Ferryl and Ferrate Species: Mössbauer Spectroscopy Investigation

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INTRODUCTION

IRON is an earth abundant element that occurs in +2 and +3 oxidation states, Fe(II) and Fe(III), in the natural environment. However, more electron-deficient iron with oxidation states of +4, +5, and +6 have been invoked in biological, industrial, and waste treatment reactions.[1-4] For example, the activation of dioxygen to oxidize unactivated C–H bond of the substrates involved oxoiron(IV) (FeO4=O) and oxoiron(V) (FeO5=O), generally called ferryl species.[5-9] Figure 1 represents characterization of the three kinds of high-valent iron-oxo active sites in haem and non-haem enzymes. In the first kind, haem-comprising enzymes contain mononuclear iron-protoporphin IX active sites.[10] The second and the third kinds of enzymes involve mononuclear and diiron centers. The Fe(II) centers in the enzymes can perform different two-electron oxidation reactions. These high-valent iron species may also be participating in other reactions including epoxidation, decarboxylation, and cyclization.[6,10]

In addition to iron oxo species, high-valent iron nitride complexes have also been suggested to play significant role in nitrogenase enzyme reactions and industrial Haber-Bosch process.[11-13] Figure 2 depicts the postulated intermediates with iron-nitrogen multiple bonds.[14] At the single iron site, reduction of nitrogen may occur. Importantly, binding of nitrogen and also the cleavage happen at this iron site, which ultimately lead to the generation of ammonia and the formation of multiple iron-nitrogen bonds (Figure 2). Similar imidioiron(IV) porphyrin species as intermediates have also been suggested in enzyme-catalyzed nitrogen-atom transfer reactions.[14]

Because of implication of high-valent iron species in biological transformation, many model iron(IV) and iron(V) complexes have been synthesized to comprehend mechanisms of the catalytic cycles of enzymatic reactions. Most of these complexes have organic ligands around the iron center. Investigation of such high-valent iron species have been carried out in non-aqueous environment. Because of
their instability at room temperature, studies have been restricted to low temperatures.\cite{1,10}

Other high-valent iron compounds are tetraoxyanions, which are commonly called ferrates (Fe$^{\text{VI}}$(O$_4$$^{2-}$), Fe(VI), Fe$^{\text{V}}$(O$_4$$^{3-}$), Fe(V), and Fe$^{\text{IV}}$(O$_4$$^{4-}$), Fe(IV)). Ferrates are soluble in water and have a broad portfolio of applications in energy materials, green organic synthesis, and treating merging contaminants and toxins in water.\cite{15–19} Fe(VI) can perform multimodal actions in water treatment which include oxidation, disinfection, and coagulation. The benign nature of ferrates has advantages over other conventional disinfectants (e.g. chlorine, chloramine, and chlorine dioxide).\cite{20}

This paper presents some examples on the application of Mössbauer spectroscopy to characterize ferryl and ferrate species. A progress on their identification as intermediates of oxidative reactions is also given.

**IRON-OXO COMPLEXES**

A series of high-valent oxoiron(IV) and oxoiron(V) have been synthetized in the last three decades.\cite{6,9} Complexes of oxoiron(IV) containing haem and non-haem ligands have spin (S) of 1 and 2. $^{57}$Fe Mössbauer spectroscopy was utilized to probe locally the iron center. The isomer shifts ($\delta$), which relates to oxidation state (or electron density) of the iron nucleus of haem and non-haem oxoiron species, are shown in Figure 3. The quadrupole splitting ($\Delta E_Q$) is determined by the electric field gradient at the iron center. Information on molecular geometry and electronic ground state can be obtained from the values of $\Delta E_Q$. Other parameter

**Figure 1** The structures in the boxes depict high-valent intermediates of the enzymatic reactions; TauD-J: intermediate J of taurine dioxygenase, Q: intermediate Q of soluble methane monoxygenase, Cpd-I: Compound-I intermediates of iron containing haem enzymes, like catalase, peroxidase or cytochrome-P450; His: histidine; Hisdist: distal histidine; Asp: asparagine; Glu: glutamic acid. Adapted from Hohenberger et al.\cite{1} with the permission of Macmillan Publishers Limited.

