HERFD-XANES probes of electronic structures of iron\textsuperscript{II/III} carbene complexes\textsuperscript{†}

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Iron centered N-heterocyclic carbene (Fe-NHC) complexes have shown long-lived excited states with charge transfer character useful for light harvesting applications. Understanding the nature of the metal-ligand bond is of fundamental importance to rationally tailor the properties of transition metal complexes. The high-energy-resolution fluorescence detected X-ray absorption near edge structure (HERFD-XANES) has been used to probe the valence orbitals of three carbene complexes, [Fe\textsuperscript{II}(bpy)(btz)\textsubscript{2}](PF\textsubscript{6})\textsubscript{2} (bpy = 2,2\textsuperscript{′}-bipyridine, btz = 3,3\textsuperscript{′}-dimethyl-1,1\textsuperscript{′}-bis(4-toly)-4,4\textsuperscript{′}-bis(1,2,3-triazol-5-ylidenel)), [Fe\textsuperscript{II}(btz)\textsubscript{3}](PF\textsubscript{6})\textsubscript{5}, and [Fe\textsuperscript{II}(phtmeimb)\textsubscript{2}]PF\textsubscript{6} (phtmeimb = phenyl(tris(3-methylimidazol-2-ylidene))borate\textsuperscript{−}).

1 Introduction

Transition metal complexes using ruthenium, osmium, and iridium metals are popular choices as photosensitizers or photocatalysts due to their long-lived charge transfer (CT) excited states combined with photo- and chemical stability.\textsuperscript{1–5} However, their widespread deployment has been limited due to the rare and sometimes harmful nature of their central metal. The abundance (and low price) as well as the relatively non-toxic properties of first-row transition metals like iron make them attractive choices to replace noble metals for large-scale light-harvesting and light-emitting devices. The application of first-row transition metals has so far been hampered by the short lifetimes of CT states, which usually deactivate ultra-fast through lower-energy metal-centered (MC) states.\textsuperscript{5–11} Visible light excitation generates in the here discussed complexes a metal-to-ligand charge transfer (MLCT) state or a ligand-to-metal charge transfer (LMCT) state depending on the oxidation state of iron. The relative energy of the CT states and the MC states can be manipulated by e.g. tuning the metal-ligand interactions, for instance, by tuning the σ-donating and \pi-accepting strengths of the ligands.\textsuperscript{12} Our recently developed iron complexes show long-lived (up to ns) excited states with lifetime of 13 ps and 528 ps have been achieved for iron carbene complexes [Fe\textsuperscript{II}(bpy)(btz)\textsubscript{2}](PF\textsubscript{6})\textsubscript{2} and [Fe\textsuperscript{II}(btz)\textsubscript{3}](PF\textsubscript{6})\textsubscript{5}, respectively,\textsuperscript{13,15} and LMCT state lifetimes of 100 ps and 2 ns have been achieved for [Fe\textsuperscript{II}(btz)\textsubscript{3}](PF\textsubscript{6})\textsubscript{5} and [Fe\textsuperscript{II}(phtmeimb)\textsubscript{2}]PF\textsubscript{6}, respectively.\textsuperscript{14,16} These long lifetimes make iron carbene...
complexes interesting candidates as photo-sensitizers and photo-redox catalysts.\textsuperscript{14,16}

For the design of new generations of iron carbene complexes, improved knowledge about their electronic structures is required such as how the ligands affect the valence orbitals, and how they influence the relative energies of CT and MC states involved in the excited state decay processes. X-ray absorption spectroscopy (XAS) directly probes the unoccupied and partially occupied valence orbitals through transitions from strongly localized deep core orbitals and is thus a useful method to elucidate these details.\textsuperscript{17,18} Metal K edge XAS is usually subdivided into two regions, X-ray absorption near edge structure (XANES) and the extended X-ray fine structure (EXAFS). The transitions probed with XANES give electronic-structure information about the occupation of valence orbitals, the orbital covalency, oxidation state of metal center, and the site symmetry, which this paper is focused on. Particularly we will focus on the energetic position and intensity of the pre-edge features, as these are a direct probe of the valence orbitals. The intensity of the transition metal K pre-edge features is usually associated with dipole-forbidden electronic transitions from the 1s to the 3d orbitals. The K pre-edge intensity is obviously dependent on the number of holes in the 3d shell, another strong, sometimes even stronger influence is the character of the target molecular orbitals. Their intensity is thus a measure for the metal 3d character of the valence orbitals involved in the transitions and of the covalency of the metal–ligand bonds.\textsuperscript{17} Apart from the metals 3d characters, the intensity can be significantly deviated by including electric-dipole transitions, back-donation charge transfers.\textsuperscript{17,19,20} The electric quadrupole transitions are in general 2 orders of magnitude weaker than electric dipole allowed transitions, which has been shown in complexes with distorted centrosymmetry.\textsuperscript{21} The distortion allows for the delocalization of metal 4p character into metal-3d character orbitals through orbital interaction with the ligand.

