Theoretical insight into highly durable iron phthalocyanine derived non-precious catalysts for oxygen reduction reactions

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N4-chelate macrocycles comprise the foundation for non-precious metal oxygen reduction reaction (ORR) catalyst research, where the main electrochemical process occurs in polymer electrolyte membrane (PEM) fuel cells. Although iron–nitrogen–carbon (M–N–C) complexes are known to be the most active non-precious ORR catalysts to date, a fundamental understanding of the ORR mechanisms of these materials is still in its nascent stage and needs further investigation. In this work, ab initio density functional theory (DFT) calculations have been applied to unveil the underlying principles for the electrocatalytic activity and structural stability of Fe–N4 chelates exposed to acidic media. Therefore, we compared the electronic structures of ferrous phthalocyanine (Fe-Pc) and an in-house developed Fe-Pc modified with diphenylphenthioether substituent species (Fe-SPc). The results of these DFT simulations directly correlate with the results of the half-cell ORR activity and stability electrochemical testing in 0.1 M HClO4. The results indicate that the relative energetic position of the dzz-orbital with respect to the Fermi level can induce a Fe redox couple potential shift and modulate the catalytic activity towards the ORR. Furthermore, our combined DFT calculations and empirical observations highlight that the relative position of the dzz-orbital can be controlled by the incorporation of functional groups, resulting in the ability to tune the ORR activity of these complexes. Structural stability of the materials, as predicted by the DFT-calculated cohesive energies of Fe and FeO, can also be readily tuned by modulating Fe-Pc with the substituent species. This study, coupling rigorous experimental observations with DFT investigations, thereby provides a fundamental insight that can aid in the design of future generations of non-precious ORR catalysts with improved activity and stability.

1. Introduction

Accurate understanding of oxygen reduction reaction (ORR) mechanisms is vital for improving the efficiency of many devices, including metal-air batteries,1,2 fuel generation devices3,4 and fuel cells.5,6 This issue is highly pertinent to proton exchange membrane (PEM) fuel cells, which are non-polluting electrochemical power generation devices that convert the chemical energy of fuels into electricity, heat and water. Despite the clean and thermodynamically efficient operation of these devices, there are many technical and economic challenges that still prevent their widespread application.9,11,15,17 The main obstacles include the inherently sluggish ORR kinetics and the low durability of catalysts under the harsh operating conditions.18-28 Until now, the best and most widely used catalysts are expensive Pt-based materials, owing to their relatively high activity and stability in acidic media19-22 in comparison to other metal and non-metal based catalysts. The reliance on high-priced Pt catalysts has provided economical motivation to develop various non-precious ORR catalyst materials, including transition metal–N4 chelate macrocycles, nitrogen-doped carbons, and metal-nitrides, oxides and chalcogenides.23-27

Transition metal–nitrogen–carbon (M–N–C) complexes based on iron, first prepared by the heat treatment of Fe–N4 chelate macrocycles, or more recently, relatively inexpensive inorganic and organic precursors,26-32 have emerged as the most promising non-precious catalysts. It is a well-established fact that heat treatment at temperatures ranging from ca. 700 to 1000 °C is important for imparting high catalyst activity and stability. However, significant structural reconfiguration and the heterogeneous complexity of the M–N–C materials resulting

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from the heat treatment have led the majority of studies to rely on trial-and-error-based approaches. These investigations involve the empirical optimization of various thermodynamic and kinetic variables, and they are performed by systematically varying the catalyst synthesis procedures (i.e. precursor types and concentrations, mixing/blending strategies, heat-treatment temperatures and treatment durations, post-treatment procedures, etc.). In addition, a poor understanding of the active sites in M–N–C catalysts has further made the fundamental mechanisms underlying the ORR activity and stability immensely challenging to elucidate. Without this knowledge, it is impossible to apply rational design strategies to prepare non-precious ORR catalysts with improved activity and stability in PEM fuel cells.

Previously, we reported a new iron phthalocyanine (Fe-Pc)-based material functionalized with four diphenylthioether substituent species (Fe-SPc) as a non-precious ORR catalyst and investigated the half-cell electrochemical properties in 0.1 M HClO₄. Fe-SPc was prepared in lieu of a pyrolysis process, enabling us to adequately control the atomic structure and surface properties of the Fe-SPc materials. A significantly improved electrochemical stability of Fe-SPc was observed in comparison to the Fe-Pc catalyst. This could be attributed to the incorporation of thiether substituent groups that were speculated to provide electron-donating capabilities to the metal-ion center. Conclusively explaining the electrochemical performance variations between Fe-Pc and Fe-SPc and understanding the electronic structure–property relationships of this class of materials will provide valuable fundamental insights that could guide future catalyst design endeavors.

