Computational Screening of Single-Metal-Atom Embedded Graphene-Based Electrocatalysts Stabilized by Heteroatoms

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Metal-N-doped carbon is a promising replacement for non-precious-metal catalysts such as Pt for the oxygen reduction reaction (ORR) in polymer electrolyte membrane fuel cells (PEMFCs). Although these materials have relatively good catalytic activity and are cost-effective, they still have lower ORR activity than Pt, and so improving their performances is greatly required. In this study, high-throughput screening was employed based on density functional theory (DFT) calculations to search for good candidate catalysts with a transition metal atom coordinated by heteroatoms (B, N, S, O, and P) embedded in a graphene structure. In addition, coordinating a transition metal with two types of heteroatom dopants in a graphene structure was also considered. We calculated the binding energies of ORR intermediates on metal-heteroatom-based graphene structures because they are known to play a key role in ORR. Based on our results, the new group of electrocatalysts imparts excellent ORR activity for PEMFCs, and we suggest that our approach provides useful insight into exploring other promising candidate catalysts.

Keywords: heteroatom doping, metal-nitrogen-doped carbon, single-metal-atom catalysts, oxygen reduction reaction, electrocatalysts, computational screening

INTRODUCTION

Developing high-performance oxygen reduction reaction (ORR) catalysts is a key for clean energy technologies such as fuel cells and metal-air batteries. ORR activity can be determined using the adsorption energy of oxygen intermediates as a descriptor. Although Pt invokes a high reaction rate due to its appropriate adsorption strength of oxygen intermediates (Nørskov et al., 2004), its high price and scarcity are impediments to its practical use in industry. Therefore, numerous efforts have been made to develop catalysts composed of cheap and abundant elements to replace Pt and Pt-based catalysts.

Metal single atom materials coordinated to nitrogen (M-N-Cs), especially the Fe-N-C catalyst (Lefèvre et al., 2009; Chen et al., 2019; He et al., 2020; Lu et al., 2020), have attracted research interest as the most promising alternative to Pt. Numerous studies of M-N-Cs have focused on modifying the electronic structure of the active site by controlling the coordination environment or changing the d-block element at the metal center. For example, Xu et al. computationally screened M-N-Cs by tuning the metal species and number of nitrogen atoms to be coordinated with the metal atom (Xu et al., 2018). They found that Fe-pyridine/pyrrole-N₄ catalysts lying near the top of a volcano plot showed the highest ORR activity, which indicates that it would be
challenging for M-N-C catalysts with only nitrogen ligands to outperform Fe-N-C catalysts. Therefore, a new strategy is required to improve ORR activity beyond the Fe-N-C catalyst.

Recently, a new approach of introducing a p-block element (B, N, O, P, or S) into M-N-Cs that can adjust the adsorption strength of intermediates by changing the electronic and geometric structures has been suggested. For example, Jung et al. found that hydrogen peroxide (H₂O₂) production rate and selectivity can be regulated by adding electron-rich or electron-poor species into Co-N-C (Jung et al., 2020). Their theoretical and experimental results demonstrated that the adsorption energy of *OOH was increased with the addition of electron-rich species such as oxygen, but decreased with the addition of electron-poor species such as protons, thereby affecting the charge amount of the metal center. Similarly, Mun et al. modulated the adsorption strength of Fe atoms in an Fe-N-C catalyst by controlling the charge density of the carbon plane via electron withdrawing and donating properties (Mun et al., 2019). In addition, Shang et al. showed that Cu coordinated with N and S (Cu-N/S-C) formed an unsymmetrical structure with a higher ORR activity than either Cu-N-C or Fe-N-C (Shang et al., 2020). Due to their asymmetric structure, additional π-bonds between the Cu atom and the oxygen intermediates were formed, thereby enhancing the adsorption strength. Although the evidence from many studies suggests that M-N-C catalysts in which a single metal atom is coordinated with two other elements can enhance ORR activity, no one has yet examined all possible combinations.

In this work, we computationally screened all possible atomic combinations in MX₄ and MYX₃ structures, where M is a transition metal atom (3d, 4d, or 5d) and X and Y are heteroatoms (B, N, O, P, or S). First, we obtained structural stability based on the formation energy, and then the stable structures were filtered out and their limiting potentials for ORR were calculated. We found a linear scaling relationship to establish an ORR activity volcano plot from which we were able to derive several potential candidates.

