Biomolecule-guided cation regulation for dendrite-free metal anodes

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Lithium (Li) or zinc (Zn) metal anodes have attracted interest for battery research due to their high theoretical capacities and low redox potentials. However, uncontrollable dendrite growth, especially under high current (>4 mA cm⁻²), precludes reversible cycling in Li or Zn metal batteries with a high-loading (>4 mA cm⁻²), precludes reversible cycling in Li or Zn metal batteries with high-loading (>4 mA cm⁻²) cathode. We report a cation regulation mechanism to address this failure. Collagen hydrolysate coated on absorbed glass mat (CH@AGM) can simultaneously induce a deionization shock inside the separator and spread cations on the anode to promote uniform electrodeposition. Employing 24 mAh cm⁻² cathodes, Li and Zn metal batteries with CH@AGM delivered 600 cycles with a Coulombic efficiency of 99.7%. In comparison, pristine Li and Zn metal batteries only survive for 10 and 100 cycles, respectively. This approach enabled 400 cycles in a 200 Ah-class Zn metal battery, which suggests a scalable method to achieve dendrite-free anodes in various batteries.

INTRODUCTION

The increasing demand for battery systems for portable electronics, electric vehicles, and other electric applications has motivated intensive research seeking high energy density, high unit capacity, and decently stable energy storage devices (1). Because of high theoretical capacity with low oxidation-reduction potential, lithium (Li) and zinc (Zn) metals are considered the most promising anode materials for rechargeable metal-ion batteries (2). However, the main obstacle for the practical application of Li and Zn metal batteries is the dendrite formation (3). Dendrites not only penetrate the separator to induce short circuit in the battery (4) but also generate high surface area on the anode to accelerate metal ion depletion, resulting in active material loss and subsequent battery failure (5). Great efforts have been made to stabilize Li and Zn anodes in lithium ion batteries in the last 10 years. Conductive hosts, such as graphene (6), carbon nanotubes (7), porous copper (8), and graphite fibers (9), have been developed to control the deposition of Li and Zn ions to achieve a uniform and stable metal anode. The large surface area of these materials decreases the local current density, which effectively improves battery safety and extends Sand’s time. Moreover, the ion distribution in the electrolyte can be regulated by surface functional groups and pore structures of these conductive materials. Simultaneously, electrolyte additives (10), porous separators (11), and artificial protective films (12) are used to manipulate ion diffusion to achieve a homogeneous Li and Zn deposition.

Although good cycling life and excellent specific capacity have been achieved in Li and Zn metal batteries (13, 14), these results were still lacking actual practicalities for a reliable battery system. For fundamental study, the batteries are normally tested with an extremely low mass loading of cathode material to minimize side reactions in the cathode and avoid metal ion consumption and electrolyte depletion. However, the low cathode loading (1 mAh cm⁻² or less) used in most previous studies leads to very “shallow” stripping/deposition of metal in the same cell, which artificially leads to long cycle life that will not translate to high-energy density cell designs (15). For example, Li||nickel-cobalt-manganese batteries assembled under laboratory-scale, a low cathode mass setup delivers a low specific cathode capacity of 1.5 mAh cm⁻², which leads to a unit capacity of only 1.4 mAh (per 0.95 cm² cathode area). By considering the total weight of the cell unit, the energy density of such a battery would be less than 15 Wh kg⁻¹ (15), which is far smaller than the energy density requirements for commercial batteries. It is well known that the cathode areal capacity should be above 4.0 mAh cm⁻² to achieve the desired unit capacity and specific energy density (16). However, under such conditions, the high current density during charge-discharge due to the high mass loading of cathode materials always leads to fast heterogeneous nucleation and deposition of Li or Zn ions and, hence, dendritic metal growth, which results in an extremely short battery life span (17). Fundamental challenges associated with accelerated dendrite formation, fast metal depletion, and rapid electrode passivation under a high-loading intercalation cathode have so far presented formidable barriers to progress in achieving large-scale rechargeable batteries based on the Li and Zn metal anodes (16, 18).

Several models have been proposed to assess the stability of the electrodeposition in batteries using metal anodes, which provide feasible strategies to suppress the dendrites (19–21). Chazalviel (22) suggests that the planarity of the metal surface is destabilized at high current densities by a large electric field in the immediate vicinity of the metal electrode, caused by electrode anion depletion. Therefore, any protrusion of the electrode surface bears enhanced electrodeposition, leading to the growth of dendrite tips. On the basis of this model, Tikekar et al. (23) showed a weak effect of background charge theoretically on the linear stability of electrodeposition in steady state and proposed that a structured electrolyte with immobilized anions can stabilize the electrodeposition at large overpotentials. The group of M. Z. Bazant also investigated the influence of charged porous media in copper electrodeposition from copper sulfate solution. The negative surface charge in anodized aluminum oxide (AAO) membrane enables overlimiting current faster than diffusion. The surface conduction enhances the growth of copper metal along the nanopore surfaces, leading to an array of nanotubes. However, with

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a positive surface charge in AAO, the opposing surface conduction blocks dendrite penetration upon ion depletion. Dendrites are channeled along the pore centers, avoiding the electric double layers (24). In addition, it is also found that the positive separator (cellulose nitride or polyethylene) shows reduced flux of cation that is opposed by surface conduction, while the negative separator shows over-limiting current retained by surface conduction that leads to a dendrite-free metal deposition under high current density (25). The same effect of surface transport in charged porous media resulting in “deionization shockwaves” and stable shock electrodeposition have also been exploited for water desalination by “shock electrodialysis” (26, 27).

Inspired from the above fundamental studies, here we propose that collagen hydrolysate (CH) homogeneously coated on absorbed glass mat (CH@AGM) between the anode and the AGM separator can induce a similar shock electrodeposition as reported by Bazant’s group, which fundamentally stabilizes metal anodes and suppresses dendrite formation in lithium and zinc metal batteries with a high areal capacity cathode. Specifically, we found that CH shows a negative surface charge in both aqueous and organic electrolytes, which enables a surface conduction in CH@AGM and a deionization shock in random, charged porous media under overlimiting current, which promotes uniform Li⁺ and Zn²⁺ deposition. We also found that the binding of CH with Li⁺ or Zn²⁺ cations further spreads the distribution of cations over the whole surface of the metal anode, which navigates metal deposition along the surface/horizontal direction of the anode and significantly stabilizes Li and Zn growth. Such superimposed regulation effects in metal electrodeposition produce a dendrite-free metal deposition under high current density and enable the construction of stably cycled lithium and zinc metal batteries, where a Li or Zn metal anode is coupled with a high-loading cathode [LiMn₂O₄ (LMO)]. Compared with pristine batteries with a 24 mAh cm⁻² cathode that can only survive for 10 and 100 cycles: eabb1342 (28), CH@AGM exhibited a newly appeared peak located at 1650 cm⁻¹, which corresponds to the peptide bond from CH (Fig. 1E). The porosity of CH@AGM is investigated by measuring the air permeance using a Gurley densometer as well as by porosimetry. As shown in Fig. 1F, the air permeance and pore size of CH@AGM are superior to the Celgard separator and similar to those of the AGM separator. The electrolyte uptaking of CH@AGM is also investigated in electrolyte wetting test using a 1 M ZnSO₄ and 2 M Li₂SO₄ aqueous electrolyte. As shown in fig. S2, CH@AGM exhibits superior electrolyte-uptaking performances in both vertical and horizontal electrolyte-wetting setups in comparison to the AGM separator. Given that the CH@AGM shows a similar porosity to the AGM separator (Fig. 1F), this result indicates an intrinsic physicochemical affinity of CH to aqueous solutions containing Li and Zn ions, which may be due to the surface charge of CH@AGM. To evaluate the mechanical property of CH@AGM, we performed puncture tests on CH@AGM with different thicknesses. As shown in fig. S3, the force required to penetrate the sample films depends on the film thickness. Increasing the film thickness resulted in a significant improvement in puncture resistance. By comprehensively considering the separator cost, electrical resistance, and the total weight of the final battery, we intentionally use a 1.3-mm-thick CH@AGM in both laboratory-scale and commercialized batteries in this work, which is strong enough for manufacturing operations.

