Supporting Information

Probing the Oxidation State of Transition Metal Complexes: A Case Study on How Charge and Spin Densities Determine Mn L-Edge X-ray Absorption Energies

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**X-ray Dose Accumulated in a Probed Sample Volume**

X-ray dose is the ratio of absorbed energy and absorbing mass. For flowing solution samples the choice of the absorbing volume and thus mass is often ambiguous. In order to provide an upper limit for the dose absorbed under our experimental conditions we herein calculate the “skin dose” (see refs. 22,40 in the main paper). This is the dose absorbed by a sample volume spanned by the surface area that exposed to the x-ray beam and the attenuation length $\Lambda$ of the sample (here $\Lambda=1.44$ µm for ethanol and $\Lambda=1.14$ µm for acetylacetone at 640 eV photon energy). We experimentally determined the horizontal liquid jet diameter to $d_{jet}=20\pm4$ µm and $22\pm4$ µm for the Mn$^{II}$(acac)$_2$ and Mn$^{III}$(acac)$_3$ solution samples, respectively, the x-ray focus size to $H\times V=100\times90$ µm$^2$ (horizontal x vertical, FWHM), and the beamline flux to $\Phi=(5.4\pm0.1)x10^{12}$ photons/s at $h\nu=640$ eV photon energy. Only a portion $\eta=d_{jet}/H$ of the photons hit the sample due to the smaller horizontal size of the liquid jet as compared to the horizontal focus size. For the Mn$^{II}$(acac)$_2$ and Mn$^{III}$(acac)$_3$ solution samples we measured sample flow rates $F$ of $8.5\pm1.5$ µl/min and $12\pm0.1$ µl/min, respectively, corresponding to jet velocities $v_{jet}=4F/(\pi\cdot d_{jet}^2)=0.45\pm0.20$ m/s and $0.53\pm0.20$ m/s and illumination times $t=V/v_{jet}=200\pm90$ µs and 170±70 µs, respectively. The resulting skin dose $D_s$ is calculated via

$$D_s = \frac{\Phi \cdot \eta \cdot h\nu \cdot t \cdot (1 - 1/e) \cdot 0.761}{\rho \cdot \Lambda \cdot d_{jet} \cdot V} = \frac{\Phi \cdot h\nu \cdot d_{jet}^2 \cdot \pi \cdot (1 - 1/e) \cdot 0.761}{4 \cdot F \cdot H \cdot \rho \cdot \Lambda}$$

where we used the above defined identities and where the factors $(1-1/e)$ and 0.761 account for the portions of photons absorbed within $\Lambda$ and contained within the vertical focus size $V$ (FWHM), respectively. With $\rho$(ethanol)=$789$ kg/m$^3$ and $\rho$(acetylacetone)=$980$ kg/m$^3$ the skin doses estimated for solution samples of Mn$^{II}$(acac)$_2$ in ethanol and Mn$^{III}$(acac)$_3$ in acetylacetone are $5200\pm2300$ Gy and $4600\pm1700$ Gy, respectively (1 Gy= 1 J/kg).
Structures and Orbitals of Mn\textsuperscript{II}(acac)\textsubscript{2}, Mn\textsuperscript{III}(acac)\textsubscript{3}, and (Mn\textsuperscript{II}(acac))\textsuperscript{1-}

For the approximately tetrahedral (T\textsubscript{d}) Mn\textsuperscript{II}(acac)\textsubscript{2} complex we find for the four Mn-O bonds, Mn to O distances of 2.06 Å (Figure 1 (a) in the main paper). The point group is D\textsubscript{2d} but as MOLCAS only handles Abelian point groups, the subgroup C\textsubscript{2v} is used for all calculations and for labeling the orbitals. The six-coordinated Mn\textsuperscript{III}(acac)\textsubscript{3} complex has C\textsubscript{2} symmetry and the octahedral ligand environment is affected by a strong Jahn-Teller distortion (refs. Forman (1959) and Stults (1979) in this document) leading to approximate D\textsubscript{4h} symmetry (Figure 1 (b) in the main paper).

