



Cite this: *Nanoscale*, 2021, **13**, 19133

Coordination environment engineering on nickel single-atom catalysts for CO₂ electroreduction†

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Coordination engineering has recently emerged as a promising strategy to boost the activity of single atom catalysts (SACs) in electrocatalytic CO₂ reduction reactions (CO₂RR). Understanding the correlation between activity/selectivity and the coordination environment would enable the rational design of more advanced SACs for CO₂ reduction. Herein, via density functional theory (DFT) computations, we systematically studied the effects of coordination environment regulation on the CO₂RR activity of Ni SACs on C, N, or B co-doped graphene. The results reveal that the coordination environments can strongly affect the adsorption and reaction characteristics. In the C and/or N coordinated Ni–B_XC_YN_Z (B-free, X = 0), only Ni acts as the active site. While in the B, C and/or N coordinated Ni–B_XC_YN_Z (X ≠ 0), the B has transition-metal-like properties, where B and Ni function as dual-site active centers and concertedly tune the adsorption of CO₂RR intermediates. The tunability in the adsorption modes and strengths also results in a weakened linear scaling relationship between *COOH and *CO and causes a significant activity difference. The CO₂RR activity and the adsorption energy of *COOH/*CO are correlated to construct a volcano-type activity plot. Most of the B, C, and/or N-coordinated Ni–B_XC_YN_Z (X ≠ 0) are located in the left region where *CO desorption is the most difficult step, while the C and/or N coordinated Ni–B_XC_YN_Z (X = 0) are located in the right region where *COOH formation is the potential-determining step. Among all the possible Ni–B_XC_YN_Z candidates, Ni–B₀C₃N₁ and Ni–B₁C₁N_{2-N-oppo} are predicted to be the most active and selective catalysts for the CO₂RR. Our findings provide insightful guidance for developing highly effective CO₂RR catalysts based on a codoped coordination environment.

Received 1st September 2021,

Accepted 19th October 2021

DOI: 10.1039/d1nr05742a

rsc.li/nanoscale

Introduction

With the excessive consumption of non-renewable fossil fuels (coal, oil and natural gas), the increasing emission of carbon dioxide (CO₂) into nature has caused severe ecological and environmental problems.^{1–3} As an alternative way, CO₂ can be reduced into useful chemical products, such as carbon monoxide, methane, ethylene and methanol. However, the carbon element in CO₂ is in the highest +4 oxidation state and the C=O bond is highly stable, which makes it very difficult to be reduced.^{4–7} Among the variety of developed solutions, the electrochemical reduction of CO₂ is one of the most promising strategies and has attracted widespread attention due to its low energy consumption, mild reaction conditions, and simple

operation, which relies on efficient catalysts to accelerate the reaction.^{8–10} The promising electrocatalysts for the electrochemical CO₂ reduction reaction (CO₂RR) should be able to operate at a low overpotential and can effectively control the product selectivity, inhibit the competitive hydrogen evolution reaction (HER) and maintain high electrochemical stability.^{11–14}

Among the various types of investigated electrocatalysts,^{15,16} single-metal-atom catalysts have developed rapidly and emerged as a promising class of catalysts for the CO₂RR.^{17–20} Compared to the metal bulk and nanoparticle counterparts, the atomically dispersed single metal catalysts have largely exposed active sites and exhibit effective atom utilization and high selectivity for the electrochemical CO₂RR.^{21,22} Moreover, their well-defined structures serve as an ideal model to establish the relationship between the structure and catalytic properties at an atomic level. To stabilize the single metal center from agglomeration, the single-metal-atom catalysts are usually coordinated and immobilized on nitrogenated carbon-based conductive substrates, such as carbon nanotubes, graphene and amorphous or porous carbon.^{23–25} In particular, the graphene matrix has been widely used due to its high electronic conductivity and large surface area for metal loading.

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† Electronic supplementary information (ESI) available: Computed formation energy, dissolution potential, adsorption free energy, charge transfer, charge density difference, reaction free energy change of the elementary reaction step of CO₂RR, projected density of states, and Bader charge analyses of Ni–B_XC_YN_Z systems. See DOI: 10.1039/d1nr05742a

Many transition metals, such as Mn, Fe, Co, Ni, and Cu,^{26–30} have been embedded into nitrogen-doped carbon systems and investigated as CO₂RR electrocatalysts. The results demonstrate that the intrinsic CO₂RR activity of Fe–N–C and especially Ni–N–C catalysts to yield CO is higher than that of the Co-, Mn-, and Cu-based moieties, which even rival the state-of-the-art Au- and Ag-based catalysts.^{26,31–34} However, the relatively strong binding of CO* over Fe–N_x and Mn–N_x single sites could lead to lower selectivity for CO formation.

Particularly, apart from being cost effective, Ni single atom catalysts have displayed exceptional activity, selectivity and high stability.^{6,35–37} Additionally, several recent experimental advances have demonstrated that the CO₂RR activity of Ni–N–C entities can be effectively manipulated by controlling the local coordination environment of the Ni active center.^{19,38} For example, Li *et al.*²² realized a specific Ni–N₄ structure through a topo-chemical transformation strategy by carbon layer coating, which shows excellent activity and remarkable stability for the CO₂RR to CO. Bao *et al.*³⁹ reported that coordinatively unsaturated Ni–N sites within porous carbon had higher selectivity and activity for the CO₂RR than the Ni–N₄ sites. Joo *et al.*⁴⁰ and Lu *et al.*⁴¹ both showed that the Ni–N₃V (V: vacancy) sites with a shrunk Ni–N–C local structure incorporated into the graphene lattice exhibit enhanced CO₂RR performance compared to the Ni–N₄ sites. Moreover, Jiang *et al.*⁴² prepared a series of Ni single atom catalysts with controlled Ni–N coordination numbers (Ni–N_x–C, $x = 2, 3, 4$) by varying the pyrolysis temperature of the polypyrrole@MgNi-MOF-74 precursor. They revealed that the Ni–N₂–C catalyst with Ni coordinated by two N and two C shows far superior CO faradaic efficiency and turnover frequency compared to the Ni–N₃–C and Ni–N₄–C counterparts. Cheng *et al.*⁴³ fabricated Ni-based catalysts with various N/C coordination numbers (Ni@N_xC_y) through pyrolysis of carbon substrates at different temperatures and achieved an optimal catalytic performance for the



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Ni@N₂C₂ catalysts. Theoretically, Zhang *et al.*¹¹ suggested that the graphene embedded Ni–N₅ site with additional ligated axial N atoms exhibits lower CO₂RR onset overpotential than the Ni–N₄ site.

