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## Long-life sodium–sulfur batteries enabled by super-sodiophilic seeds†

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Sodium–metal batteries (SMBs) are an appealing sustainable low-cost alternative to lithium–metal batteries due to their high theoretical capacity ( $1165 \text{ mA h g}^{-1}$ ) and abundance of sodium. However, the practical viability of SMBs is challenged by a non-uniform deposition and uncontrollable growth of dendrites at the Na–metal anode. We employ here a super-sodiophilic  $\text{BiF}_3$  as a pre-planted seed to guide a uniform Na nucleation and deposition along the current collector. With the assistance of  $\text{BiF}_3$ , Na metal infiltrates easily into a carbon nanotube (CNT) paper matrix within a second. Such a Na anode exhibits a superior cycle life of  $>2000 \text{ h}$  with an extremely low overpotential of only  $13.5 \text{ mV}$  and a high coulombic efficiency of 99%. A full cell assembled with this anode and a sulfur cathode with a low negative to positive electrode capacity (N/P) ratio of  $\sim 2$  exhibits good cycling stability over 700 cycles. This work demonstrates a promising approach for the development of sustainable, low-cost SMBs.

### Broader context

Advanced energy-storage technologies are urgently needed to meet the growing demands for clean energy. It will be challenging to meet the projected future market demand with lithium-ion batteries (LIBs) due to their high cost. As a cost-effective alternative, sodium-based batteries are garnering significant attention due to the high abundance of sodium. Like sodium, sulphur is also abundant and inexpensive, and it offers an order of magnitude higher charge-storage capacity than the insertion-reaction cathodes currently employed in LIBs. However, room-temperature sodium–sulfur batteries generally suffer from low practical capacity and serious cycle life problems due to the severe sodium polysulfide shuttling and uncontrolled sodium dendritic growth. Herein, we report that commercial  $\text{BiF}_3$  particles serving as pre-planted seeds assist a uniform nucleation of sodium metal.  $\text{BiF}_3$  seeds facilitate an ultrafast loading of sodium metal into a CNT paper current collector within a second of immersion in molten sodium metal due to the super-sodiophilic property of  $\text{BiF}_3$ . Based on these features, symmetric cells assembled with the obtained anode exhibit a superior cycle life of  $>2000 \text{ h}$  with an extremely low overpotential of only  $13.5 \text{ mV}$ . This work provides a simple, scalable approach to Na-anode protection and is also instructive for other metal batteries.

## Introduction

The burgeoning demand for renewable energy storage and electric vehicles underscores the need for innovative battery technologies to sustain the growth.<sup>1–4</sup> Lithium-ion batteries (LIBs), despite being the most prevalent energy storage technology, fall short of meeting the evolving demands of modern portable smart grids, electric vehicles, and electronic devices.<sup>5–9</sup> As an alternative to LIBs for storing energy, sodium-based batteries offer great potential as next-generation energy storage systems due to the abundance and low cost of sodium-based minerals.<sup>10–12</sup> Sodium metal with a low redox potential ( $-2.71 \text{ V}$  vs. standard hydrogen

electrode) and a high theoretical specific capacity ( $1166 \text{ mA h g}^{-1}$ ) has garnered considerable research attention as an anode for sodium–metal batteries (SMBs) in recent years.<sup>13–16</sup>

For decades, significant endeavors have been undertaken to elucidate the mechanisms underlying sodium (Na) deposition, mitigate dendrite formation, and stabilize the solid electrolyte interphase (SEI) in pursuit of unleashing the complete potential of sodium–metal anodes.<sup>17–20</sup> Innovative approaches employed previously are use of solid-state electrolytes,<sup>21–24</sup> optimization of liquid electrolytes,<sup>25–28</sup> engineering of sodiophilic hosts/current collectors,<sup>29–32</sup> and constructing interfacial protective layers.<sup>33–36</sup> Of the various strategies, the engineering of sodiophilic hosts/current collectors is among the most promising, which can guide a uniform Na nucleation, enabling SEI stabilization, dendrite suppression, and volume change accommodation.<sup>37–39</sup> However, the preparation procedures of many reported hosts are complicated, posing a challenge to practical viability.

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Herein, we report commercial  $\text{BiF}_3$  particles serving as pre-implanted seeds to guide a uniform Na nucleation and deposition in a matrix. The pulverization of  $\text{BiF}_3$  particles, often regarded as a significant drawback when employed as a cathode material, is envisioned in this study as a notable advantage. This behavior facilitates the rearrangement of nucleating seeds, consequently simplifying the process of introducing pre-implanted seeds. In addition,  $\text{BiF}_3$  exhibits super-sodiophilic property, enabling an ultrafast loading of Na metal into carbon nanotube (CNT) paper current collector within a second of immersion in molten Na metal. Benefiting from these two distinctive features, symmetric cells assembled with the obtained anode exhibit a superior cycle life of  $>2000$  h with an extremely low overpotential of only 13.5 mV. A pouch cell paired with a sulfur cathode and the obtained sodiophilic anode with a low negative to positive electrode capacity (N/P) ratio of  $\sim 2$  exhibits stable cycling performance over 50 cycles with a capacity of 661  $\text{mA h g}^{-1}$ .

## Results and discussion

### Synthesis route and mechanism

The synthetic route for the  $\text{Na}@\text{CNT-BiF}_3$  self-supporting electrodes is schematically shown in Fig. 1. It can be easily prepared by simply mixing CNT and  $\text{BiF}_3$  via ultrasonication, followed by filtering, drying, and diffusing molten Na. It is worth noting that the CNT- $\text{BiF}_3$  electrode is super-sodiophilic and can quickly adsorb molten Na metal into the material. As shown in Fig. S1 (ESI<sup>†</sup>), since commercialization is based on Cu foil current collectors rather than simple self-supporting electrodes, we have also achieved excellent electrochemical performance by coating commercial  $\text{BiF}_3$  paste onto a Cu foil. In addition, the  $\text{BiF}_3$  particles in the  $\text{Na}@\text{CNT-BiF}_3$  and  $\text{Na}@\text{Cu-BiF}_3$  electrodes gradually pulverize and uniformly cover CNT after alloying/dealloying of  $\text{BiF}_3$  with increasing number of cycles,<sup>40,41</sup> effectively improving the uniformity and electrochemical properties of the  $\text{Na}@\text{CNT-BiF}_3$  and  $\text{Na}@\text{Cu-BiF}_3$  electrodes materials.

