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Editorial

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Perspectives

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DOI: 10.1039/b927120c

Advances in high brilliance energy dispersive X-ray absorption spectroscopy
Sakura Pascarelli and Olivier Mathon, Phys. Chem. Chem. Phys., 2010
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Communication

$\mu$-XANES mapping of buried interfaces: pushing microbeam techniques to the nanoscale
Paolo Ghigna, Sonia Pin, Giorgio Spinolo, Mark A. Newton, Michele Zema, Serena C. Tarantino, Giancarlo Capitani and Francesco Tatti, Phys. Chem. Chem. Phys., 2010
DOI: 10.1039/c000195c

Papers

L-edge XANES analysis of photoexcited metal complexes in solution
Renske M. van der Veen, Joshua J. Kas, Christopher J. Milne, Van-Thai Pham, Amal El Nahhas, Frederico A. Lima, Dimali A. Vithanage, John J. Rehr, Rafael Abela and Majed Chergui, Phys. Chem. Chem. Phys., 2010
DOI: 10.1039/b927033g

EXAFS as a tool to interrogate the size and shape of mono and bimetallic catalyst nanoparticles
Andrew M. Beale and Bert M. Weckhuysen, Phys. Chem. Chem. Phys., 2010
DOI: 10.1039/b925206a

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Matthias Bauer and Christoph Gastl, Phys. Chem. Chem. Phys., 2010
DOI: 10.1039/b926385c

Combined TPRx, in situ GISAXS and GIXAS studies of model semiconductor-supported platinum catalysts in the hydrogenation of ethene
DOI: 10.1039/b926493k

Near sulfur L-edge X-ray absorption spectra of methanethiol in isolation and adsorbed on a Au(111) surface: a theoretical study using the four-component static exchange approximation
Sebastien Villaume, Ulf Ekström, Henrik Ottersson and Patrick Norman, Phys. Chem. Chem. Phys., 2010
DOI: 10.1039/b926109e

Influence of additives in defining the active phase of the ethylene oxychlorination catalyst
DOI: 10.1039/b926502n

First-principles calculations of X-ray absorption spectra at the K-edge of 3d transition metals: an electronic structure analysis of the pre-edge
DOI: 10.1039/b926886c

First steps in combining modulation excitation spectroscopy with synchronous dispersive EXAFS/DRIFTS/mass spectrometry for in situ time resolved study of heterogeneous catalysts
Davide Ferri, M. Santosh Kumar, Ronny Wirz, Armin Eysler, Oxana Korsak, Paul Hug, Anke Weidenkaff and Mark A. Newton, Phys. Chem. Chem. Phys., 2010
DOI: 10.1039/b926886c

Novel opportunities for time-resolved absorption spectroscopy at the X-ray free electron laser
DOI: 10.1039/c003406a
Spatially resolved 3D micro-XANES by a confocal detection scheme
DOI: 10.1039/c004103n

Wavelet transform EXAFS analysis of mono- and dimolybdate model compounds and a Mo/HZSM-5 dehydroaromatization catalyst
DOI: 10.1039/b926474d

Electronic structure of alumina-supported monometallic Pt and bimetallic PtSn catalysts under hydrogen and carbon monoxide environment
DOI: 10.1039/c000403k

Determination of CO, H$_2$O and H coverage by XANES and EXAFS on Pt and Au during water gas shift reaction
DOI: 10.1039/c000240m

Complementarity between high-energy photoelectron and L-edge spectroscopy for probing the electronic structure of 5d transition metal catalysts
DOI: 10.1039/b926414k

In situ time-resolved DXAFS for the determination of kinetics of structural changes of H-ZSM-5-supported active Re-cluster catalyst in the direct phenol synthesis from benzene and O$_2$
Mizuki Tada, Yohei Uemura, Rajaram Bal, Yasuhiro Inada, Masaharu Nomura and Yasuhiro Iwasawa, Phys. Chem. Chem. Phys., 2010
DOI: 10.1039/c000843p

