INTRODUCTION
The steadily growing global energy storage market is driving demand for innovative science and technology solutions that enable high-performance and economical batteries, beyond the capabilities of state-of-the-art lithium-based technology (1–3). Batteries based on multivalent redox chemistries (including Zn/Zn2+, Mg/Mg2+, Al/Al3+, etc.) at the anode have long been heralded as the next generation of energy storage technology solutions because of their ultrahigh theoretical capacity and abundant natural supply of the raw materials (4–6). A large and growing body of work has shown, however, that irrespective of their chemistry, all metal anodes are plagued by the propensity of metals to crystallize at planar electrodes in loose, irregular, and extreme cases of dendritic assemblies, which lead to poor reversibility and highly inefficient utilization of the electrochemically active metal anode (7–9). An additional body of work has shown that randomly oriented electrodeposits with high porosity have strong tendency to become electrically isolated, which also leads to poor reversibility (10–12).

The interphase generated by chemical and electrochemical transformations between any battery electrode and electrolyte is now understood to play a crucial role in regulating ion transport and electrodeposition morphology at a metal anode (13, 14). Well-designed solid electrolyte interphases (SEIs) in lithium batteries are known, for example, to facilitate ion desolvation, to repel free electrolyte molecules at the electrified interface, and to manipulate the local ion transport electrokinetics (15–17). Preformed SEIs anchored on metal anodes have consequently been reported to be highly effective in tuning the morphologies of metal deposits (18–20). Unfortunately, at the high anode capacities encountered during deep cycling of practical batteries, the artificial SEIs typically break, enabling electrolyte components to freely access and be reduced by the anode, consuming electrolyte and markedly lowering the anode reversibility. Building a conductive interfacial layer with low lattice mismatch of metal deposits has been reported recently as an effective strategy to achieve compact metal electrodeposition by regulating the epitaxial electrodeposition with preferred facet orientation (21, 22). This interfacial layer is conductive and thus has to be anchored to a substrate. Its influence on electrodeposited morphology decays with distance from the substrate, and as a consequence, the regulation is limited to moderate to low anode capacities (e.g., <5 mA-hour cm−2) (23–25).

A strategy that would enforcing similar strong constraints on metal crystallization but can be dynamically deployed using freely diffusing agents inside a liquid electrolyte is obviously and potentially far-reaching importance. If the designed interphase can itself assemble and disassemble on time scales that are fast relative to typical metal electrodeposition rates in batteries, then the metal electrodeposits growth can be regulated to arbitrarily high extents. We note that dynamic regulation via interlaminar interphases is a versatile methodology for natural organisms to produce hierarchically structured minerals (26). For example, nacre with laminated microstructures can pregnerate layered biopolymeric interphases, as interlaminar templates, to dynamically bind ion precursors and promote oriented crystallization between the biopolymeric interphases (27). Dynamic assembly process can also be applied to deposit or mineralize diverse kinds of inorganic crystals with hierarchical superstructures through the delicate interaction between building blocks and designed interphases (28, 29).

In order for dynamic interphases formed inside a battery cell to regulate metal deposition at low or high capacities, the interphase must satisfy at least three criteria: (i) it must be nonconductive to facilitate metal electrodeposition without concern of automatic short circuit; (ii) it must be anisotropic and exhibit preferred orientation under the electric fields; and (iii) it must exhibit strong interaction with specific metal facets to template/direct the deposit growth in a preferred orientation. We choose to study Zn metal anode as the model system due to its already well-known merits including the
use of water as the electrolyte solvent, the relatively high volumetric capacity (5855 A·hour liter⁻¹ for zinc), in comparison to other anode technologies (e.g., 790 A·hour liter⁻¹ for graphite and 2066 A·hour liter⁻¹ for lithium), as well as the intrinsic safety and cost efficiency characteristics (30–32). The higher Young’s modulus of zinc (108 GPa) than other metals (Mg, 45 GPa; Al, 69 GPa) at the liquid/solid interface also means that batteries based on Zn have much lower tolerance for nonuniform/noncompact deposition of the metal, as the strong Zn deposits will easily break through any barriers to short-circuit the battery cell (33), in addition to driving chemical/electrochemical side reaction toward aqueous electrolytes (34, 35).

