Autogenous Production and Stabilization of Highly Loaded Sub-Nanometric Particles within Multishell Hollow Metal–Organic Frameworks and Their Utilization for High Performance in Li–O₂ Batteries

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is a chemically designable system providing manipulative formation and simultaneous stabilization of SNPs. In order that the system is effectively implemented, it should have a high porosity and a large surface area for high SNP loading. Metal–organic frameworks (MOFs) provide chemical flexibility and high porosity, such that stable MOFs allow the controlled mass transfer and quantitative stabilization of atomic clusters, while the constituents of decomposable MOFs can work as precursors for SNPs. Accordingly, there were many great efforts to utilize MOFs as materials for preparing various shaped structures with advanced functionalities. In this work, we develop multilayer MOFs with alternating stable and decomposable layers for the autogenous production and stabilization of SNPs in scalable mass loadings. Interestingly, we also show that the single water molecule transfer throughout multilayer MOFs plays a very important role to activate the autogenous synthesis of SNPs. The step-by-step deterioration of a decomposable MOF layer by the single water molecule leads to the construction of uniform atomic clusters, thus promoting the synthesis of SNPs in the stable MOF layer. It is also investigated how to introduce the single water molecules in multilayer MOFs and study the interaction between the single water molecule and its environment as a key factor. Finally, the higher mass loading of SNPs in multishell MOFs is realized by the autogenous production and stabilization of dinuclear SNPs with multishell hollow MOFs, and they are applied as electrocatalysts to realize high performance in Li–O

Figure 1. Strategy for the synthesis of SNPs and the structural characterization of H-ZIF-8[nS]. a) Schematic of the formation process of SNP-embedded MOFs, where different colors signify different metal nodes. b1,c1) Illustrations of multilayer MOFs and multishell with SNPs inside MOFs, where different colors signify different metal complexes. b2–d4) TEM images of H-ZIF-8[1S, 2S, 3S, 4S, 5S]. a) Strategy for autogenously producing and stabilizing SNPs inside MOFs is shown in Figure 1a. First, alternating water-stable and water-decomposable MOFs were stacked in multilayers using the same organic linkers. Next, water molecules need to be transferred to the decomposable MOF layer throughout the stable MOF layer to activate the autogenous production of SNPs. The high hydrogen bonding affinity between ethylene glycol (EG) and water molecules was used to isolate each water molecule, thus providing the manipulative formation of SNP precursors while the water clusters formation is blocked. Owing to their hydrophobicity, the stable MOF layers effectively transferred water molecules to the decomposable MOF layers without adsorption of water molecules. Subsequently, SNPs derived from the decomposable MOFs were stabilized inside the pores of the water-stable MOFs. We chose zeolitic imidazolate framework-8 (ZIF-8) and ZIF-67 as water-stable and water-decomposable MOFs, respectively. ZIF-8 enabled water molecules to be fed one by one, because the penetration of the isolated water molecule could be manipulated by sieving EG–water complexes through hydrophobic micropores. The X-ray diffraction (XRD) patterns of ZIF-8 after hydrolysis (Figure S1, Supporting Information) and the unchanged XRD patterns of ZIF-8 (Figure S2, Supporting Information) demonstrate that all the metal complexes in ZIF-67 are disconnected from the organic linkers when exposed to water. ZIF-67, which is isostructural with ZIF-8 because both MOFs are composed of 2-methylimidazole (2-mim), was seeded for the epitaxial growth of ZIF-8 to construct multilayer ZIFs (ML-ZIFs[nL], where n is the number of layers) (Figure 1b1). The boundary between ZIF-67 and ZIF-8 is indicated by the dotted line in the high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) image (Figure 1b1II) and by the distinct colors corresponding to Co in ZIF-67 and Zn in ZIF-8 revealed by the energy-dispersive X-ray spectroscopy (EDX) (Figure 1bIII; Figure S3, Supporting Information). Figure 1c1 shows that ML-ZIFs[nL] was transformed into SNP-embedded multishell hollow ZIF-8 (H-ZIF-8[nS], where n is the number of shells). The disassembly of the bonds between the Co ions and 2-mim in ZIF-67 generates SNP precursors and interspaces, as identified by the elemental mapping and HAADF-STEM images (Figure 1cII,III; Figure S4, Supporting Information). The line profiling analysis (Figure S5, Supporting Information) clarifies that the Co species migrated into the ZIF-8 shell. The inductively coupled plasma optical emission spectroscopy (ICP-OES) and EDX analyses (Tables S1–S3, Supporting Information) show that H-ZIF-8[1S] has a Co content of 10.4 wt%, indicating that the water-decomposable MOFs allow the high mass loading of Co particles in the water-stable MOFs. Moreover, the transmission electron microscopy (TEM) images (Figure 1d; Figures S6–S10, Supporting Information) demonstrate that H-ZIF-8[2S, 3S, 4S, 5S] are created after sieving EG–water complexes through ML-ZIFs[4L, 6L, 8L, 10L]. These
results reveal that multishell hollow MOFs could be produced via the controlled transfer of isolated water molecules from EG–water complexes through multilayer MOFs.

