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Evaluation of CH_4 oxidation activity of high-valent iron-oxo species of a μ -nitrido-bridged heterodimer of iron porphycene and iron phthalocyanine†

Yasuyuki Yamada, [®]** Yusuke Miwa,^a Yuka Toyoda,^b Quan Manh Phung, [®]** Kin-ichi Oyama [®]* and Kentaro Tanaka [®]**

A μ -nitrido-bridged dimer of iron phthalocyanine is one of the most potent molecule-based CH₄ oxidation catalysts reported to date. The reactive intermediate is a high-valent iron-oxo species generated through reaction with H₂O₂ in an acidic aqueous solution. However, there are few reports on the synthesis and catalytic CH₄ oxidation activity of a μ -nitrido-bridged heterodimer of two different iron porphyrinoids, despite that there are a variety of iron porphyrinoids with coordination and electronic structures different from those of iron phthalocyanines or iron porphyrins. Herein, we report the synthesis of a novel μ -nitrido-bridged heterodimer of an iron phthalocyanine and iron porphycene and examine its CH₄ oxidation activity. Porphycenes are an important class of porphyrinoids with a smaller coordination sphere than phthalocyanines or porphyrins. Single crystal structural analyses revealed that the heterodimer possessed a Fe-N=Fe core structure similar to that of the phthalocyanine homodimer. The heterodimer showed catalytic CH₄ oxidation activity in an acidic aqueous solution in the presence of H₂O₂ at 60 °C through the high-valent iron-oxo species as in the case with the phthalocyanine homodimer. This was in clear contrast to the result that the high-valent iron-oxo species of a μ -nitrido-bridged iron porphycene dimer was so unstable that it decomposed quickly in the same reaction conditions.

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Introduction

Several decades have passed since some methane monooxygenases (MMOs) were found to utilize high-valent iron-oxo species as reaction intermediates for the catalytic conversion of CH₄ into CH₃OH under ambient reaction conditions. CH₄ has long been expected to be the next-generation carbon resource because it is abundant in nature as natural gas or methane hydrate. However, CH₄ is a stable organic compound with a particularly high C-H bond dissociation energy of *ca.* 105 kcal mol⁻¹. Therefore, the development of an efficient catalyst is indispensable for low-cost and efficient conversion of CH₄ into industrially useful

However, from the viewpoint of catalytic activity, a very limited number of artificial molecular iron-oxo-based biomimetic catalysts exist that activate the C-H bonding of CH₄ at temperatures lower than 100 °C because of the high stability of CH₄. 3,18,20 A. B. Sorokin *et al.* found that a μ -nitrido-bridged iron phthalocyanine (Pc) dimer (1 or 1⁺) reacts with H₂O₂ in an acidic aqueous solution to afford a high-valent iron-oxo species $\mathbf{1}_{oxo}$, which shows a particularly high CH₄ oxidation activity among various molecular CH₄ oxidation catalysts at a lower temperature than 100 °C (Fig. 1a). $^{20-22}$ Similarly, μ -nitrido-bridged iron porphyrin dimer 2 was reported to indicate CH₄ oxidation activity

chemical raw materials such as CH_3OH . Therefore, it is quite understandable that the fact that some MMOs achieve efficient catalytic conversion of CH_4 into CH_3OH at ambient temperature by utilizing high-valent iron-oxo species has stimulated the interest of chemists to fabricate a large variety of biomimetic iron-oxo-based molecular catalysts. The detailed reaction mechanisms of natural MMOs have been clarified as a result of utilizing these biomimetic catalysts. The CH_4 oxidation reaction by high-valent iron-oxo species is facilitated by the proton-coupled electron transfer (PCET) pathway, enabling low-temperature and efficient CH_4 oxidation. 18,19

^a Department of Chemistry, Graduate School of Science, Nagoya University, Furocho, Chikusa-ku, Nagoya 464-8602, Japan. E-mail: kentaro@chem.nagoya-u.ac.jp, yy@chem.nagoya-u.ac.jp

^b Research Center for Materials Science, Nagoya University, Furo-cho, Chikusa-ku, Nagoya 464-8602, Japan

^c Institute of Transformative Bio-Molecules (ITbM), Nagoya University, Furo-cho, Chikusa-ku, Nagoya 464-8601, Japan

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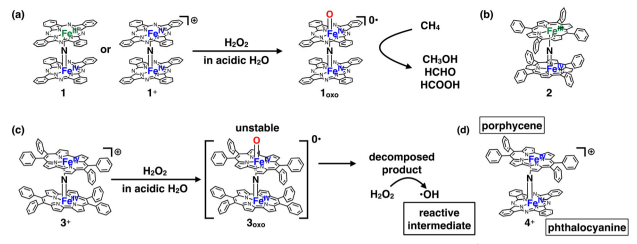


Fig. 1 (a) Formation of high-valent iron oxo species $\mathbf{1}_{oxo}$ having potent CH_4 oxidation ability, from $\mathbf{1}$ or $\mathbf{1}^+$ through the reaction with H_2O_2 . (b) A μ-nitrido-bridged iron porphyrin dimer (2) showing potent CH₄ oxidation ability. (c) Structure of a monocationic μ-nitrido-bridged iron porphycene dimer (3⁺) and its proposed CH₄ oxidation mechanism. (d) A monocationic μ-nitrido-bridged heterodimer of an iron porphycene and an iron phthalocyanine (4+) synthesized in this work.

comparable to that of µ-nitrido-bridged iron phthalocyanine dimer (Fig. 1b).²³ According to some theoretical estimates, the core µ-nitrido-bridged dinuclear iron structure (Fe-N=Fe) is believed to be crucial in the reaction of these high CH₄ oxidation activities. 24-27

Inspired by these previous works, we recently synthesized μ-nitrido-bridged dinuclear iron porphycene (Ppc) dimer 3⁺, a structural analog of 1⁺, and investigated its CH₄ oxidation activity (Fig. 1c).28 Porphycene is an important class of porphyrinoids and has a rectangular-shaped narrower cavity than phthalocyanine, which renders the coordination structures and reactivities of the metal complexes different from those of phthalocyanine. 29,30

Although 3⁺ has a μ-nitrido-bridged dinuclear iron core (Fe-N=Fe) similar to that of 1⁺, it was found that 3⁺ showed less CH4 oxidation activity, and the reaction proceeded through a different mechanism (Fenton-type reaction; the reaction intermediate is 'OH). This is presumably because the high-valent iron-oxo species 3_{oxo} is less stable than 1_{oxo} and decomposes during the CH₄ oxidation reaction. The decomposed product could act as a catalyst for 'OH production. These results indicate that the difference in the structure of the porphyrinoid moiety strongly affects the reactivity of the high-valent iron-oxo species.