We report that dense and vertically aligned Zn electrodeposition can be achieved via dynamic adsorption of artificial interphases formed in a simple colloidal electrolyte containing graphitic C₃N₄ (g-C₃N₄) nanosheets (fig. S1). The g-C₃N₄ nanosheets are of specific interest because of their high crystallographic match for hexagonal close-packed (HCP) Zn and, as will be shown later, can be dynamically adsorbed/desorbed on (0002) Zn facet during the depositing/stripping process. In situ quantitative experiments reveal that the ordered Zn deposits result in a notably decreased consumption of Zn and markedly lower accumulation of by-products during each cycle. These merits result in unprecedented Zn reversibility (~99.8%) and high cumulative capacity (2520 mA·hour cm⁻²) suitable for practical aqueous Zn metal batteries (AZMB) with areal capacities as high as 20 mA·hour cm⁻².

RESULTS AND DISCUSSION

Aligned Zn electrodeposits under dynamic interphase mediation

An improved anode reversibility is associated with the dense and oriented assembly of Zn electrodeposits. It is known that HCP crystallography of Zn plays a central role in Zn electrodeposition growth by facilitating the formation of hexagonal Zn platelets exposing (0002)Zn facet with the lowest surface energy (fig. S2) (36). Focused ion beam scanning electron microscopy (FIB-SEM) analysis was carried out to provide insights into the inner microstructural details of Zn growth in aqueous electrolyte (Fig. 1A). Hexagonal Zn platelets with various sizes (1 to 5 μm) are observed to form loosely connected building blocks assembled to create a porous electrodeposition structure in the control materials. The Zn deposits are revealed to be randomly oriented using the reconstructed stack of cross-sectional slices under slice and view modes (Fig. 1B and fig. S3). We hypothesize that the increased surface area associated with these structures exposed in aqueous electrolyte media caused by the irregular interstices inside the bulk deposits is the main cause of uncontrolled interfacial side reactions and poor Zn anode reversibility. In contrast, dense Zn electrodeposits without inner interstices formed in electrolytes containing the g-C₃N₄ (fig. S4) are observed. Hexagonal Zn platelets with uniform size are seen to be vertically stacked together (~100% compactness) and induce a sharp decrease in contact area with aqueous electrolytes (Fig. 1, C and D).

The growth of vertically aligned Zn electrodeposits is triggered by the orientated nanosheets with facet-dependent adsorption ability under the electric field. As shown in Fig. 2A, the critical steps to facilitate the torsion and alignment of nanosheets in aqueous electrolyte are as follows: (i) Electron-donating functional groups on the g-C₃N₄ nanosheets can induce an ordered electric double layer (EDL) in the colloidal electrolyte. (ii) Under the electric field, excess counterions inside the EDL migrate and lead an electro-osmotic flow along the nanosheet surface, inducing an EDL polarization (37). Because of the anisotropic structure of nanosheets, excess polarization parallel to nanosheet surface can induce a torque to align the

![Fig. 1. Densely packed Zn electrodeposits featuring vertically aligned hexagonal platelets.](https://example.com/f1.png)

Cross-sectional images produced by FIB and three-dimensional (3D) morphology reconstructions of Zn deposits in (A and B) control 2 M ZnSO₄ (aq) liquid electrolytes. Randomly oriented Zn electrodeposits assemble to form porous and loose microstructures; and (C and D) aqueous 2 M ZnSO₄ colloidal electrolytes prepared through g-C₃N₄ addition. A fixed concentration of g-C₃N₄ of 1.0 mg ml⁻¹ was used in these studies. Vertically aligned and dense electrodeposits are observed in the colloidal electrolytes under the dynamic interphase mediation. The current density was 4 mA cm⁻².
nanosheets with the electric field (text S1) (38). (iii) The vertically aligned nanosheets at the liquid-solid interface can further influence the orientation of Zn electrodeposits by preferential adsorption on (0002)Zn facet.

