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# Electrochemical reduction of nitrate to ammonia on ultra-stable amorphous Co-P electrocatalyst†

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Electrocatalytic reduction of nitrate (NO<sub>3</sub><sup>-</sup>) to ammonia (NH<sub>3</sub>) is garnering increasing interest due to its potential to reduce CO<sub>2</sub> emissions as a substitute for the Haber-Bosch process, while also mitigating  $NO_3$  pollution. However, it remains a challenge to achieve a current density exceeding 300 mA cm<sup>-2</sup> while maintaining the stability of catalysts. Additionally, the anodic oxygen evolution reaction, characterized by slow kinetics and high energy barriers, severely impedes the widespread adoption of  $NH_3$  formation from  $NO_3^-$  reduction. Therefore, in this study, we introduce amorphous phosphorusdoped cobalt catalysts (Co-P@NF) prepared via a facile electrodeposition process for efficient NO<sub>3</sub><sup>-</sup> reduction and hydrazine oxidation. The incorporation of phosphorus in Co-P@NF facilitates electron migration from phosphorus to cobalt, enhancing \*H provision for efficient hydrogenation of the intermediate  $*NO_2^-$ . This results in a current density of 2 A cm<sup>-2</sup> at -0.3 V, with a faradaic efficiency for  $NH_3$  of 91% in an electrolyte containing 1 M  $NO_3^-$ . Moreover, the Co-P@NF catalyst exhibits remarkable long-term stability, maintaining an NH<sub>3</sub> faradaic efficiency exceeding 90% and a current density of 799 mA cm<sup>-2</sup> after 82 hours of electrolysis. Furthermore, Co-P@NF displays high catalytic activity in promoting the rate-determining step of hydrazine oxidation, from \*N<sub>2</sub>H<sub>2</sub> to \*N<sub>2</sub>H. The incorporation of the HzOR (hydrazine oxidation reaction)-assisted NO<sub>3</sub>-RR (nitrate reduction reaction) unit significantly reduces the cell voltage to 0.34 V at 300 mA cm<sup>-2</sup>.

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# 1. Introduction

Ammonia (NH<sub>3</sub>) is a vital chemical utilized in various industrial applications, such as chemical production, fuel synthesis, and as a fertilizer. Approximately 50% of global food production depends on NH3 fertilizers.1 The industrial-scale production of NH<sub>3</sub> predominantly relies on the Haber-Bosch process, which operates at high temperatures (400-600 °C) and high pressures (200-350 atm).2 This production process is responsible for approximately 1.2% of annual global CO<sub>2</sub> emissions, surpassing emissions from other industrial chemical processes.3 To address decarbonization challenges in the ammonia sector, an alternative approach known as the dinitrogen reduction reaction has been developed [4]. However, the dinitrogen reduction reaction exhibits a significantly low reaction rate (<10 mmol g<sub>cat</sub><sup>-1</sup> h<sup>-1</sup>),<sup>2</sup> which is two to three orders of magnitude lower than that of the Haber-Bosch process. In marked contrast, nitrate anions (NO<sub>3</sub><sup>-</sup>) require only 204 kJ mol<sup>-1</sup> of energy to break the N=O bond, a significantly lower energy requirement compared to the N≡N bond in dinitrogen gas (941 kJ mol<sup>-1</sup>). Additionally, NO<sub>3</sub> anions exhibit excellent solubility in aqueous solutions.4 Consequently, the electrochemical reduction reaction of NO<sub>3</sub><sup>-</sup> (NO<sub>3</sub><sup>-</sup>RR) for NH<sub>3</sub> production has garnered increasing attention.

NO<sub>3</sub> pollution is a pressing environmental issue, resulting from agricultural runoff, wastewater discharge, and industrial processes. It poses risks to human health and ecosystems, making the development of effective NO<sub>3</sub><sup>-</sup> removal strategies critical. The electroreduction of NO<sub>3</sub><sup>-</sup> aims to produce ammonia, serving the dual purpose of recycling waste materials and mitigating the adverse environmental and ecological impacts of excessive NO<sub>3</sub><sup>-</sup> discharge.<sup>5</sup> Thanks to these advantages, various metal-based electrocatalysts, including Cu, Ru, Ag, Fe, Pd, and Co, have been developed to facilitate the reduction of NO<sub>3</sub><sup>-</sup> to NH<sub>3</sub>.<sup>2,6,7</sup> However, the reduction of NO<sub>3</sub><sup>-</sup> to NH<sub>3</sub> is a complex eight-electron reaction (eqn (1)), inevitably resulting in the formation of undesired byproducts such as nitrogen oxyanions and dinitrogen.8,9 Due to significant progress in electrocatalyst development, achieving high faradaic efficiencies towards NH<sub>3</sub> (FE<sub>NH<sub>3</sub></sub> > 90%) is feasible at pseudoampere-lever current densities. Nevertheless, the long-term stability of catalysts operating at currents exceeding 300 mA cm<sup>-2</sup> with FE<sub>NH<sub>2</sub></sub> > 90% remains challenging, falling short of the standards set by the U.S. Department of Energy. 10 Meanwhile, the anodic oxygen evolution reaction (OER,  $2OH^- \rightarrow H_2O + 1/V$  $2O_2 + 2e^-$ , 1.23 V vs. RHE), suffers from slow reaction kinetics and high energy barriers, necessitating an applied potential exceeding 1.6 V.11 The high voltage requires substantial energy expenditure thereby significantly restricting the widespread application of NH<sub>3</sub> formation from NO<sub>3</sub> reduction. Substituting sluggish reactions with thermodynamically favorable ones, such as hydrazine oxidation reaction (HzOR, N<sub>2</sub>H<sub>4</sub> +  $4OH^- \rightarrow N_2 + 4H_2O + 4e^-$ ,  $-0.33~V~\nu s.~RHE$ ), to assist  $NO_3^-RR$ could greatly lower the electricity consumption for NH3 formation. Moreover, the employment of HzOR-assisted NO<sub>3</sub> RR offers environmental benefits, converting toxic, carcinogenic N<sub>2</sub>H<sub>4</sub> in wastewater into dinitrogen and water.

