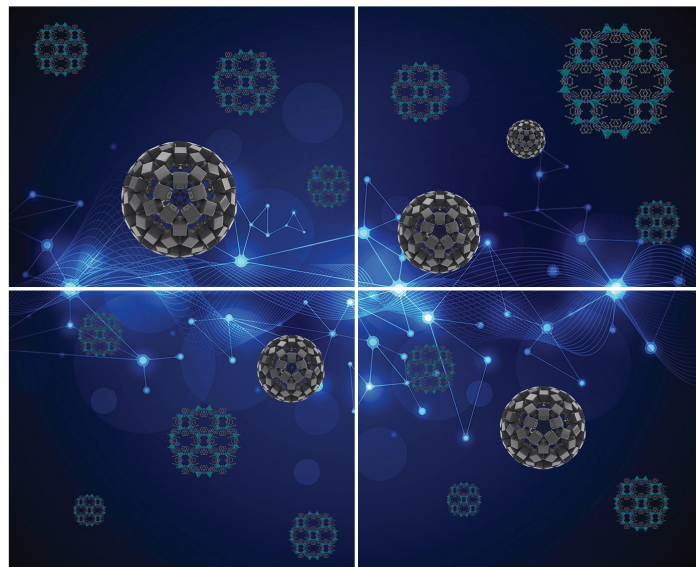


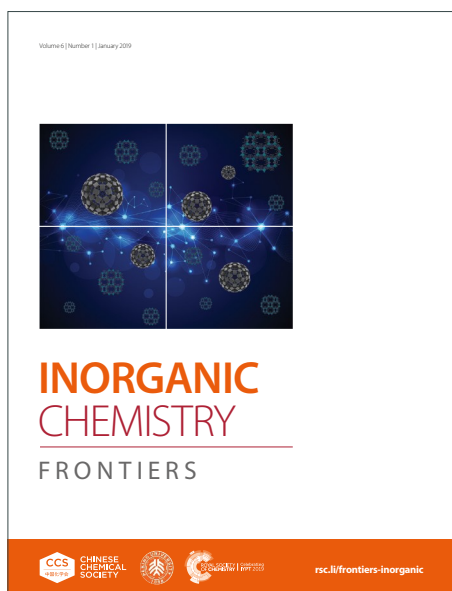
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ARTICLE

Catalyst Design Strategies for NO_x-Involved Electrocatalytic C-N Coupling ReactionsRuhan Wang,^{a,b} Xiaofu Sun^{*a,b} and Buxing Han^{*a,b}Received 00th January 20xx,
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Electrocatalytic C-N coupling reaction involving NO_x species (NO₃⁻, NO₂⁻, NO) have emerged as a sustainable approach for synthesizing high-value nitrogen-containing chemicals. This review provides a comprehensive overview of recent advances in catalyst design strategies for enhancing the efficiency and selectivity of NO_x-involved C-N bond formation. Five key strategies are categorized and discussed: defect engineering, coordination environment design, interface engineering, dual-site synergy, and emerging architectures. For each strategy, representative literature cases are summarized to illustrate mechanistic insights and practical applications. By addressing challenges such as intermediate instability, low selectivity, and competing side reactions, these strategies demonstrate great potential for advancing electrocatalytic C-N coupling toward practical implementation. Finally, future directions are also proposed, including dynamic catalyst design, microenvironment regulation, and data-driven catalyst screening.

1 Introduction

In the context of accelerating the global transition toward sustainable chemistry, electrocatalysis and electrosynthesis have garnered extensive attention as green, efficient, and versatile platforms¹⁻³ for producing high-value chemicals under mild ambient conditions⁴⁻⁹. These processes leverage renewable electricity and water as energy sources and benign hydrogen sources, providing an efficient and clean alternative to traditional thermochemical methods^{10, 11}. Over the past several decades, significant advances have been made in the electrocatalytic transformation of abundant and renewable small-molecule feedstocks (such as CO₂, N₂ and NO₃⁻) into a wide energy-rich/functionalized compounds^{9, 12-22}. Beyond the independent transformation of individual small molecules, increasing attention has recently shifted toward electrocatalytic strategies that integrate multiple substrates (particularly nitrogen and carbon sources) into a single reaction framework²³⁻³¹.

Electrocatalytic C-N coupling reactions, particularly involving nitrogen oxides (NO_x, including NO, NO₂⁻, and NO₃⁻)³²⁻³⁶, have emerged as a transformative research frontier in green chemical synthesis³⁷⁻⁴⁰. Unlike conventional thermal catalytic processes that typically require harsh reaction conditions⁴¹, high energy inputs¹⁹, electrocatalytic pathway leverage renewable electricity to directly couple reactive nitrogen

species with diverse carbon-based substrates, including CO₂ and its electroreduction-derived intermediates, aldehydes and ketones, and biomass-derived active intermediates, which offering a sustainable alternative to traditional thermochemical approaches. The resulting products encompass a broad spectrum of important chemicals, such as oximes⁴²⁻⁴⁴, amino acids⁴⁵⁻⁴⁷, amines⁴⁸⁻⁵⁰, amides⁵¹⁻⁵³, urea⁵⁴⁻⁵⁷ and heterocyclic compounds (Fig. 1), which are essential intermediates in agriculture, pharmaceuticals, materials, and environmental remediation⁵⁸.

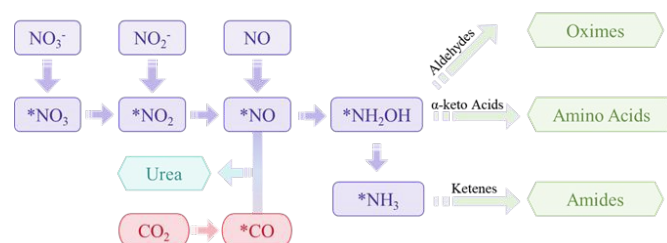


Figure 1. Reaction pathways for NO_x-involved C-N coupling.

Among these versatile transformations, several representative NO_x-involved C-N coupling reactions have been intensively studied. For instance, the reductive amination reactions involving nitrate or nitrite and carbonyl substrates afford high-value oximes, amines, amino acids and etc⁵⁹⁻⁶¹. Similarly, electrochemical synthesis of urea from CO₂ and NO_x provides an attractive route to sustainably produce urea fertilizers⁶². These C-N coupling reactions expanding the scope of green electrosynthesis to complex organic molecules traditionally synthesized by multi-step pathways⁶³⁻⁶⁷.

In recent years, substantial progress has been made in identifying suitable substrates and demonstrating the feasibility of electrochemical C-N coupling reactions⁶⁸. Meanwhile, aided by advances in theoretical modelling, computational

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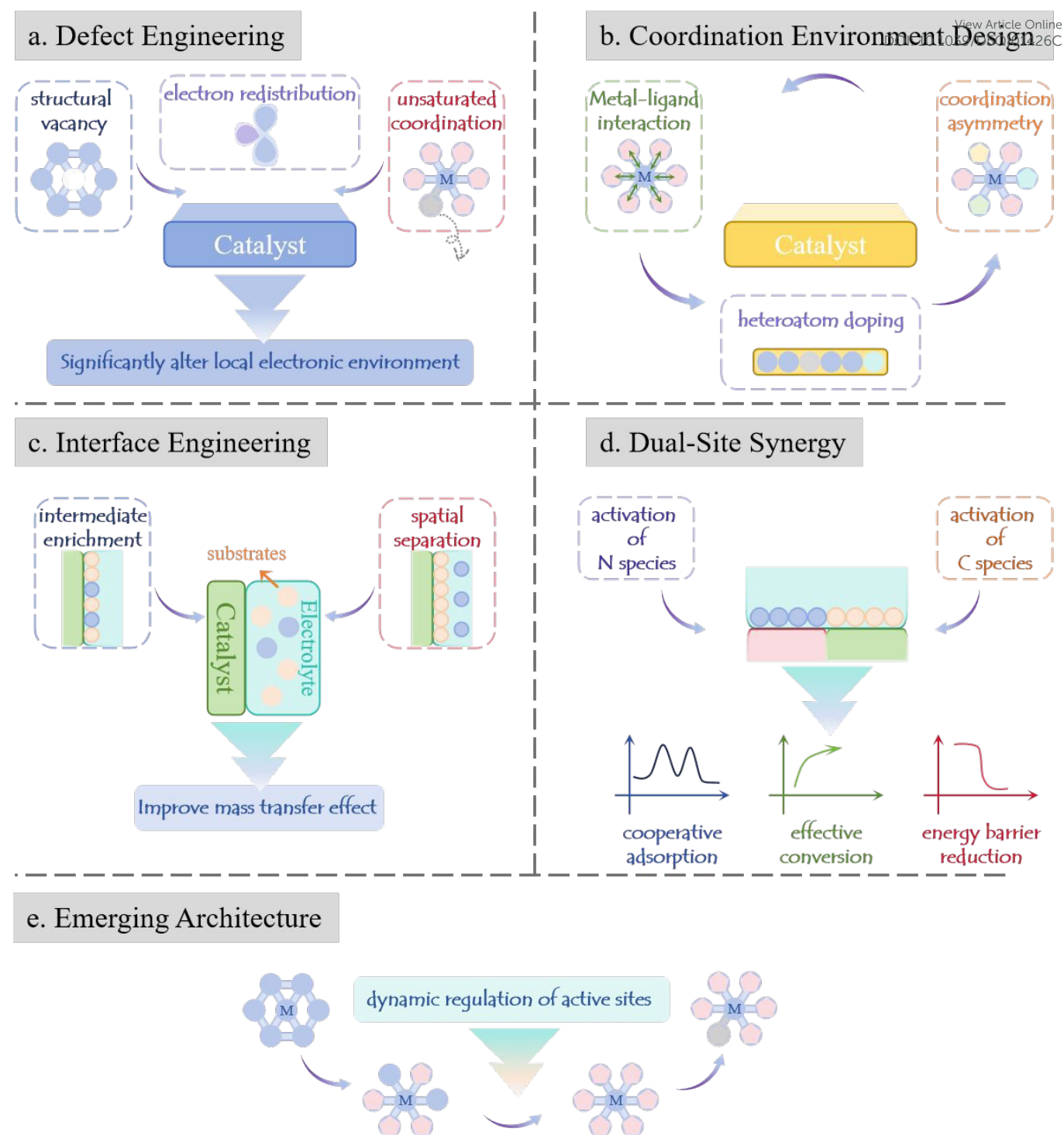


Figure 2. Schematic of five catalyst design strategies for electrocatalytic C-N coupling. (a) Defect engineering: Modulating local electronic environments via structural vacancies, unsaturated coordination, and electron redistribution. **(b) Coordination environment design:** Optimizing active sites through metal-ligand interactions and asymmetric coordination. **(c) Interface engineering:** Enhancing mass transfer by spatial separation and substrate-enriched microenvironments. **(d) Dual-site synergy:** Cooperatively activating N/C species to reduce C-N coupling barriers. **(e) Emerging architectures:** Dynamically regulating active sites for adaptive reaction pathways. Collectively, these strategies address bottlenecks in electron transfer, mass transport, and dynamic instability during NO_x conversion.

simulations and in situ/operando characterization techniques, electrochemical catalysis has enabled researchers to gain deeper insights into the composition and configuration of key intermediates, evaluating their binding affinities, and regulate catalytic activity and product selectivity within these systems⁶⁹⁻⁷⁶. However, due to the multi-step electron/proton transfer

processes, as well as various reactive chemical bonds, active intermediates, and competing reaction pathways, which make these reactions become more complex^{32, 71, 77}. A central challenge lies in controlling reaction selectivity and preventing side reactions such as hydrogen evolution reaction (HER), over-hydrogenation, or undesired side-reaction⁷⁸⁻⁸¹. Consequently,



minor fluctuations in the electrode surface properties⁸²⁻⁸⁵, active intermediates binding energies⁸⁶, or local reaction microenvironments⁸⁷⁻⁸⁹ can drastically alter product distributions and overall reaction efficiency^{8, 71}. Furthermore, achieving simultaneous optimal adsorption and activation of disparate reactants (such as NO_x and carbon sources) poses significant challenges⁹⁰⁻⁹². Different reactants typically require different binding sites and distinct electronic environments, making it difficult to align reaction pathways on single-site or homogeneous catalyst surfaces⁹³. At the same time, conventional catalyst designs rely on static structures, which fail to accommodate the dynamic structural and electronic changes occurring during electrocatalysis⁹⁴. If these mechanistic and catalytic challenges are not effectively solved, it would severely hinder the applicability of electrocatalytic C-N coupling processes transition from laboratory-scale demonstrations to practical, large-scale applications.