### Electrochemical performance of sodium–metal anode

To further assess the sodiophilicity of CNT@- $\text{BiF}_3$  and CNT electrodes for molten Na, the time required for the electrodes to fully adsorb molten Na metal was compared macroscopically (Fig. 2a and Videos S1 and S2, ESI<sup>†</sup>). Notably, the CNT@ $\text{BiF}_3$  electrode fully adsorbed molten Na metal within a second, whereas the CNT electrode took 52 s only to adsorb a small

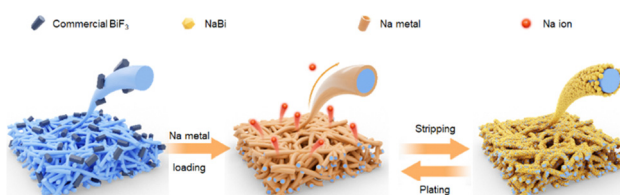


Fig. 1 Synthetic route and mechanism. Illustration of the synthetic route and mechanism after the cycling of  $\text{Na}@\text{CNT-BiF}_3$ .

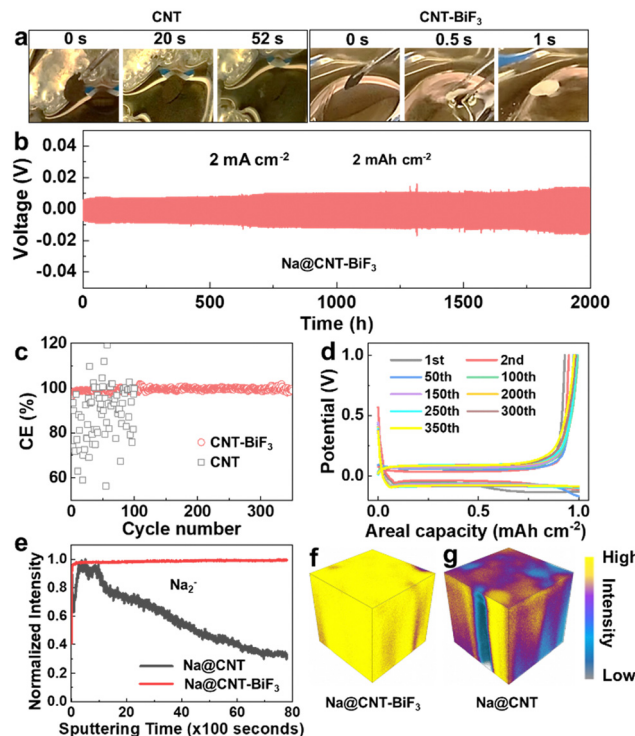


Fig. 2 Characterization and electrochemical performance of  $\text{Na}@\text{CNT-BiF}_3$ . (a) Digital images of CNT and CNT- $\text{BiF}_3$  electrodes, comparing the sodiophilicity for molten Na metal. (b) Voltage profiles of the  $\text{Na}@\text{CNT-BiF}_3||\text{Na}@\text{CNT-BiF}_3$  cells measured at  $2 \text{ mA cm}^{-2}$ . (c) Coulombic efficiency of Na plating/stripping using Cu electrodes with  $\text{Na}@\text{CNT-BiF}_3$  or bare  $\text{Na}@\text{CNT}$  at  $1 \text{ mA cm}^{-2}$ . (d) The voltage–capacity curves of Na plating/stripping using Cu electrodes with  $\text{Na}@\text{CNT-BiF}_3$  at  $1 \text{ mA cm}^{-2}$ . (e) TOF-SIMS depth profiling of  $\text{Na}_2^-$  secondary ions (a) after cycling  $\text{Na}@\text{CNT}$  and  $\text{Na}@\text{CNT-BiF}_3$ . TOF-SIMS 3D visualization of after cycling (f)  $\text{Na}@\text{CNT-BiF}_3$  and (g)  $\text{Na}@\text{CNT}$ .

amount of molten Na, demonstrating the high sodiophilicity of  $\text{BiF}_3$  particles in greatly improving the sodiophilicity of CNT@ $\text{BiF}_3$ . The cycling stability of the  $\text{Na}@\text{CNT-BiF}_3$  anode was further assessed with symmetric cells at a current density of  $2 \text{ mA cm}^{-2}$  and an areal capacity of  $2 \text{ mA h cm}^{-2}$ . As seen in Fig. 2b and Fig. S2 (ESI<sup>†</sup>), the symmetric cell with  $\text{Na}@\text{CNT-BiF}_3$  displays an overpotential of only 6 mV initially, followed by a slight increase of up to 13.5 mV, but shows a remarkable lifespan of over 2000 h. The coulombic efficiency (CE) is a very vital parameter to reflect the side reactions between electrodes and electrolytes. The CE of the cell with the  $\text{Cu}||\text{Na}@\text{CNT}$  electrode exhibits strong irregular fluctuations (Fig. 2c), which may be due to the CNT surface having poor sodiophilicity. The lack of an effective nucleation surface leads to a loss of active Na during plating/stripping, accompanied by a reconstruction and cracking of the SEI. As a comparison, the  $\text{Cu}||\text{Na}@\text{CNT-BiF}_3$  cell presents a well-maintained CE of 99% after 350 cycles, demonstrating a great improvement in Na plating/stripping reversibility. Additionally, the corresponding discharge–charge curves of the  $\text{Cu}||\text{Na}@\text{CNT-BiF}_3$  cell show a low Na nucleation overpotential with constant, overlapping plateaus during repetitive cycling (Fig. 2d).

To further investigate the interphase composition formed on the surface of the Na@CNT and Na@CNT–BiF<sub>3</sub>, time-of-flight-secondary ion mass spectrometric (TOF-SIMS) depth profiling was employed, which is an advanced technique to identify the chemical composition of the solid surface within its depth.<sup>42,43</sup> As shown in Fig. 2e, the depth profiles reveal the uniformity of Na infiltration into the matrix in Na@CNT and Na@CNT–BiF<sub>3</sub> anode *via* the Na<sub>2</sub><sup>−</sup> secondary-ion as a representative species. The Na<sub>2</sub><sup>−</sup> intensity in Na@CNT anode shows two peaks consecutively and then decreases continuously, which indicates that the poor sodiophilicity of Na@CNT results in a non-uniform incorporation of metallic Na. The 3D reconstruction of the elemental depth profiles of Na@CNT in Fig. 2g further reaffirms the inhomogeneous Na metal distribution. It is reasonable to posit that the non-uniformity leads to dendrite formation and irregular cycling, as observed in Fig. 2c. On the other hand, the continuous signal for Na<sub>2</sub><sup>−</sup> intensity in Na@CNT–BiF<sub>3</sub> indicates a uniform infiltration of Na metal, despite the presence of large BiF<sub>3</sub> particles in the CNT. In further support, the 3D reconstruction of the Na@CNT–BiF<sub>3</sub> electrode in Fig. 2f shows a homogeneous Na<sub>2</sub><sup>−</sup> signal over the whole investigated depth. The continual abundance of seeding sites leads to a consistent performance of the cell, observed in Fig. 2c and d.