Developing bifunctional electrocatalysts toward anodic HzOR and cathodic NO<sub>3</sub><sup>-</sup>RR with high activity and durability is pivotal to promote the practical applications.

$$NO_3^- + 6H_2O + 8e^- \rightarrow NH_3 + 9OH^-,$$
  
 $pH = 14, E^0 = 0.69 \text{ V vs. RHE}$  (1)

NO<sub>3</sub><sup>-</sup>RR is a typical hydrodeoxygenation reaction. Electrocatalysts, like the first-row transition metal phosphides, with good performance in hydrogen evolution and hydrogenation reactions have been reported to have good NO<sub>3</sub><sup>-</sup>RR catalytic activities.2,6 Javier Vela et al.12 found that over 96% selectivity of NH<sub>3</sub> from NO<sub>3</sub><sup>-</sup> reduction was achieved on Ni<sub>2</sub>P, which was previously reported as a good electrocatalyst for hydrogen evolution reaction (HER).13 Yang et al.,14 found that the Co nanocrystals that could generate plenty of \*H had an exceptional performance of electrochemical NO<sub>3</sub><sup>-</sup>RR to NH<sub>3</sub>. Li et al.15 also proved that the generated \*H with moderate adsorption energy on the catalyst can behave as important reactive species for the hydrogenation of reaction intermediates to NH<sub>3</sub> and suppress the competing HER. Notedly, the good modulation of the adsorption energy of \*H and \*NO3 on the catalyst surface is critical for the improvement of the catalytic activity and product selectivity.16 Meanwhile, a synergistic benefit was also proposed by Simpson and Johnson for the CuNi alloys: the Ni sites acted as the active sites for the adsorption of \*H, and the Cu sites facilitated the adsorption of NO<sub>3</sub><sup>-</sup> intermediates on these binary alloy electrodes.17

On the other hand, amorphous materials with long-range disordered structures have exhibited superior electrocatalytic performance and durability compared to their crystalline counterparts. 18,19 The coordination of surface atoms on amorphous materials is highly unsaturated. Also, amorphous alloy has a high degree of structure flexibility, both of which benefit catalysis. Due to the advanced properties, the amorphous-structure catalysts were widely used in oxygen reduction,20 carbon dioxide reduction,21 and hydrogen evolution reaction.22 However, to our best knowledge, few literature reports exist on the amorphous catalysts for the NO<sub>3</sub><sup>-</sup>RR process. Inspired by the exceptional performance of the Co-based catalyst for HER and hydrodeoxygenation reaction,23 the amorphous Co-P was prepared via a facile one-step electrodeposition procedure for HzOR-assisted NO<sub>3</sub><sup>-</sup>RR. The as-prepared amorphous Co-P exhibited superior catalytic performances, which is one of the most advanced NO<sub>3</sub><sup>-</sup>RR catalysts so far, as (1) a high partial current density for  $NH_3$  formation  $(j_{NH_3})$  was observed at 2 A cm<sup>-2</sup>, along with a faradaic efficiency ( $FE_{NH_2}$ ) of 91% at -0.3 V versus RHE. The corresponding yield of  $NH_3$  production was 2.9 mol  $g_{cat}^{-1} h^{-1}$ , which is almost 11-fold higher than that achieved by the Haber-Bosch process (200 mmol g<sub>cat</sub><sup>-1</sup> h<sup>-1</sup>); (2) a good stability for NH<sub>3</sub> formation with FE<sub>NH<sub>3</sub></sub> over 90% at current density around 800 mA cm<sup>-2</sup> for more than 82 h electrolysis; (3) the cell voltage of HzOR-assisted NO<sub>3</sub><sup>-</sup>RR at 300 mA cm<sup>-2</sup> over Co-P@NF||Co-P@NF was only 0.34 V. Electrochemical in situ shell-isolated nanoparticle enhanced Raman spectroscopy (SHINERS) was conducted to clarify the pathways and mechanisms of NO<sub>3</sub><sup>-</sup>RR.

### 2. Materials and methods

#### 2.1. Materials

Paper

Nickel foam (NF) used as electrode substrate was supplied from Suzhou JiaShide Foam Metal Co., Ltd. Trisodium citrate dihydrate ( $C_6H_5Na_3O_7\cdot 2H_2O$ ) and sodium hypophosphite (NaH<sub>2</sub>-PO<sub>2</sub>·H<sub>2</sub>O) were obtained from Meryer Chemical Technology Co., Ltd. CoCl<sub>2</sub>· $6H_2O$ , KOH and (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> were purchased from Sinopharm Chemical Reagent Co., Ltd. NaNO<sub>3</sub> and NaNO<sub>2</sub> were bought from Aladdin Co., Ltd. All reagents were analytical grade without any further purification and all solution were prepared by Milii-Q water (MQ).

#### 2.2. Synthesis of Co@Ni foam and Co-P@Ni foam

Phosphorus doped cobalt on the Ni foam substrate (Co–P@NF) was prepared by a simple constant-potential electrodeposition method in a three-electrode system, where Ni foam, Pt foil, and saturated calomel electrode (SCE) were applied as the working electrode, counter electrode and reference electrode, respectively. Before the electrodeposition process, Ni foam (0.5  $\times$  1.0 cm²) was ultrasonically cleaned in acetone solution, ethanol solution and deionized (DI) water several times. An aqueous solution (55 mL) containing 0.2 M  $\rm C_6H_5Na_3O_7, 0.5~M~(NH_4)_2SO_4, 0.6~M~NaH_2PO_2$  and 0.1 M  $\rm CoCl_2\cdot 6H_2O$  were used as the electrolyte. Co–P was deposited on the Ni foam at  $-2.0~\rm V~\nu s$ . SCE in the above electrolyte saturated with N2 at room temperature for 45 min. The preparation of Co@NF followed the same process but without the addition of NaH2PO2.