In order to fundamentally elucidate the structural properties and to explain the empirical results with a fundamental mechanistic understanding of the underlying catalysis for a variety of different materials and reactions, ab initio computational methods using density functional theory (DFT) have emerged as an extremely useful tool. For example, ab initio computational studies have been used to confirm Sabatier’s principle, where a good electrocatalyst should moderate the chemical interactions with the reaction intermediates. This has led to the proposition that a simple electronic structure parameter, the d-band center, could successfully dictate the ORR kinetics occurring on a wide array of materials. According to this principle, the N₄-chelate macrocycles should also provide finite chemisorption energies between the oxygen intermediates and the ORR active surface sites, and a certain value exists that can maximize the ORR turnover frequencies.

It is known that the electronic structure of a metal ion can be modified by the surrounding π-conjugated ligand topologies, thereby causing a change in its redox potential. This indicates that variations in the electronic properties of the overall macrocycle structures are governed by contributions from both the metal-ion center as well as the surrounding ligand structure. As such, it is necessary to prepare well-defined model systems and to develop accurate molecular orbital analysis to acquire a fundamental understanding of the ORR catalytic activity. In the present work, we utilize first principles DFT calculations to understand the particular mechanisms governing the catalytic activity and structural stability of Fe-SPc and Fe-Pc materials from the perspective of macrocycle electronic structure. Empirical electrochemical measurements published previously were used in combination with molecular orbital analysis to identify the key fundamental factors controlling the ORR activity and long-term stability, most importantly the electron transfer mechanisms occurring at the molecular scale.

2. Computational design and experimental procedures

Computational details

Density functional theory (DFT) calculations implemented in the Vienna ab initio simulation package (VASP) program were utilized for the present study. Exchange-correlation energies of electrons are described by the Perdew–Burke–Ernzerhof (PBE) functional for generalized gradient approximation (GGA). Core electrons were replaced by the projector augmented wave (PAW) pseudo-potentials and Kohn-Sham wave functions of valence electrons were expanded by a plane wave basis set with a cutoff energy of 520 eV. The designed Fe-Pc and Fe-SPc are located at the center of the vacuum cube box, having lengths of 40 Å for the x, y and z directions. The atoms and cell parameters of each structure were fully relaxed, and spin polarized calculations were subsequently performed. A Gamma point mesh of 1 × 1 × 1 was used to calculate the density of states (DOS) and for finding the thermodynamically stable Fe-SPc and Fe-Pc structures with a Gaussian-smearing width of 0.2 eV.

Synthesis of ferrous 2,9,16,23-tetra(2,6′-diphenylphenothen) phthalocyanine (Fe-SPc)

Fe-SPc was prepared according to the method reported in our previous study. In brief, and depicted schematically in Fig. 1. 4-Nitrophthalonitrile and Fe(OAc)₃ were dissolved in 100 ml of quinoline and stir-fried at 483 K under N₂ protection. After 24 hours, methanol was added to precipitate the materials, and a green powder consisting of ferrous 2,9,16,23-tetranitro phthalocyanine (Fe(nitro)Pc) was obtained by purification with column chromatography on a silica gel, with pyridine as the eluent. The 2,6-diphenylthiophenol, Cs₂CO₃, N-methylpyrrolidone (NMP), and toluene were stirred at 418 K under N₂ atmosphere until the water and toluene evaporated, and the
mixture was cooled down subsequently. The prepared Fe-(nitro) Pc was then added into the reaction mixture under N₂ protection and heated at 453 K, with constant stirring for 12 h. After cooling, the mixture was poured into methanol to precipitate the resulting green powder, which was then purified by column chromatography on silica gel with dichloromethane.

