Direct Mechanistic Evidence for a Nonheme Complex Reaction through a Multivariate XAS Analysis

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ABSTRACT: In this work, we propose a method to directly determine the mechanism of the reaction between the nonheme complex FeII(tris(2-pyridylmethyl)amine) ([FeII(TPA)(CH3CN)2]2+) and peracetic acid (AcOOH) in CH3CN, working at room temperature. A multivariate analysis is applied to the time-resolved coupled energy-dispersive X-ray absorption spectroscopy (EDXAS) reaction data, from which a set of spectral and concentration profiles for the reaction key species is derived. These “pure” extracted EDXAS spectra are then quantitatively characterized by full multiple scattering (MS) calculations. As a result, structural information for the elusive reaction intermediates [FeIII(TPA)(κ2-OOAc)]2+ and [FeIV(TPA)(O)(X)]+/2+ is obtained, and it is suggested that X = AcO− in opposition to X = CH3CN. The employed strategy is promising both for the spectroscopic characterization of reaction intermediates that are labile or silent to the conventional spectroscopic techniques, as well as for the mechanistic understanding of complex redox reactions involving organic substrates.

1. INTRODUCTION

The full understanding of a given reaction mechanism, defined as the sequence of elementary steps leading reactants to products, is vital for chemical knowledge. In fact, unveiling the identity, the concentration time evolution, and the structural properties of the reaction intermediates provides essential insight into the process and paves the way for its rational optimization. Innovative experimental and theoretical approaches are required to tackle the complexity of chemical systems dealt with by contemporary researchers and to acquire accurate information on how these transformations take place.

Nonheme iron complexes are a class of bioinspired catalysts that are gaining special interest for their capacity of oxidizing C–H and C=C bonds with high regio- and stereoselectivity.1–3 A special attention has been dedicated to the use of the environmentally friendly H2O2 oxidant in association with acetic acid, which is able to increase both catalytic activity and reaction selectivity.4 In a previous investigation, we employed time-resolved energy-dispersive X-ray absorption spectroscopy (EDXAS) to qualitatively identify the sequence of oxidation states during the reaction between the nonheme iron complex FeII(tris(2-pyridylmethyl)amine) ([FeII(TPA)(CH3CN)2]2+) and peroxyacetic acid (AcOOH) in CH3CN/AcOH (99.6:0.4 (v/v)) at 25 °C.5 Investigating this transformation at −40 °C, a seminal study showed that AcOOH oxidizes [FeII(TPA)(CH3CN)2]2+ to the relatively stable oxo-dimeric product [Fe2III(TPA)(μ-O)(μ-OAc)]3+.6 The complex [FeII(TPA)(O)(X)]+/2+ was studied through a combination
of electrospray ionization (ESI) mass spectrometry, UV-vis and Mössbauer spectroscopies, and an extended X-ray absorption fine structure (EXAFS) experiment, which however could not establish the identity of the sixth coordinating ligand X, maintaining X to be a molecule with a terminal oxygen or nitrogen atom bound to the Fe metal cation. In that same work, the authors advanced the hypothesis that the Fe(IV) species derived from an unobserved Fe(II)(TPA)-acyl peroxy complex. While the structures of the initial Fe(II) and final dimeric Fe(III) complex have been solved through X-ray crystallography some time ago, extensive spectroscopic studies have been performed to determine the true oxidation state and geometry of the reaction intermediate arising immediately after the initial Fe(II) species, but a definite answer has not yet been obtained. Talsi et al. measured new S = 1/2 electron paramagnetic resonance (EPR) signals at g = 2.71, 2.42, and 1.53 in the reaction of 40 mM [FeII(TPA)2(CH3CN)2]2+ in 1:1:1 CH3CN/CH2Cl2 with either H2O2/AcOH, peracetic acid or m-chloroperbenzoic acid at ~60 °C. On the basis of this observation, the authors claimed to have identified a putative FeIV(O)(OAc) species, which has been predicted to be the true oxidant in the reaction of [FeII(TPA)(CH3CN)2]2+ with C–H bond containing substrates. The same authors excluded the intermediate to be a low-spin acylperoxo complex of the kind [(L)FeIII(OOCR)]2+, because the EPR parameters of [(L)FeIII(OOCR)]2+ coordination complexes are sensitive to the identity of the R′ group, and the same intermediate was observed in all reactions. However, this assignment was based on the isolation of the intermediate in only 7% yield, and it was not supported by additional spectroscopic characterizations. Subsequently, Que and co-workers replaced TPA with the variant TPA* (where six −CH3 and three −OCH3 groups substitute hydrogen atoms in the aromatic moieties of the three pyridine ligands of TPA) to generate a g = 2.7 intermediate in 50% yield at ~40 °C. This strategy enabled the isolation and characterization of the intermediate as a low-spin acylperoxoiron(III) complex by combining UV–vis, EPR, resonance Raman, Mossbauer, and ESI-mass spectrometry data relative to the reactions of [FeII(TPA)(CH3CN)2]2+ with either H2O2/AcOH, AcOOH, or meta-chloroperbenzoic acid (mCPBA). Specifically, the intermediate generated from the reaction of [FeII(TPA*)(CH3CN)2]2+ with AcOOH yielded UV–vis and EPR spectra similar to those of the intermediate produced from the reactions of [FeII(TPA*)(CH3CN)2]2+ with other oxidants, but it was not characterized by any other experimental techniques. The Fe(III) species was found to evolve to the corresponding Fe(IV)(TPA*)(O) complex in the reaction of [FeII(TPA*)(CH3CN)2]2+ with H2O2/AcOH or AcOOH. The proposed mechanism for the reaction between [FeII(TPA)(CH3CN)2]2+ and AcOOH is presented in Figure 1.