To elucidate the chemical state of Co SNPs in the ZIF-8 shell, the X-ray photoelectron spectroscopy (XPS) measurements were conducted. No Co 2p characteristics are observed in ML-ZIFs[2L] (Figure S11, Supporting Information), but the XPS peaks of H-ZIF-8[1S] confirm the existence of divalent Co atoms comparable to those in a bulk Co(OH)₂. To further clarify the different structures of ML-ZIFs[2L] and H-ZIF-8[1S], the Co K-edge X-ray absorption near-edge structure (XANES) spectra were collected (Figure 2a), which revealed three noticeable changes. First, the pre-edge feature of H-ZIF-8[1S], marked by * around 7709 eV, is significantly reduced. Second, the shoulder absorption edge caused by the partially localized excitation of Co–N bonds[21] disappears, as indicated by an arrow around 7720 eV. Finally, the main absorption peak shifts to higher photon energy. The pre-edge peak of ML-ZIFs[2L] corresponds to the 1s–3d electric dipole forbidden transition originating from tetrahedrally coordinated Co in ZIF-67, whereas that of bulk Co(OH)₂ becomes almost flat with the formation of centrosymmetric edge-shared CoO₆ octahedra. Thus, the geometric transformation from ML-ZIFs[2L] to H-ZIF-8[1S] results in a transition of Co coordination from tetrahedral to distorted octahedral, attributable to 3d–4p orbital mixing initiated by the slightly tilted centrosymmetric coordination.[22] Moreover, Co atoms are more oxidized through the formation of Co–O bonds, thereby enhancing the 1s–4p transition on back scattering[23] induced by the structural transformation, resulting in the disappeared shoulder edge and higher energy shift. The red area in the 2D contour map (Figure S12, Supporting Information) indicates the formation of a higher oxidation state after

Figure 2. Characterization of Co(OH)₂ SNPs. a) Normalized Co K-edge XANES spectra. b) Radial distribution function obtained by the Fourier transformation of k⁴-weighted Co EXAFS spectra. c) Co K-edge EXAFS simulation curves with Co(OH)₂ size obtained by back Fourier transformation of the radial structural function. d) XRD patterns with simulated patterns of pristine ZIF-8. e) Normalized Co L-edge NEXAFS spectra. f) Normalized N K-edge NEXAFS spectra. g) Illustration of the autogenous production and stabilization of Co(OH)₂ SNPs in a micropore.
the structural transformation of ML-ZIFs[2L] to H-ZIF-8[1S]. The coordination environment of Co was explored using the K-edge extended X-ray absorption fine structure (EXAFS) spectroscopy (Figure 2b). The main peak of ML-ZIFs[2L] at 1.65 Å is attributable to Co–N bonds and the peaks at 2.23, 2.67, 2.94, and 3.71 Å correspond to 2-mim. However, H-ZIF-8[1S] has two peaks corresponding to Co–OH bonds at 1.70 Å, as supported by the IR and XPS spectra[24,25] (Figures S13 and S14, Supporting Information), and to oxygen edge-sharing Co–Co bonds at 2.90 Å. The changes in the coordination environment during the transformation of ML-ZIFs[2L] to H-ZIF-8[1S] can be confirmed by the disappearance of the 2-mim signals and the appearance of only two peaks (Figure S15, Supporting Information). The Co–Co peak is significantly weaker than the Co–OH peak, which indicates the existence of atomic particles.[26] In the Co K-edge simulation curves (Figure 2c; Figure S16, Supporting Information), the ratio between Co–OH peak and Co–Co peak demonstrates that Co(OH)2 is dinuclear in H-ZIF-8[1S] (Table S4, Supporting Information). The XANES and EXAFS spectra (Figures S17 and S18, Supporting Information) verify that Zn is not affected in the process. These results are consistent with the effective disconnection of all the coordinate bonds of ZIF-67 and the generation of dinuclear Co(OH)2 SNPs inside the micropores of H-ZIF-8[1S]. In addition, the “disordered-to-crystalline” phenomenon[27] involves agglomeration of SNPs upon exposure to a highly enhanced electron beam, thereby corroborating the existence of SNPs inside the damaged multishell hollow MOFs (Figure S19 and Video S1, Supporting Information).

