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Introduction

Lithium-ion is currently the go-to high-energy battery technology, but its inherent safety issues persist and represent a continued challenge, particularly for large-format applications. To enable wider implementation in electric vehicles and grid-level energy

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Capacity and phase stability of metal-substituted α -Ni(OH)₂ nanosheets in aqueous Ni–Zn batteries†

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Batteries that offer high specific energy and energy density coupled with improved safety and lower cost will affect applications ranging from electric vehicles, portable electronic devices, and grid-level energy storage. Alkaline nickel-zinc (Ni-Zn) batteries use nonflammable aqueous electrolyte and nonstrategic, low-cost electrode materials; however with a two-electron anode, a cathode that stores more than one electron per Ni atom would increase energy density. Herein, we report the effect of substituting metal ions (aluminium, cobalt, manganese, or zinc) into α -Ni(OH)₂, a phase that can accommodate more than one-electron charge storage, but which typically converts to lower-capacity β -Ni(OH)₂ upon cycling in alkaline electrolytes. We adapt a microwave-assisted process that expresses α -Ni(OH)₂ as a high surface-area nanosheet morphology and find that we retain this morphology with all metal-ion substituents. The series is characterized using scanning electron microscopy, transmission electron microscopy, X-ray diffraction, and Raman spectroscopy. Metal-ion substitution influences aggregate growth, interlayer distance, and vibrational frequencies. We test powder-composite cathodes prepared using the substituted α -Ni(OH)₂ series versus zinc sponge anodes in alkaline electrolyte under devicerelevant mass loadings and using an intentionally aggressive charging protocol to determine onset voltage for oxygen evolution. The electrochemical charge-storage behaviour is established using galvanostatic cycling and differential capacity analysis. The substituents significantly influence both Ni-centred redox and oxygen-evolution voltages (vs. Zn/Zn^{2+}). The incorporation of Al³⁺ within α -Ni(OH)₂ nanosheets provides higher capacity and phase stability compared to the divalent substituents and unsubstituted α -Ni(OH)₂. The presence of ordered free nitrates in the interlayer of Al³⁺-substituted α -Ni(OH)₂, not seen with Co^{2+} or Mn^{2+} substituents, correlates with the improved electrochemical performance.

> storage, non–lithium-based batteries with high specific energy and energy density, improved safety, and lower cost are essential. The alkaline nickel–zinc (Ni–Zn) battery is an attractive alternative to Li-ion due to its use of nonflammable aqueous electrolyte.

> The three-dimensional (3D) aperiodic zinc architecture ("sponge") developed at the U.S. Naval Research Laboratory solves long-standing performance limitations for zinc anodes¹ and paves the way for next generation Ni–Zn batteries competitive with the specific energy of Li-ion systems.² Rewiring the zinc anode as a porous 3D monolith enables more uniform charge and discharge at greater zinc utilization while preventing dendritic growth at the zinc anode. The architected metal opens the door to zinc-based battery chemistries that approach the system-level specific energy and energy density of Li-ion without the accompanying weight and volume of ancillary structures required to mitigate system-failures. A practical Ni–Zn battery provides a specific energy of up to 135 W h kg⁻¹ (energy density of ~ 300 W h L⁻¹) depending on the Zn depth of

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discharge. The projected specific energy of a fully packaged Ni–Zn system is comparable or superior to a Li-ion pack that includes catastrophe management systems.²

Several fabrication iterations of the Zn sponge anode have further improved electrochemical performance, scalability, and mechanical durability such that the capacity of the cathode limits cell performance.^{3–8} Deep utilization of the zinc sponge in a Ni–Zn cell for 150 cycles requires swapping out not one, but four commercially harvested nickel hydroxide cathodes per Zn sponge anode.^{5,8} Matching the performance of next-generation zinc anodes in terms of capacity and rate requires improving alkaline cathode materials and architectures.

Commercial Ni-Zn batteries use β -Ni(OH)₂-based cathodes that have practical capacities of 237–263 mA h g^{-1} ,^{9,10} which capture up to 90% of the theoretical $Ni^{2+/3+}$ one-electron capacity of 289 mA h g⁻¹. Challenges inherent to β -Ni(OH)₂ include a steep drop in capacity at high rates and generating γ -NiOOH upon overcharge, which results in large volumetric changes leading to battery failure.¹¹ Compared to β -Ni(OH)₂, α -Ni(OH)₂ has a higher theoretical capacity (~487 mA h g⁻¹) storing \sim 1.67 electrons per Ni center, which partially poises Ni as Ni⁴⁺ upon charging.^{12,13} Electrochemical and spectroscopic support for Ni⁴⁺ within charged Ni(OH)₂ thin films has been previously reported.^{14,15} Prior work suggests the oxidation state of Ni is influenced by interlayer ions.^{14,16–18} At high pH, α -Ni(OH)₂ is unstable converting to β -Ni(OH)₂,¹⁹ thus cycling α-Ni(OH)2 in alkaline electrolyte inevitably leads to lower capacity.20

Metal substituents can influence the phase stability and electrochemical performance of α -Ni(OH)₂. Prior studies find that the stability and charge-storage capacity of α -Ni(OH)₂ is affected by such cation substituents as aluminium,^{9,21–28} cobalt,^{24,25,29–31} copper,^{31–34} chromium,^{31,35} iron,³¹ manganese,^{31,35–37} tin,³⁸ and zinc.^{31,39,40} Aluminum (Al) substitution was determined to increase the interlayer spacing and increase discharge capacity relative to unsubstituted α -Ni(OH)₂.^{9,27,41} Cobalt-substitution into α -Ni(OH)₂ accompanied by ethylene glycol intercalation between the layers is reported to stabilize the α -phase when evaluated as an active material in supercapacitors⁴² and alkaline batteries.⁴³ Interlayer distance and electrochemical performance are also affected by anions that accompany the cation substituent.²⁶

In addition to the capacity of the cathode, redox potentials are an important consideration when operating within an aqueous alkaline electrolyte because $Ni(OH)_2$ electrocatalyzes the oxygen-evolution reaction (OER). In alkaline electrolyte, OER occurs near the $Ni(OH)_2/NiOOH$ redox potential, and as a parasitic reaction, it decreases charge efficiency and lowers discharge capacity.¹¹ Coprecipitation with a variety of metals can influence the Ni redox potential and either catalyze or poison OER.³¹

The route used to synthesize α -Ni(OH)₂ plays a significant role in the resulting morphology, structure, and electrochemical properties.⁴⁴ Common routes include chemical precipitation, electrochemical precipitation, sol–gel synthesis, chemical aging, hydrothermal/solvothermal synthesis, and conversion of a nickel alloy.⁴⁵ Microwave chemistry provides a rapid synthesis method and can influence the kinetics, selectivity, and yield of chemical reactions.⁴⁶ We recently reported that microwave-synthesized α -Ni(OH)₂ nanosheets serve as a high surface-area precursor to high-activity bimetallic OER electrocatalysts.^{47,48} Microwave synthesis using neutral/mild pH conditions generates α -Ni(OH)₂ with a 3D flowerlike morphology⁴⁹ unlike the low surface-area aggregates characteristic of highly basic conditions.^{44,45,50}