Compared to the published x-ray structure of Mn\textsuperscript{III}(acac)\textsubscript{3} (ref. Stults (1979) in this document), the deviation between Mn-O distances is only 0.01 Å, except along the Jahn-Teller axis where the distance is overestimated by 0.05 Å. For Mn\textsuperscript{II}(acac)\textsubscript{2} we are not aware of any crystallographic data since the compound crystallizes as Mn\textsuperscript{II}(acac)\textsubscript{2} trimers (ref. Shibata (1984) in this document) resulting in a six-coordination of each Mn with Mn-O bond lengths between 2.1 and 2.2 Å. The Mn-O bond length of 2.06 Å from DFT optimization for Mn\textsuperscript{II}(acac)\textsubscript{2} (Figure 1 (a) in the main paper) is consistent with this. The difference in Mn-O bond lengths of Mn\textsuperscript{II}(acac)\textsubscript{2} and Mn\textsuperscript{III}(acac)\textsubscript{3} as well as the Jahn-Teller distortion of Mn\textsuperscript{III}(acac)\textsubscript{3} are also reflected in the results of infrared (IR) spectroscopy, published in refs. Forman (1959) and Shapkin (2006).
Decomposition of the RAS Spectra into Orbital Components

Figure S2. Calculated RAS absorption spectra (XAS not PFY-XAS) decomposed according to the relative orbital contributions in the final core-excited states for (a) Mn\textsuperscript{II}(acac)\textsubscript{2} (T\textsubscript{d} labels) and (b) Mn\textsuperscript{III}(acac)\textsubscript{3} (O\textsubscript{h} labels).

In Mn\textsuperscript{II}(acac)\textsubscript{2} from the sextet e\textsuperscript{2}t\textsubscript{2}\textsuperscript{3} (T\textsubscript{d}) ground state, excitations to both e and t\textsubscript{2} orbitals contribute with similar relative intensity over almost the entire range of the spectrum (Figure S2 (a)). The one-electron picture of excitations to distinct orbitals is consequently not appropriate for interpreting individual features. The orbital analysis of Mn\textsuperscript{III}(acac)\textsubscript{3} (t\textsubscript{2g}\textsuperscript{3}e\textsubscript{g}\textsuperscript{1} in O\textsubscript{h}) looks different (exclusive excitation to e\textsubscript{g} and partial depopulation of t\textsubscript{2g}, Figure S2 (b)), but the conclusion is very similar: Strong mixing of different electron configurations in the final states gives close to unity occupation numbers for all orbitals over the entire spectral range, which corresponds to a change in occupation number of zero for all orbitals except for the initially unoccupied e\textsubscript{g} (b\textsubscript{1g} or d\textsubscript{x2-y2}, see Figure 1 (b) of the main paper) orbital in Mn\textsuperscript{III}(acac)\textsubscript{3} where it is unity.

Spin Multiplicity Analysis of (Mn\textsuperscript{II}(acac)\textsubscript{3})\textsuperscript{1-}

We further plot in Figure S3 the calculated RAS spectrum of (Mn\textsuperscript{II}(acac)\textsubscript{3})\textsuperscript{1-} decomposed according to the spin multiplicity components sextet, quartet, and doublet. As could be expected, the spin multiplicity analysis of the (Mn\textsuperscript{II}(acac)\textsubscript{3})\textsuperscript{1-} RAS spectrum also gives very similar results as for Mn\textsuperscript{II}(acac)\textsubscript{2} (Figure 4 (c) in the main paper).