One challenge for these type of studies is that conventional metal K pre-edge spectra measured with XAS are broadened significantly due to the short lifetime of the initial 1s core hole excitation (core-hole lifetime broadening $\sim 1.25$ eV for Fe).\textsuperscript{22} This spectral broadening can often be circumvented by collecting the emitted X-ray fluorescence in a narrow energy bandwidth in a technique called high-energy-resolution fluorescence detected X-ray absorption spectroscopy (HERFD-XAS) and the measured spectrum is in many cases closely related to the XAS.\textsuperscript{23-26}

### 3 Computational details

Calculations of molecular systems have been performed on ground state geometries (see Fig. 1) optimized using density functional theory (DFT)\textsuperscript{27-28} with the modified hybrid functional B3LYP (15% Hartree–Fock exchange)\textsuperscript{29} and the 6-311G(d) basis set using Gaussian 09.\textsuperscript{30}

The calculations of X-ray absorption spectra have been performed using OpenMolcas.\textsuperscript{31} The method for metal K pre-edge XAS has shown its validity on 3d transition metal complexes.\textsuperscript{20,32,33} The active space used for spectra calculations could be designated as $\text{RAS}(n, l, m; i, j, k)$, where $i, j, k$ are the number of orbitals in $\text{RAS1}$, $\text{RAS2}$, and $\text{RAS3}$ spaces respectively, $n$ is the total number of electrons in the active space, $l$ the maximum number of holes allowed in $\text{RAS1}$, and $m$ the maximum number of electrons in $\text{RAS3}$. For the metal K pre-edge XAS calculation, the 1s core orbital is placed in the $\text{RAS1}$, the five metal 3d character orbitals together with two ligand-character $\sigma$ donation orbitals and three empty orbitals of $\pi$ symmetry are placed in the $\text{RAS2}$, the active space is shown in Fig. 2, this gives active space $\text{RAS}(12,1,0; 1,10,0)$ for $\text{Fe}^{\text{III}}(\text{bpy})(\text{btz})_2^{2+}$, and $\text{RAS}(11,1,0; 1,9,0)$ for $\text{Fe}^{\text{III}}(\text{btz})_3^{3+}$ and $\text{Fe}^{\text{III}}(\text{phtmeimb})_3^{3+}$.

RASSCF wavefunction optimizations have been performed using the state average (SA) formalism,\textsuperscript{34} which means that the same orbitals are used for all states of a specific spin and symmetry. Scalar relativistic effects have been included by using a second-order Douglas–Kroll–Hess Hamiltonian,\textsuperscript{35,36} in combination with the ANO-RCC basis set and the use of a Cholesky decomposition approach to approximate the two-electron integrals.\textsuperscript{17-19} The choice of basis set and number of final states included in which the calculations are performed is constant on all complexes. For further information on their dependency see e.g. Fig. S2 and S3 in the ESI.\textsuperscript{13}