This recent progress has evidenced the strong potential of coordination environment regulation in tuning the CO₂RR activity of a single metal center. In principle, due to the difference in electronegativity and atomic size, changing the surrounding coordination elements will modify the electronic structure of the metal center, which would essentially affect the adsorption strength of the reaction intermediates and in turn, modify the activity and selectivity. It is noteworthy that, in addition to the widely studied Ni–N_xC_y catalysts with Ni–N and Ni–C coordination environments, many recent studies have shown that boron can be doped into the carbon matrix and the incorporation of boron can greatly accelerate the electrocatalytic reactions.^{44–46} The boron (~0.82 Å) has a comparable covalent radius with C (~0.77 Å) and N (~0.75 Å), which can form rich compounds with C (e.g., BC₃),⁴⁷ N (BN)^{48,49} and Ni (e.g., NiB, Ni₂B, Ni₃B, Ni₄B₃).⁵⁰ Moreover, B, in many cases, functions like a transition metal since the hybridization of its non-fully occupied valence electrons supplies a great chance to accept and donate lone-pair electrons. Hence, B can also be a potential element to dope and adjust the coordination environment.

In this research, we investigate the CO₂RR performance of Ni-centered graphene with various B/C/N coordination environments from density functional theory (DFT) calculations. Recently, Goddard and Luo *et al.*⁵¹ reported a comprehensive study of the CO₂RR mechanism on C/N coordinated Ni SAC by taking kinetics, the solvation effect, and experiment comparison into consideration. The CO₂RR performance varies remarkably by differences in the number of C or N bonded to Ni. Herein, we used the computational hydrogen electrode model with a number of simplifications and approximations for fast screening the potential catalysts by calculating the thermodynamic energetics. This simplified model, in most cases, allows the reliable alignment of theoretical electrochemical potentials to those measured in experiment. Compared to the prior theoretical studies that mainly focus on the N and C coordination,^{51,52} the hybrid coordination by non-metal C, N and transition-metal-like B leads to more versatile adsorption characteristics and diverse activities. Our results showed that most of the Ni–B_XC_YN_Z ($X + Y + Z = 4$) catalysts (25 out of 27 possible candidates) have high thermodynamic and electrochemical stability. The adsorption modes of CO₂, *COOH and *CO are versatile. In the case of C and/or N coordinated Ni–B_XC_YN_Z (B-free, $X = 0$), only the Ni center acts as the active site, and the CO₂ hydrogenation to *COOH is the potential-determining step. While in the case of B-coordinated Ni–B_XC_YN_Z ($X \neq 0$), the B can actively participate in the reaction. As a result, the Ni and B function as dual sites to flexibly tune the adsorption of CO₂RR intermediates, which facilitate the CO₂ hydrogenation but increase the adsorption strength of *CO, making *CO desorption the most difficult step. Because of the versatile single- and dual-site adsorption modes, the

scaling relationship between the adsorption energy of $^*\text{COOH}$ and $^*\text{CO}$ is significantly weakened. Moreover, we build the correlation between the adsorption energy and the limiting potential to describe the activity and selectivity. Compared to the pure N-coordinated Ni, the hybrid coordination by C–N, B–N or B–C–N can lead to much higher electrocatalytic activity, and several highly promising candidates stand out with high CO_2RR activity and selectivity, including $\text{Ni-B}_0\text{C}_3\text{N}_1$, $\text{Ni-B}_1\text{C}_0\text{N}_3$, $\text{Ni-B}_2\text{C}_0\text{N}_{2-\text{B-hex}}$, $\text{Ni-B}_1\text{C}_1\text{N}_{2-\text{N-oppo}}$, and $\text{Ni-B}_1\text{C}_1\text{N}_{2-\text{N-hex}}$. These results provide useful insights into the understanding of the coordination effect on the CO_2RR and offer a reference for further research on advanced electrocatalysts.

Computational details

All the spin-unrestricted DFT computations are carried out in the DMol³ code.⁵³ The generalized gradient approximation (GGA)⁵⁴ of the Perdew–Burke–Ernzerhof (PBE)⁵⁵ functional is used to describe the exchange–correlation interactions and the core electrons are treated with semi-core pseudopotentials (DSPPs).⁵⁶ The double numerical plus polarization (DNP) basis is adopted and the global orbital cut off value is set as 4.0 Å. The convergence tolerance of geometry optimization is set as 2.0×10^{-5} Ha in energy, 0.004 Ha Å⁻¹ in force, and the force exerted on each atom is set to be 0.005 Å. To simulate the aqueous solvent environment, a conductor-like screening model (COSMO) with a dielectric constant of 78.54 is used.⁵⁷ The k -points mesh of $5 \times 5 \times 1$ and $9 \times 9 \times 1$ using the Monkhorst–Pack method⁵⁸ is utilized for the structural optimization and electronic property calculations, respectively.

The absorption free energy of CO_2 on $\text{Ni-B}_x\text{C}_y\text{N}_z$ is calculated by: $\Delta G_{\text{ads}} = \Delta E_{\text{ads}} + \Delta ZPE - T\Delta S$, where ΔE_{ads} is the adsorption energy obtained by subtracting the total energy of the isolated $\text{Ni-B}_x\text{C}_y\text{N}_z$ catalyst and free CO_2 molecule from the adsorbed system. Due to the inaccurate description of CO_2 molecule by the PBE functional, we added a correction of +0.10 eV for CO_2 .⁵⁹

The change of Gibbs free energy (ΔG) for each elementary reaction step is calculated using the equation: $\Delta G = \Delta E + \Delta ZPE - T\Delta S + \Delta G_{\text{U}}$, where ΔE is the electronic reaction energy of a certain reaction step directly obtained from DFT energies, $\Delta G_{\text{U}} = -eU$, with U being the electrode potential of the electrochemical step. ΔZPE and $T\Delta S$ are the contributions of the zero-point energy and entropy, respectively, which originate from the calculation of vibrational frequencies of the adsorbed intermediates at $T = 298.15$ K. The entropy of gas phase molecules is derived from the NIST database. The detailed data are provided in the ESI.†

Results and discussion

Catalyst models and stability

As shown in Fig. 1, we use a $6 \times 6 \times 1$ graphene supercell with two C vacancies as the original substrate to construct the

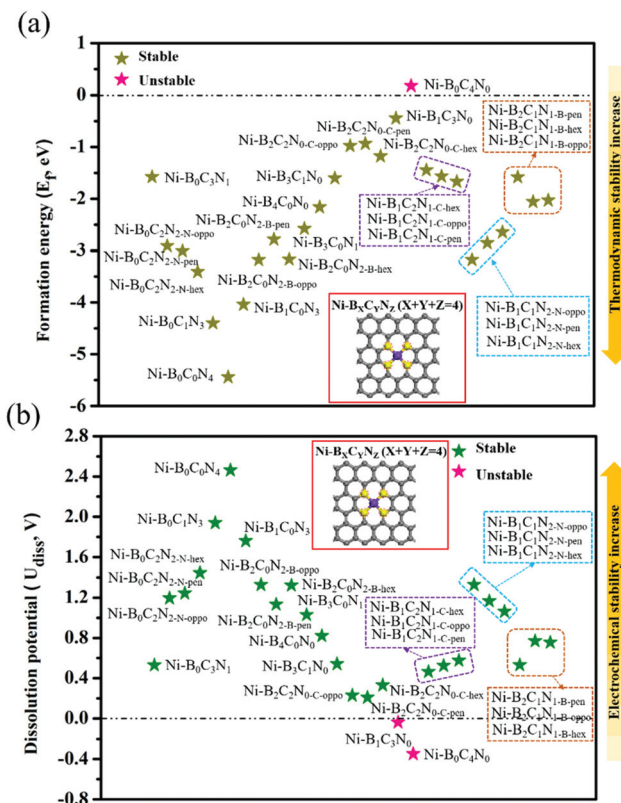


Fig. 1 The DFT-computed formation energy (a) and dissolution potential (b) of $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X + Y + Z = 4$). The inset shows the top view of the catalyst model, where the highlighted yellow atoms represent the coordination atoms of the Ni center (purple atom).