#### 2.3. Assembly of Zn-nitrate flow battery

The Zn-nitrate flow battery was assembled in a membrane electrode assembly flow reactor wherein Co-P@NF (2 cm  $\times$  2 cm) was used as both the anode and cathode, respectively. The anodic and cathodic chambers were separated by a Nafion N115 proton exchange membrane. Bipolar plates were highpurity titanium plates with a single serpentine flow field and guard plates were Perspex sheets. The catholyte was 1 M NaOH/0.1 M KNO<sub>3</sub> while the anolyte was 6 M NaOH, and their flow rates were controlled to be 150 mL min<sup>-1</sup> by a peristaltic pump. Chronopotentiometry was recorded to plot the polarization curves. The power density (P) was calculated by the following equation:  $P = j \times E$ , where j is the current density and E is the cell voltage.

#### 2.4. Electrochemical measurements

The electrocatalytic performances of Co@NF and Co-P@NF were investigated in a three-electrode system connected to a DH7000C electrochemical workstation (Donghua, Jiangsu) in an H-type cell with a Nafion 115 membrane as a separator. The Co@NF and Co-P@NF with the same size  $(0.5 \times 0.5 \text{ cm}^2)$  were used as working electrodes, while a platinum foil and a saturated calomel electrode were used as the counter electrode and reference electrode. 1 M KOH aqueous solution (30 mL) containing different NaNO<sub>3</sub> concentrations (*i.e.*, 10, 20, 50, 100 and 1000 mM) and 1.0 M N<sub>2</sub>H<sub>4</sub> were used as the cathodic and anodic

electrolyte, respectively. The electrochemical linear voltammetry (LSV) analysis was carried out in a single undivided electrochemical cell. The Tafel slope was obtained by fitting the liner part of the Tafel plots according to the Tafel equation ( $\eta = a + b \times \log(j)$ ). The electrochemical impedance measurement was performed in a frequency range of 0.01 Hz to 1000 Hz with an amplitude of 5 mV. The double layer capacitance ( $C_{\rm cll}$ ) of the obtained samples was calculated from CV curves at different scan rates of 10–100 mV s<sup>-1</sup> in the non-Faraday area. The current density was normalized to the geometric electrode area (0.25 cm²) unless otherwise specified. All recorded electrode potentials  $\nu$ s. Hg/HgO were transformed to potentials against the reversible hydrogen electrode (RHE) according to the calibration equation expressed as  $E_{\rm RHE} = E_{\rm Hg/HgO} + 0.098 + 0.0591 \times \rm pH$ .

## Results and discussions

#### 3.1. Catalyst composition and microstructure

The Co-P@NF exhibits a much rougher surface than Co@NF (Fig. 1a and b), as indicated by the irregular nanograins formed on the surface of Co-P@NF (Fig. 1c), compared to the flat surface of Co@NF (Fig. 1d). As shown in the high-resolution transmission electron microscopy (HRTEM) images, the nanograins on Co-P@NF were amorphous (Fig. 1e), while the lattice fringe of Co@NF was observed (Fig. 1f). There was also no diffraction pattern for Co-P@NF, demonstrating the amorphous feature of this catalyst (Fig. S1†). The lattice distance of 1.91 Å was well ascribed to the (101) plane of metallic Co. The element mapping analysis (Fig. 1g) indicated that Co and P were evenly dispersed in the nanograins of Co-P@NF. The different structure morphologies of Co-P@NF and Co@NF were also confirmed by the XRD pattern (Fig. 1h). There was no crystalline peak observed in the XRD pattern of Co-P@NF, except the two peaks belonging to the Ni substrate. In comparison, two small peaks at 41.6° and 44.5° were observed in the XRD pattern of Co@NF, which were assigned to Co(100) and Co(101). The elemental compositions and electronic structure of Co-P@NF were determined using XPS (Fig. 1i and j). For Co-P@NF, P 2p showed a doublet with a binding energy of 128.3 (P 2p3/2) and 130.2 eV (P 2p<sub>1/2</sub>), assigned to elemental P (P<sup>0</sup>) and oxidized P  $(P^{5+})$ , while Co 2p depicted two peaks at 781.8 eV (Co  $2p_{3/2}$ ) and 797.6 eV (Co  $2p_{1/2}$ ) assigned to oxidation of Co resulting from the exposition of the catalyst to air. There was no P 2p peaks found in the Co@NF spectra. It was well-fitted with the results from the HRTEM-EDS analysis. There were about 13 at% P in Co-P nanograins (Fig. S2 and Table S1†). Compared to Co-P@NF, the Co  $2p_{3/2}$  and  $2p_{1/2}$  peaks of Co@NF exhibited a small shift towards higher binding energies (Fig. 1i). This shift indicated that the introduction of P promotes electron migration from the P to the Co, which makes Co electron-rich and becomes more favorable for the NO<sub>3</sub><sup>-</sup>RR processes. The electron-rich property of Co-P@NF was also demonstrated by EIS analysis (Fig. S3†). Thanks to the electron transfer from P to Co, the impedance of Co-P@NF (0.32  $\Omega$ ) was lower than that of the Co@NF  $(0.52 \Omega)$ .

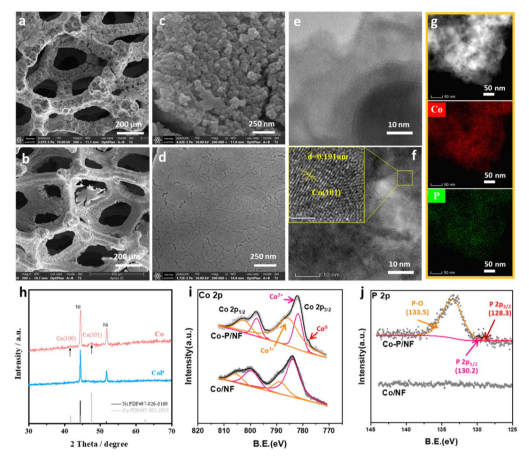


Fig. 1 SEM imagines of Co-P@NF (a and c) and Co@NF (b and d). HR-TEM imagines of Co-P@NF (e) and Co@NF (f). TEM-EDS mapping of Co-P@NF (g). (h) XRD pattern of Co-P@NF and Co@NF. XPS spectra: Co 2p (i) and P 2p (j) of Co-P@NF and Co@NF.

