A general strategy for preparing pyrrolic-\(\text{N}_4\) type single-atom catalysts via pre-located isolated atoms

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Single-atom catalysts (SACs) have been applied in many fields due to their superior catalytic performance. Because of the unique properties of the single-atom-site, using the single atoms as catalysts to synthesize SACs is promising. In this work, we have successfully achieved Co\(_1\) SAC using Pt\(_1\) atoms as catalysts. More importantly, this synthesis strategy can be extended to achieve Fe and Ni SACs as well. X-ray absorption spectroscopy (XAS) results demonstrate that the achieved Fe, Co, and Ni SACs are in a M\(_1\)-pyrrolic \(\text{N}_4\) (M=Fe, Co, and Ni) structure. Density functional theory (DFT) studies show that the Co(Cp)\(_2\) dissociation is enhanced by Pt\(_1\) atoms, thus leading to the formation of Co\(_1\) atoms instead of nanoparticles. These SACs are also evaluated under hydrogen evolution reaction (HER) and oxygen evolution reaction (OER), and the nature of active sites under HER are unveiled by the \textit{operando} XAS studies. These new findings extend the application fields of SACs to catalytic fabrication methodology, which is promising for the rational design of advanced SACs.
single-atom catalysts (SACs) have attracted considerable interest due to their superior catalytic activity and unique selectivity towards different chemical reactions, including CO oxidation, hydrogenation, dehydrogenation, and electrochemical reactions. Recently, the application of SACs has been widely extended to various research areas including energy storage systems like Li-S batteries, enzyme catalysis, photocatalysis, and even cellular NO sensor. Therefore, SACs are continuously showing great potentials for wide applications.

The most significant distinction between SACs and other types of catalysts is their unique single-atom-site, which both increases the atom utilization efficiency and tailors the interaction between the reactant and metal atoms through the adsorption and activation process. For example, Yan et al. reported the Pd/graphene SACs showed a high butane selectivity in the hydrogenation of 1,3-butadiene, where the adsorption of 1,3-butadiene on Pt was mainly in a mono-π mode, which differs from that on Pt bulk.

Duchesne and co-workers showed that the Pt single atoms on Au nanoparticles (NPs) exhibited superior catalytic performance in formic acid oxidation. Such a high activity of Pt4Au138 is attributed to the weakened CO adsorption on single and few Pt atoms than that on Pt bulk. Li and co-workers demonstrated that the Pt1/Co3O4 exhibited a much weaker H2 adsorption energy than Pt that on Pt bulk. Li and co-workers demonstrated that the Pt1/Co3O4 exhibited a much weaker H2 adsorption energy than Pt bulk, thus leading to its high catalytic performance in the hydrogenation of ammonia borane.

However, because of their highly unsaturated coordination environment, heterogeneously supported single atoms possess a dramatically increased surface-free energy, which can lead to their aggregation into NPs, particularly at high metal loadings. Recently, some strategies have been developed to achieve relatively high-loading SACs. These methods include zeolite or mesoporous carbon to stabilize single atoms through a confinement strategy, metal-organic frameworks (MOFs) to achieve SACs through high-temperature pyrolysis, metal NPs to achieve SACs through high-temperature migration, and a high-temperature shock wave treatment to achieve high loading SACs. However, there are yet some restrictions on these newly developed methods such as general applicability. Therefore, developing a general approach for the synthesis of various types of SACs is highly desired.

Atomic layer deposition (ALD), a sequential surface reaction relying on a self-limiting process is used to synthesize non-noble single atoms such as Fe and Gr. Unfortunately, it results in a low metal loading and the formation of NPs, especially on carbon-based supports due to the high dissociation barrier and predominantly physical adsorption of metal precursors. Therefore, a new reasonable route to produce SACs by ALD may be achieved by lowering the dissociation energy of the ALD precursor. For instance, this high dissociation energy might be achieved by lowering the dissociation energy of the ALD precursor to decrease the dissociation energy of Co(Cp)2, which may increase the Co loading and more importantly to achieve Co single atoms.

