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Recent developments and prospects for engineering first-row transition metal-based catalysts for electrocatalytic NO_x⁻ reduction to ammonia

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Immense interest in the electrocatalytic reduction of nitrate (NO₃⁻)/nitrite (NO₂⁻) to ammonia has been driven by promising prospects as an eco-friendly and energy-efficient approach for wastewater treatment and ammonia synthesis. Currently, a variety of transition metal-based catalysts have been developed, but these still suffer from inferior selectivity and low energy conversion efficiency. Among diverse transition metal-based catalysts, first-row transition metal-based catalysts exhibit the advantages of suitable adsorption energies for NO_x⁻ species and high abundance. As a consequence, this review firstly discusses the reaction mechanisms and hurdles for reaching high-efficiency NH₃ production and analyses the principles for constructing superior NO_x⁻ reduction catalysts. Then, the focus of this review is placed on recent advances in first-row transition metal-based electrocatalysts, including Cu, Fe, Co, Ni, Ti-based electrocatalysts, for electrochemical NO₃⁻/NO₂⁻ reduction. Finally, the challenges and opportunities are highlighted for future studies and applications. This review provides novel perspectives and ideas for developing first-row transition metal-based NO_x⁻ reduction electrocatalysts.

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1. Introduction

Ammonia (NH₃) exerts a significant influence on economic development and demographic growth.^{1–3} NH₃ currently accounts for approximately 5% of the market value of chemicals, occupying a market size reaching 175 million tons and a market value approaching \$67 billion.⁴ Lim *et al.* have speculated that with the global population growing on a moderate fecundity assumption model, the market size of NH₃ could reach 220–402 million tons in 2050, while the market value would expand to \$374–\$495 billion.⁵ However, current ammonia production is primarily dependent on the Haber–Bosch process. The Haber–Bosch process is characterized by high energy consumption (5.5 EJ yr⁻¹), accompanied by significant carbon dioxide emissions exceeding 450 million metric tons per year.⁶ Apparently, with the unceasing employment of the industrial Haber–Bosch route for NH₃ production, the global energy crisis will inevitably be aggravated and the achievement of carbon neutrality will be hindered. In order to

achieve a flourishing ammonia economy, it is highly imperative to explore alternative routes for sustainable ammonia production.

Among these alternative approaches, the electrocatalytic reduction of N-containing species has emerged as an attractive approach under ambient conditions on account of its appealing cost and environmental friendliness.⁷ The advantages of electrocatalytic ammonia production compared to the Haber–Bosch process (N₂ + 3H₂ = 2NH₃) include the following two points: (1) It is operated under ambient conditions, which avoids the high temperature (>450 °C) and high pressure (>10 MPa) conditions of the Haber–Bosch process.⁸ (2) It is more conducive to achieving carbon neutrality due to the elimination of massive CO₂ production. However, since aqueous phase conditions are usually adopted for electrocatalytic ammonia production, it is difficult to escape competition from the hydrogen evolution reaction (HER).⁹ In addition, the lack of prominent catalysts results in large-scale electrocatalytic ammonia production with uncompetitive cost and inferior efficiency compared to the sufficiently mature Haber–Bosch process in industry. For example, the aqueous electrochemical N₂ reduction reaction (NRR) has been extensively studied under laboratory conditions,¹⁰ but currently reported aqueous NRR experiments are limited by low NH₃ yield (10⁻¹¹ mol cm⁻² s⁻¹) and an undesirable faradaic efficiency (FE) below

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20%.¹¹ The bleak performance of the aqueous NRR system is apparently insufficient to realize sustainable ammonia production.¹²

Recently, the electrocatalytic nitrate (NO_3^-)/nitrite (NO_2^-) reduction reaction has shed new light on renewable NH_3 synthesis as NO_x^- exhibits substantially increased solubility in water and lower cleavage energy compared to N_2 .^{13–15} The stable supply of NO_3^- and NO_2^- is a crucial factor for achieving electrocatalytic green ammonia generation. It might be advisable to obtain NO_x^- from wastewater.¹⁶ The enrichment of NO_x^- in ground water is primarily caused by the runoff of artificial nitrogenous fertilizers and acid rain formed by nitrogen oxides emitted from factories and automobiles, which results in jeopardized public health, devastated aquatic ecosystems and disturbed global nitrogen cycle.¹⁷ Indeed, the electrocatalytic reduction of NO_x^- to produce NH_3 is also a favorable avenue for removing NO_x^- pollution and restoring the global nitrogen cycle.¹⁸ Compared with the most widely employed conventional biotechnology constrained by prohibitive start-up costs and cumbersome technical procedures,¹⁹ the electrocatalytic reduction of NO_x^- to NH_3 holds a more favourable prospect for the treatment of NO_3^- and NO_2^- in wastewater, due to its uncomplicated operating conditions, excellent efficiency and appealing products. However, for direct electrocatalytic reduction or advanced collection of NO_x^- from wastewater, it is difficult to escape from the effects of unknown NO_x^- concentration, other ions and pH of the solution.^{20,21} Time-consuming and labor-intensive wastewater pretreatment is required due to the numerous unknown factors of wastewater, which hinder the realization of electrocatalytic NH_3 production.

In addition to gaining NO_3^- from wastewater, the electrocatalytic oxidation of N_2 to produce NO_3^- opens up novel insights into the green industrial synthesis of NO_3^- and restoration of the global nitrogen cycle.²² Diverse electrocatalysts including Fe– SnO_2 ²³ and $\text{Pd}_{0.9}\text{Ru}_{0.1}\text{O}_2$ ²⁴ have been developed for the electrooxidation of N_2 to NO_3^- over the past three years. However, accelerating the complex five-electron transfer process and breaking the robust $\text{N}\equiv\text{N}$ bonds remain major bottlenecks for the development of this technology.²⁵

With regard to electrocatalytic NO_x^- reduction reactions, they are multi-step processes involving multiple electron/proton transfers and complex intermediate evolution.²⁶ Moreover, NO_x^- reduction reactions face the challenge of competing with other side reactions under aqueous conditions including the hydrogen evolution reaction (HER).²⁷ Consequently, considerable efforts are currently being spent on exploring highly selective and active catalysts for the electrocatalytic reduction of NO_x^- to NH_3 , including noble-metal catalysts,^{28,29} single-atom catalysts,³⁰ transition metal-based compounds,³¹ non-metal-based nanomaterials.³²

The number of publications during the past decade on different elements applied to the electrocatalytic reduction of NO_x^- to NH_3 according to the Web of Science database are illustrated in Fig. 1. Obviously, first-row transition metals including Ti, Fe, Co, Ni and Cu have gained broad attention

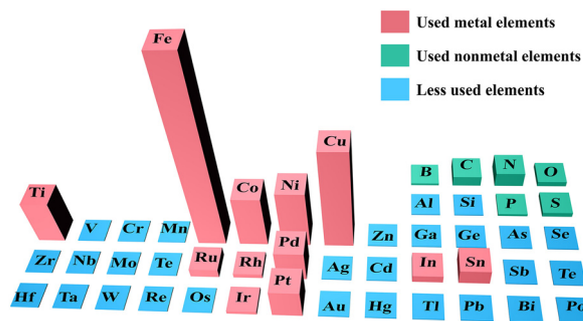


Fig. 1 Publications on various elements for NO_x^- reduction in the last decade (the height of the column is positively correlated with the number of relevant publications).

due to their affordability and easily tunable structures. Some reviews focus on NO_3^- reduction,^{33–35} but few contributions concentrate on NO_2^- reduction to NH_3 . Indeed, NO_2^- is more easily reduced compared to NO_3^- , and NO_3^- in the natural environment is also easily transformed into NO_2^- by microorganisms.³⁶

Hence, this review summarizes the recent progress of studies on first-row transition metal-based electrocatalysts for aqueous $\text{NO}_3^-/\text{NO}_2^-$ reduction to NH_3 under ambient conditions. This review firstly presents a brief demonstration of the differences in mechanisms of NO_3^- and NO_2^- reduction, while hurdles and corresponding strategies are analysed toward electrocatalytic NO_x^- reduction to NH_3 . Then, the focus is placed on recent advances in first-row transition metal-based electrocatalysts, including Cu, Fe, Co, Ni, and Ti-based electrocatalysts, for electrochemical $\text{NO}_3^-/\text{NO}_2^-$ reduction. Finally, the challenges and opportunities for future studies and applications are discussed with the aim of shedding light on the preparation of highly active and selective transition metal-based electrocatalysts for electrochemical $\text{NO}_3^-/\text{NO}_2^-$ reduction.