**Electrode preparation and electrochemical measurements**

The electrode preparation and electrochemical analysis carried out previously are summarized in this section. Electrochemical measurements were performed via cyclic voltammetry (CV) using a bipotentiostat (PINE research instrumentation). The half-cell system consisted of a pre-treated glassy carbon working electrode (0.196 cm²), a platinum wire counter electrode and a reference electrode (Reversible Hydrogen Electrode, Hydroflex™, RHE). The Fe-SPc was refuxed for 2 h on KJ300, which was sonicated in 1,2-dichloroethane. The Fe-SPc/KJ300 was then collected after the solvent was distilled off. 20 mg of Fe-SPc was dissolved in 5 mL of 1,2-dichloroethane at the same time. Catalyst inks were prepared as follows: 2 mg catalyst samples were suspended in 1 mL of ethanol and a Nafion® ionomer solution (5 wt%) and sonicated. Subsequently, 60 μL of the catalyst solution was pipetted out and transferred to the working electrode. N₂ or O₂ gas was purged into the electrolyte for at least 30 min before performing electrochemical measurements. The ORR polarization curves for the RDE were obtained in an O₂-saturated 0.1 M HClO₄ solution at room temperature. A positive sweep was performed at a scan rate of 10 mV s⁻¹ and at the electrode rotation rates of 400, 900, 1600 and 2500 rpm.

**3. Results and discussion**

To understand the mechanistic origin of ORR activity and structural stability of Fe-SPc in acidic conditions, four possible positional isomers (e.g. Cᵥᵥ, Cᵥ₂ᵥ, Dᵥ₂ and Cᵥ₄ᵥ symmetry) were designed by replacing the H atoms on each of the four external heterocycles in Fe-Pc with diphenylthiophenol substituent groups. The Cᵥ₄ᵥ structure showed the most stable structure, which is consistent with our previous reports. Fig. 2a and b show the DFT optimized Fe-Pc and Fe-SPc structures, respectively. The structure of Fe-Pc was optimized as a nearly planar macrocycle structure, whereas Fe-SPc appeared to undergo structural distortion owing to the diphenyl thiophenol groups. The bond lengths between the central Fe atom and surrounding N atoms were denoted as N₁, N₂, N₃ and N₄ in Fig. 2 and were calculated to be ca. 1.93–1.94 Å in Fe-Pc as summarized in Table 1. This was also consistent with previous reports. It has been proposed that the changes in the molecular geometry, particularly the Fe–N bonding lengths, arise due to the modulated electronic properties of the metal-center, induced by the effect of the diphenyl thiophenol groups on the surrounding ligand structures in Fe-SPc. We verified this speculation by calculating the atomic partial charges on Fe-Pc and Fe-SPc and by relating the structural change to the nature of the Fe metal-center. We believe that the central Fe-ion may have significant orbital hybridizations with the surrounding N atoms, thereby affecting the catalytic performance. Fig. 3a and b display the differences in the charge density, which are defined by eqn (1) given below,

\[ \rho = \rho_{Fe-SPc} - \rho_{Fe-Pc} - \rho_{N} - \rho_{H} - \rho_{Fe} - \rho_{S} \]  

The yellow and blue areas in Fig. 3 indicate the electronic charge accumulation and depletion, respectively. Both Fe-Pc and Fe-SPc show that the electronic charge distribution is symmetric, and that the charges are accumulated around the nitrogen (N) atoms owing to its high electronegativity (3.04) on the Pauling scale. The central Fe-ion in Fe-Pc demonstrates a depletion of charge, whereas the addition of electron-donating diphenyl thiophenol functional groups on Fe-SPc leads to charge accumulation. The projected density of states (PDOS) of the valence Fe electrons are provided in Fig. 4, demonstrating that the electronic structure of Fe-SPc changes significantly with respect to Fe-Pc. The DOS of Fe in Fe-Pc and Fe-SPc displayed in Fig. 4a and b, respectively, demonstrate bonding-antibonding splitting, which provides indication of the strong chemical bonds between the metal and non-metal atoms. The octahedral environment gives rise to doubly and triply degenerated \( \epsilon_g \) and \( \epsilon_u \) sets, which consist of the axial orbitals of \( d_{xy}, d_{yz} \) and \( d_{xz} \), and the interaxial of \( d_{xy}, d_{xz} \) and \( d_{yz} \) orbitals. Fig. 4 demonstrates occupied and unoccupied molecular orbitals for the ground states of the Fe-Pc and Fe-SPc molecules. For the Fe-Pc and Fe-SPc with a local octahedral symmetry (or square planar symmetry, \( C_{4v} \)) coordination of the Fe center, the uppermost band is made up of \( d_{xy}, d_{xz} \), and \( d_{yz} \) levels, an observation consistent with previous reports. In the PDOS for the 3d electrons of Fe in Fe-Pc (Fig. 4a), the electrons of Fe-\( t_{2g} \)-like bands (e.g., \( d_{xy}, d_{xz} \), and \( d_{yz} \)) in the up-spin state are almost fully occupied, whereas those in a down-spin state are half-filled near the Fermi level. In contrast, adding a diphenyl thiophenol functional group on Fe-Pc shifts the Fermi energy to a more positive value, and the \( t_{2g} \)-like orbital near the Fermi level is fully filled for Fe-SPc. This can be attributed to the
additional electrons provided by the ring ligands, as demonstrated in Fig. 3.