In this work, we firstly verified the strategy by using supported Pt single atoms on N-doped carbon nanosheets (Pt1/NCNS) as the catalyst to synthesize Co SAC through ALD. X-ray absorption spectroscopy (XAS) results reveal singly dispersed Co1 atoms. Interestingly, the Pt1 atoms are still atomically dispersed after Co deposition, as confirmed by XAS and high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM). Furthermore, we also show that this synthesis strategy is general and can also be easily extended to achieve Fe and Ni SACs. X-ray absorption near edge structure (XANES) simulation and extended X-ray absorption fine structure (EXAFS) analysis results demonstrate that the achieved Co, Fe, and Ni SACs are in a M1-pyrrolic-N4 structure. Density functional theory (DFT) calculations show that Pt1 atoms promote the dissociation of Co(Cp)2 into CoCp and Cp fragments. The Co(Cp)2 product further deposited on the substrate through strong chemisorption, thus leading to a higher metal loading and formation of Co1 atoms. These SACs were evaluated under hydrogen evolution reaction (HER) and the nature of single-atom sites are unveiled by the operando XAS studies. Moreover, in the oxygen evolution reaction (OER), the Ni1 SAC showed much higher activity than Fe, and Co1 SACs, which is in line with the DFT predictions.

Results and discussion

Direct fabrication of Pt and Co single atoms on NCNS using ALD. Graphene like NCNS (Fig. 1a and Supplementary Fig. 1) was firstly achieved using C3N4 as the template and glucose as the carbon source. N K-edge XANES results indicate that there are various types of N species including pyridinic, pyrrolic, and graphitic N on NCNS (Fig. 1b). According to the literature, the N defects can coordinate with metal species and form a stable SAC. Therefore, the NCNS substrate shows high potential for the synthesis of SACs. After performing one cycle of Pt ALD on NCNS, well-dispersed Pt single atoms were confirmed by HAADF-STEM (Supplementary Fig. 2) and XAS results (Supplementary Fig. 3). The loading of Pt is as high as 2.0 wt% based on the inductively coupled plasma optical emission spectrometer (ICP-OES) results.

Since the NCNS shows great potential to support Pt1 atoms, it has the possibility to act as a suitable substrate for other types of SACs. Using ALD, we performed one cycle of Co ALD on NCNS with Co(Cp)2 and O2 as the precursors. However, unlike the formation of Pt1 atoms, we found obvious Co NPs/clusters in HAADF-STEM (Supplementary Fig. 4a) and Co-Co coordination from the EXAFS results at the Co K edge (Fig. 1c), indicating the formation of Co NPs (Co-NPs/NCNS). Besides the formation of Co NPs, we also noticed a very low metal loading of Co (0.5 wt%). Such a low loading of Co indicates limited physisorption after the dose of Co(Cp)2 precursor, and the unreacted Co(Cp)2 would be purged away by N2 as illustrated in (Supplementary Fig. 5), leading to a low effective Co deposition. Unfortunately, the weak adsorption effect of Co(Cp)2 on NCNS will form Co NPs due to the weak interaction between Co species and substrate when O2 is introduced. Therefore, the direct utilization of the Co ALD process to produce Co single atoms on NCNS is not practical.

Synthesizing Co, Fe, and Ni SACs using pre-located isolated Pt1 atoms. Inspired by previous research, the dissociation of Co(Cp)2 will be enhanced on the metal surface. In our system, the Pt single atoms are easy to achieve, and the Pt-based catalysts are active in many chemical reactions. Therefore, the singly dispersed Pt1 atoms may have a similar ability and are likely capable to decrease the dissociation energy of Co(Cp)2, which may increase the Co loading and more importantly to achieve Co single atoms. With this idea in mind, we firstly conducted one cycle of Pt ALD on NCNS to achieve Pt1/NCNS and followed up with one cycle of Co ALD to verify our assumption (Fig. 1d). After performing one cycle of Co ALD on Pt1/NCNS substrate, the Co K edge EXAFS of this sample exhibits only one single peak at 1.57 Å (Fig. 2a), which can be attributed to the first shell of Co-Co coordination. Meanwhile, the coordination of Co-Co is...
negligible, indicating that the Co species mostly exist as single atoms from the new synthesis strategy. In the following parts, this sample is denoted as Co1Pt1/NCNS.

In order to further extend the application of this synthesis strategy to other SACs such as Fe and Ni, we have chosen to use Fe and Ni ALD precursors with similar chemical structures as that of Co, i.e., ferrocene and nickelocene, respectively. Similarly, Fe and Ni single atoms also cannot be achieved by ALD on pristine NCNS as revealed by Fe-Fe and Ni-Ni coordination in the EXAFS (Fig. 2b, c) and TEM results (Supplementary Fig. 4). In contrast, the Fe-Fe and Ni-Ni coordination are significantly depressed when performing one cycle of Fe and Ni ALD on Pt1/NCNS (Fe1Pt1/NCNS and Ni1Pt1/NCNS) (Fig. 2b, c). Wavelet-transform EXAFS results further verified the atomically dispersed Co, Fe, and Ni single atoms, which differs from that of their foil and oxide (Fig. 2d, f).