The role of each micropore was also confirmed by the changes in the physical and chemical properties of H-ZIF-8[1S]. The XRD patterns of ZIF-8 agree with the simulated patterns of H-ZIF-8[1S] at diffraction angles lower than 20° (Figure 2d; Figure S20, Supporting Information). However, the peaks associated with the (011) facets shift toward higher angles and exhibit reduced intensities. The scanning electron microscopy (SEM) images (Figures S21 and S22, Supporting Information) reveal that ZIF-8 has a rhombic dodecahedron morphology with exposed (011) facets that form 1.1 nm micropores connected to 0.34 nm apertures.[28] The micropore is induced by the (011) facets allowing the passage of only a single water molecule.[29] Thus, the formation of Co(OH)2 SNPs in micropores results in a shift to higher angles owing to the slightly reduced pore sizes, with the reduced intensities for the (011) facets. The decreased pore volumes and pore sizes in the Brunauer–Emmett–Teller (BET) isotherms as well as the pore size distribution curves also indicate the micropore filling by the Co(OH)2 SNPs (Figures S23 and S24, Supporting Information). The near-edge X-ray absorption fine structure (NEXAFS) spectra showed a changed dipole transition from core electrons to unoccupied molecular orbitals caused by the micropore filling. The trigonal or tetragonal distortions of octahedral bulk Co(OH)2 lead to e_g orbital splitting, as represented by Co L2,3-edge multipllets.[30] (Figure 2e). This splitting completely disappears at 776.9 and 779.7 eV in the H-ZIF-8[1S] spectrum, but a dominant dipole transition at 778.1 eV, which is related to the polarized orbital occupation resulting from the spatial strain in confined spaces,[31,32] is observed. Interestingly, we observe a narrow, strong peak at 779.4 eV owing to the transition of more oxidized Co+ (t2g_6 e_g^0, low-spin) to the higher energy e_g orbital, which is distinguishable from Co+ (t2g_5 e_g^2, high-spin). The blue-shifted IR spectrum (Figure S25, Supporting Information) supports the presence of Co+ in H-ZIF-8[1S].[33] The electron-deficient Co+ strongly attracts hydroxyl group lone pair electrons to fill the completely empty e_g orbitals, as confirmed by the strong σ*-transition at 781.7 eV. The enriched σ bonds also result in shorter bond lengths (Figure 2b), as indicated by the shoulder below 1.70 Å.[34] Generally, the donor atoms involved in σ bonds become more electropositive, thereby allowing π-backbonding from the completely filled metallic t_2g orbital to the empty π* orbital of the donor atom.[35] Commonly, π-backbonding occurs between transition metals and organic ligands, but the predominant π* transition at 784.8 eV for H-ZIF-8[1S] indicates an increase in π-backbonding[36] not observed in bulk Co(OH)2 and ML-ZIFs[2L]. The N K-edge NEXAFS spectrum of H-ZIF-8[1S] (Figure 2f) shows a split π* transition, which is typically observed in extended π–π* systems, with π orbitals accessible to the K-edge excited electrons[37] at 400.3 and 399.7 eV (marked by arrows). Imidazole rings are stronger π-acceptors than hydroxyl groups, so that the ~0.7 eV shift of the π* transition is derived from the increased electron density resulting from π-backbonding by the surrounding Co(OH)2 SNPs. Furthermore, preferential π-backbonding between Co(OH)2 SNPs and 2-mim lowers the energy level of the t_2g orbital. The NEXAFS spectra show 22.5% Co+ and 44.6% π-backbonding, suggesting that Co(OH)2 SNPs are stabilized in H-ZIF-8[1S] (Figures S26 and S27, Supporting Information). The splitting of the π* transition indicates that Co is not substituted at the Zn tetrahedral site in ZIF-8, as supported by the fingerprint region of the IR spectrum (Figure S28, Supporting Information), which shows the N–Zn–N stretching mode at 420 cm⁻¹ without the N–Co–N stretching mode at 424 cm⁻¹. It has been reported that the lower electron density of Co results in a shift of 0.3 eV toward higher energies relative to Zn, which is more electron dense,[38] as observed in the case of ZIF-67 and ML-ZIFs[2L]. These observations are also supported by the absence of changes in the C 1s, N 1s, and Zn 2p XPS spectra and the Zn L-edge NEXAFS spectra (Figures S29–S32, Supporting Information).