To overcome these intrinsic mechanistic limitations and to further enhance the performance of electrocatalytic NO_x-involved C-N coupling reactions, researchers have proposed five representative catalyst engineering strategies (**Fig. 2**), including defect engineering, coordination environment design, interface engineering, dual-site synergy and emerging architectures. By resolving challenges such as the instability of reactive intermediates, the occurrence of competing side reactions, and the mismatch in substrate adsorption requirements, these approaches collectively enhance the effectiveness of C-N bond construction under electrochemical conditions.

In this paper, we provide a comprehensive overview of electrocatalytic C-N coupling reactions involving NO_x species from the perspective of catalyst design strategies. We systematically categorize recent advances into five major approaches: defect engineering, coordination environment design, interface engineering, dual-site synergy, and emerging architectures. These strategies can regulate structural and electronic characteristics, helping to resolve key issues in intermediate stabilization, substrate adsorption and reaction selectivity. In addition, the last section outlines the challenges and development prospects for NO_x-involved C-N coupling. We believe that this review can inspire new exploration and principles for catalyst design for electrocatalytic C-N coupling reactions.

2 Catalyst Design Strategies for NO_x-Involved C-N Coupling

Defect engineering represents a fundamental approach wherein deliberate introduction of structural vacancies, unsaturated coordination sites, or electronic redistributions significantly alters local electronic environments⁶⁰ of catalysts (**Fig. 2a**). These engineered defects can act as highly active adsorption and activation centers, precisely tune the binding energy of key intermediates and lower activation barriers for critical C-N coupling steps. Such precise electronic control at defect sites provides an effective solution for stabilizing the

elusive intermediates, thereby significantly improving overall catalytic selectivity and efficiency. DOI: 10.1039/D5QI01426C

The second category is coordination environment design, emphasizing the adjustment of the local atomic arrangement around the active metal center (**Fig. 2b**). Through rational tuning of metal-ligand interactions, heteroatom incorporation, and coordination asymmetry, catalysts are able to influence intermediate adsorption behavior and energy barriers, thereby promoting more selective and efficient C-N bond formation⁹⁵. Interface engineering, as the third major strategy, focuses on regulating the physicochemical properties at the boundary between different phases to construct favorable local microenvironments at the interface between the electrode and electrolyte (**Fig. 2c**). By tuning interfacial characteristic such as surface polarity, wettability, and spatial distribution of reactants, this approach can facilitate selective intermediate enrichment, improve mass transport, and spatially separate reaction steps⁴⁴. As a result, interface engineering electrocatalytic systems often demonstrate improved reaction selectivity, enhanced stability, and better compatibility with complex multi-step C-N coupling processes.

Dual-site synergy enhancing the performance of NO_x-involved C-N coupling reactions by integrating two types of catalytic active centers with complementary effects in a single system (**Fig. 2d**). Typically, one site promotes the activation of nitrogen-containing species, while the other is responsible for the activation of carbon-containing species⁹⁶. The spatial proximity and electronic complementarity between the two sites promote synergistic adsorption, efficient intermediate transfer, and selective C-N bond formation⁹⁷. This cooperative mechanism improves overall reaction coordination, lowers energy barrier, and helps to suppress undesirable side reactions, thereby contributing to improved activity and selectivity in complex multistep transformations.

Emerging architectures introduce a new design perspective, emphasizing the adaptability and responsiveness (**Fig. 2e**). Rather than relying solely on rigid structural parameters, this strategy focuses on dynamic regulation of active sites and structural evolution during the reaction process⁹⁸, which offering opportunities to balance activity and selectivity across complex pathways and open up new directions for intelligent, programmable and multi-step coupling catalytic platforms.

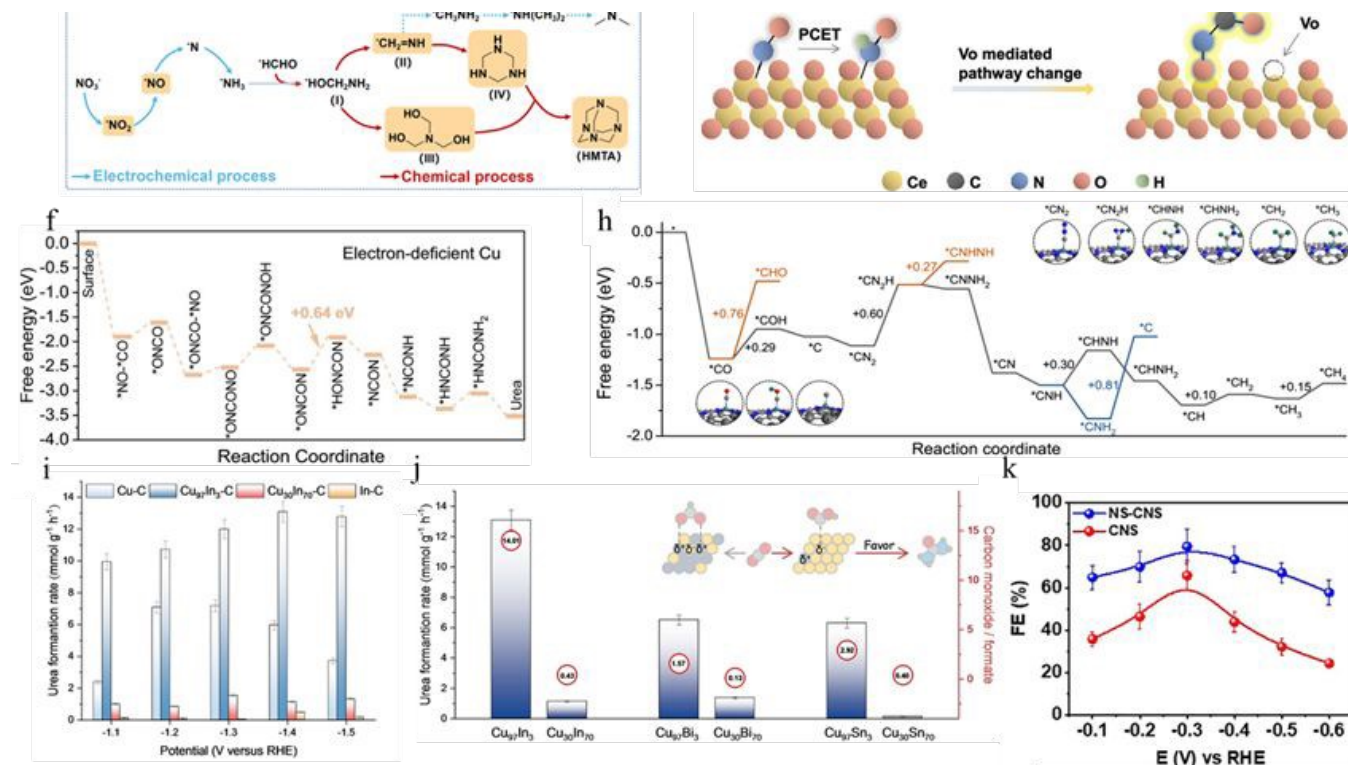
3 Representative Cases in the Literature

3.1 Defect Engineering

Defect engineering offers a powerful strategy to tailor the local electronic environment of electrocatalysts by introducing vacancies, unsaturated coordination sites, and charge redistributions. These defect-induced modulations profoundly influence the adsorption configurations, intermediate stabilization, and activation barriers associated with NO_x and carbon-based species. Particularly, defect sites can serve either as highly active centers for C-N bond formation or as electronic reservoirs that stabilize key intermediates, thereby suppressing



Figure 3. (a) In situ Fe K-edge XANES spectra of AD-Fe/NC under applied potentials from OCV to -1.0 V vs. RHE. The white-line intensity attenuation at -1.0 V indicates Fe valence reduction during reductive amination⁶⁰. Reproduced with permission from ref. 60, Copyright © 2023 Wiley-VCH GmbH. (b) Gibbs free energy change (ΔG) for CO₂ reduction to *OCHO intermediate and subsequent C-N coupling on Pd/Cu-V_{Cu}⁹⁹. (c) Comparative Gibbs free energy profiles for DMF synthesis on pure Cu, Cu-V_{Cu}, and Pd/Cu-V_{Cu} surfaces⁹⁹. Reproduced with permission from ref. 99, Copyright © 2024 Elsevier Inc. (d) Electron paramagnetic resonance (EPR) spectra of c-OD-Cu and. The distinct signal at $g = 1.997$ for e-OD-Cu indicates the presence of unpaired electrons trapped at copper vacancy sites, while no significant signal is observed for c-OD-Cu¹⁰⁰. (e) Schematic illustration of the tandem electrochemical-chemical pathway for electrocatalytic HMTA synthesis from nitrate and formaldehyde¹⁰⁰. Reproduced with permission from ref. 100, Copyright © 2024 American Chemical Society. (f) Free energy profiles for urea synthesis pathways on electron-deficient Cu (Cu/PI-500)¹⁰¹. Reproduced with permission from ref. 101, Copyright © 2024 Wiley-VCH GmbH. (g) Schematic illustration of oxygen vacancy (V_O)-mediated reaction pathway switching on CeO₂¹⁰². Reproduced with permission from ref. 102, Copyright © 2022, American Chemical Society. (h) Free energy profile for nitrogen reduction reaction (NRR) elementary steps on *C-modified ReMn-NC¹⁰¹. Reproduced with permission from ref. 101, Copyright © 2024 Elsevier Inc. (i) Urea yield rates under applied potential for Cu-C, Cu₉₇In₃-C, Cu₃₀In₇₀-C, and In-C electrocatalysts⁶². (j) Urea yield rates and CO/HCOO⁻ molar ratios across bimetallic Cu-In, Cu-Bi, and Cu-Sn electrocatalysts⁶². Reproduced with permission from ref. 62, Copyright © 2023 Wiley-VCH GmbH. (k) FE of the production for NS-CNS (blue), CNS (red)¹⁰⁴. Reproduced with permission from ref. 104, Copyright © 2023 Elsevier.



competing reactions and steering the reaction toward selective coupling pathways. By precisely tuning the type, density, and spatial distribution of defects, recent studies have demonstrated enhanced coupling efficiency, optimized reaction energetics, and improved selectivity across various electrocatalytic systems for value-added nitrogenous product synthesis.