**Figure 2** The proposed structures have inspired studies on model iron–nitrido and imido complexes. (a) Structure of the iron(IV) imido porphyrin intermediate that has been invoked for cytochrome-P450 catalysed nitrogen group transfer reactions; (b) The iron molybdenum cofactor of the nitrogenase enzyme with a central carbide, (C) and the proposed intermediates (scheme in the box) of the dinitrogen activation process occurring at a single iron site. Adapted from Hohenberger et al.\cite{1} with the permission of Macmillan Publishers Limited.
of the Mössbauer spectrum is nuclear hyperfine tensor (A), which is helpful in understanding the electronic structure of paramagnetic species. Values of δ, collected under zero-field for both haem and non-haem, belong to iron(IV) oxidation state (Figure 3). However, collection of Mössbauer spectra under applied magnetic field revealed different spin states of iron(IV) complexes. In the haem system values of ΔE_Q were positive (0.90–2.06 mm s⁻¹) whereas non-haem systems had some negative values of ΔE_Q (−0.22–1.53 mm s⁻¹). The difference between S = 1 and S = 2 of oxoiron(IV) complexes was elucidated from their A tensor parameters. Haem enzymes with S = 1 had “two large negative values” and “one small negative value”. Comparatively, non-haem systems having S = 2 had “three large negative” A tensors.[1] This S = 2 configuration may correspond to weak ligand field on the iron center.[2]

The oxoiron(IV) (d⁵) complex was obtained by oxidizing Fe³⁺ complex ([TAML]Fe³⁺(H₂O))²⁻, TAML = tetraamido macrocyclic ligand with meta-chloroperbenzoic acid in n-butyronitrile at −60 °C.[22] The low negative isomer shift of −0.42 mm s⁻¹ was obtained (Figure 3). A much larger value of ΔE_Q = 4.24 mm s⁻¹ for oxoiron(V) species than that of oxoiron(IV) complexes was evaluated from the Mössbauer spectrum. The spin of S = 1/2 for oxoiron(V) was revealed by the electron paramagnetic resonance (EPR) spectrum.

**IRON-NITRIDO COMPLEXES**

Studies on the synthesis of iron-nitro compounds having valence states of +4, +5, and +6 have been reported.[1,11,13] Figure 4 shows the example of the scheme in which different high-valent species have been demonstrated. Details of the scheme has been given elsewhere.[13] Briefly, this scheme shows the photolysis of species 1 yielded compound 2 (Fe(V)-nitrido complex). When species 3 was reduced, the compound 4 having +4 oxidation state of iron (Fe(IV)-nitrido complex) was obtained. The photochemical reaction of compound 4 yielded species 5. The parameters of the Mössbauer spectrum of compound 5 were δ = 0.11 mm s⁻¹ and ΔE_Q = 1.92 mm s⁻¹; indicated +6 formal oxidation state (i.e. Fe(VI)-nitrido complex) (Figure 4). The assignment of +6 oxidation state to species 5 was further confirmed by correlating the isomer shifts to the oxidation state of octahedral complexes having similar ligands, but different electronic configurations ranging from d⁶ to d⁰. Almost linear relationship between isomer shifts and oxidation state of these complexes was observed confirming assignments of different iron valence states of species in the scheme (Figure 4).