More interestingly, the Pt L3 edge XANES (Fig. 2g) and X-ray Photoelectron Spectroscopy (XPS) (Supplementary Fig. 6) results show inconspicuous differences, indicating the electronic properties of Pt on these M1Pt1/NCNS (M=Fe, Co, and Ni) samples are close to that on Pt1/NCNS. Pt L3 edge EXAFS results further confirm the stable local structure of the Pt1 atom (Fig. 2h). Typical EXAFS fitting on Co1Pt1/NCNS at Pt L3 edge also demonstrates the unchanged atomic nature of Pt single atoms (Supplementary Fig. 7 and Supplementary Table 1). These results indicate that the Pt1 atoms are still atomically dispersed after the deposition of Co, Fe, and Ni. This observation provides strong evidence that the Pt1 atoms only act as the catalyst, while they change the deposition process of Co, Fe, and Ni, but does not lose the atomic dispersion or form a Pt-Co/Fe/Ni cluster, because of the lack of metal-metal coordination in Pt L3, Fe, Co, and Ni K edges EXAFS results. Subsequently, we examined the Co loading through ICP-OES after performing one cycle of Co ALD on pristine NCNS and Pt1/NCNS (Fig. 2i). The ICP-OES results show that the Co loading of Co1Pt1/NCNS (1.8 wt%) is more than three times higher than that of Co-NPs/NCNS (0.5 wt%). Similar results are also observed on Fe and Ni, showing low Fe and Ni loadings on Fe-NPs/NCNS and Ni-NPs/NCNS (0.6 and 0.5 wt%) but much higher on Fe1Pt1/NCNS and Ni1Pt1/NCNS (1.6 and 2.2 wt%), respectively. These results unambiguously demonstrate that the Pt1 atoms change the adsorption model of M precursors to predominantly chemisorption, which promotes the degree of effective deposition. However, when the M ALD was performed on pristine NCNS, physisorption is dominant, thus leading to a low metal loading because of less effective deposition.

Morphology of M1Pt1/NCNS catalysts. The morphology of M1Pt1/NCNS catalysts is characterized by HAADF-STEM. As shown in Fig. 3a, e, i, no observable Pt or M NPs can be found at low magnification. Energy dispersive X-ray (EDX) mapping results show the Pt and M elements are well-dispersed on the NCNS substrate (Fig. 3b, f, j and Supplementary Figs. 8 to 10), indicating their uniform dispersion and high density. At high magnification, high-density bright Pt single atoms (white circles) can be found on Co1Pt1/NCNS (Fig. 3c and Supplementary Fig. 8f), Fe1Pt1/NCNS (Fig. 3g and Supplementary Fig. 9f), and Ni1Pt1/NCNS (Fig. 3k and Supplementary Fig. 10f). All the visible Pt atoms are singly dispersed without any obvious clusters or NPs. Meanwhile, some isolated atoms with less brightness (red circles) were also found to reside nearby the bright Pt1 atoms, which can be assigned to Fe, Co, and Ni single atoms due to their lower Z-contrast. Furthermore, when the electron energy loss spectroscopy (EELS) was measured at the atoms with less brightness on Co1Pt1/NCNS, the signals of Co L2,3 and Ni L2,3 could also be detected (Fig. 3d), confirming the existence of Co and Ni atoms. Similar results can be also found on Fe1Pt1/NCNS (Fig. 3b) and Ni1Pt1/NCNS (Fig. 3i).
Fig. 2 XAS characterizations for \(M_iPt_1/NCNS\) catalysts. a–c Radial distribution curve from FT-EXAFS of Co-NPs/NCNS, CoPt/NCS, Fe-NPs/NCNS, FePt/NCS, Ni-NPs/NCNS, and NiPt/NCS, respectively, as well as their corresponding foil and oxide reference at Co, Fe, and Ni K edges. d–f Wavelet-transformed spectra of Co1Pt1/NCNS, Fe1Pt1/NCNS, Ni1Pt1/NCNS, as well as their corresponding foil and oxide reference at Co, Fe, and Ni K edges. g and h XANES and FT-EXAFS of Pt1/NCNS, CoPt/NCNS, FePt/NCNS, and NiPt/NCNS, respectively, as well as Pt foil and PtO\(_2\) reference at Pt L\(_3\) edge. i ICP-OES results of M (Fe, Co, and Ni) metal loadings.
Fe, Co, and Ni-Pt pairs are absent. Taken together, the EXAFS and HAADF-STEM results demonstrate the single-atom nature of both Pt and M atoms on NCNS.

