A Synthetic Model of Enzymatic [Fe₄S₄]–Alkyl Intermediates

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ABSTRACT: Although alkyl complexes of [Fe₄S₄] clusters have been invoked as intermediates in a number of enzymatic reactions, obtaining a detailed understanding of their reactivity patterns and electronic structures has been difficult owing to their transient nature. To address this challenge, we herein report the synthesis and characterization of a 3:1 site-differentiated [Fe₄S₄]⁺−alkyl cluster. Whereas [Fe₄S₄]⁺ clusters typically exhibit pairwise delocalized electronic structures in which each Fe has a formal valence of 2.5+, Mössbauer spectroscopic and computational studies suggest that the highly electron-releasing alkyl group partially localizes the charge distribution within the cubane, an effect that has not been previously observed in tetrahedrally coordinated [Fe₄S₄] clusters.

Iron–sulfur (Fe–S) proteins are found in all kingdoms of life and perform myriad functions in the cell.¹⁻⁴ Those that utilize [Fe₄S₄] cofactors are the most ubiquitous and have well-documented roles as electron-transfer and Lewis acid catalysts.⁵⁻¹² Recently, several classes of [Fe₄S₄] enzymes have been proposed to operate via organometallic intermediates. In particular, [Fe₄S₄]–alkyl species have been invoked in mechanisms of reductive dehydroxylation reactions in isoprenoid biosynthesis,¹³⁻¹⁴ reductive coupling of CO and CO₂ to higher-order hydrocarbons,¹⁵ and radical reactions by the >100 000 members of the radical S-adenosyl-l-methionine (SAM) superfamily of enzymes.¹⁶⁻¹⁸

Despite the emerging significance of [Fe₄S₄]–alkyl intermediates, little is known about how these species form, their spectroscopic features, or for their electronic structures. Moreover, no [Fe₄S₄]–alkyl species have been structurally characterized and their identification as intermediates (Chart 1) has relied on EPR/ENDOR,⁵⁻¹¹,E₁³⁻¹⁵ or DFT studies. And although reliable models for the electronic structures of [Fe₄S₄] clusters have been developed,¹⁷⁻¹⁹ these models were derived for clusters ligated by relatively weak-field, moderately donating ligands (e.g., cysteine thiolates). How binding of a strong-field, highly electron-releasing alkyl ligand perturbs the electronic structure of [Fe₄S₄] clusters—to what extent it induces valence localization, how (if at all) it affects Fe–S interactions, and how these effects impact the reactivity of the Fe–C bond—is unknown.

Synthetic chemistry will play an important role in answering these questions. Synthetic analogues of [Fe₄S₄]–alkyl intermediates would, for example, allow for the structures of intermediates to be elucidated (by linking their structures with their spectroscopic features) and for their electronic structures to be interrogated. These efforts are hampered by the lack of synthetic access to alkylated [Fe₄S₄] clusters. To date, only one alkylated Fe–S cluster has been structurally characterized—an [Fe₄S₄] cluster in which the alkyl group is derived from decamethylcobaltocene.²¹ We herein describe the synthesis of the first [Fe₄S₄]–alkyl cluster and investigations into how the alkyl ligand perturbs the electronic structure of the [Fe₄S₄] core.

We reasoned that synthetic [Fe₄S₄]–alkyl clusters could be accessed by reaction of a nucleophilic alkylating reagent with a 3:1 site-differentiated [Fe₄S₄]–halide cluster. This route requires that the clusters be stable and soluble in solvents compatible with alkylating reagents, and we therefore pursued clusters with low overall charge. Inspired by the chelating trithiolate architecture (LS₃) developed by Holm,²²,²³ and adapted by others,²⁴,²⁵ we prepared a structurally analogous ligand, L(N=PTol), commonly referred to as three iminophosphorane donors. Iminophosphoranes are strongly basic and, like thiolates, can serve as both σ- and π-donors due to the presence of two lone pairs on the nitrogen atom.²⁶ In addition, iminophosphoranes are neutral (allowing for compounds with low overall charge) and, hence, increased solubility in unreactive solvents), sterically demanding, and tunable at their P-substituents.