#### 3.2. Electrocatalytic performances of NO<sub>3</sub><sup>-</sup>RR

The linear sweeping voltammetry was firstly applied to investigate the catalytic activity over Co-P@NF and Co@NF during NO<sub>3</sub> RR (Fig. 2a). In the absence of NO<sub>3</sub>, Co-P@NF has a smaller onset potential ( $\eta_{HER}$ ) and a much higher current density  $(j_{HER})$  of hydrogen evolution reaction than that of Co@NF ( $\eta_{\text{HER}}$ : -0.05 V vs. -0.15 V;  $j_{\text{HER}}$  at -0.1 V: 69 mA cm<sup>-2</sup> vs. 12 mA cm $^{-2}$ ), indicating the superior catalytic activity of HER on Co-P@NF. The Tafel slop of HER over Co-P@NF (136 mV dec<sup>-1</sup>) was also smaller than that of Co@NF (164 mV dec<sup>-1</sup>) (Fig. 2b), demonstrating that Co-P@NF had a faster kinetic for HER and provided more \*H for NO<sub>3</sub> reduction, according to Volmer step  $(H_2O + e^- \rightarrow *H + OH^-)$ . In the presence of  $NO_3^-$ , increased current densities (j) were observed on both Co-based electrodes, since Co was a good candidate for NO<sub>3</sub><sup>-</sup>RR. The larger j on Co-P@NF indicated the enhanced catalytic activity of NO<sub>3</sub><sup>-</sup> reduction after the incorporation of P into Co. Two current reduction peaks were appearing after the addition of  $NO_3^-$ , 0.09 V and -0.03 V on the curve of Co-P@NF, and 0.01 V and -0.15 V on the curve of Co. According to previous studies,24,25 the first peak was assigned to NO<sub>3</sub> reduction into \*NO<sub>2</sub> following a 2-electrons transfer process (eqn (2)), while the second peak was allocated to the \*NO<sub>2</sub><sup>-</sup> reduction into NH<sub>3</sub> following a 6-electrons transfer process (eqn (3)).

$$NO_3^- + H_2O + 2e^- \rightarrow *NO_2^- + 2OH^-$$
 (2)

$$*NO_2^- + 5H_2O + 6e^- \rightarrow NH_3 + 7OH^-$$
 (3)

Co-P@NF displayed positive shifts of 80 mV for the  $NO_3^- \rightarrow$  $NO_2^-$  peak and 120 mV for the  $NO_2^- \rightarrow NH_3$  peak, compared to Co@NF, suggesting a drop of the energy barrier of \*NO<sub>3</sub><sup>-</sup> reduction to NH<sub>3</sub> on Co-P@NF. According to the Tafel slop in the NO<sub>3</sub><sup>-</sup>-containing electrolyte, Co-P@NF (32 mV dec<sup>-1</sup>) had a higher kinetic activity for NO<sub>3</sub><sup>-</sup>RR than that of Co (160 mV dec<sup>-1</sup>). EIS results also revealed the incorporation of P into Co would decrease the electron transfer resistance that was beneficial to facilitate NO<sub>3</sub><sup>-</sup>RR process. The constant-potential electrolysis was performed and the partial current density of  $NH_3$  formation  $(j_{NH_3})$  at different applied potential was obtained (Fig. 2c). The concentration of NH<sub>3</sub> and NO<sub>2</sub><sup>-</sup> formed was detected by colorimetric methods using a UV-vis absorbance spectrophotometer (DR3900) and ion chromatography (ICS-600), respectively (Text S1, Text S2, Fig. S4 and S5†). The substrate of NF was relatively inactive for NO<sub>3</sub><sup>-</sup>RR (Fig. S6†), since the  $j_{NH_3}$  on Ni foam was only 7 mA cm<sup>-2</sup>, compared with the electrodeposited Co-based catalysts (1296 mA cm<sup>-2</sup> on Co-P@NF and 212 mA cm<sup>-2</sup> on Co@NF). Significantly higher  $j_{NH_0}$ was obtained with Co-P@NF, surpassing 1 A cm<sup>-2</sup> at potential

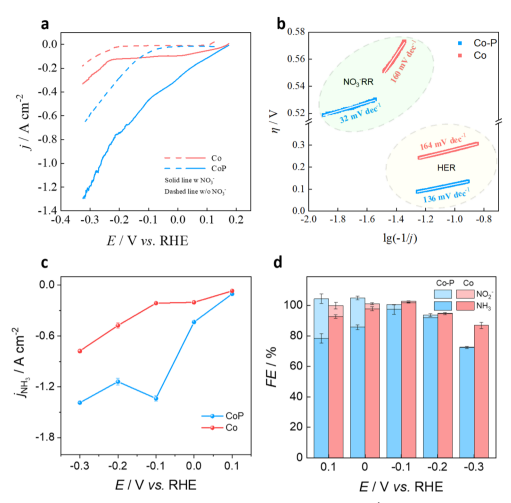


Fig. 2 (a) Linear sweeping voltammetry over Co@NF and Co-P@NF in the electrolyte of 1 mol  $L^{-1}$  KOH with or without 100 mmol  $L^{-1}$  NO $_3^{-}$ . (b) Tafel slop of Co@NF and Co-P@NF in the presence of NO<sub>3</sub><sup>-</sup>. (c) Partial current density of NH<sub>3</sub> production over Co@NF and Co-P@NF. (d) FE of  $NH_3$  and  $NO_2$  over Co-P@NF and Co@NF in the potential range of 0.1--0.3 V.

