2D transition metal–TCNQ sheets as bifunctional single-atom catalysts for oxygen reduction and evolution reaction (ORR/OER)

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Two-dimensional Transition Metal–Tetracyanoquinodimethane (TM–TCNQ) Sheets as High-Activity Bifunctional Single-Atom Catalysts toward ORR and OER

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ABSTRACT

Searching for stable, low-cost and active electrocatalysts for the oxygen reduction reaction (ORR) and oxygen evolution reaction (OER) to reduce use of precious platinum family is of great significance for large-scale commercialization of fuel cells and electrolyzers, but still remains a big challenge. Based on density functional theory (DFT), we systematically investigate two-dimensional transition metal based tetracyanoquinodimethane (referred to as TM–TCNQ, TM = Cr – Zn, Ru, Ag, Pt, Ir) monolayers, whose metal centers are distributed orderly and separately. After considering stabilities, conductivities and reaction mechanisms in acid solutions, the synthesized Fe–TCNQ monolayer is found to exhibit excellent catalytic activity for ORR with a very low overpotential of 0.35 V via four-electron reaction, which is lower than that of Pt metal of 0.48 V. The catalytic performances of Ir–, Co–TCNQs for OER are identified as the highest two among the 12 candidates. By applying grafting axial ligands and external strain, the adsorption strength of reaction intermediates on TM reactive sites can further be tuned and resulting in enhanced activities of Fe–, Co– and Mn–TCNQs for the ORR/OER. Most importantly, Fe–TCNQ–Cl (η=0.32/0.34 V), then following Mn–TCNQ–Cl (η=0.41/0.44 V) and Mn–TCNQ–CO (η=0.33/0.54 V) are predicted as high-activity bifunctional catalysts for the ORR/OER reactions with modified scaling relations of adsorbates, whose catalytic activities are even higher than the two best commercial electrocatalysts Pt (ORR:η=0.48V) and RuO₂ (OER:η=0.41V). In addition, as novel synthesized single-atom catalysts, Fe–TCNQ and Mn–TCNQ possess excellent conductivity with fast electron transfer during the catalytic processes. These results might revive such 2D monolayer materials to design durable, low-cost, and efficient catalysts of ORR/OER in metal-air batteries and fuel cells.

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1. Introduction

The rising energy demand and depletion of fossil fuels have led to increasing interest in alternative energy sources. Additionally, fossil fuels are not renewable and generate deleterious combustion products such as CO, CO₂, NO, SO₂, and even hazards of heavy metal contamination posing serious challenges to human health, and environmental protection¹. Sustainable power generation technologies are regarded as a promising solution, and a good example of such technologies is the proton exchange membrane (PEM) fuel cell,² ³ where hydrogen is oxidized at the anode producing protons which are transported through the membrane to the cathode where the oxygen reduction reaction (ORR) takes place during the production of water. However, slow oxygen kinetics of cathode and the use of precious metals based electrocatalysts, such as platinum (Pt), hinder widespread commercialization⁴. Likewise, the oxygen evolution reaction (OER) in water splitting electrocatalysts also requires noble metals namely ruthenium or iridium-based catalysts⁵–⁷. Thus, scientists are continuously seeking new families of low-cost electrocatalysts whose catalytic performances are comparable or even higher than those of noble metals based materials⁶. Moreover, finding bifunctional ORR and OER electrocatalyst is a key step in rechargeable metal-air batteries and regenerative fuel cell technologies⁸. Any technical breakthrough would therefore be greatly beneficial for easing the contradiction between supply and demand of energy and solving the environmental problems with