2. Reaction mechanisms of electrocatalytic NO_x^- reduction to ammonia

2.1. Reaction mechanisms

Recognizing and mastering the mechanisms of the electrocatalytic conversions of NO_3^- and NO_2^- to NH_3 are a prerequisite for exploring highly selective and active electrocatalysts. The electrochemical reduction of NO_3^- is a complex process with multielectron reactions,³⁷ which involve various nitrogen-containing intermediates and products ranging from -3 to $+5$ valence states. N_2 and $\text{NH}_3/\text{NH}_4^+$ are widely recognized as the most thermodynamically stable products,²⁶ but the final result can be altered for several reasons, such as cathode materials and the pH value of the solution. The indirect autocatalytic reduction mechanism and the direct electrocatalytic reduction mechanism comprise the currently accepted mechanisms of

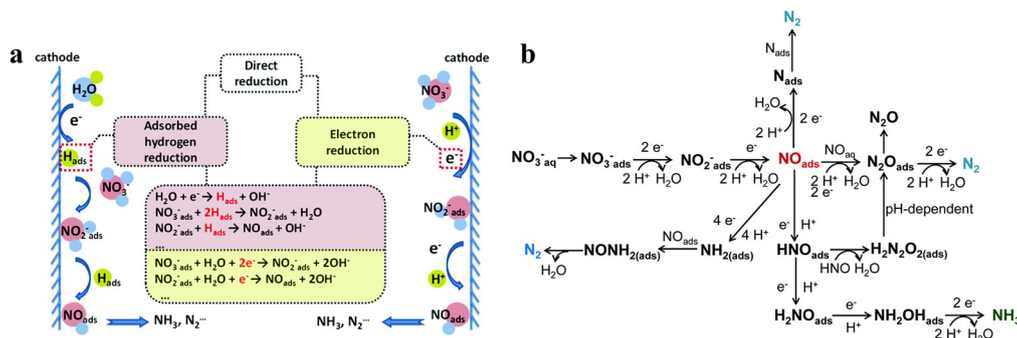
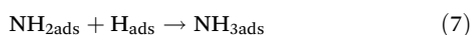
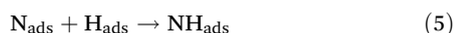
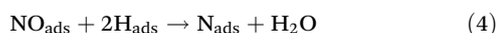
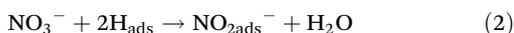


Fig. 2 (a) The proposed direct and indirect pathways of nitrate electroreduction. (b) The electron-mediated pathway of nitrate electroreduction copied with permission.³⁸ Copyright 2021, Royal Society of Chemistry.

NO_3^- electroreduction.³⁸ The conversion of NO_3^- to NH_3 generally follows the direct electrocatalytic reduction mechanism occurring in alkaline media with comparatively low concentrations of NO_3^- .³⁹ The active adsorbed hydrogen atom (H_{ads})-mediated route and the electron-mediated pathway are pursued simultaneously in the direct mechanism (Fig. 2a), which leads to the complexity of electrocatalytic NO_3^- reduction.

The electroreduction of NO_3^- is initiated by the adsorption of NO_3^- ions onto the cathodic electrodes. Adsorbed NO_3^- is transformed into NO_2^- by a tripartite electrochemical–chemical–electrochemical process, which is the dominant rate-controlling step.⁴⁰ Later, the important nitric oxide (NO_{ads}) intermediate is obtained by NO_2^- conversion. As depicted in Fig. 2b, NO_{ads} can be reduced to NH_3 as the ultimate product and occupy a dominant position in the N_2 formation pathway.

In addition, the reduction process of NO_3^- can proceed *via* the intermediate H_{ads} . NO_2^- , NO_3^- , and NO_{ads} can be reduced by H_{ads} .⁷ The predominant final product in this H_{ads} -mediated process is ammonia, which is caused by the fact that the formation of N–N bonds mediated by H_{ads} is kinetically more challenging than the formation of N–H bonds. The specific H_{ads} -mediated pathways are described in reactions (1)–(7).



The electrocatalytic reduction of NO_2^- to NH_3 is roughly identical to the reduction process following the conversion of NO_3^- to NO_2^- .⁴¹ It is worth noting that during the reduction of NO_{ads} generated by NO_2^- to NH_3 , NO_2^- may generate hydroxyl-

amine (NH_2OH) as a by-product. This specific reaction is as follows:



2.2. Hurdles and strategies for reaching high-efficiency NH_3 production

2.2.1. Selectivity requiring improvement.

The various nitrogen-containing products involved in the NO_x^- reduction process were initially discussed in section 2.1. During the NO_x^- reduction process, the hydrogenation path of N_{ads} may be hampered by the coupling reaction with the adjacent N_{ads} , which leads to the production of N_2 , N_2H_4 and N_2O , blocking the generation of NH_3 . Therefore, avoiding coupling reactions between N_{ads} and N_{ads} is essential to design high selectivity electrocatalysts for NH_3 synthesis. Simultaneously, competition from the HER is also a major obstacle in the process of reducing NO_x^- to ammonia.³⁴ Although the standard electrode potentials of the NO_x^- reduction reactions are larger than that of the HER (Fig. 3a), it is still difficult to circumvent competition from the HER, which only involves two-electron transfers in an aqueous system.

DFT calculations have been employed to explore the thermodynamics of NH_3 selectivity on different transition metal elements. The Gibbs free energy of N_{ads} ($G_{\text{ad}}(\text{N})$) of different transition metal elements is correlated with the Gibbs free energy of the NO_x^- reduction reaction. Fig. 3b and c demonstrates theoretical estimates based on the $G_{\text{ad}}(\text{N})$ of various transition metal elements.⁴¹ Cu is more favourable for conducting NORR compared to HER, which simultaneously possesses a suitable adsorption energy scaling relationship. Thereby, Cu is theoretically considered to be a decent catalyst for ammonia production. The selectivity of catalysts can also be affected by different potentials. Liu *et al.* predicted the activity and selectivity for NO_3^- reduction on transition metals at various voltages;²⁶ this was accomplished by exploring the adsorption energies of O and N atoms under different potentials. The study reveals that catalysts with moderate ΔE_{O} and ΔE_{N} are inclined to exhibit more remarkable NH_3 selectivity at higher negative potentials. As shown in Fig. 3d, Co and Rh

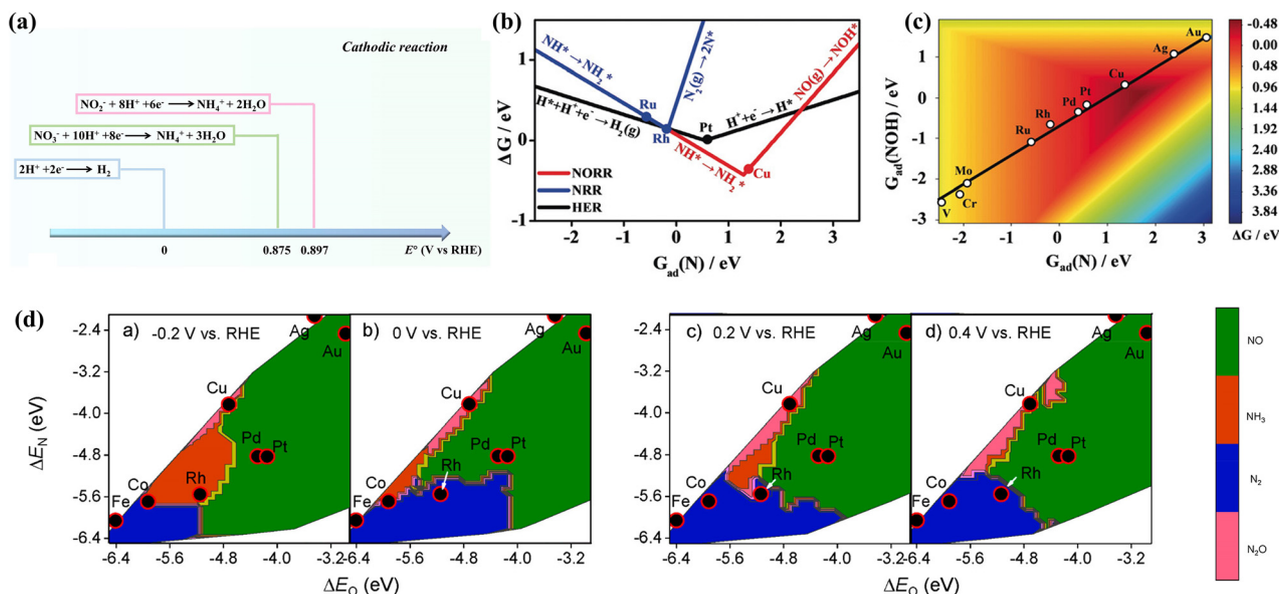


Fig. 3 (a) Comparison of standard reduction potentials for the HER and NO_3^- reduction reactions. (b) Comparison of the ΔG -determining steps between diverse reactions. (c) Activity planar graph for ammonia production copied with permission.⁴¹ Copyright 2020, John Wiley and Sons. (d) Theoretical selectivity maps of various nitrogen-containing products from electrocatalytic NO_3^- reduction on the basis of ΔE_{O} and ΔE_{N} under different applied voltages copied with permission.²⁶ Copyright 2019, American Chemical Society.

demonstrate more outstanding selectivity for ammonia at -0.2 V vs. RHE.

It is obvious from the above content that the adsorption capacities of N or O-containing species are essential to enhance NH_3 selectivity. According to d-band theory,^{42,43} the adsorption energy of intermediates is related to the location of the d-band center of the catalyst. Obviously, ΔE_{O} and ΔE_{N} can be modulated by adjusting the d-band center of the catalyst, thus regulating the selectivity of ammonia generation.⁴⁴ For example, Cu is doped in Co_3O_4 nanowires to regulate the d band center position of Co_3O_4 due to the interactions between Cu and Co.⁴⁵ The density of states reveals that the d-band center position of Cu- Co_3O_4 (-1.57 eV) is downwardly shifted compared to Co_3O_4 (-1.45 eV). Correspondingly, the adsorption energy of $^*\text{NO}_2$ on the surface of Cu- Co_3O_4 is also reduced compared with that of Co_3O_4 . The hydrodeoxygenation free energy of the NO_3^- reduction intermediate is optimized in Cu- Co_3O_4 with an appropriate d-band center position and suitable $^*\text{NO}_2$ adsorption capacity, and the optimal NO_3^- reduction performance of Cu- Co_3O_4 is also experimentally confirmed to reach a decent yield rate of $36.71 \text{ mmol h}^{-1} \text{ g}^{-1}$ at -0.6 V vs. RHE with an excellent FE of 86.5%.