below -0.1 V vs. RHE. At this potential, the highest faradaic efficiency for NH<sub>3</sub> formation (FE<sub>NH.</sub>, 97.5%) was achieved with Co-P@NF (Fig. 2d). Consequently, the Co-P@NF electrode shows superior performance with high  $FE_{NH_3}$  and  $j_{NH_3}$  for  $NO_3^$ electroreduction to NH3 as compared to other reported electrodes, as exhibited in Table S2.† The low FE<sub>NH</sub>, over Co-P@NF at the potential of 0.1 V and 0 V, where the HER did not occur and the \*H coverage ( $\theta_{\rm H}$ ) was too low to further hydrogenation of  $NO_2^-$  to  $NH_3$ . Until the potential negatively shifted to -0.1 V, the  $\theta_{\rm H}$  increased, based on eqn (4), and therefore the maximum FE<sub>NH</sub>, was obtained. As potential continued to decreasing, the FE<sub>NO</sub> significantly declined but the slight decreased FE<sub>NH</sub> was also observed due to the strong HER competition on the Co-P@NF (Fig. 2d). The competitive HER also led to a decrease in  $j_{\rm NH_3}$  from 1336 mA cm<sup>-2</sup> to 1138 mA cm<sup>-2</sup>, when potential was negatively shifted from -0.1 V to -0.2 V. The electrochemically active surface area (ECSA) was also measured (Fig. S7a-c†), revealing that Co@NF and Co-P@NF had comparable ECSA. However, a higher ECSA-normalized current density for NH<sub>3</sub> production  $(j_{NH_2(ECSA)})$  was observed with Co-P@NF (Fig. S7d†), indicating that this catalyst had higher intrinsic activity for NH3

production. To clarify the possible N pollution caused by nitrogen reduction reaction, the electrolysis in the electrolyte free of NO<sub>3</sub><sup>-</sup> ions was performed and little NH<sub>3</sub> was produced (Fig. S8 $\dagger$ ).  $j_{NH_3}$  in the solution with NO<sub>3</sub> ions was over 133-fold higher than that in the electrolyte free of NO<sub>3</sub>, where the FE<sub>NH</sub>, was only about 1%. In addition, the typical two peaks of <sup>15</sup>NH<sub>4</sub> after the electrolysis of 15NO<sub>3</sub> also suggested that the NH<sub>3</sub> product indeed came from the electrocatalytic reduction of  $NO_3^-$  (Fig. S9†).

$$\eta = E - E_{\rm eq} = \frac{RT}{F} \ln \frac{\theta_{\rm H}}{\theta_{\rm H}^0} \tag{4}$$

where  $\theta_{\rm H}$  and  $\theta_{\rm H}^0$  are the \*H coverage at E and E°, respectively; F is the Faraday constant, 96 485C  $\text{mol}^{-1}$ ; R is the gas constant, 8.314 J mol<sup>-1</sup> K<sup>-1</sup>; T is the kelvin temperature, K.

It should be noted that the sum FE of NO<sub>2</sub><sup>-</sup> and NH<sub>3</sub> formation was over 100% (Fig. 2d). The NO<sub>3</sub><sup>-</sup>RR over Co-P@NF and Co@NF were then performed at open circle potential (OCP) and the NO<sub>2</sub> and NH<sub>3</sub> were both detected in the electrolyte (Fig. 3a and S10†), indicating that there was a spontaneous reduction of NO<sub>3</sub><sup>-</sup> to NO<sub>2</sub><sup>-</sup> and NH<sub>3</sub> on Co-P@NF and Co@NF.

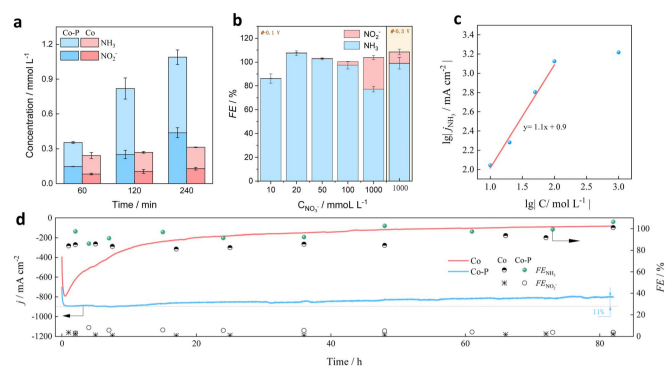


Fig. 3 (a) Concentration of  $NO_2^-$  and  $NH_3$  over Co-P@NF and Co@NF at open circle potential. (b) FE of  $NO_2^-$  and  $NH_3$  and (c) lg(-j) vs. lg C plots for Co-P@NF were calculated at -0.1 V vs. RHE. (d) Long-term stability of Co-P@NF in the electrolyte of 1 mol L<sup>-1</sup> with 100 mmol L<sup>-1</sup> NO<sub>3</sub><sup>-</sup> at

This spontaneous reduction process may explain the over-100% FE. A similar phenomenon was reported by Han and his colleagues:26 oxidation of Co to Co(OH)2 by NO3 (eqn (5)) and the electroreduction of Co(OH)<sub>2</sub> to Co (eqn (6)) proceed simultaneously during the NO<sub>3</sub><sup>-</sup>RR (eqn (2) and (3)), and a dynamic Co valence cycle was achieved. The OCP over Co-P@NF was obviously lower than that of Co@NF (Fig. S11†), indicating that Co-P@NF had a superior ability of the spontaneous reduction of NO<sub>3</sub>-, thanks to the electron-rich of Co atom in Co-P@NF, giving that the NO2 and NH3 formation on Co-P@NF was approximate 3-times higher than that of Co@NF at OCP.

$$NO_3^- + Co + H_2O \rightarrow *NO_2^- + Co(OH)_2$$
 (5)

$$Co(OH)_2 + 2e^- \rightarrow Co + 2OH^-$$
 (6)