### 2 Experimental section

The synthesis of the three carbene complexes discussed here has been described previously.\textsuperscript{13,14,16} The powder samples were ground into a fine powder and measured undiluted in a template of homogeneous thickness between two Kapton foils. The HERFD-XAS were recorded at the ID26 beamline of the European Synchrotron Radiation Facility (ESRF) by scanning the incident beam energy with the Si(311) monochromator crystal at a fixed undulator gap, while the energy of the fluorescence detector was kept fixed at the maximum energy of Fe K\textsubscript{2,3} emission line. An iron foil was used to calibrate the monochromator energy to 7112 eV at the first maximum of the derivative spectrum in transmission before the first sample was measured. The absorption edge of the last sample was measured prior to this experiment and found coinciding with this measurement, excluding energy drifts within the $\sim 15$ min total measurement time for all samples. The incoming flux was $\sim 10^{13}$ photons per second at a beam footprint on the sample of 0.1 (vertical) by 1 (horizontal) mm$^2$. The K\textsubscript{2,3} fluorescence intensity was recorded with an emission spectrometer employing the (440) reflection of four spherical ($r = 50$ mm) Ge analyzer crystals with spherically bent 1 m radius. The width of the elastic peak giving the combined energy bandwidth was 0.65 eV. All samples were checked for damage induced by the X-rays. The statistical quality of experimental spectra are evaluated by using Poisson distributed photon statistics of the number of counts and adding an (estimated) additional 2% of the local intensity to include possible variations during the normalization. In the plots the double of this error is presented as $2\sigma$ confidence environment.
The inclusion of dynamical correlation through second-order perturbation theory (RASPT2) calculation has been performed on \([\text{Fe}^{\text{III}}(\text{phtmeimb})_2]^{1+}\), the spectrum is presented in Fig. S4 (ESI †). The inclusion of RASPT2 calculation only has a minor influence on the data, while significantly increasing the calculation costs.\(^{40}\) Spin–orbit coupling is included by a RASSI approach.\(^{41,42}\) Overall, the expanded basis set and the inclusion of dynamical correction have pronounced effects on the absolute incident energy, and negligible effect on the spectral shape, see Fig. S2 and S4 in the ESI.\(^{†}\)

We thus use the usual octahedral labels of \(e_g\) and \(t_{2g}\) for the metal 3d-like orbitals. The \([\text{Fe}^{\text{II}}(\text{bpy})(\text{btz})_2]^2+\) has six electrons in the three \(t_{2g}\) orbitals, which gives a \(t_{2g}^6e_g^0\) ground state configuration. Both \([\text{Fe}^{\text{III}}(\text{btz})_3]^3+\) and \([\text{Fe}^{\text{III}}(\text{phtmeimb})_2]^3+\) have five electrons in three \(t_{2g}\) orbitals and thus \(t_{2g}^5e_g^0\) as ground state configuration.

The \([\text{Fe}^{\text{II}}(\text{bpy})(\text{btz})_2]^2+\) contains one bpy ligand and two btz ligands (also Fig. 1). Here the introduction of strongly \(\sigma\)-donating NHC ligands destabilized the lower-lying triplet and quintet MC states, while pushing away the minimum of their potential energy surface from the \(3\text{MLCT}\) state, hence making an intersystem crossing in to quintet multiplicity unlikely as \(3\text{MLCT}\) deactivation path as we have shown in our recent publication.\(^{11}\) In \([\text{Fe}^{\text{III}}(\text{btz})_3]^3+\), the bpy ligand is replaced by a third btz ligand. This replacement improves the orthoaxiality and leads to \(\text{trans}\) C–Fe–C angles of 179° compared to the 159° of the previously reported homoleptic NHC complexes with bis-tridentate ligand environment.\(^{43}\) The improved orthoaxiality could contribute to a larger ligand field splitting, and result in a further destabilization of the MC states.\(^{44}\) At the same time, the btz ligand is a stronger acceptor than bpy, which does stabilize the \(t_{2g}\) orbitals further, and gives a stronger ligand field strength. The \([\text{Fe}^{\text{III}}(\text{phtmeimb})_2]^3+\) has a near-perfect 180° for the C–Fe–C \(\text{trans}\) angle. The two monononic facial tris-carbene ligands have a more pronounced \(\sigma\)-donating ability and can further destabilize the \(e_g\) anti-bonding molecular orbitals of metal character, and hence destabilize the quartet and sextet MC states relative to the complex \([\text{Fe}^{\text{III}}(\text{btz})_3]^3+\). The MC state destabilization can contribute to the increase of activation barrier for the \(2\text{LMCT}\) state decay through the MC states.

The orbital covalency analyses for the selected valence orbitals from the RASSCF ground state calculations of the three carbene complexes are presented in the Table 1. These orbital covalency analyses are performed using the Multiwfn package.\(^{45}\) The characters of the selected orbitals are connected with the observed spectral features in the measurements and simulated metal K pre-edge XAS (and will be discussed in the following sections). The metal 3d and 4p characters in the targeted orbitals do reflect the electric quadrupole and dipole transition contribution intensity differences in the XAS, respectively.