Structure identification of M SACs. Using this synthesis strategy, we have successfully achieved Co, Fe, and Ni SACs. Unveiling the local structure of these SACs is critical to understand the synthesis mechanism and their catalytic applications. XANES spectroscopy is very sensitive to the local geometric structure of the photon-absorbing atom, which provides a better option to understand the local atomic structure. As shown in Fig. 4a to c, the CoPt1/NCNS, FePt1/NCNS, and NiPt1/NCNS at Co, Fe, and Ni K edge XANES spectra, respectively, show significant differences from their corresponding foil and oxide references. We also notice that the XANES spectra of CoPt1/NCNS, FePt1/NCNS, and NiPt1/NCNS are nearly identical, indicating a similar local coordination environment of M1Pt1/NCNS catalysts. This can be delineated since the k-dependent absorber phase shift of Fe, Co, and Ni, adjacent in the periodic table, is very similar. Taking the first derivative XANES of M1Pt1/NCNS, the energy position of the point of first inflection suggests that the oxidation states of Fe, Co, and Ni are close to +3, +2, and +2, respectively. These observations are perfectly in line with the XPS results (Supplementary Fig. 11).

Focusing on the CoPt1/NCNS catalyst, we first compared the modeling XANES result based on a recently reported Co-C4-O2 type Co 1 atom on graphene using ALD (Supplementary Fig. 12a). Significant differences can be observed between the theoretical simulated XANES and experimental results, indicating the necessity to consider the critical role of N atoms. We then further compared the reported pyridinic-N4-based Co 1-N4 structure, which still shows the obvious difference with our experimental XANES results (Supplementary Fig. 12b), suggesting the existence of defects in this model as well. Then we progressively add one and two end-on oxygen moieties along the axial direction of the pyridinic Co1-N4 structure (Supplementary Fig. 12c, d). Although parts of the experimentally resolved features can be reproduced and achieve a satisfactory agreement with the Co1Pt1/NCNS Co K edge XANES results, the short Co-N...
bonding distance (less than 1.90 Å) disagrees with our EXAFS results (Fig. 2a). Considering that there is a large proportion of pyrrolic-N in the NCNS support, the possibility of a pyrrolic-N4 type Co1-N4 structure cannot be ruled out. As shown in Supplementary Fig. 12e, the simulated XANES result of pyrrolic-N4-based Co1-N4 showed much better conformity than the pyridinic-N4 structure. We then further added one molecule of dioxygen with end on coordination on Co1 atom, the features from “b” to “d” are well reproduced but the pre-edge feature “a” is exaggerated (Supplementary Fig. 12f). However, when we added another molecule of dioxygen on the Co1 atom at the opposite side, all the features from “a” to “d” can be correctly reproduced (Fig. 4d). Similarly, the simulated 2O2-Fe1-pyrrolic N4 and 2O2-Ni1-pyrrolic N4 XANES spectra (Fig. 4e, f) show the most satisfactory results comparing with other DFT optimized structures (Supplementary Figs. 13 and 14). It is important to note, this is the first time that general M1-pyrrolic N4 type SACs can be achieved using ALD, as this type of SACs is usually achieved through high-temperature pyrolysis and ball-milling methods.39,40. Besides the above XANES simulation, we also carried out the EXAFS fitting and the results consistently demonstrate the 2O2-M1-pyrrolic N4 moiety (Fig. 4g to i, Supplementary Figs. 15 to 18, and Table 1). It is worth noting that the EXAFS results perfectly match the DFT results (Supplementary Table 2). In summary, a close local structure of Co1Pt1/NCNS, Fe1Pt1/NCNS and Ni1Pt1/NCNS from simulated...
XANES and EXAFS fitting results strongly suggest the same deposition mechanism for Co, Fe, and Ni ALD on Pt1/NCNS, which indicates that this synthesis strategy can be a universal approach to achieve M1-pyrrolic N4 type SACs using Pt1 atoms.