Xian and co-workers developed a catalyst based on atomically dispersed Fe-N₄ sites anchored in a nitrogen-doped carbon matrix (AD-Fe/NC), aiming to achieve precise control over amino acid electrosynthesis via defect-engineered local electronic redistribution⁶⁰. In the AD-Fe/NC catalyst, the Fe-N₄ centers act as electron-deficient sites, where potential-driven orbital rearrangement enhances NO adsorption and facilitates the formation of key intermediates (**Fig. 3a**), thereby promoting the C-N coupling step. Under an applied potential of -0.6 V vs. reversible hydrogen electrode (RHE), the catalyst achieved a valine yield of 32.1 $\mu\text{mol mg}_{\text{cat}}^{-1}$ with a selectivity of 11.3%, demonstrating excellent capability for the high-value conversion of nitrogen oxides.

Fan et al. developed a Pd/Cu-V_{Cu} electrocatalyst based on defect engineering by introducing Cu vacancies to modulated the local electronic structure of Cu nanosheets⁹⁹, which exhibited

remarkable activity for the electrochemical synthesis of N,N-dimethylformamide (DMF) from CO₂ and dimethylamine (DMA). The introduction of Cu vacancies (Cu-V_{Cu}) into Cu nanosheets modulated the local electronic structure, significantly enhancing CO₂ adsorption and facilitating spontaneous coupling with DMA to form the C-N bonds (**Fig. 3b**). Concurrently, Pd nanoparticles accelerated the electrochemical reduction of the key intermediate *OCN(CH₃)₂OH to OCHN(CH₃)₂OH, thereby markedly improving the overall efficiency of DMF formation (**Fig. 3c**). Under an applied potential of -0.6 V vs. RHE, the catalyst achieved a high DMF yield of 385 $\text{mmol h}^{-1} \text{g}_{\text{cat}}^{-1}$ and a Faradaic efficiency (FE) of 37.5%. Density functional theory (DFT) calculations further revealed that the introduction of copper vacancies led to electronic redistribution, which reduced the energy barriers for C-N bond formation, facilitated proton-coupled electron transfer, and enhanced the kinetics of intermediate hydrogenation.

Pan and co-authors developed a vacancy-rich electrochemically oxidation-derived copper (e-OD-Cu) for the efficient synthesis of hexamethylenetetramine (HMTA) via electrochemical C-N coupling between nitrate and formaldehyde¹⁰⁰. The presence of Cu vacancies significantly enhanced the adsorption of NO₃⁻ and HCHO, thereby facilitating the initial steps of the reaction



cascade (**Fig. 3d**). The nitrate was first reduced to $^*\text{NH}_3$ through a multi-electron pathway, which then reacted with $^*\text{HCHO}$ to form imine-type intermediates (**Fig. 3e**). Notably, the Cu vacancies played a critical role in suppressing the over hydrogenation of $^*\text{CH}_2=\text{NH}$ intermediates, effectively steering the reaction away from undesired byproducts and enabling the progressive formation of multi-substituted amine intermediates. This promoted the tandem condensation and ring-closing reactions that ultimately yielded HMTA with a FE of 74.9% and a yield of 76.8% at -0.30 V vs. RHE.

Wang et al. constructed electron-deficient Cu sites via strong metal-polyimide (PI) semiconductor interactions (named Cu/PI-X catalysts, where X represents the thermal treatment temperature on PI), enabling efficient urea synthesis through the electrocatalytic coupling of nitrate and carbon dioxide¹⁰¹. The electron-deficient Cu/PI-500 significantly enhanced co-adsorption of $^*\text{NO}$ and $^*\text{CO}$ intermediates and stabilized key intermediates (e.g. $^*\text{ONCONO}$), thus reducing the reaction energy barriers for sequential C-N bond formation (**Fig. 3f**). Cu/PI-500 delivered a high urea yield rate of $255.0\text{ mmol h}^{-1}\text{ g}^{-1}$ and a FE of 14.3% at -1.4 V vs. RHE, with outstanding electrochemical durability.

Wei et al. developed a CeO_2 -supported Cu catalyst enriched with oxygen vacancies (V_O - CeO_2 -750) to enable the electrocatalytic coupling of nitrate and carbon dioxide for urea synthesis¹⁰². The introduction of Vo sites significantly reshaped the local electronic structure of CeO_2 , enabling strong stabilization of the $^*\text{NO}$ intermediate and favoring its coupling with $^*\text{CO}$ to form the key $^*\text{OCNO}$ species (**Fig. 3g**). By anchoring $^*\text{NO}$ intermediates via V_O , the catalyst raises $^*\text{NO}$ hydrogenation barriers by 0.12 eV while reducing $^*\text{NO}$ - $^*\text{CO}$ coupling barriers to 0.27 eV (48% decrease). As a result, the catalyst achieved a high urea yield of $8.81\text{ mmol h}^{-1}\text{ g}_{\text{cat}}^{-1}$ and a FE of 12.2% at -0.7 V vs. RHE.

Zhang and co-workers proposed a “Janus C-N coupling” strategy for the selective synthesis of ammonia and urea under ambient conditions, using nitrogen, carbon dioxide, water and electricity as feedstocks¹⁰³. Specifically, a Re-Mn dual-atomic electrocatalyst (ReMn-NC) with strong CO adsorption was constructed, facilitating the formation of $^*\text{C}$ intermediates, which acting as electron reservoirs. These electron-rich $^*\text{C}$ species promoted end-on adsorption and subsequent C-N coupling with nitrogen ($\text{N}_2 + ^*\text{C} \rightarrow ^*\text{CN}_2$, **Fig. 3h**). Subsequent hydrogenation of $^*\text{CN}_2$, accompanied by C-N dissociation, selectively produced ammonia at a remarkable yield of $48.9\text{ mg g}^{-1}\text{ h}^{-1}$ and a nitrogen selectivity of 98.4%. Conversely, a Zn-Mn dual-atomic electrocatalyst (ZnMn-NC) exhibiting low Co binding strength facilitated the release and migration of CO, enabling its coupling with adsorbed nitrogen ($\text{CO} + ^*\text{N}=\text{N}^* \rightarrow ^*\text{NCON}^*$). Subsequent hydrogenation of nitrogen atoms efficiently yielded urea, achieving an average yield rate of $36.7\text{ mg g}^{-1}\text{ h}^{-1}$ and nitrogen selectivity of 89.1%. Notably, modulating the local electronic states of catalysts significantly influences the electronic density distribution at the active sites, directly affecting the stability and activity of adsorbed species, thereby facilitating the synergistic development of nitrogen reduction and C-N coupling reactions.

Beyond the aforementioned strategies, modulation of local electronic states (such as redistribution of localized charge density) constitutes defect engineering at the electronic-structure level. By profoundly altering the adsorption configurations of intermediates and steering catalytic reaction pathways, this strategy enables efficient electrocatalytic synthesis of target chemicals. Liu et al. demonstrated efficient electrocatalytic urea synthesis under ambient conditions via tuning the local electronic state of bimetallic electrocatalysts using CO_2 and NO_3^- reactant⁶². They found that the localized surface charge of catalysts significantly influenced the adsorption configurations of CO_2 intermediates and subsequent reaction pathways. Specifically, the negatively charged $\text{Cu}_{97}\text{In}_{3-\text{c}}$ surface preferentially induced the formation of C-bound $^*\text{COOH}$ intermediates, greatly enhancing the subsequent C-N coupling reactions and resulting in an impressive urea yield rate of $13.1\text{ mmol g}^{-1}\text{ h}^{-1}$, which is about 13 times higher than the positively charged $\text{Cu}_{30}\text{In}_{70-\text{c}}$ surface (dominated by O-bound $^*\text{OCHO}$ intermediates, shown in **Fig. 3i**). Mechanistic studies revealed that negatively charged surfaces facilitated stable adsorption of reactive intermediates through the formation of C-bound configurations, significantly improving urea synthesis performance. Conversely, positively charged surfaces promoted O-bound intermediate formation, creating a dead-end for subsequent C-N coupling and thus substantially reducing urea yield (**Fig. 3j**). This precise regulation of adsorption configurations and reaction pathways via localized electronic state tuning exemplifies defect engineering's critical role in electrocatalytic urea synthesis.

Jia et al. utilized a defect engineering strategy by developing nitrogen and sulfur co-doped lignin-derived carbon nanosheets (NS-CNS) for the efficient electrocatalytic reductive amination of pyruvate to alanine¹⁰⁴. Co-doping with nitrogen and sulfur effectively modulated the localized electronic states on the catalyst surface, significantly enhancing the local electron density distribution at carbon active sites, thereby improving adsorption affinity toward critical imine intermediates and lowering the reaction energy barrier. Specifically, the localized electron-rich sites facilitated the rapid reduction of imine intermediates to alanine while simultaneously suppressing competing side reactions, such as hydrogen evolution. Under optimized reaction conditions (-0.3 V vs. RHE), the NS-CNS catalyst exhibited an impressive FE of 79.5% with an alanine yield reaching up to $199\text{ mmol h}^{-1}\text{ cm}^{-2}$ and a selectivity exceeding 99.9% (**Fig. 3k**). Additionally, the catalyst demonstrated remarkable stability for long-term electrocatalytic processes and showed excellent potential for practical applications, as evidenced by successfully converting real-world polylactic acid waste into value-added alanine with selectivity greater than 75%.

3.2 Coordination Environment Design

The coordination environment surrounding active metal centers plays a fundamental role in determining the activity, selectivity, and stability of electrocatalysts involved in NO_x-related C-N coupling reactions. Rather than merely relying on



the elemental composition of catalysts, recent advances have demonstrated that adjusting the coordination configuration, such as through heteroatom incorporation, low-coordination site construction, and asymmetry induction, which can effectively tune the local electronic structure and reactivity. These coordination design strategies regulate the adsorption behavior of NO_x species and stabilize key nitrogen-containing intermediates such as *NH₂OH and oximes, while suppressing side processes like HER and over-reduction. By leveraging diverse coordination motifs across nanostructured, alloyed, and molecular catalyst systems, researchers have created favorable reaction microenvironments that promote efficient multi-electron transfer and selective C-N bond formation. The following section highlights several representative studies where coordination environment engineering serves as a powerful means to guide catalytic transformations at the atomic level.

Kang and co-workers proposed an interfacial coordination strategy by anchoring a Zn-based metal-organic framework (ZIF-7) onto carboxyl-functionalized graphite felt (CGF), which named ZIF-7/CGF⁴³, where Zn-O bridges formed between the MOF and the substrate. This modification induced a transformation of the Zn coordination environment from a symmetric Zn-N₄ tetrahedral configuration to an asymmetric Zn-N₃O structure. In situ XAFS analysis revealed that the Zn sites underwent in situ reconstruction, characterized by a reduction in Zn coordination number from 4.0 to 3.3 and an increase in the average Zn-ligand bond length (**Fig. 4a**). Compared with the conventional Zn-N₄ sites, the Zn-N₃O configuration not only enhanced NO adsorption capacity but also facilitated the formation and desorption of the NH₂OH intermediate, thereby promoting the key C-N coupling pathway.