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burning fossil fuels. Metal-free electrodes have recently attracted
attention as alternative to rare metal containing catalysts that can
be used to catalyse the ORR or OER effectively\textsuperscript{10}. For instance,
vertical aligned nitrogen-doped carbon nanotube (VA-NCNT)
arrays can catalyse a 4e\textsuperscript{−} ORR process with a 3-times higher
electrocatalytic activity and better long-term operational stability
than that of widely used platinum/C electrodes\textsuperscript{11}. Another way to
reduce the usage of precious metals is mixing or alloying with
cheaper metals, as the adsorption of reaction intermediates can be
tuned to enhance the ORR or OER activity. The third approach
involves the replacement of noble metals with much cheaper 3d
transition-metal catalysts supported on one or two dimensional
materials. Typical examples are the transition metal-N\textsubscript{4}
macrocycles (TM-N\textsubscript{4}), e.g. phthalocyanine (TM-Pc)\textsuperscript{12-17}, TM-
porphyrin-like complexes (TM-PP)\textsuperscript{18,20} and TM coordinate
hexaaminobenzene-base polymer (TM-HAP)\textsuperscript{21, 22}, whose
catalytic performances have been extensively studied both
experimentally and theoretically. One big advantage of two
dimensional single-atom catalysts (SACs) is that the various kinds
of single metal atom sites are ordered and strongly anchored into
the pores offering great potential for achieving high-activity
catalysts with maximal usage of the transition metals. Besides,
most of such two dimensional metal-organic frameworks are
metallic allowing fast electron transport, which is particularly
important for high activity electrocatalysts.

Analogous to the TM-N\textsubscript{4} complexes discussed above, two-
dimensional transition metal–tetracyanoquinodimethane (TM–
TCNQ) with four-coordinated metal structure have been
synthesized with unique electronic, magnetic, and photoactive
properties\textsuperscript{23-27}. Wang \textit{et al.} predicted Fe-TCNQ can serve as an
efficient electrocatalyst for the ORR in acidic and alkaline
conditions by DFT\textsuperscript{28}. The activity range of the TM-TCNQs
remains, however, largely unexplored, and their capacity as
bifunctional ORR/OER electrocatalyst is still unknown. Inspired
by the realized structures and the present simulations, we therefore
systematically investigate twelve TM-TCNQs\textsuperscript{1} stabilities,
structures, and catalytic behaviors for the ORR and the OER by
using first-principle calculations (TM=Cr, Mn, Fe, Co, Ni, Cu, Ru,
Rh, Ir, Pt, Pd, Ag). Our results reveal that Fe-TCNQ possesses the
highest catalytic activity for the ORR with a rather low
thermodynamic overpotential (\(\eta\)) of 0.35 V via a four-electron
reduction; this overpotential is noticeably lower than on the widely
used Pt metal\textsuperscript{20, 29, 30}. The catalytic performances of Ir-, Co-TCNQs
for the OER are identified as the highest among 12 candidates.
Grafting axial ligands and applying external strain can further
enhance the activity, especially towards the OER, attributed to
modify scaling relations of intermediates. Most importantly, Fe-
TCNQ and Mn-TCNQ with ligands of Cl and Mn-TCNQ with CO
are predicted as bifunctional ORR/OER catalysts whose catalytic
activities are comparable or even higher than the two best
electrocatalysts Pt\textsuperscript{20, 29, 30} and RuO\textsubscript{2}\textsuperscript{31}.

2. Computational Methods
All calculations in this study are carried out using spin-polarized
density function theory (DFT) calculations as implemented in the
Vienna \textit{ab initio} simulation package (VASP)\textsuperscript{7- 22} to explore the
activities for the ORR and the OER on TM-TCNQ monolayers.
The exchange-correlation interactions are approximated by the
RPBE functional\textsuperscript{32} within the generalized gradient approximation
(GGA)\textsuperscript{33}, which provide better chemisorption energies than PBE.
The kinetic energy cutoff of the plane-wave basis set is 500 eV,
and dipole corrections are applied in the direction perpendicular to
the TM-TCNQ plane. The thickness of the vacuum layer between
TM-TCNQ planes is set to be more than 20 Å, which is sufficiently
large to avoid interlayer interactions. The electronic SCF tolerance
is set to 10\textsuperscript{-5} eV. Fully relaxed geometries and lattice constant are
obtained by optimizing all atomic positions until the Hellmann–
Feynman forces are below 0.02 eV/Å. The k-point sampling is
4x4x1 in the first Brillouin zone for structural optimizations, while
a 8x8x1 sampling is used for the density of states (DOS)
calculations. VASP input files are prepared by the atomistic
simulation environment (ASE)\textsuperscript{34}. The structures of transition state
(TS) are identified by the climbing image nudged elastic band
method (CI-NEB)\textsuperscript{35}. At least eight intermediate images are
constructed along the reaction pathway between the initial state (IS)
and the final state (FS) with a spring force of 5.0 eV/Å. The
minimum energy path is optimized with the fast inertial relaxation
engine (FIRE)\textsuperscript{36} until the maximum force on the band is below
0.05 eV/Å.