Numerous spectroscopic techniques have been applied to follow fast chemical reactions with half–life times lower than seconds. Among them, X-ray absorption spectroscopy (XAS) is a unique and versatile tool that allows one to follow the variations in both the local electronic and structural configuration of a selected photoabsorbing atom. We recently used a coupled EDXAS/UV–vis approach to measure pseudo-first-order kinetic constants in a reaction involving a nonheme iron-oxo complex and a series of aromatic sulfides and benzyl alcohol, demonstrating the suitability of EDXAS to extract quantitative kinetic information for a bimolecular process on the millisecond to second time scale. Here, we show that it is possible to use a multivariate approach for the analysis of the EDXAS spectral data relative to the reaction of [FeII(TPA)(CH3CN)2]2+ and AcOOH occurring at room temperature. This procedure enables one to extract the X-ray absorption near edge structure (XANES) spectra belonging to the reaction key species, to assess their oxidation states and lifetimes, and to quantitatively shed light on their elusive structures. The decomposition is achieved through a multivariate curve resolution (MCR) method, a class of algorithms that has been applied extensively to the analysis of spectroscopic data coming from the monitoring of chemical reaction processes, such as UV–vis, fluorescence, nuclear magnetic resonance, circular dichroism, near-infrared (NIR), Fourier transform IR (FTIR), time-
resolved FTIR, and Raman. MCR techniques have been increasingly applied also to time- and space-resolved XANES with studies investigating doped V₂O₅ lithium batteries, ZnO Q-dot formation, degradation of chloride layered double hydroxide (LDH) upon heating, and a variety of catalytic systems in the solid phase. To the best of our knowledge, herein we report the first application of the MCR approach to XANES spectra pertaining to a bimolecular reaction in solution on organic substrates evolving on the millisecond time scale. In the presented framework, the direct in situ determination of the full mechanistic picture for the reaction involving the TPA substrate and the geometrical characterization of the reaction intermediates are achieved.

2. MATERIALS AND METHODS

2.1. Materials. All reagents and solvents were employed at the highest commercial quality and used without additional purification. TPA and peracetic acid (36–40 wt % in acetic acid, stored at 4 °C) were purchased from Sigma-Aldrich. Iron(II) bis(trifluoro-methanesulfonate)bis(acetonitrile), [Fe(OTf)₂(CH₃CN)₂], was prepared according to a literature procedure from anhydrous Fe(II) chloride (Sigma-Aldrich). [Fe⁶⁺(TPA)(CH₃CN)₂](OTf)₂ was prepared by metatation of the ligand TPA (Sigma-Aldrich) with [Fe(OTf)₂(CH₃CN)₂] in dry CH₃CN, and crystallization was performed by slow diffusion of dry diethyl ether in a dry dichloromethane solution as described in a literature method. Preparation and handling of air-sensitive materials were performed in an inert atmosphere by using a standard Schlenk and vacuum line techniques or a glove bag under N₂ atmosphere. Subsequently, the complex was stored under inert atmosphere. When the [Fe⁶⁺(TPA)-(OTf)] complex is dissolved in CH₃CN, two solvent molecules enter the iron first coordination sphere giving rise to the [Fe⁶⁺(TPA)-(CH₃CN)] complex. From now on, solely the cation [Fe⁶⁺(TPA)(CH₃CN)]⁻ will be mentioned.

2.2. Methods. 2.2.1. Reaction Details. For every stopped-flow mixing experiment, stock solutions of 70 mM [Fe⁶⁺(TPA)-(CH₃CN)]⁻ in CH₃CN and 70 mM AcOOH in CH₃CN (diluted from the commercially available 36–40 wt % AcOOH solution in acetic acid, Sigma-Aldrich) were inserted into the reservoirs of the stopped-flow instrument. They were mixed in a 1:1 volume ratio at room temperature to obtain final concentrations of 35 mM for both [Fe⁶⁺(TPA)(CH₃CN)]⁻ and AcOOH. For all measurements, 100 μL of each solution was shot by the instrument into the cell.