In this study, we report the synthesis of a μ-nitridobridged heterodimer of an iron phthalocyanine and an iron porphycene (4⁺, Fig. 1d) and its CH₄ oxidation activity. Thus far, there have been very few reports investigating the CH₄ oxidation activity of µ-nitrido-bridged heterodimers of iron porphyrinoids. We previously confirmed supramolecular μ-nitrido-bridged heterodimer of an iron porphyrin and an iron phthalocyanine provided sufficiently stable high-valent iron-oxo species for catalytic CH₄ or CH₃-CH₃ oxidation reactions. 31,32 It has also been reported that H₂O₂ preferentially reacts with the electron-rich iron phthalocyanine moiety of the µ-nitrido-bridged heterodimer

of an electron-rich iron phthalocyanine and an electrondeficient iron phthalocyanine to afford a high-valent iron-oxo species, although the reactivity of the heterodimer has not been reported.³³ Therefore, the fact that a μ -nitrido-bridged iron phthalocyanine dimer 1 (or 1⁺) indicated potent CH₄ oxidation activity through the reaction of high-valent iron-oxo species, whereas a u-nitrido-bridged dinuclear iron porphycene dimer 3⁺ showed a different CH₄ oxidation mechanism, prompted us to clarify the relationship between the structure and reactivity of 4⁺.

Results and discussion

Synthesis and spectroscopic characterization of 4⁺

A mixture of iron porphycene 5 and iron phthalocyanine 6 was heated in 1-chloronaphthalene in the presence of excess

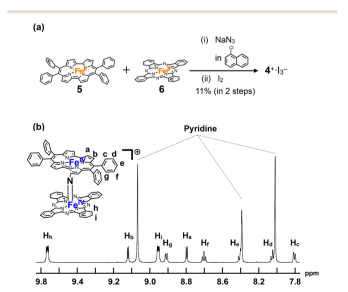


Fig. 2 (a) Synthesis of a monocationic heterodimer 4⁺·I₃⁻. (b) ¹H-NMR spectrum of $4^+ \cdot l_3^-$ in pyridine- d_5 .

NaN₃ at 280 °C, followed by oxidation with iodine to yield a monocationic μ-nitrido-bridged heterodimer of an iron phthalocyanine and iron porphycene (4⁺·I₃⁻, Fig. 2a). When an equimolar mixture of iron porphycene 5 and iron phthalocyanine 6 was heated, the isolated yield of the desired compound, 4+I₃, was relatively low (ca. 10%). This is because a significant amount of the μ-nitrido-bridged iron phthalocvanine dimer was obtained (ca. 40%), suggesting that the formation of the µ-nitrido-bridged iron phthalocyanine dimer was faster than the formation of the desired heterodimer under these reaction conditions. Therefore, heating a 2:1 mixture of iron porphycene 5 and iron phthalocyanine 6 with NaN3 improved the isolated yield of $4^{+}\cdot I_{3}^{-}$ to 71%. Elemental analysis suggested that the counter anion of the monocationic heterodimer was triiodide, which was further confirmed using single-crystal X-ray structural analysis.

¹H-NMR and MALDI-TOF MS analyses were performed to characterize 4+·I₃-. In the MALDI-TOF MS spectrum shown in Fig. S1 in the ESI,† a signal corresponding to 4⁺ was observed at m/z = 1250.26, which corresponds to the theoretical isotope distribution pattern of 4⁺. As shown in Fig. 2b, all the ¹H-NMR peaks of 4+I₃ were observed as sharp signals in the range of 6-10 ppm. A neutral μ-nitrido-bridged FeTPP dimer containing Fe(III) and Fe(IV) was reported to indicate a significant broadening of the ¹H-NMR signals, whereas its monocationic species containing two Fe(IV) centers showed sharp ¹H-NMR signals because the two Fe(IV) centers interacted with each other in an antiferromagnetic manner. $^{34-36}$ Therefore, it is considered that $\mathbf{4}^+\cdot\mathbf{I}_3^-$ includes two Fe(IV) ions interacting with each other in an antiferromagnetic manner. Moreover, considering that two sets of signals corresponding to those of the o- and m-protons of the peripheral phenyl groups were observed, the rotation of the peripheral phenyl groups of the porphycene ring was restricted owing to steric repulsion between the phenyl groups and the porphycene ring, as in the case of the previously reported µ-nitrido-bridged dinuclear iron porphycene dimer 3+·PF₆-. However, the rotation of the porphycene ring along the Fe-N=Fe axis was sufficiently fast

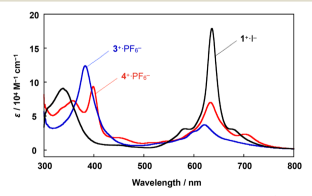


Fig. 3 Comparison of UV-vis spectra of a heterodimer 4+.PF₆ and two homodimers, 1+·l- and 3+·PF₆-, in pyridine at 20 °C.

compared to the NMR timescale because the porphycene ring was observed as a C_2 symmetrical structure.

A comparison of the UV-vis spectrum of $4^+ \cdot PF_6^-$ with those of μ-nitrido-bridged iron phthalocyanine dimer 1+·I and μ-nitrido-bridged iron porphycene dimer 3⁺·PF₆⁻ is shown in Fig. 3. The bands at 357 and 399 nm were assignable to the absorptions by the phthalocyanine and porphycene units, respectively, both of which showed significant shifts compared to those of the homoleptic dimers 1+.I- and 3⁺·PF₆⁻. ^{28,37} The Q-band of the phthalocyanine unit of 4⁺·PF₆⁻ at 634 nm showed an apparent hypochromic shift compared to that of the phthalocyanine dimer 1+·1-, whereas the Q-band of the phthalocyanine unit of 4+·PF₆ at 705 nm appeared at a higher wavelength than that of $\mathbf{1}^+\cdot\mathbf{I}^-$. These results implied an apparent electronic interaction between the two porphyrinoids in $4^+ \cdot PF_6^-$.

Single crystal X-ray structural analysis

Crystal structural analysis of $4^{+}\cdot I_{3}^{-}$ demonstrated that $4^{+}\cdot I_{3}^{-}$ crystallized in the monoclinic space group P2₁/c from a solution of 1:1 (v/v) mixture of pyridine and CHCl₃. The molecular structures of $\mathbf{4}^+ \cdot \mathbf{I_3}^-$ are shown in Fig. 4a and b.

The distance between the μ-nitrogen and Fe(IV) ion in porphycene (1.62(9) Å) is almost identical to that of the μ-nitrogen and Fe(IV) ion in phthalocyanine (1.66(4) Å), suggesting that the Fe- μ -N-Fe core has a π -conjugated structure. The mean bond distances between the four porphycene nitrogens and the coordinated Fe(IV) ion (1.93 Å) almost identical to those between the four phthalocyanine nitrogens and the coordinated Fe(IV) ion (1.94 Å). The average bond angles of the μ-nitrogen–Fe– nitrogens(porphyrinoid) in 4⁺·I₃⁻ were 93.45(8)° for iron phthalocyanine and 93.13(3)° for iron porphycene, both of

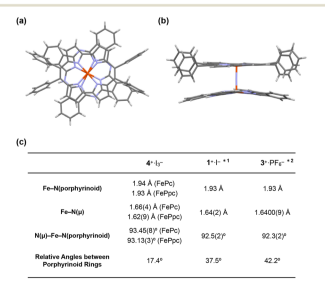


Fig. 4 (a) Side and (b) top views of the single crystal X-ray structure of $4^+ \cdot l_3^-$. l_3^- , coordinating pyridines, and crystalline solvents were omitted for clarity. (c) Comparison of some mean bond lengths and angles of $4^{+} \cdot l_{3}^{-}$, $1^{+} \cdot l_{3}^{-}$, and $3^{+} \cdot PF_{6}^{-}$. *1: ref. 28, *2: ref. 38.

which were slightly larger than 90°. The core-coordinating N4 planes of the two porphyrinoid rings are located in an almost parallel configuration. The phthalocyanine ring was slightly distorted from the planar structure, presumably because of steric repulsion between the phthalocyanine ring and peripheral phenyl groups of the porphycene ring. As suggested by ¹H NMR spectroscopy, the rotation of the peripheral phenyl groups of the porphycene ring appeared to be significantly suppressed by steric repulsion between the phenyl groups and porphycene rings.