The electro-orientation of nanosheets in the electrolyte upon applying the electric field can be verified by small-angle x-ray scattering (SAXS) measurements (Fig. 2, B and C). The isotropic scattering pattern and nearly flat azimuthal angle plot of the colloidal dispersion without electric field suggest the random distribution of nanosheets. Stronger equatorial scattering patterns in SAXS images and peak intensity in azimuthal angle plots reflect improved alignment of nanosheets with the increase in electric field intensity. Dynamic adsorption of the artificial interphases on specific exposed facet can further promote the growth of aligned Zn electrodeposits. Free energy diagram is plotted to reveal the ion coordination process and preferential adsorption of g-C₃N₄ nanosheets (Fig. 2D). A unit of g-C₃N₄ can spontaneously adsorb at least four zinc ions and form an ionized nanosheet with high structural symmetry in the aqueous electrolyte. Moreover, density functional theory (DFT) calculations reveal that ionized nanosheets exhibit stronger binding affinity with (0002)Zn facet (−16.05 eV) compared to (1010)Zn or (1011)Zn, and thus facet-dependent adsorption encourages the vertical growth of oriented Zn electrodeposits by blocking (0002)Zn facets (fig. S5). To further demonstrate the synergistic effect of electro-oriented interphases and facet-dependent adsorption, we tested Zn electrodeposition using fixed g-C₃N₄ nanosheets parallel to the separator as a comparison (fig. S6). Distinctly different morphologies featuring horizontally aligned Zn electrodeposits can be observed at 1 mA·hour cm⁻² and become random-oriented at 3 mA·hour cm⁻² because of the lack of dynamic facet-dependent regulation.

The vertically aligned Zn electrodeposits exhibit a (1011) texture, benchmarking against the random orientation in bare electrolyte (Fig. 2E). The nucleation and growth patterns of Zn electrodeposits under the dynamic interphase mediation were further investigated (Fig. 2, F to H, and fig. S7). Hexagonal Zn platelets cofacially stack and form the uniform distributed nucleus at the nucleation stage, and then ordered crystal coarsening and lateral spreading mediated by electro-oriented interphases can be achieved during the growth stage. Under deep depositing condition with a high capacity of 6 mA·hour cm⁻², the thickness of the (1011) textured Zn electrodeposits is 13.79 μm, in sharp contrast to the loose and nonuniform electrodeposits in bare electrolyte (22.25 μm in thickness) (fig. S8). In addition to aqueous electrolytes containing ZnSO₄, dynamic interphase-mediated Zn electrodeposition can be readily applied to other aqueous electrolytes of interest, such as ZnCl₂, Zn(OAc)₂, and Zn(CF₃SO₃)₂ (fig. S9).

Dynamic assembly/disassembly of Zn electrodeposits
To clarify the nanoscale spatial structure and chemical information of dynamic interphase over the vertically aligned Zn electrodeposits,
we performed atomic force microscopy–based infrared (AFM-IR) spectroscopy. The localized IR absorption spectra at different sites over the sample can be measured by AFM cantilevers (Fig. 3A), based on the instantaneous thermal expansion stems from local absorption of IR radiation (39). The characteristic absorption peak at 1637 cm$^{-1}$ can be assigned to the $\text{C}=$-$\text{N}$ stretching vibrations, which exhibits an excellent agreement with the spectra of g-$\text{C}_3\text{N}_4$ powders measured by conventional Fourier transform IR spectrometer (fig. S10). IR signals with higher intensities at sites 1 and 3 suggested that g-$\text{C}_3\text{N}_4$ nanosheets may directly interact with side walls of the Zn electrodeposits. AFM topography image (Fig. 3B) and IR absorption mapping image at 1637 cm$^{-1}$ (Fig. 3C and fig. S11) further reveal the linear distribution of nanosheets correlated with the aligned Zn platelets. A cross-sectional lamellae of aligned Zn deposits were prepared by FIB lift-out method, and a clear crystalline–amorphous crystalline heterogeneous structure can be observed (figs. S12 and S13). The aligned Zn platelets are strongly bounded by intercalated g-$\text{C}_3\text{N}_4$ nanosheets (2 to 4 nm in thickness) and stacked to form a dense laminated microstructure.