model catalysts, and then a single Ni atom is placed into the vacancy center and binds to four pyridine-like coordination atoms (B, C or N). The corresponding catalyst models are named $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X + Y + Z = 4$). Note that there are three possible coordination configurations when X , Y or $Z = 2$, denoted as $\text{Ni-B}_x\text{C}_y\text{N}_z\text{-oppo}$, $\text{Ni-B}_x\text{C}_y\text{N}_z\text{-pen}$, and $\text{Ni-B}_x\text{C}_y\text{N}_z\text{-hex}$, in which the same two atoms occupy the opposite coordination sites or neighboring coordination sites in penta-atomic and hex-atomic rings, respectively. Ultimately, 27 catalyst structures are constructed.

In order to characterize the thermodynamic stability, we calculated the formation energy (E_f) of $\text{Ni-B}_x\text{C}_y\text{N}_z$ systems, defined as $E_f = E_{\text{total}} - n_B\mu_B - n_C\mu_C - n_N\mu_N - E_{\text{Ni}}$. Here, E_{total} is the total energy of $\text{Ni-B}_x\text{C}_y\text{N}_z$. μ_B , μ_C and μ_N represent the chemical potential of B, C, and N, which corresponds to the energy of a single carbon atom in graphene, a single boron atom in planar hexagonal B_{36} ,⁶⁰ and half of the energy of N_2 molecules, respectively. While n_B , n_C , and n_N stand for the number of corresponding coordination atoms in $\text{Ni-B}_x\text{C}_y\text{N}_z$, E_{Ni} is the energy of the isolated Ni atom. From Fig. 1a, one can see that except for the pure carbon coordinated $\text{Ni-B}_0\text{C}_4\text{N}_0$, the calculated E_f values of the other 26 $\text{Ni-B}_x\text{C}_y\text{N}_z$ catalysts are all negative (−0.94 to −5.44 eV), indicating their high thermodynamic stabilities and high synthetic potential in experiments. In addition, the dissolution potential U_{diss} is another

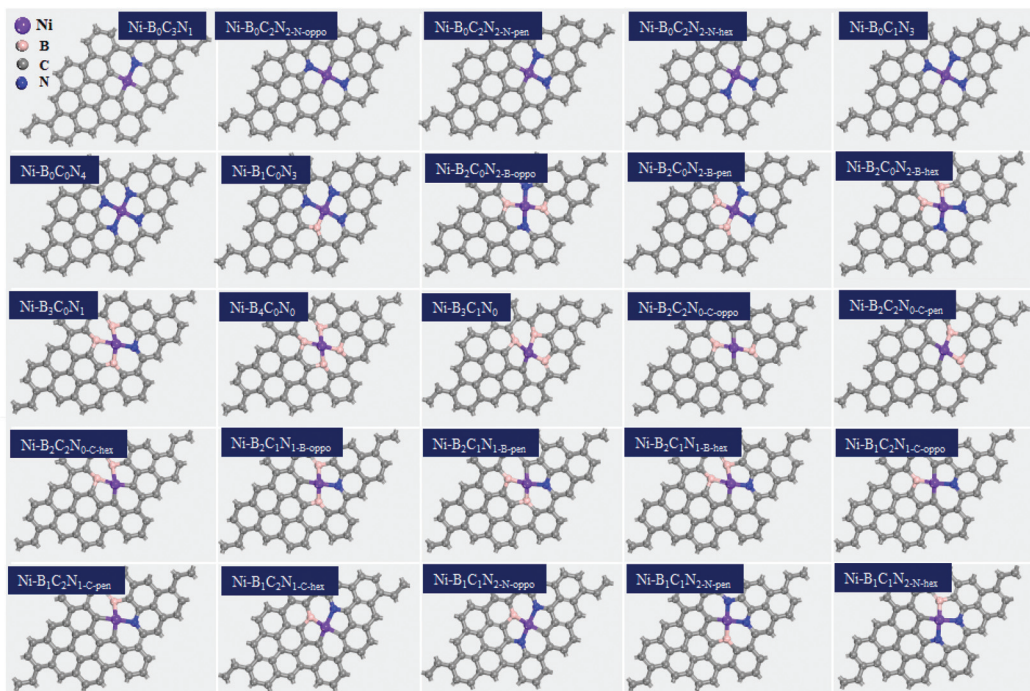


Fig. 2 The optimized structures (top view) of the screened 25 $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X + Y + Z = 4$), the color modes: purple for Ni, pink for B, blue for N, and grey for C.

important parameter to evaluate the electrochemical stabilities in a realistic electrochemical environment.^{61,62} The U_{diss} is calculated as $U_{\text{diss}} = U_{\text{diss(metal)}}^{\circ} - E_f/ne$, where $U_{\text{diss(metal)}}^{\circ}$ is the standard dissolution potential of the Ni metal, E_f is the formation energy, and n is the number of electrons involved in the dissolution (herein $n = 2$). According to this definition, only a catalyst with $U_{\text{diss}} > 0$ V vs. SHE can stably exist under acidic electrochemical conditions (the exact values of E_f and U_{diss} are listed in Table S1†).^{61,62} Among the 27 $\text{Ni-B}_x\text{C}_y\text{N}_z$ systems, only $\text{Ni-B}_1\text{C}_3\text{N}_0$ and $\text{Ni-B}_0\text{C}_4\text{N}_0$ are excluded due to their negative U_{diss} and electrochemical instability (Fig. 1b). Finally, after the stability screening, there are about 25 $\text{Ni-B}_x\text{C}_y\text{N}_z$ structures that both meet the criteria of thermodynamic and electrochemical stability. The corresponding structures of the 25 $\text{Ni-B}_x\text{C}_y\text{N}_z$ are displayed in Fig. 2, which are further investigated as the candidate catalysts of the CO_2RR .