**Theoretical understanding of deposition process.** Having resolved the structures of M1Pt1/NCNS and Pt1/NCNS, we carried out DFT calculations to elucidate the deposition mechanism. Considering the similar structure of Co, Fe, and Ni ALD precursors and local structures of CoPt1/NCNS, FePt1/NCNS, and NiPt1/NCNS catalysts, typical Co ALD process is chosen for the following theoretical investigation. As revealed by the calculated Gibbs free energies shown in Fig. 5a, the Co(Cp)2 has moderate adsorption ($\Delta G_{\text{ad}} = -0.32$ eV) but very high dissociation energy ($\Delta G_{\text{de}} = 4.91$ eV) on the pristine NCNS. Therefore, the physisorption of Co(Cp)2 should be dominant on pristine NCNS. This physically and weakly adsorbed Co(Cp)2 will lead to a low metal loading from inefficient deposition (Fig. 2i and Supplementary Fig. 5). Owing to the weak physisorption, Co species will easily aggregate because of the lack of strong chemical bonding with the substrate. Therefore, Co NPs were observed in the direct Co ALD process on NCNS (Fig. 1c). However, when Pt1 atoms are involved in the Co ALD process, the deposition model differs considerably. Although the adsorption of Co(Cp)2 on Pt1 atom is slightly weaker ($\Delta G_{\text{ad}} = +0.06$ eV), the dissociation energy of Co(Cp)2 is reduced to $\Delta G_{\text{de}} = 3.54$ eV. The lower dissociation energy can be attributed to the electron transfer from the Cp ligand to the Pt1/NCNS, thus weakening the interactions between Co and Cp, which facilitates the dissociation process (Supplementary Fig. 19 and Supplementary Table 3). Considering it is a gas phase reaction, the Co(Cp) likely diffuses and deposits on the unoccupied pyrrolic N4 sites through chemisorption. Such a strong chemical bonding of Co with pyrrolic N4 sites will form a stable Co1 structure after the O2 pulse to remove the Cp ligand. It is worth noting that the introduction of O2 may also remove the attached Cp ligand on the Pt1 atom and recover the structure of Pt1/NCNS as shown in the Pt L3 XANES and EXAFS results (Fig. 2g, h). As a result, when the Co ALD is performed on Pt1/NCNS, Co single atoms will form and Pt1 atoms will remain atomic dispersion (Fig. 5b). The principle revealed by the theoretical calculation results indicates that this synthesis strategy could be universal for the rational design of SACs (Supplementary Fig. 20).

**Theoretical simulation and electrocatalytic performance of M1Pt1/NCNS.** The capability of application for the achieved M1Pt1/NCNS catalysts was further evaluated in electrochemical reactions. Prior to the experiment, we first studied the projected adsorption ($\Delta G_{\text{ad}} = -0.32$ eV) but very high dissociation energy ($\Delta G_{\text{de}} = 4.91$ eV) on the pristine NCNS. Therefore, the physisorption of Co(Cp)2 should be dominant on pristine NCNS. This physically and weakly adsorbed Co(Cp)2 will lead to a low metal loading from inefficient deposition (Fig. 2i and Supplementary Fig. 5). Owing to the weak physisorption, Co species will easily aggregate because of the lack of strong chemical bonding with the substrate. Therefore, Co NPs were observed in the direct Co ALD process on NCNS (Fig. 1c). However, whenPt1 atoms are involved in the Co ALD process, the deposition model differs considerably. Although the adsorption of Co(Cp)2 on Pt1 atom is slightly weaker ($\Delta G_{\text{ad}} = +0.06$ eV), the dissociation energy of Co(Cp)2 is reduced to $\Delta G_{\text{de}} = 3.54$ eV. The lower dissociation energy can be attributed to the electron transfer from the Cp ligand to the Pt1/NCNS, thus weakening the interactions between Co and Cp, which facilitates the dissociation process (Supplementary Fig. 19 and Supplementary Table 3). Considering it is a gas phase reaction, the Co(Cp) likely diffuses and deposits on the unoccupied pyrrolic N4 sites through chemisorption. Such a strong chemical bonding of Co with pyrrolic N4 sites will form a stable Co1 structure after the O2 pulse to remove the Cp ligand. It is worth noting that the introduction of O2 may also remove the attached Cp ligand on the Pt1 atom and recover the structure of Pt1/NCNS as shown in the Pt L3 XANES and EXAFS results (Fig. 2g, h). As a result, when the Co ALD is performed on Pt1/NCNS, Co single atoms will form and Pt1 atoms will remain atomic dispersion (Fig. 5b). The principle revealed by the theoretical calculation results indicates that this synthesis strategy could be universal for the rational design of SACs (Supplementary Fig. 20).

### Table 1 Structural parameters of the M1Pt1/NCNS (M=Fe, Co, and Ni), foil, and oxide references extracted from quantitative EXAFS curve-fittings at Co, Fe, and Ni K edges.