Circumventing the direct coupling of $^*\text{N}$ is also a valid approach for reducing the generation of N_2 and N_2O by-products and amplifying the NH_3 selectivity. Compared with the inevitable production of N_2O and N_2 by transition metal-based catalysts possessing two adjacent active sites, single-atom metals are more promising for inhibiting the direct coupling of N owing to the absence of adjacent active sites.⁴⁶ The isolated Au atoms are distributed into Cu nanoparticles to modify the selectivity and activity of NO_3^- reduction.²⁹ The introduc-

tion of Au single atoms on Cu has been proved to observably accelerate NO_3^- adsorption; the deoxidation process of $^*\text{NO}_2$ is also boosted due to the formed Au-Cu bond. In addition, more outstanding NH_3 selectivity is obtained for Cu doped with Au single atoms due to the difficult direct coupling of $^*\text{N}$ on single atoms, reaching remarkable FEs up to 99.69% at -0.80 V vs. RHE.

Furthermore, strong competition from the HER is hard to neglect in the aqueous system.⁴⁷ However, it is unconscionable to impede the HER by inhibiting the formation of active hydrogen species, since active hydrogen species are indispensable reactants for the subsequent hydrogenation process in NO_x^- reduction. In an environment with sufficient active hydrogen species, employing metal-based catalysts such as Mo and W (Fig. 4c) with difficult release of H_{ads} intermediates may be a decent approach to reduce hydrogen formation and enhance NH_3 selectivity.⁴⁸ The strategy of employing two sites can be used for reference to simultaneously realize active hydrogen species production and reduce the direct coupling of active hydrogen species. In this strategy, one site acts as a proton warehouse to promote proton generation, and the other site temporarily stores active hydrogen species for the subsequent hydrogenation reaction.

In addition, the hydrophilicity and aerophobicity of catalysts can also have a bearing on the selectivity of catalysts. The outstanding hydrophilicity of catalysts can accelerate mass transfer in an aqueous system. For example, the super-hydrophilic surface of Fe-based cyano-coordination polymer nanosheets (Fe-cyano NSSs) is more conducive to enhancing the contact between the electrolyte and the electrode surface for faster conversion to Fe^0 active sites. Abundant Fe^0 active

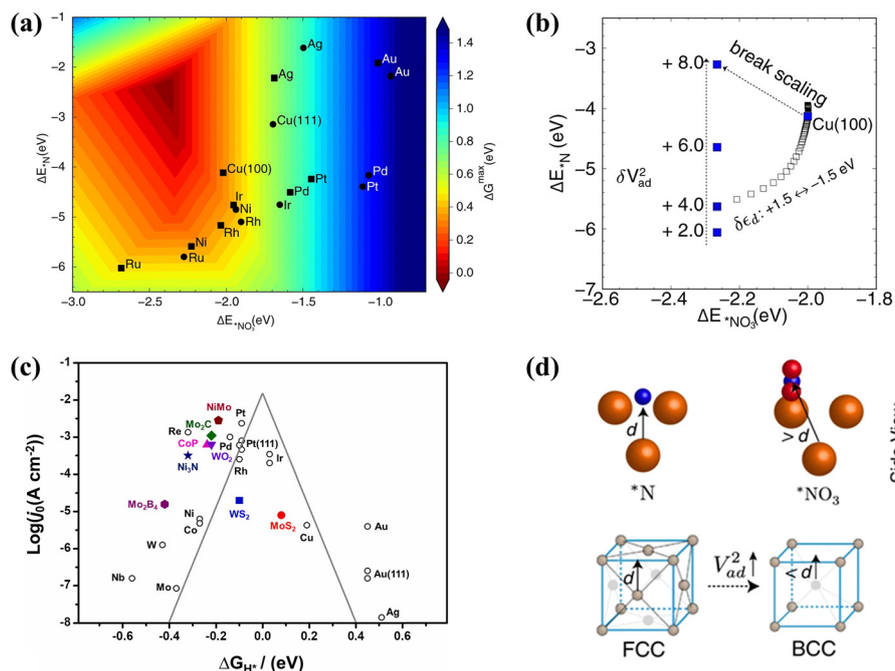


Fig. 4 (a) The activity volcano plot of various metal elements for the conversion of NO_3^- to NH_3 . (b) Adsorption energies of *NO_3 and *N on Cu (100) under different V_{ad}^2 . (d) Models of the (100) facet and the (111) facet copied with permission.⁵³ Copyright 2022, Springer Nature. (c) The volcano plot of Gibbs free energy of hydrogen (ΔG_{H^+}) on different metal-based compounds copied with permission.⁴⁸ Copyright 2019, Elsevier.

sites in Fe–cyano-R NSs are more favorable for the adsorption of NO_x^- ions.⁴⁹ Moreover, considering that the strongly competing reaction with the NO_x^- reduction reaction is the HER, which is accompanied by bubble release, the super-aerophobic nature of a material is more conducive to boosting NO_x^- reduction, which can effectively avoid blockage of the active sites caused by the accumulation and adhesion of gas bubbles on the surface.⁵⁰

2.2.2. Inferior energy conversion efficiency. Although decent selectivity has been shown in the currently developed catalysts, little emphasis has been placed on the undesirable energy conversion efficiency. Indeed, due to the multiple electron transfer processes of NO_x^- , the majority of ammonia-producing systems require considerably negative potentials (< -0.2 V) to exhibit promising performance, accompanied by suboptimal energy consumption (21–38 kW h kg^{-1}).⁵¹ Therefore, it is of major concern to reduce the overpotential in the NO_x^- reduction process, which is tightly related to the energy conversion efficiency of the ammonia production process.

Exploring the factors influencing the activation energy of the elemental reaction is of significance to optimize the overpotential, which is the potential determining step (PDS) in the conversion of NO_x^- to NH_3 . Despite the increased adsorption of reactants causing accelerated reactant activity, the dilemma of enhanced activation energy still exists due to the corresponding increase in intermediate adsorption energies restricting the desorption of products. According to Sabatier's principle, scaling relationships between the adsorption energy and activation energy are obstacles to overcome for enhanced

reduction activity, causing a volcano plot for the adsorption of reactants in NO_x^- reduction.⁵² The design of catalysts with dual active sites is an effective strategy to break the scaling relationships by simultaneously promoting the adsorption of reactants and accelerating the desorption of products. The scaling relationships are broken in $\text{Co}_3\text{O}_4/\text{Cu}$ catalysts with dual active sites, reaching enhanced adsorption of NO_3^- (-2.91 eV) with a low desorption energy barrier for NH_3 (0.13 eV).⁴⁵

Imposing nanoscopic confinement⁵⁴ and adjusting strain⁵⁵ can also be considered to break scaling relationships. However, constructing strategies based on theoretical guidance is more important for circumventing scaling relationships. The adsorption energies of hollow *N and bridge-bidentate *NO_3 are employed as reactivity descriptors for constructing an activity volcano plot of NO_x^- reduction to NH_3 (Fig. 4a).⁵³ The (100)-oriented B2 CuPd nanocubes are developed with increased *NO_3 adsorption and attenuated *N binding; this has been confirmed to break the scaling relationship. The enhanced bridge-bidentate *NO_3 adsorption is caused by the upshift of the d-band center position in Cu after introducing Pd. More significantly, regulating the Pauli repulsion between the metallic d-state and the adsorbate frontier orbital has been found to promote the hydrogenation process of hollow *N . As described in Fig. 4b, the adsorption strength of *N at the (100) hollow site increases with the enhancement of the interatomic coupling strength (V_{ad}^2); this is caused by the obtained dominant position of Pauli repulsion with decreasing adsorbate-metal antibonding states. Fig. 4d displays the advantage of the

(100) facet compared to the (111) facet; stronger interatomic coupling can be achieved for the (100) facet due to the closer distance between the subsurface metal–ligand and *N. Hollow *N is more easily destroyed due to the dominant effect of Pauli repulsion, which promotes the hydrogenation of *N to form NH₃. This intriguing strategy can be employed for reference to break the scaling relationship through modifying the Pauli repulsion between the metallic d-state and the adsorbate frontier orbital.

In summary, the design of high-efficiency NO_x⁻ reduction electrocatalysts requires consideration of selectivity enhancement and overpotential reduction, which are of significant importance for achieving large-scale and green electrocatalytic ammonia production routes. According to the current development of aqueous NO_x⁻ reduction technology, the following points should be observed when designing electrocatalysts with excellent selectivity and remarkable overpotential. (1) Increasing the adsorption of NO₃⁻ and NO₂⁻ on catalysts is beneficial for the reaction. However, the problem of attenuated product desorption should also be taken into account accompanied by enhanced NO_x⁻ adsorption. It is an important strategy to drastically improve the NO_x⁻ reduction performance of catalysts though breaking the scaling relationship between the adsorption energies of intermediates and reactants. (2) It is difficult to escape competition from the HER under aqueous conditions. Suppressing the formation of active hydrogen species is unadvisable to reduce HER competition because active hydrogen species are indispensable reactants for the subsequent hydrogenation process in NO_x⁻ reduction. The strategy of employing two sites can be used for reference to simultaneously realize active hydrogen species production and reduce the direct coupling of active hydrogen species. In this strategy, one site acts as a proton warehouse to promote proton generation, and the other site temporarily stores active hydrogen species for the subsequent hydrogenation reaction. (3) In addition to the HER side reaction, other side reactions generating N₂ or NO should not be ignored. Improved stability of intermediates including *NO₂ or *NOH facilitates more efficient ammonia production.