The DFT formation energies of OOH*, O*, and OH* are
 calculated with reference to H\textsubscript{2}O(g) and H\textsubscript{2}(g) according to the
following equations (1)-(3):

\[
\Delta E_{\text{ads}}(\text{OOH}* ) = E_{\text{TCNQ-OOH}} - E_{\text{TCNQ}} - (2E_{\text{H2O}} - 3/2E_{\text{H2}}) \tag{1}
\]

\[
\Delta E_{\text{ads}}(\text{O}*) = E_{\text{TCNQ-O}} - E_{\text{TCNQ}} - (E_{\text{H2O}} - E_{\text{H2}}) \tag{2}
\]

\[
\Delta E_{\text{ads}}(\text{OH}*) = E_{\text{TCNQ-OH}} - E_{\text{TCNQ}} - (E_{\text{H2O}} - 1/2E_{\text{H2}}) \tag{3}
\]

where \(E_{\text{TCNQ-OOH}}, E_{\text{TCNQ-O}}, E_{\text{TCNQ-OH}}\) are the energies of adsorbate
binding to the transition metal of TCNQ complexes, \(E_{\text{H2O}}\) and \(E_{\text{H2}}\)
are the total energy of single isolated H\textsubscript{2}O and H\textsubscript{2} molecule,
respectively.

The computational hydrogen electrode (CHE) model\textsuperscript{29} is
applied to evaluate the change in Gibbs free energy of adsorbates
by equation (4).
where $E_{\text{ad}}$ is calculated from equation (1) to (3). $\Delta E_{\text{PE}}$ and $T\Delta S$ are the zero point energy change and entropy change upon adsorption, respectively. Here, $\Delta E_{\text{PE}} + T\Delta S$ is approximated to be 0.4, 0.05 and 0.35 eV for OOH, O, and OH adsorbates, respectively.\(^{29,37,38}\) $\Delta G_n = n e U$, where $n$ is the number of electrons transferred and $U$ is the electrode potential versus the reversible hydrogen electrode.\(^{19}\) Solvation energies are added to $\Delta G$ by -0.3 eV for OH and OOH adsorbates followed by previous studies.\(^{37}\)

3. Results and Discussion

Irreversible detachment of the central transition metal from the catalyst significantly limits catalyst durability. To elucidate the stabilities of TM-TCNQ monolayers, we firstly examine the energetic difference ($E_{\text{diff}}$) between the embedding energy of the metals ($E_{\text{emb}}$) and the cohesive energy ($E_c$) of the bulk metal according to the equation:

$$E_{\text{diff}} = E_{\text{emb}} - E_c = (E_{\text{TM-TCNQ}} - E_{\text{TCNQ}} - E_{\text{M}}) - (E_{\text{emb}} - NE_M)/N$$

where $E_{\text{TM-TCNQ}}$, $E_{\text{TCNQ}}$, $E_{\text{emb}}$, $E_M$ are total energies of the TM-TCNQ system, the metal-free TCNQ, the bulk metal unit cell containing $N$ metal atoms and the isolated single metal in the same slab, respectively. Negative values of $E_{\text{diff}}$, here in the range from -0.81 to -2.89 eV (Table.S1), indicate the metal atoms prefer to be embedded into the TCNQ rather than forming bulk metal or clusters likely because of the very strong coordination bonds formed as nitrogen atoms donate 2p electron pairs to the vacant d orbitals of metal. Thus, Cr and Mn-TCNQ are expected to have the largest $E_{\text{diff}}$ due to the relative high fraction of vacant d orbitals of metal. Secondly, the removal energy ($E_{\text{rev}}$) is calculated to evaluate the feasibility of metal substitution by hydrogenation of TCNQ in solution by the following equation:

$$E_{\text{rev}} = E_{\text{TM-TCNQ}} + E_{\text{H2-TCNQ}} - E_{\text{bulk}}/N$$

where $E_{\text{TM-TCNQ}}$, $E_{\text{H2-TCNQ}}$, $E_{\text{bulk}}$ are total energies of the TM-TCNQ system, hydrogenated TCNQ, $H_2$ molecule and basic unit cell containing $N$ metal atoms, respectively. Large negative values of $E_{\text{rev}}$ for Cr-, Mn-, Fe-TCNQ ($\approx$-0.28eV) reveal that these metal atoms can coordinate strongly with four nitrogen atoms in a square pyramidal configuration rather than being substituted by $H_2$ (or protons and electrons) in Table.S1. However, the central metal atoms are more easily substituted by protons in Ru-, Pd-, Ag-, and Pt-TCNQs because of their larger positive values of $E_{\text{rev}}$ ($\approx$0.28eV). Hydrogen substitution of the central metal atom in Co-, Rh-, Ni-, Cu-, and Ir-TCNQs are moderate due to small removal energies from -0.11 to 0.16 eV. Thirdly, in acidic electrochemical electrolyte transition metals might oxidize and the most stable form of the transition metal can be a cation rather than the bulk metal. To access the stability in acid of these corroding transition metals, we also calculate the acidic stability as

$$E_{\text{acid}} = E_{\text{diff}} + ne(U - U_0)$$

for the corroding elements. Here $n$ is the number of electrons transferred in the oxidation, $e$ is the elementary charge, $U$ is the catalyst potential, and $U_0$ is the experimental reversible potential for the metal redox reaction.\(^{19}\) We see (in Table.S1, Supporting Information) that early 3d transition metals in TM-TCNQs are thermodynamically unstable at a representable operating potential of 0.75 V vs RHE, and that Fe and Co are less unstable than Cr and Mn. In brief, by applying three descriptors to evaluate the stability of the TM-TCNQ monolayers, the stability of these candidates towards metal atoms clustering follow the trend Cr > Mn > Fe > Co > Rh > Cu > Ni > Ir for TCNQ monolayers in aqueous solution. At oxidizing operating potentials, the trend is largely reversed Ir > Rh > Cu > Co > Fe > Ni > Mn > Cr. While, the unlisted candidates should be ruled out due to unsatisfying the stability criterion.

The optimized structure of Fe-TCNQ is shown in Fig. 1, as an example. The TM-TCNQs have rectangular unit cells with lattice constants listed in Table S1. Similar to graphene and boron nitride sheets, the structures of all the catalysts are all perfect 2D atomic sheets without buckling. The bond length between Fe and the nearest neighboring N atom is 1.853 Å which is 5% shorter than the value of 1.946 Å in FePc. Except Cu, Pd, and Ag, the TM-N bond lengths of TCNQ are all shorter than the TM-Pc bond length as shown in Table S1. Bader charge analysis suggests each highly electronegative nitrogen atom carries $-1.24 \cdot e^\dagger$, and the neighbouring Fe and C atoms carry 1.19 $\cdot e^\dagger$ and 0.77 $\cdot e^\dagger$ positive...
charge, indicating that the bonding between N and Fe/C is largely covalent. In most cases, the spin and oxidation states of TM-TCNQ complexes listed in Table S1 are quite similar to other metal-N₄ coordinate materials, such as TM-Pc²¹, TM-N₄-gra²⁷, and TM-CAP²¹. Interestingly, Ni in the TCNQ monolayer as a “high-spin” complex carries 1.14 μₗ magnetic moment leading to a +2 oxidation state, whereas Ni-CAP is nonmagnetic (based on a previous study²¹ and our calculations). Thus, small differences in atomic geometries of TM-N₄ moieties can have quite different electronic structure leading to great influence on adsorbates’ binding energies and therefore their catalytic activities for the ORR and OER.

As proposed by Nørskov et al.²⁹, the activity of the ORR and the OER is mainly governed by the adsorption energy of key reaction intermediates, because the kinetics of proton coupled electron transfer to the reaction intermediates is fast even at room temperature. According to the Sabatier principle, if intermediates interact too strongly with TM-TCNQ monolayer, the active sites of the catalyst are poisoned; whereas, if the interactions are too weak, it is difficult to activate reactants or intermediates and launch the catalytic process. Therefore, identifying promising materials with “just right” adsorption energies of the reaction intermediates is a challenge. Via the four-electron reaction pathway, the overall formation energy toward ORR is 4.92 eV, so the ideal adsorption free energies of OOH*, O* and OH* are 3.69, 2.46, and 1.23 eV with consecutive intermediates differing by 1.23eV. In Table S2, the adsorption free energies of 12 candidates are reported. The trend in intermediate binding strength is Cr > Ru > Mn > Fe > Co > Ni > Ir > Rh > Cu > Pd > Pt > Ag. It is well known the adsorption energies correlate strongly with d-band centres thirty-two, thirty-nine. Negative d-band centers correspond to low reactivities of the transition metal when only a small proportion of d orbitals lie above the Fermi level. From figure S1 and table S2, we can see that the trend in d-band centers follows the trend in adsorption energies with the increase of the atomic number of the TM atoms. Thus, the performance of the ORR and the OER can be well described by the trend in d-band centres, which may guide better design of electrocatalysts. Besides, the electronic local density of states (LDOS) projected on the TM and the total density of states (DOS) reveal the TM-TCNQs are all found to be metallic at the applied level of theory (GGA-DFT), except Pt-TCNQ whose band gap reaches 0.71eV. Metallic conductivity guarantees fast electron transfer during the electrochemical reactions, a feature consistent with TM-CAPs²¹.