| Sample | Path     | CNs | R (Å) | $\sigma^2 (\times 10^{-3} \text{Å}^2)$ | $\Delta E_0$ (eV) |
|--------|----------|-----|-------|----------------------------------|------------------|
| Co foil | Co-Co   | 12.0 | 2.49  | 6.1                              | 6.9              |
| Co$_2$O$_4$ | Co-O  | 4.0  | 1.91  | 17.0                             | 1.0              |
| CoPt1/NCNS | Co-O    | 2.0  | 1.98  | 3.3                              | 2.2              |
| NCNS   | Co-N    | 4.0  | 2.08  | 6.4                              | -                |
| Fe foil | Fe-Fe   | 8.0  | 2.46  | 5.3                              | 5.3              |
| Fe$_2$O$_3$ | Fe-O  | 6.0  | 2.85  | 10.0                             | -6.0             |
| FePt1/NCNS | Fe-O   | 2.2  | 1.95  | 2.0                              | -4.2             |
| NCNS   | Fe-N    | 4.2  | 2.02  | 2.9                              | -                |
| Ni foil | Ni-Ni   | 12.0 | 2.48  | 6.1                              | 7.4              |
| NiO    | Ni-O    | 6.0  | 2.05  | 4.2                              | -3.7             |
| NiPt1/NCNS | Ni-O   | 2.2  | 1.99  | 1.5                              | 2.1              |
| NCNS   | Ni-N    | 4.2  | 2.11  | 2.5                              | -                |

CNs coordination numbers, R bonding distance, $\sigma^2$ Debye-Waller factor, $\Delta E_0$ inner potential shift. Errors in the fitting parameters are CN ± 20%, R ± 0.02, $\sigma^2$ ± 20%, and $\Delta E_0$ ± 3.0.

**Fig. 5 Illustration of theoretical understanding of the Co ALD process.**

(a) Theoretical reaction coordinate for Co(Cp)$_2$ deposition on pristine NCNS and Pt1/NCNS. The Gibbs free energies were calculated at $T = 523$ K and $P = 1$ atm. (b) Illustration of Co ALD process on Pt1/NCNS. The white, brown, blue, orange, and silver spheres represent H, C, N, Co, and Pt, respectively.
density of state (pDOS) of various M1-pyrrolic N4 sites. As shown in Fig. 6a, the Co1-pyrrolic N4 and Fe1-pyrrolic N4 show higher DOS near the Fermi level than that of Ni1-pyrrolic N4. As is known, the high electron densities near the Fermi level could facilitate the adsorption of the proton (H\(^+\))\(^{41-43}\). Further calculated hydrogen adsorption free energy on Co1-pyrrolic N4 catalyst is closer to zero than that on Fe1-pyrrolic N4, indicating its relatively higher catalytic performance in HER (Supplementary Fig. 21). The Ni1-pyrrolic N4 shows the weakest H adsorption ability, indicating its unsatisfactory HER activity. This theoretical prediction is confirmed by the experiment, in which the Co1Pt1/NCNS showed higher activity than Fe1Pt1/NCNS and Ni1Pt1/NCNS (Fig. 6b). The higher catalytic activity of Co1Pt1/NCNS than the Pt1/NCNS could be attributed to the contributions of Co atoms in HER (Supplementary Figs. 22 and 23)\(^{44-46}\). The Co1Pt1/NCNS also exhibits good stability without showing any obvious activity decrease after 8,000 cycles durability test (Supplementary Fig. 24). Although the Ni1-pyrrolic N4 site exhibits low catalytic activity in HER, the high d-band center near the Fermi level on the Ni1 atom (Fig. 6a and Supplementary Table 4) shows the promising application in OER\(^{47}\). In the experiment, the Ni1Pt1/NCNS shows the best catalytic performance among various SACs in OER (Fig. 6c), which confirms the theoretical prediction (Supplementary Figs. 25 and 26). The NCNS shows no catalytic activity in OER, indicating the carbon corrosion is not obvious (Supplementary Fig. 27).

For the first time, the synthesis of M1-pyrrolic N4 type SACs has been successfully achieved using heterogeneously supported Pt\(_1\)
atoms as catalysts. This general M$_1$-pyrrolic N$_4$ framework is confirmed by XANES simulation and EXAFS analysis. DFT calculations show that the Pt$_1$ atoms act as the catalyst to modulate the adsorption behavior and promote the dissociation of Co(Cp)$_2$ in Co ALD, which leads to the chemical binding of CoCp on the pyrrolic N$_4$ site and formation of Co single atoms. More importantly, this synthesis strategy can be extended to achieve Fe and Ni SACs. These SACs are evaluated under HER and the nature of single-atom sites are further unveiled by the operando XAS studies. In OER, the Ni$_n$Pt$_1$/NCNS catalyst show much better catalytic activity than the Fe and Co SACs counterparts, which is in excellent agreement with DFT prediction. These new findings extend the application fields of SACs to catalytic fabrication methodology, which is promising for the rational design of advanced SACs. The prominent covalent metal-support interaction bonding of M$_1$ atom (M = Fe, Co, Ni) with the local coordinated N-atoms provides a unique opportunity to further tune the catalytic properties through substrate design.