Microwave synthesis has been used to synthesize a number of metal-substituted α -Ni(OH)₂ materials such as Mn/ Co-substituted α -Ni(OH)₂ as a precursor for Ni/Mn/Co oxide materials for supercapacitors,⁵¹ 3D flowerlike Co-substituted α -Ni(OH)₂ for supercapacitors,^{52,53} and Al/Zn substituted α -Ni(OH)₂.⁵⁴ These compositions were not explored, however, as active cathode materials in aqueous rechargeable Ni-Zn batteries. report microwave-synthesized, metal-ion substituted We α -Ni(OH)₂ and the effects of aluminium, cobalt, manganese, or zinc substitution on morphology, structure and electrochemical performance when formulated into cathodes and cycled versus zinc sponge anodes in alkaline Ni-Zn cells. To determine the effect of substituents on oxygen evolution at the Ni cathode, we do not use a charging protocol optimized for coulombic efficiency and long-term cycling; instead, we design an intentionally aggressive one that allows us to evaluate OER onset voltage.

Experimental

Microwave synthesis of Ni(OH)₂ nanosheets

The α -Ni(OH)₂ nanosheets were synthesized using a microwavesynthesis technique modified from our previously reported microwave-assisted hydrothermal method,^{47,48} by adapting the protocol to a large-format microwave reactor at the Naval Research Laboratory (CEM Mars 5 Digestion Microwave System microwave reactor). In the modified synthesis, Ni(NO₃)₂·6H₂O (3.0 g, Sigma-Aldrich) and urea (2.46 g, Sigma-Aldrich) were added to a solution of ethylene glycol (63 mL, Sigma-Aldrich) and ultrapure water (9 mL, \geq 18 M Ω cm; used for all subsequent reactions). The unstirred solution was exposed to variable microwave radiation at 120 °C for 39 min. The green precipitate was collected by centrifugation and washed five times with water and three times with ethanol (200 proof, Pharmco), and then dried overnight at 60 °C (Fig. 1a).

Metal-substituted α -Ni(OH)₂ nanosheets (notated as Ni-M₁₀; M = Al³⁺, Co²⁺, Mn²⁺, or Zn²⁺) were synthesized by partially substituting M for Ni²⁺ at a Ni-to-M ratio of 9:1 (*i.e.*, 10 atomic % substitution). For the metal-substituted α -Ni(OH)₂ synthesis, either Al(NO₃)₃·9H₂O (0.387 g, Sigma-Aldrich), Co(NO₃)₂·6H₂O (0.30 g, Alfa Aesar), or Zn(NO₃)₂·6H₂O (0.306 g, Sigma-Aldrich) was added to a solution of Ni(NO₃)₂·6H₂O (2.7 g, Sigma-Aldrich), urea (2.46 g, Sigma-Aldrich) ethylene glycol (63 mL, Sigma-Aldrich) and 9 mL of ultrapure water. The respective solutions were then microwaved and washed as described above. The manganese-substituted (Ni-Mn₁₀) reaction was synthesized using a small-scale reactor (CEM Discover SP) at Texas State University, and Ni(NO₃)₂·6H₂O (0.9 g, Ward's



Fig. 1 (a) Photograph of microwave-synthesized powders–25 mg each of unsubstituted α -Ni(OH)₂, aluminum-substituted (Ni–Al₁₀), cobalt-substituted (Ni–Co₁₀), manganese-substituted (Ni–Mn₁₀), and zinc-substituted (Ni–Zn₁₀) nanosheets; (b–f) scanning electron micrographs of microwave-synthesized nanosheets at three different magnifications (6500×, 25000×, 100000×); (b) unsubstituted α -Ni(OH)₂ nanosheets, (c) aluminum-substituted nanosheets (Ni–Al₁₀); (d) cobalt-substituted nanosheets (Ni–Co₁₀); (e) manganese-substituted nanosheets (Ni–Mn₁₀); and (f) zinc-substituted nanosheets (Ni–Zn₁₀).

Science), urea (0.82 g, VWR Analytical) and $Mn(NO_3)_2 \cdot 4H_2O$ (0.086 g, Alfa Aesar) were added to a solution of ethylene glycol (21 mL, VWR Analytical) and ultrapure water (3 mL). The solution was then exposed to variable microwave radiation at 120 °C for 13 min while magnetically stirring. The precipitate was collected and processed as described above (Fig. 1a).

To evaluate potential phase conversions from α -Ni(OH)₂ to β -Ni(OH)₂, a hydrothermal treatment method was used to treat unsubstituted α -Ni(OH)₂ and Ni–Al₁₀. The hydrothermal treatment was adapted from a previously reported method⁵⁵ and consisted of

placing a suspension of 70 mg of the material and 40.0 mL of 0.01 M NaOH (Alfa Aesar) in a 50 mL Teflon-lined hydrothermal pressure reactor (Parr Instruments Co., model 276AC2) and then heating in a 180 °C-oven for 2 h. The solid was separated by centrifugation and rinsed three times with water and twice with ethanol. The powder was dried overnight at 60 °C.

Materials characterization

Morphology of the microwave-synthesized solids was probed using scanning electron microscopy (SEM; FEI-Helios Nanolab 400).

Nitrogen physisorption measurements were conducted using a Micromeritics ASAP 2020, where the sample was degassed under vacuum at 120 °C for 16 h prior to analysis. Specific surface area was determined by Brunauer–Emmett–Teller (BET) analysis while pore diameters and pore volumes were determined by Barrett–Joyner–Halenda (BJH) analysis of the desorption isotherm. Elemental analysis of the amount of Ni and M within Ni–Al₁₀, Ni–Co₁₀, Ni–Mn₁₀, and Ni–Zn₁₀ was determined from inductively coupled plasma-atomic emission spectroscopy (ICP-AES) of digested samples performed by Galbraith Laboratories (Knoxville, TN).

Powder X-ray diffraction measurements (XRD) were performed using a Bruker AXS D8 Advanced powder X-ray diffractometer with a Cu K α radiation source, operating at 40 kV and 25 mA and a high-resolution energy-dispersive 1D Linxeye XE detector with a 2θ scan range between 5–80° with a 0.01 increment. Raman spectroscopy experiments were carried out using a Horiba LabRam HR evolution confocal Raman spectrometer using a backscattering geometry and a 600 grooves per mm grating. A 514 nm argon-ion laser line focused through a 50×-objective Olympus microscope was used for the excitation source. A laser power of <1 mW at the sample was maintained using a neutral density filter. The Raman instrument was calibrated using the 520.7 cm⁻¹ band of Si. The sample was not visibly damaged by the incident laser.