3. First-row transition metal-based electrocatalysts

3.1. The unique advantages of first-row transition metal elements

The development of transition metal-based catalysts is a popular topic due to their low price and controllable structures.^{56–58} In accordance with the above exploration of the design principles for NO_x⁻ reduction catalysts, a promising catalyst should possess moderate interactions with the reactants; either overly weak or excessively strong reactant adsorption is unfavourable for the reaction to proceed. Fig. 5a reveals the adsorption energies of NO₃⁻ species and protons on transition metal atoms. Obviously, overly strong adsorption strength with NO₃⁻ species will hinder the production of NH₃ due to dead-end species formation.⁵⁹ Therefore, Zr, Hf, W and Re atoms are obviously unsuitable for employment as NO_x⁻ reduction catalysts. The difference between the adsorption energies of NO₃⁻ species and protons can provide rough indications of the selectivity degree. Au, Rh, and Ag atoms are found to possess strong adsorption for protons; this is clearly detrimental for escaping competition from the HER. HER competition is likely to be reduced with first-row transition metals, including Ti, Fe, Co, Ni and Cu, which possess more suitable adsorption energies for NO₃⁻ species and protons. In addition, the unclosed d-orbital shells of first-row transition metals are conducive to the injection of charge into the lowest unoccupied molecular π* orbital of NO_x⁻.⁶⁰ The first-row transition metal elements also satisfy the requirements of desirable abundance and competitive costs (Fig. 5b).⁶¹ Since few or no reports are found on the application of V, Cr, and Mn as NO_x⁻ reduction catalysts, Ti, Fe, Co, Ni and Cu-based catalysts are mainly discussed in this paper.

3.2. Cu-based electrocatalysts

Considerable endeavors have been directed towards the development of Cu-based catalysts (Table 1) with competitive costs and superior intrinsic activity,⁵³ for example, the alloying of

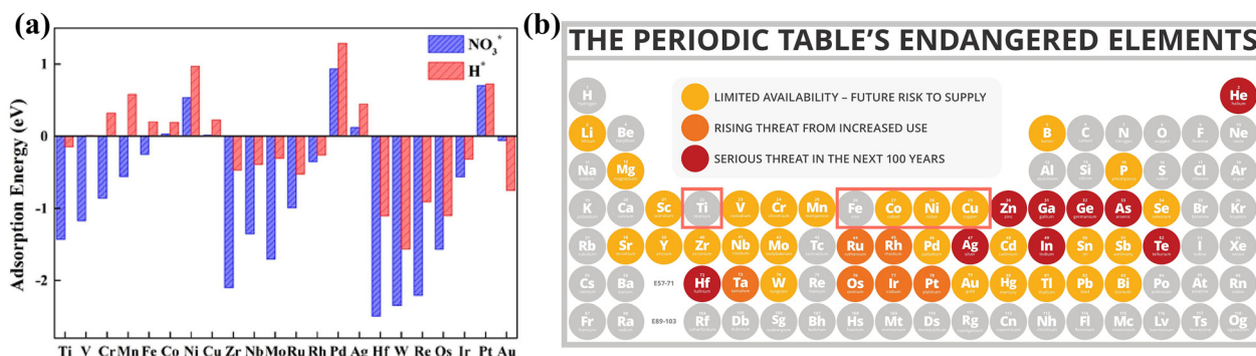


Fig. 5 (a) Adsorption energies for NO₃⁻ species and protons on different metals copied with permission.⁵⁹ Copyright 2022, Elsevier. (b) The predicted elemental content of the periodic table within the next 100 years (gray represents high abundance) copied with permission.⁶⁰ Copyright 2021, Elsevier.

Table 1 Summary of NO_x^- reduction performance of Cu-based electrocatalysts

Electrocatalyst	Electrolyte	Potential (V vs. RHE)	NH_3 yield	Faradaic efficiency (%)	Ref.
Cu SAC	0.1 M OH^- + 0.1 M NO_3^-	-1.0	$4.5 \text{ mg cm}^{-2} \text{ h}^{-1}$	84.7	104
Cu nanosheets	0.1 M OH^- + 0.1 M NO_3^-	-0.15	$390.1 \text{ } \mu\text{g mg}^{-1} \text{ h}^{-1}$	99.7	105
$\text{Cu}_{50}\text{Ni}_{50}$ alloy	1 M KOH + 10 mM KNO_3	-0.15	—	99 ± 1	63
$\text{Cu}_{50}\text{Co}_{50}$	1 M KOH + 100 mM KNO_3	-0.2	$4.8 \text{ mmol cm}^{-2} \text{ h}^{-1}$	100	64
Cu-CuO	0.1 M KOH + 0.1 M NO_3^-	-0.8	$3.17 \text{ mol h}^{-1} \text{ g}^{-1}$	98.7	106
$\text{Cu}@ \text{Cu}_{2+1}\text{O}$ nanowires	0.5 M K_2SO_4 + 50 $\text{mg L}^{-1} \text{ NO}_3^-$	-0.545	$3.53 \text{ } \mu\text{g h}^{-1} \text{ mg}^{-1}$	87.7	107
$\text{Cu}@ \text{TiO}_2/\text{TP}$	0.1 M Na_2SO_4 + 0.1 M NO_2^-	-0.6	$760.5 \text{ } \mu\text{mol h}^{-1} \text{ cm}^{-2}$	95.3	108
$\text{CF}@ \text{Cu}_2\text{O}$	0.1 M PBS + 0.1 M NaNO_2	—	$441.8 \text{ } \mu\text{mol h}^{-1} \text{ cm}^{-2}$	94.2	109
$\text{Cu}_3\text{P NA}/\text{CF}$	0.1 M PBS + 0.1 M NaNO_2	—	$95.6 \text{ } \mu\text{mol h}^{-1} \text{ cm}^{-2}$	91.2	110

Cu.⁶² The introduction of other metal elements contributes to modulating the d-band center of Cu and thus regulating the adsorption energies of intermediates. The $\text{Cu}_{50}\text{Ni}_{50}$ alloy obtained by introducing Ni shows a nitrate reduction activity six times higher than that of pure copper at 0 V.⁶³ By alloying Ni, the half-wave potential of Cu is shifted upwards by 0.12 V and the adsorption energies of intermediates including $^*\text{NO}_3$, $^*\text{NO}_2$ and $^*\text{NH}_2$ are regulated. The bimetallic electrocatalysts can also synergistically promote the NO_x^- reduction process by combining the advantages of different metals. The CuCo alloy nanosheets exhibit FEs exceeding 90% for ammonia over a wide potential window (from -0.1 to -0.4 V) through emulating the bifunctional nature of nitrite reductase.⁶⁴ $^*\text{H}$ species can be more easily adsorbed by Co, resulting in a decrease in ΔG of the hydrogenation reaction of $^*\text{NO}_x$. The electronic structure of Cu is also adjusted by introducing Co, which is conducive to reducing the energy barrier of the initial $^*\text{NO}_3$ adsorption step on Cu. In addition to transition metal elements, combinations of Cu with precious metals to form

high-performance alloy catalysts are also not unusual. The CuRu alloy demonstrates a decent nitrate reduction activity and significantly inhibits the production of nitrite.⁶⁵ The superior performance of CuRu alloy can be attributed to the synergistic effects between the Ru and Cu sites through relay catalysis. The Cu active sites dramatically promote the reduction of NO_3^- to NO_2^- , while the Ru sites exhibit outstanding activity for the subsequent conversion of NO_2^- to NH_3 . However, the cost of producing such bimetallic catalysts is undesirable due to the scarce reserves and prohibitive price of Ru.

Among various Cu-based catalysts, Cu and Cu_2O have been gaining considerable attention. Revealing the reactivity of different exposed facets on Cu and Cu_2O is crucial for investigating the kinetics of NO_x^- reduction and designing catalysts with superior performance. The NO_3^- reduction behaviour of the (111), (100) and (110) facets on Cu is probed, while competition from the HER on various facets is also investigated at different pH values (Fig. 6a).⁶⁶ Theoretical calculations

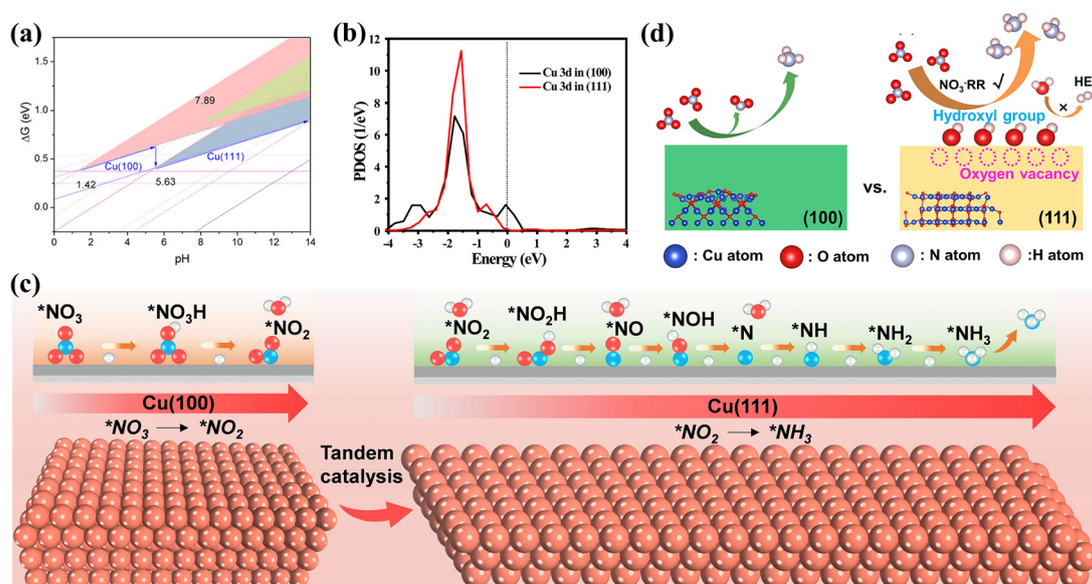


Fig. 6 (a) Comparison of different facets under various pH conditions copied with permission.⁶⁶ Copyright 2021, American Chemical Society. (b) Projected density of states of Cu 3d states on different Cu_2O surfaces copied with permission.⁶⁷ Copyright 2022, American Chemical Society. (c) Tandem catalytic process of Cu(100) and Cu(111) surfaces copied with permission.⁶⁹ Copyright 2023, John Wiley and Sons. (d) Effects of surface oxygen species on the NO_3^- reduction process at different surfaces of Cu_2O copied with permission.⁷¹ Copyright 2023, Elsevier.