The concentration of NO<sub>3</sub><sup>-</sup> in real wastewater can vary from 1 mmol L<sup>-1</sup> to 2000 mmol L<sup>-1</sup>.<sup>27</sup> Therefore, NO<sub>3</sub><sup>-</sup>RR on Co-P@NF was performed in a wide NO<sub>3</sub><sup>-</sup> concentration range. The  $FE_{NH_2}$  was maintained above 95% in the range of 20 mmol  $L^{-1}$ to 100 mmol  $L^{-1}$  at -0.1 V (Fig. 3b). The low  $FE_{NH_3}$  in the solution of 10 mmol  $L^{-1}$  and 1000 mmol  $L^{-1}$  was due to the competitive adsorption between \*H and \*NO3. For example, in the case of 10 mmol L<sup>-1</sup>, NO<sub>3</sub> concentration was so low that catalytic sites were occupied by \*H which would be formed into H<sub>2</sub> via Volmer-Tafel or Volmer-Heyrovsky mechanism and decrease the FE<sub>NH<sub>3</sub></sub>. When the NO<sub>3</sub><sup>-</sup> concentration increased, the HER would be gradually hindered so that the  $FE_{NO_2}$ increased. FE<sub>NO<sub>2</sub></sub>- was about 27% in the case of 1000 mmol L

NO<sub>3</sub><sup>-</sup>, where \*H was insufficient for \*NO<sub>2</sub><sup>-</sup> reduction (eqn (3)), leading to the accumulation of \*NO2 on the catalytic surface. A quasi-first-order reaction relationship between j and the NO<sub>3</sub> concentration was obtained (Fig. 3c), revealing that the ratedetermining step of  $NO_3$  RR ( $RDS_{NO_3}$  RR) was  $NO_3$   $\rightarrow$   $NO_2$ step. Therefore, supplement of sufficient \*H was significant for efficient reduction of  $NO_2^-$ . When a potential of -0.3 V was applied in the solution of 1000 mmol L<sup>-1</sup>, an increase in FE<sub>NH</sub>. and a decrease in FE<sub>NO</sub>, were observed, respectively. This is attributed to a higher  $\theta_H$  (according to eqn (4)) and a stronger affinity of \*H at more negative potentials,28 which promotes further hydrogenation of  $NO_2^-$  to  $NH_3$ . Meanwhile,  $j_{NH_3}$ increased to 2 A cm  $^{-2}$  with the FE $_{\mathrm{NH_3}}$  as 91% (Fig. S12†). The corresponding NH<sub>3</sub> yield rate was 2.9 mol g<sub>cat</sub><sup>-1</sup> h<sup>-1</sup> that was almost 11-fold higher than that of Haber-Bosch route (200 mmol  $g_{cat}^{-1} h^{-1}$ ). Long-term stability tests at  $-0.1 \text{ V} \nu s$ . RHE assessed the durability of Co-P@NF against Co@NF (Fig. 3d). For Co-P@NF, FE<sub>NH</sub>, (>90%) remained stable after the 82 h electrolysis. The  $j_{NH_2}$  experienced a slight decrease from 890 mA cm<sup>-2</sup> to 799 mA cm<sup>-2</sup> (about 89% of the initial  $j_{NH}$ ), which still exceeded U.S. Department of Energy's standard requirement by more than two fold (300 mA cm $^{-2}$ ). For Co@NF, the  $j_{\rm NH_3}$  quickly decayed from 798 mA cm $^{-2}$  to 198 mA cm $^{-2}$ after 20 h electrolysis and its FENH, was also lower than that of Co-P@NF. To further check the stability of the Co-P@NF catalyst, the cyclic electrolysis was also conducted for NO<sub>3</sub> reduction. The results demonstrated that the faradaic efficiency of NH<sub>3</sub> formation remained consistently at 100% throughout the 11 electrolysis cycles. Moreover, the yield of NH<sub>3</sub> formation

of the 11th was still two-times higher than that of Haber-Bosch route (200 mmol  $g_{cat}^{-1} h^{-1}$ ) (Fig. S13†). Above stability test results indicated the amorphous-structured Co@P indeed had a satisfactory durability during NO<sub>3</sub><sup>-</sup>RR. According to the TEM analysis, the morphology of Co-P was well kept after the consecutive recycling tests (Fig. S14†). Furthermore, XRD (Fig. S15†) and XPS (Fig. S16†) analysis was also performed on the samples after the consecutive recycling tests, and the results demonstrated negligible changes in the chemical compositions and oxidation states. The excellent stability of the catalysts was possibly ascribed to amorphous feature of the Co-P catalysts.

Tert-butanol (TBA), used as a specific \*H radical quenching reagent, was added during the process to evaluate the role of active \*H (Fig. 4a). Upon the addition of TBA, the j decreased sharply from 546 mA cm<sup>-2</sup> to 373 mA cm<sup>-2</sup> on Co@NF and from 886 mA cm<sup>-2</sup> to 628 mA cm<sup>-2</sup> on Co-P@NF. Meanwhile, the FENH, also decreased from 96% to 88% on Co@NF and from 106% to 100% on Co-P@NF. The quenching test results demonstrated that the \*H radicals truly improved the reaction rate and FE<sub>NH</sub>. We further used dimethyl-1-pyrroline-N-oxide