**METHODS**

Spin-polarized density functional theory (DFT) calculations were performed by using the Vienna Ab Initio Simulation Package (VASP) (Kresse and Hafner, 1993; Kresse and Hafner, 1994; Kresse and Furthmüller, 1996a; Kresse and Furthmüller, 1996b) with projector augmented wave (PAW) pseudopotentials (Blochl, 1994; Kresse and Joubert, 1999). Electron-exchange correlation energy values were treated with the Perdew-Burke-Ernzerhof (PBE) functional of the generalized gradient approximation (Perdew et al., 1996). In the expansion of the plane wave, the cutoff energy was set as 400 eV. Geometry relaxation was stopped when the difference in the total force was less than 0.03 eV/Å. To avoid an artificial electrostatic field, dipole corrections were used to compute all of the energy values reported here (Neugebauer and Scheffler, 1992). In our calculations, we used semi-empirical dispersion correction of the DFT-D3(BJ) method (Grimme et al., 2011).

The MX₄ and MYX₃ structures were constructed based on a (5 × 5) supercell graphene with a vacuum layer of 20 Å by using Monkhorst-Pack 3 × 3 × 1 k-point meshes (Monkhorst and Pack, 1976). We adopted a computational hydrogen electrode (CHE) model including proton-coupled electron transfer (Nørskov et al., 2004) for the ORR free energy pathway. Based on this model, the reference electrode potential was a standard hydrogen electrode (SHE) where the free energy of protons can be related to that of 1/2 H₂(g) under conditions of pH = 0 and 1 bar of H₂ gas at 298 K. The Gibbs free energy of the proton-electron transfer step was given by

\[
\Delta G = \Delta E + \Delta ZPE - T \Delta S + \Delta G_U + \Delta G_{sol},
\]

where \(\Delta E\) is the reaction energy obtained from the DFT calculations, \(\Delta ZPE\) is the change in zero-point energy, \(T\) is the temperature (298 K), \(\Delta S\) is the change in entropy, \(U\) is the applied potential to the SHE (when a potential is applied, the free energy is shifted by \(\Delta G_U = -eU\), where \(e\) is the elementary charge of an electron), and \(\Delta G_{sol}\) is a solvation correction of 0.3 eV for *OH and *OOH (Xu et al., 2018).

**RESULTS AND DISCUSSION**

**Computational Screening Procedure for Metal-Heteroatom-Doped Carbon**

As shown in Figure 1, we only focused on the porphyrin-like active sites where the metal center atom is coordinated with four heteroatoms in a carbon plane with two carbon vacancies due to the high computational cost. In the case of dual heteroatom doping, one of the four heteroatoms was changed, resulting in a structure with a 3:1 ratio. From the results, there were 700 possible combinations (= 28 transition metals × 5 host heteroatoms × 5 dopant heteroatoms). The variation of the d-orbital for transition metals ultimately increases the catalytic performance by controlling...
the binding strength of electrocatalytic reaction intermediates in the hydrogen evolution reaction (HER), ORR, the oxygen evolution reaction (OER), and the carbon dioxide reduction reaction (CO2RR) (Xu et al., 2018; Zhu et al., 2019; Park et al., 2021). Therefore, we considered 28 d-block transition metal elements, including 3d (from Sc to Zn), 4d (from Y to Cd), and 5d (from Hf to Au). Heteroatoms including B, N, O, P, and S acted as glue atoms for the transition metal in the heteroatom-doped carbon structures by changing the covalent bonding in the coordinating environment when their p-orbital positions were dissimilar. Moreover, local strain in the carbon lattice plane was introduced at different levels because of the different atomic sizes between the host and dopant heteroatoms. Various forms of metal-heteroatom-doped carbon (M-YX3, M = transition metal, Y = dopant heteroatom, and X = host heteroatom) were considered. For example, the Co-ON3 structure comprises a Co atom fourfold coordinated with one O atom and three N atoms in the presence of two C vacancies within the carbon matrix. In the computational screening process, we first estimated the stability of a metal-heteroatom-doped carbon structure by considering the formation energy and single atom binding energy. Second, the free energy changes for ORR on the stable metal-heteroatom-doped carbon structures were calculated to construct a 2-D volcano plot with ΔG_OOH and ΔG_OH as descriptors to measure ORR activity.

**Figure 2** | (A) Formation energies (E_{form}) of 700 M-YX3. (B) Metal-binding energies (E_b) of 222 M-YX3 with negative E_{form} as a function of transition metals. (C) Stable 66 M-YX3 structures with both negative E_{form} and negative E_b.