Wu et al. reported a sulfur-modified copper electrocatalyst (Cu-S) that leverages interface engineering to precisely regulate the surface reaction environment for electrochemical C-N coupling between nitrite and cyclohexanone, which attaining a remarkable product selectivity of 99% and yield of 92% at -0.9 V vs. Ag/AgCl⁴². The superior performance originated from the tailored Cu-S coordination structure, which modulated the electronic configuration of the Cu active sites. The appearance of the Cu-S configuration weakened the over-reduction of NH₂OH*. In situ ATR-SEIRAS spectroscopy (**Fig. 4b**) further demonstrated that NH₂OH* remains stably adsorbed on the surface and undergoes direct coupling with cyclohexanone, enabling selective oxime formation.

Sheng et al. constructed a porous high-entropy alloy metallene catalyst (HEA-PdCuAgBiInene) that demonstrates an effective coordination environment modulation strategy for promoting C-N coupling electrosynthesis⁹⁵. The incorporation of p-block metals (Bi, In) into the PdCuAg matrix induces unconventional p-d orbital hybridization, which leads to local electron enrichment and d-state localization at Pd active sites (**Fig. 4c**). This tailored coordination environment fine-tunes the adsorption energies of hydrogen and NO₂-derived intermediates, suppresses overhydrogenation of NH₂OH*, and facilitates its selective condensation with surface-enriched cyclohexanone (C₆H₁₀O*). The HEA catalyst achieves a FE of 47.6%

and nearly 100% yield under mild electrochemical conditions (-0.9 V vs. Ag/AgCl), offering a compelling example of active-site engineering via orbital-level coordination control.

Xiang et al. developed an aluminum-containing nanofiber membrane catalyst (Al-NFM), derived from NH₂-MIL-53(Al)¹⁰⁵, for the electrosynthesis of pyridine oximes via in situ reduction of NO₂⁻ to NH₂OH*. The catalyst possesses a highly disordered carbon structure with abundant defect sites and a mixture of six-, five-, and four-coordinated Al centers, of which approximately 20% are coordinatively unsaturated. These Al-N motifs modulate the local electronic environment, promote NO₂⁻ activation, and stabilize NH₂OH* intermediates, as supported by charge distribution analysis and energy profile calculations (**Fig. 4d**). The absence of Al or nitrogen doping led to significantly lower NH₂OH yields and reduced product selectivity, underscoring the essential role of coordination tuning. Operating at -0.9 V vs. Ag/AgCl under ambient conditions, Al-NFM achieves a FE of 42.1% and 92.1% selectivity. Wang and co-authors reported a one-pot electrosynthesis strategy for the production of oxime ether by NO_x reduction and aldehyde activation using ultrafine MgO nanoparticles embedded in nitrogen-doped carbon nanofiber membranes (MgO-SCM)¹⁰⁶. The electrosynthesis process targets the efficient one-pot conversion of NO_x and 4-cyanobenzaldehyde (4-CB) to 4-cyanobenzaldoxime (4-CBO), and subsequently to its oxime ether (4-CBOE), under ambient conditions, shown in **Fig. 4e**. In this system, the catalyst is derived from Mg-MOFs, and the Mg-O coordination structure was largely preserved during thermal decomposition, forming well-dispersed MgO active sites. This preserved coordination environment generated



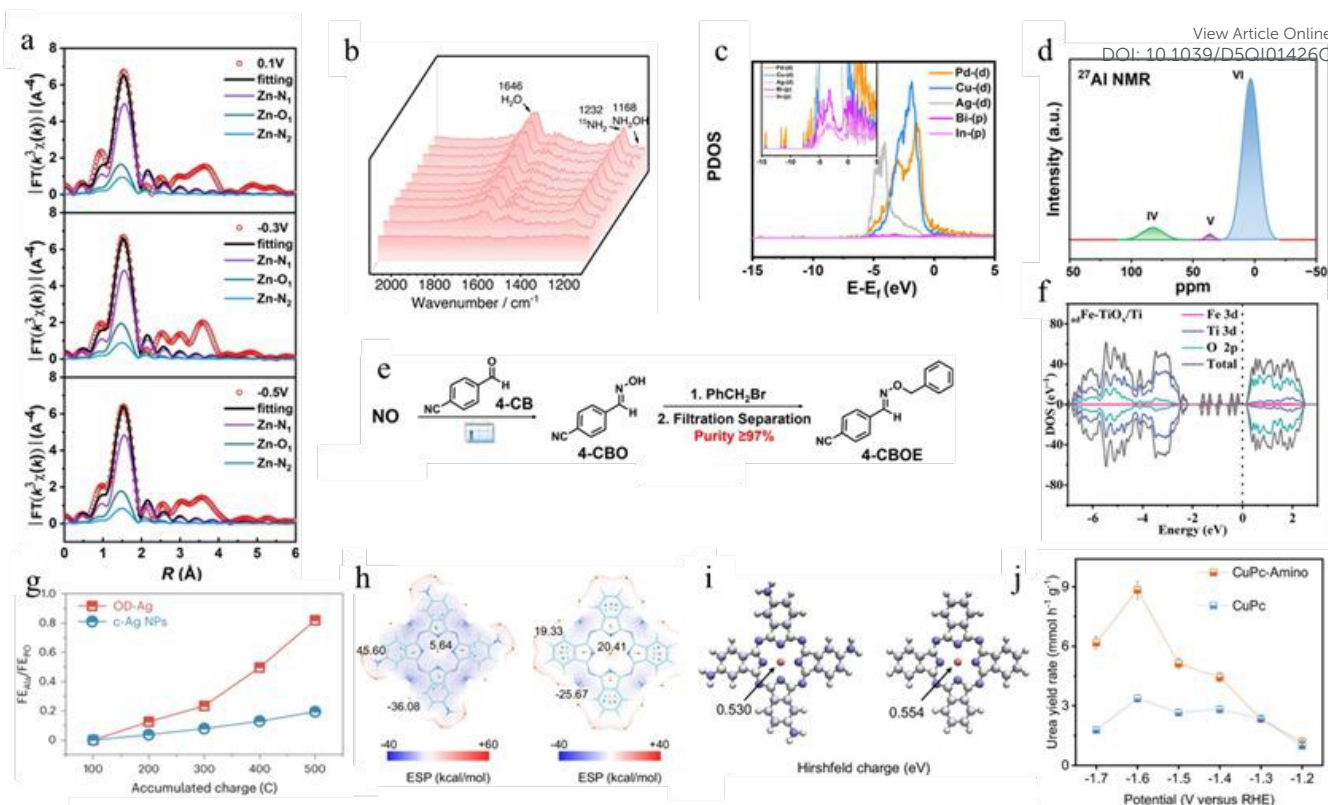


Figure 4. (a) The fitting curves recorded at the Zn K-edge under working conditions⁴³. Reproduced with permission from ref. 43, Copyright © 2025 Springer Nature. (b) Time-dependent in situ ATR-SEIRAS was carried out with isotopically labeled $^{15}\text{NO}_2$ as the N source and H_2O as the H donor⁴². Reproduced with permission from ref. 42, Copyright © 2023 Springer Nature. (c) PDOS of HEA-PdCuAgBiIn with p-d orbital hybridization inset⁹⁵. Reproduced with permission from ref. 95, Copyright © 2024 Wiley-VCH GmbH. (d) Distribution of aluminum coordination states in Al-NFM¹⁰⁵. Reproduced with permission from ref. 105, Copyright © 2023 Wiley-VCH GmbH. (e) Schematic diagram of synthesizing 4-CBOE from NO and 4-CB¹⁰⁶. Reproduced with permission from ref. 106, Copyright © 2024 Wiley-VCH GmbH. (f) Projected density of states (PDOS) and total density of states (TDOS) plots of $\text{adFe-TiO}_x/\text{Ti}$ ¹⁰⁷. Reproduced with permission from ref. 107, Copyright © 2024 Wiley-VCH GmbH. (g) Comparison of $\text{FE}_{\text{Ala}}/\text{FE}_{\text{PO}}$ ratios over OD-Ag and commercial Ag nanoparticles (c-Ag NPs) during NO and pyruvic acid electrolysis. The gradually increasing $\text{FE}_{\text{Ala}}/\text{FE}_{\text{PO}}$ ratio over OD-Ag with accumulated charge highlights its superior activity in oxime (PO) reduction, while c-Ag NPs show limited conversion efficiency⁴⁷. Reproduced with permission from ref. 47, Copyright © 2023 Springer Nature. (h) Electrostatic potential (ESP) analysis for CuPc-Amino and CuPc¹⁰⁸. (i) Electronic charge redistribution patterns were examined through Hirshfeld charges analysis for CuPc-Amino relative to its unsubstituted CuPc¹⁰⁸. (j) Urea synthesis partial current densities for both CuPc-Amino and CuPc catalysts were measured across a range of applied potentials¹⁰⁸. Reproduced with permission from ref. 108, Copyright © 2024 Springer Nature.

abundant acid-base pairs on the surface of MgO, which played a key role in steering the NO_x reduction pathway, while the highly dispersed MgO nanoparticles embedded in the N-doped carbon matrix facilitate preferential NO adsorption limiting HER activity. This fine control over surface coordination enables the production of 4-CBOE with 93% selectivity and 65.1% FE at 12 mA.