In situ electrochemical quartz crystal microbalance (EQCM) and cyclic voltammetry (CV) tests were combined to provide a quantitative understanding on dynamic interphase–mediated depositing/stripping behavior (fig. S14). EQCM can precisely monitor the mass changes on the quartz crystal working electrode with a function of potentials applied by CV (40). Coulombic efficiency (CE) and mass efficiency (ME), defined as the electron transfer and mass change ratio between Zn stripping and Zn depositing during each cycle, respectively, are introduced to quantitatively monitor the irreversible consumption of active Zn and by-product accumulation. As shown in Fig. 3 (D and E), severe mass accumulation originated from electrochemically inactive Zn (average CE, 90.8%) and by-products (average ME, 48.3%) after stripping repeatedly occurred in bare ZnSO$_4$ electrolyte. However, highly reversible depositing/stripping process with an average CE of 99.6% can be observed under the mediation of dynamic interphase. Side reactions associated with the lack of interfacial protection can be resolved (average ME, 98.5%) because of the existence of the dynamic interphase adsorbed on Zn platelets. Moreover, the weight of intercalated g-$\text{C}_3\text{N}_4$ nanosheets in the aligned Zn deposits is approximately 3.28 weight %, which can be quantified on the basis of the value difference between actual mass increase in quartz crystal and the theoretical mass change according to electron transfer numbers in CV (Fig. 3F).

**Performance of practical metal batteries**

Practical energy-dense AZMB require an extremely high CE to minimize the active Zn loss during each cycle. Long-term CE tests were first carried out in electrochemical cells to assess the utility function of dynamic interphase mediation. At a moderate areal capacity of 1 mA-hour cm$^{-2}$, the average CE reaches 99.6% over 1600 cycles under the dynamic interphase mediation, while the CE drops below 90% after 160 cycles for bare electrolyte. As shown in Fig. 4B, Zn cyclability was further evaluated under deep cycling...
condition (e.g., 6 mA·hour cm\(^{-2}\)), which is equivalent to the ideal areal capacity in practical full batteries. Significantly improved CE (averaging 99.8% over 400 cycles) demonstrates the high-efficiency Zn utilization during the vertically aligned depositing/stripping process. By contrast, the cell without modifications short-circuited because of irregular Zn deposits after just 36 cycles. Under the mediation of dynamic interphase, well-maintained Zn reversibility and stable voltage hysteresis can be achieved at various current densities from 1 to 10 mA cm\(^{-2}\) (figs. S15 and S16). It is also noteworthy that the CE (average, 99.5%; Fig. 4C and fig. S17) was not compromised under ultrahigh current density (40 mA cm\(^{-2}\)) and ultrahigh areal capacity condition (20 mA·hour cm\(^{-2}\)).

The superiority of dynamic interphase–mediated high-efficiency Zn depositing/stripping can be further demonstrated in practical full batteries. We note that most of the AZMB reported in literature were tested using huge excess Zn metal anodes (>100 \(\mu\)m in thickness), low mass loading cathodes (<1 mA·hour cm\(^{-2}\)), and flooded electrolytes (>100 \(\mu\)l mA·hour\(^{-1}\)), which will markedly reduce the actual energy density and conceal the anode challenges in full batteries. From application perspectives, the use of thin Zn anode (<30 \(\mu\)m) under deep cycling conditions (>4 mA·hour cm\(^{-2}\)) is the necessary parameters for practical AZMB designs (24, 41). Here, we design the Zn/Zn\(_x\)V\(_2\)O\(_5\)/H\(_2\)O (ZnVO) full batteries tested under highly challenging conditions: ultrathin zinc anode (20 \(\mu\)m), high mass loading ZnVO cathode (16 mg cm\(^{-2}\); fig. S18), and lean electrolyte (10 \(\mu\)l mA·hour\(^{-1}\)). ZnVO nanobelts with high specific capacity and reversible Zn\(^{2+}\) intercalation/deintercalation ability (fig. S19) were chosen as a candidate for cathode materials (42). The rational designed negative-to-positive capacity (N/P) ratio of 1.85 and deep cycling condition enable a high energy density of 280 W·hour liter\(^{-1}\) (table. S1; based on anode, cathode, and separator). As shown in Fig. 4C, Zn/ZnVO full batteries using bare electrolyte can only run 29 cycles before a sudden internal short circuit, while approximately 13 times longer cycling life span (400 cycles) and continuous operation time (1100 hours) can be achieved via the application of dynamic interphase. Postcycling characterizations of dynamic interphase–mediated Zn anodes reveal that excellent cycling stability of full batteries can be attributed to the high-efficiency and ordered stripping pattern over the vertically aligned electrodeposits (fig. S20). The performance of Zn/ZnVO full batteries under the dynamic interphase mediation is among the best to date for practical AZMB (table S2). Compared with previous literature, our work stands out for the highest cumulative capacity (up to 2520 mA·hour cm\(^{-2}\)), highest per-cycle areal capacity (6.3 mA·hour cm\(^{-2}\)) and lowest N/P ratio (1.85), providing a promising route for practical AZMB (Fig. 4E).