2.2.2. Energy Dispersive X-ray Absorption Measurements. EXDAS were collected at the ID24 beamline of the European Synchrotron Radiation Facility (ESRF), Grenoble (the ring energy was 6.0 GeV, and the current was 150–200 mA). The X-ray source consists of two undulators, whose gaps were tuned to place the first harmonic at 7100 eV. The beam was focused horizontally to an 8 μm full width at half-maximum (fwhm) spot on the sample by the curved Si(111) polychromator crystal in Bragg geometry. In the vertical direction, the beam was focused using a bent Si mirror at a glancing angle of 3 mrad with respect to the direct beam. To minimize sample radiation damage, the vertical spot size was set at 40 μm fwhm. Spectra were recorded in transmission mode using a fast read out low noise (FReLoN) high frame-rate detector based on charge coupled device (CCD) cameras optically coupled with a scintillator screen. Acquisition time was 40 ms for each spectrum. Sequences of 50–100 individual spectra were acquired, covering a total time span of 2–4 s during the reaction. Each sequence was repeated three times, and the data were averaged to obtain a better signal-to-noise ratio. The energy calibration was made by measuring the absorption spectrum of an Fe foil, and the first inflection point was set at 7111 eV. All measurements were performed at 25 °C. EXDAS spectra were recorded with a Bio-Logic SFM-400 stopped-flow device equipped with a flow-through quartz capillary cell. The quartz capillary cell had a diameter of 1.3 mm and wall thickness of ~10 μm. The dead time of the stopped-flow device is ~2.0 ms for the flow rate of 8 mL/s as calibrated using the procedure described elsewhere, and it defines the shortest kinetic time that is accessible for spectroscopic measurements.

2.2.3. EDXAS Data Treatment. The stopped-flow apparatus used to perform the reaction requires a quartz capillary cell that worsens the quality of the EDXAS spectra due to scattering by quartz. For each measurement the EDXAS spectrum of the cell containing pure acetonitrile was collected after the sample spectrum, using the same statistic. The cell spectrum was subtracted from the sample spectrum to gain a better signal-to-noise (S/N) ratio and a higher resolution for the structural oscillations and a more defined Fe K-edge position. The spectra were then subjected to a smoothing procedure using the Savitzky-Golay Smoothing filter, as described in refs 57 and 58.

2.2.4. Decomposition of EDXAS Data into the Spectra and Relative Concentrations of Key Components. XANES time-resolved measurements yield a large series of spectroscopic data that may be arranged in a spectral matrix D, where each column of D is a spectrum measured at time t. Following the Lambert–Beer law, each experimental spectrum may be seen as the superposition of a number N of “pure” and uncorrelated components multiplied by their relative concentration. The decomposition of the experimental EDXAS data into the N spectra associated with the key reaction species and the relative concentration profiles was performed using the PyFitIt code. To do so, this software employs a strategy belonging to the class of the MCR methods. The decomposition’s starting point is the Singular Value Decomposition (SVD) expression:

\[ D = U \Sigma V^T + E \]

where the product \( U \Sigma \) contains, on its N columns, a set of values associable to the normalized absorption coefficients, \( \Sigma \) is a diagonal matrix called singular values term, whose elements are sorted in decreasing order, while V can be interpreted as the concentration matrix associated with the N-selected components. Finally, the error matrix E represents the lack of fit between the experimental data matrix D and the reconstructed one \( \mu = U \Sigma V \). The SVD decomposition depends on the correct estimation of the number of components N present in the experimental spectral matrix. This may be achieved by combining different statistical and empirical evidence. Among them, in this work we chose to use the scree plot analysis as shown afterward in Figure 2a, since it is easily and
effectively interpreted. At this stage, all matrices present in eq 1 are mere mathematical solutions of the spectral separation problem and do not possess any chemical meaning. Once N is established, the approach implemented by PyFitt requires the introduction of a transformation \( N \times N \) matrix \( T \) in eq 1, using the relation \( I = T T^T \)

\[
D = U \Sigma T^{-1} V + E
\]

(2)

where the spectra belonging to the key reaction species are given by \( S = U \Sigma T \), and their concentration profiles are given by \( C = T^{-1} V \). The matrix elements \( T_{ij} \) of matrix \( T \) are then modified by sliders to achieve \( S \) and \( C \), which are chemically and physically interpretable. Once this step is achieved, one can finally write

\[
D = S C + E
\]

(3)

In this work, to reduce the unknown number of elements of \( T \), which is in principle equal to \( N^2 \), the normalization of all spectral components contained in matrix \( S \) and the mass balance condition for the concentrations contained in matrix \( C \) were imposed. Further, the first spectrum assigned to the reaction’s initial species, complex \([Fe^{II}(TPA)(CH_3CN)]^{2+}\), was constrained to be equal to the EDXAS spectrum recorded on a CH_3CN solution containing complex \([Fe^{II}(TPA)(CH_3CN)]^{2+}\). This EDXAS spectrum is superimposable with the XAS spectrum recorded in transmission mode of a \([Fe^{II}(TPA)(CH_3CN)]^{2+}\) solution at 25 °C (see Figure 1 in ref S) and therefore belongs to the initial Fe(II) species. This spectral profile represents the \( t = 0 \) s starting point for the matrical decomposition. For the detailed explanation of how these constraints are imposed, see ref 17.