Sulfur poisoning mechanism of steam reforming catalysts: an X-ray absorption near edge structure (XANES) spectroscopic study
Yongsheng Chen, Chao Xie, Yan Li, Chunshan Song and Trudy B. Bolin, Phys. Chem. Chem. Phys., 2010
DOI: 10.1039/b925910b

Peroxide-like intermediate observed at hydrogen rich condition on Pt(111) after interaction with oxygen
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First-principles calculations of X-ray absorption spectra at the \(K\)-edge of \(3d\) transition metals: an electronic structure analysis of the pre-edge

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We first present an extended introduction of the various methods used to extract electronic and structural information from the \(K\) pre-edge X-ray absorption spectra of \(3d\) transition metal ions. The \(K\) pre-edge structure is then modelled for a selection of \(3d\) transition metal compounds and analyzed using first-principles calculations based on the density functional theory (DFT) in the local density approximation (LDA). The selected compounds under study are presented in an ascending order of electronic structure complexity, starting with the Ti \(K\)-edge of rutile and anatase, and finishing with the Fe \(K\)-edge of the cyanomet-myoglobin. In most cases, the calculations are compared to polarized experimental spectra. It is shown that DFT-LDA methods enable us to reproduce satisfactorily the experimental features and to understand the nature of the electronic transitions involved in the pre-edge region. The limiting aspects of such methods in modelling the core–hole electron interaction and the \(3d\) electron–electron repulsion are also pointed out.

1 Introduction and overview

Transition metal elements play a crucial role in coordination chemistry (catalysis, metalloproteins), Earth sciences (\(3d\) elements are major constituents of the Earth) and solid state physics (magnetic materials, superconductors, etc.). X-Ray absorption near-edge structure (XANES) spectroscopy, thanks to its chemical and orbital selectivity, is a powerful technique to obtain precise structural and electronic information of \(3d\) transition metal compounds. In particular, at the \(K\) edge of transition metal elements, some features, which probe the \(3d\) empty orbitals, arise in the pre-edge region. Thus, the pre-edge features of transition metals are related to the coordination number, oxidation state and spin state of the absorbing atom, and to the point symmetry of the absorbing atom site (see the recent review paper of Yamamoto\textsuperscript{1}). The latter point concerning the site symmetry is essential in the sense that the pre-edge features can be interpreted using group theory. Indeed, the analyses of the pre-edge features usually make use of the character table of the irreducible representations of the absorbing atom site symmetry point group (the more often, \(O_d\) and \(T_d\) point group symmetries are considered for a six-fold and four-fold coordinated absorbing atom, respectively, even if the absorbing atom site polyhedron is not regular). The various methods used for pre-edge analyses can be classified into two groups: the fingerprint approach and the calculations. The fingerprint approach consists of a comparison between the pre-edge spectrum of the material under study with the ones of reference model compounds, including eventually fitting procedures of the spectra by pseudo-Voigt functions.\textsuperscript{2–4} Three kinds of theoretical approaches to calculate the \(K\) pre-edge are distinguished: the multielectronic approach based on the Ligand Field Multiplet theory (LFM), the single-particle (or monoelectronic) approach based on the Density Functional Theory (DFT), and the many-body Green’s function methods. This paper is focused on single-particle calculations of the \(K\) pre-edge structure for \(3d\) transition metal bearing compounds. Before presenting the outline of the paper, we first draw up an overview of various pre-edge analyses using the methods mentioned above.

