INORGANIC CHEMISTRY







FRONTIERS

RESEARCH ARTICLE

View Article Online
View Journal | View Issue



Cite this: *Inorg. Chem. Front.*, 2023, **10**, 5044

Two-dimensional conductive metal—organic frameworks as efficient electrocatalysts for oxygen evolution and reduction reactions†

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It is vital to search for efficient and stable oxygen evolution reaction (OER) and oxygen reduction reaction (ORR) electrocatalysts for the development of metal—air batteries. Herein, we systematically investigated a series of TMN $_x$ O $_{4-x}$ -HTC (TM = Fe, Co, Ni, Ru, Rh, Pd, Ir and Pt; x = 0-4; HTC = hexatribenzocyclyne) analogs of two-dimensional (2D) electrically conductive metal—organic frameworks (MOFs) as potential electrocatalysts for the OER and ORR by using density functional theory calculations. The calculated results exhibit good thermodynamic and electrochemical stabilities of the designed TMN $_x$ O $_{4-x}$ -HTC. The OER and ORR catalytic activity of the designed catalyst is governed by the interaction strength between the intermediates and the catalyst, and this interaction can be tuned by adjusting TM atoms and the local coordination number of N/O atoms. CoN $_3$ O $_1$ -HTC is found to be the best OER catalyst with an overpotential η^{OER} of 0.29 V, and RhN $_2$ O $_2$ -HTC exhibits the lowest ORR overpotential η^{ORR} of 0.20 V. Importantly, RhO $_4$ -HTC, RhN $_2$ O $_2$ -HTC and CoN $_1$ O $_3$ -HTC are predicted as efficient bifunctional catalysts for the OER and ORR. Moreover, the kinetics simulation verifies the four-electron ORR pathway with high activity and selectivity toward H $_2$ O production. The results not only contribute to designing and searching for efficient OER and ORR electrocatalysts but shed light on the opportunities to explore electrochemical applications based on 2D MOE materials

Received 13th June 2023, Accepted 14th July 2023 DOI: 10.1039/d3qi01112g

rsc.li/frontiers-inorganic

1. Introduction

The development of sustainable and renewable energy technologies is of great significance to solve the increasingly serious global energy crisis and environmental pollution problems.^{1,2} Among various technologies, rechargeable metal-air batteries have attracted much attention owing to their high theoretical energy density and environmental compatibility.³ Charging

and discharging processes are driven by the electrochemical oxygen evolution reaction (OER) and oxygen reduction reaction (ORR) processes.4 The OER and ORR with sluggish kinetics require high overpotential and thus lead to low energy efficiency which hinders the application of metal-air batteries.3 Ir/Ru oxides^{5,6} and Pt-based materials^{7,8} are known as the state-of-theart catalysts for the OER and ORR, respectively. Generally, only the metal atoms on the surface of the bulk material could react with reactive species.9 Therefore, only a small portion of metal atoms in the bulk material could participate in the catalytic reaction. It should be noted that the uniformly exposed metal active sites on two-dimensional (2D) materials compared with that of bulk materials could provide promising avenues for exploring alternative catalysts. Moreover, the current studies on 2D material-based catalysts for the OER and ORR are usually investigated one at a time, 10-12 thus, making it difficult to understand the overall catalytic performance of the catalyst. Specifically, to a certain extent, the use of bifunctional electrocatalysts could also reduce costs and simplify procedures since the working conditions of bifunctional catalysts are the same. Hence, exploring efficient 2D material-based bifunctional electrocatalysts for both the OER and ORR is highly desirable for the development of metal-air batteries.