2.2.5. XANES Data Analysis. Each XANES spectra extracted by the matrical decomposition was assigned to a reaction key species and analyzed using the MXAN code.59,60 This code is based on the matrical decomposition was assigned to a reaction key species and their concentration profiles are given by \( C = T^{-1} V \). The matrix elements \( T_{ij} \) of matrix \( T \) are then modified by sliders to achieve \( S \) and \( C \), which are chemically and physically interpretable. Once this step is achieved, one can finally write

The analysis of the XANES spectrum assigned to complex \([Fe^{ii}(TPA)](\mu-O)(\mu-OAC)\)^{2+} was performed starting from its crystal structure. In this structure there are two Fe atoms each coordinated by a TPA ligand and an oxygen atom belonging to an acetate molecule, and they are linked through a bridging oxygen atom (Obridge). Because of the symmetry of the two Fe sites, the minimization procedure was performed by optimizing three bond lengths (Fe–NTPA, Fe–Ooxo, and Fe–Obridge). Theoretical XANES spectra were calculated including scatterers within 5 and 6 Å around a selected Fe atom, and it was found that scattering atoms do not contribute significantly to the theoretical spectrum outside a cutoff radius of 5 Å.

Hydrogen atoms were not included in all MXAN analyses. For all spectra, five nonstructural parameters were refined, namely, the threshold energy \( E_p \) the Fermi energy level \( E_F \) the energy and amplitude of the plasmon \( E_p \) and \( A_p \), and the experimental resolution \( \Gamma_{exp} \). The quality of the fits was estimated with the residual function \( R_{exp} \).

3. RESULTS AND DISCUSSION

Figure 2 shows the experimental EDXAS spectra recorded with a time resolution of 40 ms during the reaction of \([Fe^{ii}(TPA)](\text{CH}_3\text{CN})_2\)^{2+} (35 mM) with AcOOH (35 mM) CH_3CN/AcOH (99.6:0.4 (v/v)) at 25 °C, where the \( t = 0.00 \) s spectrum was fixed to the EDXAS spectrum of a CH_3CN solution containing \([Fe^{ii}(TPA)(\text{CH}_3\text{CN})]^{2+}\) (35 mM). One can note that the most apparent variations in the spectra are contained in the spectra between \( t = 0.00 \) s and \( t = 0.40 \) s from reaction start. Notably, between \( t = 0.04 \) s and \( t = 0.20 \) s the energy edge progressively shifts to higher energies, while between \( t = 0.20 \) s and \( t = 0.40 \) s the energy edge moves to lower energies. In this same time interval one may note the appearance of a 1s → 3d transition located at ~7113 eV. This transition is visible for spectra between \( t = 0.12 \) s and \( t = 0.20 \) s before decaying to zero as the reaction proceeds. After \( t = 0.60 \) s, the visible spectral variations are greatly abated. These results are consistent with the reaction mechanism shown in Figure 1, where the initial Fe(II) species undergoes a first oxidation to the Fe(III) complex, which is further oxidized to the Fe(IV) oxo complex, which returns by decay to an Fe(III) state. Iron acquires three different oxidation states (assigned to complexes stable enough to be isolated) during the reaction, and therefore one expects the number \( N \) of independent components present in the data mixture to be \( N = 3 \) or greater.

Principal component analysis (PCA) was applied to the EDXAS data set to confirm this qualitative analysis and to identify the number of chemical components present in the reaction data mixture.61 The results are presented in Figure 3. The singular values, extracted from SVD method, are the diagonal elements of matrix \( \Sigma \) reported in eq 1. These quantities are proportional to the data variance explained by each component. It follows that each of them can be properly plotted against the related component number, generating the so-called scree plot, as reported in Figure 3a. One can note from the plot the existence of an elbow indicating the presence of three relevant components. Conversely, for numbers of components greater than three, the related singular values decrease slowly with approximately the same decaying slope,
indicating that these components contribute to the data set reconstruction in the same way and are, for this reason, associated with noise. This statistical evidence suggests there are three principal components present in the data set. This result is in accordance with the chemical knowledge of the reaction mechanism that predicts the succession of three distinct oxidation states for Fe. The percentage residual error committed in reconstructing the data set with three components is shown in Figure 3b. The percentage error function was calculated with the following expression

\[
R(t) = \frac{\sum_{i=1}^{K} \sum_{j=1}^{m} |d_{ij} - \mu_{ij}^{PC=3}|}{\sum_{i=1}^{K} \sum_{j=1}^{m} |d_{ij}|} \times 100
\]

(4)

where \(d_{ij}\) and \(\mu_{ij}^{PC=3}\) are the normalized absorbance values for the data set and for the data set reconstructed with \(N = 3\), respectively (\(K\) and \(m\) represent the number of acquired time-resolved spectra and of the energy points, respectively). Interestingly, one may observe an increase in the percentage error in proximity of the spectra recorded between \(t = 0.04\) s and \(t = 0.16\) s. Since the main EDXAS spectral variation in the experimental data is observed in the same time interval, this finding suggests the presence of a diluted and transient species that contributes in small percentage to the overall measured signal. It is probable that by including an ulterior fourth (or fifth) component in the decomposition this error would diminish. However, relying on the knowledge of the reaction mechanism, on the scree plot analysis, and on the relatively small error (inferior to 1.2%) committed in the reconstruction with \(N = 3\), we decided to employ only three PCs for the subsequent analysis.