confirm the strongest contribution from the (100) and (111) facets in the reduction of NO_3^- to NH_3 , excluding the (110) face. The Cu(111) surface exhibits the greatest NO_3^- to NH_3 performance in near-neutral and alkaline environments, while the Cu(100) surface demonstrates better performance in strongly acidic environments. The effects of Cu_2O (100) and (111) surfaces on NO_3^- reduction are also explored. The higher NH_3 yield of $743 \mu\text{g h}^{-1} \text{mg}_{\text{cat}}^{-1}$ on the Cu_2O (100) surface is attributed to the partially unfilled Cu 3d state possessed by the (100) facet (Fig. 6b), simultaneously accepting electrons from Cu_2O and combining with NO_x^- . In contrast, the excess filling of the Cu 3d state on the (111) surface inhibits the formation of NH_3 .⁶⁷ Based on the overview of the roles of the various crystalline planes, it is evident that constructing electrocatalysts with synergistic multiple crystalline planes is more favourable for NO_x^- reduction.⁶⁸ For example, the Cu nanosheets benefit from the tandem interactions between the Cu(100) and Cu(111) facets, obtaining an excellent NH_3 yield of $1.41 \text{ mmol h}^{-1} \text{cm}^{-2}$ and remarkable stability reaching 700 h at -0.59 V . As depicted in Fig. 6c, NO_2^- formed on the Cu(100) surface is subsequently hydrogenated on the Cu(111) surface; such tandem catalysis accelerates the hydrogenation of $^*\text{NO}$ to $^*\text{NOH}$ and the generation of NH_3 .⁶⁹

Meanwhile, defects have been proved to promote NO_x^- reduction processes in synergy with diverse crystal planes. Ultrathin copper oxide nanoribbons are obtained by *in situ* electrochemical reduction with exposed Cu(100) surfaces and abundant surface defects, achieving 2.3 times higher NH_3 yield in the NO_3^- reduction process than that *via* the Haber-Bosch process. The key to their outstanding property is the upshifting of the Cu d-band center due to the synergistic effect of the Cu(100) facets and defects; this facilitates the increased adsorption intensities of $^*\text{NO}_3$ and $^*\text{H}$, thus reducing the NO_3^- reduction potential and inhibiting the HER.⁷⁰ The contributions of surface oxygen species (oxygen vacancies and hydroxyl groups) on Cu_2O have also been demonstrated to

promote NO_3^- reduction. Chen's group obtained Cu_2O (111) surfaces with more oxygen vacancies and hydroxyl groups than the Cu_2O (100) facets.⁷¹ It was also confirmed that the adsorption of reactants and intermediates was accelerated by oxygen vacancies on the Cu_2O (111) surfaces, while the HER competition could be inhibited by surface hydroxyl groups (Fig. 6d).

Exposing specific facets can effectively enhance the activity of Cu-based catalysts, but the robustness of the material falls far short of expectations. The pure Cu catalysts are rapidly deactivated during NO_x^- reduction by poisoning from adsorbed nitrogenous species and hydrogen, with current densities attenuating by over half in six minutes under potentials exceeding -0.5 V .⁷² Gao *et al.* proposed a strategy of pulsed electrolysis of Cu to obtain Cu/ Cu_2O through peroxidation and self-repair processes,⁷³ and the Cu/ Cu_2O catalysts obtained by pulsed electrolysis were more favourable for both NO_3^- and NO_2^- reduction processes than the static electroreduction on Cu. However, the accumulation of NO_2^- during NO_3^- reduction is still not ignored; this is attributed to the excessively aggressive HER competition hampering subsequent hydrogenation processes. The regulated pH and enhanced cathodic potential naturally facilitate the hydrogenation process to a certain extent, but the degree of enhancement is inadequate. Consequently, pulsed electrolysis of CuNi alloy is employed to introduce $\text{Ni}(\text{OH})_2$ promoting H intermediate formation, which contributes to the improvement of the H adsorption capacity (Fig. 7a). The tandem catalysis of Cu-based and Ni-based materials breaks the awkward scenario (Fig. 7b) of the simultaneous existence of rapid NO_3^- transformation to NO_2^- (reaction (1)) and sluggish subsequent hydrogenation (reaction (2)), enabling a more efficient and stable catalytic NO_3^- reduction process at lower potentials.

In addition to the cathode potential and pH, CO_2 has also been confirmed to affect the NO_2^- reduction process on copper catalysts. Sun's group demonstrated that the faradaic efficiencies of copper catalysts could reach approximately

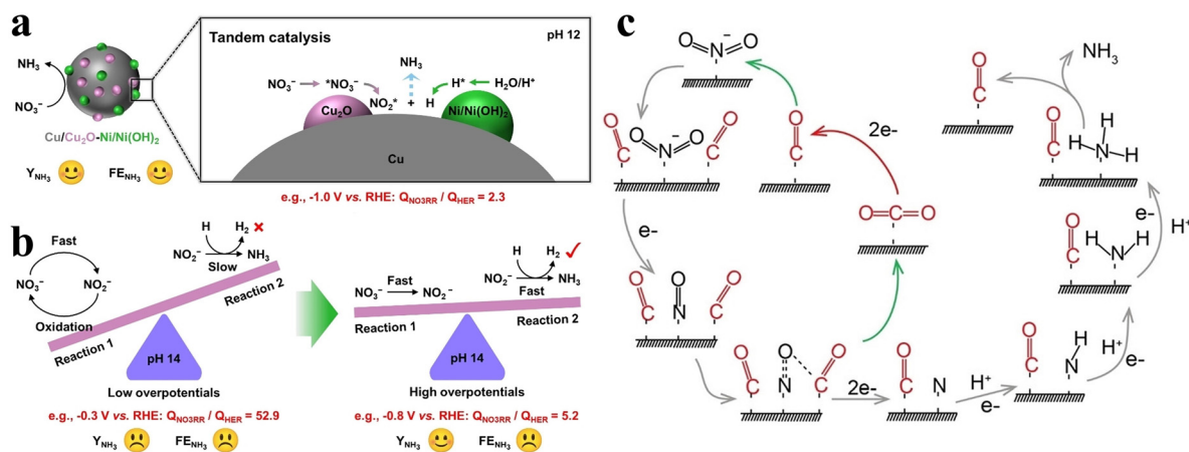


Fig. 7 (a) Tandem catalysis of CuNi alloy under pulsed electrolysis. (b) Kinetic regulation process of NO_3^- reduction copied with permission.⁷³ Copyright 2023, John Wiley and Sons. (c) NO_2^- reduction reaction pathway assisted by CO_2 copied with permission.⁷⁴ Copyright 2022, John Wiley and Sons.

100% for NO_2^- reduction within a wide range of potentials assisted by CO_2 .⁷⁴ The causes of the enhanced performance due to CO_2 are described in Fig. 7c. The *CO species obtained from CO_2 reduction simultaneously facilitate deoxygenation of the *NO intermediates and hydrogenation of the *NH_2 intermediate in the NO_2^- reduction process. In the facilitated *NO conversion process to *N , *CO is oxidized to CO_2 by *NO . CO_2 assisted strategies also prove to be beneficial for nitrate electroreduction.

3.3. Fe-based electrocatalysts

In nature, anammox bacteria have been proved to exploit $\text{Fe}^0/\text{Fe}^{2+}$ as electron donors for the denitrification of NO_3^- and NO_2^- .⁷⁵ In addition, the presence of Fe as the active site in nitrogenase enzymes has also promoted considerable exploration of Fe-based electrocatalysts with abundance on Earth.⁴⁴ The performances of currently developed Fe-based electrocatalysts are listed in Table 2. Recently, a deeper understanding has emerged of the intrinsic catalytic behavior on Fe isolated sites for NO_x^- reduction. The nitrogen-coordinated Fe single atoms on carbon were obtained *via* a polymer-hydrogel strategy.⁷⁶ The turnover frequency of synthesized Fe single sites is 12 times larger than that of metallic Fe nanoparticles. DFT cal-

culations confirm that the NO_3^- -preemption mechanism may occur in the catalytic process for Fe single atoms, effectively preventing strong water adsorption. Despite extensive studies on single-atom catalysts, which maximize the utilization of active atoms, the performance of single-atom catalysts is still limited in the multi-step NO_x^- reduction process due to the intrinsic adsorption energy scaling relationship. Dual-atom catalysts exhibit more favourable performance in multi-step reactions compared with single-atom catalysts, which inherit the advantages of single-atom catalysts and possess the synergistic effect of two sites.⁷⁷ For example, FeMo@g-CN has been theoretically proved to exhibit a promising limiting potential (-0.34 V) for forming NH_3 through NO_3^- reduction.⁷⁸ The activation of NO_3^- reduction in FeMo@g-CN is verified to originate from the synergistic interaction of FeMo dimer d orbitals rather than specific d suborbitals.