To understand the ORR activity mechanism from molecular orbital analysis, we designed model systems including either O2 or O adsorbed on Fe-Pc and Fe-SPc, as shown in Fig. 5. From the optimized structures, the bonding lengths between Fe and the nearest neighboring O atom of O2, which were directly associated with the adsorption energy of O2 on the Fe present in the macrocycle complexes, were evaluated to be 1.813 Å and 1.812 Å for Fe-Pc and Fe-SPc, respectively. Fig. 6a and b illustrate that the d_{x^2-y^2} orbital in the projected-density of states (PDOS) does not directly link with the O2 adsorption on both Fe-Pc and Fe-SPc, showing that the peaks for the d_{x^2-y^2} orbital are in fact formed without the effect of split-off bands, thereby indicating strong chemisorption. However, Fig. 6c and d show that the electronic d_{z^2} orbital is split into bonding and antibonding states before and after the adsorption of O2, suggesting that the d_{z^2} orbital in the PDOS is more or less a dominant factor influencing the ORR activity. This in turn highlights that the d_{z^2} orbital is closely linked with the redox potential of the Fe-ion, a value that has been well established to govern the onset potential for ORR on these macrocycle materials.\(^{34,60-62}\) The triangles in Fig. 6c and d demonstrate that the level of the d_{z^2} orbitals in the up-spin state and under the Fermi level in Fe-SPc is shifted upwards in comparison to the Fe-Pc shift from \(-2.158\) eV to \(-2.083\) eV.

To truly identify the relationship between the d_{z^2} orbital and the ORR activity, the Fe-SPc and Fe-Pc supported on KJ300 were subjected to rotating disk electrode (RDE) voltammetry, as shown in our previous work.\(^{33}\) The ORR is a multi-electron transfer reaction with two primary pathways.\(^{63-65}\) The first step involves the transfer of two electrons to produce hydrogen peroxide,\(^{63-65}\) \(\text{H}_2\text{O}_2\), and the second is the direct four-electron pathway that produces water, \(\text{H}_2\text{O}\), described by the following set of reactions (2 and 3):\(^{63,65}\)

\[
\text{O}_2 + 2\text{H}_2 + 2e^- \rightarrow \text{H}_2\text{O}_2 \tag{2}
\]
The ORR polarization curves of the catalysts were obtained at rotation rates varying from 400 to 2500 rpm in an O2-saturated 0.1 M HClO4 electrolyte. After removing the background currents to eliminate capacitive contributions, the number of electrons transferred for the reduction of O2 on both Fe-Pc and Fe-SPc was calculated to be ca. 4, using the Koutecky–Levich eqn (4) (ref. 66 and 67) mentioned below:

\[
i_{\text{id}} = 0.62n_e F S D_{\text{O}_2}^{2/3} \rho^{-1/6} c_0^{1/2} \sigma^{1/2} = B \sigma^{1/2}
\]

where, \(i_{\text{id}}\) is the disk current density, \(n_e\) is the number of electrons transferred per molecule of O2, \(F\) is Faraday’s constant, \(S\) is the geometric surface area of the electrode, \(D_{\text{O}_2}\) is the diffusion coefficient of the dissolved \(\text{O}_2\) in the electrolyte, \(\rho\) is the electrode rotation rate in radian s\(^{-1}\), \(c_0\) is the concentration of the dissolved \(\text{O}_2\) in the electrolyte and \(B\) is the Levich constant. In Fig. 7a, as previously reported,\(^{11}\) the polarization curves were obtained in O2-saturated 0.1 M HClO4 at 298 K using a potential scan rate of 10 mV s\(^{-1}\) from 0 to 1.0 V vs. RHE and at the rotation speed of 400 rpm. Here, Fe-Pc/KJ300 demonstrates an increased on-set potential and a half-wave potential (0.62 V vs. RHE), which is 20 mV higher than that of Fe-SPc/KJ300 (0.60 V vs. RHE), indicating better ORR performance without the addition of the diphenyl thiophenol substituent groups.