Adsorption of reaction species

The activation of CO_2 as well as the adsorption of reaction intermediates (*e.g.*, $^*\text{COOH}$, $^*\text{CO}$) onto the surface of catalysts plays a critical role during the electrocatalytic CO_2RR . Hence, we firstly investigated the adsorption behaviors, and the most stable adsorption configurations for $^*\text{CO}_2$, $^*\text{COOH}$ and $^*\text{CO}$ are shown in Fig. 3 (the detailed adsorption free energy and charge transfer values are listed in Table S2†). From Fig. 3a, the $^*\text{CO}_2$ adsorption has two types of adsorption modes: physical adsorption and chemical adsorption. Among them, about 22 $\text{Ni-B}_x\text{C}_y\text{N}_z$ compositions form physisorption. On $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 0$), $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 1, 2$ (pen); $Y = 0$), $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 1; Y = 2; Z = 1$) and $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 1; Y = 1; Z = 2$ (hex)), the

CO_2 is physically adsorbed with the C atom of CO_2 pointed above the Ni center. While for $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 2$ (oppo, hex), 3, 4; $Y = 0$), $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($Y = 1, 2; Z = 0$) and $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 2$ (oppo, hex); $Y = 1; Z = 1$), the CO_2 is physically adsorbed with the O atom of CO_2 pointed above the Ni center. In the case of physical adsorption, the calculated CO_2 adsorption free energy is around $-0.09\text{--}0.20$ eV, and the charge transfer between CO_2 and $\text{Ni-B}_x\text{C}_y\text{N}_z$ is around $0.004\text{--}0.019|e|$ from Bader charge analysis. In the case of chemical adsorption, there also exist two types of binding modes. On $\text{Ni-B}_2\text{C}_1\text{N}_1\text{-B-pen}$, CO_2 is chemically adhered with both the C and O atoms bonded to the Ni atom, leading to a favorable adsorption free energy of -0.18 eV, substantial charge transfer of $0.36|e|$, and curved O-C-O bond angle of 145.8° . Differently, on $\text{Ni-B}_1\text{C}_1\text{N}_2\text{-N-oppo}$ and $\text{Ni-B}_1\text{C}_1\text{N}_2\text{-N-pen}$, the C atom of chemisorbed CO_2 is bonded to the Ni atom, while the O atom is bonded to the coordinated B atom. The dual-site activation of CO_2 results in stronger adsorption energies (-0.25 and -0.39 eV), larger charge transfer (0.76 and $0.96|e|$), and stronger deviation of the O-C-O bond angle from linearity (130.8° and 129.3°). The charge density difference of $\text{Ni-B}_2\text{C}_1\text{N}_1\text{-B-pen}$, $\text{Ni-B}_1\text{C}_1\text{N}_2\text{-N-oppo}$ and $\text{Ni-B}_1\text{C}_1\text{N}_2\text{-N-pen}$ with chemically captured CO_2 is shown in Fig. S1.† One can see that the Ni center and the directly coordinated N, C or B atom are the main player in the activation and orbital interaction with the CO_2 reactant.

From Fig. 3b, the $^*\text{COOH}$ has versatile adsorption modes, which can be divided into five types. Specifically, (i) on $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 0$), $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 2$ (hex), 3; $Y = 0$) and $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 2$ (hex); $Z = 0$), the C atom of $^*\text{COOH}$ is singly bonded to the Ni active center; (ii) on $\text{Ni-B}_4\text{C}_0\text{N}_0$, $\text{Ni-B}_3\text{C}_1\text{N}_0$, Ni

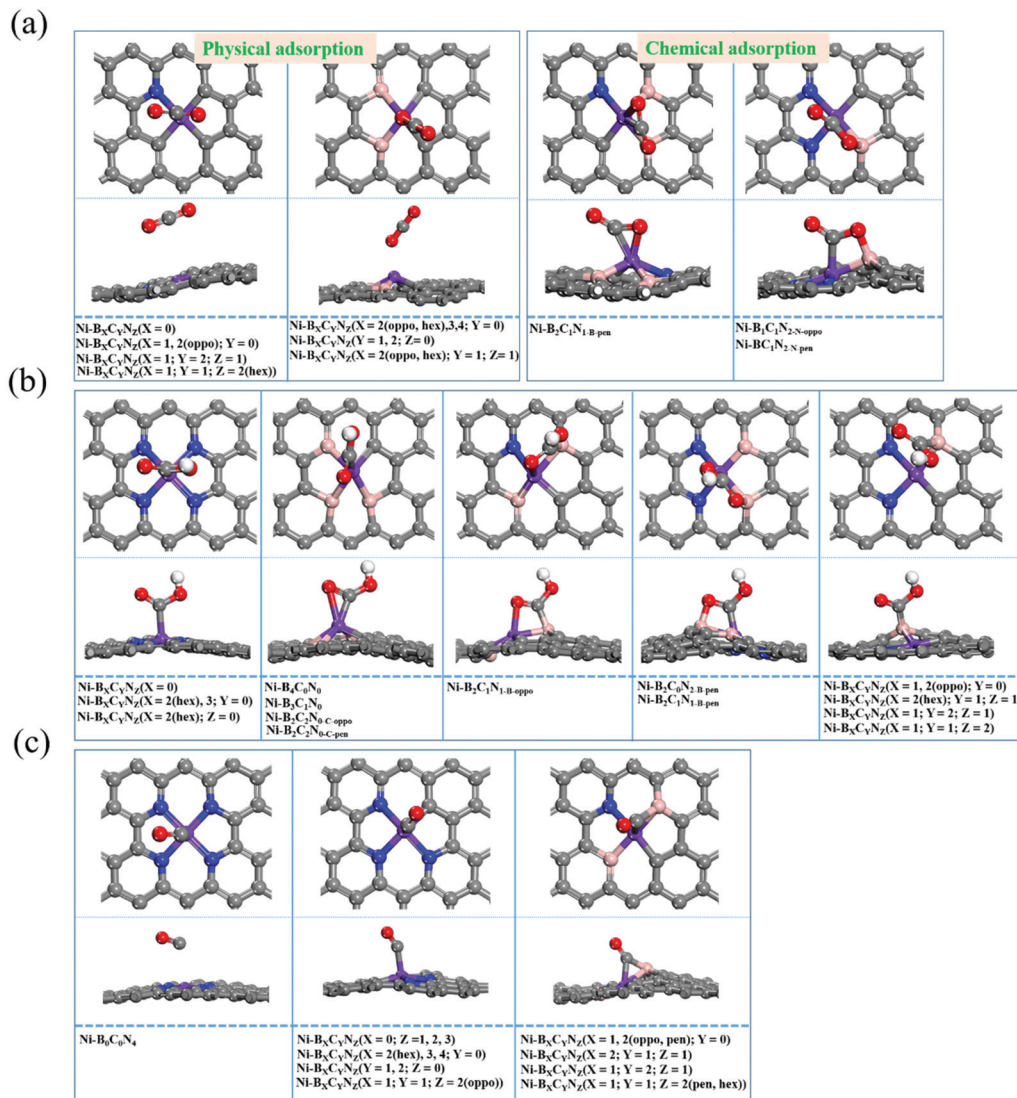


Fig. 3 The different adsorption modes of $^{*}\text{CO}_2$ (a), $^{*}\text{COOH}$ (b), and $^{*}\text{CO}$ (c) intermediates on $\text{Ni}-\text{B}_x\text{C}_y\text{N}_z$.