The synthesis of L(N=PTol)₃ (3) is shown in Scheme 1. Trianiline 2 was generated via Buchwald–Hartwig coupling...
between tribromide 1 and Ph3C=NH followed by hydrolysis. The iminophosphorane groups were then installed in a Kirsanov reaction using Tol3PCI, and excess Et3N.

The 3:1 site-differentiated cluster \((\{\text{L(N=P}^\text{Tol})_3\})\text{Fe}_4\text{S}_4\text{Cl}^2\) can be synthesized on a multigram scale in 65% yield by reaction of \(\text{L(N=P}^\text{Tol})_3\) with \([\text{PPh}_4]_2[\text{Fe}_4\text{S}_4\text{Cl}_4]\) and 3 equiv of \(\text{NaBPh}_4\) (Scheme 1). The molecular structure of 4 as determined by single-crystal X-ray diffraction (XRD) shows the anticipated 3:1 site differentiation with three Fe atoms bound by \(\text{L(N=P}^\text{Tol})_3\) and the apical Fe (Feapical) site bound by Cl (Figure 1a). The molecule has pseudo-C3 symmetry with one p-tolyl group of each iminophosphorane aligned with the pseudo-C3 axis, forming a protective cavity around the unique Fe site (Figure 1a). The Fe–S distances in 4 are similar to those observed in the 3:1 site-differentiated cluster \((\{\text{LS}_3\})\text{Fe}_4\text{S}_4\text{Cl}^2\).

Treatment of 4 with Et3Zn generates the \([\text{Fe}_4\text{S}_4]\)–alkyl complex \((\{\text{L(N=P}^\text{Tol})_3\})\text{Fe}_4\text{S}_4\text{Et}\) [BPh4] (5), which is isolated as brown solids in 85% yield. The structure of 5 (Figure 1b) was confirmed by single-crystal XRD and is similar to that of 4. The Feapical–C bond length of 5 (2.05 Å) is shorter than that in the \([\text{Fe}_4\text{S}_4]\)–decamethylcobaltocenyl cluster (2.12 Å)21 and comparable to that in a tris(thioether)-borate-ligated Fe2+–Me complex (2.03 Å).28 In THF solution, 5 slowly decomposes to unidentified products; further reactivity studies of 5 are underway.

The 57Fe Mössbauer spectrum of solid 4 at 90 K (Figure 2a) was simulated as three quadrupole doublets in a 2:1:1 ratio with identical isomer shifts of 0.47 mm/s (Figure 2a, Table 1). This simulation is in accordance with the canonical electronic structure of an \([\text{Fe}_4\text{S}_4]\)2+ cluster: an \(S = 0\) ground state arising from antiferromagnetic coupling of two \(S = 9/2\) \([\text{Fe}_4\text{S}_4]\)2+ rhombs, each of which consists of spin-aligned, high-spin Fe2.5+ ions engaged in a double-exchange interaction.29 Thus, we assign the doublet comprising 50% of the spectral area to the \([\text{Fe}_4\text{S}_4]\)2+ rhomb bound by two iminophosphorane donors and the second pair of doublets to the Cl-ligated Feapical center and its spin-aligned, iminophosphorane-ligated partner (Figure 2c). That identical isomer shifts are observed for all sites in this spectrum indicates that each Fe site possesses similar core charge density, and hence we can assign to each a formal oxidation state (FOS) of 2.5+. Moreover, the similarity between the isomer shifts of 4 and those reported for \([(\text{LS}_3)\text{Fe}_4\text{S}_4\text{Cl}^2]^{2-}\) (δ = 0.46 mm/s at 80 K)30 and for protein-bound \([\text{Fe}_4\text{S}_4]\)2+ clusters (δ ≈ 0.42 mm/s)31 underscores the utility of the L(N=P)3 ligand in modeling a trithiolate donor set.