The first electron transfer to the adsorbed \(\text{O}_2\) to form the superoxide species is generally considered the factor that governs the onset potential for ORR.\(^{61,68,69}\) The mechanism can be expressed by eqn (5) and (6):

\[
\text{O}_2 + 4\text{H}^+ + 4e^- \rightarrow 2\text{H}_2\text{O}
\]

Here, \(O_2^*\) is the adsorption of \(\text{O}_2\) onto catalyst surface. The energy difference (\(\Delta O2p\)) between the Fermi level and the peak positions of the unoccupied \(O2p\) orbital, denoted by an asterisk in the PDOS for the adsorbed \(\text{O}_2\) (Fig. 7b and c), could provide insight into the first electron transfer reaction (5), wherein the smaller \(\Delta O2p\) values indicate an easier electron transfer. The energy level of the \(O2p\) orbital in the antibonding states of \(\text{O}_2\) with down spin states are 0.477 eV and 0.481 eV for Fe-Pc and Fe-SPc, respectively. Therefore, the higher onset potential of Fe-Pc in Fig. 7a could be interpreted from the computational results, which indicate that the \(\Delta O2p\) value of Fe-Pc is smaller than that of Fe-SPc, leading to a relatively easier electron transfer to the O atom, thereby improving the ORR kinetics. From these observations, it can be inferred that the lower ORR onset potential of Fe-SPc is attributed to the electronic structure changes of the central Fe atom, induced by adding the electronic donating substituents, which result into an upward shift of the \(c_g(\text{d}_{z^2})\) orbital and limit the electron transferability to the adsorbed \(\text{O}_2\) molecules.
Using the calculated thermodynamics of different intermediates, we make direct comparisons to voltammetric and rotating ring disk electrode experiments, to evaluate the electrochemical activity of the Fe-Pc and Fe-SPc surface for the ORR. With the DOS studies, thermodynamic free energy diagrams were calculated to define a rate-determine step of the ORR on the Fe-Pc and Fe-SPc surfaces. Four electron reaction pathways were considered, as shown in eqn (7)–(10):

\[
\begin{align*}
O_2(g) + 4H^+ + 4e^- &\rightleftharpoons OOH^* + 3H^+ + 3e^- \\
OOH^* + 3H^+ + 3e^- &\rightleftharpoons O^* + H_2O(l) + 2H^+ + 2e^- \\
O^* + H_2O(l) + 2H^+ + 2e^- &\rightleftharpoons [OH]^* + H_2O(l) + H^+ + e^- \\
[OH]^* + H_2O(l) + H^+ + e^- &\rightleftharpoons 2H_2O
\end{align*}
\]

(7)–(10)

The details regarding the ORR free energy diagrams are well-described in the previously published reports. Briefly, the free energy of any ORR intermediate with respect to liquid H₂O (ΔG) was calculated using eqn (11):

\[
\Delta G = \Delta E + \Delta ZPE - T\Delta S
\]

(11)

where ΔE, ΔZPE and ΔS denote the DFT-calculated ground state energy, zero point energy, and entropy corrections, respectively. The ΔZPE for intermediates were adopted from Nørskov et al., which used ΔS at T = 298 K, as presented by a thermodynamic table. The effect of the potential was evaluated at pH = 0 by shifting the electronic chemical potential according to Nernst’s law. Fig. 8 demonstrates the free energy change of each ORR step for the ideal, Fe-Pc and Fe-SPc at zero electrode potential (Φ = 0 V vs. RHE), and at the potential for which all steps become energetically negative. At the equilibrium potential (U = 1.23 V), the O* formation is energetically negative (exothermic) and all other step are positive (endothermic) for both Fe-Pc as well as Fe-SPc. By applying the overpotential, all the steps become negative for the ORR at 0.67 V and 0.59 V vs. the RHE on the Fe-Pc and Fe-SPc, respectively. The calculated thermodynamic free energy diagram predicts that the Fe-SPc activity is lower than that of the Fe-Pc activity, consistent with the RDE experimental results. With even a slightly lower overpotential than the potential at which every reaction step is energetically negative (i.e., 0.67 and 0.59 V vs. RHE for Fe-Pc and Fe-SPc), the reduction step of O* to OH* will become energetically positive, revealing that the O*/OH* exchange is the rate-determining step for the ORR.