(DMPO) as the radical trapping reagent to monitor the formation of \*H radicals with and without using NO<sub>3</sub><sup>-</sup> (Text S3). As shown in Fig. 4b, the ESR spectra display typical 9 signals with intensity ratios of 1:1:2:1:2:1:2:1:1 without adding  $NO_3$ in the catholyte, which confirms the formation of DMPO-H. When NO<sub>3</sub> was added, the typical 9 signals nearly disappeared, indicating that the produced \*H is consumed by the intermediates in the NO<sub>3</sub> RR process. These results strongly prove the key role of the \*H radicals in NO<sub>3</sub> reduction into NH<sub>3</sub>. The electrochemical in situ Raman spectra of Co-P and Co catalysts depicted in Fig. 4c and d, were recorded across a potential window of -0.7--1.13 V versus the RHE with increments of 50 mV. Three distinct peaks, corresponding to  $A_{1g}$ ,  $E_g$  and  $F_{2g}$  vibrations models of Co-O band, were visible on Co-P and Co catalysts at around 686.9, 480.5 and 524.9 cm<sup>-1</sup>.29 The formation of Co-O band was ascribed to the oxygen vacancies over the two catalysts. The coordination of surface atoms on amorphous materials is highly unsaturated.30 According to the electron paramagnetic resonance (EPR) analysis (Fig. S17†), the Co-P@NF had a much higher density of

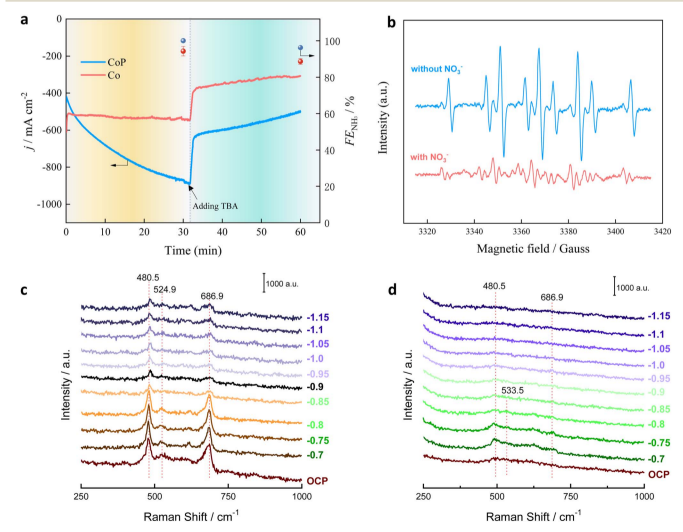


Fig. 4 (a) j and  $FE_{NH_3}$  over Co-P@NF and Co@NF at potential of -0.1 V vs. RHE. (b) The DMPO-H\* obtained on Co-P@NF after 5 min of electrocatalysis at -0.1 V vs. RHE in 1 M KOH solution with  $0.1 \text{ M NO}_3^-$  (bottom) or (b) without  $NO_3^-$  (top). In situ electrochemical–Raman on Co-P (c) and Co (d) in the potential of -0.7 V–-1.15 V vs. Hg/HgO.

oxygen vacancies. The intensity of Co–O bands on Co–P was much higher than that Co. The Raman spectra were well fitting with the EPR results that the Co–P with amorphous structure had a higher density of oxygen vacancies. As the potential negatively shifted, the Co–O bands gradually decreased. For Co–P catalyst, the Co–O band could be preserved, revealing that the amorphous structure had more stable oxygen vacancies; while no signal could be observed on Co catalyst after the potential lower than –0.9 V. The oxygen vacancies can be used as the adsorption site for the charge transfer, which is conducive to the adsorption/desorption of the reaction intermediates speeding up the NO<sub>3</sub><sup>-</sup>RR.<sup>31</sup>

#### 3.3. Electrocatalytic performances of HzOR

The HzOR electrochemical performance of Co–P@NF was then evaluated in 1.0 M KOH + 1 M  $N_2H_4$  electrolyte, compared to Co@NF and NF. Co–P@NF showed a higher HzOR catalytic activity with small overpotentials of -67, -16 and 486 mV to reach 10, 100 and 1000 mA cm<sup>-2</sup> (Fig. 5a), respectively, compared to Co@NF and NF. The smallest Tafel slope of 24.2 mV dec<sup>-1</sup> was obtained in Co–P@NF, indicating the favorable catalytic oxidation kinetics toward HzOR (Fig. 5b). This ultra-low Tafel slope also indicated that the rate-determining step over Co–P@NF was the dehydrogenation

from \*N2H2 to \*N2H. In addition, a comparison between the OER and HzOR performances over Co-P@NF was performed (Fig. 5c and d). Much lower potentials of 16 and 207 mV were needed in HzOR than those in N2H4-free electrolyte (1.66 and 1.95 V) to reach the current densities of 100 and 500 mA cm<sup>-2</sup>, respectively, indicating the potential application to replace OER for HzOR-assisted NO<sub>3</sub><sup>-</sup>RR. Therefore, thanks to the excellent NO<sub>3</sub><sup>-</sup>RR and HzOR catalytic performances on Co-P@NF, an HzOR-assisted NO<sub>3</sub> RR unit with two-electrode system was assembled by employing Co-P@NF as both anode and cathode. As shown in Fig. 5e, the HzOR-assisted NO<sub>3</sub> RR unit exhibits significantly enhanced catalytic activity compared to the stand alone NO<sub>3</sub>-RR unit, only requiring voltages of 93, 308, and 516 mV to achieve current densities of 100, 300, and 500 mA cm<sup>-2</sup>, respectively. These values are 1.45, 1.53, and 1.67 V lower than those required by the OER-assisted NO<sub>3</sub><sup>-</sup>RR system. The cell voltage of Co-P@NF||Co-P@NF at 300 mA cm<sup>-2</sup> dramatically dropped from ca. 2.0 V to ca. 0.34 V after addition of N<sub>2</sub>H<sub>4</sub> in the electrolyte containing NO<sub>3</sub><sup>-</sup> ions (Fig. 5f). For comparison, NF as electrodes for the HzOR-assisted NO<sub>3</sub> RR unit was also assembled (Fig. S18†). For a current density of 300 mA cm<sup>-2</sup>, and the cell voltage of NF||NF (2.40 V) are much higher than that of Co-P@NF||Co-P@NF (0.34 V), indicating the greatly reduced electricity consumption over Co-P catalysts.