### Sodium storage mechanism and the behavior of sodium–metal anode

In addition, to observe the morphological changes of Na metal at the CNT–BiF<sub>3</sub> electrode microscopically, the top-surface and cross-sectional SEM images of CNT–BiF<sub>3</sub>, Na@CNT–BiF<sub>3</sub> (before cycling), and Na@CNT–BiF<sub>3</sub> (after 200 cycles) electrodes

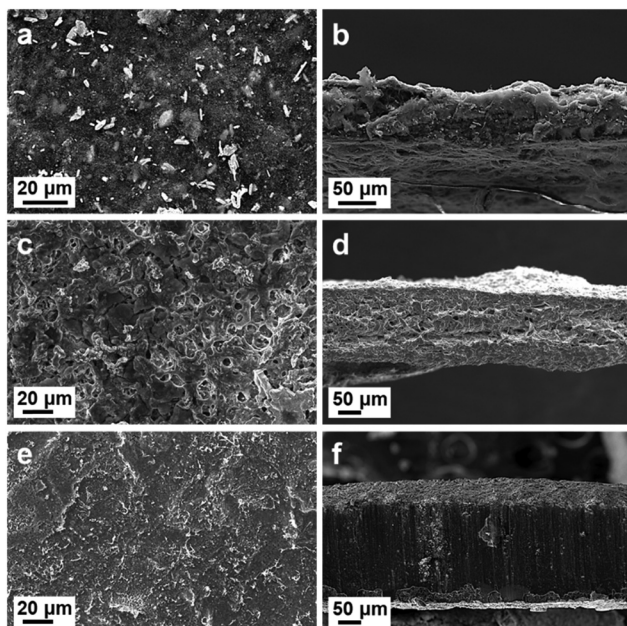


Fig. 3 Morphological characterization of the anode. Top-surface SEM of (a) BiF<sub>3</sub>–CNT, (c) Na@CNT–BiF<sub>3</sub> after cycling, and (e) Na@CNT–BiF<sub>3</sub> (after 200 cycles). Cross-sectional morphologies of (b) BiF<sub>3</sub> before cycling, (d) Na@CNT–BiF<sub>3</sub> (before cycling), and (f) Na@CNT–BiF<sub>3</sub> (after 200 cycles).

are compared in Fig. 3. Fig. 3(a and b) shows uniformly distributed, large (micron-sized) BiF<sub>3</sub> particles in the CNT–BiF<sub>3</sub> electrode. The top-surface and cross-sectional morphologies of the CNT–BiF<sub>3</sub> electrode after adsorbing molten Na are shown in Fig. 3(c and d) shows the top-surface and cross-sectional morphologies of the CNT–BiF<sub>3</sub> electrode after adsorbing molten Na (before cycling). As seen, there are no obvious BiF<sub>3</sub> particles on the top surface, due to the rapid pulverization and quick coverage by Na metal during the adsorption of molten Na. In addition, from the cross-sectional SEM image, it can be seen that there are still many pores and gap structures after the adsorption of Na metal, which can effectively buffer the volume expansion during the cycling process. To this end, the structural changes of the Na@CNT–BiF<sub>3</sub> electrode after 200 cycles can be observed in Fig. 3(e and f). The SEM images show that the size of the surface BiF<sub>3</sub> particles is smaller and more uniformly distributed. In addition, the pores in the cross-sectional SEM are filled and thus become denser, which may be attributed to the continuous alloying/de-alloying of BiF<sub>3</sub> during the cycling process. The repeated alloying process causes BiF<sub>3</sub> to break down into small particles that get homogeneously distributed such that they occupy more volume compared to the initial stage. Thus, the void spaces that existed in Na@CNT–BiF<sub>3</sub> electrode before cycling are filled during the Na plating process.

To better understand the Na storage mechanism, Na plating behavior on Cu, CNT, and CNT–BiF<sub>3</sub> substrates was studied. The potential–capacity curves in Fig. 4a show a deep dip in the potential caused by the Na–metal nucleation barrier. This nucleation potential for Cu, CNT, and CNT–BiF<sub>3</sub> are, respectively, 428, 383, and 129 mV. The low overpotential for CNT–BiF<sub>3</sub> indicates that it is a highly favorable substrate to plate Na–metal with a fast Na<sup>+</sup> diffusion rate. It also means that BiF<sub>3</sub> is labile for conversion by easily reacting with Na. The conversion was confirmed by comparing the X-ray photoelectron spectroscopy (XPS)

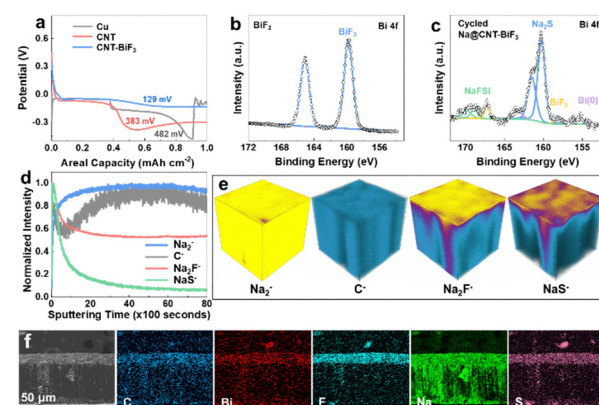


Fig. 4 Electrochemical behaviors and characterization. (a) Galvanostatic discharge curves (<0 V) at a current rate of 1 mA cm<sup>−2</sup> in Cu, CNT, and CNT–BiF<sub>3</sub> electrodes. High-resolution Bi 4f XPS data of (b) BiF<sub>3</sub> and (c) Na@CNT–BiF<sub>3</sub> after cycling. (d) Depth profiles of Na<sub>2</sub><sup>−</sup>, C<sup>−</sup>, Na<sub>2</sub>F<sup>−</sup>, and NaS<sup>−</sup> secondary ions and (e) 3D visualization after many cycles of Na plating in Na@CNT–BiF<sub>3</sub>. (f) Cross-sectional SEMs of C, Bi, F, Na, and S elements mapping in selected areas of Na@CNT–BiF<sub>3</sub> after many cycles.

data of CNT-BiF<sub>3</sub> (Fig. 4b) and Na@CNT-BiF<sub>3</sub> (Fig. 4c). As expected, the spectrum of CNT-BiF<sub>3</sub> shows the Bi 4f<sub>7/2</sub> peak at 160.8 eV. On plating Na on the CNT-BiF<sub>3</sub> electrode, a new Bi 4f<sub>7/2</sub> peak at 155.8 eV corresponding to metallic Bi is observed. This peak appears at a lower binding energy than expected for metallic Bi (156.9 eV), potentially due to the formation of Na-Bi alloy, indicating that the Na is not just physically plated, but rather chemically reacting with and converting BiF<sub>3</sub>. Furthermore, the peak for BiF<sub>3</sub> persists even after sodium plating, implying that BiF<sub>3</sub> progressively undergoes pulverization followed by conversion to metallic Bi with cycling. This steady conversion process ensures continuous availability of BiF<sub>3</sub>, which acts as seeds and thus ensures uniform plating.