In addition to Fe single-atom catalysts, oxides of iron have received considerable attention. Wang *et al.* synthesized Fe_3O_4 doped with appropriate amounts of Cu,⁷⁹ achieving an ammonia yield of 222.75 ± 12.15 $\text{mg h}^{-1} \text{mg}_{\text{cat}}^{-1}$ and an excellent FE of $95.7\% \pm 4.8$ at -0.7 V. More intriguingly, in addition to enhancing the adsorption of various intermediates (Fig. 8a), the doping of Cu alters the state of *NO adsorbed on the metal

Table 2 Summary of NO_x^- reduction performance of Fe-based electrocatalysts

Electrocatalyst	Electrolyte	Potential (V vs. RHE)	NH_3 yield	Faradaic efficiency (%)	Ref.
Fe-PPy SACs	0.1 M KOH + 0.1 M NO_3^-	-0.2	$2.75 \text{ mg h}^{-1} \text{ cm}^{-2}$	100	76
Cu- Fe_3O_4	0.1 M KOH + 0.1 M KNO_3	-0.6	$179.55 \text{ mg h}^{-1} \text{ mg}_{\text{cat}}^{-1}$	100	79
Fe-cyano NSs	1.0 M KOH + 10 mM KNO_3	-0.5	$42.1 \text{ mg h}^{-1} \text{ mg}_{\text{cat}}^{-1}$	90	49
P- FeCo_2O_4 @CC	0.1 M KOH + 0.1 M KNO_2	-0.4	$5.29 \text{ mg h}^{-1} \text{ cm}^{-2}$	99.99	111
FeP	0.1 M KOH + 0.1 M KNO_2	-0.5 V	—	85	112

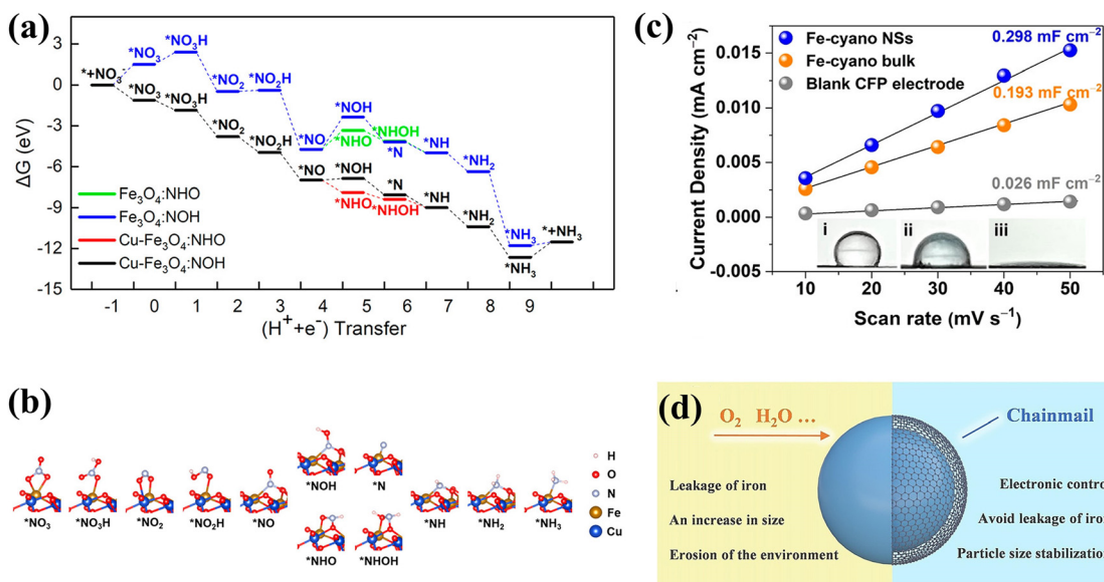


Fig. 8 (a) Gibbs free energy plot of NO_3^- reduction on the electrocatalyst surfaces. (b) Optimized structures of the intermediates on the Cu-doped Fe_3O_4 surface during NO_3^- reduction copied with permission.⁷⁹ Copyright 2023, American Chemical Society. (c) ECSA and contact angles of different catalysts copied with permission.⁴⁹ Copyright 2021, American Chemical Society. (d) Schematic diagram of the action of graphene armour copied with permission.⁸⁰ Copyright 2022, John Wiley and Sons.

atoms. *NO is susceptible to reaction poisoning due to strong adsorption. The *NO in Fe₃O₄ is adsorbed vertically on the Fe atom. After Cu doping, *NO adsorption is carried out by the bridge between Fe and O atoms, thus *NO preferentially undergoes the route of reduction to *NHO rather than the route to *NOH with higher ΔG (Fig. 8b).

Indeed, the hydrophilicity of materials is also worth considering, and hydrophilic materials are more conducive to the adsorption of NO_x⁻ ions. However, the hydrophobic surface of the bulk metal catalyst inhibits mass transport and diminishes the overall electrochemical surface area due to the dramatically reduced contact of the electrocatalyst with the electrolyte. Metalloxyano polymers are a class of superhydrophilic materials suitable for efficient electrocatalysis due to the abundance of hydrophilic groups and surface vacancies. The Fe-cyano NSs with a two-dimensional structure display a high NH₃ yield reaching 42.1 mg h⁻¹ mg_{cat}⁻¹ and FE exceeding 90% at -0.5 V vs. RHE.⁴⁹ Furthermore, an electrolyzer based on Fe-cyano NSs as the cathode and anode has been constructed with an energy efficiency of 26.2%. As illustrated in Fig. 8c, Fe-cyano NSs possess a superhydrophilic surface and distinctly larger active area owing to the nature of the cyano groups and two-dimensional structure, which facilitates enhanced molecular adsorption and provides speedier transport avenues for electrolyte diffusion and electron transport.

The Fe-based catalysts exhibit decent nitrate reduction activity, but the catalysts still suffer from the limitations of inferior long-term robustness and poor kinetic durability in the practical catalytic process.²⁰ The recent exploitation of chainmail-protected catalysts has offered innovative insights into the design of high-efficiency and robust Fe-based catalysts. Fe nanoparticles suffer from poor durability due to their greater susceptibility to leaching and oxidation during the electrocatalytic process. Zhang *et al.* prepared Fe nanoparticles protected by ultrathin graphene nanosheet layers (Fe@Gnc) for NO₃⁻ conversion to N₂; the NO₃⁻ conversion rate and N₂ selectivity of Fe@Gnc remained in excess of 96% after 960 hours of cycling.⁸⁰ The contribution of graphene nanochainmail is depicted in Fig. 8d. In the absence of graphene nanochainmail protection, active site agglomeration and toxic inactivation would occur on Fe particles. With the protection of graphene nanochainmail, the aggregation issue is not only tackled, but electron/ion transport is also accelerated by favorably conducting graphene for highly efficient and more sustained NO₃⁻ conversion. Despite the fact that Fe@Gnc is employed for NO₃⁻ reduction to N₂, it is still valid to draw

inspiration from the construction of Fe@Gnc for NH₃ production. Substituting the sites requiring protection for other high NH₃ selectivity sites might be a promising strategy.

3.4. Co-based electrocatalysts

Co has attracted extensive interest due to its low activation energy barriers for dissociating intermediates such as *NO₃, *NO₂, *NO and *N₂O.⁸¹ The Co based materials in Table 3 all show decent performances. Recently, the Gibbs free energies (ΔG_{rxn}) of the associated rate-determined step were compared for NO₃⁻ reduction on commonly available metals through theoretical screening. As shown in Fig. 9a and b, ΔG_{rxn} of the rate-determining step on Co exhibits the lowest value of 0.3 V.⁸² Furthermore, Shao *et al.* demonstrated through DFT calculations that surface oxygen on Co sites could stabilize hydrogen adsorbed on the Co oxides, thus inhibiting the HER and causing enhanced activity of electrocatalytic NO_x⁻ reduction to NH₃.⁸³ Consequently, Co and Co oxides have also received considerable attention for nitrate reduction in recent years.⁸⁴ Hybrid nanoparticles of metallic Co and Co oxides containing Co, Co²⁺ and Co³⁺ were prepared by a simple electrodeposition strategy and the catalysts exhibited excellent NO_x⁻ reduction performance (FE: 100% at -0.2 V) and OER activity (η_{10} = 310 mV).⁸² A zinc nitrate cell assembled from the catalysts demonstrated an outstanding power density of 25 mW cm⁻².

Cobalt phosphide has also been probed extensively for NO_x⁻ reduction.⁸⁵ The introduction of P has been confirmed to improve the stability of the catalysts by mitigating the reconstitution of Co to form Co(OH)₂ during NO₃⁻ reduction, while reducing ΔG_{rxn} of the rate-determining step and optimizing the required energy barriers for the reaction. In addition, the Co 4p orbitals in CoP proved to be engaged directly in the NO₃⁻ adsorption and electron transfer steps. The mechanism of CoP action in NO₃⁻ reduction is shown in Fig. 9c. Initially, empty Co 4p orbitals in CoP are coupled with O 2p orbitals in NO₃⁻ to form the Co-O-N bond for NO₃⁻ adsorption on Co sites. With the employment of a strong electrical field, the Co 4p orbitals obtain excited electrons from the Co 3d orbitals. Subsequently, electrons in the Co 4p orbitals are shifted to O 2p orbitals *via* the Co-O-N covalent bond and infused into the π^* orbitals of NO₃⁻, causing considerably reduced NO₃⁻ adsorption for accelerated reduction to NH₃. The CoP nanoarrays also exhibit promising performance in the NO₂⁻ electroreduction process, reaching a NH₃ yield of 2260.7 ± 51.5 μg h⁻¹ cm⁻² and a FE of 90.0 ± 2.3% at -0.2 V vs. RHE.⁸⁶ DFT calculations have confirmed the dominance of NO₂⁻ electroreduc-

Table 3 Summary of NO_x⁻ reduction performance of Co-based electrocatalysts

Electrocatalyst	Electrolyte	Potential (V vs. RHE)	NH ₃ yield	Faradaic efficiency (%)	Ref.
DM-Co	1 M KOH + 1 M KNO ₃	-0.2	1.2 mmol h ⁻¹ cm ⁻²	100	82
CoOx	0.1 M KOH + 0.1 M KNO ₃	-0.5	82.4 mg h ⁻¹ mg _{cat} ⁻¹	93.4	83
CoP NAs/CFC	1.0 M NaOH + 1.0 M NaNO ₃	-0.3	9.56 mol h ⁻¹ m ⁻²	100	85
Ru _x Co _y alloy	0.1 M KOH + 0.1 M KNO ₃	0	3.21 mol g _{cat} ⁻¹ h ⁻¹	97	87
C/Co ₃ O ₄	50 mM K ₂ SO ₄ + 50 mM KNO ₂	-0.6	4.1 mg h ⁻¹ cm ⁻²	100	12
CoP NA/TM	0.1 M PBS + 500 ppm NO ₂ ⁻	-0.2	2,26 mg h ⁻¹ cm ⁻²	90	86