The transformation matrix approach implemented in PyFitit\(^1\) was used to decompose the data set, employing a \(3 \times 3\) \(T\) transformation matrix. Furthermore, by imposing the set of constraints described in Section 2.2.4, the number of \(T_{ij}\) elements was reduced from nine to four. Each of these four terms was varied preserving the mass balance condition and the non-negativity of the extracted spectra and concentration profiles. A solution to the decomposition expressed by eq 3, possessing a sound chemical meaning, was achieved through the matrix

\[
T = \begin{pmatrix}
1/\sigma & 1/\sigma & 1/\sigma \\
T_{21} & +0.10 & +0.50 \\
T_{31} & -0.10 & -0.15
\end{pmatrix}
\]

(5)

where \(\sigma\) is the normalization coefficient, \(1/\sigma = -0.17\), \(T_{21} = 0.51\), and \(T_{31} = 0.57\).

Figure 4a shows the isolated EDXAS spectra, and Figure 4b shows their fractional components in the reaction mixture. The first spectral component (blue) belongs to complex \([\text{Fe}^{II}(\text{TPA})(\text{CH}_3\text{CN})_2]^+\). The second (red) and third (green) components are assigned to complexes in which iron has the oxidation states of Fe(IV) and Fe(III), respectively. In fact, the oxidation state of each spectrum is identified by the relative energy position of the main absorption edge. The first reflection point of the spectrum belonging to the initial Fe(II) reactant lies at lower energy than those assigned to the Fe(IV) and Fe(III) species, while that of the Fe(IV) is found at the highest energies. Interestingly, the Fe(IV) complex shows a 1s \(\rightarrow\) 3d dipole-forbidden transition centered at \(\sim 7113\) eV. This feature is absent in the spectrum of the Fe(II) reactant and weak in that of the Fe(III) compound. This finding further supports the proposed identification of the reaction species. It is known that Fe(IV) oxo complexes show a relatively intense 1s \(\rightarrow\) 3d transition due to their noncentrosymmetry, and it has been reported that this is also the case for complex...
On this basis, the second XANES spectral component is assigned to complex \([\text{Fe}^{\text{IV}}(\text{TPA})(\text{O})(X)]^{+}/2+\).

Looking at Figure 4b one may note that the fractional concentration of the initial \([\text{Fe}^{\text{II}}(\text{TPA})(\text{CH}_3\text{CN})_2]^2+\) complex rapidly decays to zero, while the concentration of the \(\text{Fe}^{\text{IV}}\) species shows an accumulation between \(t = 0.12\) s and \(t = 0.20\) s. Conversely, the concentration of the \(\text{Fe}^{\text{III}}\) component is prevalent before the formation of the oxo complex at \(t = 0.04\) s and \(t = 0.08\) s. It decreases to almost zero when the concentration of \(\text{Fe}^{\text{IV}}\) reaches its maximum, and then it gradually increases to become the reaction product from \(t = 0.24\) s until the end of the process. These results confirm the sequence of the oxidation states that \(\text{Fe}\) assumes during the reaction shown in Figure 1 and prove, through the direct analysis of room-temperature reaction EDXAS spectra, that the starting \(\text{Fe}^{\text{II}}\) species initially evolves to an \(\text{Fe}^{\text{III}}\) intermediate.

The transformation matrix-based approach implemented in this investigation inherently suffers of rotational ambiguity. It follows that the solutions of the decomposition problem shown in Figure 4 are not unique.\(^{15,17}\) To address the validity of the extracted spectra and concentrations for the reaction components, the time evolution of the area belonging to the pre-edge 1s \(\rightarrow\) 3d transition at 7113 eV and of the edge energy position of the EDXAS spectra were evaluated turning to the raw EDXAS time-resolved spectra.

Figure 5a presents the variation during the reaction of the area of the dipole-forbidden transition measured on the raw XANES spectra. One can note that there is a maximum localized at \(t = 0.16\) s, which is indicative of the formation of the noncentrosymmetric oxo complex. The time evolution of the \(\text{Fe}E\) energy (shown Figure 5b) was qualitatively evaluated by measuring the energy at \(\mu(E) = 0.40\) for each raw spectrum, an approach that we have shown to be successful in the analysis of EDXAS spectra acquired during a chemical reaction in solution.\(^{14}\) The result of this procedure is shown in