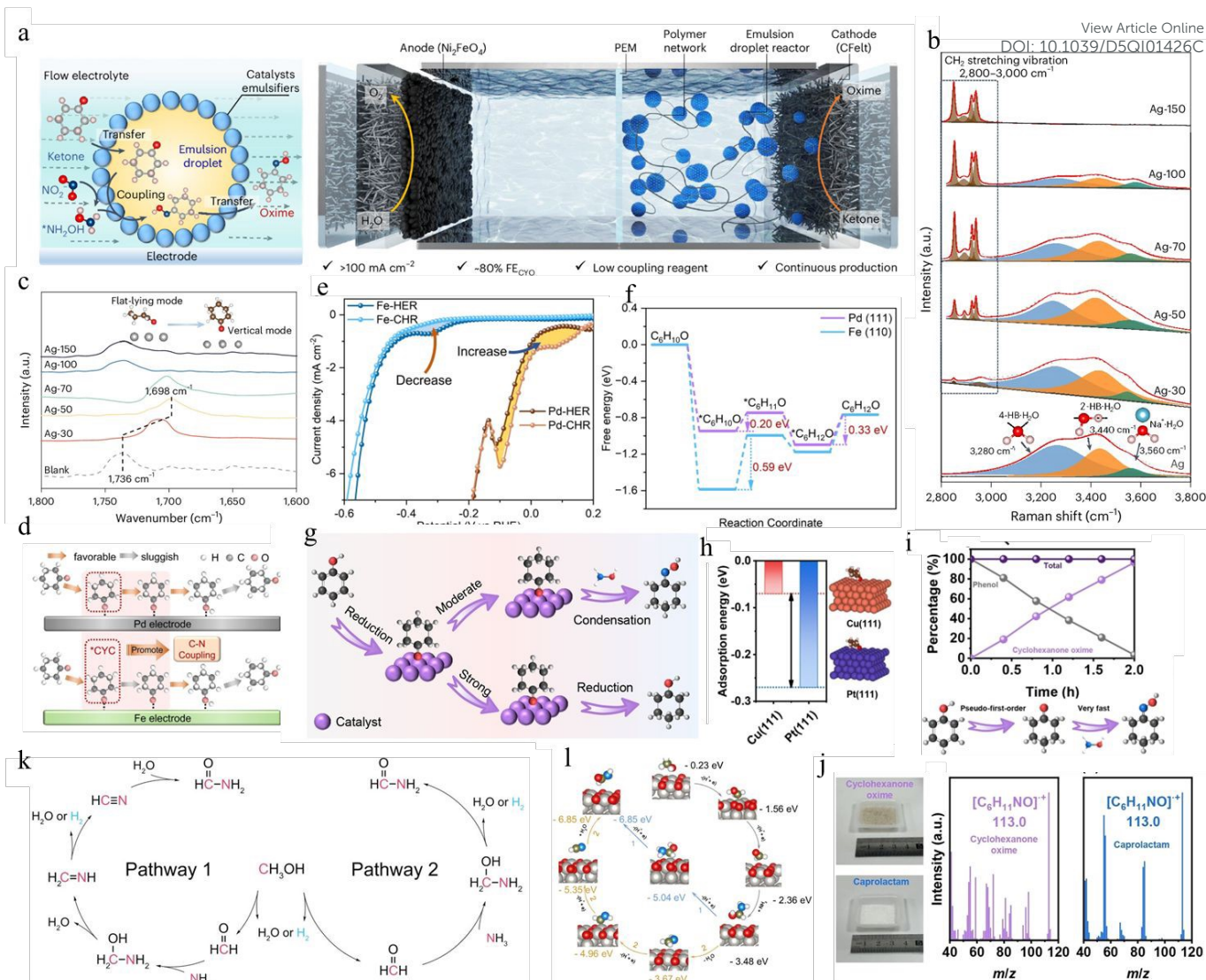
Zhu and co-authors developed an atomically dispersed iron-supported defect TiO_2 electrocatalyst ($\text{adFe-TiO}_x/\text{Ti}$) that synergistically integrates oxygen vacancies (OVs) and isolated Fe sites to facilitate C-N coupling reactions for α -amino acid synthesis¹⁰⁷. The defect rich TiO_2 matrix was designed to introduce a large number of OVs, while Fe atoms were anchored at these vacant positions to form a uniformly dispersed Fe-O coordination structure. The formation of Fe-O₃ complexes enhances the electronic interaction between Fe and defective oxide support (Fig. 4f). OVs promoted the adsorption of nitric acid (NO_3^-), glyoxylic acid (GA) and glyoxylic oxime (GO), while adFe significantly reduced the energy barrier for the conversion of NO_3^- to NH_2OH and GO to glycine. A glycine yield of 80.2%

was achieved using the catalyst $\text{adFe-TiO}_x/\text{Ti}$ with GA conversion close to 100%.

Li et al. Developed a coordination-modulated Ag-based catalyst to realize the electrosynthesis of alanine from NO and pyruvic acid⁴⁷. The oxide-derived Ag (OD-Ag) exhibits a reduced average coordination number ($\text{CN} \approx 7.2$), which leads to under-coordinated Ag sites with altered surface electronic structures. These modified coordination environments enable stronger adsorption of oxime intermediates and lower the energy barriers for the key transformation step, including N-O bond cleavage and C=N hydrogenation. Compared to commercial Ag nanoparticles, OD-Ag significantly boosts the pyruvate oxime (PO)-to-alanine conversion rate while suppressing side HER processes (Fig. 4g). The coordination-induced activity enhancement is further supported by DFT calculations showing favorable charge redistribution and reduced activation energy on low-coordinated sites.

The research of Li and co-workers utilized the ligand engineering strategy exemplified by CuPc-Amino demonstrating how coordination environment modulation enhances electrocatalytic urea synthesis¹⁰⁸. Amino substitution





strengthens intramolecular Cu-N coordination (coordination number increased from 3.9 to 4.2), while simultaneously optimizing the electronic structure through reduction of the Cu site electrostatic potential from 20.41 to 5.64 kcal mol⁻¹ (Fig. 4h). This coordination environment modification effectively suppresses electrochemical demetallation and promotes adsorption of key *CO and *NO intermediates for C-N coupling (Fig. 4i). The engineered CuPc-Amino catalyst achieves a remarkable urea yield rate of 103.1 ± 5.3 mmol h⁻¹ g⁻¹ at -1.6 V vs. RHE (Fig. 4j), representing a 2.6-fold improvement over

unmodified CuPc, while maintaining near-zero activity decay over 10 cycles compared to the 67.4% decay observed for CuPc.

3.3 Interface Engineering

Interface engineering involves the rational design and control of regions where two or more phases intersect, which including solid-liquid interfaces and multiphase interface such as solid-liquid-gas or oil-water-gas interfaces^{83, 109}. These interface regions often exhibit unique electron transfer, reaction environment, reaction environments, and mass transfer properties. In C-N coupling reaction, multiphase interfaces can



enhance reactant enrichment, stabilize key intermediates, and provide spatial separation of reaction steps. Therefore, interface engineering serves as a powerful strategy to promote multi-step catalytic pathway/multi-electron transfer pathway and improve the efficiency of C-N bond formation.

A representative example of multiphase interface design is the Pickering emulsion droplet-integrated electrode developed by Zhang et al., which enables the continuous-flow electrocatalysis of cyclohexanone oxime from NO and cyclohexanone⁴⁴. In this system, the catalyst Ag-50 is composed of silver nanoparticles modified by polypyrrole, which has both the functions of a catalyst and a Pickering emulsifier. It assembles at the oil-water interface and stabilizes the emulsion microreactor on the electrode surface (**Fig. 5a**). The interface environment promotes selective enrichment and alignment of oil phase cyclohexanone and water phase NH_2OH . In situ ATR-FTIR (**Fig. 5b**) and Raman spectroscopy (**Fig. 5c**) revealed asymmetric hydrogen-bonding network formed by interfacial water, which aligning cyclohexanone molecules vertically on the surface and promoting nucleophilic attack by NH_2OH . This unique interfacial configuration enhances the nucleophilic attack of NH_2OH and suppresses its over-reduction to NH_3 , thereby improving C-N coupling efficiency. Under industrial-level current densities (100 mA cm^{-2}), the system achieved 83.8% FE and $0.78 \text{ mmol h}^{-1} \text{ cm}^{-2}$ yield. Further immobilization of the emulsion droplets using polypyrrole cross-linking allowed for gram-scale oxime synthesis in continuous flow, highlighting the power of interfacial engineering in biphasic electrocatalysis.

Wu et al. designed a biphasic electrochemical system for the synthesis of cyclohexanone oxime from NOx and cyclohexanone (CYC), in which achieving a nearly 100% yield on Fe-based electrocatalysts under flow conditions⁷⁸. In this system, NOx and CYC from two immiscible phases accumulate simultaneously on the Fe surface, forming $^*\text{NH}_2\text{OH}$ and $^*\text{CYC}$ as adsorbed intermediates (**Fig. 5d**). Fe exhibits a stronger adsorption to CYC but weaker hydrogenation ability compared to Pd (**Fig. 5e**), thus preventing over-reduction and ensuring $^*\text{CYC}$ availability for C-N coupling. Meanwhile, DFT calculations show that $^*\text{NH}_2\text{OH}$ desorption is energetically favored over its reduction to NH_3 (**Fig. 5f**), enabling its coupling with $^*\text{CYC}$ to form the oxime product. This spatial separation and enrichment effect at the interface enhances C-N coupling selectivity.

Jia and co-workers reported a well-designed interfacial electrocatalytic system using oxide-derived Cu catalyst for the synthesis of cyclohexanone oxime from phenol and hydroxylamine⁶¹. The Cu catalyst effectively balances the adsorption and activation of H_2O , phenol, and NH_2OH at the liquid-solid interface. Moderate adsorption of in situ formed cyclohexanone enables efficient coupling with NH_2OH while suppressing its over-reduction to cyclohexanol (**Fig. 5g**). In addition, weak binding of the oxime product facilitates rapid desorption, avoiding further hydrogenation (**Fig. 5h-i**). These features enable a high FE (69.1%) and a phenol-to-oxime conversion rate of 97.5% at -0.9 V . In addition to its high selectivity and FE, this system demonstrated excellent practical applicability under industrially relevant conditions. Using a PEM flow cell at 50 mA cm^{-2} , the production rate of cyclohexanone

oxime reached $54.0 \text{ g h}^{-1} \text{ g}_{\text{cat}}^{-1}$ with a yield exceeding 90.0% and a carbon selectivity above 99.9%. The oxime product could be readily purified by simple separation and drying, affording a 99.9% isolated yield with high purity (**Fig. 5j**).

Chen et al. proposed an interfacial regulation strategy based on the cooperative enrichment and coupling of key intermediates to enable the efficient aqueous electrocatalysis of oximes from NOx and aldehyde substrates⁵⁹. By tailoring the microenvironment at the Fe-based catalyst-electrolyte interface, the strategy facilitates the simultaneous surface accumulation and spatial proximity of NH_2OH and R-CHO, thereby promoting their preferential C-N coupling. The interfacial-specific enrichment of NH_2OH , combined with the strong adsorption of R-CHO (desorption energy increased by 23%), effectively suppresses competing hydrogenation pathways. Moreover, the activation barrier for interfacial C-N coupling (0.32 eV) is significantly lower than that for NH_2OH reduction (0.68 eV), enabling a high oxime yield of up to 99%. In the subsequent flow reactor system, enhanced interfacial mass transport further boosts the oxime production rate to $22.8 \text{ g h}^{-1} \text{ g}_{\text{cat}}^{-1}$.

Jia et al. engineered a MOF-derived Cu-Cu₂O heterojunction catalyst ($\text{Cu}_x\text{C}_y\text{O}_z@600$) for efficient hydroxylamine (HA) electrocatalysis via in situ electrochemical reconstruction²⁹. In this structure, the cooperative interaction between Cu^0 and Cu^+ at the interface optimizes key steps in the reaction pathway, particularly facilitating the protonation and desorption of intermediates such as $^*\text{NO}_2$ and $^*\text{NH}_2\text{OH}$. The Cu(111)-Cu₂O(111) heterointerface plays a pivotal role in modulating the local electronic environment. This interface lowers the desorption energy of $^*\text{NH}_2\text{OH}$ and facilitates the protonation of $^*\text{NO}$ to $^*\text{NHO}$ via a barrierless process, effectively overcoming the rate-determining step in HA formation. The coexistence of Cu^0 and Cu^+ species further promotes $^*\text{NO}_2$ protonation and intermediate stabilization, enabling highly selective CP-O generation.

Shao et al. developed a scalable electrocatalytic system for converting methanol and ammonia into formamide via C-N coupling over a boron-doped diamond (BDD) electrode¹¹⁰, achieving a FE of 41.2% and selectivity of 73.2% at a current density of 120 mA cm^{-2} . The BDD electrode provides an inert liquid-solid interface that supports the oxidation of methanol to aldehyde-like species. Systematic tuning of $\text{CH}_3\text{OH}/\text{NH}_3$ molar ratios and electrolyte pH demonstrated that modulating the local interfacial environment is key to enhancing formamide selectivity and suppressing side reactions. The reaction pathway was further elucidated by analyzing the electrochemical behavior of model intermediates, where **Fig. 5k** confirms the oxidation of methanol to formaldehyde-like species as a prerequisite for C-N coupling.