A comparison of the single-crystal structure of $4^+ \cdot I_3^-$ with those of 1+·I- and 3+·PF₆- is summarized in a table shown in Fig. 4c.^{28,38} Although slight differences were observed in the distance between nitrogen and Fe(IV) and the average bond angles of the μ-nitrogen-Fe-nitrogen (porphyrinoid), these differences are presumably derived from the heteroleptic structure of $4^+ \cdot I_3^-$. Overall, it could be concluded that $4^+ \cdot I_3^$ has a similar Fe-N-Fe core structure to those of 1+·I- and 3⁺·PF₆. The relative angle between the phthalocyanine ring and the long axis of the porphycene ring was calculated to be 17.4°, which was much smaller than the relative angles between the two porphyrinoid rings in 1+·I⁻ (37.5°) and $3^{+}\cdot PF_{6}^{-}$ (42.2°). We attributed this difference in the relative angle between the two porphyrinoids to the difference in the packing structure, because both rings in 4⁺·I₃⁻ are rotatable along the Fe-N=Fe axis.

Cyclic voltammetric analysis

Cyclic voltammograms of 4+PF₆ were recorded in a pyridine solution (200 μ M) containing 100 mM n Bu₄N ${}^{+}$ ·PF₆ ${}^{-}$ (Fig. 5a). A comparison of the voltammograms of the μ-nitrido-bridged iron phthalocyanine dimer 1+·I⁻ and μ-nitrido-bridged iron porphycene dimer 3⁺·PF₆⁻ is shown in Fig. 5b and c. 4⁺·PF₆⁻

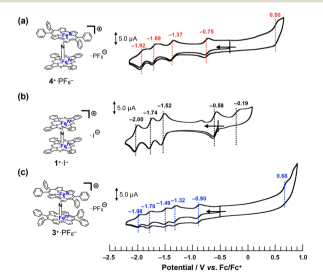


Fig. 5 Comparison of cyclic voltammograms of (a) $4^+ \cdot PF_6^-$, (b) $1^+ \cdot I^-$, and (c) 3+.PF₆ in a pyridine solution containing 100 mM of n Bu₄N⁺·PF₆ at room temperature. [Substrate] = 200 μ M, [scan rate] =

showed four quasi-reversible redox waves at -0.75, -1.37, -1.69 and -1.92 V vs. Fc/Fc⁺ and one irreversible oxidation wave at 0.50 V.

In the case of a voltammogram of a μ-nitrido-bridged iron phthalocyanine dimer 1+1 (shown in Fig. 5b), Ercolani et al. carefully investigated the redox processes to conclude that the four redox waves at -0.58, -1.52, -1.74 and -2.00 V should be assignable to the redox in the iron cores of 1+J-39 For the voltammogram of a μ-nitrido-bridged iron porphycene dimer $3^+ \cdot PF_6^-$ shown in Fig. 5c, we assigned the 1st (-0.90 V vs. Fc⁺/ Fc), 2nd (-1.32 V), 3rd (-1.49 V), and 4th (-1.78 V) reduction waves of the porphycene dimer to those of Fe(IV)Fe(IV)/Fe(III) Fe(π), Fe(π)/Fe(π)/Fe(π)/Fe(π)/Fe(π)/Fe(π), and Fe(π) $Fe(\pi)/Fe(\pi)Fe(\pi)$, respectively. This is based on the fact that Gross et al. investigated the detailed redox properties of a 2,7,12,17-tetra-n-propylporphycene iron(III) chloride (ClFeTprPpc), a monomeric porphycene, by electrochemical and spectroelectrochemical measurements to report that the reduction of the iron center (Fe(III)/Fe(II)) occurred at more positive potential than that of the porphycene center (TPrPpc/ TPrPpc'-).40 Similarly, it is assumed that at least the four reduction waves of $4^+\cdot PF_6^-$ at -0.75, -1.37, -1.69, and -1.92 V could be assignable to those of the iron centers of $4^+ \cdot PF_6^-$. Thus, it was demonstrated that 4+PF₆ has a completely different electronic structure from those of 1+·I- and 3+·PF6owing to the different types of porphyrinoid rings.³⁸

As for the oxidation of 4⁺·PF₆, theoretical calculations suggested that the phthalocyanine ring is more easily oxidized than the porphycene ring (vide infra, see "DFT calculations" section). In addition, the μ -nitrido-bridged iron phthalocyanine dimer 1⁺·I[−] showed a lower oxidation potential at -0.19 V than that of a μ-nitrido-bridged iron porphycene dimer 3⁺·PF₆⁻ at 0.68 V, both of which are assignable to the oxidations of the porphyrinoid cores. Therefore, the oxidation wave at 0.50 V is assignable to that of the phthalocyanine ring. Overall, 4+PF₆ showed a first reduction wave higher than that of $\mathbf{1}^+\cdot\mathbf{I}^-$ and lower than that of 3+PF₆, whereas the first oxidation wave of **4**⁺·PF₆[−] was observed at potentials higher than **1**⁺·I[−] and lower than $3^+ \cdot PF_6^-$.

ESI-TOF MS analysis of high-valent iron-oxo species

To confirm the generation of the high-valent iron-oxo species, the reactive intermediate for the oxidation reaction, by treating $4^+ \cdot I_3^-$ with H_2O_2 in an aqueous solution, ESI-TOF MS measurement of an acetonitrile solution of $4^+ \cdot I_3^-$ in the presence of H₂O₂ was performed.