### Results

All three complexes have pseudo-octahedral ligand coordination and are low spin in their ground state configuration.
4.1 Observations in the experimental HERFD-XANES

The [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2} has a $t_{2g}^{0}$ ground state configuration with fully filled $t_{2g}$ orbitals. The experimental K pre-edge of [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2} thus shows one dominant peak at 7113.6 eV which originates from the 1s to $e_{g}$ transitions, see Fig. 3. The very small feature at 7111.6 eV, a fingerprint for the $t_{2g}$ peak of the Fe \text{II} complexes, has not been resolved in the conventional metal K pre-edge. The $e_{g}$ peak of the Fe \text{III} complexes is located at lower energy compared to the respective peaks in the other two Fe \text{III} complexes. The analysis of the orbital character of the valence orbitals in the ground state complexes showed that the molecular orbitals with $e_{g}$ character of [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2} complex has $C_{2}$ symmetry in the calculation, the 4p orbitals can mix with $d_{x^2}$ and $d_{y^2}$ orbitals under $C_{2}$ symmetry.

Both Fe \text{III} complexes show the classical strong two-peak features originating from the 1s core electron transition to $t_{2g}$ and $e_{g}$, respectively. The $t_{2g}$ peak of [Fe\text{III}(phthmeimb)\text{2}](PF_{6})\text{2} shifts by $\sim 0.2$ eV compared to the $t_{2g}$ peak of [Fe\text{III}(btz)\text{3}](PF_{6})\text{3} complex, see Fig. 3. Both Fe \text{III} complexes also have clearly separated and broad $e_{g}$ features which originate from the multiplet structures of the valence shell. The energy difference between the different $e_{g}$ peaks is a measure of the difference of the attraction between the hole in a $t_{2g}$ orbital and the electron in an $e_{g}$ orbital but is also influenced by the orbital covalency.20 This fine splitting is usually not resolved in the conventional metal K pre-edge XAS due to the large lifetime broadening. The $e_{g}$ peak of [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2} is located at lower energy compared to the $e_{g}$ peak of the Fe \text{III} complexes. Recently this energy shift has been explained by the difference in contraction of the 3d shell in the core-excited states.46

4.2 Intensity contribution analysis

The distortion from centrosymmetry delocalizes some metal 4p orbitals into metal 3d orbitals through their interactions with the ligand orbitals. The heteroleptic [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2} complex has two btz ligands and one bpy ligand, which creates a less centrosymmetric environment around the iron center compared to the other two homoleptic Fe \text{III} complexes that are coordinated with tris-bidentate or bis-tridentate all-NHC ligands. The [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2} complex has $C_{2}$ symmetry in the calculation, the 4p orbitals can mix with $d_{x^2}$ and $d_{y^2}$ orbitals under $C_{2}$ symmetry. The near-perfect octahedral symmetry for the Fe \text{III} complexes is purposefully designed to achieve large octahedral ligand field splitting.14,16 The intensity of transitions with $e_{g}$ character peak for the [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2} at almost double intensity compared to the respective peaks in the other two Fe \text{III} complexes. The analysis of the orbital character of the valence orbitals in the ground state complexes showed that the molecular orbitals with $e_{g}$ character of [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2} have $\sim 2\%$ Fe 4p character. The larger number of iron 4p components in [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2} are expected as it has less orthoaxiality, which allows the iron 4p to delocalize into the 3d-like orbitals. This has an substantial effect on the intensity of this feature due to the now allowed 1s to $e_{g}$ transitions in the Fe \text{II} 4p to $e_{g}$ electric dipole transitions. These features were well reproduced by the RAS calculations of the three iron carbene complexes (Fig. 4), which show a clear intensity difference in the peak originating from transitions into orbitals with $e_{g}$ character between [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2} and the Fe \text{III} complexes as can be seen in Fig. 4 and 5.