Meng et al. developed a sustainable electrooxidation strategy to synthesize formamide under ambient conditions, offering an alternative to traditional high-temperature, high-pressure synthesis routes, achieving a FE of 40.39% and product



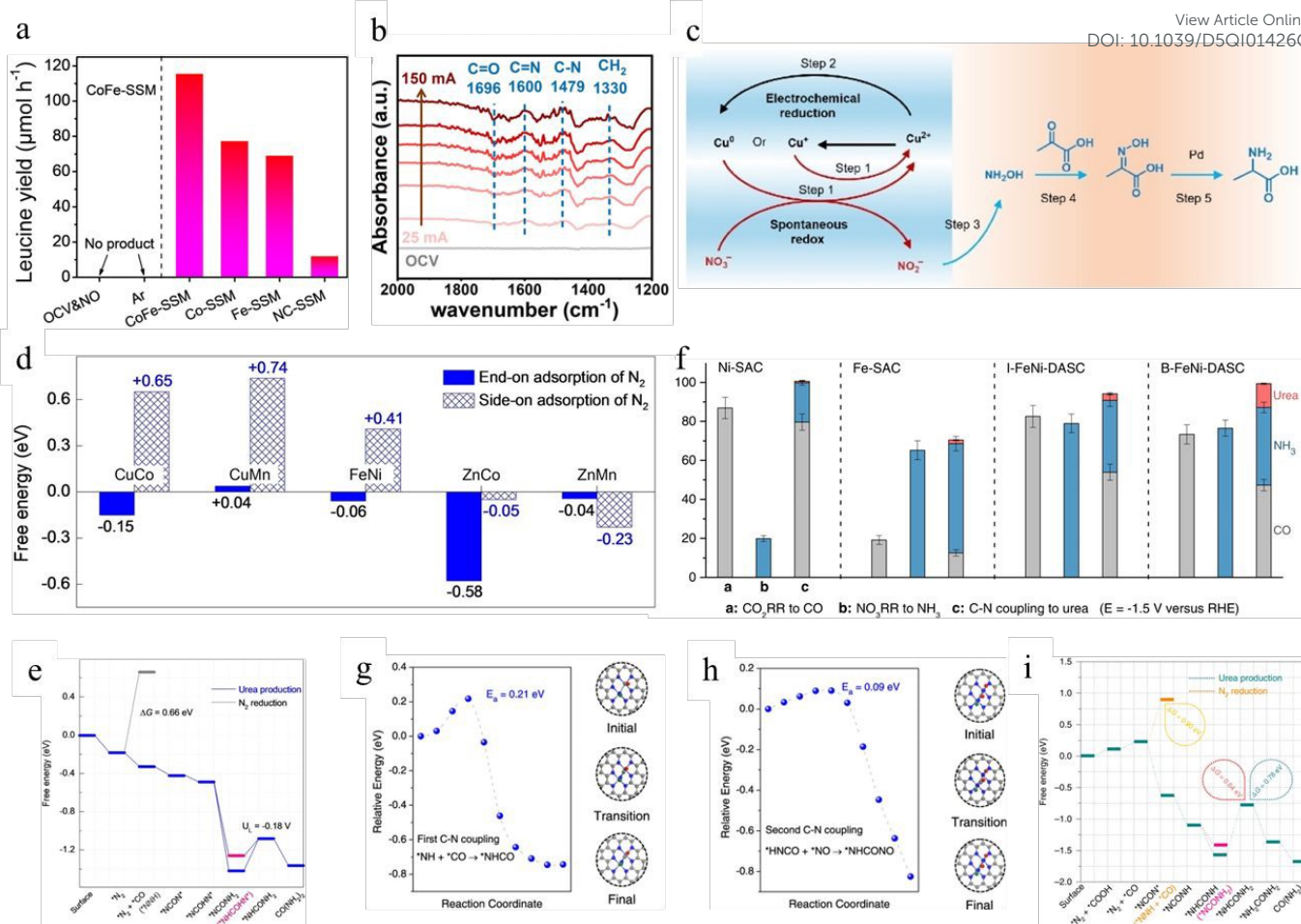


Figure 6. (a) Comparative leucine production yields of CoFe-SSM catalyst and control samples under varied reaction conditions¹¹². Reproduced with permission from ref. 112, Copyright © 2023 Wiley-VCH GmbH. (b) In situ ATR-FTIR spectra of the electrosynthesis of glycine on Cu/Bi-C@CF⁹⁷. Reproduced with permission from ref. 97, Copyright © 2024 Wiley-VCH GmbH. (c) The mechanism of nitrate and glyoxylate C-N coupling reaction⁷⁹. Reproduced with permission from ref. 79, Copyright © 2023 Wiley-VCH GmbH. (d) Comparison of free energies for side-on versus end-on N₂ adsorption on diatomic catalysts¹¹³. (e) Energy barrier of C-N coupling on ZnMn-Ni-Cl¹¹³. Reproduced with permission from ref. 113, Copyright © 2023 Wiley-VCH GmbH. (f) Comparative analysis on Ni-SAC, Fe-SAC, I-FeNi-DASC, and B-FeNi-DASC catalysts at -1.4 V vs. RHE¹¹⁴. (g) Pathway for the first C-N coupling forming *NHCO intermediate¹¹⁴. (h) Pathway for the second C-N coupling forming *NHCONO intermediate¹¹⁴. Reproduced with permission from ref. 114, Copyright © 2022 Springer Nature. (i) Reaction pathway of Pd₁Cu₁/TiO₂-400⁵⁷. Reproduced with permission from ref. 57, Copyright © 2020 Springer Nature.

selectivity of 74.26% at 100 mA cm⁻² reaction condition¹¹¹. By employing a Pt electrocatalyst, particularly its oxidized α -PtO₂ surface phase, the system facilitates the formation and stabilization of aldehyde-like intermediates from methanol oxidation (Fig. 5I). These intermediates are subsequently attacked by ammonia through a nucleophilic pathway to generate formamide. The spatial confinement and reactive environment at the electrode-electrolyte interface effectively direct the multi-step conversion and suppress overoxidation, underscoring the vital role of interface engineering in modulating C-N coupling efficiency.

3.4 Dual-Site Synergy

Single active sites may not be sufficient to perform multi-substrate complex reactions involve both NO_x and carbon-containing species. In contrast, dual-site catalysts, which contain two types of active centers with different functions, can offer new active sites to facilitate the adsorption/activation/conversion of different substrates. For

example, site A to active NO_x, while site B to active carbon-containing species. The close distance between these sites can help transfer key intermediates and encourage the formation of C-N bonds. This cooperative mechanism is useful for improving both activity and selectivity. Therefore, dual-site catalyst design has become a promising strategy for the construction of more efficient NO_x-involved C-N coupling reactions.

As a typical example of dual-site catalysis, Xian et al. constructed a CoFe alloy-decorated self-standing carbon fiber membrane (CoFe-SSM) that promoted the electrosynthesis of α -amino acids from NO and α -keto acids via a dual-site synergy strategy¹¹². In this system, the CoFe alloy provides two distinct but synergistic metal centers: the Co sites facilitated the reduction of NO to hydroxylamine (NH₂OH), while the Fe sites assisted in the transformation of oxime intermediates into α -keto acids via hydrogenation. This dual-site synergy strategy led to improve C-N coupling selectivity and FE compared to single-site catalyst systems (Fig. 6a).

Liao et al. developed a Cu-Bi bimetallic catalyst derived from MOF arrays grown on copper foam (Cu/Bi-C@CF) that enables



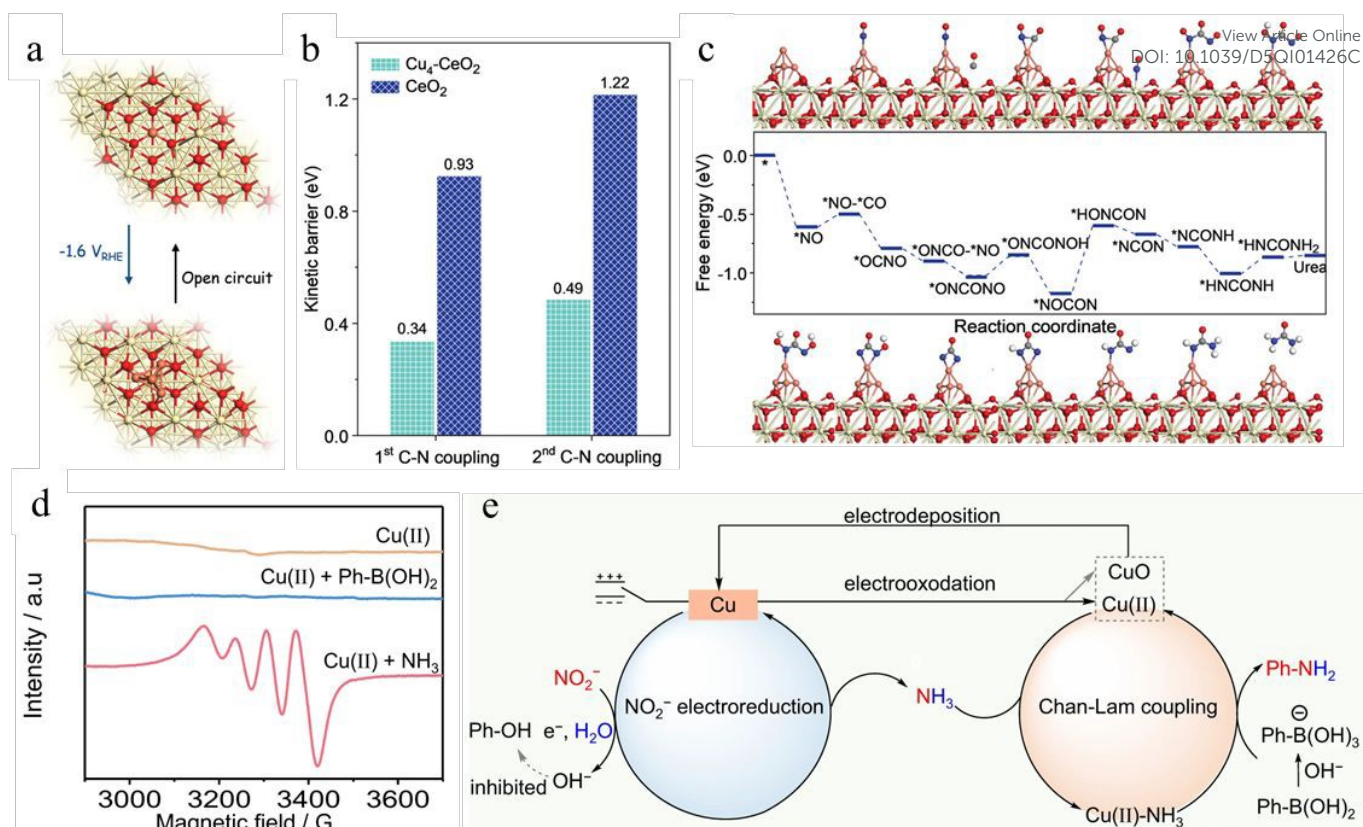


Figure 7. (a) Free energy changes of urea production on Pd₁Cu₁/TiO₂-400⁹⁴. (b) Kinetic barriers of *OCNO and *ONCONO on CeO₂ vs. Cu₄-CeO₂⁹⁴. (c) Free-energy profiles and optimized geometry of C-N coupling on Cu₄-CeO₂⁹⁴. Reproduced with permission from ref. 94, Copyright © 2023 Wiley-VCH GmbH. (d) EPR spectrum of the Cu(II)-NH₃⁹⁸. (e) Catalytic mechanism⁹⁸. Reproduced with permission from ref. 98, Copyright © 2023 Springer Nature.

high-efficiency electrosynthesis of glycine from nitrate and glyoxylate under ambient conditions⁹⁷. Cu and Bi act as two functionally distinct but spatially adjacent active sites: Cu reduces nitrate to hydroxylamine (NH₂OH), while Bi modulates the local electronic environment of Cu, weakening *NO adsorption and enhancing NH₂OH desorption. XPS and XANES data show electron transfer from Bi to Cu, weakening excessive *NO adsorption and enhancing NH₂OH selectivity. In situ ATR-FTIR spectra (**Fig. 6b**) show the formation of C=N-OH and CH₂ groups, supporting oxime formation and subsequent hydrogenation. Compared with monometallic Cu or Bi catalysts, the Cu-Bi system demonstrates significantly improved FE and product selectivity, confirming the importance of dual-site synergy in directing the C-N coupling pathway.