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### Table 1. Fe K-Edge XANES Best-Fit Structural Parameters\(^a\)

| \(N\) | \(R_{\text{Fe-NTPA}}\) (Å) | \(R_{\text{Fe-NACN}}\) (Å) | \(R_{\text{Fe-O,O}^{\text{oxo}}}\) (Å) | \(R_{\text{Fe-O,O}^{\text{pe}}}\) (Å) | \(R_{\text{Fe-O,O}^{\text{OA}}_c}\) (Å) | \(R_{\text{Fe-O,O}^{\text{bridge}}}\) (Å) |
|------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| 1    | 4 1.91(1) − 1.99(1) 2 1.92(1) − 1.93(1) | 2 1.92(1) − 1.93(1) | 1.80(5) | 1.80(5) | 1.80(5) |
| this work | 4 2.04(5) 2 1.97(5) | 2 1.97(5) | 1.80(5) | 1.80(5) | 1.80(5) |
| 2    | DFT\(^b\) 4 1.94−2.01 | 1 2.00 | 1 1.81 | 1 1.86(5) |
| this work | 4 2.03(5) | 1 2.00(5) | 1 1.86(5) |
| 3    | EXAFS\(^c\) 4 1.99 | 1 2.00| 1 1.67(2) | 1 2.00 | 1 2.05(5) |
| this work | 4 2.04(5) | 1 1.77(5) | 1 2.05(5) |
| 3a   | 4 2.05(5) 1 2.06(5) 1 1.77(5) | 1 2.05(5) |
| 3b   | this work | 4 2.05(5) 1 2.06(5) 1 1.77(5) | 1 2.05(5) |
| 4    | cryst\(^d\) 8 2.105(6) − 2.242(6) | 2 1.972(6) − 2.038(6) 2 1.779(3)−1.815(3) | 2 1.972(6) − 2.038(6) 2 1.779(3)−1.815(3) | 1 2.00(5) 1 1.80(5) |
| this work | 4 2.06(5) | 1 2.00(5) | 1 1.80(5) | 1 2.00(5) | 1 1.80(5) |

\(^a\)Fe K-edge XANES best-fit structural parameters of \([\text{Fe}^{\text{II}}(\text{TPA})(\text{CH}_3\text{CN})_2]^2+\) (1), \([\text{Fe}^{\text{III}}(\text{TPA})(\kappa^2-\text{OAc})]^2+\) (2), \([\text{Fe}^{\text{IV}}(\text{TPA})(\text{O})(X)]^{+}/2+\) (3), \([\text{Fe}^{\text{IV}}(\text{TPA})(\text{O})(\text{OAc})]^+\) (3a), \([\text{Fe}^{\text{IV}}(\text{TPA})(\text{O})(\text{CH}_3\text{CN})]^{2+}\) (3b), and \([\text{Fe}^{\text{IV}}(\text{TPA})(\mu-\text{O})(\mu-\text{OAc})]^3+\) (4) compared to the available literature: crystallographic, EXAFS, and DFT data. \(R_{\text{Fe-NTPA}}\) and \(R_{\text{Fe-NACN}}\) are the average distances between the metal cation, the TPA, and the solvent nitrogen atoms, respectively, \(R_{\text{Fe-O,O}^{\text{oxo}}}\) is the distance between the metal cation and the coordinating oxygen of the oxo group in \([\text{Fe}^{\text{IV}}(\text{TPA})(\text{O})(\text{OAc})]^+\) and \([\text{Fe}^{\text{IV}}(\text{TPA})(\text{O})(\text{CH}_3\text{CN})]^{2+}\), \(R_{\text{Fe-O,O}^{\text{pe}}}\) is the distance between \(\text{Fe}\) and the negatively charged peracetate oxygen in complex \([\text{Fe}^{\text{III}}(\text{TPA})(\kappa^2-\text{OAc})]^2+\), \(R_{\text{Fe-O,O}^{\text{OA}}_c}\) is the average distance between the metal cation and the acetate group, and \(R_{\text{Fe-O,O}^{\text{bridge}}}\) is the average distance between the metal cation and the bridging oxygen atom in complex \([\text{Fe}^{\text{IV}}(\text{TPA})(\mu-\text{O})(\mu-\text{OAc})]^3+\), while \(N\) is the coordination number.

\(^b\)A nitrogen or an oxygen atom located at 2.20 Å from the central cation is reported in ref 6.

\(^c\)A nitrogen or an oxygen atom located at 2.20 Å from the central cation is reported in ref 6.
Figure 5b, where one can note a shift of the main absorption edge to higher energies of ~0.8 eV from $t = 0.08$ s to $t = 0.12$ s. Such a change is consistent with the oxidation of the Fe(III) present in the reaction mixture to Fe(IV). This analysis confirms also that the lifetime of the Fe(IV) complex is in the range within $t = 0.12$ s and $t = 0.24$ s. Finally, the edge moves to lower energies upon the reduction of Fe(IV) to Fe(III) and reaches a constant value, which does not exactly coincide with the edge value prior to the Fe(III) $\rightarrow$ Fe(IV) oxidation. This is due to the presence at $t = 0.04$ s and $t = 0.08$ s of the Fe(II) component and to the contribution of the Fe(IV) component at $t > 0.24$ s. Such independent methods fully support the validity of the mathematical solutions shown in Figure 4.