Selected samples were imaged by high-angle annular darkfield (HAADF) imaging in a Nion UltraSTEM 200-X scanning transmission electron microscope (STEM) operated at 60 kV with a detector inner semi-angle of 60 mrad. Thickness of the nanosheets was measured using electron energy-loss spectroscopy (EELS) data and calculating the log-ratio absolute thickness using the algorithm in DigitalMicrograph and the following parameters: 10 mrad convergence semi-angle, 50 mrad collection semi-angle, and 24.6 effective atomic number (Z_{eff}).⁵⁶ Thickness accuracy is estimated to within 10%.

Electrode and cell fabrication

Ni-foam current collectors (MTI Corp.) were punched out (round, 1/2 inch diameter) and cleaned as recommended by the manufacturer, which consisted of sonicating in an ice bath for 5 min each in concentrated hydrochloric acid (12 M, Fisher Scientific), water, and ethanol (200 proof, Warner-Graham Company). The current collectors were then dried for 10 min under vacuum at 60 °C to remove residual ethanol. Composite cathodes were fabricated by preparing a slurry composed of the active material (a-Ni(OH)2, Ni-Al10, Ni-Co10, Ni-Mn10, or Ni-Zn₁₀), carbon (100% compressed, 99.9%+ acetylene black, Alfa Aesar), and binder (PTFE dispersion in water, Sigma-Aldrich, item number 665800) using a weight percentage ratio of 70:25:5 for the active material, conductive carbon, and binder, respectively. For preparing the slurry, the active material (60 mg) and conductive carbon (21.4 mg) were mixed by hand with an agate mortar and pestle and then combined with 12-15 drops of water in a vial. The slurry was sonicated for 10 min. The binder was then added, and the mixture was

sonicated an additional 10 min. The Ni-foam current collectors were immersed in the slurry and treated under vacuum for 30 min. The cathodes were dried overnight under ambient conditions and then further dried at 60 °C for 2 h. Following drying, the cathodes were pressed by rolling through a pasta roller (Imperia) which produced an electrode 1.26 cm in diameter and 0.4 mm-thick. The average mass loading of active material in the carbon-nanocomposite vacuum-infiltrated Ni-foam electrodes is 11.5 mg cm⁻². Achieving >10 mg cm⁻² represents an important metric to evaluate the performance of Ni(OH)₂ electrodes under practical loadings.⁵⁷

Nickel-zinc coin cells were constructed in a similar manner to that previously reported using a "Gen 3" zinc sponge as the anode inside a gas-tight laser-cut acrylic alkaline-resistant cell.⁵ Briefly, the cells were constructed (from the anode up) using a tin-foil current collector (0.25 mm thick, 99.8% Alfa Aesar), Gen-3 Zn-sponge anode, Celgard separator (3501), Freudenberg separator (700/28), carbon composite cathode embedded in Ni foam, expanded Ni-mesh current collector (Dexmet), and a second Freudenberg separator within gas-tight ethylene propylene diene monomer gaskets. Before cell assembly, the cathode and the separators were vacuum infiltrated with a 6 M KOH/1 M LiOH electrolyte for 30 min (electrolyte volume: 0.3-0.4 mL);⁵ the zinc sponge was vacuum infiltrated for the same amount of time in a dispersion of 11 wt% Ca(OH)₂ in 6 M KOH/ 1 M LiOH. The coin-cell experiments were designed such that the capacity of the active cathode material in the cell was significantly lower than the Zn anode capacity enabling a study of the cathode material within a cathode-limited cell configuration. The cells were fabricated in the discharged state using Zn sponge anodes fabricated with ~ 30 wt% ZnO.⁵ Within the cathode-limited cell configuration, the depth of discharge of the Zn anode is < 3%.

Electrochemical characterization

Galvanostatic experiments were carried out using an Arbin battery cycler. The cells were charged and discharged at a mass-normalized current rate of 84.4 mA g_{active}^{-1} (0.2C based on a theoretical capacity of 422 mA h g_{active}^{-1} for α -Ni(OH)₂) between 1.3 and 2.0 V. During charging, if the cell's theoretical capacity was reached before the upper voltage limit, the cell automatically began discharging. The cycling conditions were not optimized for battery performance but chosen to study the effect that the metal substituents impart on the redox and OER onset voltages of α -Ni(OH)₂. We note that the cycling protocol we used to purposefully induce OER will lower coulombic efficiency and contribute to lower electrochemical discharge capacity compared to previous studies.^{5,26,34,44}

The cell was charged and discharged four times to activate the cathode material and nickel current collector; these cycles are designated as formation cycles and are not included during cycling analysis.^{26,58} The contribution of the Ni-foam current collector to the discharge capacity is negligible (<1%) relative to the active material, in agreement with prior studies.^{59,60} The 5th cycle was designated as the starting cycle to evaluate cycling stability. The mass of the active material (α -Ni(OH)₂, Ni–Al₁₀, $Ni-Co_{10}$, $Ni-Mn_{10}$, or $Ni-Zn_{10}$) was calculated to be 70% of the weight of the electrode after subtracting the mass of the Ni foam. Reported capacities represent an average obtained from multiple tests. The capacity retention upon cycling was determined by dividing the specific discharge capacity of the 35th cycle by the specific discharge capacity of the 5th cycle.

Differential capacity analysis $(\partial Q/\partial V)$ was performed in Origin by taking the derivative of the mass-normalized capacity with respect to the cell voltage. The resulting $\partial Q/\partial V$ plots were smoothed within Origin (FFT Filter) removing all highfrequency signals. The subsequent mass-normalized $\partial Q/\partial V$ plots were fit with Origin's Fit Peak Pro which uses a Levenberg– Marquardt algorithm. Without the use of high-precision $\partial Q/\partial V$ analysis, the raw data required a line smoothing and peak fitting analysis in order to compare samples.⁶¹

Results and discussion

Analysis of metal-ion substituent composition

During microwave synthesis, selected metal ions $(Al^{3+}, Co^{2+},$ Mn^{2+} , and Zn^{2+}) are substituted into α -Ni(OH)₂ at nominal Ni:M atomic ratios of 9:1 (notated as either $Ni_{0.9}M_{0.1}$ or Ni $-M_{10}$). For comparison to the precursor stoichiometry, the Ni:M ratios measured by ICP are nominal for Ni_{0.90}Co_{0.10}, higher than targeted M incorporation for Ni_{0.87}Al_{0.13} and Ni_{0.84}Zn_{0.16}, and significantly below target value for Ni_{0.98}Mn_{0.02} (Table 1). Factors that may influence the relative incorporation of metal-ion substituents within the Ni(OH)₂ structure include differences in charge density, ionic radii, metal ligand-field strength, solubility product constants, and pH, any of which can influence solution-phase speciation and rates of hydrolysis and condensation.⁶² Metal ions with larger ionic radii, such as Zn²⁺ are reported to have higher olation rates while cations with higher ligand-field strength have lower rates of olation due to a greater orbital overlap between metallic d-orbitals and H₂O.⁶² Olation rates affect the rate of condensation; rates that are too different between the Ni and the metal substituent may influence the degree of substitution within the Ni(OH)₂ structure. Some of these factors likely influence the low relative incorporation of manganese (Ni_{0.98}Mn_{0.02}) compared to the targeted synthetic ratio (Ni_{0.9}Mn_{0.1}).