The transmission of isolated water molecules via controlled hydrogen bonding affinity through the micropores is the key factor in enabling the autogenous production and stabilization of SNPs within multishell MOFs (Figure 3a). When water clusters react without EG, H-ZIF-8[1S] is not generated (Figure 3b) because the water clusters are too large to pass through the ZIF-8 aperture.[39] Solvated Co ions also do not penetrate the aperture because fully solvated Co ions are larger than the aperture (Figure S33, Supporting Information). For this reason, the strong hydrogen bonding affinity between EG and water molecules was exploited to isolate water molecules and allow them to pass through the 0.34 nm aperture of ZIF-8. Using the isolation approach, exposure to relatively large or small amounts of water produces sheet-like materials similar in structure to bulk Co(OH)2 or voids owing to the lack of water in ML-ZIFs[2L] (Figure 3b). With the optimal amount of water, the disassembled Co ions aggregate into Co ion clusters via ion–dipole interactions,[40] as verified by the bright spots in the HAADF-STEM image during the intermediate stage (Figure S34, Supporting Information). Then, Co atoms in the clusters are
laminated on the inner shell surface and pass through the micro pores, suppressing the growth of sheet-like structures (Figure S35, Supporting Information). The limited amount of water prevents complete hydration and regulates Co ion nucleation. Co ions are fed into the micro pores prior to bulk Co(OH)₂ formation and then form dinuclear Co(OH)₂ SNPs as...
water molecules are supplied one by one into the micropores. To confirm the isolation of water molecules by EG, we used the nuclear magnetic resonance (NMR) spectroscopy. EG has exchangeable hydroxyl groups that are indistinguishable on the $^1$H-$^1$H correlation spectroscopy (COSY), which explains a spin–spin coupling between EG and water molecule. However, a strong peak is observed at 5–6 ppm when water molecules are added to EG (Figure S37, Supporting Information). The $^1$H–$^1$H correlation spectroscopy (COSY), which can explain a spin–spin coupling between EG and water molecule, was conducted to find the reason for newly appeared peak by interpretation of the nature of the EG–water complex (Figures S38 and S39, Supporting Information). The cross-peaks, denoted as “OH–H$_2$O” (Figure 3c), demonstrate that the formation of bonds between water and EG occurs via the donation of water hydrogen atoms to the oxygen of EG. The $^1$H-NMR spectra (Figure 3d) show a broad peak at 63.3 ppm for the EG–water complex (intermolecular form), which is distinct from the sharp peak at 63.7 ppm for a pure EG (intramolecular form). We found that the intramolecular form transforms into the intermolecular form in the presence of water molecules and water isolation proceeds effectively, as evidenced by the fixed hydroxyl group and the interaction between EG and water molecule (Figures S40 and S41, Supporting Information). The two forms behave differently owing to their distinct structures. The broadness of the $^1$C-NMR peak originates from a slower tumbling rate owing to a large molecular size. The $^1$C diffusion-ordered spectroscopy (DOSY) was also utilized to determine the cluster size. The diffusion coefficient calculated for the intramolecular form indicates minor interactivity, as the value is not affected significantly by the presence of water. In contrast, the value determined for the intermolecular form is approximately 20-fold higher than that for the intramolecular form, indicating that the intermolecular form behaves like a huge cluster similar to a structure of consecutively bridged water molecules (Table S5, Supporting Information). As a result of this huge EG–water cluster, the isolated water molecules come into contact with the decomposable MOFs one by one, which prevents supersaturation of the reactants and thus avoids rapid nucleation. This approach not only makes hydrolysis controllable by regulating molecular water transfer, but also effectively suppresses nucleation, resulting in the autogenous production of uniform SNPs at high mass loadings. This method can be used to synthesize multishell MOFs via selective detachability, even in the presence of SNPs, which has previously been difficult to implement. After water molecule transfer, the $^1$H-NMR and $^1$C-NMR spectra show weakening of the hydroxyl group peak and disappearance of the water and intermolecular peaks (Figure 3e; Figure S42, Supporting Information), indicating that the intermolecular form returns to the original intramolecular form through reversion of the conformational changes caused by the presence of a water molecule (Figure S43, Supporting Information).