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[†]Electronic supplementary information (ESI) available. See DOI: https://doi.org/10.1039/d3qi01112g

2D metal-organic frameworks (MOFs) are a class of layerstacked materials consisting of well-organized metal centers and organic ligands and have exhibited great potential for application in catalysis and energy storage due to their large surface area, exposed metal active sites and tunable chemical functionality. 13-17 Particularly, 2D electrically conductive MOFs are a newly emerging class of electronic materials that not only inherit most merits from conventional 2D MOFs but also show electrical conductivity due to the extended conjugation. 18-20 Recently, a new 2D conductive MOF, Co/Nibased 2,3,8,9,14,15-hexahydroxyltribenzocyclyne (namely Co/ NiO₄-HTC), was successfully synthesized.²¹ The active centers for the two materials are transition metal-oxygen (TM-O) linkages. Topologically, 2,3,8,9,14,15-hexahydroxyltribenzocyclyne resembles 2,3,6,7,10,11-hexaiminotriphenylene that we have studied in our previous work,²² an archetype ligand with the same 3-fold symmetry, but possesses large surface area due to its bigger size. Therefore, it is worth investigating the electrochemical catalytic activity of this kind of material. Importantly, a systematic theoretical investigation of using this conductive 2D MOF material as a catalyst for electrochemistry is lacking. Such a theoretical investigation is crucial and necessary given the rapid progress of experimental work in this field. As we demonstrated in our previous work based on 2,3,6,7,10,11-hexaiminotriphenylene,²² due to the structural tunability of 2D MOF materials, the catalytic activity of 2D MOF materials can be tuned by substituting the central active transition metal atoms and the organic ligand to adjust the electronic properties. Hence, different TMO₄-HTC materials with TM-O linkage and TMN₄-HTC materials with TM-N linkage as well as materials with different local coordination environments between TM and N/O atoms (TMN_xO_{4-x}-HTC, x = 0-4) were designed.

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In this work, a series of TMN_xO_{4-x} -HTC monolayers were constructed via tuning the TM atoms and the local coordination number between TM atoms and N/O linkages, and their reaction mechanisms and catalytic activity for the OER and ORR were systematically investigated by using density functional theory (DFT) calculations. The results show that all the catalysts could exhibit good thermodynamic and electrochemical stabilities. Notably, CoN_3O_1 -HTC and RhN_2O_2 -HTC are predicted to be promising electrocatalysts for the OER and ORR with the calculated overpotential η^{OER} and η^{ORR} of 0.29 and 0.20 V, respectively. Moreover, RhO_4 -HTC and RhN_2O_2 -HTC are found to be efficient bifunctional catalysts for the OER and ORR.

2. Computational methods

All calculations were carried out by using the Vienna *ab initio* Simulation Package (VASP)^{23,24} based on the spin-polarized density functional theory method. The ion–electron interactions were described by using the projector augmented wave (PAW) method,²⁵ and the electronic exchange–correlation interactions were determined using the Perdew–Burke–

Ernzerhof (PBE)²⁶ functional of the generalized gradient approximation (GGA).²⁷ The van der Waals (vdW) interactions were described by using Grimme's DFT-D3 correction method.²⁸ A plane-wave cutoff energy of 500 eV was adopted for all the computations. The convergence criterion for energy and force during geometrical optimization was set to 10^{-5} eV and 10^{-2} eV Å⁻¹, respectively. The Brillouin zone was sampled using a $3 \times 3 \times 1$ k-point²⁹ during geometry optimization. A vacuum space of 20 Å was applied to avoid the interaction between the periodic images. Throughout all the calculations, we used an implicit solvent model to consider the solvent effect of the water environment through the polarized continuum model as implemented in VASPsol with a dielectric constant of 78.4.30 Ab initio molecular dynamics (AIMD) simulations were performed to demonstrate the thermodynamic stability of the designed catalyst, and the algorithm of the Nose thermostat was carried out to simulate a canonical ensemble³¹ for 10 ps with a time step of 2 fs. Bader charge analysis was adopted to investigate the charge transfer process.32 The calculation details for the OER and ORR are listed in the ESI† as in our previous study. 33,34 The adsorption Gibbs free energy is defined as $G_{ads} = G_{adsorbent+catalyst}$ – G_{catalyst} - $G_{\text{adsorbent}}$, here $G_{\text{adsorbent+catalyst}}$, G_{catalyst} , and $G_{\text{adsorbent}}$ refer to the Gibbs free energies of the adsorbent on the catalyst, the isolated catalyst, and the isolated adsorbent, respectively.