Effect of substituents on morphology and porosimetry

Microanalysis of microwave-synthesized α -Ni(OH)₂ using highangle annular dark-field (HAADF) and scanning transmission electron microscopy-energy dispersive spectroscopy (STEM-EDS) confirms a uniform distribution of Ni and O (Fig. S1a and b, ESI†). The material is expressed as stacks of nanosheets with a thickness of ~2 nm (6–7 layers) to ~12 nm (~40 layers) as determined by electron energy-loss spectroscopy (Fig. S2 and Table S1, ESI†). The morphology of α -Ni(OH)₂ as evaluated using scanning electron microscopy (SEM) (Fig. 1b) comprises large, round aggregates (diameter ~1.5 µm) of interconnected nanosheet stacks. The lower pH and solution composition of our synthetic protocol, which includes urea and ethylene glycol, leads to wispy, less densely packed layers compared to previously reported nanostructured α -Ni(OH)₂.^{44,45,50}

Upon partial substitution of Al³⁺, Co²⁺, Mn²⁺, or Zn²⁺ for Ni²⁺, the overall nanosheet morphology is maintained (Fig. 1c–f); however, the metal-ion substituents influence the size of the aggregated nanosheets. Compared to unsubstituted α -Ni(OH)₂, substitution with Al³⁺ affects sheet growth by producing smaller, more densely interwoven nanosheets (Fig. 1c) that aggregate into smaller particulates (~0.5 µm). The remaining substituents result in larger aggregates relative to unsubstituted α -Ni(OH)₂: Ni–Co₁₀ (~4 µm, Fig. 1d), Ni–Mn₁₀ (~3 µm, Fig. 1e) and Ni–Zn₁₀ (~10 µm, Fig. 1f). The changes observed in the aggregate size of the microwave-synthesized nanosheets upon substituent on solution-phase speciation and hydrolysis and condensation reaction rates, as discussed above.

Nitrogen physisorption analysis tracks the influence of the substituents on BET surface area, average pore diameter, and cumulative pore volume (Table 1). Unsubstituted α -Ni(OH)₂ nanosheets exhibit a BET surface area of 87 m² g⁻¹, expressing a much higher surface area than materials derived from a microwave-assisted hydrothermal route using water, urea, and NiCl₂ (9.2 m² g⁻¹).⁵² The greatest impact on BET surface area occurs upon substitution with Al³⁺ (40 m² g⁻¹), consistent with its more dense aggregates of interconnected nanosheets (Fig. 1c), but which is still higher than NiAl materials synthesized in the presence of EtOH and urea (24.1 m² g⁻¹).⁶³ The surface areas of Ni-Co₁₀ (55 m² g⁻¹), Ni-Mn₁₀ (76 m² g⁻¹), and Ni-Zn₁₀ (77 m² g⁻¹) are lower overall compared to unsubstituted α -Ni(OH)₂, but not to the same extent as Ni-Al₁₀.

All substituents decrease the pore volume relative to native α -Ni(OH)₂ (Table 1). The pore size distribution plots (Fig. S3, ESI†) indicate that the majority of the free volume for all samples is in the mesopore (>2–50 nm) and macropore range (>50 nm), which is advantageous for electrochemical

Table 1 Physiochemical characteristics of unsubstituted α -Ni(OH)₂, aluminum-substituted (Ni–Al₁₀), cobalt-substituted (Ni–Co₁₀), manganesesubstituted (Ni–Mn₁₀), and zinc-substituted (Ni–Zn₁₀) nanosheets obtained from elemental analysis (ICP-OES), nitrogen porosimetry, X-ray diffraction, and Raman spectroscopy; details are provided in the text

	Relative atomic % from synthesis	Relative atomic % from ICP analysis	BET surface area $(m^2 g^{-1})$	Pore volume $(cm^3 g^{-1})$	Interlayer <i>d</i> -spacing (Å)	$\nu(E_g)$	ν(M(O/OH))	ν(-OH stretch)
α-Ni(OH) ₂ Ni–Al ₁₀ Ni–Co ₁₀	— Ni _{0.90} Al _{0.10} Ni _{0.90} Co _{0.10}	— Ni _{0.87} Al _{0.13} Ni _{0.90} Co _{0.10}	87 40 55	0.579 0.370 0.279	7.70 8.72 8.17	458 461 459	 537 521	3652 3653 3656
Ni-Mn ₁₀ Ni-Zn ₁₀	$\frac{Ni_{0.90}Mn_{0.10}}{Ni_{0.90}Zn_{0.10}}$	$\begin{array}{c} Ni_{0.98}Mn_{0.02} \\ Ni_{0.84}Zn_{0.16} \end{array}$	76 77	0.380 0.391	7.88 7.81	447 448	524 522	3656 3648

applications because this size range improves electrolyte flux to the charge-storing interfaces.^{47,64} Substitution with Co²⁺ results in the lowest mesopore volume (Fig. S3, ESI[†]).

X-ray diffraction

The effects of Al³⁺, Co²⁺, Mn²⁺, or Zn²⁺ substituents on the crystal structure of α -Ni(OH)₂ nanosheets are studied by powder X-ray diffraction (Fig. 2). Unsubstituted α -Ni(OH)₂ nanosheets exhibit three prominent diffraction peaks at 11.28°, 33.32°, and 59.56° 2 θ , which are consistent with the alpha phase of Ni(OH)₂ and are assigned as the $\langle 001 \rangle$, $\langle 110 \rangle$, and $\langle 300 \rangle$ planes respectively.⁴⁵ The $\langle 001 \rangle$ plane corresponds to the reflections of Ni(OH)₂ layers within the crystal structure. Prior work on



Fig. 2 X-ray diffraction of unsubstituted α -Ni(OH)₂, aluminum-substituted (Ni–Al₁₀), cobalt-substituted (Ni–Co₁₀), manganese-substituted (Ni–Mn₁₀), and zinc-substituted (Ni–Zn₁₀); (a) diffraction pattern with a 2 θ scan range between 5–80°; (b) expanded region for a 2 θ scan range between 8–14° showing changes of the $\langle 001 \rangle$ plane associated with the interlayer spacing of α -Ni(OH)₂.

