubscript{2}/SKJ-batteries power a mini-fan equipped with light-emitting diode (LED), which operates at a minimum voltage of 3.6 V.

All of the above results highlight the excellent bifunctional electrocatalytic activity and long-term durability of CoS\textsubscript{2}/SKJ, which profit from the following aspects. First, the pore spatial confinement effect makes fine active CoS\textsubscript{2} nanoparticles root inside graphitized carbon pores, which not only presents a high surface-to-volume ratio to facilitate the exposure of more catalytically active sites but also effectively inhibits overgrowth and agglomeration of nanocatalysts during a long-cycling catalytic process. Second, the formation of the C−S−C bond derived from S doping into the carbon skeleton acts as an important catalytically active site for promoting ORR and OER.\textsuperscript{36} As per previous reports,\textsuperscript{52} it can break the electro-neutralitity of carbon to create positively charged sites favorable for O\textsubscript{2} adsorption, and this parallel diatomic adsorption can effectively weaken the O−O bonding to further promote the direct reduction of oxygen to OH\textsuperscript{−} via a four-electron process. Additionally, the S doping within the carbon framework can also enhance the spin density, which plays a pivotal role in determining the catalytically active sites.\textsuperscript{53} Third, the defect-rich interfaces function as atomic traps, which can effectively boost the localization of oxygen species toward active sites, promoting the catalytic activity and stability. Moreover, the graphitized carbon framework not only intimately enfolds the CoS\textsubscript{2} nanoparticles but also interpenetrates over the whole composite, building up an interconnected conductive network to offer fast electron transfer. More importantly, the interpenetrating porous structures erect a robust 3D architecture as the nanoreactors for ion and oxygen species transfer inside and through the CoS\textsubscript{2}/SKJ catalyst, which significantly improves the kinetics for electrocatalytic processes. This architecture capitalizes on the synergistic interactions between CoS\textsubscript{2} nanoparticles, carbon defects, and intimate covalent bonds (i.e., C−S bonding) from S doping under the steric pore environment, which endow CoS\textsubscript{2}/SKJ with prominent catalytic performance.

CONCLUSIONS

In this work, an exquisite bifunctional catalyst consisting of fine CoS\textsubscript{2} nanoparticles encapsulated inside the defective carbon pores is designed and successfully accomplished via precursor impregnation, hydrolysis and subsequent thermal sulfurization. This methodology realizes the combination of excellent catalytic kinetics and electrical conductivity provided by the interconnecting porous carbon framework acting as a nanoreactor, which effectively restricts the nucleation (i.e., overgrowth) and agglomeration of nanocatalysts during catalysis. Therefore, this exquisitely designed catalyst, harvesting the synergistic effect between defect-rich interfaces (i.e., S-doped pores) and effective catalytic active sites (CoS\textsubscript{2} and C−S−C bonding) guarantees superior electrochemical performance. The as-developed CoS\textsubscript{2}/SKJ catalyst exhibits excellent catalytic activity and durability for both ORR and OER with a ΔE of 0.74 V, unveiling its excellent bifunctional catalytic ability for rechargeable Zn−air batteries. Encouragingly, when integrated within an air electrode, CoS\textsubscript{2}/SKJ exhibits low charge and discharge over-potential (i.e., high energy efficiency), large peak power density, and exceptionally stable cyclability of over 340 h at a very high current density of 25 mA cm\textsuperscript{−2}, out-
performing noble-metal benchmarks and other recently reported results. This work provides a guide to designing efficient and durable non-precious-metal bifunctional electrocatalysts for metal–air batteries, and we believe that such a universal “ship in a bottle” design of impregnating fine metal nanoparticles inside pores will deliver an instructive model of material engineering for implementation in various fields.

METHODS

Preparation of CoS2/SKJ. Ketjen Black EC600JD (KJ) was purchased from MTI Corporation, China. The other chemicals were purchased from Sigma-Aldrich and of analytical grade without further purification. In a typical synthesis, 80 mg KJ carbon with adsorbed water vapor was placed inside a fused quartz tube under ammonium gas flow for several minutes at room temperature. Subsequently, the sample was immersed into saturated cobalt(II) acetylacetonate/ethyl acetate solution for 10 h. Then, the solid sample was filtrated and washed with ethyl acetate several times. After drying at 80 °C for 8 h, the collected sample was transferred to the center of a tube furnace. An alumina boat containing 2 g of sulfur powder was placed at the farthest upstream position in the tube furnace. The tube was purged with air with a steady flow of argon gas at 30 sccm, and the sulfurization process was performed at 400 °C for 2 h, resulting in the formation of the CoS2/SKJ sample.

Preparation of CoS2 Plus SKJ. KJ was first mechanically mixed with commercial CoS2 nanoparticles with the same CoS2 content as CoS2/SKJ. Then, the as-prepared sample underwent the above sulfurization process to obtain CoS2 plus SKJ.

Preparation of SKJ. KJ carbon was directly subjected to the above sulfurization process to produce SKJ.

Materials Characterizations. Microscopic morphology, EDS, and EELS data of materials were collected by SEM using an FEI Quanta FEG 250 ESEM microscope and TEM using a JEOL 2010F TEM/STEM equipped with a large solid angle for high X-ray throughput, scanning, scanning and transmission, and a Gatan imaging filter for energy filtered imaging. XRD was carried out using a Rigaku MiniFlex 600 X-ray diffractometer equipped with a Cu Kα irradiation. To collect Raman spectra, a 532 nm laser source was employed. The Thermal Scientific K-α spectrometer was used to collect XPS spectra. TGA was performed on a TA Instruments Q500 under air atmosphere. Nitrogen adsorption–desorption isotherms were measured at liquid nitrogen temperature (−196 °C) by using the Micromeritics ASAP 2020 analyzer; the Brunauer–Emmet–Teller equation was used to calculate the specific surface area, and the Barrett–Joyner–Halenda model was used to determine the pore size distribution.

Electrochemical Measurements. The electrocatalytic performance was investigated using an electrochemical workstation (Biologic VSP 300) with a three-electrode configuration and rotating disk electrode (RDE). A glassy carbon electrode (GC; 0.196 cm2) and a graphite rod were used as the working and counter electrodes, respectively. All of the results were obtained with reference to a graphite rod as the working and counter electrodes, and Pt/C and Ir/C catalysts were also prepared via the same procedure as recently reported results. This work provides a guide to designing efficient and durable non-precious-metal bifunctional electrocatalysts for metal–air batteries, and we believe that such a universal “ship in a bottle” design of impregnating fine metal nanoparticles inside pores will deliver an instructive model of material engineering for implementation in various fields.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.9b02315.

Calculation of the loading content of CoS2 in CoS2/SKJ composite, the morphology and structure of experimental materials, the size distribution of CoS2 nanoparticles within carbon pores, XRD patterns for burning products of CoS2/SKJ in air; entire LSV curves for bifunctional electrocatalytic activities of catalysts and cyclability of Zn–air batteries between this work and recently reported results (PDF)

AUTHOR INFORMATION

Notes

The authors declare no competing financial interest.

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