Results and discussion

Fig. 1 shows the stable geometric configurations of 2D TMN_xO_{4-x} -HTC (x = 0-4). The optimized unit cell of TMN_xO_{4-x} -HTC contains three equivalent TM atoms, and each TM atom is surrounded by four nitrogen or oxygen atoms with different ratios. In this work, three 3d transition metals (Fe, Co and Ni), three 4d transition metals (Ru, Rh and Pd), and two 5d transition metals (Ir and Pt) were considered to build catalysts, as these transition metals are commonly used to design OER electrocatalysts.33,34 It should be mentioned that the stability of the designed catalysts is of significant importance for their long-term use. Hence, we calculated the formation energy $(E_{\rm f})$ and dissolution potential $(U_{\rm diss})$ of all the designed catalysts to evaluate their thermodynamic and electrochemical stabilities. The E_f and U_{diss} are defined as $E_f = (E_{TMN_*O_{d-x}} - E_f)$ $E_{N_xO_{4-x}} - 3E_{TM}$)/3 and $U_{diss} = U_{diss}$ (bulk) - E_f /ne, respectively, where $E_{\text{TMN}_{r}O_{4-r}}$ and $E_{\text{N}_{r}O_{4-r}}$ are the total energies of the TMN_xO_{4-x} system and the N_xO_{4-x} substrate, respectively. E_{TM} is the total energy of a metal atom in its most stable bulk structure. U_{diss} (bulk) is the standard dissolution potential of the bulk metal and n is the number of electrons involved during the dissolution process. Since E_{TM} is referenced with respect to that of the bulk metal, systems with negative values of $E_{\rm f}$ are evaluated to be thermodynamically stable against the clustering of TM atoms. Systems with positive values of $U_{\rm diss}$ vs. the standard hydrogen electrode (SHE) are considered to be electrochemically stable. The calculated results of $E_{\rm f}$ and $U_{\rm diss}$ are

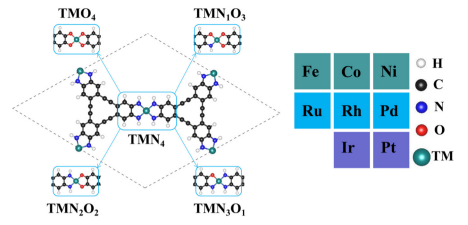


Fig. 1 Optimized geometric configurations of 2D TMN_xO_{4-x}-HTC (x = 0-4) and the considered dopant transition metal atoms.

shown in Fig. 2, which suggest that all the designed catalysts could exhibit good thermodynamic and electrochemical stabilities that meet the stability criteria for electrocatalysts.

Importantly, the distinct electronic properties of all the designed catalysts were studied to obtain insight into their catalytic performance. As presented in Fig. S1-S5,† all the designed catalysts exhibit metallicity, indicating their good electrical conductivity and ensuring efficient electron transfer during the OER and ORR processes. Additionally, in Fig. S6-S10,† the calculated results of the partial density of states (PDOS) suggest that the different dopant TM atoms show different contributions to the electronic states of the designed catalysts around the Fermi level, and the electronic states of the catalysts across the Fermi level are mainly contributed by the d orbitals of the dopant TM atoms. Moreover, around the Fermi energy, the hybridization between the p orbitals of the O/N atoms and the d orbitals of the TM atoms further demonstrates their strong binding interaction. The charge transfer between the substrates and TM atoms could also reveal their strong interaction. Fig. S11† shows a large amount of charge (0.76–1.33e) transfer from the TM atoms to substrates, making the dopant TM atoms positively charged. These positively charged TM atoms are considered to be active sites in the OER and ORR catalytic processes. In previous literature, the d-band