There are two strategies to increase the active sites for electrocatalysis, which include i) exposing more active sites or/and ii) loading more active materials. We increase the number of active sites by increasing the exposed active sites through the synthesis of dinuclear Co(OH)$_2$ SNPs, while simultaneously loading more active materials into the multishell structures (Figure 4a). On the basis of the strategically-designed system, we evaluated the dependence of the electrocatalytic performance of a Li–O$_2$ battery on Co(OH)$_2$ SNPs and multishell structures. For pristine ZIF-8, the charging curve shows a high overpotential with a potential gap of 1.494 V at half-capacity (Figure 4b). The Co(OH)$_2$ SNPs in H-ZIF-8[1S] remarkably improve the overpotential for the oxygen evolution reaction (OER) upon charging. The potential gap of 0.540 V at half-capacity is approximately 63.9% lower than that of pristine ZIF-8. Notably, the oxo groups between the Co ions promote the rapid oxidation of Co$^{2+}$ to Co$^{4+}$, facilitating the OER. The Co$^{4+}$ content of 22.5% enhances the OER kinetics by providing a shortcut to Co$^{4+}$ and more active sites. Although bulk Co(OH)$_2$ exhibits an improved overpotential, the discharge potential gradually decreases, whereas H-ZIF-8[1S] sustains a higher voltage (Figure S44, Supporting Information). This result indicates that bulk Co(OH)$_2$ is a good electrocatalyst for the OER upon charging, but unsuitable for the oxygen reduction reaction (ORR) upon discharging. Meanwhile, H-ZIF-8[1S] not only provides a large surface area for the accumulation of the discharge product Li$_2$O$_2$ during the ORR, but also helps to decompose Li$_2$O$_2$ efficiently during the OER. The specific gravimetric capacity clarifies the importance of Co(OH)$_2$ SNPs, but a smaller amount of H-ZIF-8[nS] is loaded as the number of shells increases at the same weight. In this reason, the specific geometric performance was utilized to determine the trend as the number of shell increases. The geometric performance (Figure 4c; Figure S45, Supporting Information) shows improved overpotentials under a constant areal current density, and the capacity also increases proportionally (Figure 4d; Figure S46 and Table S6, Supporting Information) to elucidate the origins for these enhancements, electrochemical impedance spectroscopy (EIS) measurements were performed. The Nyquist plot (Figure 4e) reveals two distinct changes that indicate improved performance. The charge-transfer resistance ($R_{ct}$) of H-ZIF-8[5S] is significantly smaller than that of pristine ZIF-8, as is the solution resistance ($R_s$) (Figure S47 and Table S7, Supporting Information). The reduced $R_s$ suggests that Co(OH)$_2$ SNPs enhance the poor electrical conductivity of ZIF-8 through the hopping transport mechanism. Furthermore, the reduced $R_{ct}$ indicates that the hollow structure minimizes transport resistance, as the diffusion time is inversely proportional to the diffusion length. The π-backbonding between Co(OH)$_2$ SNPs and micropores enhances stability owing to strong adhesion. Thus, a larger amount of Co(OH)$_2$ SNPs within multishell ZIF-8 should give excellent cycling stability, as demonstrated by the superiorof H-ZIF-8[5S] in Li–O$_2$ batteries (Figure 4f). Notably, excellent cycling in Li–O$_2$ batteries is also attributable to enhanced ORR and OER efficiencies. To investigate the structural stability of H-ZIF-8[1S] after electrochemical reactions, the XRD patterns were compared before and after 20 cycles (Figure S48, Supporting Information). The patterns were well maintained after 20 cycles, while the intensity associated with 011 facets was decreased due to the Li$_2$O$_2$ products of electrochemical reactions. This indicates that the structure of H-ZIF-8[1S] is stable during electrochemical reactions. Figure 4g shows that the capacity increases and the overpotential decreases at a higher mass loading with the same electrode volume, demonstrating that the SNP-embedded multishell ZIF-8 allows the scalable synthesis of electrocatalysts into electrodes.
Dinuclear Co(OH)$_2$ SNPs within multishell MOFs were autogenously synthesized and stabilized in chemically designable system using multilayer MOFs. The strong hydrogen bonding affinity between EG and water effectively prevented the formation of water clusters and acted as a mediator to balance the nucleation and the growth of SNPs within multilayer MOFs. It was confirmed that the transfer of isolated water molecules was a decisive factor in achieving SNP-embedded multishell hollow MOFs. In addition, alternating MOF layers with adjustable water stability enabled the step-by-step deterioration, resulting in the production of SNPs and the formation of multishell MOFs. In this process, we found that the π-backbonding between organic ligands and dinuclear SNPs could stabilize SNPs in the MOFs. Also, the π-backbonding and hollow structure led to enhanced electrical conductivity and minimized transport resistance. Moreover, the increased multishell number of H-ZIF-8[$n$S] allowed the scalable mass loadings of SNPs. Hence, these features of SNP-embedded multishell hollow MOFs resulted in high capacities and low overpotentials in Li–O$_2$ batteries. Consequently, this work provides a new method to produce and stabilize SNPs from alternating MOF layers so that it could be expanded to fabricate SNPs within various other types of frameworks. Additionally, the hopping charge transfer between SNPs stabilized by π-backbondings presents new possibilities to introduce electrical conduction in MOFs, thereby enabling to exploit the numerous advantages of MOFs likely to be applicable in a broad range of future technologies such as solar cells and sensors systems in addition to energy storage and conversion systems.