Overview

In Earth and environmental sciences, the fingerprint approach is widely used to determine the oxidation state of the probed \(3d\) element in complex minerals and natural/synthetic glasses. For instance, the oxidation state of Fe in synthetic and volcanic glasses has been investigated, by fitting the corresponding Fe \(K\) pre-edge spectra with those of reference compounds.\textsuperscript{5} Such an analysis is based on the 2 eV chemical shift existing between the pre-edge structure of the ferrous and ferric ions in the chosen reference compounds. According to Wilke et al.,\textsuperscript{3} the most useful characteristics of the Fe \(K\) pre-edge to determine Fe oxidation state and coordination number are the position of its centroid and its integrated intensity. By measuring the Fe \(K\) pre-edge of 30 model compounds, it has been established that the separation between the average pre-edge centroid positions for \(\text{Fe}^{2+}\) and \(\text{Fe}^{3+}\) is

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1.4 ± 0.1 eV. This result has been used to measure the average Fe oxidation state for mixed Fe$^{2+}$–Fe$^{3+}$ compounds, the pre-edge structure arising between the positions of that of Fe$^{2+}$ and Fe$^{3+}$. In such a way, the Fe$^{3+}$/Fe$^{2+}$ ratio has been estimated in more than ten minerals containing variable/unknown amounts of Fe$^{2+}$/Fe$^{3+}$ (with an accuracy of ±10 mol% in the oxidation state determination). More recently, an experimental XANES study of the oxidation state of chromium in a variety of Cr-bearing model compounds containing Cr$^{2+}$, Cr$^{3+}$, Cr$^{4+}$, Cr$^{5+}$, and Cr$^{6+}$, in which the Cr-site symmetry is $D_{5h}$, $O_h$, or $T_d$, has been reported. In particular, it is shown that the centroid position of the pre-edge feature is again a better indicator of the Cr valence than the edge position. As chromium, vanadium occurs in nature under various oxidation states (i.e., V$^{2+}$, V$^{3+}$, V$^{4+}$ and V$^{5+}$), and vanadium K pre-edge XANES spectra vary systematically with valence state and site symmetry. These V K pre-edge properties have been used to determine the valence state of vanadium in natural vanadiferrous titanomagnetites and the proportion of vanadium occurring as V$^{4+}$ has been calculated by comparison with octahedral V$^{3+}$ and V$^{4+}$.6,7 All the examples cited above attest to the interest of the fingerprint approach for the analysis of the pre-edge, at least in Earth and Environmental sciences. However, the pre-edge region contains much more precise information, the extraction of which requires calculations. Before introducing the theoretical approaches used to model the pre-edge, the different electronic transitions, which may occur in the K pre-edge region of transition metal elements have to be established. There are three kinds of electronic transitions used to describe the pre-edge features, as recently explained by de Groot: (i) local electric quadrupole transitions (E2), $1s \rightarrow 3d$; (ii) non-local electric dipole transition (E1), $1s \rightarrow p$, where the empty p states of the absorbing atom are hybridized with the empty 3d states of the nearest metal neighbours via the p empty states of the ligands; (iii) local electric dipole transition (E1), $1s \rightarrow p$, where the empty p states of the absorbing atom are hybridized with the empty 3d states of the absorbing atom. In the latter, the $p$–$d$ mixing is possible only if the absorbing atom site is not centrosymmetric or if the absorbing atom site centrosymmetry is broken by the atomic vibrations. The use of calculation tools enables to determine the contribution of each kind of electronic transitions to the pre-edge features. In any case, the 3d empty states in the transition metal elements of the compound under study are probed in the pre-edge region, either directly through E2 transitions, or indirectly through E1 transitions. The latter can be either local (intratrace) or non-local (intersite, off-site). Therefore, the final states probed in the pre-edge are rather localized states, where the electron–electron repulsion is supposed to be important. In such cases, the appropriate method to calculate the pre-edge is a priori the Ligand Field Multiplet (LFM) one. For instance, the remarkable work by Wester et al.9 shows that the K pre-edge features of iron ions can be interpreted in the LFM approach. In this study, a fitting procedure is first used to identify the pre-edge features. The allowed multielectronic final states are then determined using the LFM theory, and the relative energies and intensities of the pre-edge features are simulated using molecular orbitals calculations based on DFT. Thanks to this methodology, a detailed understanding of the Fe K pre-edge features of ferrous and ferric model complexes (comprising tetrahedral, octahedral, and square pyramidal Fe environment, as well as low/high spin states of Fe) has been obtained. However in ref. 9, the transition matrix elements are not calculated in the LFM framework. This has been done for the first time in the case of the Fe K pre-edge by Arrio et al.10 More precisely, the eigenstates of the ions and the absolute intensities of the E2 and E1 transitions involved in the K pre-edge of Fe$^{2+}$ and Fe$^{3+}$ in minerals, where the iron ions are either in octahedral or tetrahedral sites, have been calculated. The LFM calculated spectra of the 3d$^{n} \rightarrow 1s^{2}3d^{n+1}$ E2 transitions with a crystal field value $10Dq$ of 1.2 eV for $n = 0, \ldots, 9$ have been recently reported, and it has been shown that charge-transfer effects have a weak influence on the $1s \rightarrow 3d$ pre-edge spectral shapes.11