E_{form} = E_{M-YX3} - (4\mu_C + \mu_Y + 3\mu_X + E_M),

where E_{M-YX3} is the total energy of the optimized M-YX3 structure; \(\mu_C\), \(\mu_Y\), and \(\mu_X\) are the chemical potentials of carbon, the dopant heteroatom, and the host heteroatom, respectively; \(E_M\) is the energy per one atom of the most stable bulk metal. A negative E_{form} value indicates that the M-YX3 structure is thermodynamically stable. Figure 2A shows the results of plotting the E_{form} values for M-YX3 for each host heteroatom. The metal-heteroatom-doped carbon with N and O host heteroatoms (M-YN3 and M-YO3) had a greater number of negative E_{form} values compared to the other host heteroatoms. The differences in stability between the host heteroatoms are mostly attributed to the differences in their atomic sizes. Since the van der Waals atomic radii of N and O (1.55 and 1.52 Å, respectively) are smaller than C (1.70 Å) (Bondi, 1964; Mantina et al., 2009), it is possible to minimize the local strain in the carbon lattice plane, thus preventing its disruption. Because the van der Waals atomic radii of B (1.92 Å), P (1.80 Å), and S (1.80 Å) are greater than that of C, they cause local strain and disruption of the carbon lattice plane. As a result, from a total of 222 stable structures, 6 of M-YB3, 95 of M-YN3, 107 of M-YO3, 7 of M-YP3, and 7 of M-YS3 are stable.

Next, we calculated the metal-binding energy (E_b) for the 222 thermodynamically stable M-YX3 structures based on the formation energy. Under the applied potential for ORR, metal atoms embedded in a carbon matrix can dissolve into the electrolyte, thereby causing the matrix to lose its active sites as follows:

\[ M - YX_3 \rightarrow YX_3 + M^{n+} + ne^- \]
Thus, the corresponding dissolution reaction energy at applied potential $U$ was calculated by using the followed equation:

$$\Delta G_{M-YX_3 \rightarrow YX_3} = G_{YX_3} + G_{M^\text{aq}} + n\epsilon U - G_{M-YX_3},$$

where $G_{M^\text{aq}}$ is calculated from experimental standard dissolution potential $U_0^{\text{VRHE}}$ and $n$ is the number of electrons transferred during the dissolution reaction. The free energy of the bulk metal ($E_{M(bulk)}$) was computed as

$$G_{M^\text{aq}} = E_{M(bulk)} + n\epsilon U_0^{\text{VRHE}}.$$

Therefore, we define the metal-binding energy using the dissolution reaction energy at operating ORR condition $U = 0.8 \ V_{RHE}$ as follows:

$$E_b = -\Delta G_{M-YX_3 \rightarrow YX_3} = G_{M-YX_3} - \left( G_{YX_3} + G_{M^\text{aq}} + n\epsilon U \right),$$

where $\Delta G_{M-YX_3 \rightarrow HX_3}$ is the dissolution reaction energy. Figure 2B shows the metal-binding energy of 222 M-YX$_3$ at 0.8 $V_{RHE}$ as a function of the various transition metals. The $E_b$ value of M-YO$_3$ is more positive than that of M-YN$_3$, which means that the N host heteroatoms can stabilize the metal center more readily than the O host heteroatoms. In the M-YN$_3$ structures, the binding stability with the 3d transition metals was better than with the 4d and 5d transition metals. As a result, 66 M-YX$_3$ structures were identified as stable based on the criteria of both $E_{\text{form}} < 0$ and $E_b < 0$ (Figure 2C).

Considering the adsorption characteristics of key reaction intermediates OOH, O, and OH during ORR is highly important for understanding the difference in electrocatalytic performance between different metal-heteroatom-doped carbon structures, and thus offers guidelines for identifying the electrocatalyst with the best ORR activity. Supplementary Table S1 summarizes all of the free energy levels of the ORR intermediates in structurally stable metal-heteroatom-doped carbon. To reduce the computational cost, we only considered the metal atom center as the active site and assumed that the associative mechanism of ORR occurs on the M-YX$_3$ structure. The four-electron transfer pathway for ORR under acidic conditions, which has been well studied previously (Kulkarni et al., 2018), proceeds as follows:

\[
\begin{align*}
* + O_2 + H^+ + e^- & \rightarrow OOH^+ \\
OOH^+ + H^+ + e^- & \rightarrow O^+ + H_2O \\
O^+ + H^+ + e^- & \rightarrow OH^- \\
OH^- + H^+ + e^- & \rightarrow * + H_2O
\end{align*}
\]