Figure S3. Contribution of (spin) multiplicities 2S+1 in the core-excited final XAS states for (Mn\textsuperscript{II}(acac)\textsubscript{3})\textsuperscript{1-} as obtained from the RAS calculations. The absolute photon energies of the spectrum are calibrated as in the main paper.
Initial Ground-State and Core-Excited State Configurations for Mn$^{II}$ and Mn$^{III}$ with $\Delta S=0$ within the LS Coupling Scheme and Number of Coulomb and Exchange Integrals

Table S1. Electron spin configurations and number of Coulomb (J) and exchange (K) integrals of the Mn$^{II}$ 3d$^5$ and Mn$^{III}$ 3d$^4$ initial ground state configurations and the Mn$^{II}$ 2p$^5$3d$^6$ and Mn$^{III}$ 2p$^5$3d$^5$ final core-excited state configurations (black: paired spins, red: unpaired spins).

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<th>Mn($^{III}$)</th>
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<td>2p$^5$ 3d$^5$</td>
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Slater Integrals

To quantify the energies of different configurations we use Coulomb (J) and exchange (K) integrals, where the total interaction between electrons of opposite spin is J, and the interaction between electrons of parallel spin is (J-K). As the effect of the ligand environment is small for the complexes studied here, it is reasonable to start with an analysis based on atomic theory, in which case the integrals J and K can be derived from the Slater integrals for direct and exchange Coulomb interactions, $F_k$ and $G_k$ (ref. Slater (1929)) in the terminology of R. D. Cowan (ref. 78 of the main text). In Table S2 we list the average J and K integrals for ionic Mn$^{II}$ and Mn$^{III}$ initial ground and Mn$^{II}$ 2p$^5$3d$^6$ and Mn$^{III}$ 2p$^5$3d$^5$ final core-excited states for respective p (2p) and d (3d) electron interactions (the corresponding Slater integrals were also calculated and are identical to the values reported in ref. 26 of the main paper).
Table S2. Calculated energies of electron-electron interactions for Coulomb (J) and exchange (K) integrals for p (2p) and d (3d) electrons relevant to L-edge XAS of MnII and MnIII for initial ground states (IS) and final core-excited states (FS). Averaged values of J and K integrals are given where averages were performed over all pairs of possible interactions in (d,d) and (p,d) orbitals.

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<th>MnIII IS</th>
<th>MnII FS</th>
<th>MnIII FS</th>
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<tr>
<td>J(d,d)</td>
<td>22.56</td>
<td>24.43</td>
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<td>J(p,d)</td>
<td>31.34</td>
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<td>K(d,d)</td>
<td>0.85</td>
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<td>K(p,d)</td>
<td>0.70</td>
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The J and K integral values in Table S2 represent averages over interactions of electrons in all different tuples of the different orbitals spreading over ranges up to 9.6% (J(d, d)), 4.5% (J(p, d)), 65% (K(d, d)) and 162% (K(p, d)), relative to these average values. For example, in the excited states of MnII (MnIII) the J integrals spread from 23.3 to 25.6 eV (24.9 to 27.4 eV) for J(d, d) and from 32.9 to 34.4 eV (34.7 to 36.3 eV) for J(p, d), while the K integrals spread from 0.55 to 1.15 eV (0.61 to 1.26 eV) for K(d, d) and from 0.16 to 1.52 eV (0.18 to 1.71 eV) for K(p, d).

The number of direct and exchange Coulomb interactions (number of J and K integrals) in the final core-excited states are given in Table S1. Together with the corresponding numbers in the initial ground states of the two systems (also given in Table S1) we can now calculate how the number of J and K integrals changes for going from initial to final states and for going from MnII to MnIII. For MnII the 2p-3d excitation process changes the number of exchange Coulomb interactions K(p, d) and K(d, d) by 2 and 0, respectively (see Table S1). For MnIII and upon 2p-3d excitation the number of K(p, d) and K(d, d) integrals as well changes by 2 and 0, respectively (ΔS=0 final states with parallel 2p and 3d spins). The net difference of changes in the number of exchange interactions between MnII and MnIII upon 2p-3d excitation is thus zero.