center (ε_d) was used to analyze the interaction strength between catalysts and adsorbates. 37-39 Thus, we calculated the $\varepsilon_{\rm d}$ values of the designed catalysts and plotted them in Fig. S6– S10.† From Fig. S12a,† it can be concluded that the ε_d values shift to a lower energy position than the Fermi level with the increase of the number of d-electrons in the TM atoms at least when the TM atoms are in the same row of the periodic table. Generally, a larger d-electron number of the TM atom and lower energy of ε_d could result in weaker interaction strength between catalysts and adsorbates. 40 For the OER and ORR, the calculated adsorption Gibbs free energies of the HO*, O* and HOO* intermediates ($\Delta G_{\text{HO*}}$, $\Delta G_{\text{O*}}$, and $\Delta G_{\text{HOO*}}$) with the corresponding d-electron numbers of TMN_xO_{4-x}-HTC catalysts are plotted in Fig. S12b-f.† Moreover, ΔG_{HO^*} , ΔG_{O^*} , and ΔG_{HOO^*} with the corresponding ε_{d} values of all TMN_xO_{4-x}-HTC systems are plotted in Fig. S13.† It can be concluded that the adsorption Gibbs free energies of the intermediates decrease with the increase of d-electron numbers of the TM atoms when they are in the same row of the periodic table, which also agrees with the position of ε_d values. Therefore, the adsorption Gibbs free energies of intermediates are negatively correlated with the ε_d values when the TM atoms are in the same row of the periodic table. This phenomenon was also observed in reported experimental and

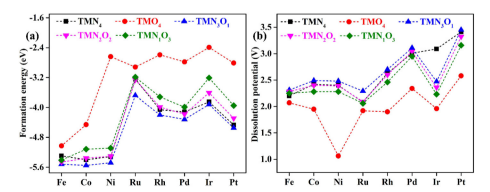


Fig. 2 Calculated (a) formation energy and (b) dissolution potential of transition metal atoms for the designed TMN_xO_{4-x}-HTC catalysts.

literature. 41,42 As a consequence, by tuning the doping of TM atoms on the substrate, the catalyst could exhibit the optimal interaction strength with the intermediates for the OER and ORR.