Dynamic interphase mediation is a versatile strategy for high-capacity metal anodes. Similar to dynamic function of g-C\(_3\)N\(_4\) nanosheets in the aforementioned electrochemical cells, boron nitride (BN) nanosheets with high aspect ratio and six-membered ring structure can also work as dynamic interphase to regulate the Zn electrodeposition. EQCM revealed that significantly improved plating/stripping reversibility can also be achieved under the mediation of BN nanosheets (fig. S21). Moreover, this strategy can be readily extended to other energy-dense metal batteries systems. Metallic Mg with the same HCP crystal structure as Zn exhibited a vertically aligned morphology with (10\(_{11}\)) texture, and an improved CE of 98.6% under the dynamic interphase mediation, in contrast to the nonuniform, random assembled Mg electrodeposits (average CE of 79.9% for control cells), formed in bare electrolytes (fig. S22). We further evaluated the utility of our concept in multivalent Al anodes. As shown in fig. S23, the dynamic interphase provides a large driving force between Al electrodeposit grains, enabling a densely
packed morphology and exceptional reversibility (average CE, 98.74%) at high areal capacity (3 mA-hour cm⁻²).

In summary, we have developed a dynamic interphase strategy to promote ordered assembly of metal electrodeposits under deep cycling condition. Using Zn anode as an example, exceptionally reversible Zn depositing/stripping (>99.8%) under deep cycling conditions (6 to 20 mA-hour cm⁻²) can be achieved by the dynamic mediation of artificial interphases. Artificial interphases adsorbed on selective facet can promote cofacially stacked Zn electrodeposits and hinder the electrolyte-related interfacial side reactions. The dynamic interphase concept can be extended to achieve densely assembled Mg and Al electrodeposits in multivalent metal batteries, providing a brand-new route for the development of deep cycling metal batteries.

**MATERIALS AND METHODS**

**Preparation of electrolytes**

**Zn electrolyte**

ZnSO₄·7H₂O was dissolved in the deionized water to prepare the 2 M ZnSO₄ electrolyte for Zn electrochemical cells.

**Mg electrolyte**

AlCl₃ powder was slowly added into phenyl magnesium chloride Mg electrolyte 600°C for 4 hours under air. The g-C₃N₄ nanosheet or BN nanosheet H₂O suspension with a concentration of ~1.0 mg ml⁻¹. The suspension is centrifuged at 4000 rpm for 10 min and gives nanosheets/suspension of 5°C min⁻¹. The as-prepared powder was further heated at 900 to 1900 cm⁻¹ with a resolution of 5 cm⁻¹. The AFM-IR spectra absorption rates of the AFM cantilever. The resonant amplitudes of the AFM cantilever are positively correlated with the thermal induced resonance of the AFM cantilever. The resonant frequency change, 2 f_q, is the frequency change, f₀ is the resonant frequency

**Preparation of nanosheet suspension**

Bulk g-C₃N₄ powder was synthesized by thermal polymerization of melamine. Briefly, 10 g of melamine was placed in a covered crucible and then heated to 550°C for 2 hours in Ar atmosphere with a heating rate of 5°C min⁻¹. The as-prepared powder was further heated at 600°C for 4 hours under air. The g-C₃N₄ nanosheet or BN nanosheet suspensions were fabricated by liquid phase exfoliation of the above bulk powders. Typically, the bulk powder was dispersed in H₂O with a concentration of 5 mg ml⁻¹. The dispersion was sonicated for 6 hours in an ultrasonic cell crusher (700 W), after which the suspension is centrifuged at 4000 rpm for 10 min and gives nanosheets/H₂O suspension with a concentration of ~1.0 mg ml⁻¹.