**RESULTS**

**Structure and morphology of CH and CH@AGM**

CH originated from animal tissue was first used as glue as far back as 6000 BC. It is now widely applied in the photographic and food industries. CH consists of three α chains that form a triple-helix structure. The α chain is composed of repetitions of glycine-proline-hydroxyproline amino acid sequences, which are stabilized by hydrogen bonds between every three amino acids. These strong interactions are situated regularly on the chain and retain a triple-helical structure (Fig. 1A) (28). Typical CH consists of 19 amino acids, predominantly glycine, proline, and hydroxyproline, which together represent around 50% of the total amino acid content (Fig. 1B). Raman spectroscopy offers efficient characterization to investigate heterogeneous materials as it provides submicrometer spatial resolution with high sensitivity. As shown in Fig. 1C, the amide I band at 1654 cm⁻¹ indicates the α helix structure (29). The amide I absorption is due to the stretching vibration of the C=O bond, which corresponds to the conformation of protein. Other characteristic peaks around 1440 and 1250 cm⁻¹ are the amide III with C–N stretching and N–H bending vibrations of the peptide bond (30).

The excellent solubility of CH in hot water leads to a facile one-step fabrication of CH@AGM, using porous AGM networks as supporting skeleton (see Materials and Methods). As characterized by scanning electron microscopy (SEM) (Fig. 1D) and corresponding energy-dispersive x-ray mapping (fig. S1), CH results in homogeneous deposition of thin adherent membranes on the immersed AGM. The obtained CH@AGM was further characterized by Fourier transform infrared spectra. While the pristine AGM exhibited only Si—O—Si and Si—O peaks corresponding to typical glass fibers, CH@AGM exhibited a newly appeared peak located at 1650 cm⁻¹, which corresponds to the peptide bond from CH (Fig. 1E). The porosity of CH@AGM is investigated by measuring the air permeance using a Gurley densometer as well as by porosimetry. As shown in Fig. 1F, the air permeance and pore size of CH@AGM are superior to the Celgard separator and similar to those of the AGM separator. The electrolyte uptaking of CH@AGM is also investigated in electrolyte wetting test using a 1 M ZnSO₄ and 2 M Li₂SO₄ aqueous electrolyte. As shown in fig. S2, CH@AGM exhibits superior electrolyte-uptaking performances in both vertical and horizontal electrolyte-wetting setups in comparison to the AGM separator. Given that the CH@AGM shows a similar porosity to the AGM separator (Fig. 1F), this result indicates an intrinsic physicochemical affinity of CH to aqueous solutions containing Li and Zn ions, which may be due to the surface charge of CH@AGM. To evaluate the mechanical property of CH@AGM, we performed puncture tests on CH@AGM with different thicknesses. As shown in fig. S3, the force required to penetrate the sample films depends on the film thickness. Increasing the film thickness resulted in a significant improvement in puncture resistance. By comprehensively considering the separator cost, electrical resistance, and the total weight of the final battery, we intentionally use a 1.3-mm-thick CH@AGM in both laboratory-scale and commercialized batteries in this work, which is strong enough for manufacturing operations.

**Suppression of Zn and Li dendrite by shock electrodeposition in charged porous media**

Zeta potential experiments were performed to study the surface charge of CH@AGM and the pristine AGM. At a pH range of 2 to 6, which corresponds to the acidic electrolyte used in Zn and Li metal batteries, CH@AGM exhibited a negative surface charge derived from the oxygen functional groups in CH backbones (Fig. 2A) (31), while pristine AGM shows a positive surface charge (Fig. 2B) that may be attributed to the protective sizing used in fiber industries. On the basis of these surface charge results, it can be concluded that coating of CH on the AGM surface is realized via electrostatic self-assemblying of the negatively charged CH and the positively charged AGM. Voltammetry measurements were further performed in symmetric Zn and Li cells with and without CH in the AGM separator, to fully understand the influence of background charge in metal electrodeposition. Figure 2C shows current-voltage curves of Zn|AGM + CH@AGM|Zn and Zn|AGM|Zn cells in the 1 M ZnSO₄ electrolyte at a scan rate of 1 mV s⁻¹. Although the current in Zn|AGM|Zn reaches ~46 mA at ~0.2 V and retains a current of ~43 mA, Zn|AGM + CH@AGM|Zn exhibits a linear current increase...
starting from $-0.34\ V$. Similar current-voltage profiles are also observed in the voltammetry of Li|AGM + CH@AGM|Li and Li|AGM|Li cells in 1 M LiPF$_6$ in 1:1:1 dimethyl carbonate: ethyl carbamate: diethyl carbonate (Fig. 2D). These data clearly show that the sign of surface charge in porous media is responsible for either enhancement or suppression of overlimiting current between metal electrodes, which may control the metal electrodeposition behavior under high current (25). As determined from the plateau of voltammetry in Zn|AGM|Zn and Li|AGM|Li symmetric cells, the limiting current density of Zn and Li electrodeposition in this work is 11.3 and 2.06 mA cm$^{-2}$, which is consistent to the reported value in similar battery systems (32, 33).

Figure 2 (E and F) shows mechanistic sketches of the effect of surface conduction on Zn or Li deposition in a charged random porous medium. It is well known that during metal electrodeposition, there is an ion depletion region as cations are continually reduced and deposited on the electrode surface (24, 25, 34). Owing to the reduced bulk conductivity, the axial electric field is amplified to sustain the current (34). With pristine AGM corresponding to a positive surface charge (Fig. 2E), the oppositely directed surface conduction greatly suppresses the overlimiting current and leads to a reduced cation flux (25). Meanwhile, the SO$_4^{2-}$ or PF$_6^-$ counterions move away from the anode, further blocking the diffusion of Zn$^{2+}$ or Li$^+$ ions to maintain neutrality. Above a critical voltage, Zn or Li metal tends to grow avoiding the pore walls and forms large and irregular dendrites owing to the electro-osmotic flow returning to the pore centers (Fig. 2G, top). In comparison, with CH@AGM corresponding to a negative surface charge (Fig. 2F), Zn$^{2+}$ or Li$^+$ counterions provide surface conduction to maintain electrodeposition, and surface conduction eventually dominates bulk diffusion in carrying the current through the depleted region, leading to a significantly enhanced overlimiting current (25). Meanwhile, SO$_4^{2-}$ or PF$_6^-$ counterions are repelled by the large electric field, thus further enhancing bulk depletion and resulting in a “deionization shock” analogous to pressure shocks in gases, leading to a uniform “shock electrodeposition” in the wake and suppressing dendritic growth (Fig. 2G, bottom). The mechanisms of surface-driven overlimiting current in a microchannel, which was shown to influence electrodeposition in this work, were predicted theoretically by Dydek and co-workers (35). Subsequently, Nam and co-workers performed in situ particle tracking and current-voltage measurements in a micro-/nanofluidic device to investigate the transition from surface conduction to electro-osmotic flow (36). All these reports provide strong evidence to support the surface conduction-controlled metal growth proposed in this work.