As shown in Fig. 6, the signals assignable to those of the high-valent iron-oxo species ($[4 + O]^+$) were observed at approximately m/z = 1266 after the addition of aqueous H_2O_2 at room temperature, with its isotopic distribution pattern corresponding to that of the calculated pattern. The signals observed at approximately 1284 correspond to those of the hydroperoxo species ([4 + HOOH]⁺). We also confirmed that treatment of $\mathbf{4}^+ \cdot \mathbf{I_3}^-$ with $\mathbf{H_2}^{18} \mathbf{O_2}$ resulted in the generation of the peaks assignable to [4 + 18O] as shown in the ESI.† These

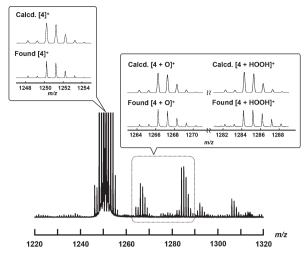


Fig. 6 ESI-TOF MS spectrum of 4⁺·I₃⁻ in an aqueous CH₃CN solution containing excess amount of H2O2 at room temperature (the detail of the experimental procedure is shown in the Experimental section). Insets: Comparison of calculated (top) and experimental (bottom) isotopic distribution patterns of molecular peaks assignable to [4]⁺, [4 + O]⁺, and [4 + HOOH]⁺.

results demonstrate that high-valent iron-oxo species were generated after the threat of 4⁺ with H₂O₂ via the corresponding hydroperoxo species, as in the case with the other µ-nitrido-bridged iron phthalocyanine dimers. Sorokin et al. reported that treatment of heteroleptic u-nitrido-bridged dimer of iron phthalocyanine and iron porphyrazine with m-chloroperbenzoic acid (mCPBA) resulted in the attachment of oxo moiety either on the iron phthalocyanine and iron porphyrazine. In this case, they concluded that the attachment of oxo moiety occurred with slight preference for the iron phthalocyanine unit based on the results of cryospray collision induced dissociation MS/MS technique.41 However, in the case ESI-TOF MS analysis of $[4 + O]^+$, it was difficult to determine the accurate position of the Fe=O moiety (whether it is on the iron porphycene or the iron phthalocyanine in 4⁺) even after MS/MS analysis of the $[4 + O]^+$ peak because the ionization efficiency of the Fe-phthalocyanine fragments was much smaller than that of the Fe-porphycene fragments.

DFT calculations

Density functional theory (DFT) calculations were performed to gain further insight into experimental observations. The calculation details are presented in the Experimental section. We first focused on the electronic structure of $\mathbf{4}^{+}$ as well as its oxidized and reduced species. The electronic structure of 4⁺ is very similar to that of [{(porphyrin)Fe}₂N]⁺, as described by Ghosh et al. 42 (Fig. 7). The predicted Fe-N(Pc) and Fe-N(Ppc) average bond distance is 1.948 and 1.939 Å, respectively, which reasonably agrees with the experimental values of 1.94 and 1.93 Å, respectively (see Tables S1 and S2†). The short Fe-N bond distances were consistent with the low-spin character of the Fe centers. The predicted Fe-N(µ) bond lengths are 1.582 and 1.607 Å, respectively, which are systematically smaller than the X-ray data by approximately 0.05 Å.

Upon reduction, both Fe-N(μ) bonds were lengthened as an antibonding orbital comprising $Fe(3d_{z^2})$ and N(2s)orbitals. The difference between the two Fe-N(μ) bonds is only 0.03 Å, indicating that the two Fe centers are similar and the complex could be formally characterized as Fe^{3.5}-N-Fe^{3.5}. ²⁰ Spin populations also reveal that both Fe centers are involved in the reduction, with Fe(Pc) being slightly more reduced than Fe(Ppc). Upon oxidation, the Fe-N(μ) bond lengths change by less than 0.01 Å, indicating that the oxidation occurs mainly at the macrocycles. The spin population of Pc is 0.76, whereas Ppc has a spin population of 0.15. This difference in the spin density implies that oxidation primarily occurs at Pc. The higher tendency of phthalocyanine to be oxidized is related to its longer conjugation length compared to that of porphycene.

We now compare the two oxo isomers, O=Fe(Pc)-N-Fe(Ppc) and O=Fe(Ppc)-N-Fe(Pc) (DFT structures are shown in Fig. 7). Similar to a previous work, 27 two doublet states were calculated: the ground state in which one of the macrocycles was oxidized Fe^{IV}Fe^{IV}(P*+) and a low-lying excited state Fe^{IV}Fe^V (Fig. S4†). Selected bond lengths are listed in Table S3.† The Fe-N-Fe fragment is no longer symmetric. The general trend is that the formation of Fe-oxo significantly increases the length of the adjacent Fe-N(µ) bond above 1.7 \mathring{A} . DFT predicts that O=Fe(Pc)-N-Fe(Ppc) and O=Fe(Ppc)-N-Fe(Pc) are close in energy. The former isomer (in pyridine at 298 K) was slightly more stable (2-4 kcal mol⁻¹) than the latter (Table S5†).

Considering that the phthalocyanine ring is more easily oxidized than the porphycene ring, H₂O₂ might coordinate to Fe-porphycene more strongly than to Fe-phthalocyanine because the porphycene ring is more electron deficient than the phthalocyanine ring. This suggests that the oxo species of

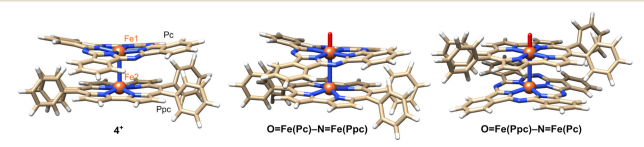


Fig. 7 DFT structures of 4⁺ and its isomeric oxo complexes.

4⁺ could be preferentially generated on the Fe-porphycene moiety. This prompted us to compare the stability of the hydroperoxo species on the Fe-porphycene ring (HOOFePpc-N=FePc) with that of the Fe-phthalocvanine ring (HOOFePc-N=FePpc). We found that the difference between these two species is quite small (ca. 3 kcal mol⁻¹; HOOFePpc-N=FePc is more stable than HOOFePc-N=FePpc). Moreover, we compared the O-O bond dissociation free energy in HOOFePpc-N=FePc and HOOFePc-N=FePpc dissociation of the O-O bond is indispensable for the generation of iron-oxo species from hydroperoxo species. However, it was demonstrated that both hydroperoxo species have similar O-O bond dissociation free energy (37.2 kcal mol⁻¹ for HOOFePpc-N=FePc and 36.7 kcal mol⁻¹ for HOOFePc-N=FePpc in pyridine).

Considering the uncertainty in DFT calculations, the results obtained from DFT calculation propose that both oxo isomers (O=Fe(Pc)-N-Fe(Ppc) and O=Fe(Ppc)-N-Fe(Pc)) could be produced. However, it is also important to consider that the formation of oxo isomers should be kinetically controlled. The complete reaction mechanisms for the production of high-valent iron-oxo species are outside the scope of this study.

CH₄ oxidation activity

We adopted the method developed by Sorokin et al. to evaluate the catalytic CH₄ oxidation activity of 4⁺·I₃⁻.²² H₂O was used as the solvent for CH4 oxidation because it is sufficiently stable against oxidation by the high-valent ironoxo species. For this purpose, we first prepared a silica gelsupported catalyst because $4^+ \cdot I_3^-$ is not soluble in H_2O . The CH₄ oxidation reaction was performed in an acidic aqueous solution containing 189 mM H₂O₂ and 51 mM trifluoroacetic acid (TFA) at 60 °C under a CH₄ atmosphere of 1.0 MPa. After the oxidation reaction, MeOH, HCHO, and HCOOH were observed and quantified by GC-MS spectroscopy. The time dependence of each oxidized product is shown in Fig. 8a and Table S7.† The amount of oxidized products was significantly larger than that observed in the absence of CH₄, suggesting that the observed MeOH, HCHO, and HCOOH were mostly derived from CH4. The small amount of oxidized products observed in the absence of CH₄ (under N₂ atmosphere) was mainly derived from the organic solvents adsorbed on the SiO₂ surface of the catalysts. ^{21,28,31,32,43}