Methods

Synthesis of g-C$_3$N$_4$ Twelve grams of urea was put into an alumina crucible with a cover and then heated to 850 °C at a rate of 3 °C/min in a muffle furnace and maintained at this temperature for 4 h. After cooling down to room temperature, ~600 mg of bulk g-C$_3$N$_4$ was obtained.

Synthesis of N-doped carbon nanosheet. The synthesis method for the N-doped carbon nanosheets is based on a previous report. Typically, 500 mg g-C$_3$N$_4$ was mixed with 2.16 g glycerol and dispersed in 40 mL of deionized water under sonication for 5 h. After that, the suspension was transferred to a Teflon autoclave and heated at 140 °C for 11 h in an oven. The product was collected by centrifugation and washed with water and ethanol several times, then dried under vacuum at 60 °C overnight. After that, the dried powder was heated up to 900 °C in Ar at a rate of 5 °C/min and maintained at this temperature for 1 h to achieve the N-doped carbon nanosheet (NCNS).

Synthesis of Pt$_1$/NCNS. Pt was deposited on the as-prepared NCNS by ALD (Savannah 100, Cambridge Nanotechnology Inc., USA) using trimethyl(methylcyclopentadienyl)platinum (IV) (MeCpPtMe$_3$) and O$_2$ as precursors. High-purity N$_2$ (99.9999%) was used as both a purging gas and carrier gas. The deposition temperature was kept at 150 °C, while the container for MeCpPtMe$_3$ was heated to 65 °C to provide a steady flux of Pt to the reactor. The manifold was kept at 115 °C to avoid any condensation. The timing sequence of one cycle Pt ALD was 15s and 30s for MeCpPtMe$_3$ exposure and N$_2$ purge respectively. Subsequently, the ALD chamber was heated up to 250 °C, with 30s O$_2$ exposure and 30s N$_2$ purge to remove the ligand.

Synthesis of M$_n$Pt$_1$/NCNS (M = Fe, Co, and Ni). The M ALD is conducted on the as-prepared Pt$_1$/NCNS. The precursor used for Co, Fe and Ni are Ferrocene (Fe(Cp)$_2$), Cobaltocene (Co(Cp)$_2$) and Nickelocene (Ni(Cp)$_2$). High-purity N$_2$ (99.9999%) was used as both a purging gas and carrier gas. The deposition temperature was kept at 250 °C, while the container for M(Cp)$_2$ was heated to 90 °C to provide a steady flux of precursor to the reactor. The manifold was kept at 140 °C to avoid any condensation. The timing sequence of one cycle M ALD was 15s and 30s for M(Cp)$_2$ exposure and N$_2$ purge respectively. Subsequently, the ALD chamber was heated up to 300 °C with 30s O$_2$ exposure and 30s N$_2$ purge to remove the ligand.

Instrumentation. TEM samples were prepared by drop-casting an ultrasonicated solution of dilute high-performance liquid chromatography grade methanol solution onto a lacey carbon grid. The TEM characterization was carried out on a FEI Titan Cubed 80–300 kV microscope equipped with spherical aberration correctors (for probe and image forming lenses) at 200 kV. Atomic-resolution high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) images were taken using a double spherical aberration-corrected FEI Themis microscope operated at 300 kV. The corresponding inner and outer collection semi-angles for HAADF imaging were 48–200 mrad. The Imaging was performed at a collection semi-angle of 28.7 mrad with a dispersion of 0.5 eV/channel. The metal loadings were analyzed using an inductively coupled plasma optical emission spectrometer (ICP-OES) with samples dissolved in hot fresh aqua regia overnight and filtered.

Electrochemical measurements. The electrochemical measurements were performed using a glassy carbon rotating-disc electrode (Pine Instruments) as the working electrode with carbon paper and a standard hydroxide electrode as the counter and reference electrodes, respectively. Ink for the electrochemical measurement was prepared by adding 2 mg of the catalyst into 1 mL ethanol, and Nano (5% solution, Sigma-Aldrich, 20 µL), followed by sonication for 10 min. A working electrode was prepared by spreading the ink (100 µL) formed by the glassy carbon electrode. The HER test was carried out in 0.5 M H$_2$SO$_4$ with a scan rate of 0.01 V s$^{-1}$. The durability test was carried out in 0.5 M H$_2$SO$_4$ between −0.1 and 0.4 V at a scan rate of 0.1 V s$^{-1}$. The OER LSV polarization curves were measured in an O$_2$-saturated electrolyte at a sweep rate of 5 mV s$^{-1}$. The durability test was carried out on a constant current density of 10 mA/cm$^2$. The measured potential against the reference electrode was converted to a reversible hydrogen electrode (RHE).