To investigate the electrochemical evidence for such deionization shock-controlled metal growth, we performed SEM analyses of the Zn and Li anode after a rapid chronoamperometry experiment (10 min) in Zn and Li metal batteries at $-135\ mV$ overpotential, which continuously sustain overlimiting current in the cell. To eliminate the effect of mechanical stress from CH@AGM on Zn and Li deposition, a hollow rubber film was stacked in the battery assembly. Owing to the short deposition time, no sharp dendrites can be seen on all the anodes. However, quantities of Zn and Li bushy-like deposits, which are the precursor of large dendrites, are observed on the anodes in pristine LMO|AGM|Zn and LMO|AGM|Li batteries (Fig. 2, H and J). In contrast, the surface of Zn and Li deposited beneath CH@AGM shows a dense layer with uniform metal hills (Fig. 2, I and K). Clearly, the positive surface charge in pristine AGM blocks cation penetration and induces the formation of dendrites avoiding the pore walls, while negative surface charge in CH@AGM leads to a uniform metal electrodeposition stabilized by a deionization shock. Khoo and co-workers have studied the linear stability of transient electrodeposition in a charged random porous medium through sophisticated modeling and concluded that after diffusion limitation is reached, a negative background...
charge significantly stabilizes the electrode surface instability, while a positive background charge destabilizes the system (37). This theory agrees well with our experimental phenomenon, which is a critical evidence to support the proposed mechanism in this work.

We performed galvanostatic electrochemical impedance spectroscopy (EIS) in symmetric Zn and Li cells at different dc biases to study how charged porous media affect electrodiffusion. As shown in fig. S4 (A and B), when applying a dc bias, the Warburg-like arc for Zn|AGM + CH@AGM|Zn in 1 M ZnSO₄ shrinks as the current increases because of the surface conduction-driven overlimiting current as confirmed in Fig. 2C, while the low-frequency response of Zn|AGM|Zn diverges along with the increased current, which is due to the blocking of Zn²⁺. Consistent trend was also observed in the impedance spectra of Li|AGM + CH@AGM|Li and Li|AGM|Li
in 1 M LiPF₆ in 1:1:1 DMC:EC:DEC (fig. S4, C and D). These data demonstrate good agreement with galvanostatic impedance spectra of copper deposition using a negative or a positive porous separator as reported by Han et al. (24, 25), indicating the generality of background charge in cation diffusion.

We further used Song’s theory in electrochemical impedance under charged medium to interpret the galvanostatic EIS data in this work, which clearly describe the diffusion and electromigration of ions in a charged background (38). In Song’s model, a one-dimensional (1D) system from \( x = 0 \) to \( x = L \) is considered, which contains two oppositely charged mobile species and another charged immobile species (38). Two common cell configurations are considered for boundary conditions. In the reservoir configuration, an ideal reservoir maintains dimensionless concentration of charge carriers \( \tilde{c} \) and electric potential \( \tilde{\phi} \), while in the symmetric configuration, the boundary at \( x = 0 \) has the same semi-blocking condition (dimensionless flux of negatively charged carriers \( \tilde{F} = 0 \)) identical to the boundary at \( x = L \) (38). After solving the boundary value problem and calculating the impedance for any combination of steady-state dimensionless direct current bias \( \tilde{j}^{(0)} \) and charge density of immobile species \( \tilde{\rho} \), the limiting behaviors of dimensionless impedance \( \tilde{Z} \) as a function of \( \tilde{j}^{(0)} \) in both reservoir and symmetric configurations are obtained. The calculation shows that both low-frequency resistance \( \tilde{R}_L \) and high-frequency resistance \( \tilde{R}_H \) increase exponentially under the diffusion-limiting regime and then converge to \( \tilde{Z}^{\infty} \) over the diffusion-limiting regime, when the background charge is negative \( (\tilde{\rho} < 0) \). When \( \tilde{\rho} > 0 \), \( \tilde{R}_L \) and \( \tilde{R}_H \) diverge before the diffusion-limiting bias.

To compare our experimental data to this theory, we fitted the impedance spectra of Zn and Li symmetrical cells through the equivalent circuit models (fig. S5) and obtained the solution resistance \( (R_s) \) and mass transfer resistance \( (R_m) \), corresponding to the sum of \( R_m \) charge transfer resistance \( R_c \) and bulk diffusion resistance \( R_d \). Figure S6 shows the \( R_s \) and \( R_m \) as a function of applied current in Zn and Li symmetrical cells. It is found that \( R_s \) in both Zn|AGM + CH@AGM|Zn and Li|AGM + CH@AGM|Li cells increases in the regime below limiting current \( (11.3 \, \text{mA cm}^{-2} \, \text{for Zn and } 2.06 \, \text{mA cm}^{-2} \, \text{for Li metal batteries}) \) and then converges in overlimiting current (fig. S6, A and B). As \( R_s \) corresponds to high-frequency resistance \( (\omega \rightarrow \infty) \), these results show close agreement with Song’s calculations in the trend of \( R_m \) demonstrating the transition in the impedance behavior around the diffusion-limiting bias with a charged immobile species. On the other hand, \( R_m \) decreases with increasing current. It can be attributed to the shrinking diffusion layer as the depletion zone expands (24). In the case of Zn|AGM |Zn and Li|AGM |Li cells, however, both \( R_s \) and \( R_m \) diverge as the current is increased (fig. S6, C and D), which is a strong evidence of severe ion depletion.

We also used Khoo’s model to interpret the voltammetry data in Fig. 2 (C and D). Khoo and Bazant (39) proposed a system of a charged nanoporous medium filled with a binary electrolyte flanked on the left by anode and on the right by cathode. Through numerically solving the steady-state equations under different boundary conditions, steady-state current-voltage relations are obtained (39). With negative surface charge, \( I \) varies linearly with \( V \) for a sufficiently large \( V \) and the overlimiting conductance \( \sigma_{OLC} \) is the gradient of this linear relationship. At a positive surface charge, \( I \) tends to a finite maximum value as \( \rightarrow \infty \). Experimental voltammetry data in symmetrical Zn and Li cells (fig. C and D) agree well with this model, indicating the sensitivity of surface conduction to the sign of the surface charge. By performing a linear fit of the linear portion of the current-voltage relation, the experimental \( \sigma_{OLC} \) of Zn|AGM + CH@AGM|Zn and Li|AGM + CH@AGM|Li cells can be obtained, which shows a value of 0.164 and 0.188 ohm⁻¹, respectively. Moreover, Yan and co-workers also simulated an increasing trend of current in response to a voltage ramp in a liquid electrolyte with one electrode and negative background charge (40), which further supports the proposed mechanism in this work.

On the basis of the surface conduction-driven theory, for a current smaller than the limiting current, dendrites will not occur anyway, and there is little effect of charge surface on metal electrodeposition. To verify this, we performed a 72-hour galvanostatic deposition of Zn and Li in symmetric cells at a current density of \(-1.5 \, \text{mA cm}^{-2} \), which is below the limiting current density from Fig. 2 (C and D). No sharp dendrite can be found from the SEM images of Zn and Li electrode after deposition, regardless of the CH in the separator, demonstrating that stable growth of Zn and Li is possible below the limiting current (fig. S7). However, root-growing mossy structures are still formed during electrodeposition in both Zn and Li electrodes (fig. S7). Similar to dendrites, mossy structures are mechanically instable and may also affect the battery life, which appears below the limiting current in Zn and Li metal batteries (41, 42). It should be noted that the moss-structured area in the Zn and Li electrode with CH in the separator (fig. S7, A and C) is much smaller than the ones in the sample without CH (fig. S7, B and D). Clearly, there should be another mechanism to stabilize the anode in Zn and Li metal batteries using CH@AGM, which will be discussed in the next section.