As proposed by Nørskov et al., 43 the adsorption Gibbs free energies of intermediates govern the intrinsic OER and ORR activity of a catalyst. The calculated corresponding adsorption Gibbs free energy values of intermediates on all the designed TMN_xO_{4-x}-HTC catalysts are listed in Fig. S14-S18.† According to the Sabatier principle, 44 too weak or too strong interaction strength of the intermediates on the catalysts could lead to an adverse effect on the catalytic activity. Therefore, one of our goals is to identify efficient OER/ORR catalysts with moderate intermediate interaction strength. For an ideal OER/ORR catalyst under conditions where the applied potential U equals zero, the adsorption Gibbs free energy value difference between two adjacent intermediates for all the four-electron transfer steps should be 1.23 eV. In other words, for an ideal catalyst, the adsorption Gibbs free energy values of HO*, O* and HOO* intermediates should be 1.23, 2.46 and 3.69 eV, respectively, which makes the four-electron OER/ORR occur at the thermodynamic limit and the overpotential η is zero. However, the reality is that the adsorption Gibbs free energy value difference between two adjacent intermediates is not equal. The OER overpotential (η^{OER}) is determined using the maximum adsorption Gibbs free energy difference of two adjacent intermediates, while the ORR overpotential (η^{ORR}) is determined using the minimum adsorption Gibbs free energy difference of two adjacent intermediates. The calculated Gibbs free energy diagrams for the OER and ORR on all the designed electrocatalysts are shown in Fig. S14-S18,† and the potentialdetermining step (PDS) is colored in yellow for the OER and in pink for the ORR. Moreover, the calculated η^{OER} and η^{ORR} values on all the designed catalysts are summarized in Fig. 3. Notably, among all the designed TMN_xO_{4-x} -HTC catalysts, CoN₃O₁-HTC is predicted to be the best OER catalyst with a calculated η^{OER} value of 0.29 V, followed by RhN₃O₁-HTC (η^{OER} = 0.32 V), CoN_2O_2 -HTC ($\eta^{OER} = 0.33$ V), RhO_4 -HTC ($\eta^{OER} = 0.33$ V), CoN_4 -HTC ($\eta^{OER} = 0.35 \text{ V}$), CoO_4 -HTC ($\eta^{OER} = 0.38 \text{ V}$), $\text{CoN}_1\text{O}_3\text{-HTC}$ ($\eta^{\text{OER}} = 0.39 \text{ V}$), and $\text{RhN}_2\text{O}_2\text{-HTC}$ ($\eta^{\text{OER}} = 0.43 \text{ V}$). Importantly, the OER overpotential values of all the abovementioned electrocatalysts are lower than that of the IrO₂ (110) catalyst ($\eta^{OER} = 0.52 \text{ V}$), 45 indicating their efficient OER catalytic activity. Additionally, the formation of HOO* from O* is the potential-determining step for all the above-mentioned designed efficient catalysts. As known, the ORR is the reverse reaction of the OER. The calculated Gibbs free energy diagrams of all the designed TMN_xO_{4-x} -HTC catalysts toward the ORR are also displayed in Fig. S14-S18.† From Fig. 3b, it can be seen that RhN₂O₂-HTC is predicted to be the best ORR electrocatalyst with a calculated η^{ORR} value of 0.20 V, followed by RhN_1O_3 -HTC ($\eta^{ORR} = 0.29 \text{ V}$), IrO_4 -HTC ($\eta^{ORR} = 0.35 \text{ V}$), FeN_1O_3 -HTC ($\eta^{ORR} = 0.37 \text{ V}$), RhO_4 -HTC ($\eta^{ORR} = 0.39 \text{ V}$), $\text{FeN}_2\text{O}_2\text{-HTC }(\eta^{\text{ORR}} = 0.44 \text{ V}), \text{ CoN}_1\text{O}_3\text{-HTC }(\eta^{\text{ORR}} = 0.49 \text{ V}), \text{ and}$ IrN_2O_2 -HTC ($\eta^{ORR} = 0.50 \text{ V}$), suggesting their efficient ORR catalytic activity. Remarkably, their ORR overpotentials are lower than or comparable to that of Pt (111) ($\eta^{ORR} = 0.48 \text{ V}$). In addition, the formation of HO* is the potential-determining step for RhN₂O₂-HTC, RhN₁O₃-HTC, IrO₄-HTC, RhO₄-HTC, FeN2O2-HTC and IrN2O2-HTC catalysts, and the formation of HOO* is the potential-determining step for FeN1O3-HTC and CoN₁O₃-HTC catalysts. Hence, based on the above results, it

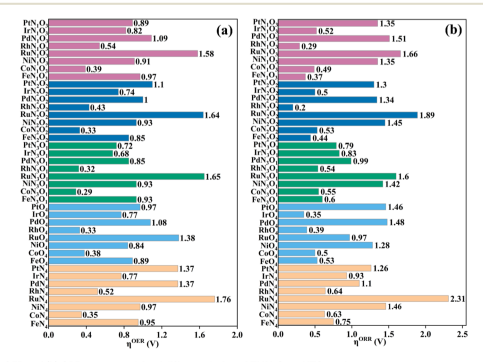


Fig. 3 Calculated (a) OER and (b) ORR overpotentials of different designed TMN_xO_{4-x}-HTC catalysts.