To better understand the surface and bulk characteristics of the Na@CNT-BiF<sub>3</sub> anode cycled in a localized high-concentration electrolyte (LHCE), ToF-SIMS on the cycled anode was performed. The LHCE consists of sodium bis(fluorosulfonyl) imide (NaFSI):1,2-dimethoxyethane (DME):1,1,2,2-tetrafluoroethyl 2,2,3,3-tetrafluoropropyl ether (TTE) in 1:1.2:1 (molar ratio). Fig. 4d shows the depth profiles and Fig. 4e shows the 3D reconstruction of the secondary-ion species, such as Na<sub>2</sub><sup>-</sup>, C<sup>-</sup>, Na<sub>2</sub>F<sup>-</sup>, and NaS<sup>-</sup>, which are representative of, respectively, metallic Na, CNT, BiF<sub>3</sub>, and LHCE electrolyte. It can be seen from the depth profiles and 3D model that the signals for Na<sub>2</sub><sup>-</sup> and C<sup>-</sup> gradually increase to reach a uniform and stable level by the end of sputtering, which is similar to the trend of secondary ion Na<sub>2</sub><sup>-</sup> signal in the original Na@CNT-BiF<sub>3</sub> anode (Fig. 2f), suggesting that the structure of CNT matrix and the homogenous Na distribution are maintained across cycling. In addition, the SEI layer was analyzed by the secondary ions Na<sub>2</sub>F<sup>-</sup> and NaS<sup>-</sup>. First, the signal of Na<sub>2</sub>F<sup>-</sup> gradually decreases until about 2000 s of sputtering time and then it stabilizes, indicating that a compact NaF-rich SEI is formed, as seen in the 3D reconstruction. This layer suppresses the growth of dendrites, while providing fast ion transport, which is responsible for the high CE observed. The NaS<sup>-</sup> signal from the electrolyte follows a trend similar to that of Na<sub>2</sub>F<sup>-</sup> in the SEI layer, which indicates that the inorganic-rich NaF and Na<sub>2</sub>S layers protect the underlying Na-metal from continuous electrolyte exposure and decomposition. Fig. 4f displays the cross-sectional SEM micrograph along with the elemental mapping distributions of C, Bi, F, Na, and S elements across a cycled Na@CNT-BiF<sub>3</sub> anode. The carbon signal is uniformly distributed in the whole cross-section, which confirms that the structure of the electrode remains intact and stable. The concentration of F and S near the surface aligns with the ToF-SIMS data, reaffirming the NaF and Na<sub>2</sub>S layers shield the Na-metal from electrolyte degradation.

The above studies show that the Na@CNT-BiF<sub>3</sub> forms and maintains a stable interface through cycling.

To better understand the mechanism by which Na-metal can easily diffuse into a matrix of BiF<sub>3</sub> and to understand the chemical difference in the Na stripping/plating behavior between the CNT and CNT-BiF<sub>3</sub> anodes, synchrotron-based *in operando* X-ray diffraction (XRD) experiments were performed (the test mechanism schematic is shown in Fig. S3, ESI<sup>†</sup>). Fig. 5 presents the *in operando* XRD results of Na||CNT and Na||CNT-BiF<sub>3</sub> cells cycled with the LHCE in custom coin

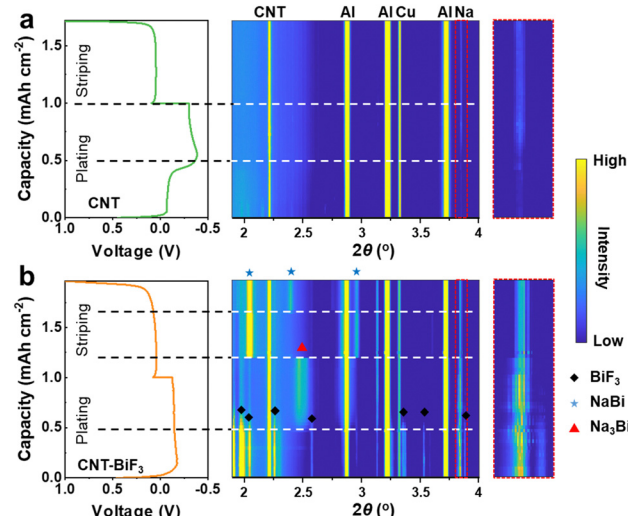


Fig. 5 *In operando* X-ray diffraction (XRD) characterization. Charge and discharge curves of (a) Na@CNT||Na@CNT and (b) Na@CNT||Na@CNT-BiF<sub>3</sub> coin cell and the corresponding diffraction patterns.

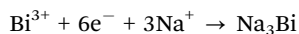
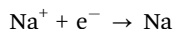
cells with a window, as described in our previous work.<sup>44</sup> The cells were tested at 1 mA cm<sup>-2</sup> for one full plating/stripping process, during which the XRD pattern was simultaneously recorded to probe the Na plating/stripping with a small volume where the X-rays are focused as presented by the contour plot next to the voltage profile.