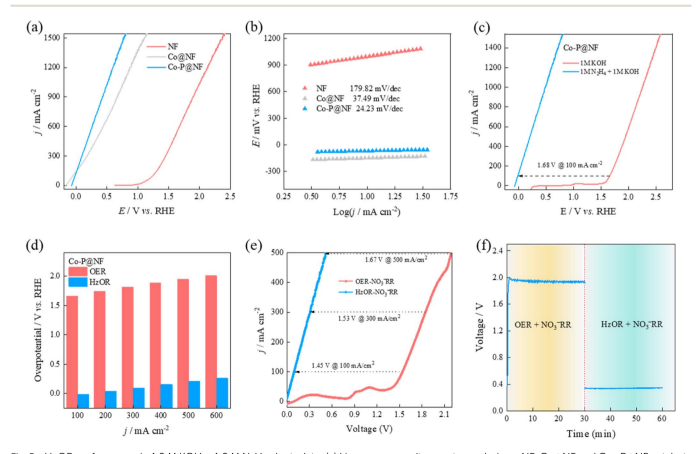


Fig. 5 HzOR performance in 1.0 M KOH + 1.0 M  $N_2H_4$  electrolyte. (a) Linear sweep voltammetry analysis on NF, Co@NF and Co-P@NF catalysts and the corresponding (b) Tafel slopes. (c) Compared LSV curves of Co-P@NF between OER (1.0 M KOH) and HzOR (1.0 M KOH and 1.0 M  $N_2H_4$ ), and (d) the corresponding histogram catalytic performance comparison. (e) Comparison LSV curves of Co-P@NF between OER-NO $_3$ -RR and HzOR-NO $_3$ -RR. (f) Cell voltage of Co-P@NF at 300 mA cm<sup>-2</sup> in OER-NO $_3$ -RR and HzOR-NO $_3$ -RR system.

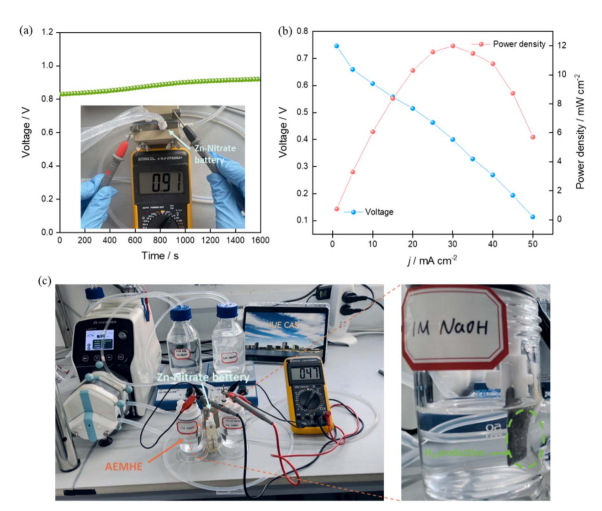


Fig. 6 Zn-nitrate flow battery application. (a) The open circuit voltage. The cathodic and anodic electrolytes used in Zn-nitrate flow battery were 0.1 M NO<sub>3</sub><sup>-</sup> + 1 M NaOH and 6 M NaOH, respectively. (b) Polarization and power density curves. (c) The digital photograph of the tandem Zn-nitrate flow battery and AEMHE devices for hydrogen production.

The Zn-nitrate flow battery was assembled with an open circuit voltage over 0.8 V (Fig. 6a). The polarization curves by steady-state chronopotentiometry were obtained. The Zn-nitrate flow battery is practically feasible with the discharging power density up to 12 mW cm<sup>-2</sup> (Fig. 6b). To confirm the efficacy of the electricity supply of this battery, the low-voltage-driven anion exchange membrane hydrazine electrolyzer (AEMHE), which used the bimetallic Co-P@NF as both the anode and cathode for the HzOR and HER, respectively (Fig. 6c). The Zn-nitrate flow battery can well drive the AEMHE with only a cell voltage of 0.47 V and give high current density over 35 mA cm<sup>-2</sup> for H<sub>2</sub> production (Fig. S19†), the corresponding H<sub>2</sub> production rate as 0.64 mmol h<sup>-1</sup> cm<sup>-2</sup>. This result shows that the Zn-nitrate flow battery possesses great potential in electricity generation.

#### 4. Conclusions

In summary, an amorphous P doped Co (Co-P@NF) was prepared via a facile electrodeposition process. Co-P@NF with a higher density oxygen vacancies sites showed high catalytic activities for both NO<sub>3</sub><sup>-</sup>RR and HzOR. They show a high current density over 2 A cm<sup>-2</sup> for NH<sub>3</sub> formation and high FE up to 91%

at -0.3 V versus RHE in the electrolyte of 1 M  $NO_3^-$ . Thanks to the amorphous structure, the Co-P could preserve the oxygen vacancies sites and give a high stability for NO<sub>3</sub> reduction over 82 h electrolysis with the current density over 799 mA cm $^{-2}$  and the FE over 90%. A low cell voltage of HzOR-assisted NO<sub>3</sub><sup>-</sup>RR at 300 mA cm<sup>-2</sup> was also obtained over Co-P@NF||Co-P@NF and was only 0.34 V. A Zn-nitrate flow battery was assemble using Co-P@NF as electrodes can deliver a peak power density of 12 mW cm2 and well drive an AEMHE, accompanied with an H2 yield rate of ca. 0.64 mmol h<sup>-1</sup> cm<sup>-2</sup>. This work demonstrates the feasibility for boosting the activity and stability toward NO<sub>3</sub><sup>-</sup>RR by constructing the amorphous structure.

# Data availability

All relevant data are within the manuscript and its additional files.

# **Author contributions**

Jin-Long Fan: conceptualization, investigation, writing - original draft. Sheng-Bo Liu: investigation. Ming-Liang Chen: writing – review & editing. Zhangxiong Wu: writing – review & editing. Sheng-Peng Sun: investigation, supervision. Yao-Yin Lou: conceptualization, investigation, supervision, writing – review & editing.

# Conflicts of interest

No potential conflicts of interest were reported by the authors.

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