**Preparation of electro-oriented hydrogels**

First, 3 g of acrylamide, 15 mg of N,N'-methylenebis(acrylamide), 15 mg of ammonium persulfate, and 30 μl of N,N,N',N'-tetramethylethylenediamine were sequentially added into 15 ml of 2 M ZnSO₄ electrolyte (with ~1.0 mg ml⁻¹ of g-C₃N₄ nanosheets). The precursor solution was quickly injected into a glass mold (between two electrodes), and then the cell was charged at 80°C for 30 min. The distance between the electrodes is 1 mm. The obtained hydrogels were rinsed in deionized water and can be used in SAXS tests.

**Orientation-confined tests**

The g-C₃N₄ nanosheet–coated separator was prepared by a vacuum-assisted infiltration method. g-C₃N₄ nanosheet suspension (~1.0 mg ml⁻¹; with 5% Nafion solution as a binder) was filtrated on a cellulose separator (Shanghai Xinya Purification Equipment Co. Ltd.). The g-C₃N₄ nanosheet–coated separators were dried in a vacuum oven at 60°C for 12 hours. The lamellar membrane with fixed g-C₃N₄ nanosheets were used as separator in the orientation-confined tests.

**Material characterization**

Field-emission SEM measurements were conducted on Hitachi SU8000. X-ray diffraction (XRD) patterns were obtained using a PANalytical X’Pert Powder with Cu-Kα radiation. SAXS measurements were performed on a Jeol SAXS system (Xenocs SA). Transmission electron microscopy (TEM) lamella sample was prepared by a dual-beam FIB-SEM (FEI Quanta 3D FEG) with a nanomanipulator, using a standard lift-out procedure. The initial lamella was milled with a Ga ion beam (30 kV and 1 nA) and then mounted on a TEM grid. Thinning was first conducted with a beam current of 500 pA and then final polishing with a beam current of tens of picoamperes to remove amorphization. TEM measurements were performed on JEOL 2100F with a Gatan K2 direct detection camera to identify the crystallography of electrodeposits.

**FIB and three-dimensional morphology reconstructions**

The cross sections and bulk structures of electrochemically deposited Zn were analyzed by dual-beam SEM (Thermo Fisher Scientific, FIB Helios G3 UC). To maintain the structural integrity of Zn deposits in bare 2 M ZnSO₄ during FIB milling, platinum was sputtered on the samples using the gas injection system. A Ga ion beam (30 kV and 0.79 nA) was used to serially section and polish the sample. The data were collected until up to 400 cross-sectional images were captured to provide a full size of Zn deposits. Alignment corrections of the image stack and digital reconstruction three-dimensional (3D) morphology were visualized by IMOD (Boulder Laboratory) and Amira 6.5 (Thermo Fisher Scientific).

**AFM-IR spectroscopy measurements**

The localized nanoscale mid-IR spectra and absorption maps were carried out using a NanoIR2-Fs instrument (Anasys Instruments Inc.) under contact mode. These measurements are based on the photothermal induced resonance of the AFM cantilever. The resonant amplitudes of the AFM cantilever are positively correlated with the absorbed IR radiation of samples. IR spectra were acquired over the 900 to 1900 cm⁻¹ with a resolution of 5 cm⁻¹. The AFM-IR spectra and 3D intensity maps were generated using Analysis Studio software for the NanoIR2 system.

**Electrochemical quartz crystal microbalance**

The zinc deposition/stripping happened in an EQCM cell (QCM922, Princeton Applied Research, USA). The EQCM experiments carried out at a scan rate of 10 mV s⁻¹ a specially designed electrolytic cell using a standard three-electrode configuration. A gold-coated quartz crystal electrode, a platinum electrode, and an Ag/AgCl electrode were used respectively as the working electrode, counter electrode, and reference electrode.