**CH-guided Zn and Li electrodeposition on the anode surface**

It is reported that during electrochemical tests, the extra heat generated from the side reactions accumulates in the electrode and results in large temperature rise within the cell (43). It is also well known that CH can be slowly dissolved in water and organic solvents when the temperature is above 40°C (44). As a result, there should be some CH fine particles stripped from CH@AGM, dispersed in both aqueous and organic electrolytes in CH@AGM-based batteries, and then absorbed on the anode surface during the long-term chronocamperometry test. This assumption was confirmed by the chemical compositions of the Zn and Li metal anodes with CH@AGM in the separator through x-ray photoelectric spectroscopy (XPS), where the N component from CH was detected after 10- and 30-min chronocamperometry at \(-135 \, \text{mV overpotential in the Zn and Li anode, respectively (fig. S8 and tables S1 and S2). However, such CH dispersing in the electrolyte is only partial from this electrochemical condition. As evidenced in fig. S9, even after 1-hour chronocamperometry, a large amount of CH coatings can still be observed on the AGM separator.}

The interaction between Zn²⁺ and Li⁺ cations with CH was first studied using isothermal titration calorimetry (ITC), which is the most direct and quantitative approach for characterizing the thermodynamic properties of protein-ligand interactions. In CH, glycine (Gly), proline (Pro), and alanine (Ala) hold more than 50% of the total amino acids (Fig. 1B). As a result, an aqueous solution of Gly-Pro-Ala peptide (10⁻⁴ M) is used as titrate to represent CH in ITC, while 10⁻³ M aqueous ZnSO₄ or Li₂SO₄ acts as titrant. Strong interaction between Gly-Pro-Ala and Zn²⁺/Li⁺ is confirmed through the exothermicity of the calorimetry peaks in Fig. 3 (A and B),
which indicates the released heat in one injection of titrant into the amino acids. As the sites available on Gly-Pro-Ala to interact Zn\(^{2+}\)/Li\(^+\) are occupied during titration, the exothermicity of the peaks decreases and finally saturates. For comparison, aqueous solution of \(10^{-4}\) M colloidal SiO\(_2\) is used as titrate to investigate the Zn\(^{2+}/Li^+\) cations binding to AGM, as SiO\(_2\) is the main component of glass fiber in AGM. Figure 3 (C and D) shows the calorimetric response of colloidal SiO\(_2\) with the reaction of ZnSO\(_4\) or Li\(_2\)SO\(_4\) at the pH 7.4. The data show a significant scatter with no obvious trend. A comparison of the heats evolved in the injection of Zn\(^{2+}/Li^+\) to Gly-Pro-Ala (Fig. 3, A and B) with that in the reaction of Zn\(^{2+}/Li^+\) with SiO\(_2\) (Fig. 3, C and D) shows that the interaction of Zn\(^{2+}/Li^+\) with SiO\(_2\) is quite weak.

Figure 3 (E and F) shows plots of the integrated heat response from calorimetry data in Fig. 3 (A to D) against the molar ratio of Zn\(^{2+}/Li^+\) cations to amino acids in the reaction vessel. The heat data, combined with the known quantity of titrant and titrate, are fit using a nonlinear least squares method to a binding model, which can yield the best fit values of the stoichiometry (n), binding constant (K), and heat of binding (enthalpy, ΔH). As shown in table S3, the binding constant from the injection of Zn\(^{2+}\) and Li\(^+\) cations to Gly-Pro-Ala solution is 1.13 \(\times 10^4\) M\(^{-1}\) and 4.38 \(\times 10^7\) M\(^{-1}\), respectively, which is two orders of magnitude higher than the values of SiO\(_2\) with Li\(^+\) and Zn\(^{2+}\). These characterizations thus exhibit an obvious interaction between amino acids and Zn\(^{2+}\) or Li\(^+\) cations, which is consistent with previous theoretical studies (45).

Inspired by the strong interactions between CH and Zn\(^{2+}/Li^+\) cations as quantified from ITC characterization, we speculate that the CH fine particles absorbed on the anode surface can also bind to Zn\(^{2+}/Li^+\) cations, which may affect the dendrite growth on the anode surface as well. To verify this hypothesis, we intentionally designed a battery for chronoamperometry experiment, in which CH fine particles was dispersed in the aqueous and organic electrolyte [2% (w/w)] and absorbed on the anode surface, using the pristine AGM as separator. Figure 3 (G and H) shows the SEM images of the Zn anode after 10-min chronoamperometry, which enables continuous overlimiting current to induce dendrite, with and without CH in the aqueous electrolyte, respectively. Without the dispersing effect from CH@AGM on cation transport through the separator, adding only CH into the electrolyte cannot guarantee a uniform electrodeposition in the cell. Dendritic growth is evident on both Zn anodes. However, there are still big differences in their dendrite morphology. In the aqueous electrolyte with 2% CH, the anode surface was only partially covered with Zn coating, with a grain diameter smaller than 1 μm. In comparison, continuous granular microstructure with grains up to 10 μm in diameter was formed on the anode surface in the pristine aqueous electrolyte. An analogous phenomenon is also observed after 30-min chronoamperometry for the Li anode (Fig. 3, I and J). The surface of Li deposits in the electrolyte with 2% CH exhibits a densely packed Li metal flake with smooth morphology, while abundance of uneven and dead Li deposits embedded with AGM residues is detected on the Li anode in the pristine electrolyte. Obviously, CH fine particles absorbed on the anode surface can significantly refine grain size, obtain uniform structure, and affect the texture and dendrite growth direction.

The binding structures of Zn\(^{2+}/\)CH fine particles in the aqueous electrolyte at various mass ratios were investigated using Raman spectroscopy in 1550 to 1780 cm\(^{-1}\) (Supplementary note 2). The pure CH in water shows a band around 1590 to 1670 cm\(^{-1}\) that is associated with amide I C═O stretching (46). After adding ZnSO\(_4\), a new band at 1630 to 1740 cm\(^{-1}\) is probably due to the binding of CH with the Zn\(^{2+}\) ions. By deconvoluting into two Gaussian-Lorentzian bands, the Raman bands at 1650 cm\(^{-1}\) (I\(_{free}\)) and 1680 cm\(^{-1}\) (I\(_{bind}\)) can be attributed to the concentration of free CH molecules ([CH]\(_{free}\)) and CH molecules bonded to Zn\(^{2+}\) ([CH]\(_{bind}\)) (fig. S10). The plot of I\(_{bind}/[CH]\) versus I\(_{free}/[CH]\) in the ZnSO\(_4\)-CH electrolyte with various ZnSO\(_4\)-CH mass ratios shows a linear feature, and the molar scattering coefficient ratio of bonded and free CH (I\(_{bind}/[CH]\)) can be obtained from the slope of the plot (Fig. 3K). By knowing I\(_{free}/[CH]\) and I\(_{bind}/[CH]\), the binding number (n\(_{bind}\)) of CH with Zn\(^{2+}\) can be calculated from eqs. S2 to S7 in Supplementary note 1 (47).

As mass ratio increases, the concentration of CH decreases, and its binding number with Zn\(^{2+}\) decreases, probably due to the depletion of the CH.