Single-particle methods are quite widely used to calculate 3d element K pre-edge spectra, although they are in principle not fully appropriate for such applications. The analysis of the pre-edge features can be carried out by both cross-section and local density of states calculations. Single-particle calculations are performed either in a cluster approach, or using periodic boundary conditions. A cluster approach using the finite difference method of the FDMNES code12 has been successfully used to calculate the Ti K pre-edge structure in TiO$_2$-rutile, providing a Mulliken-type electron population analysis of the titanium atoms (either excited or not).13 The cluster approach of the Feff code14 has been applied to the Mn K pre-edge, in order to distinguish the spectral signature of Mn$^{2+}$ from the one of Mn$^{3+}$ in Mn-bearing oxides.15 The calculations also helped to determine the E1 contribution arising from the metal–metal pairs, as previously described in point (ii), in order to include it in the baseline that is used to extract the pre-edge features. Indeed, such E1 contributions complicate the determination of the absorbing atom oxidation state and symmetry when using fingerprint fitting procedures.15 Similarly, Feff single-particle calculations have been performed at the Cr K-edge to support the experimental fingerprint analysis carried out on a large series of pre-edge spectra in Cr-bearing oxide minerals.4 Next to cluster approaches, band structure codes are often used to interpret the pre-edge features. Many band structure codes now include the explicit calculation of the absorption cross-section in the electric dipole and quadrupole approximations, and several applications can be found in the literature.16–21 Recently, a plane-wave DFT method including GGA + U has been developed for XANES calculations and successfully applied to the Ni K pre-edge polarized spectra.22 The method initially devoted to norm-conserving pseudopotentials was also extended to ultrasoft pseudopotentials.23 Since single-particle methods are formally dedicated to ground state calculations, many body Green’s function approaches have been developed in order to model electronic excitations.24 These approaches include time-dependent DFT (TD-DFT), $GW$ calculations and the Bethe–Salpeter equation. TD-DFT calculations have been performed at the V K-edge of V$_2$O$_5$,25 and at the Fe K-edge of a series of ten iron model complexes, giving a correct agreement with experiment in the pre-edge region.26 The Bethe–Salpeter approach has been
applied to the calculation of the Ti K pre-edge structure in rutile\textsuperscript{27} and in SrTiO\textsubscript{3}.\textsuperscript{28} Although promising, such methodologies still remain quite too challenging to be applied to complex materials.