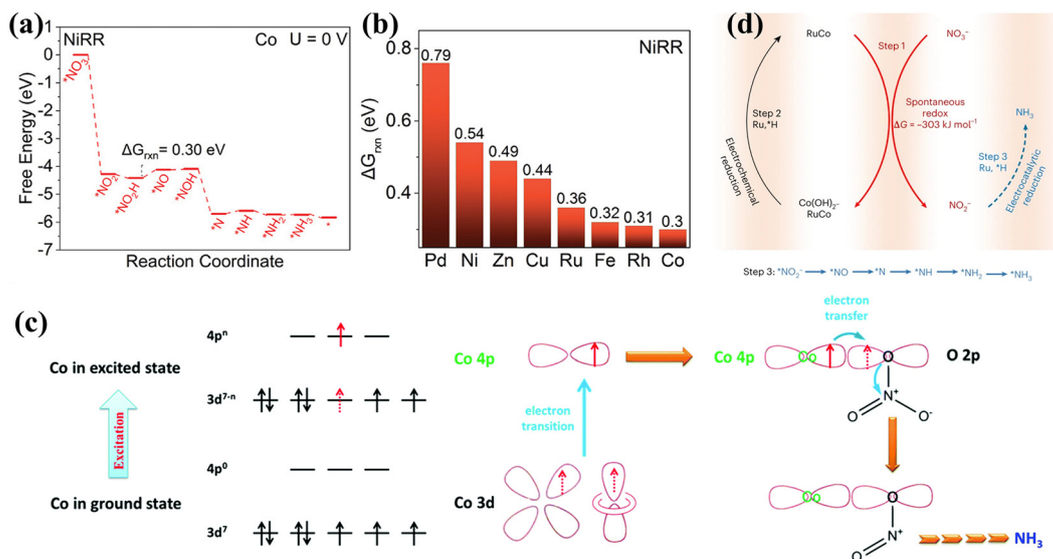


Fig. 9 (a) The Gibbs energy diagram of NO₃⁻ reduction on the Co surface. (b) ΔG_{rxn} on different metals copied with permission.⁸² Copyright 2022, John Wiley and Sons. (c) Mechanism of NO₃⁻ reduction on CoP copied with permission.⁸⁵ Copyright 2022, Royal Society of Chemistry. (d) The three-step relay route of Ru_xCo_y alloy copied with permission.⁸⁷ Copyright 2023, Springer Nature.

tion over the HER, owing to the much smaller adsorption energy (−0.6 eV) of NO₂⁻ on CoP(112) compared to that of H (−0.12 eV).

The doping of P can suppress the reconstruction of Co, while the introduction of Ru can achieve the reconversion of Co(OH)₂ to Co for more efficient and durable ammonia production.⁸⁷ Co and NO₃⁻ are spontaneously converted to form Co(OH)₂ and NO₂⁻, and the conversion of Co(OH)₂ to Co and NO₂⁻ reduction to NH₃ can be facilitated by active hydrogen species. In view of Ru having decent adsorption energies for hydrogen atoms, Ru_xCo_y alloy hollow nanostructures have been developed to achieve NH₃ production with a remarkable energy efficiency of 42 ± 2%. As indicated in Fig. 9d, the existence of Ru facilitates the formation of active hydrogen species, promoting a dynamic Co valence cycle and achieving an innovative three-step relay route involving spontaneous redox reactions, electrochemical reduction and electrocatalytic reduction.

3.5. Ni-based and Ti-based electrocatalysts

Aiming to eliminate competition from the HER in NO_x⁻ reduction, scientists have generally been drawn to metals that are relatively inert in the water splitting process, like Cu.⁸⁸ However, the hydrogenation process of NO_x⁻ reduction is also urgently required for the generation of H* *via* water splitting.

HER-active metals including Ni can facilitate significant H* generation,^{89–91} which can also be promising catalysts for ammonia production after effectively obviating the direct coupling path of H*. The Ni based materials in Table 4 also show promising performance. For example, the self-activated Ni(OH)₂@Ni cathode was tested on a pilot scale for the treatment of wastewater containing nitrate (Fig. 10a) showing no decay in the reaction rate after two months of successive operation (Fig. 10b).⁹² The impressive properties of Ni(OH)₂@Ni are conjectured to stem from inhibiting the HER in the H*-mediated NO₃⁻ reduction pathway, thus promoting the reduction of NO₃⁻ to NH₃.

The construction of grain boundary defect engineering on nickel nanoparticles (GB Ni-NPs) is also well-suited to simultaneously achieving the availability of sufficient H* and suppressing competition from the HER.⁹³ As illustrated in Fig. 10c and d, the increased H* retention potential results from the GB region having a higher energy barrier for H₂ formation from H*. Compared with pristine Ni, abundant H* in GB Ni-NPs is inclined to combine with adjacent adsorbed intermediates for NH₃ production, rather than coupling directly to produce H₂. Furthermore, the appropriate adsorption energies of GB Ni-NPs for key species (Fig. 10e) facilitate the conversion of NO₃^{*} to NO₂^{*} and accelerate the dissociation of NH₃.

Table 4 Summary of NO_x⁻ reduction performance of Ni-based electrocatalysts

Electrocatalyst	Electrolyte	Potential (V vs. RHE)	NH ₃ yield	Faradaic efficiency (%)	Ref.
Ni(OH) ₂ @Ni	0.1 M Na ₂ SO ₄ + 70 mg L ⁻¹ NO ₃ ⁻	3.67	—	95.5	92
GB Ni-NPs	1 M NaOH + 1 M NaNO ₃	−0.93	15.49 mmol h ⁻¹ cm ⁻²	90	93
Ni-TiO ₂	0.1 M NaOH + 0.1 M NO ₂ ⁻	−0.7	727 μmol h ⁻¹ cm ⁻²	90	94
Ni@JBC	0.1 M NaOH + 0.1 M NO ₂ ⁻	−0.5	4.1 mg h ⁻¹ mg _{cat} ⁻¹	83.4	113
Ni-NSA-VN	50 mM K ₂ SO ₄ + 50 mM KNO ₂	−0.6	—	95.5	114

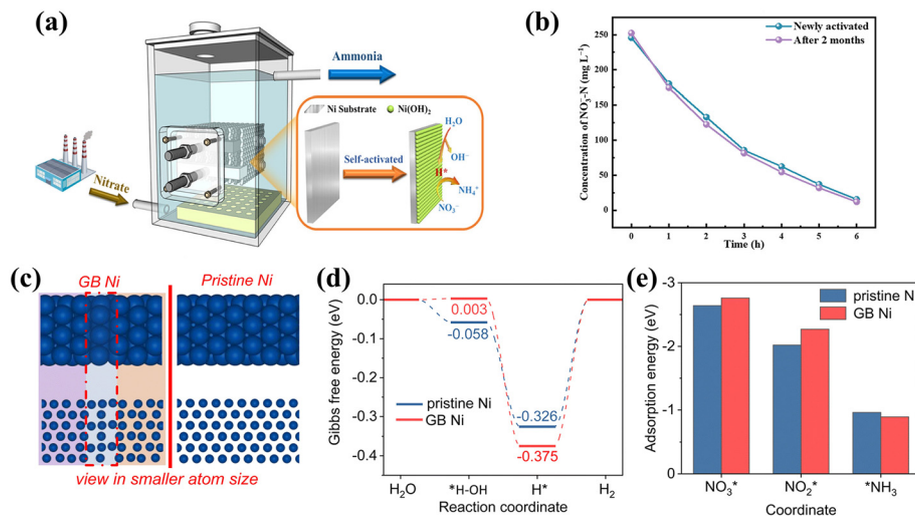


Fig. 10 (a) Diagram of an electrochemical reactor on the pilot scale and (b) comparison of initial NO_3^- reduction performance and that after two months copied with permission.⁹² Copyright 2021, American Chemical Society. (c) The simulated structure, (d) Gibbs free energy diagrams of the HER and (e) adsorption energies of important N-containing species on GB Ni and pristine Ni copied with permission.⁹³ Copyright 2023, Royal Society of Chemistry.

For the design of NO_x^- reduction catalysts, facilitating the hydrogenation process of the intermediate is essential, and the enhanced adsorption of NO_x^- should also not be neglected. The introduction of Ni into TiO_2 can significantly boost the nitrite reduction performance of TiO_2 .⁹⁴ The improved material has an excellent FE as high as 94.89% at -0.5 V and a maximum NH_3 yield reaching $727 \mu\text{mol h}^{-1} \text{cm}^{-2}$ at -0.7 V. Theoretical calculations have confirmed that the introduction of Ni lowers the energy barrier in the critical conversion step of $^*\text{NO}$ to $^*\text{N}$, thus enhancing the adsorption of NO_2^- and promoting the conversion of NO_2^- to NH_3 .

TiO_2 is the most broadly studied Ti-based material due to its high abundance, low cost, non-toxicity and excellent stability, nevertheless it also encounters deficiencies such as poor electrical conductivity and insufficient active sites.^{95,96} A variety of strategies including vacancy engineering,⁹⁷ introducing metal atoms⁹⁸ and non-metal atom doping⁹⁶ have been employed to modify TiO_2 ; these have achieved positive results (Table 5). More interestingly, TiO_2 is also a typical n-type semiconductor with significant chemical and structural stability,⁹⁹ which can be employed for the construction of Mott-Schottky junctions and p-n heterojunctions.