We further used \(^{15}\)N and \(^{7}\)Li nuclear magnetic resonance (NMR) with pulsed-field gradient (PFG) to study the association of CH with Li\(^+\) ions (Supplementary note 2) (48). For an ideal NMR sample, all the compounds should be fully dissolved in the solvent to obtain a reliable result. Unfortunately, the solubility of CH in 1:1:1 DMC:EC:DEC mixed solvent is very low. Therefore, it is not possible to prepare a clear NMR sample in the mixture of Li salt and CH under an organic electrolyte. To overcome this challenge, here we studied the binding behavior of Li\(_2\)SO\(_4\) with collagen in aqueous solution to simulate the association of Li\(^+\) with CH in an organic electrolyte, by reasonably assuming that solvent will not affect the interaction between Li\(^+\) and CH. Figure S11 shows that along with Li\(_2\)SO\(_4\)-CH mass ratio increases, the \(^{15}\)N NMR peak from CH, \(\delta^{(15)N}\), shifts upfield due to the screening of Li\(^+\) ions on the lone electron pair density of the oxygen in C═O groups from amino acids in CH (49). Owing to the Li\(^+\) ion complexation by CH, the upfield shift of the \(^{7}\)Li NMR peak, \(\delta^{(7)Li}\), is proportional to mass ratio. Diffusion coefficients of CH (D\(_{CH}\)) and Li\(^+\) ions (D\(_{Li^+}\)) were obtained using \(^{15}\)N and \(^{7}\)Li NMR, respectively. As shown in Fig. 3M, diffusion coefficients decrease along with the mass ratio increase. In all mass ratios, D\(_{CH}\) is higher than D\(_{Li^+}\), and both D\(_{CH}\) and D\(_{Li^+}\) decrease with the mass ratio increase. The values of D\(_{Li^+}\) and D\(_{CH}\) become the same after the mass ratio reaches 1, indicating the firm binding of Li\(^+\) with CH.

The above molecular-level analysis suggests a clear binding mechanism between CH fine particles and Zn\(^{2+}\) or Li\(^+\) cations in aqueous solution and indicates a possible guiding effect of CH in cation movement and nucleation on the anode surface. As shown in Fig. 3N (right), Zn or Li will be preferentially deposited around the tips of protuberances on the CH-free anode rather than on smooth regions to form large dendrites, due to the strong electrical field on the sharp edge. In contrast, the strong binding of CH fine particles absorbed on the anode to Zn\(^{2+}\) or Li\(^+\) cations imposes a competing effect to the deposition of Zn\(^{2+}\) or Li\(^+\) cations on the protuberances, which spreads Zn or Li to adjacent regions of the initial growth tips of protuberances and prevents the formation of large dendrites (Fig. 3N, left). The computer simulation (density functional theory) also indicates that Zn tends to grow into 2D clusters around -NH-C═O in CH due to the more favorable energy, which facilitates Zn deposition along the surface/horizontal direction of the anode and leads to flake-shaped small dendrites (fig. S12). However, on the pristine anode surface, Zn favors the vertical direction of growth.
to minimize the surface area exposed to the electrolyte (fig. S12), which results in the large dendrites on the flat anode (Fig. 3N, right). Of course, the actual electrochemical cation deposition is much more complicated than the nucleation phenomena discussed here and deserves in-depth study in the future.

**Long-term electrochemical cycles**

The long-term interfacial stability of the Zn anode was first investigated through symmetric Zn|AGM + CH@AGM|Zn and Zn|AGM|Zn cells using an aqueous 1 M ZnSO₄ and 2 M Li₂SO₄ electrolyte. During cycling, Zn ions are transported back and forth between the
two Zn metal electrodes at 1.0 mA cm$^{-2}$, and the corresponding voltage versus time plot is shown in Fig. 4A. Amplified voltage profiles in Fig. 4A for 0 to 4 hours (16,000 s) and 36 hours (130,000 s) to 43 hours (155,000 s) of cycling are provided in Fig. 4 (C and D). Lower mass transfer–controlled potential ($\mu_{\text{mtc}}$) values and better potential retentions are realized for the Zn|AGM + CH@AGM|Zn cell than for the Zn|AGM|Zn cell, indicating a homogeneous Zn metal electrode with a stable electrolyte/metal interface. The electrochemical impedances of Zn symmetrical cells were measured at the end of 0-, 20-, and 40-hour deposition/dissolution cycles in the 1 M ZnSO$_4$ and 2 M Li$_2$SO$_4$ electrolyte at 1.0 mA cm$^{-2}$. As shown in Fig. 4G, the initial charge transfer resistance ($R_{\text{ct}}$) of Zn|AGM + CH@AGM|Zn is equal to that of Zn|AGM|Zn (29.3 versus 27.8 ohms, after fitting the Nyquist plots through equivalent circuit in fig. S13), indicating the well-retained pore structure after CH coating. The $R_{\text{ct}}$ remains stable at around 23.6 ohms in 40-hour cycling, indicating the stabilized Zn growth and little consumption in the electrolyte. However, with pristine AGM separators, the $R_{\text{ct}}$ exhibits large fluctuations during cycling. The value significantly drops to 12.9 ohms after 20-hour cycling and further reduces to 6.7 ohms in 40 hours (Fig. 4H). Such large fluctuations in $R_{\text{ct}}$ suggest unstable electrochemical environments during Zn depositing and stripping, which induces breakage of native oxide on the surface and exposes the fresh Zn to the electrolyte.

Furthermore, the Li migrations for the symmetrical Li|AGM + CH@AGM|Li and Li|AGM|Li cells using the organic electrolyte (1 M LiPF$_6$ in 1:1:1 DMC:EC:DEC mixed solvent) are measured at 1 mA cm$^{-2}$ (Fig. 4B). Magnified voltage profiles in Fig. 4B for 0 to 9 hours (35,000 s) and 70 hours (255,000 s) to 78 hours (280,000 s) of cycling are shown in Fig. 4 (E and F). Compared with the cell using routine AGM separators, the Li|AGM + CH@AGM|Li cell renders a tighter voltage hysteresis (200 to 500 mV versus >5000 mV for the pristine AGM) and much more extended cycle life (111 hours versus <85 hours for AGM), indicating the depletion of Li ions in the electrolyte resulting from the dead Li. Figure 4 (I and J) shows typical impedance spectra measured at the end of 0-, 30-, and 60-hour cycling compared to the value (from 18.2 to 7.3 ohms) in the Li|AGM|Li cell, implying the stable Li growth. Clearly, the CH in Li and Zn significantly stabilizes Zn and Li growth, which affords new insights into the realization of stable cycling in Zn and Li metal batteries using both aqueous and organic electrolytes.

We also assembled coin-type Li|AGM + CH@AGM|Cu and Li|AGM|Cu half-cells using 1 M LiPF$_6$ in 1:1:1 DMC:EC:DEC as...
electrolyte, to further evaluate the long-term stability of Li plating/stripping behavior on a metal substrate. High Li reversibility and CE (~99%) are achieved in the Li|AGM + CH@AGM|Cu cell at a current density of 1 mA cm⁻², while the pristine Li|AGM|Cu cell fails after about the 80th cycle (fig. S14). The polarization profiles in different cycles were also recorded to gain the kinetic behavior during Li stripping/plating (fig. S15). As shown in fig. S15A, Li|AGM + CH@AGM|Cu exhibited a similar Li migration potential at the 10th and 80th cycle from the charge/discharge curves in 1 mA cm⁻² (126 mV versus 139 mV), indicating a stable Li electrodeposition. In comparison, large polarization was observed in the Li|AGM|Cu cell, which resulted in a much larger Li migration potential at the 80th cycle (182 mV; fig. S15B).