To appropriately evaluate the catalytic CH₄ oxidation activity, the effective total turnover number (TTNeff) and effective methane conversion number (MCNeff) were defined according to eqn (i)-(iv), based on the idea that CH₄ is oxidized in a stepwise manner, as shown in Fig. 8b.

$$TTN_{eff} = TTN_{(CH_4)} - TTN_{(N_2)}$$
 (i)

$$TTN_{(CH_4)}$$
 or $TTN_{(N_2)} = (C_{MeOH} + 2 \times C_{HCHO} + 3 \times C_{HCOOH})/C_{Cat}$ (ii)

$$MCN_{eff} = MCN_{(CH_A)} - MCN_{(N_2)}$$
 (iii)

$$MCN_{(CH_4)}$$
 or $MCN_{(N_2)} = (C_{MeOH} + C_{HCHO} + C_{HCOOH})/C_{Cat}$ (iv)

Fig. 9a shows the time dependence of TTN_{eff}. Although TTN_{eff} increased almost linearly at the initial stage, the rate of increase increased, particularly after 32 h of oxidation. It should also be mentioned that the bluish-green color of the catalyst apparently became thinner, especially after 32 h, and

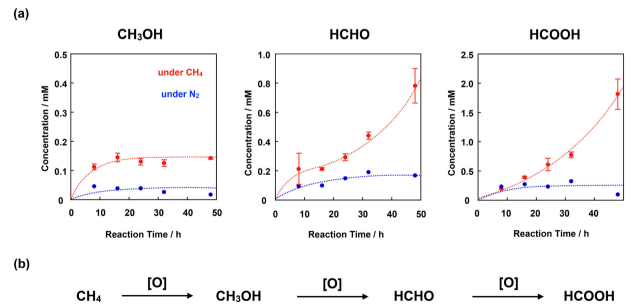


Fig. 8 (a) Time dependence of the concentrations of each oxidized product (MeOH, HCHO, and HCOOH) observed in the reaction under a CH₄ atmosphere of 1.0 MPa (red filled circle) or in the absence of CH₄ (under 1.0 MPa of N₂, blue filled circle) in an aqueous solution (3.0 mL) containing 4+I₃-/SiO₂ (55 µM as 4+I₃-), H₂O₂ (189 mM), and TFA (51 mM) at 60 °C. Error bars indicate standard deviations of three independent oxidation reactions. (b) Stepwise CH₄ oxidation reaction.

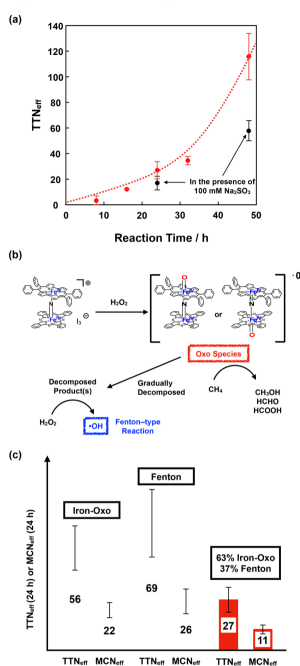


Fig. 9 (a) Time dependence of TTN_{eff} for CH₄ oxidation by $4^+ \cdot l_3^-$ in the absence (red circle) and presence (black circle) of 100 mM Na₂SO₃. Error bars indicate standard deviations of three independent oxidation reactions. (b) Proposed reaction mechanism for the CH₄ oxidation by $4^+ \cdot l_3^-$. (c) Comparison of TTN_{eff} and MCN_{eff} for the CH₄ oxidation for 24 h by $1^+ \cdot l_7^-$, $3^+ \cdot PF_6^-$, and $4^+ \cdot l_3^-$.

1+·I-/SiO2

3+-PF₆-/SiO₂

almost colorless after 48 h of oxidation (Fig. S5 \dagger), suggesting that the porphyrinoid core of $4^{+}\cdot I_{3}^{-}$ gradually decomposed during the reaction. A similar bleaching of the color of the catalyst was observed in the oxidation reaction using the porphycene dimer $3^{+}\cdot PF_{6}^{-}$. However, the bleaching speed was much slower for $4^{+}\cdot I_{3}^{-}$, implying that the stability of $4^{+}\cdot I_{3}^{-}$ is higher than that of $3^{+}\cdot PF_{6}^{-}$ under these reaction conditions.

Considering that apparent bleaching was not observed in the case of the phthalocyanine dimer $\mathbf{1}^+ \cdot \mathbf{I}^-$, the higher stability of $\mathbf{4}^+ \cdot \mathbf{I}_3^-$ than $\mathbf{3}^+ \cdot \mathrm{PF}_6^-$ might be because the phthalocyanine unit in $\mathbf{4}^+ \cdot \mathbf{I}_3^-$ contributed to the stable generation of the high-valent iron-oxo species after the reaction with $\mathrm{H_2O_2}$.

To further investigate the reaction mechanism of CH₄ oxidation by 4+13, we performed the CH4 oxidation in the presence of Na₂SO₃, a radical scavenger (Fig. 9a and Table S7,† entries 6 and 7).44 It was demonstrated that the reaction after 24 h of oxidation was suppressed by approximately 63%, whereas the reaction after 48 h of oxidation was suppressed to 50%. This indicates that the oxidation reaction mainly proceeded through the high-valent iron-oxo species, at least in its initial stage. However, considering that the color of 4+I₃ gradually bleached, particularly after 32 h, oxidation by the Fenton-type reaction via 'OH gradually increased, presumably because the decomposed product acted as the catalyst for the Fenton-type reaction, as summarized in Fig. 9b. Because 'OH is so reactive that it can even decompose the porphyrinoid cores of 4+I₃, the accelerated decomposition of the catalyst is not only because of the intrinsic instability of the catalyst, but also because of the decomposition by 'OH.

Fig. 9c shows a comparison of TTN_{eff} and MCN_{eff} of 1⁺·I⁻, 3⁺·PF₆, and 4⁺·I₃⁻ after 24 h of oxidation under the same reaction conditions. The reason why $\mathbf{4}^+ \cdot \mathbf{I_3}^-$ showed less CH_4 oxidation activity than the porphycene dimer 3⁺·PF₆⁻ is that 3⁺·PF₆⁻ decomposed more rapidly and CH₄ oxidation for 24 h mostly proceeded via a Fenton-type reaction. However, both TTN_{eff} and MCN_{eff} for oxidation by 4⁺·I₃⁻ were much less than in the case of oxidation by 1+.I-, which mostly proceeded via the high-valent iron-oxo species. Therefore, it can be concluded that the catalytic methane oxidation activity via the high-valent iron-oxo species of 4⁺·I₃⁻ (O=FePpc-N=FePc and/or O=FePc-N=FePpc) is lower than that of the high-valent iron-oxo species of 1'·I (O=FePc-N=FePc). This result indicates that the reactivity of the μ-nitrido-bridged iron porphyrinoid dimer strongly reflects the difference in the porphyrinoid skeleton, although the structures of the Fe-N=Fe cores are quite similar. We consider that one possible explanation for the lower catalytic activity of $\mathbf{4}^+ \cdot \mathbf{I}_3^-$ than $\mathbf{1}^+ \cdot \mathbf{I}^-$ could be the slow generation of the high-valent iron-oxo species of $4^+ \cdot I_3^-$ than that of $1^+ \cdot I^$ because of the lower efficiency of the push effect, which facilitates the heterolytic O-O bond cleavage of the hydroperoxo species (Fe-O-OH) owing to the lower electrondonating ability of the porphyrinoid ring of 4⁺, as can be estimated from the fact that the oxidation potential of the phthalocyanine ring of $4^+ \cdot I_3^-$ is higher than that of $1^+ \cdot I_3^-$.