In the contour plot of the cell with Na@CNT anode (Fig. 5a), peaks corresponding to Na metal plating on the CNT matrix can be detected when about 0.5 mA h cm<sup>-2</sup> is plated, and the peak intensity grows stronger with continuous plating. While stripping, the intensity reduces, but a faint intensity is still detected at the end of the stripping process, which can act as the nucleation site during the subsequent plating steps. In addition, the intensity of the characteristic Na peaks after complete desodiation is significantly stronger than that before initial plating, indicating significant irreversibility of Na plating/stripping, which may be responsible for the growth of dendrites. In contrast, as shown in Fig. 5b, during the sodium plating process of CNT-BiF<sub>3</sub> anode, strong Na metal, and BiF<sub>3</sub> characteristic peaks are present at the beginning of the plating step. The intensity of the Na-metal peak is much stronger in the case of CNT-BiF<sub>3</sub> compared to pure CNT, indicating that the probed volume quickly fills with Na metal, owing to BiF<sub>3</sub> acting as a strong seeding layer, while the sparse Na distribution in the case of CNT is reflected in a weaker signal. This also points to the good sodiophilicity of the matrix with BiF<sub>3</sub>. By the time 0.5 mA h cm<sup>-2</sup> is plated, the signal for Na<sub>3</sub>Bi can be observed, while the Na metal signal intensifies. This indicates that the conversion-alloying process drives the dense plating of Na in the matrix. The formation of Na<sub>3</sub>Bi and NaBi, as indicated by the *in-situ* XRD along with the presence of NaF signal in the ToF-SIMS and SEM data in Fig. 4, suggests that the facile conversion of BiF<sub>3</sub> into Na-Bi alloys and NaF is the reason for the high sodiophilicity of a BiF<sub>3</sub> containing matrix, which allows for the ultra-quick melt-infusion of Na metal. In addition, the intensity

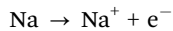
of the characteristic Na peak after complete sodium removal is significantly weaker than that before initial plating, suggesting that Na plating/stripping is reversible.

During the stripping process, the characteristic peak of  $\text{Na}_3\text{Bi}$  is gradually transformed into  $\text{NaBi}$ , and the characteristic peak of Na metal is obviously and gradually weakened. The remaining Na metal and the  $\text{NaBi}$  alloy act as an excelling seeding layer in the subsequent plating step, thus guiding a dendrite-free nucleation and growth of Na.<sup>45,46</sup> In essence, the addition of  $\text{BiF}_3$  improves the sodiophilicity of CNT, resulting in a highly reversible Na plating/stripping and alloying/de-alloying process for  $\text{Na}@\text{CNT}-\text{BiF}_3$  anode. Overall, from the above XRD data, in a melt-infiltrated  $\text{Na}@\text{CNT}-\text{BiF}_3$  anode, the following (electro)chemical process can be envisioned:

Plating process:



Stripping process:



### Electrochemical performance of $\text{Na}@\text{CNT}-\text{BiF}_3$ and $\text{Na}@\text{Cu}-\text{BiF}_3$

Generally, self-supporting electrodes can provide higher energy density, but practical applications necessitate the use of Cu foil as the current collector. Hence, for further practical and scalable application, the electrochemical performance of  $\text{BiF}_3$  coated onto Cu collector was tested. Here, different contents of  $\text{BiF}_3$  electrodes were assembled into  $\text{Na}||\text{Cu}$  cells to characterize the CE and corresponding voltage profile curves for sodium plating/stripping. Fig. 6(a and b) shows the CE and the corresponding voltage profiles for the blank control group of Cu anode with 0%  $\text{BiF}_3$ . It is observed that the CE is very unstable and the voltage profiles fluctuate with high sodium–metal nucleation potentials and low capacity. These fluctuations may be due to the dendrite growth leading to a short circuit of the cell. Fig. 6(c–h) compares the CE with the corresponding voltage profiles for the Cu anodes with 5%  $\text{BiF}_3$ , 20%  $\text{BiF}_3$ , and 40%  $\text{BiF}_3$ . The CE is increased when 5%  $\text{BiF}_3$  is added, and its voltage profiles are relatively smooth with the CE being stabilized only after cycling up to 100 cycles. However, the CE becomes much more stable when  $\text{BiF}_3$  is added up to 20% and the corresponding voltage profiles are stable with a low Na metal nucleation overpotential. When  $\text{BiF}_3$  is increased to 40%, the cells cycle with relatively good stability. However, the CE fluctuations are slightly increased compared to that with 20%  $\text{BiF}_3$ . The increased content of  $\text{BiF}_3$  and its longer alloying transformation affect the CE. Therefore, in consideration of cost and economics, 20%  $\text{BiF}_3$  addition improves both the sodiophilicity of the electrode and keeps the Na loss at less than 5% in the first cycle.

To demonstrate the practical application of the conversion composite anode, high-energy C/S cathode was matched with

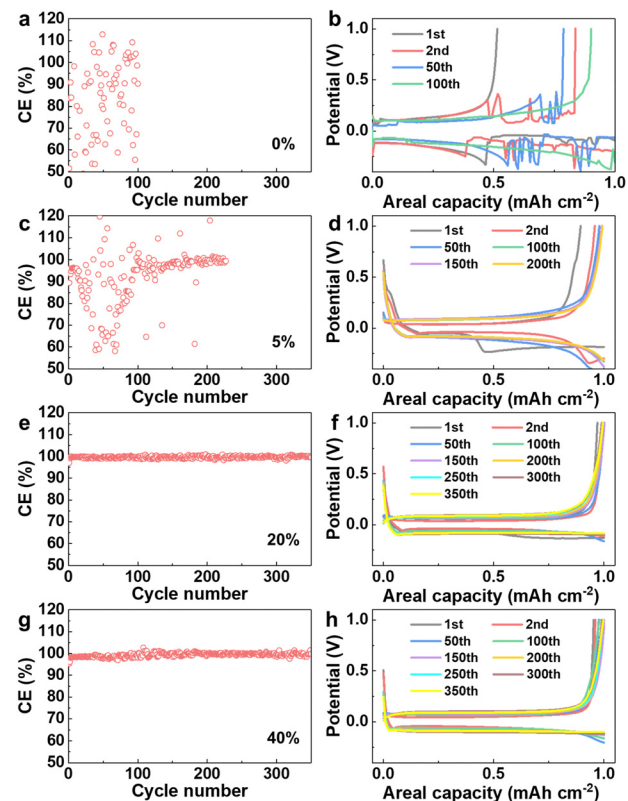


Fig. 6 Characterization and electrochemical performance of  $\text{Cu}-\text{BiF}_3$  anodes. Coulombic efficiency and voltage–capacity curves of Na plating/stripping in (a) and (b)  $\text{Cu}-0\%$   $\text{BiF}_3$ , (c) and (d)  $\text{Cu}-5\%$   $\text{BiF}_3$ , (e) and (f)  $\text{Cu}-20\%$   $\text{BiF}_3$  (g) and (h)  $\text{Cu}-40\%$   $\text{BiF}_3$  at  $1 \text{ mA cm}^{-2}$ .

bare Na and  $\text{Na}@\text{CNT}-\text{BiF}_3$  anodes to make Na–S cells. In these cells, the sulfur loading is  $3 \text{ mg cm}^{-2}$ , and the electrolyte-to-sulfur (E/S) ratio is controlled to be  $7 \mu\text{L mg}^{-1}$  with an ultra-low N/P ratio of 2.8 to showcase the stability of the anode. Fig. 7a displays the voltage profiles of  $\text{Na}@\text{CNT}-\text{BiF}_3||\text{C/S}$  cell at different cycles at 0.1C rate ( $1\text{C} = 1675 \text{ mA g}^{-1}$ ) within the voltage range of 0.8–2.8 V. It is noteworthy that there is no obvious irreversible capacity loss region with Na polysulfide (NaPSs) transformation in the voltage profiles of the initial cycles. The stability of the charge/discharge curves implies that the  $\text{Na}@\text{CNT}-\text{BiF}_3$  anode surface is resistant to the shuttle of polysulfides.