On the basis of the established reaction mechanism, the isolated spectral component associated with an Fe(III) oxidation state may be assigned to both the proposed acyl-peroxo intermediate $[\text{Fe}^{\text{III}}(\text{TPA})(\kappa^2-\text{OOAc})]^2+$ and to the dimeric product $[\text{Fe}^{\text{II}}(\text{TPA})_2(\mu-\text{O})(\mu-\text{OAc})]^3+$. In fact, the local geometry around the central Fe(III), in both complexes, is made of four nitrogen atoms belonging to the TPA chain and of two coordinating oxygen atoms. Further, as is listed in Table 1 the DFT-optimized first-shell distances of complex $[\text{Fe}^{\text{III}}(\text{TPA})(\kappa^2-\text{OOAc})]^2+$ and the crystallographic first-shell distances of complex $[\text{Fe}^{\text{III}}(\text{TPA})_2(\mu-\text{O})(\mu-\text{OAc})]^3+$ closely resemble one another. This evidence, together with the relatively small error committed in reproducing the experimental data set with $N = 3$, supports the identification of the same spectral component for both Fe(III) species.

To test these hypotheses and to obtain quantitative structural information regarding all the reaction intermediates, a full MS analysis was performed on the three isolated spectral components. The XANES spectrum of the Fe(II) complex is quite different from those of the Fe(III) and Fe(IV) species. As previously mentioned, the Fe first coordination shell is made up by the four nitrogen belonging to the TPA backbone, which were placed at the same Fe$-\text{N}_{\text{TPA}}$ distance, and by two nitrogen atoms belonging to the CH$_3$CN solvent molecules. During the fitting procedure, the Fe$-\text{N}_{\text{TPA}}$ and the Fe$-\text{N}_{\text{ACN}}$ distances were refined together with the nonstructural parameters to obtain the best agreement with the experimental spectrum. The best-fit results are shown in Figure 6a, while the molecular cluster obtained from the minimization is shown to the right. The agreement between the theoretical spectrum and the isolated component is excellent. The refined parameters are listed in Table 1. The Fe$-\text{N}_{\text{TPA}}$ and the Fe$-\text{N}_{\text{ACN}}$ distances are in good agreement with the crystallographic values within the statistical errors. It is well-known that systematic errors are present in the XANES analysis performed with MXAN and that they arise mostly because of the poor approximation used for the phenomenological broadening function $\Gamma(E)$ that mimics the electronic damping. In all cases studied until now such systematic errors did not appreciably affect the structural results, confirming how this spectroscopy is dominated by the geometry of the atomic cluster rather than by its electronic structure.$^{66-69}$ The full list of nonstructural parameters is reported in Table 2.

As far as the Fe(IV) oxo complex is concerned, the MXAN analysis was performed using two different models. In the former ($[\text{Fe}^{\text{IV}}(\text{TPA})(\kappa^2-\text{OAc})]^2+$) the central Fe cation is coordinated to an acetate molecule and to the TPA chain. In this case, the Fe$-\text{O}_{\text{exo}}$ and the Fe$-\text{O}_{\text{OAc}}$ distances were optimized independently together with the Fe$-\text{N}_{\text{TPA}}$ one. The results of this analysis are shown in Figure 6c, while the best-fit structural parameters are listed in Table 1. Also in this case the agreement between the experimental and theoretical spectra is
Figure 7. Fe K-edge XANES isolated spectrum relative to the \([\text{Fe}^{IV}(\text{TPA})(\text{O})(\text{X})]^{2+}\) complex (blue, dotted line) compared with the theoretical curve (red, full line) calculated upon an optimized geometrical model, where \(X = \text{CH}_3\text{CN}\). The best-fit geometry is also depicted, where iron, nitrogen, carbon, and oxygen atoms are in yellow, cyan, orange, and red, respectively.

satisfactory (\(R_w = 2.8\)), suggesting that an acetate molecule coordinates the Fe atom in the Fe(IV) species. Further proof of this hypothesis was gained by performing a second minimization using a structural model where the central Fe is coordinated to the four TPA nitrogens, the oxo oxygen atom, and a CH3CN ligand. In this case the Fe–O_{oxo} and the Fe–N_{TPA} distances were optimized together with the Fe–N_{CH3CN} bond length. The best-fit structural and non-structural parameters are listed in Tables 1 and 2, respectively. The comparison of the theoretical and isolated XANES spectra is presented in Figure 7, while the corresponding optimized atomic cluster is depicted below. In this case a slightly worse agreement was obtained between the two XANES spectra (\(R_w = 2.9\)). This finding supports the hypothesis that a molecule coordinating the central Fe cation with an oxygen atom, such as acetate, has a higher residence time compared to that of a molecule coordinating the metal site with a nitrogen atom, such as CH3CN. Consequently, one may suggest that the previously unidentified sixth ligand in the \([\text{Fe}^{IV}(\text{TPA})(\text{O})(\text{X})]^{2+}\) complex is the acetate anion.