hydrated layered Ni(OH)₂ has shown that the position of the $\langle 001 \rangle$ peak depends on drying conditions.⁶⁵ The broad peak at 23.72° 2 θ is assigned as a $\langle 002 \rangle$ reflection, which is related to the $\langle 001 \rangle$ plane and indicates ordering within the interlayer region. The asymmetric character of the $\langle 110 \rangle$ plane is characteristic of the turbostratic disorder of α -Ni(OH)₂.^{9,26,66}

The effects of substitution on the (001) plane of α -Ni(OH)₂ are presented in Fig. 2b and Table 1. The metal substituents shift the (001) peak relative to α -Ni(OH)₂; these shifts correspond to larger or smaller interlayer spacings arising from differences in the amounts and coordination of water, anions or other interlayer guests.^{26,27,45} In Ni-Al₁₀, the interlayer spacing increases from 7.70 Å to 8.72 Å, while STEM-EDS analysis confirms that Al is uniformly distributed within the material (Fig. S1c and d, ESI⁺). The increased interlayer spacing is attributed to the influx of negatively charged anions such as NO₃⁻, OH⁻, or CO₃²⁻ required to balance the increased Al³⁺-derived electrostatic charge.^{23,26,27} Within Ni-Al layered hydroxides, anions adopt different orientations and packing within the interlayer.^{16,67} A high degree of ordering within the interlayer region of Ni-Al₁₀ is evident from the relative intensity of the $\langle 002 \rangle$ plane (Fig. 2a). Our Raman data (discussed below) support a higher relative concentration of nitrates within Ni–Al₁₀ nanosheets compared to unsubstituted α -Ni(OH)₂.

The diffraction pattern of Ni–Zn₁₀ shows an additional lowangle peak at 9.72°, not previously observed for Zn²⁺substituted α -Ni(OH)₂.^{39,40,68} The presence of two low-angle peaks within Ni–Zn₁₀ is consistent with a two-phase material. Within Ni–Zn₁₀, the peak at 11.42° may arise from domains with no or low amounts of Zn²⁺ within the interlayer region, while the peak at 9.72° may result from domains with an interlayer pillared by Zn²⁺. A prior study of a zinc-containing layered vanadium-oxide material⁶⁹ showed that the vanadium oxide layers were pillared by interlayer water and Zn²⁺ ions.

Raman spectroscopy

Raman spectroscopy provides information on the local structure of the substituted α-Ni(OH)2 nanosheets. The substituents shift the frequency of the intralayer Ni(OH)₂ asymmetric doubly degenerate E_g lattice mode,^{70,71} which indicates that the substituent alters the local potential energy environment (Fig. 3a and Table 1). A previous study of α -Ni(OH)₂ correlates frequency shifts in the Eg mode to changes in the Ni-O bond length; shifts to higher wavenumbers indicate a shorter Ni-O bond length, while lower wavenumbers correspond to a longer bond length.⁷⁰ Al^{3+} -substitution shifts the E_g mode to a higher frequency suggesting the presence of Al3+ shortens the intralayer Ni-O bond length. Co^{2+} -substitution results in a similar $\nu(\text{E}_{\sigma})$ as α -Ni(OH)₂ within experimental error, whereas Mn²⁺ and Zn²⁺ shift the E_g mode to lower frequencies indicating Mn^{2+} and Zn^{2+} increase the intralayer Ni-O bond length. The metal-substituted samples exhibit a new Raman peak between 520 cm⁻¹ and 540 cm⁻¹, which may arise from the vibration of a bond of the substituent metal to oxygen/hydroxide $\nu(M(O/OH)_n)$.^{72–76} The presence of an additional Raman peak and the absence of an accompanying XRD reflection corresponding to a separate



Fig. 3 Raman spectra of unsubstituted α -Ni(OH)₂, aluminum-substituted (Ni–Al₁₀), cobalt-substituted (Ni–Co₁₀), manganese-substituted (Ni–Mn₁₀), and zinc-substituted (Ni–Zn₁₀): (a) 350 cm⁻¹ to 600 cm⁻¹ spectral region and assignments of lattice modes; (b) 900 cm⁻¹ to 1600 cm⁻¹ spectral region and assignments of modes from compounds incorporated into the structure.

phase (with the exception of Ni–Zn₁₀) support location of Al^{3+} , Co^{2+} , and Mn^{2+} within the plane of the Ni(OH)₂ layer.

Raman-active bands from nitrate anions (NO_3^-) are observed within the unsubstituted α -Ni(OH)₂ and metalsubstituted α -Ni(OH)₂ materials, indicating that nitrates from the synthetic precursors incorporate into the structure. Two distinct nitrate chemical environments are observed (Fig. 3b); one at ~ 1042 and ~1352 cm⁻¹ from a "free" nitrate mode and a second mode at ~986 and ~1283 cm⁻¹.⁷¹ A higher relative intensity of the free NO₃⁻ peak occurs for Ni–Al₁₀ and Ni–Zn₁₀ (see also Fig. S4a, ESI†) indicating a higher concentration of nitrates within these samples. Unlike the native and divalent-substituted materials, only free nitrate occurs in Ni–Al₁₀. Energetically, nitrate does not associate with Al³⁺ for two reasons: the highly negative free energy of hydration for Al³⁺ compared to first-row transition metals;⁶² and compensation of the increased electrostatic charge induced by substitution of Al³⁺ for Ni²⁺. Based on the increase in the interlayer spacing of Ni–Al $_{10}$ seen by XRD (Fig. 2), these nitrates are located in the interlayer region, consistent with prior studies of layered Ni(OH) $_2$.^{42,71}

The frequency of the ν (–OH) stretching mode arising from hydroxides within Ni(OH)₂ provides a delineating marker that distinguishes α -Ni(OH)₂ from β -Ni(OH)₂.⁷¹ The substituted Ni(OH)₂ nanosheets exhibit a peak at ~ 3657 cm⁻¹ (Fig. S4b, ESI†) supporting the α -Ni(OH)₂ phase assignment made from XRD (Fig. 2). Additional peaks are observed arising from ν (C–H) stretching modes and ν (–OH) bending modes of ethylene glycol,⁷⁷ carbonate,^{78,79} and cyanate from the decomposition of urea^{49,80–83} (Fig. S4, S5 and Table S2, ESI†), which indicate that compounds or derivatives from precursors used in the synthesis are incorporated into the structure.⁴⁵