The  $\text{Na}@\text{CNT}-\text{BiF}_3||\text{C/S}$  cell presents an initial discharge and charge capacities of, respectively,  $1010$  and  $972 \text{ mA h g}^{-1}$  with an initial coulombic efficiency (ICE) of 104%, as shown in Fig. 7b. This is attributed to the introduction of super-sodiophilic  $\text{BiF}_3$  particles and the rapid self-adsorption of sodium by  $\text{Na}@\text{CNT}-\text{BiF}_3$ , improving the kinetic properties of  $\text{Na}^+$ . Even through 200 cycles, the capacity remains at  $862 \text{ mA h g}^{-1}$ , corresponding to a capacity retention of 85%. The CE at the 200th cycle is nearly 100%, which demonstrates minimal polysulfide shuttling occurs during cycling. On the other hand, the cell with bare Na anode shows poor cycling performance with a rapid capacity fade within 75 cycles and finally displays a very low capacity of only  $165 \text{ mA h g}^{-1}$  after 150 cycles. Furthermore, the CE fluctuates during cycling, suggesting irreversible reactions between the

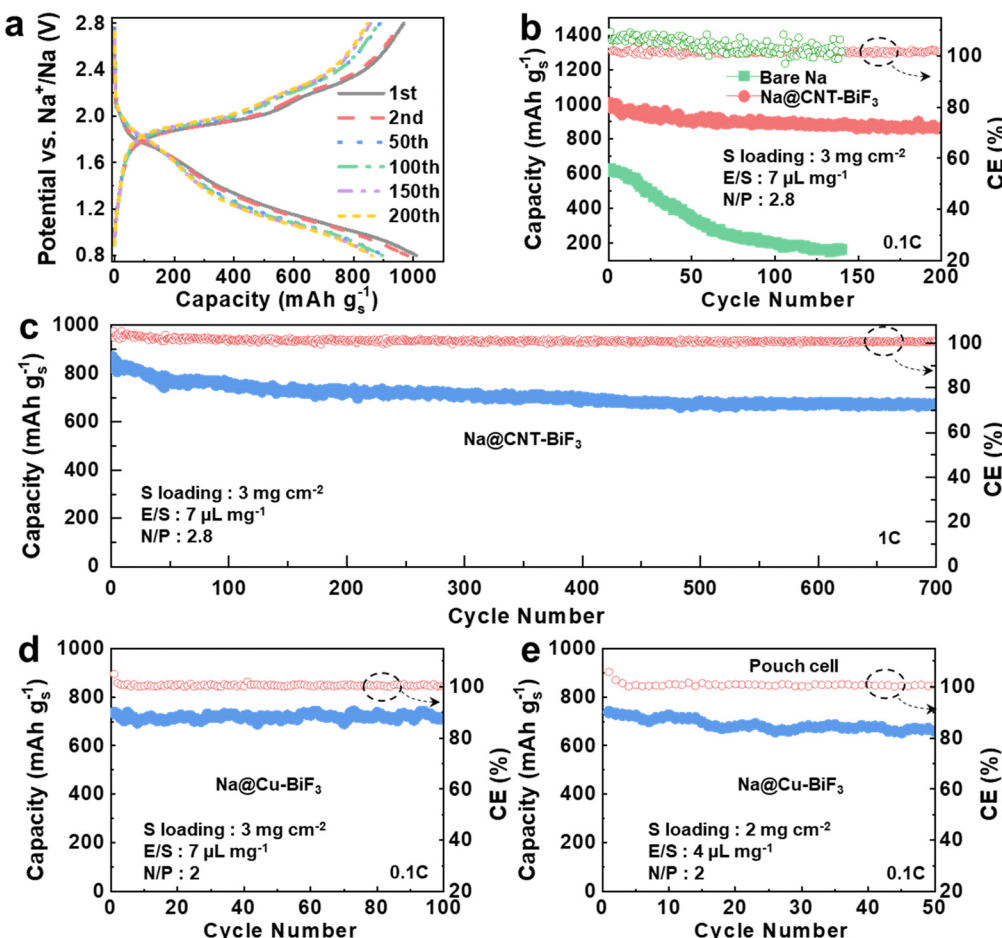


Fig. 7 Electrochemical performance of Na@CNT-BiF<sub>3</sub> and Na@Cu-BiF<sub>3</sub>. (a) Voltage profiles of Na@CNT-BiF<sub>3</sub> at various cycles. (b) Cycling performances of bare Na and Na@CNT-BiF<sub>3</sub> electrodes at 0.1C rate. (c) Cycling performance of Na@CNT-BiF<sub>3</sub> electrodes at 1C rate. Cycling performances of Na@Cu-BiF<sub>3</sub> electrodes in (d) coin cell and (e) pouch cell at 0.1C rate.

electrolyte and the bare Na–metal anode lead to capacity loss. In essence, the Na@CNT-BiF<sub>3</sub> anode demonstrates significant potential for high-performance Na–S batteries with impressive capacity, minimal capacity decay, and prolonged cycling lifespan, outperforming numerous Na–S batteries (Table S1, ESI<sup>†</sup>). The Na@CNT-BiF<sub>3</sub>||C/S cell was further tested at a higher current density of 1C rate to reveal the kinetics of the reaction and the structural stability of the anode. Fig. 7c shows that the Na@CNT-BiF<sub>3</sub>||C/S cell can still provide a discharge capacity of 674 mA h g<sup>-1</sup> even after 700 cycles, corresponding to a capacity retention of 77%, indicating a highly reversible reaction in the cell.