$\text{B}_2\text{C}_2\text{N}_0\text{-C-oppo}$ and $\text{Ni-B}_2\text{C}_2\text{N}_0\text{-C-pen}$, both the C and O atoms of $^{*}\text{COOH}$ are co-adsorbed onto the Ni atom; (iii) on $\text{Ni-B}_2\text{C}_1\text{N}_{1-\text{B-oppo}}$, the C atom of $^{*}\text{COOH}$ is bonded to a B atom while the O atom is bonded to the Ni atom; (iv) conversely, on $\text{Ni-B}_2\text{C}_0\text{N}_{2-\text{B-pen}}$ and $\text{Ni-B}_2\text{C}_1\text{N}_{1-\text{B-pen}}$, the C of $^{*}\text{COOH}$ is bonded with Ni while the O atom binds with the B atom; (v) on $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 1, 2(\text{oppo})$; $Y = 0$), $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 2(\text{hex})$; $Y = 1$; $Z = 1$), $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 1$; $Y = 2$; $Z = 1$) and $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 1$; $Y = 1$; $Z = 2$), the C atom of $^{*}\text{COOH}$ is singly bonded to the B atom, which indicates that the B acts as the active center. Moreover, from Fig. 3c, the $^{*}\text{CO}$ can have three types of adsorption modes. (i) On $\text{Ni-B}_0\text{C}_0\text{N}_4$, the $^{*}\text{CO}$ is physically adsorbed on the catalyst; (ii) on $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 0$; $Z = 1, 2, 3$), $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 2(\text{hex}), 3, 4$; $Y = 0$), $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($Y = 1, 2$; $Z = 0$) and $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 1$; $Y = 1$; $Z = 2(\text{oppo})$), the $^{*}\text{CO}$ is chemically adsorbed at the Ni center forming Ni-C single bonds; (iii) on $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 1, 2(\text{oppo, pen})$; $Y = 0$), $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 2$; $Y = 1$; $Z = 1$), $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 1$; $Y = 2$; $Z = 1$) and $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 1$; $Y = 1$; $Z = 2(\text{pen, hex})$), the $^{*}\text{CO}$ forms bridging coordination with the Ni and B atoms. These flexible and versatile adsorption modes of CO_2 , $^{*}\text{COOH}$ and $^{*}\text{CO}$ indicate that not only the Ni center but also the non-metal coordination atoms, B, in particular, play a vital role in tuning and stabilizing the reaction intermediates.

Scaling relations

The reduction of CO_2 to CO is a two-electron reaction, which includes two step hydrogenations of carbonaceous intermediates, *i.e.*, $^{*}\text{CO}_2 + \text{H}^+ + \text{e}^- \rightarrow ^{*}\text{COOH}$ and $^{*}\text{COOH} + \text{H}^+ + \text{e}^- \rightarrow ^{*}\text{CO} + \text{H}_2\text{O}$. On the traditional metal surfaces, the adsorption energies of $^{*}\text{COOH}$ and $^{*}\text{CO}$ are usually linearly correlated,⁶³ and the weak $^{*}\text{COOH}$ or $^{*}\text{CO}$ adsorption ($^{*}\text{COOH}$ formation becomes unfavorable) can lead to a low CO_2RR activity, while a strong $^{*}\text{COOH}$ or $^{*}\text{CO}$ adsorption makes the release of the CO product become difficult. Therefore, how to break or weaken the linear scaling relationship between $^{*}\text{COOH}$ and $^{*}\text{CO}$ has

long been pursued in the CO₂RR study. In our case, we also examined the correlation between *COOH and *CO. The adsorption energy (E_{ads}) of *COOH and *CO on Ni-B_XC_YN_Z is calculated using the equation:

$$E_{\text{ads}}(*\text{COOH}) = E_{(\text{Ni-B}_X\text{C}_Y\text{N}_Z + \text{COOH})} - E_{(\text{Ni-B}_X\text{C}_Y\text{N}_Z)} - E_{\text{CO}_2(\text{g})} - 1/2E_{\text{H}_2} \quad (1)$$

$$E_{\text{ads}}(*\text{CO}) = E_{(\text{Ni-B}_X\text{C}_Y\text{N}_Z + \text{CO})} - E_{(\text{Ni-B}_X\text{C}_Y\text{N}_Z)} - E_{\text{CO}(\text{g})} \quad (2)$$

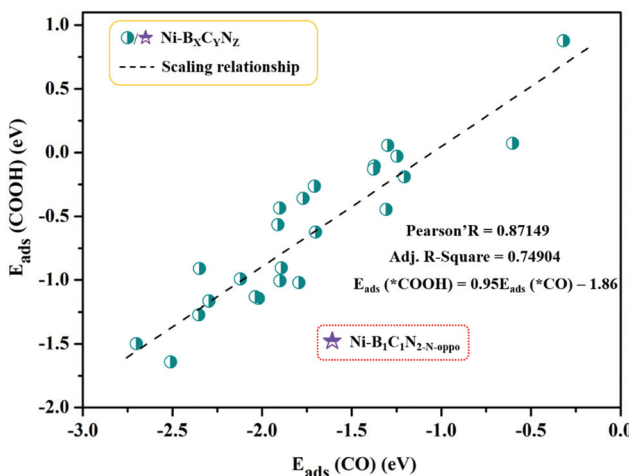


Fig. 4 Scaling relationship between the adsorption energies of *COOH ($E_{\text{ads}}(*\text{COOH})$) and *CO ($E_{\text{ads}}(*\text{CO})$).

The calculated adsorption energies are provided in Table S3,† and the correlation between E_{ads} (*COOH) and E_{ads} (*CO) is shown in Fig. 4. From Fig. 4, the adsorption energy of *COOH and *CO follows a fitted linear relationship $E_{\text{ads}}(*\text{COOH}) = 0.95E_{\text{ads}}(*\text{CO}) - 1.86$, and the scaling relations ($R \approx 0.87$) between them are slightly weakened with scattered points compared to those of pure metal surfaces ($R \approx 0.96$).⁶³ A special case is Ni-B₁C₁N_{2-N}-oppo, which deviates greatly from the overall linear correlation. Note that on the pure metal surfaces, the *COOH and *CO are uniformly adsorbed to the surface metal atoms *via* single coordination mode (C-M bond). However, in our Ni-B_XC_YN_Z systems, the change of coordination environment, especially with the introduction of B, leads to versatile coordination modes (e.g., single or dual-site coordination) that vary greatly with the type and number of coordination elements. The versatility in the adsorption structures could be the main reason for the weakened linear scaling between *COOH and *CO in Ni-B_XC_YN_Z.