Complexes 4 and 5 show similar ligand-derived resonances in their room-temperature 1H and 31P NMR spectra. Both exhibit C3 symmetry in solution as indicated by splitting of the diastereotopic Ar–CH2–Ar and Ar–CH2–CH3 resonances. Their 31P NMR resonances (at 102.1 and 96.0 ppm for 4 and 5, respectively) are shifted downfield from that of the free ligand 3 (-0.4 ppm), reflecting both the expected downfield shift upon binding a Lewis acidic metal center32,33 and the population of paramagnetic excited states as is commonly observed in \([\text{Fe}_4\text{S}_4]\)2+ clusters;34,35,38 their room-temperature solution magnetic moments (\(\mu_\text{eff} = 2.7\) and 2.8 μB, respectively) are consistent with this interpretation and typical of \([\text{Fe}_4\text{S}_4]\)2+ clusters.35,39–41 The 1H NMR signals corresponding to the –CH3 and –CH2– protons of the ethyl ligand in 5 are

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**Scheme 1. Synthesis of Iminophosphorane-Ligated \([\text{Fe}_4\text{S}_4]\) Clusters**

**Figure 1.** Thermal ellipsoid plots (50%) of (a) 4 and (b) 5. Hydrogen atoms, solvent molecules, and anions omitted for clarity. Fe (red), S (yellow), Cl (purple), N (blue), P (orange), C (gray).

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\^Conditions: (i) (a) Ph3C=NH (9 equiv), Pd2(dba)_3 (1.9 mol %), rac-BINAP (2.8 mol %), NaOBu (10.5 equiv), toluene, 80 °C; (b) 2 M HCl, THF; (c) excess NaOH, MeOH; (ii) Tol3PCI (3 equiv), Et3N (7 equiv), benzene, 80 °C; (iii) NaBPh4 (3 equiv), [PPh4]2[Fe4S4Cl4] (1.1 equiv), 1:1 THF/MeCN; (iv) Et3Zn (1.5 equiv), THF.
observed at -4.6 and 70.0 ppm, respectively, which also indicates the population of paramagnetic excited states.23,34−38

Clusters 4 and 5 exhibit reversible, one-electron reduction events in their cyclic voltammograms at -1.48 and -1.78 V vs 

Fe/Fc, respectively (see Supporting Information (SI)), reflecting the greater electron-donating ability of the ethyl ligand compared with that of the chloride.

As above, the $^{57}$Fe Mössbauer spectrum of solid 5 at 90 K (Figure 2b) was simulated with three quadrupole doublets in a 2:1:1 ratio. This approach produces two reasonable models (Table 1); our preferred model (vide infra) is shown in Figure 2bc, and alternatives are discussed in the SI. Both models feature a doublet comprising 50% of the total area with parameters that are nearly identical to those found for the iminophosphorane-bound Fe$^{2.5+}$ sites in 4 (Table 1); this doublet is therefore assigned to the analogous Fe$^{2.5+}$ sites in 5.

The quadrupole doublet with the lowest isomer shift is assigned to the alkylated Fe site because the stronger electron-donating ability (vide supra) and increased covalency of the ethyl group relative to L(N=PTol)$_3$ are expected to enhance the charge density at the $^{57}$Fe nucleus and drive the isomer shift of this site more negative.32,43 The major difference between these two models is in the magnitude of this effect ($\delta$ = 0.30 vs 0.39 mm/s) and whether the remaining iminophosphorane-bound Fe site, which is coupled to the alkylated site via a double-exchange interaction, possesses an isomer shift that is greater or less than that of the other ligand-bound pair ($\delta$ = 0.52 vs 0.44 mm/s, compared with 0.46 mm/s for the other ligand-bound pair).

In order to distinguish between these two models, we turned to broken-symmetry density functional theory (BS DFT) calculations (see SI for details). The isomer shifts calculated for 4 (Table 1) are in good agreement with the experimental values. To make the connection between $\delta$ and the DOS of the Fe sites, we further analyzed the BS determinants in terms of localized molecular orbitals (LMOs).44 For each site, the LMOs naturally partition into a set of five spin-up (or down) Fe 3d orbitals, plus an extra spin down (or up) 3d-derived orbital that is delocalized over a single additional Fe site. This picture corresponds to the canonical electronic structure of the [Fe$_4$S$_4$]$^{2+}$ cluster $^{18,19}$ in which the double-exchange interaction is mediated by the extra delocalized orbital (Figure 3). Through a population analysis, we characterized the tendency of the itinerant electron to localize on either of the two Fe sites engaging in double exchange and thereby assigned FOSs. In the case of 4, the itinerant electrons are fully delocalized, leading to FOS assignments of ~2.5+ for each site (Table 1 and Figure 3), consistent with the experimentally observed isomer shifts.