To investigate the impact of adsorption strength on the ORR activity, the adsorption energy (E_{ads}) of the O or O₂ molecules on these macrocycle complexes were calculated using eqn (12):

\[
E_{ads} = E_{O,O_2,Fe-Pc} - E_{Fe-Pc} - E_{O,O_2,Fe-SPc}
\]

(12)

Fig. 8 Standard free energy diagram of (a) ideal, (b) Fe-Pc and (c) Fe-SPc for the ORR at zero potential (U = 0), equilibrium potential for ORR (U = 1.23 V) and at the potential for which all the steps become energetically negative at pH = 0 and at T = 298 K.
SPc is too strong to obtain a high activity for the ORR. Accordingly, a downshift in the energy level of the $d_{z^2}$ orbital away from the Fermi level leads to a shift in the Fe ion’s redox potential, and by extension, results into smaller $\Delta O2p$ values and higher ORR activity potentials. In summary, the ORR activity of metal macrocycle complexes is governed by the relative position of the $d_{z^2}$ orbital of Fe in comparison to the Fermi level, leading to the peak positions of the unoccupied $O2p$ orbital in comparison to the Fermi level. The optimized adsorption strengths of the ORR intermediates could be achieved by controlling the $d_{z^2}$ orbital level, which could be downshifted by the attachment of electron withdrawing functional groups.

It has been proposed that the main degradation mechanism of Fe-N$_4$ chelate macrocycle catalysts in acidic electrolytes is the demetalation of the macrocycles.\textsuperscript{43,49} This leads to a dramatic loss in the ORR performance because the resulting H$_2$Pc materials have very limited activity.\textsuperscript{53,60} The demetalation process is highlighted in reaction (13):

\[
Fe-(S)Pc + 2H^+ \rightarrow H_2(S)Pc + Fe^{+2} \tag{13}
\]

The redox couples of Fe-Pc and Fe-SPc observed in the cyclic voltammograms (CV) of Fig. 9a and b indicate the presence of the Fe-ion centers in the macrocycle complexes. Diminishing charges of the redox couple can therefore provide evidence of the Fe-N$_4$ demetalation.\textsuperscript{11,50,51} Fig. 9 clearly demonstrates that functionalized diphenyl thiophenol groups on the Fe-Pc dramatically enhance the stability of Fe-SPc over 100 cycles in the oxygen-saturated electrolyte. After 100 cycles, the CV curve of Fe-Pc exhibits a significant loss in the redox charge transfer of Fe in comparison to the initial plot (Fig. 9a). This observation is in agreement with the significant loss of ORR activity (Fig. 9b), indicating the occurrence of Fe-Pc demetalation under these conditions. On the other hand, after 100 cycles, Fe-SPc still maintains the distinct Fe-ion redox peaks (Fig. 9c), and shows only a 20.3 mV loss in the half-wave potential for the ORR (Fig. 9d).

This pronounced increase in the electrochemical stability of the Fe-SPc macrocycle is of high interest, and has been investigated by calculating the cohesive energies ($E_{coh}$) of Fe and FeO in the ferrous phthalocyanine macrocycle complexes through the first principle calculations using eqn (14) and (15) as follows:

\[
E_{coh} = -(E_{Fe-(S)Pc} - E_{(S)Pc} - E_{Fe,g}) \tag{14}
\]

\[
E_{coh} = -(E_{FeO-(S)Pc} - E_{(S)Pc} - E_{FeO,g}) \tag{15}
\]