CO₂RR activity and selectivity

As discussed above, the adsorption characteristics of CO₂, *COOH and CO varied with the change of coordination environments, which indicates that the activity and product selectivity would be significantly affected. Thus, we further explored the activity of the CO₂RR to CO. The CO₂RR pathways can be divided into four elementary steps (Fig. 5a): (i) CO₂ adsorption: CO₂ (g) + * → *CO₂; (ii) *COOH formation: *CO₂ + H⁺ + e⁻ → *COOH; (iii) *CO formation: *COOH + H⁺ + e⁻ →

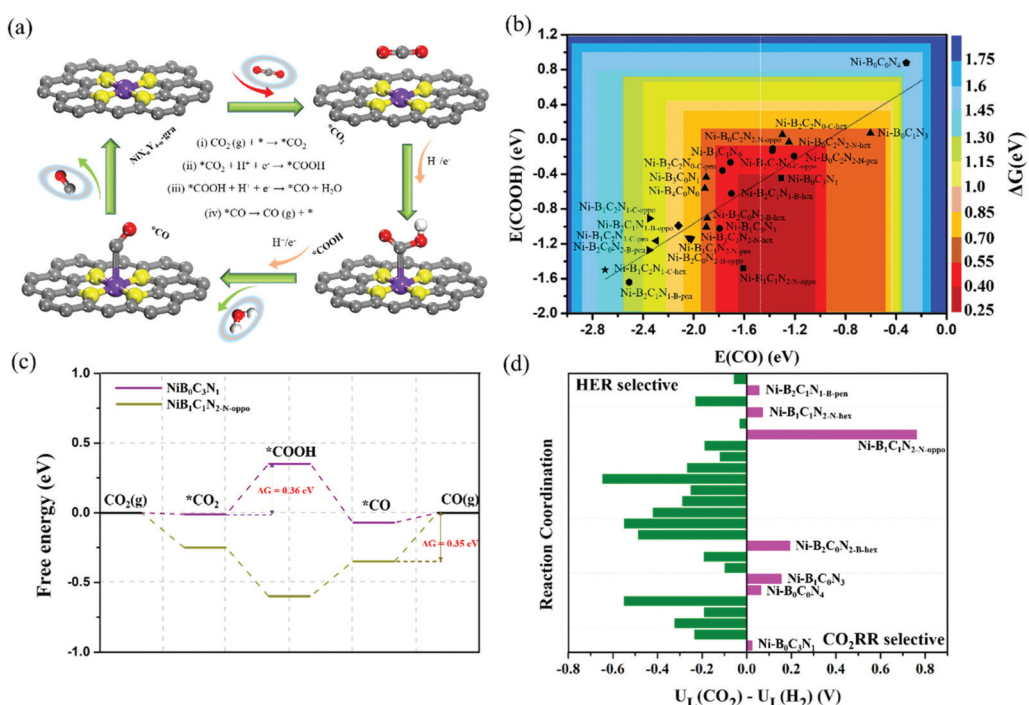


Fig. 5 Schematic of the four-step reaction processes of CO₂ reduction to CO on Ni-B_XC_YN_Z (a). 2D volcano plot for CO evolution on Ni-B_XC_YN_Z catalysts (b). Free energy diagrams for the CO₂RR on Ni-B₀C₃N₁ and Ni-B₁C₁N_{2-N}-oppo (c). The limiting potential difference between CO₂ reduction ($U_L(\text{CO}_2)$) and HER ($U_L(\text{H}_2)$) on various Ni-B_XC_YN_Z catalysts (d).

*CO + H₂O; and (iv) *CO desorption: *CO → CO (g) + *. The Gibbs free energy changes (ΔG) for each reaction step are provided in Tables S4–S6,† and the ΔG ($\Delta G = \max(\Delta G_{ii}, \Delta G_{iii}, \Delta G_{iv})$) of the most difficult step, or the most sluggish step, dictates the reaction activity. The activity as a function of the binding energy of COOH and CO is shown in Fig. 5b. It is worth noting that the activity diagram is divided into two regions. In the right region, the *CO adsorption is relatively weak, and the activity is determined by the potential-dependent protonation of *CO₂ to *COOH (step ii); while in the left region, the *CO adsorption is strong, and the activity is determined by the thermodynamic-dependent *CO desorption (*CO → * + CO_(g), step iv). Clearly, most of the B-coordinated B_XC_YN_Z ($X \neq 0$) catalysts are located in the left region, where ΔG (*CO) varies from −0.35 to −1.42 eV, and the *CO desorption becomes the most difficult step. While Ni–B₂C₂N_{0-C-oppo}, Ni–B₂C₂N_{0-C-hex}, and all the C and/or N coordinated Ni–B_XC_YN_Z ($X = 0$) systems are located in the right region, where the *COOH formation becomes the potential-determining step, and the corresponding free energy change of *CO₂ hydrogenation to *COOH varies from 0.36 to 1.49 eV. Note that for the B-free Ni–B_XC_YN_Z ($X = 0$), the CO₂RR activity is gradually decreased as the number of coordinated nitrogen atoms increases (Ni–B₀C₃N₁ > Ni–B₀C₂N_{2-N-oppo} ≈ Ni–B₀C₂N_{2-N-pen} > Ni–B₀C₂N_{2-N-hex} ≈ Ni–B₀C₁N₃ > Ni–B₀C₀N₄). Moreover, from the volcano plot, Ni–B₀C₃N₁ (the bottom of the right region) and Ni–B₁C₁N_{2-N-oppo} (the bottom of the left region) stand out as the most active catalysts for the CO₂RR. Along the right region, the Ni–B₀C₃N₁ has the lowest free energy (0.36 eV) for the reaction-limiting step (*CO₂ + H⁺ + e[−] → *COOH); and along the left region, the Ni–B₁C₁N_{2-N-oppo} has the lowest desorption energy for *CO release (−0.35 eV). The detailed free energy diagram of CO₂ reduction on Ni–B₀C₃N₁ and Ni–B₁C₁N_{2-N-oppo} is shown in Fig. 5c.

Furthermore, to compare the computational accuracy of DMol³ with the plane-wave basis method in the VASP code, we choose the six N/C coordinated Ni–B_XC_YN_Z ($X = 0$) systems to calculate the free energy diagram for CO evolution (Fig. S2†), and the free energy change for the potential-determining step (*COOH formation) is summarized in Table S7.† The free energy change predicted by the two methods differs by about 0.06–0.2 eV, and the trend of the predicted activity is similar except for Ni–B₀C₂N₂ and Ni–B₀C₁N₃ (DMol³ predicts a higher activity of Ni–B₀C₂N_{2-N-oppo}/Ni–B₀C₂N_{2-N-pen} over Ni–B₀C₁N₃, while VASP predicts the opposite). Recent studies by Luo *et al.*⁵¹ applied a grand canonical potential kinetics method to predict the reaction mechanism and rates for the CO₂RR over Ni–N₂C₂, Ni–N₃C₁, and Ni–N₄ sites in graphene. They revealed that Ni–N₂C₂ leads to the lowest onset potential (−0.84 V to achieve 10 mA cm^{−2} current density), followed by Ni–N₃C₁ (−0.92 V) and Ni–N₄ (−1.03 V). Moreover, Liu *et al.*⁵² applied a “slow-growth” sampling approach to evaluate the reaction barriers and showed that the Ni-atom coordinated with one N and three C atoms (Ni–N₁C₃) is most active and selective for the CO₂RR. These indicate that the CO₂RR activity tends to decrease with the increase of coordinated nitrogen, which is

qualitatively in good agreement with our DMol³ results based on the simplified computational hydrogen electrode model.