In MnII and upon 2p-3d excitation the number of direct Coulomb interactions J(p, d) and J(d, d) changes by 0 and 5, respectively, in (Table S1). For MnIII the number of J(p, d) and J(d, d) integrals changes by 1 and 4, respectively. The contribution to the shift when going from MnII to MnIII is then +1 J(p, d) and -1 J(d, d). As J(p, d)>J(d, d) (see Table S2) this qualitatively explains the L-edge XAS shift to higher energies for MnIII compared to MnII (for the ΔS=0 final states with parallel 2p and 3d spins).

Some of the observations from the RSD analysis can be directly related to the size of the Coulomb integrals. The smaller pairwise interactions in the initial ground state of MnII as compared to MnIII (see Table S2) is directly related to the expansion of the 3d shell upon the addition of an extra electron (Figure 5 (d) in the main paper). In the same way, the contraction of the 3d shell upon core excitation (Figure 9 (d) in the main paper) is consistent with the
larger interactions for the final core-excited states relative to the initial ground states (Table S2, and ref. 26 in the main paper).

**Comparison of Charge and Spin Density Differences for Mn\textsuperscript{II}(acac)\textsubscript{2} and (Mn\textsuperscript{II}(acac)\textsubscript{3})\textsuperscript{1-}**

![Graphs showing charge and spin density differences](image)

**Figure S4.** (a) Radial charge density (RCD) differences (RCD of core-excited final state minus RCD of ground initial state) of (Mn\textsuperscript{II}(acac)\textsubscript{3})\textsuperscript{1-}, Mn\textsuperscript{II}(acac)\textsubscript{2}, and Mn\textsuperscript{II}(acac)\textsubscript{3} (in units of one electron charge per Å, averaged over all core-excited states). (b) Differences of RCD differences from (a) between (Mn\textsuperscript{II}(acac)\textsubscript{3})\textsuperscript{1-} and Mn\textsuperscript{III}(acac)\textsubscript{2} and between Mn\textsuperscript{III}(acac)\textsubscript{2} and Mn\textsuperscript{III}(acac)\textsubscript{3}. (c) Integrated RCD differences (integrals of the RCD differences from (b), in units of one electron charge). (d) Radial spin density (RSD) differences (RSD of core-excited state minus RSD of ground state) of (Mn\textsuperscript{II}(acac)\textsubscript{3})\textsuperscript{1-}, Mn\textsuperscript{II}(acac)\textsubscript{2}, and Mn\textsuperscript{III}(acac)\textsubscript{3} (in units of one electron charge per Å, averaged over five selected core-excited states in the L-edge with \(\Delta S=0\), namely sextet for (Mn\textsuperscript{II}(acac)\textsubscript{3})\textsuperscript{1-} and Mn\textsuperscript{III}(acac)\textsubscript{2} and quintet for Mn\textsuperscript{III}(acac)\textsubscript{3}). (e) Differences of RSD differences from (d) between (Mn\textsuperscript{II}(acac)\textsubscript{3})\textsuperscript{1-} and Mn\textsuperscript{III}(acac)\textsubscript{2} and between Mn\textsuperscript{III}(acac)\textsubscript{2} and Mn\textsuperscript{III}(acac)\textsubscript{3}. (f) Integrated RSD differences (integral of the RSD differences from (e), in units of one electron charge). All properties were extracted from our RAS calculations (see Methods and Materials section in the main paper) and are plotted versus the radius of a sphere around Mn. The dashed vertical lines indicate the location of L and M shell maxima in the charge distributions, of the 2p and 3d shell maxima in the spin density distributions and approximately half the Mn-O bond length \(R_b\) (some methyl groups were replaced by H atoms, see Figure 1 of the main text, with negligible effect on the calculated charge density distributions).
References of the Supporting Information

S1  A. Forman and L. E. Orgel, Molecular Physics, 1959, 2, 362-366.