Wu and co-workers developed a PdCu nano-bead-wire (Pd₁Cu₁ NBWs) catalyst that exhibits a dual-site synergy in catalyzing the tandem coupling of biomass-derived pyruvic acid (PA) and nitrate (NO₃⁻) into alanine⁷⁹. The reaction proceeds through a three-step electrochemical-chemical-electrochemical cascade mechanism. Initially, the Cu sites catalyze the reduction of nitrate to hydroxylamine (NH₂OH), serving as the nitrogen donor. Subsequently, NH₂OH spontaneously condenses with PA to form a pyruvic oxime intermediate. Finally, the Pd sites drive the electrochemical hydrogenation of the oxime to produce alanine as the main product (**Fig. 6c**). By spatially separating the activation of nitrate and the reduction of the C=N intermediate, this dual-site system enables precise control over individual

reaction steps. As a result, a high alanine yield of 54.8% is achieved under ambient conditions, significantly suppressing the formation of side products such as lactic acid.

Zhang et al. achieved 63.5% FE urea electrosynthesis via Zn-Mn dual-site synergy with axial Cl coordination (ZnMn-N,Cl)¹¹³. By leveraging the complementary electronic characteristics of Zn and Mn, the catalyst forms local electrophilic and nucleophilic centers that enable co-adsorption and co-activation of CO₂ and N₂ molecules. This spatial and electronic cooperation facilitates a one-step C-N coupling pathway, bypassing the formation of ammonia intermediates, achieving a nearly 100% N-selectivity and a maximum FE of 63.5% under CO pre-poisoning. The N₂ molecule adopts a side-on adsorption mode on the Zn-Mn pair, which is thermodynamically more favorable than the conventional end-on configuration. The subsequent coupling with *CO species derived from CO₂ reduction directly forms a *NCON* intermediate (**Fig. 6d-e**).

Zhang et al. developed a bonded Fe-Ni diatomic electrocatalyst (B-FeNi-DASC) to achieve synergistic urea synthesis (**Fig. 6f**)¹¹⁴. By integrating Fe and Ni atoms into a well-defined diatomic configuration, this catalyst achieves concurrent adsorption and activation of nitrate and CO₂, offering spatial proximity and electronic complementarity between the two active centers. Compared to single-atom and non-bonded diatomic systems, the bonded Fe-Ni sites not only generate abundant *NO and *CO intermediates but also directly participate in the C-N bond formation steps (**Fig. 6g-h**).



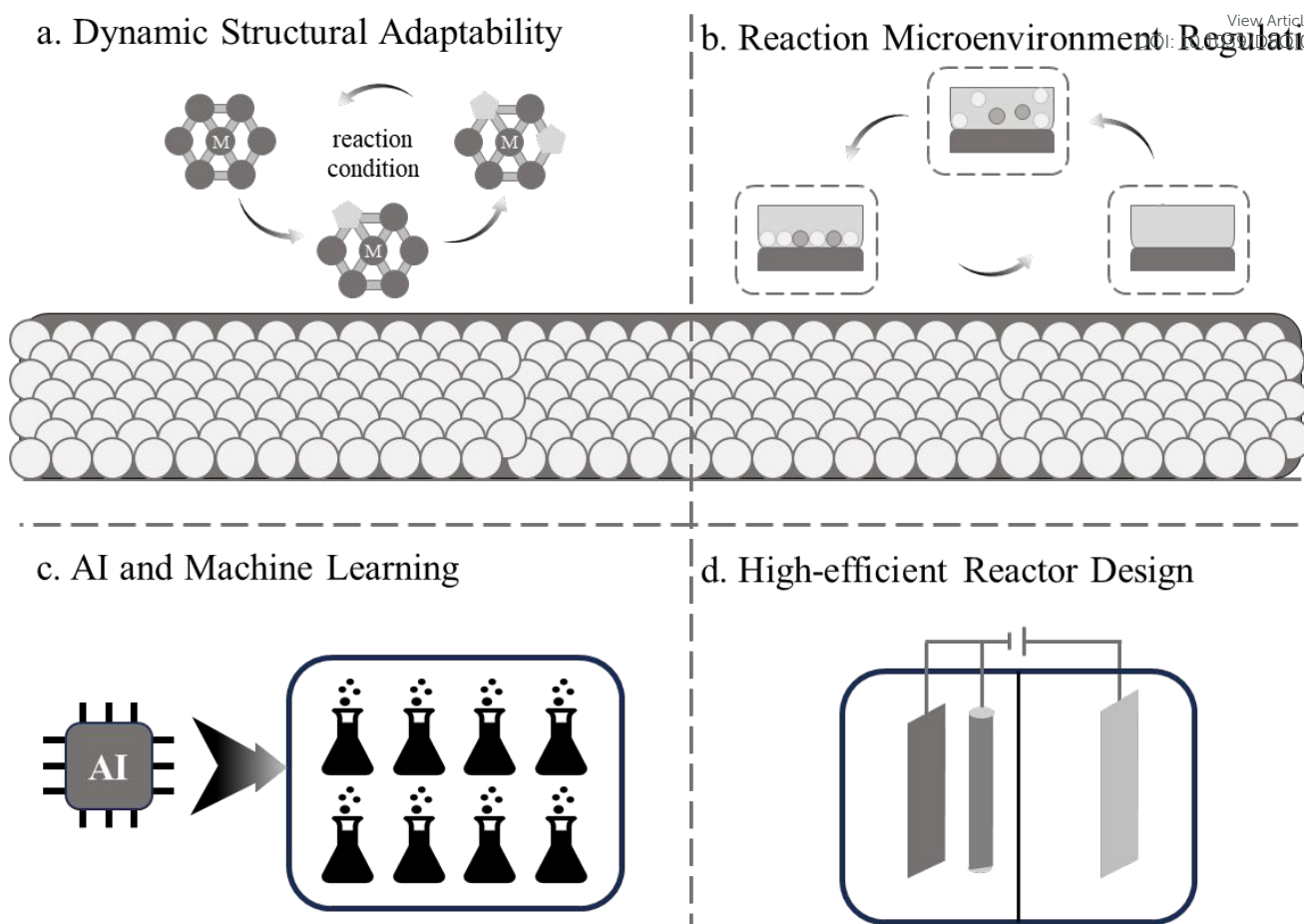


Figure 8. Four strategic directions for promoting electrocatalytic C-N bond formation. The schematic diagram outlines the strategies integrated across four dimensions to address the ongoing challenges in the electrocatalytic C-N bond formation process: (a) Dynamic structural adaptability. (b) Regulation of the reaction microenvironment to inhibit competing reactions. (c) Application of artificial intelligence and machine learning for high-throughput catalyst screening and mechanism simulation. (d) Modular flow reactors, providing a foundation for achieving industrial-scale application.

Chen and co-workers developed a dual-site synergy strategy by anchoring PdCu alloy nanoparticles onto oxygen-vacancy-rich TiO₂ nanosheets (Pd₁Cu₁/TiO₂-400), enabling the electrocatalytic coupling of N₂ and CO₂ to produce urea under ambient conditions⁵⁷. In this catalyst system, Pd enhances back-donation to weaken the N≡N bond of N₂, while Cu facilitates CO₂ activation and *CO intermediate stabilization. On the PdCu alloy surface, the activated *N=N* species couples with CO to form the key intermediate *NCON*, which proceeds through a kinetically favorable (*E*_a = 0.79 eV) and thermodynamically downhill (ΔG = -0.89 eV) pathway (Fig. 6i).

3.5 Emerging Architectures

In addition to conventional catalyst systems, emerging architectures have been developed to address the growing complexity of selective C-N coupling reactions. These include dynamically evolving structures, spatiotemporally modulated systems, and hybrid electrolyte-catalyst interfaces. Unlike the four aforementioned categories that primarily focus on spatially static structural modifications such as defect engineering, coordination tuning, interfacial design, and dual-site synergy, emerging architectures emphasize dynamic, adaptive, and process-coupled catalyst behavior. This shift reflects a transition

from material-centric design principles to those that are responsive to the reaction environment.

Wei et al. developed a dynamic reconstruction strategy by designing Cu₁ single atoms supported on CeO₂ (Cu₁-CeO₂), which undergo reversible structural transformation into Cu₄ clusters under electrocatalytic conditions⁹⁴. This dynamic evolution bridges single-atom precision with cluster-level activity, enabling a self-adjusting catalytic configuration that adapts to the electrochemical environment. The reconstituted Cu₄ clusters act as real active sites (Fig. 7a), significantly enhancing the simultaneous adsorption and activation of NO₃⁻ and CO₂. DFT calculations reveal that the C-N coupling proceeds via an Eley-Rideal mechanism, where *NO and *CO form the key intermediate *OCNO with a low energy barrier of 0.34 eV on Cu₄-CeO₂, significantly lower than the 0.93 eV required on pristine CeO₂ (Fig. 7b-c). Subsequent coupling with a second *NO yields *ONCONO, which undergoes eight electron-proton-transfer steps to produce urea.





Table 1. Progress Summary of C-N Coupling Reactions.