Thus, the overall four-electron pathway for ORR can be summarized as

$$O_2 + 4H^+ + 4e^- \rightarrow 2H_2O$$
The elementary reaction step of the ORR with the maximum free energy change ($\Delta G_{\text{max}}$) value is defined as the potential-determining step (PDS). Moreover, a PDS with the positive $\Delta G$ value is thermodynamically unfavorable for the ORR reaction, and results in a large overpotential. The relationship between the limiting potential ($U_L$) and a negative $\Delta G_{\text{max}}$ can be expressed as

$$U_L = -\frac{\Delta G_{\text{max}}}{\epsilon}$$

The free energy diagrams for metal-heteroatom-doped carbon structures having a positive $U_L$ value are shown in Figure 3A. The $U_L$ values of Co-N4, Fe-N4, Ni-N4, Zn-N4, Mn-N4, and Pt-N4 were found to be positive. The PDSs of Co-N4, Fe-N4, and Mn-N4 consist of the OH* formation step ($U_L = 0.71$, 0.33, and 0.21 V$_{RHE}$, respectively). The OOH* formation step is the PDS for Ni-N4 and Pt-N4 ($U_L = 0.45$ and 0.04 V$_{RHE}$). Thus, M-N4 structures with a strong OH* adsorption strength have the OH* removal step as the PDS, whereas those with a weak OH* adsorption strength have the OOH* formation step as the PDS because O$_2$ activation is more difficult than in the counterparts with strong OH* adsorption. The PDS of the Zn-N4 structures was the O formation step due to relatively weak O* adsorption strength compared to the other M-N4 structures. From the M-YN3 group, Co-ON3, Ni-SN3, Fe-ON3, Fe-SN3, and Fe-PN3 all showed a positive $U_L$; the PDSs of Co-ON3, Ni-SN3, Fe-ON3, and Fe-PN3 were the OH* removal step with $U_L = 0.79$, 0.46, 0.09, and 0.07 V$_{RHE}$, respectively, while the PDS of Fe-SN3 was the O formation step due to relatively strong OOH* adsorption strength compared to the other M-N4 structures.

Linear scaling correlations among the ORR intermediates on the structurally stable metal-heteroatom-doped carbon structures were observed for $\Delta G_{\text{OOH}}$ or $\Delta G_O$ vs. $\Delta G_OH$. The slope and intercept for the correlation between OH and OOH were 0.92 and 3.07, respectively, with a correlation coefficient ($R^2$) value of 0.97, which is consistent with previous research (Calle-Vallejo et al., 2017; Kulkarni et al., 2018; Wan et al., 2019) (Figure 3B). Meanwhile, the scaling relationship between OH and O+OH was highly linear with a slope value of 1.24, an intercept value of 0.37, and an $R^2$ value of 0.79. Figure 3C shows the scaling relationship between O and OH. The slope and the intercept for the correlation between OH and O were 1.16 and 1.41, respectively, with an $R^2$ value of 0.76. The O intermediates caused more oxidation of metal than OOH and OH because O species must form a double bond with the metal atom, resulting in more structural changes compared to the initial structure before the optimization. Therefore, the scaling relationship between O and OH has a larger variance than that between OOH and OH. Remarkably, for the structurally stable metal-heteroatom-doped carbon structures, the PDS is either the OOH* formation step or the OH* removal step. As shown in Supplementary Table S1, only four of the metal-heteroatom-doped carbon structures have either the O* formation step or the OH* formation step as the PDS. It is well known that the free energy changes in the OOH* formation step and the OH* removal step are determined by the relationship between $\Delta G_{\text{OOH}}$ and $\Delta G_{\text{OH}}$, and thus the $U_L$ can be predicted based on this (Kulkarni et al., 2018).