**FERRATES**

Numerous alkali and alkaline earth salts of Fe(IV), Fe(V), and Fe(VI) have been prepared and were investigated using zero-field and in-field Mössbauer spectroscopy. Isomer shifts of different salts of Fe(VI) (e.g. Na₂FeO₄, K₂FeO₄, Rb₂FeO₄, Cs₂FeO₄, and BaFeO₄) varied little (−0.87 to
The isomer shift values, shown in Figure 5, were used in studying the decomposition of salts of Fe(IV), Fe(V), and Fe(VI) under humid and thermal conditions.[25,26] The encapsulation of arsenic and heavy metals by iron(III) oxide nanoparticles (γ-Fe$_2$O$_3$), generated from Fe(VI), was investigated in detail by in-field Mössbauer spectroscopy.[27-29] In the case of arsenic, the results clearly demonstrated that a significant portion of arsenic was embedded in the tetrahedral sites of the γ-Fe$_2$O$_3$ spinel structure.

Attempts have been made to identify intermediate(s) during the aqueous reactions of Fe(VI) with sulfur and nitrogen-containing compounds in order to distinguish one- and two-electron transfer reaction mechanisms.[30,31] No direct evidence of intermediate valence state, Fe(IV) and Fe(VI), could be seen due to the highly instable character of such species, which most likely self-decomposed within ~ 3 s, a lag time of rapid-freezing off reaction mixtures (~ 3 s). However, determination of final reduced iron species, Fe(II) or Fe(III) by Mössbauer spectroscopy was able to elucidate the mechanism of oxidation of thiols (X) by Fe(VI) and is briefly described below.

The experimental stoichiometry of the oxidation of methionine (CH$_3$S(CH$_2$)$_2$CH(NH$_2$)COO$^-$ Met) and cysteine (HSCH$_2$CH(NH$_2$)$^+$COO$^-$, Cys) were described by Equations (2) and (3).[32]

\[
\text{HFeO}_4^- + \text{HSCH}_2\text{CH(NH}_2\text{)}^+\text{COO}^- + \text{H}_2\text{O} \rightarrow \text{Fe(OH)}_4^- + \text{HS(O)}_2\text{CH(\text{NH}_2\text{)}^+\text{COO}^- + \text{OH}^-}
\]

\[
2\text{HFeO}_4^- + 3\text{CH}_3\text{S(CH}_2\text{)}_2\text{CH(NH}_2\text{)}^+\text{COO}^- + 3\text{H}_2\text{O} \rightarrow 2\text{Fe(OH)}_4^- + 3\text{CH}_3\text{S(O)}_2\text{CH(\text{NH}_2\text{)}^+\text{COO}^- + 2\text{OH}^-}
\]

The Mössbauer spectra of the final frozen mixed solutions of Fe(VI) with either Cys or Met are shown in Figure 6.[41] In case of Cys, The spectrum had one singlet and three doublets (Figure 6a); the singlet component with isomer shift of ~0.82 mm s$^{-1}$ represents remaining non reacted Fe(VI), the doublet with isomer shift of 0.45 mm s$^{-1}$ and quadrupole splitting of 0.77 mm s$^{-1}$ is typical for high-spin Fe(III), and the two doublets with the same isomer shift of 0.90 mm s$^{-1}$ and a huge quadrupole splitting of 3.72 mm s$^{-1}$ and 3.00 mm s$^{-1}$ is high-spin Fe(II). Comparatively, Mössbauer spectrum of the mixed frozen solution of Fe(VI) and Met had well resolved singlet and doublet components (Figure 6b); the singlet component with isomer shift of ~0.81 mm s$^{-1}$ represents remaining non reacted Fe(VI) and the doublet with isomer shift of 0.45 mm s$^{-1}$ and quadrupole splitting of 0.76 mm s$^{-1}$ is usually for high-spin Fe(III). Importantly, no Fe(II) was seen in the oxidation of Met by Fe(VI). This indicates that Fe(II) was not involved as an intermediate during the oxidation of Met by Fe(VI).

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**Figure 5.** Relationships of isomer shifts with the Fe–O bond distances and electron density on the iron nucleus ($\rho_O$) of ferrates. Adapted from Sharma et al.[24] with the permission of Wiley Inc.