As mentioned before, Na@Cu-BiF<sub>3</sub> is a practical alternative to the freestanding Na@CNT-BiF<sub>3</sub> as it has a metallic current collector, allowing for the processing of large electrodes. To validate this, the cycling performance of the Na@Cu-BiF<sub>3</sub>||C/S cell was tested with a sulfur loading of 3 mg cm<sup>-2</sup>, a low E/S of 7 µL mg<sup>-1</sup>, a low N/P ratio of 2, and at a current of 0.1C, as shown in Fig. 7d and Fig. S4 (ESI<sup>†</sup>). The Na@Cu-BiF<sub>3</sub> anode exhibits an initial capacity of 735 mA g<sup>-1</sup> and excellent cycling stability, with a capacity retention of close to 100% after 100 cycles. The presence of a Cu current collector affords the construction of Na–S pouch cells as tabs can be easily welded.

Therefore, Na@Cu-BiF<sub>3</sub>||C/S pouch cells were assembled to validate our concept, featuring a 2 mg cm<sup>-2</sup> loading, a low E/S of 4 µL mg<sup>-1</sup>, and a low N/P ratio of 2. As depicted in Fig. 7e, the pouch cell presents a capacity of 661 mA h g<sup>-1</sup> at 0.1C rate after 50 cycles, indicating the effective utilization of sulfur operating with low E/S and limited N/P ratios. This further validates that BiF<sub>3</sub> super-sodiophilic seeds can accelerate sodium plating/stripping, improving the NaPSs kinetic performance and reducing the shuttle effect of NaPSs. These results highlight the promise of integrating Na@Cu-BiF<sub>3</sub> in the development of high-performance, practically feasible Na–S batteries.

## Conclusions

In summary, BiF<sub>3</sub> is demonstrated to be a super-sodiophilic seed for Na–metal anodes in sodium–metal batteries. When BiF<sub>3</sub> is integrated into a matrix of CNT, molten Na metal can be incorporated into the substrate in a facile and rapid manner owing to the conversion-alloying reaction of BiF<sub>3</sub>. The Na@CNT-BiF<sub>3</sub> anode thus forms a NaF<sup>-</sup> and Na<sub>2</sub>S-rich interphase with the electrolyte when cycled in an LHCE. This

interphase is compact, yet robust, to prevent continuous electrolyte degradation at the anode, which enables long-term cycling. The constant pulverization and conversion of  $\text{BiF}_3$  through cycling provides an efficient seed layer that suppresses dendrite formation. In addition to the free-standing format,  $\text{BiF}_3$  can be coated onto a metallic current collector, such as Cu foil, and mixed with molten Na metal to yield an anode. The  $\text{Na@Cu-BiF}_3$  anodes could be easily incorporated into prototype pouch cells, which showed excellent cycling stability in Na-S batteries, especially while operating at a low N/P ratio of 2. This work provides a simple, scalable approach forward to Na-anode protection and is also instructive for other metal batteries with different working conditions.

## Author contributions

J. H., A. B., and A. M. conceived the idea. J. H. performed cathode formulations and collected the electrochemical data. A. B performed the pouch cell experiments. A. M. supervised the project. All authors were involved in discussing the results and writing and reviewing the manuscript.

## Data availability

The datasets generated during and/or analyzed during the current study are available from the corresponding author upon request.

## Conflicts of interest

There are no conflicts to declare.

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## Notes and references

- Y. Liu, X. Tao, Y. Wang, C. Jiang, C. Ma, O. Sheng, G. Lu and X. W. Lou, *Science*, 2022, **375**, 739–745.
- X. Liu, J. Zhu, X. Wang, L. Yue, W. Wang, B. Wang, D. Shen and Y. Li, *Adv. Funct. Mater.*, 2023, **33**, 2209388.
- X. Liu, J. Zhu, L. Yue, X. Wang, W. Wang, T. Zheng and Y. Li, *Small*, 2022, **18**, 2204552.
- Q. Pang, J. Meng, S. Gupta, X. Hong, C. Y. Kwok, J. Zhao, Y. Jin, L. Xu, O. Karahan and Z. Wang, *Nature*, 2022, **608**, 704–711.
- L. Jiang, Y. Lu, C. Zhao, L. Liu, J. Zhang, Q. Zhang, X. Shen, J. Zhao, X. Yu and H. Li, *Nat. Energy*, 2019, **4**, 495–503.
- Y. Zou, Z. Ma, G. Liu, Q. Li, D. Yin, X. Shi, Z. Cao, Z. Tian, H. Kim and Y. Guo, *Angew. Chem., Int. Ed.*, 2023, **135**, e202216189.
- X. Liu, X. Wang, Y. Zhou, B. Wang, L. Zhao, H. Zheng, J. Wang, J. Liu, J. Liu and Y. Li, *Adv. Mater.*, 2024, **36**, 2308447.
- S. Zhang, Q. Fan, Y. Liu, S. Xi, X. Liu, Z. Wu, J. Hao, W. K. Pang, T. Zhou and Z. Guo, *Adv. Mater.*, 2020, **32**, 2000380.
- J. Meng, X. Hong, Z. Xiao, L. Xu, L. Zhu, Y. Jia, F. Liu, L. Mai and Q. Pang, *Nat. Commun.*, 2024, **15**, 596.
- T. Deng, X. Ji, L. Zou, O. Chiekezi, L. Cao, X. Fan, T. R. Adebisi, H.-J. Chang, H. Wang and B. Li, *Nat. Nanotechnol.*, 2022, **17**, 269–277.
- C. Xu, Y. Ma, J. Zhao, P. Zhang, Z. Chen, C. Yang, H. Liu and Y. S. Hu, *Angew. Chem., Int. Ed.*, 2023, **62**, e202217761.
- T. Liu, Y. Yang, S. Cao, R. Xiang, L. Zhang and J. Yu, *Adv. Mater.*, 2023, **35**, 2207752.
- X. Xia, S. Xu, F. Tang, Y. Yao, L. Wang, L. Liu, S. He, Y. Yang, W. Sun and C. Xu, *Adv. Mater.*, 2023, **35**, 2209511.
- H. Liu, X. Zheng, Y. Du, M. C. Borrás, K. Wu, K. Konstantinov, W. K. Pang, S. Chou, H. Liu and S. Dou, *Adv. Mater.*, 2024, **36**, 2307645.
- R. Zhuang, X. Zhang, C. Qu, X. Xu, J. Yang, Q. Ye, Z. Liu, S. Kaskel, F. Xu and H. Wang, *Sci. Adv.*, 2023, **9**, eadh8060.
- J. He, A. Bhargav, L. Su, J. Lamb, J. Okasinski, W. Shin and A. Manthiram, *Nat. Energy*, 2024, 1–11.
- B. Sayahpour, W. Li, S. Bai, B. Lu, B. Han, Y.-T. Chen, G. Deysher, S. Parab, P. Ridley and G. Raghavendran, *Energy Environ. Sci.*, 2024, **17**, 1216–1228.
- G. Sun, C. Lou, B. Yi, W. Jia, Z. Wei, S. Yao, Z. Lu, G. Chen, Z. Shen and M. Tang, *Nat. Commun.*, 2023, **14**, 6501.
- S. Wang, S. Weng, X. Li, Y. Liu, X. Huang, Y. Jie, Y. Pan, H. Zhou, S. Jiao and Q. Li, *Angew. Chem., Int. Ed.*, 2023, **135**, e202313447.
- S. Zhang, Y. Liu, Q. Fan, C. Zhang, T. Zhou, K. Kalantar-Zadeh and Z. Guo, *Energy Environ. Sci.*, 2021, **14**, 4177–4202.
- P. Wen, P. Lu, X. Shi, Y. Yao, H. Shi, H. Liu, Y. Yu and Z. S. Wu, *Adv. Energy Mater.*, 2021, **11**, 2002930.
- J. A. S. Oh, G. Deysher, P. Ridley, Y. T. Chen, D. Cheng, A. Cronk, S. Y. Ham, D. H. Tan, J. Jang and L. H. B. Nguyen, *Adv. Energy Mater.*, 2023, 2300776.
- X. Wang, C. Zhang, M. Sawczyk, J. Sun, Q. Yuan, F. Chen, T. C. Mendes, P. C. Howlett, C. Fu and Y. Wang, *Nat. Mater.*, 2022, **21**, 1057–1065.
- M. He, L. Zhu, G. Ye, Y. An, X. Hong, Y. Ma, Z. Xiao, Y. Jia and Q. Pang, *Angew. Chem., Int. Ed.*, 2024, **136**, e202401051.
- Q. Chen, W. Zhuang, Z. Hou, Y. Jiang, J. Wan, T. Zhang, Y. Zhao and L. Huang, *Adv. Funct. Mater.*, 2023, **33**, 2210206.
- H. Wang, C. Zhu, J. Liu, S. Qi, M. Wu, J. Huang, D. Wu and J. Ma, *Angew. Chem., Int. Ed.*, 2022, **134**, e202208506.
- X. Zhou, Q. Zhang, Z. Zhu, Y. Cai, H. Li and F. Li, *Angew. Chem., Int. Ed.*, 2022, **134**, e202205045.
- K. Doi, Y. Yamada, M. Okoshi, J. Ono, C. P. Chou, H. Nakai and A. Yamada, *Angew. Chem., Int. Ed.*, 2019, **131**, 8108–8112.
- X. Li, W. Ye, P. Xu, H. Huang, J. Fan, R. Yuan, M. S. Zheng, M. S. Wang and Q. Dong, *Adv. Mater.*, 2022, **34**, 2202898.
- P. Wang, G. Zhang, X.-Y. Wei, R. Liu, J.-J. Gu and F.-F. Cao, *J. Am. Chem. Soc.*, 2021, **143**, 3280–3283.