Conclusions

In this study, we synthesized a monocationic μ -nitridobridged heterodimer of an iron phthalocyanine and an iron porphycene $(\mathbf{4}^+\cdot\mathbf{I_3}^-)$ and investigated its electrochemical, spectroscopic, and catalytic properties. Single-crystal X-ray

structural analysis demonstrated that the Fe-N=Fe core structure of $4^+\cdot I_3^-$ is similar to that of a monocationic μ -nitrido-bridged iron phthalocyanine dimer (1⁺·I⁻) and a monocationic μ-nitrido-bridged iron porphycene dimer (3⁺·PF₆⁻). ESI-TOF MS study indicated that treatment of 4⁺·I₃⁻ with H₂O₂ gave the high-valent iron-oxo species of 4⁺. DFT calculations suggested that the high-valent iron-oxo species of $4^+ \cdot I_3^-$ could be generated on both iron-porphycene and iron-phthalocyanine rings through the reaction with H₂O₂. The catalytic CH₄ oxidation reaction by 4⁺·I₃⁻ in an acidic aqueous solution proceeded in the presence of excess H₂O₂ to afford a mixture of MeOH, HCHO, and HCOOH through the high-valent iron-oxo species at its initial stage as in the case with $\mathbf{1}^+\cdot\mathbf{I}^-$. It was demonstrated that the catalytic activity of the high-valent iron-oxo species of 1+I was higher than that of 4⁺·I₃⁻. However, as the reaction time was elongated, it was found that 4⁺·I₃⁻ was gradually decomposed. This implies that the high-valent iron-oxo species of 4+13 was less stable than that of 1⁺·I⁻. Considering that the high-valent iron-oxo species of 3+PF₆ is much less stable and readily decomposes under the same reaction conditions, the lower stability of the high-valent iron-oxo species of 4⁺·I₃⁻ compared to that of $\mathbf{1}^+\cdot\mathbf{I}^-$ is derived from the difference in the porphyrinoid ring. Thus, it was confirmed that the difference in the porphyrinoid ring of the monocationic μ-nitrido-bridged iron porphyrinoid dimer strongly affected the stability of the highvalent iron-oxo species as well as the catalytic activity of the monocationic μ-nitrido-bridged iron porphyrinoid dimer. We believe that these findings would provide some valuable insights for the future development of molecular CH4 oxidation catalysts based on μ-nitrido-bridged porphyrinoid dimers.

Experimental

General

All reagents and solvents were purchased at the highest commercial quality available and used without further purification, unless otherwise stated. A metal-free porphycene was synthesized according to the reported procedure. 45 1H NMR spectra were recorded on a JEOL JNM-ECS400 (400 MHz for ¹H) spectrometer at a constant temperature of 298 K. Elemental analysis was performed on a Yanaco MT-6 analyzer. The absorption spectrum was recorded with a Hitachi U-4100 spectrophotometer in pyridine solutions at 20 \pm 0.1 °C in 1.0 cm quartz cells. MALDI-TOF MS was performed on Bruker Daltonics ultrafleXtreme using α-CHCA as a matrix. MALDI-TOF MS was performed on Bruker Daltonics compact.

Synthesis of $4^+ \cdot I_3^-$

A mixture of 5 (201 mg, 0.30 mmol), 6 (85 mg, 0.15 mmol), NaN₃ (300 mg, 4.6 mmol) and 1-chloronaphtalene (7.5 mL) was heated under air at 280 °C for 2 h. After cooling to room temperature, the reaction mixture was diluted with hexane (200 mL), the precipitate was collected by suction filtration,

washed with hexane (400 mL) H₂O (300 mL), successively, dried under vacuum to give blackish green crude solid (392.3 mg). The crude product was dissolved in pyridine (400 mL). After iodine (70.2 mg, 0.55 mmol as I) was added, the resulting solution was stirred for 1 h at room temperature, followed by evaporation of the volatile compounds. The residue was washed with Et₂O (300 mL) to remove the remaining iodine, and then, dried under reduced pressure to give dark green solid. The crude was purified by silica gel column chromatography (4.5 cm $\phi \times 15.0$ cm, CH₂Cl₂/AcOEt 10/10, then $CH_2Cl_2/AcOEt/MeOH = <math>10/10/1$) and recrystallization from CH₂Cl₂/Et₂O to yield 4⁺·I₃⁻ as a bluish green solid (121 mg, 89% in 2 steps from a metal-free porphycene). ¹H-NMR (400 MHz, pyridine- d_5/TMS): $\delta = 9.73$ (dd, J = 5.4, 3.0 Hz, 8H), 8.84 (d, J = 4.8 Hz, 4H), 8.51 (dd, J =6.0, 2.8 Hz, 8H), 8.42 (d, J = 7.2 Hz, 4H), 8.19 (d, J = 4.8 Hz, 4H), 8.00 (t, J = 7.8 Hz, 4H), 7.59–7.62 (m, 4H), 7.22–7.26 (m, 4H), 7.01 (d, J = 7.6 Hz, 4H). MALDI-TOF MS: m/z = 1250.25: calcd for $C_{76}H_{44}Fe_2N_{13}$ ([M]⁺) found: 1250.26. Anal. calcd for $C_{164}H_{122}Fe_4I_6N_{26}O_5$ ((4⁺·I₃⁻)₂·3Et₂O·2H₂O): C; 55.93, H; 3.49, N; 10.34, found: C; 55.66, H; 3.32, N; 10.04 (0.30% error).

Synthesis of 4+PF6

 $4^+ \cdot I_3^-$ (61 mg, 38 µmol) was dissolved in 10 mL of acetonitrile. After KPF₆ (691 mg, 3.8 mmol) in CH₃CN (10 mL) was added to the solution, H2O (80 mL) was added. The resulting precipitate was collected by centrifugation, followed by decantation to yield dark green solid (54 mg). The crude product was further purified by PTLC ($CH_2Cl_2/MeOH = 50/1$) and recrystallization from CH2Cl2/Et2O to give the desired product 4⁺·PF₆⁻ as a bluish green solid (27.8 mg, 53%). ¹H-NMR (400 MHz, pyridine- d_5/TMS): $\delta = 9.72$ (dd, J = 5.6, 2.8 Hz, 8H), 8.84 (d, J = 4.8 Hz, 4H), 8.50 (dd, J = 5.6, 2.8 Hz, 8H), 8.42 (d, J = 7.6 Hz, 4H), 8.20 (d, J = 4.8 Hz, 4H), 8.00 (t, J = 7.2)Hz, 4H), 7.59-7.62 (m, 4H), 7.22-7.26 (m, 4H), 7.00 (d, J = 7.6Hz, 4H). MALDI-TOF MS: m/z = 1250.25: calcd for $C_{76}H_{44}Fe_2$ - N_{13} ([M]⁺) found: 1250.26. Anal. calcd for $C_{76}H_{44}F_{6}Fe_{2}N_{13}P$ (4⁺·PF₆⁻): C; 65.39, H; 3.18, N; 13.04, found: C; 65.45, H; 3.49, N; 13.14 (0.31% error).