Through the quantitative analysis of the XANES spectrum extracted from the decomposition, identical first-shell distances are found for both \(X = \text{AcO}^-\) and \(X = \text{CH}_3\text{CN}\), as reported in Table 1. The Fe–N_{TPA} distance is 2.04 Å, a value that coincides within the experimental errors with the EXAFS reported distance for complex \([\text{Fe}^{IV}(\text{TPA})(\text{O})(\text{X})]^{2+}\). Additionally, the theoretical calculations derive a distance between the central Fe cation and the sixth coordinating ligand of 2.05(5) and 2.06(5) Å for \(X = \text{AcO}^-\) and \(X = \text{CH}_3\text{CN}\), respectively, to be compared with the literature value of 2.20 Å.

Additionally, the Fe–O_{oxo} distance was evaluated through X-ray crystallography for some nonheme oxo complexes such as \([\text{Fe}^{IV}(\text{TPC})(\text{O})(\text{NCCCH}_3)]^{2+}\) and \([\text{Fe}^{IV}(\text{NAP})(\text{O})]^{2+}\), and it was found to be 1.646(3) Å and 1.639(5) Å, respectively. Numerous other nonheme oxo complexes have been characterized through EXAFS, with the Fe–O distance residing in the range between 1.62 and 1.70 Å. This slight discrepancy of our results with the existing literature may be because the 1.77(5) Å value is obtained on the direct analysis of the spectrum at room temperature of the Fe(IV) species, whereas other measurements have been all performed at low temperatures on a frozen solution or on the crystal, if available. Moreover, as previously underlined, a systematic error is present in the structural determinations conducted with MXAN, as evidenced in the geometrical characterization of other iron heme complexes and heme proteins.

The XANES spectrum assigned to complexes \([\text{Fe}^{III}(\text{TPA})(\kappa^2-\text{OAc})]^{2+}\) and \([\text{Fe}^{III}(\text{TPA})(\mu-\text{O})(\mu-\text{OAc})]^{3+}\) was subjected to two distinct minimization procedures. In the first one, the spectrum was analyzed starting from the DFT-optimized structure associated with complex \([\text{Fe}^{III}(\text{TPA})(\kappa^2-\text{OAc})]^{2+}\), where the Fe cation is coordinated by the four nitrogens of the TPA chain and by two oxygen atoms belonging to a peracetate molecule. The results of the analysis are shown in Figure 6b, where the experimental and theoretical curves are reported together with the molecular cluster. The agreement between the data is excellent (\(R_w = 1.1\)) and the structural results coincide with the literature data within the statistical errors (Table 1). These findings represent important structural data that confirm the identity of the reaction intermediate \([\text{Fe}^{III}(\text{TPA})(\kappa^2-\text{OAc})]^{2+}\). Note that a small percentage of the corresponding Fe(V) oxo complex may form upon heterolysis of the O–O bond of the Fe(III) peroxo species as observed at low temperature by Talsi et al. However, at room temperature the Fe(V) species is too unstable to be observed by our method given the time scale of our experimental conditions.

Finally, the XANES spectrum was analyzed starting from the crystal structure of the \(\mu\)-dimeric species \([\text{Fe}^{III}(\text{TPA})(\mu-\text{O})(\mu-\text{OAc})]^{3+}\). The same coordination environment was used for both Fe^{III} atoms in the dimer. It comprises the four nitrogen atoms of the TPA ligand, the acetate molecule, and a bridging oxygen atom. During the fitting procedure, the Fe–N_{TPA}, Fe–O_{Ac}, and Fe–O_{bridge} distances were optimized, while the TPA structure was kept fixed to the initial geometry, and all of the atoms within 5 Å of the central metal cation were included in the theoretical calculation. Figure 6d presents the experimental and theoretical spectra together with the atomic cluster. The agreement between the two curves is very good (\(R_w = 1.3\)). The structural results, listed in Table 1, highlight a slight compression of the Fe–N_{TPA} and Fe–O_{bridge} bond lengths compared to the crystal structure. One may note that the calculated first-shell distances for complexes \([\text{Fe}^{III}(\text{TPA})(\kappa^2-\text{OAc})]^{2+}\) and \([\text{Fe}^{III}(\text{TPA})(\mu-\text{O})(\mu-\text{OAc})]^{3+}\) are identical within the statistical errors, as expected, since they were optimized on the basis of the same XANES spectrum.

4. CONCLUSIONS

This work demonstrates that it is possible to derive important mechanistic insights for a reactive process occurring in solution on the millisecond scale and to structurally characterize its transient intermediates through a multivariate E DXAS analysis. The implemented approach has enabled the direct determi...
nation of the mechanism of the reaction between [Fe(III)(TPA)-(CH₂CN)]²⁺ and AcOOH using the TPA nonheme complex at 25 °C. In particular, it is confirmed that an Fe(III) acylperoxo intermediate is initially formed, which in turn evolves to a Fe(IV) oxo complex. The sixth ligand of the latter species, which was previously unidentified, is shown to be an acetate ion. This strategy allows one to characterize elusive intermediates whose geometries cannot be easily determined using the conventional experimental methods. Its combination with EDXAS holds great promise, especially for the investigation of complex redox reaction mechanisms on organic substrates that are silent to laboratory-based spectroscopies.

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Notes
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