First, we consider a dissociative mechanism where chemisorbed O₂ dissociates into two O atoms on the catalyst surface. From the structures of TM-TCNQs, because the central metal atom is isolated and embedded into the plane of the TM-TCNQ, we expect the catalyst is unable to provide enough reactive sites to accommodate two O atoms and the barrier should be quite high. A previous study²⁸ found that O₂ dissociation is difficult, as it has a large barrier of over 2 eV on Fe-TCNQ. Because stronger binding of adsorbates usually corresponds to a lower barrier of O₂ dissociation, we only need to consider O₂ dissociation on Cr- and Mn-TCNQs in the present study. With high activation energies of 1.14 eV and 1.34 eV for O₂ dissociation calculated by CI-NEB method, the dissociative mechanism is unlikely to occur, and we turn our attention to the associative mechanism in the following.

![Figure 2](image_url) Scaling relations between the adsorption energies of adsorbates on TM-TCNQ monolayers. Black: *O vs. *OH; red: *OOH vs. *OH. The dashed line shows the O₂ initial state at 4.92 eV.

The oxygen reduction reaction and oxygen evolution reaction catalyzed by TM-TCNQ.

We investigate 4-electron ORR in acidic solution via the following pathway:

\[ \Delta G_{\text{ads}} = \Delta G_{\text{ads(OOH*)}} - 4.92 \text{ eV} \]  
\[ \Delta G_s = \Delta G_{\text{ads(O*)}} - \Delta G_{\text{ads(OH*)}} \]  
\[ \Delta G_s = - \Delta G_{\text{ads(OH*)}} \]

Figure 2. Scaling relations between the adsorption energies of adsorbates on TM-TCNQ monolayers. Black: *O vs. *OH; red: *OOH vs. *OH. The dashed line shows the O₂ initial state at 4.92 eV.
electron reductions on Fe TCNQs have higher η (0.78 V, 0.67 V), which are caused by their stronger adsorption of OH*. In contrast to this, transition metals from Co to Ag provide weak binding of the ORR intermediates resulting in low activities toward the ORR with the rate-determining step being O₂ reduction to OOH*. Note that Fe almost reaches the top of the volcano, and weakening the interaction between OH* and the metal in e.g. Mn-TCNQs or Ru-TCNQs will enhance the ORR activity, whereas increasing the interaction between OH* (and OOH*) and Co-TCNQ or Ni-TCNQ will enhance activity as well.

Here, the two-electron (2e⁻) reduction of O₂ to H₂O₂ is also taken into account though formation of OOH* (eq. (21)) followed by

\[
\text{OOH}^* + \text{H}^* + e^- \rightarrow \text{H}_2\text{O}_2
\]  

(21)

where the unhydrogenated O atom in OOH* is hydrogenated to form H₂O₂. Fig. 4 displays the free energy diagram of the ORR via the two-electron and four-electron reductions on Fe-TCNQ at different electrode potentials, respectively. At U=0, all steps are exothermic, so the reaction can spontaneously occur on a Fe-TCNQ monolayer from O₂ to H₂O. However, the formation of

Figure 3. Four-electron-transfer ORR pathway through the intermediates OOH*, O* and OH*, and contour map of the ORR overpotential (η) as a function of Gibbs free energies of O*-OH* and OH*. Oxygen (O) atoms are coloured in red.