XAS measurements. XAS measurements were conducted on the 061D superconducting wiggler at the hard X-ray microanalysis (HXMA) beamline at the Canadian Light Source (CLS) and beamline 20-ID-C at the CLS@APS of the Advanced Photon Source (APS), Argonne National Laboratory (ANL). At HXMA beamline, each spectrum was collected using fluorescence yield mode with a Canberra 32 element Ge detector. The beamline initial energy calibration for the different edges was made by using the corresponding metallic foils from Exafs Materials, and the same metallic foil was further set between two FMB Oxford ion chamber detectors downstream to the sample, making in-step energy calibration available for each individual XAFS scan.

At 20-ID-C beamline, the XAS measurements were conducted at A Si (111) fixed-exit, double-crystal monochromator was used. Harmonic rejection was facilitated by detuning the beam intensity 15% at ~1000 eV above the edge of interest. The measurements were performed in fluorescence mode using four-element Vortex Si Drift detector. Details on the beamline optics and instruments can be found elsewhere. Operando XAS measurements were performed with catalyst-coated carbon paper using a custom-built cell. The carbon paper was pretreated with concentrated nitric acid at 80 °C overnight to ensure thorough electrolyte wetting. The catalyst ink was prepared with the same method as for the electrochemical measurements. The catalyst ink was dropped onto the carbon paper, and the backside of the carbon paper was taped with the Kapton film as the working electrode to ensure the entirety of the electrocatalyst had access to the H$_2$SO$_4$ electrolyte. Carbon paper and Ag/AgCl were used as the counter and reference electrodes. To collect the XAFS spectra during the HER process, the cathodic voltages were held at constant potentials during the operando experiment from 0.05 to ~0.05 V (vs RHE), respectively.

XANES modeling. To further understand the nature of experimental resolved XANES features and address them to the structural and chemical nature of M (Fe, Co, and Ni) site occupancy and their corresponding local structural environment, DFT guided XANES theoretical modeling was carried out by using code FDMNES, following standard procedure. For instance, a Co centered cluster was developed based on the structure model predicated by DFT optimized structures. The radius of the cluster is around 6 Å, corresponding roughly to the detection limit of the XAFS data.

EXAFS data analysis. The data reduction was using codes Athena. The followed procedure reported reference. In brief, the first inflection point of the XANES edge was defined as the experimental E$_0$, the post-absorption edge background was estimated and removed by cubic spline fits. Based on the local structural environment predicted by DFT theoretical modeling, the scattering amplitudes and phase shifts were calculated using FEFF and further used for R-space curve fitting. The k ranges for FT-EXAFS for Pt L$_3$, Fe, Co, and Ni K edges are 3.15–11.57, 2.50–11.16, 2.37–10.90, and 2.33 to 10.48 Å$^{-1}$, respectively.

Theoretical and computational methods. First-principles DFT calculations were performed by using the spin-polarized Kohn–Sham formalism with the generalized gradient approximation (GGA) of Perdew–Burke–Ernzerhof (PBE) as implemented in the Vienna ab initio Simulation Package (VASP). The valence electronic states of all atoms were expanded in plane-wave basis sets with a cutoff energy of 400 eV, and gamma points were used for Brillouin zone integration. All atoms were allowed to relax until the forces fell below 0.02 eV Å$^{-1}$. The energy convergence criterion was set to 10$^{-4}$ eV. The zero-point vibrational energy (ZPE) and zero-point corrections were performed by vibrational frequency analysis. We applied a graphene supercell with the surface periodicity of 8 × 8 including 128 atoms as a basis to construct the M-pyrrolic N$_4$ moieties. A vacuum region of 15 Å in the normal direction of the graphene plane was created to ensure negligible interaction between mirror images of the supercells.

Data availability. The data that support the findings of this study are available from the corresponding author upon reasonable request.

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Author contributions
Junjie Li and X.S. designed the project. Junjie Li performed the catalyst preparation, catalytic tests, XAS experiment, and wrote the manuscript; Y.J., C.X., and J. Li performed the DFT calculations. Q.W., D.W., and M.G. conducted the HAADF-STEM images acquisition and image analyses; N.C. carried out the XANES and EXAFS data analysis. M.N.B., K.A., K.D., D.M.M., Z.F., and W.L. assisted to run the XAS experiment; L.Z., R.L., and T.-K.S. gave constructive suggestions; X.S. supervised the project. All authors reviewed the manuscript.

Competing interests
The authors declare no competing interests.

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