can be noted that RhO₄-HTC, CoN₃O₁-HTC, RhN₂O₂-HTC and CoN₁O₃-HTC are predicted as promising and efficient bifunctional electrocatalysts toward both the OER and ORR, and their free energy diagrams for the OER and ORR are shown in Fig. 4. Moreover, for comparison, previous results on the OER and ORR catalytic activity of 2D-MOF materials are listed in Table S2.†

Research Article

In-depth understanding of the OER and ORR catalytic performance of different catalysts can guide us to design efficient electrocatalysts. As mentioned above, the catalytic performance for the OER and ORR is determined using the Gibbs free energy of the corresponding intermediates on the catalyst. Hence, identifying the relationship between the Gibbs free energy of the intermediates and catalytic activity is essential for the rational design of efficient catalysts. In this work, by comparing the adsorption Gibbs free energy values of the HO* and HOO* intermediates on all the designed catalysts, we found that ΔG_{HOO^*} can be expressed as a function of ΔG_{HO^*} via the equation $\Delta G_{\text{HOO}^*} = 0.92 \Delta G_{\text{HO}^*} + 3.00 \text{ eV (Fig. 5a)}$. It is suggested that the calculated adsorption Gibbs free energy values of the HO* and HOO* intermediates show a strong linear relationship mainly due to both intermediates forming single bonds between O and TM atoms (Fig. 4), and the difference between the Gibbs free energy values of the HO* and HOO* intermediates is a constant. The above results are consistent with those of previously reported carbon-based catalysts for the OER and ORR. 47,48 Given the fact that most of the OER potential-determining step occurs at the HO* to O* or O* to

HOO* step, the OER overpotential could be determined using the difference of ΔG_{O^*} – ΔG_{HO^*} . This is confirmed by the volcano plot displayed in Fig. 5b, where the overpotential values of the OER fall in a line as a function of $\Delta G_{O^*} - \Delta G_{HO^*}$. Obviously, the designed CoN₃O₁-HTC, RhN₃O₁-HTC, CoN₂O₂-HTC and RhO₄-HTC catalysts are located around the peak of the volcano curve with low OER overpotentials and stand out to be promising OER electrocatalysts. For the ORR, the potential-determining step approximately occurs at the * to HO* or HOO* to *+O2 step, and then, the ORR overpotential could be determined using the ΔG_{HO^*} value. Indeed, Fig. 5c shows the volcano plot of the ORR overpotential as a function of ΔG_{HO^*} . Apparently, RhN₂O₂-HTC, RhN₁O₃-HTC, IrO₄-HTC and FeN₁O₃-HTC catalysts with low ORR overpotentials are located around the peak of the volcano plot. Moreover, it can be concluded that the moderate interaction strength of the intermediates on the catalyst could enable good catalytic activity for the OER and ORR. What is more, RhO₄-HTC, CoN₃O₁-HTC, RhN2O2-HTC and CoN1O3-HTC catalysts are located around the top of both volcano plots and screened out to be promising and efficient bifunctional electrocatalysts for both the OER and ORR.

It is noteworthy that the four-electron ORR pathway from O₂ to H₂O is particularly important in metal-air batteries, in which hydrogen peroxide (H2O2) is an undesirable product since it could cause the degradation of the catalyst. 49 For the above screened potential ORR electrocatalysts with the calculated overpotential lower than 0.55 V (RhN₂O₂-HTC, RhN₁O₃-

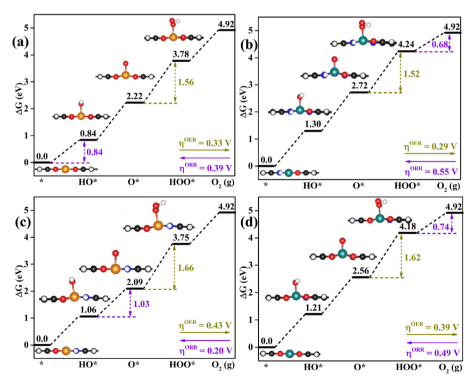


Fig. 4 Calculated free energy diagrams of the promising bifunctional catalysts for the OER and ORR: (a) RhO_4 -HTC, (b) CoN_3O_1 -HTC, (c) RhN_2O_2 -HTC and (d) CoN₁O₃-HTC. The yellow and pink values are the potential-determining step values for the OER and ORR. The optimized configurations of intermediates on the catalysts are also exhibited.