Figure 2 shows the scaling relations of \( \Delta G(\text{OH}^*) \) vs \( \Delta G(\text{O}^*) \) and \( \Delta G(\text{OH}^*) \) vs \( \Delta G(\text{OOH}^*) \). They can be expressed as a function of \( \Delta G(\text{O}^*) \) by

\[
\Delta G(O^*) = 1.88 \Delta G(\text{OH}^*) + 0.40 \text{ eV}
\]  

(17)

\[
\Delta G(\text{OOH}^*) = 0.85 \Delta G(\text{OH}^*) + 3.22 \text{ eV}
\]  

(18)

Strong linear relationships of equations (17) and (18) with high coefficients of determination (R²=0.92, 0.98) and very small root mean square error (0.09 eV, 0.005 eV) can be found for OH vs O, and OH vs OOH, respectively. The slope of 0.85 and a constant separation of 3.22 eV of OH vs OOH have been attributed to the similar bond order of M-OH and M-OOH bonds, and is consistent with those of metal and metal oxide surfaces. \( \Delta G_{\text{ads}} \) is calculated from equations (1)-(4). The Gibbs free energy of O₂ (4.92 eV) is taken from the experimental value of the reaction 2H₂O →O₂+2H₂. The first step in the ORR pathway is O₂ being hydrogenated to OOH* through coupled proton electron transfer. Then OOH* is attacked by an additional proton-electron pair to form atomic oxygen and a water molecule. O* is then further hydrogenated to OH*, before the second H₂O formed. We neglect the small barriers of proton coupled electron transfer in this work, in which case the reaction kinetics is only determined by the reaction free energy of each step. The activity of an ORR catalyst can be evaluated by the thermodynamic overpotential (η). Based on the computational hydrogen electrode (CHE) model, the limiting potential \( U_{\text{lim}} \) and the overpotential can be calculated from equations (19) and (20) by determining the greatest free energy increase of all reaction steps at the equilibrium potential 1.23 V of the 4e⁻ ORR:

\[
U_{\text{lim}} = \max \{\Delta G_1, \Delta G_2, \Delta G_3, \Delta G_4\}/4e
\]  

(19)

\[
\eta = 1.23 - U_{\text{lim}} = 1.23 + \max \{\Delta G_1, \Delta G_2, \Delta G_3, \Delta G_4\}/e
\]  

(20)

Lower overpotential represents a better ORR performance. The overpotential is evaluated on the TM-TCNQs as a function of the two variables \( \Delta G(O^*\text{-OH}^*) \) and \( \Delta G(OH^*) \) and is shown Fig. 3. Here, \( \Delta G_{\text{ads}}(\text{OOH}^*) \) is approximated as a function of \( \Delta G_{\text{ads}}(\text{OH}^*) \) from the scaling relation between OOH* and OH* in Fig. 2. From the volcano plot in Fig. 3, the minimum possible η is found as low as 0.3 V. The best TM-TCNQ for the ORR is found to be Fe-TCNQ with η=0.35 V. This thermodynamic overpotential is even lower than that on Pt(111) (η=0.48 V) calculated with the RPBE functional. The rate-determining step of the ORR on the Fe-TCNQ monolayer is the forth step of OH* reduction to H₂O* which agrees with other macrocyclic Fe-Ni materials. Mn also exhibits very good activity (η=0.51V) for ORR with the same rate-determining step as Fe. Cr and Ru-TCNQs have higher η (0.78 V, 0.67 V), which are caused by their stronger adsorption of OH*. In contrast to this, transition metals from Co to Ag provide weak binding of the ORR intermediates resulting in low activities toward the ORR with the rate-determining step being O₂ reduction to OOH*. Note that Fe almost reaches the top of the volcano, and weakening the interaction between OH* and the metal in e.g. Mn-TCNQs or Ru-TCNQs will enhance the ORR activity, whereas increasing the interaction between OH* (and OOH*) and Co-TCNQ or Ni-TCNQ will enhance activity as well.
H₂O₂ via the two-electron pathway is less exergonic than formation of O₂ in the 4e⁻ reduction, because the released free energy (0.49 eV) by H₂O₂ is much less than the free energy (1.88 eV) released in the process of converting OOH to O. Therefore, when U >0.49V, the H₂O₂ formation from OOH is endothermic, while O²⁻ formation from OOH* is still downhill. When U=Uₘₐₓ=0.88 V, the free energy of OH* to H₂O become isoenergetic, and H₂O₂ formation is supressed. At the equilibrium potential for the 4e⁻ reduction, the thermodynamic barrier is highest for the rate-determining step, which is the reduction of OH* to H₂O.