Electrochemical characterization

Differential capacity analysis of initial charge-discharge. Differential capacity analysis involves taking the derivative of the cell capacity with respect to voltage $(\partial Q/\partial V)$ thereby transposing the voltage plateau into well-defined peaks. This method can evaluate gradual changes and failure mechanisms batteries^{58,61,84} and allows for current-controlled in measurements that are more translatable to actual devices. In the present work, differential capacity plots are generated directly from the galvanostatic charge-discharge experiments in order to study the influence of substitution on the redox and oxygen evolution voltages of the nickel hydroxide cathodes. The $\partial O/\partial V$ analysis of unsubstituted α -Ni(OH)₂ (Fig. 4 and Table 2) shows an anodic peak centred at 1.80 V assigned to oxidation of Ni²⁺(OH)₂ to Ni³⁺OOH and a corresponding cathodic peak centred at 1.72 V upon reduction. The anodic current at \sim 1.94 V corresponds to onset of the parasitic oxygen evolution reaction (OER). Aging studies of α -Ni(OH)₂ in H₂O show generation of mixed phase α/β -Ni(OH)₂ in as little as four days.¹⁹ Because the formation cycles in this study last from 24-36 h (Fig. S6a, ESI⁺), unsubstituted α -Ni(OH)₂ should already have begun its transformation to β -Ni(OH)₂. Unsubstituted α -Ni(OH)₂, hydrothermally treated α -Ni(OH)₂, and commercial β -Ni(OH)₂ have comparable redox peak voltages and discharge capacities (Fig. S7b-d, ESI^{\dagger}), supporting the conclusion that α -Ni(OH)₂ nanosheets convert to β -Ni(OH)₂ during the formation cycling.

The $\partial Q/\partial V$ analysis of Ni–Al₁₀ (Fig. 4) shows that Al³⁺ substitution shifts the anodic and cathodic peaks and OER onset to more positive voltages compared to unsubstituted α -Ni(OH)₂ (summarized in Table 2). Because the OER voltage shifts positive, Ni–Al₁₀ cathodes offer a wider operational voltage window. The shift in voltage for the Ni-based redox corresponds to a change in chemical potential upon Al³⁺ substitution and is consistent with the frequency shift of the Raman-active E_g lattice mode, as discussed above. Another minor anodic peak appears at a voltage negative of the prominent oxidation peak for Ni–Al₁₀ indicating two unique electrochemical environments exist. Possible environments for the minor peak include an Al³⁺deficient Ni(OH)₂ or a portion of β -Ni(OH)₂ within the material.

Ni-Co₁₀ cathodes shift the Ni-centred oxidation and reduction peaks and OER onset to lower voltages compared



Fig. 4 Mass-normalized differential capacity ($\partial Q/\partial V$) analysis of the 5th cycle galvanostatic charge–discharge of Ni–Zn sponge cells for unsubstituted α -Ni(OH)₂, aluminum-substituted (Ni–Al₁₀), cobalt-substituted (Ni–Co₁₀), manganese-substituted (Ni–Mn₁₀), and zinc-substituted (Ni–Zn₁₀) nanosheets at a mass-normalized current of 84.4 mA g_{active}⁻¹ between 1.3–2.0 V; height of dashed lines indicates relative intensity of mass-normalized differential capacity.

to unsubstituted Ni(OH)₂ (Fig. 4 and Table 2), which is consistent with previous voltammetric studies of Co-substituted α -Ni(OH)₂.^{30,31,85} Prior work suggests that substitution with Co²⁺ can lower the band gap, increasing the conductivity of Ni_{0.5}Co_{0.5} hydroxide nanosheets relative to α -Ni(OH)₂,⁸⁶ while another group postulates that Co(OH)₂ oxidizes to form a CoOOH network.⁵² The anodic and cathodic peak positions and areas of Ni–Mn₁₀ (Fig. 4) are similar to that of unsubstituted Ni(OH)₂ (Table 2), which is not unexpected given the low relative amount of Mn within the structure (Ni_{0.98}Mn_{0.02}). The $\partial Q/\partial V$ plot for Ni–Zn₁₀ differs from the other substituents as noted by the emergence of additional anodic and cathodic peaks (Fig. 4), further supporting the existence of a multi-phase material as discussed above.

Galvanostatic discharge voltage profiles and capacities. To evaluate how substitution affects the specific discharge capacity of α -Ni(OH)₂ nanosheets, voltage vs. specific capacity plots are generated from galvanostatic measurements (Fig. 5); charging



Fig. 5 Voltage vs. specific capacity for 5th cycle discharge of Ni–Zn sponge cells for powder-composite cathodes containing unsubstituted α -Ni(OH)₂, aluminum-substituted (Ni–Al₁₀), cobalt-substituted (Ni–Co₁₀), manganese-substituted (Ni–Mn₁₀), or zinc-substituted (Ni–Zn₁₀) nanosheets at a mass-normalized current of 84.4 mA g_{active}⁻¹ between 1.3–2.0 V.

profiles are provided in Fig. S6a and b (ESI[†]). The discharge process of the Ni–Zn battery involves electroreduction of NiOOH in the cathode (eqn (1)), and electrooxidation of the Zn anode (eqn (2)), complexation of divalent Zn with hydroxide ions in the alkaline electrolyte (eqn (3)), and dehydration/ precipitation (eqn (4)) reactions at the anode to form ZnO, with an overall net reaction described by eqn (5).²

 $2NiOOH_{(s)} + 2H_2O_{(l)} + 2e^- \rightarrow 2Ni(OH)_{2(s)} + 2OH^-_{(aq)}$ (1)

$$Zn_{(s)} \rightarrow Zn^{2+}_{(aq)} + 2e^{-}$$
 (2)

$$Zn^{2^{+}}_{(aq)} + 4OH^{-}_{(aq)} \rightarrow Zn(OH)_{4}^{2^{-}}_{(aq)}$$
 (3)

$$Zn(OH)_4^{2-}(aq) \rightarrow ZnO_{(s)} + H_2O_{(l)} + 2OH_{(aq)}^{-}$$
 (4)

$$2\text{NiOOH}_{(s)} + \text{H}_2\text{O}_{(l)} \rightarrow 2\text{Ni}(\text{OH})_{2(s)} + \text{ZnO}_{(s)}$$
(5)

After the initial formation cycling (the first four cycles), all microwave-synthesized samples exhibit a 5th-cycle voltage

Table 2 Electrochemical properties of Ni–Zn sponge cells using unsubstituted α -Ni(OH)₂, aluminum-substituted (Ni–Al₁₀), cobalt-substituted (Ni–Co₁₀), manganese-substituted (Ni–Mn₁₀), and zinc-substituted (Ni–Zn₁₀) nanosheet cathodes. Differential capacity analysis of cathodic voltages, anodic voltages, and oxygen evolution reaction (OER) voltages averaged over multiple samples. Average specific discharge capacity for the 5th and 35th cycles determined using galvanostatic analysis at a mass-normalized current of 84.4 mA g_{active}⁻¹ between 1.3–2.0 V