In addition, since the HER usually competes with the CO₂RR, it is thus also important to evaluate the selectivity between CO₂ reduction and HER by comparing their limiting potential (U_L), which is defined as $−\Delta G/e$ (ΔG refers to the most sluggish step). According to this definition, a more positive ΔU_L ($U_L(\text{CO}_2) − U_L(\text{H}_2)$) means a higher selectivity toward the CO₂RR (the limiting free energy change of the competitive HER, ΔG_{HER} , is provided in Table S6†). From Fig. 5d, most of the Ni–B_XC_YN_Z catalysts are selective to the HER except for Ni–B₀C₃N₁, Ni–B₀C₀N₄, Ni–B₁C₀N₃, Ni–B₂C₀N_{2-B-hex}, Ni–B₁C₁N_{2-N-oppo}, Ni–B₁C₁N_{2-N-hex} and Ni–B₂C₁N_{1-B-pen}. Among the 7 CO₂RR selective catalysts, Ni–B₀C₃N₁ and Ni–B₀C₀N₄ are potentially determined by *CO₂ hydrogenation. However, the ΔG for *COOH formation on Ni–B₀C₀N₄ is very high (1.49 eV), making Ni–B₀C₀N₄ selective but less active for the CO₂RR. A recent report by Zhang *et al.*⁶⁴ showed that single Ni²⁺ atoms fourfold coordinated by N and dispersed over a carbon black support displayed excellent CO Faraday efficiency (above 90%) and a low faradaic efficiency to H₂. The higher CO selectivity of the Ni–N₄ catalyst over the HER is in agreement with our predictions of the CO₂RR selective Ni–B₀C₀N₄. Moreover, for the screened Ni–B₁C₀N₃, Ni–B₂C₀N_{2-B-hex}, Ni–B₁C₁N_{2-N-oppo}, Ni–B₁C₁N_{2-N-hex} and Ni–B₂C₁N_{1-B-pen}, the *CO desorption is the most sluggish step, and the corresponding ΔG (*CO) is −0.48 eV, −0.63 eV, −0.35 eV, −0.64 eV, and −1.25 eV, respectively. Since the CO₂RR process requires rapid desorption of *CO to improve the CO selectivity, Ni–B₂C₁N_{1-B-pen} is thus excluded as a viable CO₂RR catalyst due to its over strong *CO adsorption. Note that CO₂ electroreduction can produce many possible products. Previous experimental studies have verified that the N/C coordinated Ni produces CO as the main product, hence we only focus on the CO pathway for the B-free Ni–B_XC_YN_Z ($X = 0$). With regard to the B-coordinated systems, we select the 5 CO₂RR selective catalysts (Ni–B₁C₀N₃, Ni–B₂C₀N_{2-B-hex}, Ni–B₁C₁N_{2-N-oppo}, Ni–B₁C₁N_{2-N-hex} and Ni–B₂C₁N_{1-B-pen}) to further examine the possible HCOOH pathway. From Fig. S3,† the limiting potential required for HCOOH formation is much higher, which indicates that CO is the more preferred product. In addition, since the *CO adsorption on Ni–B₂C₁N_{1-B-pen} is very strong (−1.25 eV), we further studied its potential to produce more deep-reduced products. From Fig. S3e,† the *CO can be further reduced to CH₄ and CH₃OH, and the potential-determining step corresponds to hydrogenation of *CHO to *OCH₂ (0.92 eV).

Activity origin

The activity of an electrocatalyst is known to be governed by its intrinsic electronic structure. To decode the underlying activity origin, we first analyzed the electronic structures, and the projected density of states (PDOS) of Ni and the coordination atoms (B, C and N) on pure Ni–B_XC_YN_Z catalysts is shown in Fig. S4.† Since all the investigated 25 catalysts are non-magnetic, there is no spin-polarization in the calculated electronic states; we thus only plotted one of the spin channels. In the C-

and N-coordinated Ni–B_XC_YN_Z (B-free, $X = 0$), the electronic states around the Fermi level are mainly contributed by the Ni atom for Ni–B₀C₃N₁, Ni–B₀C₂N_{2-N}-oppo, Ni–B₀C₂N_{2-N}-pen, and Ni–B₀C₂N_{2-N}-hex (Fig. S4a, 2b, 2c and 2d†). While in the case of Ni–B₀C₁N₃ (Fig. S4e†) and Ni–B₀C₀N₄ (Fig. S4f†), the Ni electronic states shifted downward, and the Fermi levels are mainly contributed by the N atoms. This can qualitatively explain why Ni–B₀C₁N₃ and Ni–B₀C₀N₄ with higher N-content show lower CO₂RR activity. We further analyzed the Bader charge of the Ni active center in Ni–B_XC_YN_Z (B-free, $X = 0$) (Table S8†), where the Ni atom carries positive charge between +0.625|e| and +0.830|e|. Particularly, we note that the charge of the Ni atom in Ni–B₀C₃N₁ (+0.625|e|) is the lowest, indicating that the Ni atom has more electrons in the outer shell. The previous research studies reveal that the larger magnitude of electrons in the outer shell would facilitate the adsorption of CO₂ and *COOH intermediates thereby accelerating the CO₂RR.^{65,66}

On the other hand, the electronic structures of B-coordinated Ni–B_XC_YN_Z ($X \neq 0$) are much more complicated (Fig. S4g–2y†). Depending on the various coordination environments, the electronic states at the Fermi level are mainly contributed by the Ni atoms (Ni–B₁C₀N₃, Ni–B₂C₀N_{2-B}-pen, Ni–B₂C₀N_{2-B}-hex, Ni–B₁C₂N_{1-C}-oppo, Ni–B₁C₁N_{2-N}-pen and Ni–B₁C₁N_{2-N}-hex), the B atoms (Ni–B₂C₀N_{2-B}-oppo, Ni–B₂C₂N_{0-C}-hex and Ni–B₂C₁N_{1-B}-pen), the Ni and B atoms (Ni–B₃C₀N₁, Ni–B₄C₀N₀, Ni–B₃C₁N₀, Ni–B₂C₁N_{1-B}-oppo, Ni–B₂C₁N_{1-B}-hex and Ni–B₁C₁N_{2-N}-oppo), or the Ni and C atoms (Ni–B₂C₂N_{0-C}-oppo, Ni–B₂C₂N_{0-C}-pen, Ni–B₁C₂N_{1-C}-pen and Ni–B₁C₂N_{1-C}-hex), respectively. In the B-coordinated systems, both the Ni and B atoms are positively charged (except for Ni–B₄C₀N₀ where the Ni atom has a negative charge of −0.0411|e|), and Ni is found to carry much less positive charge (+0.07–+0.49|e|) than B (+1.22–+1.51|e|). Compared to the B-free Ni–B_XC_YN_Z ($X = 0$) (Ni atomic charge: +0.625–+0.830|e|),