The 3d–4p orbital hybridization effect on the K pre-edge XAS has been well described through the RAS calculations. The integrated intensity for the calculated electric dipole contribution is $2.72 \times 10^{-4}$, $2.76 \times 10^{-6}$ and $2.79 \times 10^{-6}$ for [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2}, [Fe\text{III}(btz)\text{3}](PF_{6})\text{3} and [Fe\text{III}(phthmeimb)\text{2}](PF_{6})\text{2}, respectively. The large difference in the electric dipole transition intensity does explain the intensity difference for the measured metal K pre-edge between the Fe \text{II} and Fe \text{III} complexes in Fig. 5.

Considering that $\sim 2\%$ 4p characters give $2.72 \times 10^{-4}$ intensity for [Fe\text{II}(bpy)(btz)\text{2}](PF_{6})\text{2}, the 4p character of both Fe \text{III} complexes should be less than 0.2%, as was indeed found. At this level small variations originating from e.g. differences between the computationally optimized and the experimentally present structure, contributions of the solvent or of the limited basis set can have an unrealistically large effect. One would e.g. expect that the near perfect orthoaxiality of [Fe\text{III}(phthmeimb)\text{2}](PF_{6})\text{2} results in the smallest dipole contribution. Conversely, if the contribution of 4p character is larger than 0.5% this is significant and differences at this level can be used to draw conclusion about symmetry effects. Future extensions and experience with this novel approach will increase the precision.

Here the relative contribution from electric dipole transitions beyond this limitation can hence be used to estimate the distortion from the centrosymmetry in the K edge measurements of octahedral complexes.

4.3 Discussion

The K pre-edge intensities depend on the metal 3d character in the probed valence orbitals, and can thus be used to evaluate
For FeIII, the peak with eg character is significantly broader due to the extra hole in the t2g orbital. After electron excitation into the empty eg orbital, the orbital occupancy of t2ge1 can give 3T1g, 2T2g, 1T1g and 1T2g valence states due to the coupling of the holes in both t2g and eg orbitals. The 3T1g and 1T2g triplets imply two electrons with parallel spin in the valence shell, the RAS calculations allow for multiple excitation within the active space and hence can give a nice description of the electron-electron interactions in the valence shell. Thus the broad feature splits into two main resonances at 7114.3 eV and 7115.9 eV for [FeIII(btz)3]3+, at 7114.3 eV and 7116.2 eV for [FeIII(phtmeimb)2]3+. The calculated spectra gave a splitting energy of 2.0 eV for [FeIII(btz)3]3+, and 2.1 eV for [FeIII(phtmeimb)2]3+, which is slightly larger than was experimentally observed. RAS calculations need an increasing number of core-excited states to fully describe the spectral features at the high energy side. Beyond a certain limit, an increasing of the number of included states only slightly shifts the observed averaged spectral features, which is taken as the cutoff point for the calculations (see ESI† for an estimation of this effect). An increase of this cutoff as well as an increase of the basis set can thus theoretically result in small additional shifts of the spectrum at the price of a significantly increased computational cost. For such large molecules as are studied here, the here employed combination of number of states and basis set is currently at the limit of our computational capabilities.

The difference in splitting between the T1g and T2g states probe the difference in attraction between the hole in the π(t2g) orbital and the electron in the σ(3d0g) compared to the σ(3dz2). The resonance at higher energy 7115.9 eV and 7116.2 eV is assigned to the 3T2g state, and it shows that the splitting of these two resonances depends on the orbital covalency.17,19,20 Considering the lower energy t2g peak for [FeIII(phtmeimb)2]3+, we can conclude that the splitting between the t2g and eg peak is larger for [FeIII(phtmeimb)2]3+. This was expected since the orbital covalency and centrosymmetry of the complexes. The metal K pre-edge analyses in terms of the chemically intuitive molecular orbitals and their occupation numbers are the starting point for the comparison. The differences in occupation numbers between the ground state and each final state allows for an interpretation of the spectrum in terms of orbital excitations.20,47 For the iron carbene molecules discussed here the contributions of individual orbitals are shown in Fig. 5.