**Experimental Section**

**Materials:** Cobalt(II) nitrate hexahydrate (Co(NO$_3$)$_2$·6H$_2$O, 98.1%), zinc(II) nitrate hexahydrate (Zn(NO$_3$)$_2$·6H$_2$O, 98.1%), 2-methyleimidazole (2-mim, 99%), polyvinylpyrrolidone (PVP, mol. wt 10 000), EG (99.8%, anhydrous), cobalt(II) hydroxide (95%), methanol (99.9%), ethanol (99.5%), poly(tetrafluoroethylene) (PTFE, 1 µm), lithium perchlorate (LiClO$_4$, 99.99%), and tetraethylene glycol dimethyl ether (TEGDME,
were performed on a Bruker Avance 400 MHz spectrometer, equipped with a double crystal monochromator and the incident beam was detuned at the appropriate rate for harmonic rejection. All the NMR experiments were performed at the 10D beamline, where the synchrotron radiation was monochromatized using a Si(111) vacuum oven at 60°C for 24 h. Following centrifugation at 6000 rpm for 10 min, the powder was washed with methanol three times. Finally, the collected powder was dried in a vacuum oven at 60°C for 12 h. After weighing, the electrode was rinsed with acetone several times and vacuum dried in a glove box at 60°C for 12 h. For the Li–O₂ battery test, a Swagelok-type cell (Wellcos, Republic of Korea) was assembled with a Li foil anode, a glass fiber filter (Whatman, GF/D) as a separator, 1 M LiClO₄ in TEGDME as an electrolyte, and the as-prepared working electrode. Water in TEGDME was removed using molecular sieves (3 Å) for 10 days. The assembled cell was purged with a 25 cm³ min⁻¹ flow of pure oxygen for 3 h at 1.1 bar. The oxygen purging pressure was maintained during the Li–O₂ battery test and confirmed at an outlet valve using an MPM12 micromanometer (KIMO, France). The EIS measurements were conducted with a PGSTAT302N potentiostat/galvanostat (Metrohm Autolab B.V., Netherlands) using an FRA32MBA module.

**Supporting Information**

Supporting Information is available from the Wiley Online Library or from the author.

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**Conflict of Interest**

The authors declare no conflict of interest.

**Keywords**

Li–O₂ batteries, metal–organic frameworks, sub-nanometric particles, water molecule transfer