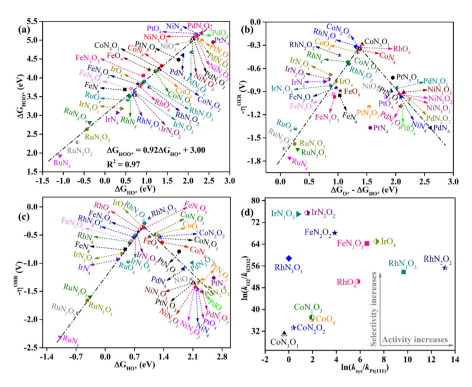


Fig. 5 (a) Scaling relationship between ΔG_{HO^*} and ΔG_{HOO^*} on all the designed TMN_xO_{4-x}-HTC catalysts. (b) Calculated OER volcano curve of $-\eta^{\text{OER}}$ as a function of $\Delta G_{O^*} - \Delta G_{HO^*}$ on all the designed catalysts. (c) Calculated ORR volcano curve of $-\eta^{ORR}$ as a function of ΔG_{HO^*} on all the designed catalysts. (d) Variations of the ORR activity vs. selectivity of the potential electrocatalysts.

HTC, IrO₄-HTC, FeN₁O₃-HTC, RhO₄-HTC, FeN₂O₂-HTC, IrN₂O₂-HTC, CoO₄-HTC, CoN₁O₃-HTC, IrN₁O₃-HTC, RhN₃O₁-HTC, CoN2O2-HTC and CoN3O1-HTC), the selectivity for the four-electron pathway is verified from the thermodynamic perspective, since the calculated ΔG_{O^*} values (2.09, 2.13, 1.84, 1.86, 2.22, 1.76, 1.57, 2.58, 2.56, 1.58, 2.00, 2.66 and 2.72) are all smaller than 3.52 eV $(\Delta G_{\rm H_2O_2} - \Delta G_{\rm H_2O})^{.50,51}$ A kinetics investigation was carried out to understand the catalytic selectivity for the ORR50 and the corresponding calculation details are listed in the ESI.† As shown in Fig. 5d, all the calculated ln $(k_{\rm O_2}/k_{\rm H,O_2})$ values are positive, suggesting that the reduction of O₂ to H₂O is prioritized on these designed catalysts. Especially, the calculated $ln(k_{sys}/k_{Pt(111)})$ values of RhN₂O₂-HTC, RhN₁O₃-HTC, IrO₄-HTC, FeN₁O₃-HTC, RhO₄-HTC and FeN₂O₂-HTC are 13.15, 9.67, 7.35, 6.57, 5.80 and 3.87, respectively, indicating that the reaction rate on these designed catalysts is much faster than that on Pt(111).

To explicitly visualize how the OER and ORR catalytic activity of the screened CoN3O1-HTC, RhN3O1-HTC, RhO4-HTC, CoO₄-HTC, RhN₂O₂-HTC, CoN₂O₂-HTC, RhN₁O₃-HTC and CoN₁O₃-HTC catalysts is in practice, the corresponding theoretical OER and ORR polarization curves were simulated based on the reversible hydrogen electrode (RHE) in comparison with those of IrO₂(110) for the OER and Pt(111) for the ORR, that is, the change in current density as a function of potential *U*. The simulation details are listed in the ESI.† As shown in Fig. 6, for the OER, at a current density of 10 mA cm⁻², the simulated polarization curves of CoN₃O₁-HTC,

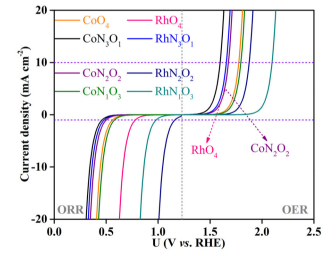


Fig. 6 Simulated polarization curves for the screened-out catalysts.