31 K. Lee, Y. J. Lee, M. J. Lee, J. Han, J. Lim, K. Ryu, H. Yoon, B. H. Kim, B. J. Kim and S. W. Lee, *Adv. Mater.*, 2022, **34**, 2109767.

32 F. Xu, C. Qu, Q. Lu, J. Meng, X. Zhang, X. Xu, Y. Qiu, B. Ding, J. Yang and F. Cao, *Sci. Adv.*, 2022, **8**, eabm7489.

33 Y. Jiang, Y. Yang, F. Ling, G. Lu, F. Huang, X. Tao, S. Wu, X. Cheng, F. Liu and D. Li, *Adv. Mater.*, 2022, **34**, 2109439.

34 L. Zhao, Z. Hu, Z. Huang, Y. Tao, W. H. Lai, A. Zhao, Q. Liu, J. Peng, Y. Lei and Y. X. Wang, *Adv. Energy Mater.*, 2022, **12**, 2200990.

35 X. Zhou, F. Liu, Y. Wang, Y. Yao, Y. Shao, X. Rui, F. Wu and Y. Yu, *Adv. Energy Mater.*, 2022, **12**, 2202323.

36 M. Zhu, G. Wang, X. Liu, B. Guo, G. Xu, Z. Huang, M. Wu, H. K. Liu, S. X. Dou and C. Wu, *Angew. Chem., Int. Ed.*, 2020, **132**, 6658–6662.

37 S. Li, H. Zhu, Y. Liu, Q. Wu, S. Cheng and J. Xie, *Adv. Mater.*, 2023, **35**, 2301967.

38 L. Yue, Y. Qi, Y. Niu, S. Bao and M. Xu, *Adv. Energy Mater.*, 2021, **11**, 2102497.

39 J. Wang, Y. Gao, D. Liu, G. Zou, L. Li, C. Fernandez, Q. Zhang and Q. Peng, *Adv. Mater.*, 2024, **36**, 2304942.

40 Z. Yu, J. Song, M. L. Gordin, R. Yi, D. Tang and D. Wang, *Adv. Sci.*, 2015, **2**, 1400020.

41 W. Li, H. Li, Z. Lu, L. Gan, L. Ke, T. Zhai and H. Zhou, *Energy Environ. Sci.*, 2015, **8**, 3629–3636.

42 J. Kim, J. Kim, J. Jeong, J. Park, C.-Y. Park, S. Park, S. G. Lim, K. T. Lee, N.-S. Choi and H. R. Byon, *Energy Environ. Sci.*, 2022, **15**, 4109–4118.

43 Y.-J. Lei, X. Lu, H. Yoshikawa, D. Matsumura, Y. Fan, L. Zhao, J. Li, S. Wang, Q. Gu and H.-K. Liu, *Nat. Commun.*, 2024, **15**, 3325.

44 Y. Xiao, F. Liu, H. Shi, L. Hou, G. Qin, C. Yuan and X. W. Lou, *Adv. Mater.*, 2023, 2301772.

45 G. Liu, Z. Sun, X. Shi, X. Wang, L. Shao, Y. Liang, X. Lu, J. Liu and Z. Guo, *Adv. Mater.*, 2023, **35**, 2305551.

46 X. Cheng, R. Shao, D. Li, H. Yang, Y. Wu, B. Wang, C. Sun, Y. Jiang, Q. Zhang and Y. Yu, *Adv. Funct. Mater.*, 2021, **31**, 2011264.