	Differential capacity analysis $(\partial Q/\partial V)$			Galvanostatic analysis			
	Cathodic peak voltage (V)	Anodic peak voltage (V)	OER onset voltage (V)	5th cycle (mA h g ⁻¹)	35th cycle (mA h g ⁻¹)	Capacity retention ^a (%)	
α -Ni(OH) ₂	1.72	1.80	1.94	249	198	79	
Ni-Al ₁₀	1.74	1.83	1.96	279	221	79	
Ni-Co ₁₀	1.69	1.77	1.93	172	137	79	
Ni-Mn ₁₀	1.72	1.80	1.94	255	192	75	
Ni-Zn ₁₀	1.73	1.81	1.94	229	167	72	

 a Capacity retention calculated as the ratio of averaged discharge capacity of the 5th and 35th cycles.

plateau between 1.65 V and 1.85 V. The higher discharge voltage for Ni–Al₁₀ relative to α -Ni(OH)₂ and the other substituted materials (Fig. 5) indicates that Al³⁺ substitution affects the electrochemical potential, an energetics argument that is supported by our Raman analysis. We find that Al³⁺ alters the local chemical environment of Ni charge-storage sites by shortening the intralayer Ni–O bond length. Note that Al³⁺-substituted α -Ni(OH)₂ synthesized under basic conditions, rather than the near pH-neutral conditions of our microwave procedure, also exhibits a higher discharge voltage compared to α -Ni(OH)₂.⁸⁷

The voltage–capacity plot of Ni–Al₁₀ shows that Al³⁺-substitution enables a greater discharge capacity of 279 mA h g_{active}⁻¹ for the 5th cycle compared to unsubstituted α -Ni(OH)₂ nanosheets (Fig. 5 and Table 2). The discharge capacity of 255 mA h g_{active}⁻¹ for Ni–Mn₁₀ is similar to that of unsubstituted α -Ni(OH)₂, but with only 2 at% of Mn present (Ni_{0.98}Mn_{0.02}), the similarity to α -Ni(OH)₂ is expected. Both Co (172 mA h g_{active}⁻¹) and Zn (229 mA h g_{active}⁻¹) lower the discharge capacity relative to α -Ni(OH)₂. Capacity suppression in the presence of Co is attributed in the literature to preventing complete oxidation to γ -NiOOH.⁵⁹ The multiphase character of Ni–Zn₁₀ complicates comparison to the single-phase materials.

The discharge capacity of our microwave-synthesized α -Ni(OH)₂ (249 mA h g_{active}⁻¹) is comparable to the discharge capacity of α -Ni(OH)₂ synthesized under basic conditions (269 mA h g_{active}⁻¹)⁸⁷ and α -Ni(OH)₂ synthesized by a sintering process (225 mA h g_{active}⁻¹),⁸⁸ but is lower compared to other studies of α -Ni(OH)₂.^{44,89} The direct comparison with prior studies is complicated by a number of factors that affect capacity including cathode material differences,⁴⁴ loading,⁵⁷ electrolyte,⁹⁰ aging time,⁹⁰ and cycling protocols, as well as our deliberately selected non-practical charging parameters resulting in extended oxygen evolution at each charging step (Fig. S6, ESI†). However, the intraseries comparison of the capacities of metal-substituted α -Ni(OH)₂ to α -Ni(OH)₂ using the same testing conditions supports our conclusion that the substituents influence the discharge capacity.

Analysis of the effect of cycling. The effect of substitution within α -Ni(OH)₂ on cycling performance is examined for 35 charge and discharge cycles (Fig. 6a and Table 2). The discharge capacity of unsubstituted α -Ni(OH)₂ initially fades within the first ten cycles, stabilizing at ~ 200 mA h g_{active}⁻¹ with modest capacity loss thereafter. Post-formation, Ni–Al₁₀ retains a higher capacity over cycles 5–10, whereas the other materials all exhibit a capacity decrease. The whole series stabilizes over cycles 10–35, with Ni–Al₁₀ retaining the highest mass-normalized capacity of the series, while Ni–Co₁₀ has the lowest capacity. After 35 cycles, the substituted materials show similar capacity retention (72–79%) upon cycling as α -Ni(OH)₂ (Table 2).

To evaluate the effect of substitution on cycling, we compared differential capacity plots of the 5th and 35th cycles (Fig. 6b). Upon cycling, the $\partial Q/\partial V$ peaks of unsubstituted α -Ni(OH)₂ shift to higher voltages, consistent with transformation to the β -polymorph and in agreement with a previous study reporting β -Ni(OH)₂ has a higher oxidation voltage.¹¹ The metal-substituted samples show



Fig. 6 Analysis of the effect of cycling of Ni–Zn sponge cells for unsubstituted α -Ni(OH)₂, aluminum-substituted (Ni–Al₁₀), cobalt-substituted (Ni–Co₁₀), manganese-substituted (Ni–Mn₁₀), and zinc-substituted (Ni–Zn₁₀) nanosheet cathodes at a mass-normalized current of 84.4 mA g_{active}⁻¹ between 1.3–2.0 V; (a) specific discharge capacity vs. cycle number; (b) comparison of 5th and 35th cycle differential capacity ($\partial Q/\partial V$) vs. cell voltage; height of dashed lines in (b) indicates relative intensity of mass-normalized differential capacity; specific discharge capacities reported in (a) were averaged from measurements of multiple samples.

similar shifts of the Ni-centred redox peak to higher voltage (Fig. 6b). In addition to shifting the Ni-centred redox peak positive, the OER onset also shifts to higher voltage. The substituents influence the drift of the OER onset voltage to differing amounts. The shift of the OER onset to higher voltage may result from an increase in internal resistance,⁹¹ differences in the OER active site,⁹² or oxygen build up within the electrode⁹³ as a result of inducing OER because of our intentionally aggressive charging protocol. In particular, the OER voltage shift to more positive values for Ni–Al₁₀ provides a wider operational voltage window; the Mn, Co, Zn-substituted materials are subjected to OER for longer duration compared to the Al material (Fig. S6, ESI†). The drift

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towards higher voltage will ultimately affect the long-term cycling of the cell. Substituents within α -Ni(OH)₂ therefore need to be considered based on their influence on electrochemical charge storage, stability, and effect on redox and OER voltages. The ideal combination is a higher Ni^{2+/3+} redox voltage (to increase the cell voltage *vs.* Zn) while simultaneously shifting OER to a higher voltage. Of the two effects, as desirable is the former, the latter is the more important.