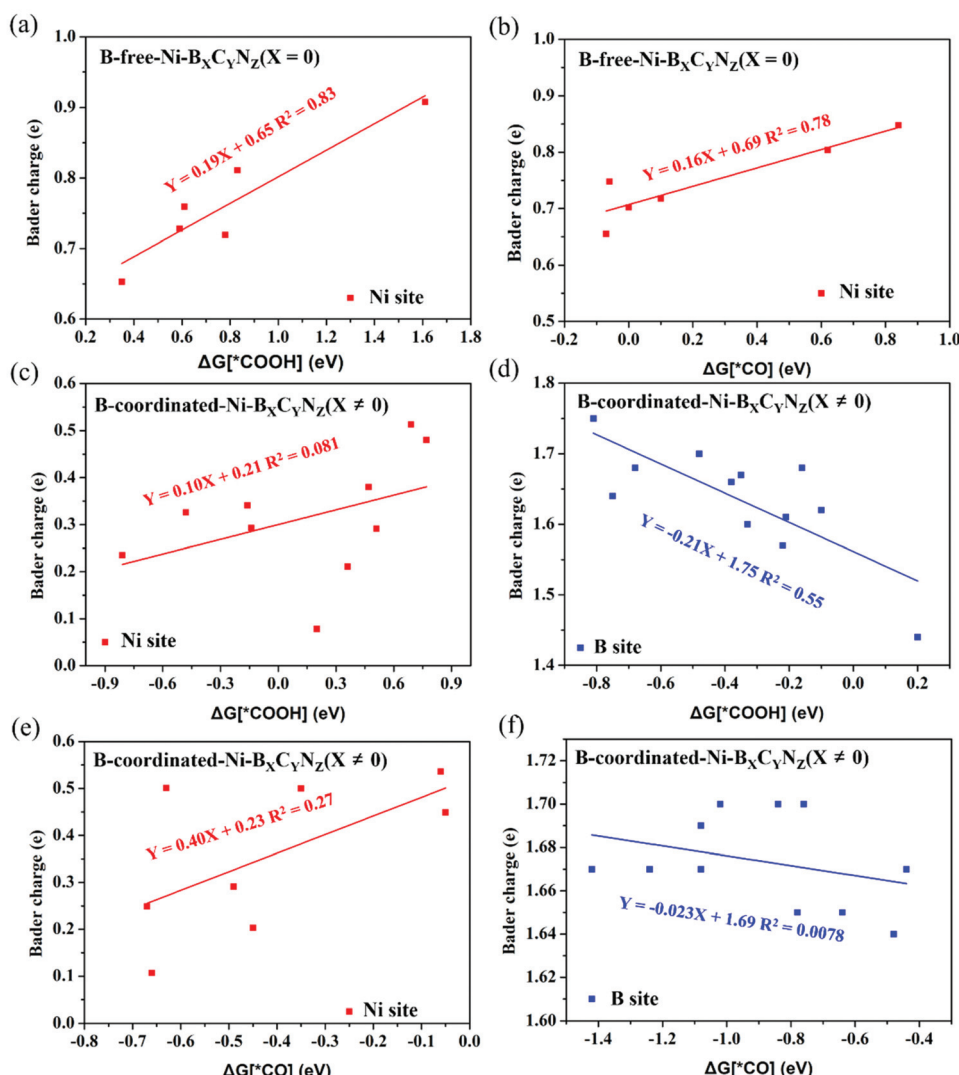


Fig. 6 Bader charge analysis of active sites (Ni or B atom) on *COOH and *CO adsorbed Ni–B_XC_YN_Z. Correlations of the adsorption free energy of *COOH (a) and *CO (b) intermediates with Ni Bader charge on B-free Ni–B_XC_YN_Z ($X = 0$). Correlations of the adsorption free energy of *COOH and *CO intermediates with Ni (c, e) and B Bader charge (d, f) on B-coordinated Ni–B_XC_YN_Z ($X \neq 0$).

the charge of the Ni atom is significantly reduced with the introduction of B (Table S8†). It is known that B behaves like a transition metal and can be the active site in electrocatalysis. The valence electronic configuration of B atoms is $2s^22p^1$ and the sp^3 hybridization of these orbitals can accept and donate the lone-pair electron, which is similar to the d orbital of transition metals.^{45,67,68} In addition, the electronegativity of B (2.04) is close to that of Ni (1.91) and lower than C (2.55) and N (3.04), which also means that the B has transition-metal-like properties. This indicates that Ni and B can work synergistically to function as dual-active sites, which can explain the dual-site adsorption of CO_2 , *COOH and *CO in some of the $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X \neq 0$) catalysts, leading to more favorable *COOH formation and stronger *CO adsorption.

Fig. 6 shows the correlation between the calculated Bader charge of the active center and the adsorption free energy of *COOH and *CO species. In the B-free $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 0$), only the Ni acts as the active site, and both the ΔG (*COOH) and ΔG (*CO) display linear correlation ($R^2(*\text{COOH}) \approx 0.83$ and $R^2(*\text{CO}) \approx 0.78$) with the Ni Bader charge (Fig. 6a and b). However, in the B-coordinated $\text{B}_x\text{C}_y\text{N}_z$ ($X \neq 0$), the active site varies with the coordination environment, and either the single Ni site, single B site or dual Ni-B site can be the catalytic active center. Due to the complexity in the bonding behaviors between CO_2RR intermediates and the active sites, the correlation between the atomic charge of Ni or B (Table S9†) with the adsorption free energy of *COOH and *CO is very poor (Fig. 6c–f). Compared to the single metal atom catalyst, the dual-site or multifunctional site catalysts need a more complex descriptor to explain the intrinsic activity, which would stimulate the future research on the dual- or multi-site electrocatalysts.

Conclusion

In summary, based on DFT computations, we explored the coordination effect on the electrocatalytic activity of Ni SACs towards CO_2 reduction, where the coordination atoms are C, N or B. Our results showed that the coordination environments of the metal atom have a remarkable influence on the adsorption and reaction characteristics. In the C and/or N coordinated $\text{Ni-B}_x\text{C}_y\text{N}_z$ (B-free, $X = 0$), only the Ni atom acts as the active site, which features as a single-site adsorption mode for CO_2RR intermediates. Differently, with the introduction of B ($\text{Ni-B}_x\text{C}_y\text{N}_z$, $X \neq 0$), the coordinated B atom and the Ni metal function as a dual-site and lead to versatile coordination modes of reaction species (single or dual-site adsorption) that varies with the type and number of coordination elements. The versatility in the adsorption modes also results in a weakened linear scaling relationship between *COOH and *CO. Moreover, we established the volcano-type activity plot. It is found that most of the B-coordinated $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X \neq 0$) catalysts are located in the left region where *CO desorption becomes the most difficult step, while all the C and/or N coordinated $\text{Ni-B}_x\text{C}_y\text{N}_z$ ($X = 0$) are located in the right region where the *COOH formation

becomes the potential-determining step. Our results predict that $\text{Ni-B}_0\text{C}_3\text{N}_1$ and $\text{Ni-B}_1\text{C}_1\text{N}_{2-\text{N-oppo}}$ stand out as the most active catalysts for the CO_2RR and have high selectivity over the HER. Hence, by precisely controlling the coordination environment, the experimentally available Ni SACs can be utilized as highly active and selective CO_2RR electrocatalysts. This work provides significant inspiration on boosting the electrocatalytic CO_2RR activity of Ni or other metal catalysts by regulating the coordination environment.

Conflicts of interest

The authors declare no competing financial interests.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (No. 21903008), the Chongqing Science and Technology Commission (cstc2020jcyj-msxmX0382), and the Fundamental Research Funds for the Central Universities (2020CDJQY-A031, 2020CDJ-LHZZ-063). This research used resources of the National Supercomputer Center in Guangzhou.

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