RhN₃O₁-HTC, RhO₄-HTC, CoN₂O₂-HTC, CoO₄-HTC, CoN₁O₃-HTC, RhN2O2-HTC and RhN1O3-HTC exhibit lower onset potentials of 1.59, 1.65, 1.67, 1.67, 1.78, 1.80, 1.88 and 2.09 V vs. RHE than that of IrO₂ (2.27 V) as reported in our previous work, respectively;²² for the ORR, at a current density of 1 mA cm⁻², the simulated polarization curves of RhN₂O₂-HTC, RhN₁O₃-HTC and RhO₄-HTC exhibit higher onset potentials of 1.16, 0.98 and 0.78, vs. RHE than that of Pt(111) (0.67 V) as

reported in our previous work, respectively,22 while the simulated polarization curves of CoN₁O₃-HTC, CoO₄-HTC, CoN₂O₂-HTC, RhN₃O₁-HTC and CoN₃O₁-HTC exhibit lower onset potentials of 0.58, 0.55, 0.51, 0.48 and 0.46 V, respectively. The above results indicate that these screened-out catalysts possess efficient OER and ORR catalytic activity, which makes them potential alternatives to IrO2 and Pt electrodes. Additionally, we performed AIMD simulations for the potential catalysts (taking CoN₃O₁-HTC, RhN₃O₁-HTC, CoN₂O₂-HTC, RhN₂O₂-HTC, CoN₁O₃-HTC, RhN₁O₃-HTC, CoO₄-HTC and RhO₄-HTC as examples) to evaluate their dynamic stabilities. The simulated results (Fig. S19-S22†) show that the energies oscillate near the equilibrium state during the 10 ps simulations and the structures have no obvious structural reconstruction, which suggests their good kinetic stability.

Conclusions 4.

In summary, a series of 2D TMN_xO_{4-x}-HTC (x = 0-4) electrocatalysts were designed and systematically investigated for their catalytic activity toward the OER and ORR based on DFT calculations. The strong interaction between TM atoms and N_xO_{4-x} HTC could guarantee the stability of TMN_xO_{4-x} -HTC. It was found that as the number of d-electrons increases, the d-band value decreases, thereby weakening the interaction between the intermediates and TM atoms. The OER overpotential η^{OER} follows a volcano plot of ΔG_{O^*} – ΔG_{HO^*} , and the ORR overpotential η^{ORR} follows a volcano plot of ΔG_{HO^*} . Among all the designed TMN_xO_{4-x}-HTC electrocatalysts, the best OER catalyst is CoN_3O_1 -HTC with a calculated η^{OER} value of 0.29 V, followed by RhN₃O₁-HTC (0.32 V), CoN₂O₂-HTC (0.33 V) and RhO₄-HTC (0.33 V); the best ORR catalyst is RhN2O2-HTC with a calculated η^{ORR} value of 0.20 V, followed by RhN₁O₃-HTC (0.29 V), IrO₄-HTC (0.35 V), FeN₁O₃-HTC (0.37 V) and RhO₄-HTC (0.39 V). Notably, RhO₄-HTC, RhN₂O₂-HTC and CoN₁O₃-HTC are predicted as promising and efficient bifunctional electrocatalysts for both the OER and ORR. Moreover, by analyzing the relationship between the ORR products H2O and H2O2, the thermodynamically favorable selectivity for H2O is elucidated from the kinetics perspective. Our results shed light on the exploration of 2D-MOF materials as promising OER and ORR electrocatalysts.

Conflicts of interest

The authors declare no competing financial interests.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (22102167 and U21A20317). The calculations were performed at the Supercomputing Center of the University of Science and Technology of China.

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