**Electrochemical performance of full cells**

It is well known that for currents beyond diffusion limitation, the concentration of salt at the electrode surface decreases to zero at a certain time. This time is known as “Sand’s time,” after which the cations preferentially deposit on the protrusions of the surface, resulting in a self-amplifying process of dendritic growth (42). By multiplying with the current density, we converted Sand’s time to Sand’s capacity and plot the current-dependent Sand’s capacity in Zn and Li metal anodes (fig. S16), which clearly shows the transition from mossy to dendritic metal. Because of the much higher limiting current in Zn deposition (11.3 mA cm⁻²) than that in Li deposition (2.06 mA cm⁻²), the Zn metal electrode exhibits a harsher experimental condition to form dendrites than Li metal. Moreover, the results in the Li metal electrode (fig. S16B) show close agreement with the previous literature revealing the conditions for dendritic versus mossy growth of lithium (42). To demonstrate the feasibility of CH@AGM in Zn metal batteries, we first investigate the cycling performances of LMO|AGM + CH@AGM|Zn and LMO|AGM|Zn laboratory-scale cells, in which a low-capacity LMO cathode (3.6 mA h cm⁻²), a thick Zn metal, and a much flooded electrolyte [electrolyte weight versus cathode capacity (E/C) of 30 g(Ah)⁻¹] were used (table S4 and fig. S17). Owing to the low cathode capacity, these cells fall into the mossy region of fig. S16. As shown in Fig. 5A and fig. S18, the LMO|AGM + CH@AGM|Zn cell exhibited a good rate capability and stable cycling over 1200 cycles at 1 C with an initial capacity retention above 90% and efficiency above 98%. When using the regular AGM as separator, the capacity retention of the cell starts to decline after 400 cycles with an obvious efficiency decay. The cycling performance in Fig. 5A is further analyzed by observing the charge-discharge curves of LMO|AGM + CH@AGM|Zn and LMO|AGM|Zn at a specific cycle. Compared to the voltage profile of LMO|AGM|Zn that shows a rapid potential spike at first charge and gradually leftward and up movement of the charge-discharge intersection (Fig. 5B), the steady potential spike at the beginning of the first charge and leftward horizontal movement of the traced intersection points are observed in the LMO|AGM + CH@AGM|Zn cell (Fig. 5C). It implies that the Zn anode is stably operated during cycling and the decrease in capacity of this cell is mainly due to the unavoidable degradation of the cathode.

In addition, we further evaluated the 60°C cycling performance of an aqueous Zn metal battery under the same C rate as in room temperature. As shown in Fig. 5D, the LMO|AGM + CH@AGM|Zn cell exhibits more than 600 stable cycles with above 99% Coulombic efficiency in such harsh conditions. In contrast, the pristine LMO|AGM|Zn cell shows a sharp capacity and efficiency decay after 400 cycles. To reveal the morphology changes of Zn metal anodes during charge-discharge interaction, ex situ SEM of Zn anodes was conducted after 400 cycles under 60°C. A uniform Zn metal layer is retained during cycling in the cells with the CH@AGM separator [Fig. 5E (left) and fig. S19]. In contrast, irregular Zn deposits are observed beneath the AGM residues in the cell using the pristine AGM separator [Fig. 5E (right) and fig. S20]. The much better cycling performance in Zn|AGM + CH@AGM|Zn than in Zn|AGM|Zn under 60°C mainly comes from the binding of dissolved CH to the Zn²⁺ on the anode surface, which spreads the distribution of cations over the whole surface of the metal anode to stabilize Zn growth (Fig. 3N, right). Previous reports revealed that larger total surface areas resulted in more metal ion consumptions in electrolytes. Therefore, stabilizing Zn growth through CH@AGM contributes to the prolonged life span, which is consistent with the long-term plating/stripping performances obtained from symmetrical cells.

Moreover, we have also investigated the cycling properties of LMO|AGM + CH@AGM|Li and LMO|AGM|Li cells, using a low-capacity LMO cathode (1.8 mA h cm⁻²) and a flooded electrolyte (table S5) that also fall into the mossy regime similar to Zn metal batteries (fig. S16). As expected, such a LMO|AGM + CH@AGM|Li full cell shows stable cycling over 800 cycles (Fig. 5F) with more stable voltage profiles (Fig. 5H). When using the regular AGM separator, the pristine LMO|AGM|Li cell (Fig. 5, F and G) shows a quick capacity drop after 300 cycles. The ex situ SEM images of the Li anode in LMO|AGM + CH@AGM|Li and LMO|AGM|Li cells after 600 and 200 cycles are shown in Fig. 5I. Rather than forming disordered Li mosses in the LMO|AGM|Li cell, the Li surface in the LMO|AGM + CH@AGM|Li cell exhibits a uniform and densely packed Li deposition layer. This phenomenon suggests a unique effect of CH in mosses’ formation on the Li and Zn anode, which is derived from the interaction between dissolved CH and Li and Zn ions in the electrolyte as discussed in the previous section. The mass of dissolved CH in aqueous and organic electrolytes during cycling can be obtained by calculating the mass of CH@AGM before and after battery cycling. As plotted in fig. S21, the highest content of CH in aqueous (13%) and organic (9%) electrolytes was reached after 200 and 400 cycles, respectively, demonstrating the maximum solubility of CH.

Although stable cyclability of the LMO cathode is verified in both Zn metal and Li metal full batteries by adding CH@AGM between anode and separator, the result is still lacking actual practicalities for a real battery as the cathode areal capacity in the above cells are extremely low (3.6 mA h cm⁻² for the Zn metal battery and 1.8 mA h cm⁻² for the Li metal battery), which greatly reduces the total capacity in a full battery. However, it is highly challenging to construct cycling stable Zn and Li metal batteries using high–mass loading cathode materials. The higher area capacity actually involves a higher current density, and a large fraction of Zn and Li metal was reacted in each charge-discharge cycle, which generally produces a fast Zn and Li consumption that causes an accelerated depletion of metal ions in the electrolyte and premature cell failure. For instance, under conventional condition with low-capacity cathode (3.6 mA h cm⁻²; table S4), after 1200 cycles, the LMO|AGM|Zn cell still shows a high capacity of 95 mA h g⁻¹. However, when tested under constrained conditions with a cathode capacity of 24 mA h cm⁻², this cell only retains 65% of its original capacity after 400 cycles (Fig. 6A). At the 450th cycle, the battery encountered an immediate short circuit, which is due to the piercing of obvious large dendrites formed on
the anode surface as this battery falls into the dendritic Zn regime in fig. S16. This performance discrepancy from cell configurations is even larger in the LMO|AGM|Li cell, which can only survive 10 cycles with high-capacity cathode (24 mAh cm\(^{-2}\)) as evidenced in Fig. 6B, in comparison to the results of the LMO|AGM|Li cell in conventional testing conditions with a cathode capacity of 1.8 mAh cm\(^{-2}\) (Fig. 5F). It should be noted that within this high-loading cathode, both ion transportation and mechanical property will be poor, and the changes in cathode structure also lead to the fading in Zn and Li metal batteries. It is reported that the diffusion of Li\(^+\) inside the electrolyte is faster than that inside the cathode (50). Abundant Li\(^+\) tends to accumulate on the surface of Li transition metal oxides and induce a partial transformation in the crystalline structure (51). This structural distortion, the so-called Jahn-Teller distortion, severely blocks the Li\(^+\) diffusion pathways and reduces the capacity retention (52). Moreover, the electrochemical oxidation of the conductive carbon in the cathode is responsible for the capacity loss of Li-ion batteries. It is induced by contacting water at low potentials and/or by oxygen evolved from the unstable delithiated materials (53, 54). As a result, we attribute the capacity fading of Zn and Li metal batteries in Fig. 6 (A and B) to the combination of structure instabilities in cathode materials, and the metal depletion in the anode, both of which can be greatly amplified under a high-loading cathode.