Single-crystal X-ray structural analysis of 4⁺·I₃⁻

A 4+·I₃ crystal suitable for single-crystal X-ray structural analysis was obtained by vapor diffusion of Et₂O into a 4⁺·I₃⁻ solution in a 1:1 (v/v) mixture of pyridine and CHCl₃. Singlecrystal X-ray diffraction measurements were performed using a Rigaku X-ray diffractometer equipped with a molybdenum MicroMax-007 and Saturn 70 CCD detector. measurement was performed at 123 K. The structure was solved via the direct method (SHELXT) and refined via fullmatrix least-squares on F_2 (SHELX-2018) using Olex2-1.3 program. All non-hydrogen atoms were refined anisotropically. Geometrical restraints were applied: DFIX, SADI, SIMU, ISOR, and OMIT. The crystal data are as follows: formula $C_{74}H_{49.6}Cl_{15.6}Fe_{1.6}I_{2.4}N_{12}$, FW = 2053.882, crystal size $0.19 \times 0.20 \times 0.22 \text{ mm}^3$, triclinic, space group $P\bar{1}$, a =

20.6469(2) Å, b = 23.6627(3) Å, c = 24.8965(3) Å, $\alpha = 64.380(1)^{\circ}$, $\beta = 69.270(1)^{\circ}$, $\gamma = 68.880(1)^{\circ}$, V = 9942.9(2) Å³, Z = 5, $R_1 = 0.1207$ (I > 2(I)), $wR_2 = 0.3227$ (all), GOF = 1.110. CCDC identification code 2217367. All the checkCIF Level-B alerts are due to the low crystal quality.

Cyclic voltammogram of 4+·PF₆

Cyclic voltammograms were measured with a BAS Electrochemical Analyzer Model 750Ds at room temperature in pyridine solutions containing 100 mM TBAPF₆ in a standard one-component cell under an N_2 atmosphere equipped with a 3 mm-O.D. glassy carbon disk working electrode, platinum wire counter electrode, and Ag/AgCl reference electrode. All solutions were deoxygenated by N_2 bubbling for at least 20 min. Obtained $E^{o\prime}$ νs . Ag/AgCl were converted to those νs . Fc/Fc⁺ based on measured redox potential of ferrocene. Tetrabutylammonium hexafluorophosphate (TBA⁺PF₆⁻) was recrystallized from 95% EtOH and dried under vacuum overnight at 100 °C.

ESI-TOF MS measurement of $4^+ \cdot I_3^-$ in CH₃CN in the presence of H₂O₂

To a solution of $4^+ \cdot I_3^-$ in CH₃CN (1.0 μ M, 2.0 mL) was added 35% aqueous H₂O₂ (0.1 mL, 1.16 mmol). After addition, the resulting mixture was subjected to ESI-TOF MS measurement immediately.

DFT calculation

DFT calculations were done with the BP86-D3(BJ)/def2-TZVP and B3LYP-D3(BJ)/def2-TVZP levels of theory implemented in the Turbomole software package. The former functional was chosen based on the studies of Sorokin *et al.* Only BP86 results were discussed, whereas the results calculated with B3LYP can be found in the ESI.† Frequency calculations were done to confirm the character of the optimized structures and to obtain Gibbs free energies. In some calculations, the solvent effect (pyridine) was determined using COSMO with a dielectric constant of 12.5. 47

Preparation of silica-supported catalyst (4⁺·I₃⁻/SiO₂)

 $4^+ \cdot I_3^-$ (10.00 mg, 6.2 μmol) was dissolved in 50 mL of mixture of pyridine and CHCl₃ (1:1 (v/v)). After the addition of silica gel (1082 mg) to the solution, solvent was evaporated. CHCl₃ (10 mL) was added to the residue. After solvent was evaporated, the residue was dried under vacuum at 60 °C over night. The catalyst was suspended in an aqueous TFA (TFA 10.0 mL + H₂O 100 mL). The mixture was sonicated for 30 min and then the solid was filtered. This TFA wash procedure was repeated twice. Finally, the resulting solid was washed with H₂O (300 mL) and dried under vacuum at 60 °C over night. The silica-supported catalyst was obtained quantitatively.

CH₄ oxidation reactions

Heterogeneous CH₄ oxidation was performed in a stainless-steel autoclave with a glass tube. A mixture containing $4^+ \cdot I_3^- / SiO_2$ (30 mg, 55 µM as $4^+ \cdot I_3^- / 35\%$ aqueous H₂O₂ (50 µL, 189 mM), and TFA (12 µL, 51 mM) in H₂O (3.0 mL) was heated at 60 °C under 1.0 MPa of CH₄ for 8–48 h with continuous stirring (900 rpm). After the autoclave was opened, the reaction mixture was filtrated through a disposable membrane filter. The filtrate was analyzed by GC-MS (system: Agilent 7890A equipped with JEOL JMS-T100GCV, detection: EI, column: Agilent DB-WAX UI, external standard: isovaleric acid (5 mM), temperature conditions: initial: 70 °C to 220 °C (10 °C min⁻¹) – hold (5 min)). The yields of CH₃OH and formic acid were determined based on the results of GC-MS. The yield of formaldehyde was examined using the method reported in our previous paper. ^{21,28,31,42}

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 M. H. Sazinsky and S. J. Lippard, Acc. Chem. Res., 2006, 39, 558–566.
- 2 S. Sirajuddin and A. C. Rosenzweig, *Biochemistry*, 2015, 54, 2283–2294.
- 3 V. C.-C. Wang, S. Maji, P. P.-Y. Chen, H. K. Lee, S. S.-F. Yu and S. I. Chan, *Chem. Rev.*, 2017, 117, 8574–8621.
- 4 C. E. Tinberg and S. J. Lippard, Acc. Chem. Res., 2011, 44(4), 280–288.
- 5 P. C. A. Bruijnincx and B. M. Weckhuysen, *Angew. Chem., Int. Ed.*, 2013, 52, 11980–11987.
- 6 Z. Guo, B. Liu, Q. Zhang, W. Deng, Y. Wang and Y. Yang, Chem. Soc. Rev., 2014, 43, 3480–3524.
- 7 S. J. Blanksby and G. B. Ellison, Acc. Chem. Res., 2003, 36, 255–263.
- 8 M. Ravi, M. Ranocchiari and J. A. van Bokhoven, *Angew. Chem., Int. Ed.*, 2017, **56**, 16464–16483.
- 9 F. T. de Oliveira, A. Chanda, D. Banerjee, X. Shan, S. Mondal, L. Que Jr., E. L. Bominaar, E. Münck and T. J. Collins, *Science*, 2007, 315, 835–838.
- 10 G. Xue, R. D. Hont, E. Münck and L. Que Jr., *Nat. Chem.*, 2010, 2, 400-405.