The OER is the reverse reaction of the ORR. Thus, the overpotential can be rewritten as the following equation (22):

\[ \eta = U_{\text{lim}} \cdot 1.23 \text{ V} = \max \{ - \Delta G_i, - \Delta G_2, - \Delta G_3, - \Delta G_4 \}/e - 1.23 \text{ V} \]  

Fig. 5 indicates the ideal minimum \( \eta \) is 0.34 V by using the scaling relation between OOH* vs OH* from Fig 2. The TM-TCNQs are distributed in three different zones corresponding to different rate-determining steps, and most are quite far from the apex of the activity volcano. The two best catalysts for OER among the 12 candidates are identified as Co- and Ir-TCNQ with \( \eta \) of 0.58 and 0.57 V respectively, which is about 0.15 V higher than RuO₂ (\( \eta=0.42 \text{ eV} \)). Ni and Fe also exhibit reasonable activity with \( \eta \) of ca. 0.65 V, which agrees quantitatively with recently identified OER overpotential of TM-Ni materials (TM=Fe, Co, Ni). According to the above discussion, there is still plenty of room to enhance the catalytic activity as none of the TM-TCNQs approaches the optimal \( \eta \) for OER. From above stability selection, Ru atoms are found easily substituted by protons in aqueous solution. Considering this and cost, we will not study Ru-TCNQ monolayer in the following part.

**Figure 5.** Four-electron-transfer OER pathway through the intermediates O⁴⁺, O²⁻ and OOH⁻, and contour map of the OER overpotential (\( \eta \)) as a function of Gibbs free energies of O²⁻OH⁻ and OH⁻.

**Figure 6.** Overpotentials of TM-TCNQ (TM=Mn, Fe, Co) towards the ORR and the OER by coordinating Cl, CO, NO, CN as axial ligands as well as ligand free monolayers for comparison, respectively.

**Improving the catalytic performance by coordinating axial ligands**

Coordinating an axial ligand to the metal centre is expected to be an effective approach⁴²,⁴³ to improve the performances of ORR and OER, which attribute to change of the properties of the metal centres and their interaction with the coordination configuration. For instance, improved OER activity for Fe₇N₄C₄ in phosphoric acid electrolyte, was recently attributed to coordination to phosphate anions⁴⁴. Moreover, ligand-field effects altering the chemical environment of the central TM sites may also lead to modified scaling relations between the reaction intermediates. In experiments, the five and six fold coordination of TM-4N complexes are common approaches to modificate their properties.⁴⁵ Therefore, inspired by this technology and a synthesised six fold coordination in Fe-TCNQ materials, we study Mn-, Fe- and Co-TCNQ sheets near the vertex of volcano with four different ligands viz. cyano (CN), carbon monoxide (CO), nitrogen oxide (NO), chloride (Cl).

The ligands are initially placed on one side of the monolayer, and the intermediates adsorbed on the other side. The ORR and OER overpotentials of Mn, Fe and Co-TCNQs with these ligands are shown in Fig. 6. Among these candidates, Fe-TCNQ-CI exhibit very good catalytic performance (\( \eta=0.32/0.34 \text{ V} \)) for bifunctional ORR/OER activity, then following Mn-TCNQ-CI with \( \eta \) less than 0.5 V, whereas Mn-TCNQ-CO is also found to be an efficient bifuncational catalysis. The superb catalytic performance is ascribed to suitably weakened adsorption energies and violating the original scaling relations of adsorbrates after grafting the appropriate ligands onto the metal centre (Fig. S2, Supporting Information). The NO ligand, however, decreases the activity toward the ORR and the OER in Fig.6. The reason is that \( \Delta G(OH^+) \) increases too much (> 0.3 eV) compared to ligand-free case.

From the discussion above, the OER overpotential can become lower with increasing \( \Delta G(OH^+) \) for Mn- and Fe-TCNQ. The CN ligand plays such role by weakening \( \Delta G(OH^+) \) by 0.15-0.35 eV.
Figure 7. Gibbs free energy diagram of bifunctional Fe-TCNQ with an axial Cl grafting ligand for the ORR and the OER at the 4e\textsuperscript{-} equilibrium potential. The overpotentials (\(\eta\)) in Volts are marked in red and blue.

and resulting in lower \(\eta\) for the OER. However, taking into account the competition between ligands and intermediates, we find a second CN ligand binds strongly to TCNQ with adsorbate formation energies in the in range from -0.71 to -1.14 eV in Fig. S3, so CN may block the active sites by forming 6-fold coordinated metal sites. Meanwhile, the binding strengths of the second ligands are relative weaker for CO (-0.54—0.02 eV) and Cl (-0.74—-0.4 eV). Thus, Cl and CO molecules are more promising ligands than CN to improve activity.