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**Figure 4** (A) Catalytic activity toward the H$_2$O production (black solid line) and H$_2$O$_2$ production (green solid line). (B) A 2-D volcano map with $\Delta G_{\text{OOH}}$ and $\Delta G_O$ as descriptors for the theoretical limiting potential of ORR on M-YX3 with a positive limiting potential.
General Activity Volcano Mapping for ORR on the Metal-Heteroatom-Doped Carbon Structures

We first investigated the ORR selectivity of 11 M-YX₃ structures with a positive limiting potential. The equilibrium potential for the four and two electron pathways are shown in the dashed line at 1.23 and 0.7 eV, respectively (Figure 4A). According to the results, Mn-N₄, Zn-N₄, Ni-N₄, and Pt-N₄ had a lower overpotential of H₂O₂ production than H₂O production. These results were consistent with previous theoretical findings (Jung et al., 2020). Figure 4B shows a 2-D volcano map for the theoretical limiting potential of ORR on various metal-heteroatom-doped carbon structures favoring the four-electron pathway for the H₂O production by using ΔGOOH and ΔGOH as descriptors. Because Uᵣ is more positive in the red area, its optimum is at ΔGOOH = 3.69 eV and ΔGOH = 1.23 eV. The dashed line represents the scaling relationship between ΔGOOH and ΔGOH (ΔGOOH = 0.92 × ΔGOH + 3.07). Interestingly, the O-dopant heteroatom shifted Co-N₄ closer to the ORR optimal value (ΔGOOH increased from 3.60 to 3.70 eV and ΔGOH from 0.71 to 0.79 eV). When an electron-rich heteroatom species is substituted for an N atom, this positively changes the charge state of the Co atom. Thus, changing the coordinating heteroatom from -N₄ to -ON₃ weakens the adsorption strength of the ORR intermediates on the structure. This is because the electronegativity of O (3.44) is higher than that of N (3.04), so the metal coordinated by O is more oxidative than the one by N, resulting in the adsorption strength being weakened.

It has already been reported that Co-N-C is a good catalyst for ORR through previous DFT computational screening studies and experimental validation (Zheng et al., 2016). Using O-dopant as a heteroatom has been reported for an O-C(Al) catalyst synthesized by using isomorphic metal-organic framework MIL-53(Al, Ga) (Yang et al., 2020) or a multiwalled carbon nanotube catalyst (Jiang et al., 2019). Moreover, the role of O-dopant heteroatoms has been elucidated in Co-NG(O) catalysts (Jung et al., 2020). The Co-ON₃ structure derived from our screening results is stable and shows the highest ORR efficiency. Based on our results and those from previous studies, we expect that it can be successfully implemented experimentally.

In this study, to simplify the procedure, we calculated the adsorption energies of the oxygen intermediates using a metal atom as the active site. However, for the materials doped with low-electronegativity heteroatoms, such as P, the oxygen may adsorb on the heteroatom rather than on the metal atom, thereby stabilizing the structure (Dipojono et al., 2019). In addition, as the applied potential increases, the oxygen intermediates can be poisoned at the active site, resulting in different adsorption configurations as well as different ORR mechanisms (Li et al., 2016). These limitations will be assessed further in the future research. Nonetheless, our work is worth noting that a large-scale screening of metal-heteroatom-doped carbon materials that include many unknown combinations was conducted and the synthesis feasibility was determined based on the set of stability criteria. Finally, we provided the ORR activity data of those stable catalysts and demonstrated an unreported promising catalyst.

CONCLUSION

We have developed a computational screening process for the efficient metal-heteroatom-doped carbon ORR catalysts. M-N₄ structures comprising 28 transition metals in conjunction with five dopant heteroatoms (B, N, O, P, and S) were constructed. Structural stability was evaluated by using the formation energy and the metal binding energy on heteroatom-doped carbon. We were able to identify 66 metal-heteroatom-doped carbon structures that were stable through rigorous stability testing. To compare the ORR activities of the 66 structurally stable metal-heteroatom-doped carbon structures, we calculated the free energy changes for ORR and selected 11 metal-heteroatom-doped carbon structures with a positive limiting potential. Finally, a 2-D volcano map constructed using ΔGOOH and ΔGOH as descriptors revealed that Co-ON₃ was located nearest to the optimal point for ORR catalysis. This high-throughput screening process provides a promising path for the rational design of heterogenous electrocatalysts for energy conversion as well as scientific insight into the effect of heteroatom doping on carbon structures.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusion of this article will be made available by the authors, without undue reservation. The coordinates of 66 structures and corresponding DFT energies used in this study are available on Catalysis-hub.org (Winther et al., 2019) under following link: https://www.catalysis-hub.org/publications/AraComputational2022.

AUTHOR CONTRIBUTIONS

JWH directed and supervised this research work. AC and BJP designed the research and performed the theoretical calculations. AC, BJP and JWH co-wrote the manuscript. All of the authors discussed the results and revised the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fchem.2022.873609/full#supplementary-material