The mass change (Δm) of Zn deposits on the quartz surface should follow the Sauertrey equation

\[ Δf = -2f_0^2 (μ_qρ_q)^{-1} Δm/A \]

Here, Δf is the frequency change, f₀ is the resonant frequency of the quartz crystal (9 MHz), μ_q is the shear modulus of quartz
(2.947 × 10^{-11} \text{ g cm}^{-2} \text{ s}^{-2}), \rho_q$ is the density of quartz (2.648 g cm^{-3}), and $A$ is the piezoelectrically active crystal area (0.196 cm^{-2}).

**Preparation of high mass loading Zn_{0.25}V_2O_5 cathode**

Zn_{0.25}V_2O_3 nanobelts were synthesized via a hydrothermal method. Typically, 0.54 g of V_2O_5 and 0.43 g of Zn(OAc)_2 were dissolved in a mixed solvent (70 ml of deionized water, 5 ml of acetonitrile, and 2 ml of 10% HNO_3). The autoclave was sealed and heated at 180°C for 24 hours. The as-synthesized product was filtrated and washed by deionized water and isopropanol and dried at 80°C for 24 hours. High mass loading Zn_{0.25}V_2O_5 cathodes were fabricated by a vacuum infiltration method. The aqueous mixture of CNT suspension (Cannd New Energy Technology Co. Ltd.) and Zn_{0.25}V_2O_3 powder (2:8 by weight) was sonicated for 10 hours and then vacuum-infiltrated. The freestanding cathodes were dried at 80°C for 12 hours.

**Electrochemical tests**

All Zn metal aqueous batteries (symmetric Zn batteries, Zn/Ti batteries, and Zn/ZnVO full batteries) were assembled in atmospheric environment using pouch-type battery model. Glass fiber membrane (GF/A, Whatman) or cellulose membrane (Shanghai Xinya Purification Equipment Co. Ltd.) was used as separator. LAND battery cycler and Solartron Analytical Electrochemical Workstation were used for the electrochemical measurements. Zn CE was measured in Zn/Ti cells. A fixed amount Zn (6 mA-hour cm^{-2}) was plated on Ti electrode and then stripped back under different current densities until the cutoff voltage increased to 1 V. Ultrathin Zn foils (20 μm; Runde Metal Co. Ltd.) were used in symmetric Zn/Zn and Zn/ZnVO tests. Long-term cycle life-span tests of Zn/ZnVO full cells were cycled at 4 mA cm^{-2} within a voltage range of 0.2 to 1.6 V versus Zn^{2+}/Zn.

**Computational methodology**

All DFT calculations were performed at the GFN1-xTB level using the xTB program, version 6.3.2. The geometry optimizations were calculated with periodic boundary conditions, while the frequency was calculated without periodic boundary conditions.

For electronic self-consistent calculations, geometry optimizations, and frequency calculations, the convergence tolerance is set as follows: integral cutoff = 0.25 × 10^{-7}, Broyden damping = 0.40, and the accuracy = 1.0.

The free energy change for different steps was determined as follows

\[
\Delta G_1 = E_{\text{bind}} - E_{\text{C3N4}} - 4 \times E_{\text{ZnSO4}} + \Delta E_{\text{ZPE}} + \Delta H_{0\rightarrow 300K} - T\Delta S
\]

\[
\Delta G_2 = E_{\text{ads}} - E_{\text{slab}} - E_{\text{bind}} + \Delta E_{\text{ZPE}} + \Delta H_{0\rightarrow 300K} - T\Delta S
\]

Here, $E_{\text{bind}}$, $E_{\text{C3N4}}$, $E_{\text{ZnSO4}}$, $E_{\text{ads}}$, and $E_{\text{slab}}$ are the electronic energy of g-C_3N_4 and ZnSO_4 binding state, g-C_3N_4 single state, ZnSO_4 single state, adsorption state, and Zn slab. $\Delta E_{\text{ZPE}}$, $\Delta H_{0\rightarrow 300K}$, and $\Delta S$ are zero-point energy change, thermal correction change from 0 to 300 K, and entropy change, which were also obtained through the xTB program, version 6.3.2.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at https://science.org/doi/10.1126/sciadv.abl3752
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