Regulation Strategy	Catalyst	Production	Electrolyte	Condition	Selectivity	Conversion	FE	Yield	Ref.
Defect Engineering	AD-Fe/NC	Amino Acids	0.1 M HCl+20 mM 3-Methyl-2-Oxobutanoic Acid	-0.6V vs. RHE	11.3%	47.4%	9.5%	32.1 $\mu\text{mol mg}_{\text{cat}}^{-1}$	60
Defect Engineering	Pd/Cu-V _{Cu}	DMF	0.5M KHCO ₃	20mA cm ⁻²	-	-	37.5%	385 mmol h ⁻¹ g _{cat} ⁻¹	99
Defect Engineering	e-OD-Cu	HMTA	0.1 M KOH+0.5 M K ₂ SO ₄	-0.3V vs. RHE	99%	-	74.9%	76.8%	100
Defect Engineering	Cu/PI-X	Urea	0.1 M KHCO ₃ +0.1 M KNO ₃	-1.4V vs. RHE	-	-	14.3%	255.0 mmol h ⁻¹ g ⁻¹	101
Defect Engineering	V _o -CeO ₂ -750	Urea	0.1 M KHCO ₃ +50mM KNO ₃	-1.6V vs. RHE	-	-	-	943.6 mg h ⁻¹ g ⁻¹	102
Defect Engineering	ReMn-NC	Urea	PBS	-0.3V vs. RHE	89.1%	-	-	48.9 ± 2.4 mg g ⁻¹ h ⁻¹	103
Defect Engineering	Cu ₉₇ In ₃ -C	Urea	0.1 M KHCO ₃	-1.5V vs. RHE	-	-	-	13.1 mmol g ⁻¹ h ⁻¹	62
Defect Engineering	NS-CNS	Alanine	1.0M KOH	-0.5V vs. RHE	>99.9%	79.5%	-	1199 $\mu\text{mol h}^{-1} \text{cm}^{-2}$	104
Coordination Environment Design	ZIF-7/CGF	1-PDO	0.1M KOH	-0.1V vs. RHE	-	-	75.9%	73.1%	43
Coordination Environment Design	Cu-S	Cyclohexanone oxime	0.5M PBS+0.2mM CYC+2mM NaNO ₂	-0.9V vs. Ag/AgCl	99%	-	26%	92%	42
Coordination Environment Design	HEA-PdCuAgBilene	Cyclohexanone oxime	0.5M PBS+0.02M CYC+0.1M KNO ₂	-0.9V vs. Ag/AgCl	-	-	47.6%	~100%	95
Coordination Environment Design	NH ₂ -MIL-53(Al)	2-Pyridinealdoxime	0.1M KOH	13 mA	-	-	49.8%	92.1%	105
Coordination Environment Design	MgO-SCM	4-CBOE	0.001M KOH/0.05M Li ₂ SO ₄	12 mA	93%	-	65.1%	94%	106
Coordination Environment Design	α _{sf} Fe-TiOx/Ti	Glycine	0.5M H ₂ SO ₄ + 0.1M GA + 1.0M NaNO ₃	-0.7 V vs. RHE	80.2%	100%	80.5%	83.4%	107
Coordination Environment Design	OD-Ag	Alanine	0.5M PBS	-0.56 V vs. RHE	-	-	17.0%	11.45 mmol h ⁻¹ g ⁻¹	47
Coordination Environment Design	CuPc-Amino	Urea	0.1 M KHCO ₃ +0.05 M KNO ₃	-1.6 V vs. RHE	-	-	11.9±0.6%	103.1 ± 5.3 mmol h ⁻¹ g ⁻¹	108
Interface Engineering	Ag-50	Cyclohexanone Oxime	0.5 M Na ₂ CO ₃ +0.5 M NaNO ₂	100 mA cm ⁻²	-	99.6%	83.8%	0.78 mmol h ⁻¹ cm ⁻²	44
Interface Engineering	Fe	Cyclohexanone Oxime	0.5 M K ₂ CO ₃ +1 M KNO ₃	500 mA cm ⁻²	100%	100%	~100%	59.5 g h ⁻¹ g _{cat} ⁻¹	78
Interface Engineering	Cu	Cyclohexanone Oxime	0.5 M KHCO ₃ + 20mM Phenol+10mM (NH ₄ OH) ₂ SO ₄	-0.9V vs. RHE	94%	97.5%	69.1%	90%	61
Interface Engineering	Fe-based	benzal doxime	0.5 M H ₂ SO ₄ +0.5 M K ₂ CO ₃	200 mA cm ⁻²	-	-	1.52%	99%	59
Interface Engineering	Cu ₃ C ₂ O ₂ @600	CP-O	0.5M KNO ₃ +0.1M CP	-1.6 V vs. Ag/AgCl	96.2%	99%	47.8%	34.9 mg h ⁻¹ cm ⁻²	29
Interface Engineering	BDD	Formamide	0.5 M NaHCO ₃	120 mA cm ⁻²	73.2%	-	41.2%	461.39 $\mu\text{mol cm}^{-2} \text{h}^{-1}$	110
Interface Engineering	Pt	Formamide	0.5 M NaHCO ₃ /0.25 M H ₂ SO ₄	100 mA cm ⁻²	74.26%	-	40.39%	305.4 $\mu\text{mol cm}^{-2} \text{h}^{-1}$	111
Dual-Site Synergy	CoFe-SSM	α -Amino Acids	0.1 M HCl	-0.7 V vs. RHE	56.7%	98.5%	32.4 %	115.4 $\mu\text{mol h}^{-1}$	112
Dual-Site Synergy	Cu/Bi-C@CF	Glycine	0.1 M HCl	75 mA	89%	>99%	65.9%	89%	97
Dual-Site Synergy	Pd/Cu ₁ NBW's	Alanine	1.0M KNO ₃ +50mM PA	-0.3V vs. RHE	-	-	-	54.8%	79
Dual-Site Synergy	ZnMn-N	Urea	KHCO ₃	-0.3V vs. RHE	100%	-	63.5%	4.0 mmol g ⁻¹ h ⁻¹	113
Dual-Site Synergy	B-FeNi-DASC	Urea	0.1 M KHCO ₃ +50 Mm KNO ₃ /KNO ₂	-1.5 V vs. RHE	-	-	17.8%	20.2 mmol h ⁻¹ g ⁻¹	114
Dual-Site Synergy	Pd/Cu ₁ /TiO ₂ -400	Urea	0.1 M KHCO ₃	-0.4 V vs. RHE	-	-	8.92%	3.36 mmol g ⁻¹ h ⁻¹	57
Emerging Architectures	Cu ₁ -CeO ₂	Urea	0.1M KHCO ₃ +50 mm KNO ₃	-1.6 V vs. RHE	-	-	-	52.84 mmol h ⁻¹ g _{cat} ⁻¹	94
Emerging Architectures	LC-Cu NC	Arylamines	0.25M PBS+MeOH	$E_{\text{on}}=-1.1 \text{ V}, E_{\text{off}}=0.4 \text{ V}$	-	-	95%	72%	98

He et al. demonstrated that pulse-engineered spatiotemporal control fundamentally transforms electrocatalytic arylamine synthesis by dynamically orchestrating three synchronized processes⁹⁸: (1) During cathodic pulses (-1.1 V), low-coordinated Cu nanocorals leverage undercoordinated sites (CN = 9.75) to reduce NO₂⁻ to NH₃ with 95% FE, accumulating critical nitrogen feedstock. (2) Switching to anodic pulses (+0.4 V) oxidizes the Cu electrode to solubilize catalytic Cu(II), which coordinates with NH₃ to form the key Cu(II)-NH₃ intermediate (Fig. 7d), while simultaneously driving electrophoretic migration of nucleophilic ArB(OH)₃⁻ toward the anode and consuming OH⁻ ions via Cu oxidation (2Cu + 2OH⁻ → Cu₂O + H₂O), thereby suppressing phenol byproducts from 28% to <5% (Fig. 7e). (3) The rapid polarity alternation maintains high local concentrations of Cu(II)/NH₃ near the electrode surface, accelerating C-N bond formation to achieve 72% arylamine yield (an 8-fold enhancement over static potential methods). This temporal strategy further enables self-regeneration of catalytic Cu species through cathodic electrodeposition and expands to ¹⁵N-labeled arylamine synthesis (98.99% isotopic purity) and click reactions, establishing pulsed electrochemistry as a paradigm-shifting approach for complex organic electrosynthesis beyond static catalyst design.

4. Conclusions and Outlook

In this paper, we have systematically reviewed recent progress in electrocatalytic C-N coupling reactions involving NO_x species and various carbon-based reactants (Table 1). By integrating NO_x species with diverse carbon sources (such as CO₂, aldehydes, or biomass derivatives), these C-N coupling reactions offer a sustainable route to synthesize high-value nitrogen-containing products, including oximes, amines, amides, amino acids, and urea.

Focusing on the underlying catalytic mechanisms and structural-function correlations, we classified catalyst engineering strategies into five representative categories: defect engineering, coordination environment design, interface engineering, dual-site synergy, and emerging architectures. These approaches address key challenges such as poor intermediate stability, weak substrate adsorption compatibility, and low selectivity.

Despite the substantial progress enabled by the five major catalyst design strategies, several fundamental challenges remain unresolved. For instance, the underlying reaction kinetics and the behavior of transient intermediates under dynamic electrochemical environment are still not clearly understood, making it difficult to rationally modulate activity and selectivity. Competing reactions such as hydrogen evolution, over-reduction and other undesired side reactions often interfere with the desired coupling steps. To further push the frontiers of NO_x-involved C-N electrosynthesis, we propose four strategic development directions (Fig. 8).

4.1 Dynamic structural adaptability

Conventional static catalyst structures are often insufficient for accommodating the dynamic evolution of active sites during

multi-step C-N coupling (Fig. 8a). Future efforts should focus on the development of adaptive catalytic systems capable of reversible reconstruction, single-atom to cluster transitions, or self-regulated site reconfiguration in response to the reaction environment. These dynamic architectures can provide spatiotemporal modulation of activity and selectivity across complex reaction pathways.

4.2 Reaction microenvironment regulation

The rational design of interfacial microenvironments, including local pH, reactant gradients, phase interfaces (e.g., solid-liquid-gas), and electric field distribution, is critical for steering intermediate adsorption and suppressing competing pathways (Fig. 8b). Engineering spatially confined or hydrophilic/hydrophobic domains on catalyst surfaces can selectively enrich reactants, promote intermediate pairing, and improve mass transport for enhanced C-N bond formation.

4.3 AI and machine learning

Artificial intelligence (AI) and machine learning (ML) are expected to play increasingly important roles in catalyst development (Fig. 8c). By integrating large-scale experimental and theoretical databases, ML algorithms can accelerate the identification of performance descriptors, predict reaction energetics, and guide the inverse design of catalysts with tailored electronic structures and adsorption characteristics. Coupling AI with high-throughput screening and automated synthesis platforms will enable more efficient exploration of the vast catalyst space.

4.4 High-efficient reactor design

To realize the practical deployment of electrocatalytic C-N coupling systems, efforts should be directed toward the design of scalable and modular electrochemical flow reactors (Fig. 8d). These systems can provide precise control over reaction residence time, electrode configuration, and mass transfer conditions, while facilitating continuous production and system integration. The combination of optimized catalysts with intelligent flow designs can bridge the gap between laboratory demonstrations and industrial applications.

In summary, electrocatalytic NO_x-involved C-N coupling represents a promising frontier in green chemical synthesis. Through advances in dynamic catalyst design, microenvironment control, AI-assisted optimization, and scalable system engineering, the field is poised to overcome current limitations and move toward high-efficiency, selective, and industrially viable electrosynthesis platforms. Future progress will rely on the integration of scalable catalyst design with modular reactor engineering, as well as cross-disciplinary collaboration spanning materials science, artificial intelligence, and process systems. Addressing these bottlenecks will be essential for translating NO_x-involved electrosynthesis from conceptual frameworks into practical, sustainable applications.

Author contributions

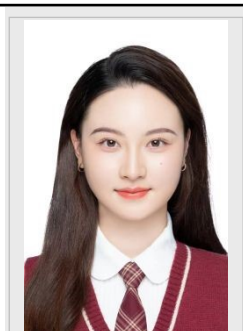


R. W., X. S., and B. H. wrote the manuscript. X. S. and B.H. supervised the project.

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

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Conflicts of interest

The authors have no conflicts of interest to declare.

Data availability

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Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