Unexpectedly, when CH@AGM was added in the Zn metal battery (LMO|AGM + CH@AGM|Zn) with a 24 mAh cm\(^{-2}\) cathode, a stable cycling over 600 cycles was realized with a capacity retention of 92% at a rate of 1 C (Fig. 6A). In addition, when CH@AGM was used, the Li metal battery using the same 24 mAh cm\(^{-2}\) cathode can maintain an excellent cycling stability of 93% capacity retention until the 100th cycle and even 68% capacity retention at the 600th cycle (Fig. 6B). Moreover, the charge/discharge curves of both batteries were well retained during cycling without an obvious overpotential increase (Fig. 6, C and D), due to the relatively stable interface between the metal anode and the electrolyte. The superimposed effects of CH-enabled cation regulation in the separator and on the anode surface are crucial to achieve such a highly stable cycling in Zn or Li metal batteries under realistic conditions. Figure S22 shows the cycling performance of the LMO|AGM|Zn cell, using CH as electrolyte additives and pristine AGM as separator under the same
battery conditions in Fig. 6A. This cell can be charged-discharged over 500 cycles without short circuit. However, compared to the cell with CH@AGM that exhibits more than 100 mAh g⁻¹ capacity at the 600th cycle (Fig. 6A), this cell only shows a low capacity of 81 mAh g⁻¹ after 500 cycles and a rapid capacity decrease after the 500th cycle. As evidenced from the SEM image of the Zn anode at the 500th cycle, although no bushy dendrites are found, large amounts of accumulated Zn grains inside the pores of the AGM separator are still formed on the surface of the anode, which can seriously affect the cycle life of a rechargeable battery (fig. S23).

A comprehensive comparative summary of cathode areal capacities and reported cycle decay per cycle of other state-of-the-art Li-ion electrochemistry systems: LiCoO₂/Li₄Ti₅O₁₂ (LCO/LTO) (55), LiFePO₄/Li (LFP/Li) (56), LiNi₀.₅Mn₁.₅O₄/Li (LNMO/Li) (57), TiS₂/Li (58), V₂O₅/Li (59), LiNi₀.₅Mn₁.₅O₄/Co₂O₃/Li (NCM811/Li) (16), LiFePO₄/Cu (LFP/Cu) (60), and LiNi₀.₅Mn₁.₅O₄/Co₂O₃/Li (NMC622/Li-C) (15). (F) Cycling performance of the discharge capacity of 5-Ah LMO|AGM + CH@AGM|Zn and LMO|AGM|Zn cells under constrained cell parameters at a rate of 1 C. (G) Photos of 5-, 65-, and 200-Ah cells. (H) Cycling performance of the discharge capacity of 5-, 65-, and 200-Ah LMO|AGM + CH@AGM|Zn cells at a rate of 1 C. Photo credit: Yunfeng Luo, Rehab Energy Co. Ltd.

Li-ion batteries is presented in Fig. 6E. It can be seen that both LMO|AGM + CH@AGM|Zn and LMO|AGM + CH@AGM|Li batteries are standouts for either cathode loading or cycling life. Clearly, CH@AGM enables the high stability and rechargeability of Zn and Li metal batteries with realistic, high-loading cathode, which suggests a significant promise for the industrial fabrication in high-capacity Li-ion batteries. For the purpose of commercialization, we have also fabricated a 5-Ah commercial high-capacity battery based on a single LMO|AGM + CH@AGM|Zn cell (1.056 Ah, with a 80-cm² cathode sheet; see fig. S24) with constrained conditions (cathode capacity of 13.2 mAh cm⁻²; table S4). Note that to seek the ease of quality assurance and industrial manufacturing, we intentionally choose a less constrained cell parameter (relatively lower cathode areal capacity compared to the battery in Fig. 6A) in a single cell. However, this battery still falls into the dendritic regime of Zn metal (fig. S16) due to the much higher cathode loading than the laboratory-scale
battery. As shown in Fig. 6E, a 5-Ah LMO|AGM + CH@AGM|Zn full cell can realize over 600 stable cycles under 100% depth of discharge. We have also fabricated 65-Ah and 200-Ah high-capacity Zn metal batteries (Fig. 6G), both of which show an excellent cyclability over 400 cycles with >90% capacity retention (Fig. 6H). We have also tried to further decrease the amount of electrolyte to increase the energy density of the commercial battery. When an extremely constrained E/C ratio (3 g Ah⁻¹) was used, the single battery unit from the 200-Ah cell, even with CH@AGM in the separator, can only survive 80 cycles (fig. S25). Obviously, the electrolyte consumption becomes a limiting factor when an extremely low amount of electrolyte was used. More optimization in electrolyte formulation is needed in this system.

DISCUSSION
In this contribution, we report a facile and general approach to resolving the stability issue in Zn and Li metal batteries with a practical, high-loading intercalation cathode, where a cation-regulating layer CH@AGM between anode and separator can effectively suppress metal dendrite formation. The surface charge of CH and the binding between CH and cations (Li⁺ or Zn²⁺) can simultaneously induce a deionized shock inside the separator and realize a homogeneous distribution of cations on the anode surface, which leads to efficient dendrite suppression and control in both Zn and Li metal batteries. Benefiting from superimposed effects of CH in cation regulation in electrolyte and on the anode surface, Zn and Li metal full cells coupled with CH@AGM and a 3.6 mAh cm⁻² cathode delivered up to 1200 dendrite-free cycles at 1 C and exhibit over 600 stable cycles with above 99% Coulombic efficiency under 60°C. At a cathode loading as high as 24 mAh cm⁻², Li and Zn metal batteries using CH@AGM can still deliver 600 dendrite-free cycles at 1 C in both organic and aqueous electrolytes, with a Coulombic efficiency of up to 99.7%. In contrast, the same batteries with the pristine AGM separator could only survive for 10 and 100 cycles, respectively. In addition, such CH-based dendrite control technology can be readily applied in large-scale, Ah class battery systems. By adding CH@AGM, 600 and 400 cycles were realized in 5-Ah class and 200-Ah class batteries at 100% depth of discharge, respectively, which show more than 30% improvement compared to the pristine Zn metal batteries. The proposed cation-biomolecule interaction for uniform metal deposition and nucleation suggests a scalable process to realize dendrite-free charge/discharge in various metal anodes, which can be facilely translated to commercial-level battery designs.

MATERIALS AND METHODS
Preparation of CH@AGM
The CH solutions are prepared by 5% [weight % (wt %)] CH (Sigma-Aldrich) mixed with deionized water with constant stirring for 2 hours at 80°C. The fabrication of CH@AGM used in the Zn metal battery is slightly different to the fabrication of CH@AGM used in Li metal batteries. For the Zn metal battery, AGM was fully immersed into CH solution for a few seconds and followed by air drying at 110°C for 12 hours to obtain CH@AGM. For the Li metal battery, 1 wt % LiOH (Sigma-Aldrich) was added into 5% aqueous CH solution. Then, AGM was immerged into the solution for a few seconds and followed by air drying at 110°C for 12 hours to obtain CH@AGM. The mass loading of absorbed CH in AGM is about 38 mg cm⁻² for the Zn metal battery and 43 mg cm⁻² for the Li metal battery.