- 11 Y. Hitomi, K. Arakawa and M. Kodera, *Chem. Commun.*, 2014, **50**, 7485–7487.
- 12 E. Y. Tshuva and S. J. Lippard, *Chem. Rev.*, 2004, **104**, 987–1012.
- 13 X. Shan and L. Que Jr., *J. Inorg. Biochem.*, 2006, **100**, 421-433.
- 14 W. Nam, Acc. Chem. Res., 2007, 40, 522-531.
- 15 A. R. McDonald and L. Que Jr., Coord. Chem. Rev., 2013, 257, 414–428.
- 16 W. Nam, Acc. Chem. Res., 2015, 48, 2415-2423.
- 17 M. Guo, T. Corona, K. Ray and W. Nam, ACS Cent. Sci., 2019, 5, 13–28.
- 18 J. E. M. N. Klein and G. Knizia, *Angew. Chem.*, *Int. Ed.*, 2018, 57, 11913–11917.
- 19 D. R. Weinberg, C. J. Gagliardi, J. F. Hull, C. F. Murphy, C. A. Kent, B. C. Westlake, A. Paul, D. H. Ess, D. G. McCafferty and T. J. Meyer, *Chem. Rev.*, 2012, 112, 4016–4093.
- 20 P. Afanasiev and A. B. Sorokin, Acc. Chem. Res., 2016, 49, 583–593.
- 21 Y. Yamada, J. Kura, Y. Toyoda and K. Tanaka, *Dalton Trans.*, 2021, **50**, 6718–6724.
- 22 A. B. Sorokin, E. V. Kudrik and D. Bouchu, *Chem. Commun.*, 2008, 2562–2564.
- 23 E. V. Kudrik, P. Afanasiev, L. X. Alvarez, P. Dubourdeaux, M. Clémancey, J.-M. Latour, G. Blondin, D. Bouchu, F. Albrieux, S. E. Nefedov and A. B. Sorokin, *Nat. Chem.*, 2012, 4, 1024–1029.
- 24 C. Colomban, E. V. Kudrik, V. Briois, J. C. Shwarbrick, A. B. Sorokin and P. Afanasiev, *Inorg. Chem.*, 2014, 53, 11517–11530.
- 25 M. G. Quesne, D. Senthilnathan, D. Singh, D. Kumar, P. Maldivi, A. B. Sorokin and S. P. de Visser, ACS Catal., 2016, 6, 2230–2243.
- 26 M. Ansari, N. Vyas, A. Ansari and G. Rajaraman, *Dalton Trans.*, 2015, 44, 15232–15243.
- 27 Q. M. Phung and K. Pierloot, Chem. Eur. J., 2019, 25, 12491–12496.
- 28 Y. Yamada, Y. Miwa, Y. Toyoda, T. Yamaguchi, S. Akine and K. Tanaka, *Dalton Trans.*, 2021, **50**, 16775–16781.
- 29 G. Anguera and D. Sánchez-García, Chem. Rev., 2017, 117, 2481–2516.
- 30 J. Waluk, Chem. Rev., 2017, 117, 2447-2480.
- 31 Y. Yamada, K. Morita, N. Mihara, K. Igawa, K. Tomooka and K. Tanaka, *New J. Chem.*, 2019, **43**, 11477–11482.

- 32 N. Mihara, Y. Yamada, H. Takaya, Y. Kitagawa, K. Igawa, K. Tomooka, H. Fujii and K. Tanaka, *Chem. Eur. J.*, 2019, 25, 3369–3375.
- 33 Ü. Íşci, A. S. Faponle, P. Afanasiev, F. Albrieux, V. Briois, V. Ahsen, F. Dumoulin, A. B. Sorokin and S. P. de Visser, *Chem. Sci.*, 2015, 6, 5063–5075.
- 34 K. M. Kadish, R. K. Rhodes, L. A. Bottomley and H. M. Goff, Inorg. Chem., 1981, 20, 3195–3200.
- 35 L. A. Bottomley and B. B. Garrett, *Inorg. Chem.*, 1982, 21, 1260–1263.
- 36 M. Li, M. Shang, N. Ehlinger, C. E. Schulz and W. R. Scheidt, *Inorg. Chem.*, 2000, 39, 580–583.
- 37 Y. Yamada, T. Sugiura, K. Morita, H. Ariga-Miwa and K. Tanaka, *Inorg. Chim. Acta*, 2019, **489**, 160–163.
- 38 T. Shimizu, K. Wakamatsu, Y. Yamada, Y. Toyoda, S. Akine, K. Yoza and H. Yoshikawa, ACS Appl. Mater. Interfaces, 2021, 13, 40612–40617.
- 39 L. A. Bottomley, J.-N. Gorce, V. L. Goedken and C. Ercolani, *Inorg. Chem.*, 1985, 24, 3733–3737.
- 40 C. Bernard, J. P. Gisselbrecht, M. Gross, E. Vogel and M. Lausmann, *Inorg. Chem.*, 1994, 33, 2393–2401.
- 41 C. Colomban, E. V. Kudrik and A. B. Sorokin, *J. Porphyrins Phthalocyanines*, 2017, 21, 346–353.
- 42 A. Ghosh, E. Tangen, E. Gonzalez and L. Que, *Angew. Chem.*, 2004, **116**, 852–856.
- 43 Y. Yamada, J. Kura, Y. Toyoda and K. Tanaka, *New J. Chem.*, 2020, 44, 19179–19183.
- 44 C. Hammond, M. M. Forde, M. H. Ab Rahim, A. Thetford, Q. He, R. L. Jenkins, N. Dimitratos, J. A. Lopez-Sanchez, N. F. Dummer, D. M. Murphy, A. F. Carley, S. H. Taylor, D. J. Willock, E. E. Stangland, J. Kang, H. Hagen, C. J. Kiely and G. J. Hutchings, *Angew. Chem., Int. Ed.*, 2012, 51, 5129–5133.
- 45 T. Ono, N. Xu, D. Koga, T. Ideo, M. Sugimoto and Y. Hisaeda, *RSC Adv.*, 2018, **8**, 39269–39273.
- 46 S. G. Balasubramani, G. P. Chen, S. Coriani, M. Diedenhofen, M. S. Frank, Y. J. Franzke, F. Furche, R. Grotjahn, M. E. Harding, C. Hättig, A. Hellweg, B. Helmich-Paris, C. Holzer, U. Huniar, M. Kaupp, A. Marefat Khah, S. Karbalaei Khani, T. Müller, F. Mack, B. D. Nguyen, S. M. Parker, E. Perlt, D. Rappoport, K. Reiter, S. Roy, M. Rückert, G. Schmitz, M. Sierka, E. Tapavicza, D. P. Tew, C. van Wüllen, V. K. Voora, F. Weigend, A. Wodyński and J. M. Yu, J. Chem. Phys., 2020, 152, 184107.
- 47 A. Klamt and G. Schüürmann, J. Chem. Soc., Perkin Trans. 2, 1993, 799–805.