For 5, the calculated isomer shift of the alkylated site, 0.22 mm/s, is most consistent with the simulated value of 0.30 mm/s. Moreover, the calculation predicts that the isomer shift of the iminophosphorane-bound Fe that is coupled to the alkylated site via double exchange increases relative to those of the remaining Fe sites—precisely what is observed in the favored simulation presented in Table 1. While the low isomer shift of the alkylated site might be expected on the basis of the properties of the alkyl ligand (vide supra), the compensatory increase in the isomer shift of the spin-aligned Fe is unusual and suggests partial charge localization within this double-exchange-coupled pair. Indeed, the calculated FOS of 2.31+ for

| Table 1. Experimental (90 K) and Computed Mössbauer Parameters for 4 and 5 |
|-----------------------------|-----------------------------|-----------------------------|
| Doublet | Site | Simulation | Calculation | Favored simulation | Calculation | Dist favored simulation | Calculation | FOS$^a$ | FOS$^a$ |
| 1 | Fe=L | 0.47 | 0.48 | 0.45 | 2.50+ | 0.46 | 0.59 | 0.46 | 0.58 | 0.46 | 2.58+ |
| 2 | Fe=L$^{\delta}$ | 0.47 | 0.75 | 0.45 | 2.54+ | 0.50 | 1.09 | 0.44 | 0.93 | 0.32 | 2.31+ |
| 3 | Fe=X$^{\delta}$ | 0.47 | 1.06 | 0.50 | 2.54+ | 0.22 | 2.79+ |

$^a$FOSs are determined from a population analysis of the LMOs of each BS determinant. See SI for details. $^b$Sites coupled via double exchange.
this site in 5 suggests increased ferrous character, and this charge localization is coupled to an increase in the FOS of the alkylated site to 2.79+ (Table 1).

Physically, this partial charge localization can be understood in terms of localized ligand field effects about the Fe sites engaging in double-exchange delocalization. Assuming that double exchange occurs through a single orbital interaction, then two Fe sites with very similar ligand fields will share the itinerant electron equally, producing an effective valence of 2.5+. This appears to be the case, both experimentally and computationally, for all sites in 4 (Figure 3). The symmetry of the double-exchange interaction is removed by alkylation in 5 whereby the electron-releasing alkyl ligand raises the average energy of the local Fe 3d manifold (Figure 3). As a result, the itinerant electron will tend to localize on the site to which the alkylated Fe is coupled, as observed. Similar electronic desymmetrization may alternatively be induced by differences in coordination number between double-exchange-coupled Fe sites. Although a two-orbital model of double exchange is likely too simplistic, these arguments should hold in the case of a more complex multiorbital picture.

In conclusion, we have reported the synthesis and characterization of 3:1 site-differentiated [Fe₅S₅]⁺−Cl and [Fe₅S₅]⁺−Et clusters that are supported by a chelating triminophosphoramidate ligand. NMR and Mössbauer spectroscopic data indicate that, although both clusters have typical diamagnetic ground states, the [Fe₅S₅]⁺−Et cluster exhibits a polarized Fe−Fe double-exchange interaction, partially localizing ferric character at the alkylated Fe site and ferrous character at its spin-aligned partner. Based on these results, we anticipate that enzymatic [Fe₅S₅]−alkyl intermediates may exhibit partial or even complete localization of Fe³⁺ at their alkylated sites. Further investigations into the reactivity of [Fe₅S₅]⁺−alkyl clusters and efforts to access [Fe₅S₅]−alkyl clusters in other redox states and coordination numbers are currently underway in our laboratory.

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