Battery assembly
For both Li and Zn metal batteries, the cathodes are the same, which were prepared by casting slurries of LMO (Wuxi Jewel Power & Materials Co., Ltd.), conductive carbon (KS-6, MTI, Co.) with styrene-butadiene rubber/carboxymethylcellulose as binding material (83:10:7 wt.%) in N-methyl-2-pyrrolidinone (Sigma-Aldrich Co.) on a conductive polyethylene film. The mass loadings on the cathode of Zn and Li metal batteries under unconstrained testing conditions are 30 and 15 mg cm⁻², corresponding to a 3.6 and 1.8 mAh cm⁻² cathode, respectively. For batteries under constrained conditions, the mass loading on the cathode of Zn and Li metal batteries is 200 mg cm⁻².

To assemble a LMO|AGM + CH@AGM|Zn battery under conventional, unconstrained conditions, a 3.6 mAh cm⁻² LMO cathode (4 cm², square) was coupled with the conventional AGM separator. Then, CH@AGM was attached on the top of the separator before assembling a 147 mAh cm⁻² Zn anode (250-μm thickness); 0.42 g of 2 M LiSO₄ and 1 M ZnSO₄ aqueous solution (pH 4) was used as the electrolyte, to make the E/C ratio equal to 30 g(Ah)⁻¹. The full battery pack is shown in fig. S17.

To assemble a LMO|AGM + CH@AGM|Li battery under unconstrained conditions, a 1.8 mAh cm⁻² LMO cathode (0.95 cm², disk) was coupled with the AGM separator, and then CH@AGM and a 51.5 mAh cm⁻² Li anode (250-μm thickness) were subsequently assembled on top of the separator to obtain the full cell; 0.09 g of mixed solution of 1 M LiPF₆ in dimethyl carbonate, ethyl carbonate, and diethyl carbonate (1:1:1 by weight) was used as the electrolyte. The E/C ratio is 60 g(Ah)⁻¹, and the battery is tested under a CR2032 coin-cell kit.

To assemble a LMO|AGM + CH@AGM|Zn battery under constrained conditions, a 24 mAh cm⁻² LMO cathode (9 cm², square) was coupled with the AGM separator, CH@AGM, and a 58 mAh cm⁻² Zn anode (100-μm thickness); 2.16 g of 1 M LiPF₆ in dimethyl carbonate, ethyl carbonate, and diethyl carbonate (1:1:1 by weight) was used as the electrolyte, to make the E/C ratio equal to 10 g(Ah)⁻¹. The battery pack is assembled in an argon glove box, and the testing pack used in this battery is the same as that of the LMO|AGM + CH@AGM|Zn battery under unconstrained conditions (fig. S17), but the cathode, anode, and separator are differently sized.

To assemble a LMO|AGM + CH@AGM|Li battery under constrained conditions, a 24 mAh cm⁻² LMO cathode (4 cm², square) was coupled with the AGM separator, CH@AGM, and a 51.5 mAh cm⁻² Li anode (250-μm thickness); 0.98 g of mixed solution of 1 M LiPF₆ in dimethyl carbonate, ethyl carbonate, and diethyl carbonate (1:1:1 by weight) was used as the electrolyte, and then CH@AGM was connected with the AGM separator. To assemble a LMO|AGM + CH@AGM|Zn battery under unconstrained conditions (fig. S17), but the cathode, anode, and separator are differently sized.

To assemble an LMO|AGM + CH@AGM|Zn battery, 1-Ah single-cell unit was first prepared. Briefly, a 13.2 mAh cm⁻² LMO cathode (80 cm² rectangular sheet; fig. S24) was coupled with the AGM separator, CH@AGM, and a 147 mAh cm⁻² Zn anode (250 μm thickness). Five, 65, and 200 repeated single-cell units were stacked together in a factory-made battery bay to assemble 5-, 65-, and 200-Ah batteries. After adding 150 g of 2 M LiSO₄ and 1 M ZnSO₄ aqueous solution (pH 4) as electrolyte, the battery pack was sealed and aged for 24 hours before charge-discharge tests.

To prepare a 2 M LiSO₄ and 1 M ZnSO₄ aqueous electrolyte with CH for LMO|AGM|Zn cells, 0.5% (weight %) CH (Sigma-Aldrich) was mixed with deionized water with constant stirring for 2 hours at 80°C. The fabrication of CH@AGM used in the Zn metal battery is slightly different to the fabrication of CH@AGM used in Li metal batteries. For the Zn metal battery, AGM was fully immersed into CH solution for a few seconds and followed by air drying at 110°C for 12 hours to obtain CH@AGM. The mass loading of absorbed CH in AGM is about 38 mg cm⁻² for the Zn metal battery and 43 mg cm⁻² for the Li metal battery.
was mixed with deionized water with constant stirring for 12 hours at 80°C. In this process, the CH was fully swollen by water and decomposed into CH. After cooling to room temperature, Li2SO4 and ZnSO4 were added to the liquid to make 2 M Li2SO4 and 1 M ZnSO4 aqueous solution (pH 4).

**Electrochemical test**

Cycling and rate tests were performed using a Neware series 3000 battery tester and maintained at 25° or 60°C under the working voltage of 1.4 to 2.1 V versus Zn/Zn for the Zn metal battery or the working voltage of 3.5 to 4.3 V versus Li/Li for the Li metal battery. Chronoamperometry measurements were performed on a Bio-Logic VMP3 electrochemical workstation at −135 mV overpotential for Zn and Li metal batteries. Chronopotentiometry and linear sweep voltammetry were also performed on the same instruments for symmetric Zn and Li cells. The cycling performance of symmetric Zn and Li cells was tested by a multichannel battery tester (BTS-5V10mA, Neware) at a 1 mA cm−2 current density. Potentiotstatic and galvanostatic EIS were conducted on a Biologic VMP3 electrochemical workstation using the symmetric cells, at an amplitude of 10 mV or various dc biases.

**Morphology and structure characterization**

The SEM images of the separator, lithium, anode and zinc anode were taken on UltraPlus field-emission scanning electron microscopes equipped with energy-dispersive spectroscopy for elemental analysis. Fourier transform infrared and Raman spectroscopy spectra were taken on UltraPlus field-emission scanning electron microscopes equipped with energy-dispersive spectroscopy for elemental analysis. Fourier transform infrared and Raman spectroscopy spectra were taken on UltraPlus field-emission scanning electron microscopes equipped with energy-dispersive spectroscopy for elemental analysis. Fourier transform infrared and Raman spectroscopy spectra were taken on UltraPlus field-emission scanning electron microscopes equipped with energy-dispersive spectroscopy for elemental analysis. Fourier transform infrared and Raman spectroscopy spectra were taken on UltraPlus field-emission scanning electron microscopes equipped with energy-dispersive spectroscopy for elemental analysis. Fourier transform infrared and Raman spectroscopy spectra were taken on UltraPlus field-emission scanning electron microscopes equipped with energy-dispersive spectroscopy for elemental analysis.

**Isothermal titration calorimetry**

ITC experiments were conducted using a MicroCal VP-ITC instrument, wherein 300 µl of an aqueous solution of 10−3 M ZnSO4 or Li2SO4 was injected in equal steps into the cell containing 10−3 M Gly-Pro-Ala peptide. For comparison, aqueous solution of colloidal SiO2 is used as titrate to investigate the Zn2+/Li+ cations binding to AGM. Each negative peak shown in the heat signal curves represents an exothermic process, which denotes the heat released in one injection.
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