Rationalizing the change in intermediate binding energies based on, e.g., Bader charge analysis does not reveal any clear correlation between binding energies and charges of the central metal atoms in the TM-TCNQ monolayers. Central metal atoms carry more positive charge after grating ligands except in the case of Mn-TCNQ-CO, while the \(E_{\text{ads}}\) values of all the adsorbates decrease consistently. Thus, coordinating various ligands can greatly change the adsorption profiles and violate the original scaling relation providing more space to rational design high active catalysts. We note that recent experiment also indicates Cl anion are not able to fully poison iron atoms of FeNiCs moieties\textsuperscript{44} but can influence the activity of ORR in the electrolyte solution suggesting moderate binding energy of Fe-Cl which is consistent with our DFT calculation. Besides, Fe-TCNQ\textsuperscript{45} and Mn-TCNQ\textsuperscript{45, 46} have been produced at present. Herein, we propose Fe-TCNQ-Cl (\(\eta=0.32/0.34\) V), then following Mn-TCNQ-Cl (\(\eta=0.41/0.44\) V) and Mn-TCNQ-CO (\(\eta=0.33/0.54\) V) as bifunctional catalysts for the ORR/OER reactions with quite high catalytic activities comparable to the two best ORR and OER electrocatalysts Pt (\(\eta=0.48\text{eV}\))\textsuperscript{30} and RuO\textsubscript{2} (\(\eta=0.42\text{eV}\))\textsuperscript{31}, respectively. Strain engineering in 2D materials is a feasible technology to achieve tunable electronic properties\textsuperscript{47-50}. Previous studies reveal that external strain can change the binding of adsorbates\textsuperscript{31} due to shifts in 2d-band centers\textsuperscript{17}. Inspired by this approach, we also perform extensive calculations of TM-TCNQ (TM=Mn, Fe, Co, Ir) to examine effect of strain on the catalytic performance by applying between -3% compressive to 3% tensile strain. The result indicates that \(\eta\) of Co-TCNQ for OER is decreased from 0.58 V to 0.35 V under 3% compressive strain. However, the catalytic performances of other candidates (TM=Mn, Fe, Ir) for ORR/OER under strain are not improved accordingly (see details in Supporting Information).

4. Conclusion
To summarize, we have systematically investigated stabilities and catalytic behaviors of twelve TM-TCNQs toward the ORR and the OER by using density functional theory. The adsorption strength of reaction intermediates on TM reactive sites is strongly correlated to the \(d\)-band centers. Except for Pt-TCNQ, the studied catalysts possess excellent conductivity for fast electron transfer during the catalytic processes. Fe-TCNQ is found to exhibit the highest catalytic activity for ORR with \(\eta\) of 0.35 V via a four-electron reaction, which is noticeably higher than the widely used Pt metal. The catalytic performances of Ir-, Co-TCNQs for the OER are identified as the highest among 12 candidates. Grafting axial ligands or applying external strain to the TM-TCNQs can further tune the adsorptive profiles of adsorbates leading to enhancement of ORR and OER activities, especially for the OER. Importantly, Fe-TCNQ and Mn-TCNQ monolayers with Cl ligands and Mn-TCNQ with a CO ligand are predicted as efficient bifunctional ORR/OER catalysts whose catalytic activities are comparable or even higher than the two best electrocatalysts Pt and RuO\textsubscript{2}. As Fe-TCNQ and Mn-TCNQ being synthesized at present, our findings open novel ways to design durable, non-precious metal, efficient catalysts of ORR/OER in metal-air batteries and fuel cells.

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Supplementary Material
Overpotentials (η) of TM-TCNQ monolayer (TM=Mn, Fe, Co, Ir) toward ORR and OER under strain. Scaling relations of the adsorption energies of *OH vs. *OH and *O vs. *OH on TM-TCNQ monolayers by strain. Coordinates of Fe-TCNQ.

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Notes and references


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