$\delta / \text{mm s}^{-1} = 1.084 - 0.326 \times \text{OS}$

Equation (1) allowed to estimate the isomer shifts of ferrate(VII) and ferrate(VIII) as $-1.18$ and $-1.40$ mm s$^{-1}$, relative to α-Fe, respectively.[23]

Significantly, density function theory calculations were carried out on ferrates with different valence states ranging from +4 to +8.[23] Figure 5 shows the relationship of isomer shift with electron density on the iron nucleus ($\rho_O$) and Fe–O bond length.[24] A decrease in Fe–O bond distance with increase in oxidation state of iron from Fe(IV) to Fe(VI) was observed. Values of isomer shifts had a negative relationship with $\rho_O$, whereas a positive linearity between isomer shift and bond distance was seen (Figure 5). The value of $\delta$ of Fe(VI) in ferrate is significantly lower than in nitride complex (~0.90 mm s$^{-1}$ versus 0.40 mm s$^{-1}$).[13,24] This is not surprising considering both species have different geometry and electronic structures. Ferrate(VI) has the tetrahedral structure while Fe(VI)-nitrido complex possess the octahedral coordination. Furthermore, ferrate(VI) anion has four strong and covalent Fe–O double bonds while in the nitride complex, only one strong Fe–N triple bond is present. A combination of these effects would influence the variation of isomer shifts in both ferrate and Fe(VI)-nitride complex species.
Overall, electron transfer steps of the oxidation of Cys by Fe(VI) may be described by Equations (4) and (5). The formation of Fe(II) as the reduced product of Fe(VI) with Cys satisfies the observation of Mössbauer spectrum of Figure 6a.

\[
\text{Fe(VI)} + \text{Cys} \rightarrow \text{Fe(V)} + \text{CysS(O)} \quad (4)
\]

\[
\text{Fe(IV)} + \text{Cys} \rightarrow \text{Fe(II)} + \text{Cys(O)} \quad (5)
\]

The electron-transfer steps of the oxidation of Met by Fe(VI) may be given as Equations (6)–(8). This reaction scheme is consistent with the observation of Mössbauer spectrum (i.e. Fe(III)) as a final reduced species of Fe(VI) without the formation of Fe(II), see Figure 6b).

\[
\text{Fe(VI)} + \text{Met} \rightarrow \text{Fe(V)} + \text{Met}^* \quad (6)
\]

\[
\text{Fe(VI)} + \text{Met}^* \rightarrow \text{Fe(V)} + \text{Met(O)} \quad (7)
\]

\[
2\text{Fe(V)} + 2\text{Met} \rightarrow 2\text{Fe(III)} + 2\text{Met(O)} \quad (8)
\]

Interestingly, the reaction schemes, given by Equations (4), (5) and Equations (6)–(8), are in agreement with the suggested initial electron transfer steps from the correlation of the rate constants with reduction potentials for the oxidation of Cys and Met by Fe(VI), respectively.\[32\]

### CONCLUSION

Mössbauer spectroscopy method was one of the main tools to determine the oxidation states of iron in ferryl, iron-nitride and ferrate species. Besides values of isomer shift (\(\delta\)), quadrupole splitting (\(\Delta E_Q\)) and nuclear hyperfine tensors (\(\mathbf{A}\)) were determined for the complexes, which enhanced knowledge on the electric field gradient at the iron center, molecular geometry and electronic ground state as well as the electronic structure of paramagnetic species. Additionally, Mössbauer spectroscopy technique was successful in studying the mechanism of the oxidation of organosulfur compounds by ferrate(VI), however, freezing of samples of oxidative reactions within three seconds did not allow to identify the reactive intermediate iron species, Fe(V) and Fe(IV). A rapid freezing of reaction samples on the time scale of milliseconds would be needed to observe the intermediate iron species to comprehend reaction mechanisms.

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