The first absorption peak for both FeIII complexes originates from the 1s → t2g transition, their integrated intensities from calculated spectra are 0.58 × 10−4 and 0.48 × 10−4 for [FeIII(btz)3]3+ and [FeIII(phtmeimb)2]3+, which is consistent with the amount of t2g character of the states (see Table 1). This orbital covalency analysis shows that the metal eg character of the molecular orbital for the three complexes follows the order [FeIII(bpy)(btz)2]2+ > [FeIII(phtmeimb)2]1+ > [FeIII(btz)3]3+ (see Table 1). The integrated intensities of electronic quadrupole transitions for the 1s → eg are 2.41 × 10−4, 2.25 × 10−4, and 2.15 × 10−4 for [FeIII(bpy)(btz)2]2+, [FeIII(phtmeimb)2]1+ and [FeIII(btz)3]3+, respectively. The consistent trend in intensity and the metal 3d characters in the probed orbitals shows that the molecular orbital based RAS approach can well interpret the XAS in terms of the orbital covalency.

For FeII, the peak with eg character is significantly broader due to the extra hole in the t2g orbital. After electron excitation into the empty eg orbital, the orbital occupancy of t2ge1 can give 3T1g, 2T2g, 1T1g and 1T2g valence states due to the coupling of the holes in both t2g and eg orbitals. The 3T1g and 1T2g triplets imply two electrons with parallel spin in the valence shell, the consistent trend in intensity and 2.15 × 10−4 for [FeIII(bpy)(btz)2]2+, [FeIII(btz)3]3+, respectively. The integrated intensities of electronic quadrupole transitions for the 1s → eg are 2.41 × 10−4, 2.25 × 10−4, and 2.15 × 10−4 for [FeIII(bpy)(btz)2]2+, [FeIII(btz)3]3+, and [FeIII(phtmeimb)2]3+, respectively. The consistent trend in intensity and the metal 3d characters in the probed orbitals shows that the molecular orbital based RAS approach can well interpret the XAS in terms of the orbital covalency.

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the adoption of [phtmeimb]⁻⁻ ligand for a more pronounced σ donating ability can further increase the relative energy of eg metal 3d character anti-bonding orbitals, which gives a larger ligand field splitting. Previous calculations have shown that the minimum energies of MC and MC states of [FeIII(phtmeimb)]⁺⁺⁺ are destabilized by 13% and 23% compared to the ones of the [FeIII(btz)]⁺⁺⁺. The 2LMCT state of [FeIII(phtmeimb)]⁺⁺⁺ (2.13 eV) is 0.17 eV lower than the [Fe III(btz)]3⁺ (2.30 eV). The increased energies of the MC states together with the decreased energy of the 2LMCT state gives an increased activation barrier for the decay of the 2LMCT state into the MC states, and consequently contributes to the increase in the experimentally observed lifetime of the 2LMCT state. The RAS calculated spectra have well reproduced the larger splitting between the 2g and the first resonance of the eg peak for [FeIII(phtmeimb)]⁺⁺⁺, see Fig. 5, and show that this method can be used to predict the influence of new ligand systems on the energetic landscape of novel molecules. This means that we can both measure the relative energies of the valence orbitals with HERFD-XANES or calculate them prior to the synthesis with RAS and can expect to achieve an at least qualitatively good agreement even if the centrosymmetry is significantly changed. With further improvements in calculation power and code optimization this tool can potentially be extended to be used on real photocatalytical systems.

5 Conclusions

The ground state characterization of three iron complexes, of [FeII(bpy)(btz)]²⁺⁺, [FeIII(btz)]³⁺⁺ and [FeIII(phtmeimb)]³⁺⁺ have been carried out through the combination of experimental measurements of HERFD-XANES and were combined with ab initio RAS simulations of the K pre-edge XAS. We observed a large intensity difference in the FeII and FeIII experimental HERFD-XANES and interpreted them by differences of the electric dipole contributions originating in a change of state mixing, covalency changes and symmetry changes. The multiplet effect features in the HERFD-XANES of FeII complexes were resolved and nicely described through the RAS calculations. We have shown that the estimation of the electric dipole contribution and quadrupole contributions can be used to probe the distortion from centrosymmetry of these hexacoordinated complexes. The slightly low-lying 2g and the relatively destabilized eg orbitals of [FeIII(phtmeimb)]⁺⁺⁺ were reproduced and their originating chemical differences identified. This nicely shows that RAS can be used as an predictive as well as an analytic tool even in these large complexes and can be verified with HERFD-XANES. We expect that by using this tool we can predict the effect of new ligand systems on the complex electronic environment of next generation of iron carbene complexes.

Conflicts of interest

There are no conflicts to declare.

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