It is often perceived that two oppositely charged regions across the interface can be created by the construction of a

semiconductor junction, along with a strong built-in field, which can modify the atomic interfacial electron density.¹⁰⁰ Mott-Schottky junctions can be formed by combining conductors (metals) with semiconducting materials. Accelerated electronic transfers at the interface and strong built-in electric field formation can be induced by discrepancies in the Fermi energy levels between metals and semiconductors, promoting local charge polarization and significantly ameliorating the adsorption of key intermediates.¹⁰¹

Co@TiO_2 heterostructures exhibit superior NO_3^- reduction performance due to the Schottky interface.¹⁰² The effects of the Schottky junction are mainly manifested in the enhanced electronic conductivity and the improved catalytic selectivity. The mapping of the electron density difference (Fig. 11a) in Co@TiO_2 also confirms electron migration from the Co side to the TiO_2 side, resulting in electron depletion regions on the Co surface. The built-in electric field can be generated due to the accumulation of net charge on the TiO_2 side, promoting the transportation of charge. DFT calculations (Fig. 11b) have also revealed that the Schottky junction of Co@TiO_2 indeed promotes NO_3^- adsorption and inhibits the appearance of by-products including NO_2 , NO , and N_2 .

The p-n junction can also be formed by coupling TiO_2 with a p-type semiconductor, thus inducing charge redistribution

Table 5 Summary of NO_x^- reduction performance of Ti-based electrocatalysts

Electrocatalyst	Electrolyte	Potential (V vs. RHE)	NH_3 yield	Faradaic efficiency (%)	Ref.
$\text{CoP/TiO}_2@\text{TP}$	0.1 M NaOH + 0.1 M NaNO_3	-0.5	$499.8 \mu\text{mol h}^{-1} \text{cm}^{-2}$	95	103
$\text{Co@TiO}_2/\text{TP}$	0.1 M PBS + 0.1 M NO_3^-	-0.93	$800 \mu\text{mol h}^{-1} \text{cm}^{-2}$	96.7	102
Cu-TiO_2	0.1 M Na_2SO_4 + 0.1 M NO_3^-	-0.3	—	84.3	115
TiO_{2-x}	0.5 M Na_2SO_4 + 0.5 M NO_3^-	-0.4	—	85	40
$\text{Cu}_3\text{P@TiO}_2/\text{TP}$	0.1 M NaOH + 0.1 M NaNO_2	-0.7	$1583.4 \mu\text{mol h}^{-1} \text{cm}^{-2}$	97.1	116
V-TiO_2	0.1 M NaOH + 0.1 M NO_2^-	-0.7	$540.8 \mu\text{mol h}^{-1} \text{cm}^{-2}$	93.2	98

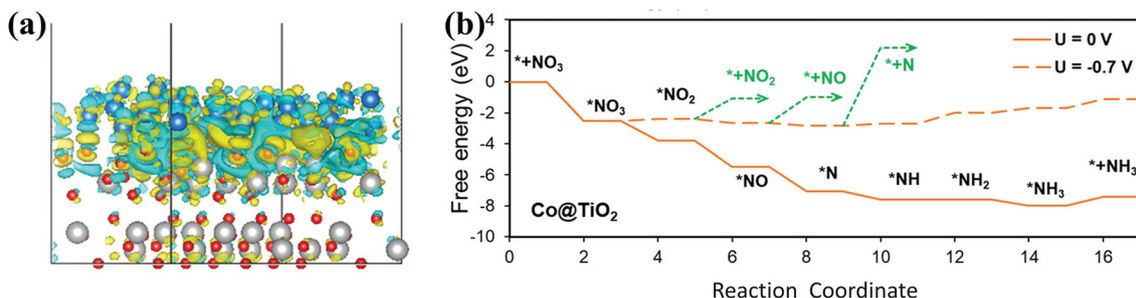


Fig. 11 (a) Mapping of the electron density difference. (b) Free energy diagram for NO_3^- conversion on Co@TiO_2 copied with permission.¹⁰² Copyright 2023, John Wiley and Sons.

in the material and optimising the adsorption free energies of the intermediates. The p-n junction of CoP/TiO_2 heterostructures has been confirmed by experimental and theoretical studies to achieve a lower energy barrier of NO_3^- conversion to $^*\text{NO}_3^-$ and increased energy barriers of NO and NO_2 production, which enhance the NO_3^- reduction activity and optimise the reaction selectivity.¹⁰³

In summary, a variety of strategies for optimizing transition metal-based catalysts have been demonstrated. Alloying and doping are common tactics to modulate the d-band center and intermediate adsorption energies of metals, but it is important to select inexpensive elements to effectively assist transition metals at accelerating NO_x^- reduction. Exposure of specific crystalline surfaces and vacancy engineering are also effective optimization strategies to effectively boost reactant and intermediate adsorption, but obtained materials show limited activity enhancement with undesired stability. The construction of heterostructures can dramatically enhance catalytic activity and selectivity caused by their intriguing physical nature due to redistributed charges and bent energy bands at the heterogeneous interface. The key to constructing superior heterostructures is the selection of coupling materials. The design of chainmail-protected catalysts offers novel insights to significantly improve catalyst stability, but the selection of suitable materials as chainmail and the exploration of gentle synthesis strategies require more attention. In addition to the optimization strategies mentioned above, other factors such as atmosphere and anions in the electrolyte may also considerably contribute to the catalytic activity and selectivity of materials, however, there are few systematic and insightful studies on the contributions of other factors to accelerate NO_x^- reduction.

4. Conclusion and future outlook

Electrocatalytic NO_x^- reduction provides innovative insights into sustainable NH_3 synthesis and eco-friendly NO_x^- removal. The review firstly outlines the mechanism of NO_x^- reduction and the challenges of catalyst design under aqueous conditions, proposing that the following points should be observed when designing electrocatalysts with excellent performance: (1) The adsorptions of NO_3^- and NO_2^- on catalysts

are vital. For NO_3^- reduction, the conversion from NO_3^- to NO_2^- is a decisive step. Suitable adsorptions of NO_3^- and NO_2^- are beneficial for the reaction. (2) It is difficult to escape competition with the HER under aqueous conditions. Suppressing the formation of active hydrogen species is unadvisable to reduce HER competition because active hydrogen species are indispensable reactants for the subsequent hydrogenation process in NO_x^- reduction. The strategy of employing two sites can be used for reference to simultaneously realize active hydrogen species production and reduce the direct coupling of active hydrogen species. In this strategy, one site acts as a proton warehouse to promote proton generation, and the other site temporarily stores active hydrogen species for subsequent hydrogenation reactions. (3) In addition to the HER side reaction, other side reactions generating N_2 or NO should not be ignored. Improved stability of intermediates including $^*\text{NO}_2$ or $^*\text{NOH}$ facilitates more efficient ammonia production.

Then, this paper focuses on widely studied first-row transition metal-based electrocatalysts including Cu , Fe , Co , Ni , Ti -based electrocatalysts, which are characterized by tempting abundance and moderate interactions with the reactants. Recent advances in engineering first-row transition metal-based electrocatalysts are presented in this paper. Hopefully these provide more inspiration for constructing remarkable NO_x^- reduction electrocatalysts. Despite the tremendous advances that have been obtained in currently developed catalysts, a series of challenges still exist in this area and are waiting to be addressed:

(i) Currently, in the field of NO_x^- reduction, the lack of benchmark catalysts makes it difficult to measure the gap between the experimentally prepared materials and industrialization. In addition, the use of isotope labeling to verify the effective transformation of NO_x^- is questionable and lacks standardization in current experiments. Researchers often choose isotopes with low purity in practical experiments, considering that isotopes with high purity will bring excessive experimental cost. Meanwhile, isotope labeling with low purity makes it difficult to exclude the interference of other N species including N_2 .

(ii) Inconsistent descriptions of ammonia production units such as $\text{mg h}^{-1} \text{cm}^{-2}$ and $\text{mmol h}^{-1} \text{mg}_{\text{cat}}^{-1}$ are employed in current articles, which is unfavourable for readers and peers to

judge the performance level of materials. Employing uniform units is more conducive to accelerating the development of NO_x^- electrocatalysts.

(iii) It is common for catalysts to undergo reconfiguration during the catalytic process, but the reconfiguration problem is rarely explored in currently developed NO_x^- reduction electrocatalysts; more advanced characterization techniques are required to be employed in the study of catalytic mechanisms.

(iv) The oxygen evolution reaction (OER) is usually employed as the anode reaction in aqueous catalytic installations for NO_x^- reduction to NH_3 . Replacing the OER with other reactions including Zn oxidation or methanol oxidation can result in more value-added product synthesis or the Zn- NO_x^- cell simultaneously producing NH_3 and electricity. These intriguing replacements can open up novel insights into the diverse applications of NO_x^- reduction.

(v) In addition to the development of catalysts with superior performance, electricity prices are also worth considering for promoting the realization of large-scale electrocatalytic NO_x^- reduction to NH_3 . Much lower electricity prices may come with the development of alternative energy sources such as plasma and nuclear fusion.

(vi) A stable NO_x^- source has long been a topic of concern. It is necessary to consider the complex composition of wastewater employed as a source. Several studies have confirmed the significant influences of ion concentration, other inorganic ions, and pH on nitrate reduction in wastewater.

(vii) When selecting wastewater as source of NO_x^- ions, complex composition of wastewater should be considered. Other ions and pH interference cannot be ignored, from the current situation, the cost of enrichment wastewater is also necessary to consider. Direct synthesis through electrocatalytic nitrogen oxidation also faces the challenge of developing highly active nitrogen oxidation catalysts.

(viii) Although the majority of catalysts currently developed exhibit decent stability, most of them are tested at the experimental scale. It is necessary to carry out stability tests on catalysts at the pilot scale or larger scale.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

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