Evaluation of aluminum substitution on stabilization of the α -Ni(OH)₂ **phase.** With the highest capacity of the series, cycling stability, and most positive overvoltage for OER, Al³⁺-substituted Ni(OH)₂ offers promise as an active cathode material over native α -Ni(OH)₂. The increased capacity of Ni–Al₁₀ occurs despite the 12% lower amount of Ni (41 wt%) compared to α -Ni(OH)₂ (48 wt%) indicating that redox-inactive Al³⁺ influences the extent of oxidation of the active Ni sites. The XRD, Raman, and



Fig. 7 Effects of hydrothermal treatment on structure of unsubstituted α -Ni(OH)₂ and aluminum-substituted (Ni–Al₁₀) nanosheets; (a) X-ray diffraction powder pattern of α -Ni(OH)₂, α -Ni(OH)₂ after hydrothermal treatment (α -Ni(OH)₂-AHT), Ni–Al₁₀, Ni–Al₁₀ after hydrothermal treatment (Ni–Al₁₀-AHT), and commercial β -Ni(OH)₂ powder (Alfa); (b) Raman spectra in the 3500–3800 cm⁻¹ range of α -Ni(OH)₂, α -Ni(OH)₂-AHT, Ni–Al₁₀, Ni–Al₁₀-AHT, and commercial β -Ni(OH)₂ powder.

electron microscopic characterization of Ni–Al₁₀ provides evidence to support the increased discharge capacity. Of the substituents, Al^{3+} is the only one that creates larger interlayer spacing, a higher concentration of ordered free nitrate between the layers, shorter intralayer Ni–O bond length, and denser packing of nanosheets within the agglomerate. This distinct physical and charge local environment for Ni–Al₁₀ correlates with the observed ability to more deeply oxidize the Ni centres.

We note that interlayer free nitrate could play at least two roles: (i) pillar the layers making it energetically unfavourable to collapse the interlayers to form β -Ni(OH)₂, and (ii) buffer the electrostatic burden near oxidized Ni sites allowing extra electron charge to be stored and released. Prior studies report that the charge, size, and structure of interlayer anions can influence the oxidation state, discharge capacity, proton transport, and cycle stability of substituted nickel hydroxides.^{14,16,26,94,95}

To evaluate whether Al-substitution stabilizes the α -Ni(OH)₂ phase, we intentionally treated Ni-Al₁₀ and α-Ni(OH)₂ at 180 °C for 2 h in 0.01 M NaOH, conditions previously shown to convert α- to β-Ni(OH)₂.⁵⁵ After hydrothermal treatment, α-Ni(OH)₂ retains the nanosheet morphology (Fig. S7a, ESI[†]). The XRD (Fig. 7a) and Raman (Fig. 7b) data indicate that hydrothermally treated α -Ni(OH)₂ (notated as α -Ni(OH)₂-AHT) fully converts to the β -phase. In comparison, the XRD pattern of hydrothermal treated Ni-Al10 (Ni-Al10-AHT) retains the major peaks associated with the α -phase (Fig. 7a); however, one additional peak at 38.65° 2θ , is consistent with the β -phase. Using the characteristic Raman hydroxyl stretching mode ν (-OH) in Ni(OH)₂ to track phase stability, we find that although some β -Ni(OH)₂ is present in Ni–Al₁₀-AHT, the α -phase predominates (Fig. 7b). The electrochemical characteristics of α -Ni(OH)₂-AHT are similar to α -Ni(OH)₂ substantiating that absent Al³⁺ substitution, α -Ni(OH)₂ converts to β -Ni(OH)₂ during formation cycling (Fig. S7b and c, ESI†).

Conclusions

Metal substituents (Al³⁺, Co²⁺, Mn²⁺, Zn²⁺) within microwavesynthesized α -Ni(OH)₂ nanosheets affect the structure, electrochemical discharge capacity, and phase stability upon electrochemical cycling. Although synthesized to substitute a nominal 10 at%, the metals incorporate within the structure to varying degrees (from 2–16 at%), related to differences in speciation in the reaction medium and rates of hydrolysis and condensation. The native nanosheet morphology is maintained upon substitution. The substituents, however, alter the size of the nanosheet aggregates, surface area, and pore volume.

A layered α -Ni(OH)₂ structure is obtained for all substituted materials, and the substituents alter the interlayer distance. Al³⁺ substitution increases the interlayer spacing, consistent with the incorporation of charge-compensating anions within the interlayer. For all substituted α -Ni(OH)₂ variants but Zn²⁺, which results in a two-phase material, the shift in the frequency of the Raman-active E_g lattice mode indicates the substituents change the local potential energy environment of intralayer Ni–O. Additional Raman peaks support substitution of Al^{3+} , Co^{2+} , and Mn^{2+} within the Ni(OH)₂ plane.

The metal-substituted α -Ni(OH)₂ cathodes *vs.* Zn-sponge anodes exhibit different behaviour upon galvanostatic charging and discharging. The substituents alter the redox and OER onset voltage, discharge capacity, and capacity retention upon cycling. In this series, Al³⁺ substitution shifts OER onset to the most positive voltage, thereby minimizing parasitic capacity losses. Cathodes prepared with Al³⁺-substituted α -Ni(OH)₂ nanosheets deliver a higher discharge capacity compared to divalent cation-substituted materials, and the higher capacity is retained upon cycling. In agreement with prior work, Al-substitution ensures retention of the α -phase after forced conversion treatment.

From this study we find that the nature of the interlayer plays an important role in the higher discharge capacity of Al^{3+} -substituted α -Ni(OH)₂. Different from the other substituents, Al substitution is accompanied by ordered free nitrate anions within the interlayer. The interlayer anions necessary to charge balance a trivalent substituent for divalent Ni stabilize the α -phase by preventing collapse of the layers and by lowering the electrostatic burden at the Ni sites. Both effects enable extra electron charge to be stored and released.

The combination of a nanosheet morphology and Al^{3+} substitution facilitates the development of higher capacity α -Ni(OH)₂ cathode materials. To realize energy dense, safe, rechargeable Ni–Zn batteries, we plan to improve cathode performance by exploring different concentrations of Al, the addition of a second substituent, the nature of the interlayer anions, deliberate pillaring of the layers, rethinking the electronic wiring within the cathode structure, and then understanding the effects of these variations on capacity, efficiency, and cycling stability.

Author contributions

C. P. R. and D. R. R. conceived the idea and designed the experiments. C. P. R., D. R. R., C. N. C., J. F. P., and J. W. L. directed the project. S. W. K. prepared the Ni(OH)₂ series using microwave synthesis and adapted the synthetic protocol to scale-up quantities of product as advised by D. R. R. N. L. S. replicated selected microwave syntheses. J. S. K., R. H. D., and S. W. K. developed the protocol used by S. W. K. to prepare powder-composite cathodes. B. J. H. fabricated the zinc sponges and designed and laser-cut the coin cells. R. H. D. developed the protocol for differential capacitance measurements. S. W. K. carried out all porosimetry, electrochemical, cycling, and Raman experiments. C. N. C. and S. W. K. performed the X-ray diffraction analyses. S. W. K. performed the scanning electron microscopy and energy-dispersive spectroscopy analyses. B. M. H. performed the transmission electron microscopy measurements and B. M. H. and R. M. S. interpreted the resulting data. S. W. K., C. P. R., and D. R. R. co-wrote the paper. All authors discussed the results, revised, and commented on the manuscript.

Conflicts of interest

There are no conflicts to declare.

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