here, $E_{FeO-(S)Pc}$ is the total energy of the O adsorbed onto Fe-Pc and Fe-SPc, $E_{Fe,g}$ and $E_{FeO,g}$ are total energy of Fe and FeO in the gas phase, and $E_{(S)Pc}$ is total energy of the phthalocyanine macrocycle frame of Fe-Pc and Fe-SPc without the central Fe atom. In the case of metals, it has been proposed that the cohesive energy and electrochemical dissolution potential of the metal are strongly correlated,\textsuperscript{22,73,75-78} and a higher cohesive energy of the metal indicates a higher dissolution potential. Thus, the cohesive energies of Fe or FeO in the macrocycle complexes are likely to be related to the durability of iron based materials under harsh conditions. DFT predicts, however, that the cohesive energy of Fe in the macrocycle complex is 9.80 eV for Fe-Pc, which is higher than 9.53 eV for Fe-SPc. These results do not align with our electrochemical measurements in acidic electrolytes discussed previously. Therefore, to provide a reasonable explanation surrounding the observed demetalation of the ferrous phthalocyanine complex, the presence of oxygen requires further consideration. In fact, it was found that the demetalation and activity losses of Fe-Pc and Fe-SPc were minimized under inert atmospheres, whereas the activity was significantly compromised in the presence of oxygen that would attack the active site structures.\textsuperscript{53,54} When oxygen is adsorbed onto Fe-Pc and Fe-SPc, the electron transfer from the central metal to the oxygen molecule could decrease the atomic radius of the central Fe, and thus the distance of the Fe-N bonds could be increased.\textsuperscript{53,54} Even in this work, the distance between the central Fe and N atoms is increased from ca. 1.93 Å to 1.96 Å for FePc and Fe-SPc upon the adsorption of oxygen, and these results agree well with previous investigations.\textsuperscript{53,79} The FeO cohesive energy of 10.22 eV for Fe-SPc is higher than 10.15 eV for Fe-Pc, as summarized in Table 1, thereby indicating that the electrochemical stability in the presence of oxygen for Fe-SPc is enhanced in comparison to Fe-Pc. Therefore, the diphenyl thiophenol electron donating groups substituted onto the phthalocyanine complex could exclusively help to suppress the demetalation and enhance the stability of Fe-SPc in the presence of oxygen, which is the necessary reactant at the cathode of PEMFCs.

4. Conclusion

Using first-principles DFT calculations, we have studied the scientifically relevant ORR processes occurring on Fe-Pc and Fe-SPc materials in terms of electrochemical activity and stability.
The results indicate that the $d_\sigma$ orbital in the DOS is closely related with ORR activity, showing electronic $d_\sigma$ orbital splitting into bonding and antibonding states before and after the adsorption of $O_2$. The position of the $d_\sigma$ orbital of Fe in comparison to the Fermi level, a factor related to the unoccupied $\Delta O_2p$ orbital level, ultimately determines the ORR onset potential and oxygen intermediate adsorption strengths of metal macrocycle complexes, owing to the modulated electron transfer capabilities to the O atom. The free energy diagram clearly indicates that a rate determining step is in the reduction step of $O^* \rightarrow OH^*$, which indicates that the overall ORR kinetics are governed by the oxygen adsorption strength on Fe-Pc and Fe-SPc. The relatively upward shift of the $d_\sigma$ orbital of the Fe atom in Fe-SPc provides stronger bonding of the oxygen intermediates, leading to a reduction in the activity towards the ORR. This prediction is consistent with the results from electrochemical testing carried out in $O_2$-saturated acidic condition, wherein the on-set potential and a half-wave potential of Fe-SPc/KJ300 was reduced in comparison to Fe-Pc/KJ300. The free energy diagram prepared using DFT also predicts the same activity trends, wherein the activity of Fe-SPc is slightly lower than that of Fe-Pc. It is therefore clear that the optimized adsorption strengths of the ORR intermediates can be achieved by carefully tuning the level of the $d_\sigma$ orbital. Interestingly, the electrochemical stability of Fe-SPc, evaluated by accelerated testing protocols, was significantly enhanced in the presence of oxygen in comparison to Fe-Pc. This observation can be explained using theoretical calculations, which show an increased cohesive energy of FeO for Fe-SPc in comparison to Fe-Pc. Furthermore, the diphenyl thiophenol electron donating groups substituted onto the phthalocyanine complex could help suppress the demetalation of the macrocycle structure by providing extra electrons to the metal ion center. Since these investigated electronic properties can be directly modulated by the appropriate selection and attachment of various substituent groups on the transition metal macrocycles, this comprehensive DFT study linked to experimental observations can provide the fundamental basis for the design of future generation non-precious ORR catalysts with improved activity and stability.

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