**Outline of the paper**

In the following, we present various single-particle band structure calculations of the $K$ pre-edge structure of 3$d$ transition metals compounds. First, the computational details are given and then the results are presented using an ascending order of electronic interaction complexity. In the selected compounds, the 3$d$ elements all sit in a distorted octahedral environment. However, a regular octahedron with $O\text{h}$ symmetry is considered in a first approximation, so that the 3$d$ levels are split into two groups: the $t\text{\textsubscript{2g}}$ and the $e\text{\textsubscript{g}}$ orbitals. The various 3$d$ shell occupations studied in this paper are illustrated in Fig. 1. Five distinct cases of 3$d$ occupation are considered. First, the simplest case of the Ti K pre-edge in titanium oxides (rutile and anatase) will be presented. Indeed, since the formal charge of Ti in TiO\textsubscript{2} is + IV, there are no 3$d$ electrons (Fig. 1a), which provides an ideal situation for single-particle calculations. Second, two examples of Fe $K$ pre-edge where Fe is divalent (3$d^6$) in a low spin state are described. In this case, the $t\text{\textsubscript{2g}}$ orbitals are fully occupied and the $e\text{\textsubscript{g}}$ ones are empty (see Fig. 1b). Thus, the 3$d$ states probed by X-ray absorption spectroscopy are the $e\text{\textsubscript{g}}$ ones only. In such a case, the photoelectron is excited to a level where the multielectronic effects are expected to be weak enough to be treated within a single-particle framework. The compounds studied are pyrite, a mineral of formula FeS\textsubscript{2}, and a model molecule, which represents the environment of iron in carbonmonoxy–myoglobin (MbCO). Third, Cr K pre-edge single-particle calculations are presented for three insulating materials (MgAl\textsubscript{2}O\textsubscript{4} : Cr\textsuperscript{3+}, Be\textsubscript{2}Si\textsubscript{6}Al\textsubscript{2}O\textsubscript{18} : Cr\textsuperscript{3+} and $\alpha$-Al\textsubscript{2}O\textsubscript{3} : Cr\textsuperscript{3+}), in which Cr occurs as an impurity and has a + III oxidation state (3$d^3$). Here, the single-particle calculations need to be spin-polarized, since in an octahedral environment, the $t\text{\textsubscript{2g}}$ orbitals of Cr\textsuperscript{3+} are occupied by three electrons with parallel spins (see Fig. 1c). The last case, corresponding to low spin Fe\textsuperscript{3+} (3$d^5$) in Mb\textsubscript{2}CN, combines the difficulties of Cr\textsuperscript{3+} for the majority spin and of V\textsuperscript{3+} for the minority spin (Fig. 1e). Hence, this last application of single-particle calculation appears to be the most critical case among all the examples treated here. For each compound, we first give the electronic transitions expected in the pre-edge region, as expected from the absorbing atom site symmetry (and crystallographic structure). Then, we compare the results of the single-particle calculations with the experimental data and finally, we give the assignment of the pre-edge features.

2 Computational details

In the monoelectronic framework, the X-ray absorption cross-section has the following general expression:

$$
\sigma(\omega) = 4\pi^2\alpha\omega \sum_{f} \left| \langle \psi_f | H_{\text{exc}} | \psi_i \rangle \right|^2 \delta(E_i + \hbar \omega - E_f)
$$

where $\alpha$ is the fine-structure constant, $\hbar \omega$ the energy of the incoming photons, $| \psi_i \rangle$ the initial state with energy $E_i$, and $| \psi_f \rangle$ the final state with energy $E_f$. For the major
the final state with $E_f$. The interaction Hamiltonian $H_{\text{int}}$ is given by $\hat{e}\cdot \mathbf{r}$ in the electric dipole approximation, and by \( \frac{1}{2}\hat{e}\cdot \mathbf{k}\cdot \mathbf{r} \) in the electric quadrupole approximation, $\hat{e}$ and $\mathbf{k}$ being the polarization vector and the wave vector of the photon beam, respectively. In the absence of coupling between the electric dipole and quadrupole terms, the total absorption cross-section $\sigma(\omega)$ is simply the sum of electric dipole and electric quadrupole contributions, as follows:

$$
\sigma(\omega; \hat{e}, \mathbf{k}) = 4\pi^2 \omega a_{\text{D}} \sum_{i} \left( |(\psi_i, \hat{e}\cdot \mathbf{r}|\psi_i)|^2 \right)
$$

$$
+ \frac{1}{4} \left( |(\psi_i, \hat{e}\cdot \mathbf{k}\cdot \mathbf{r}|\psi_i)|^2 \right) \cdot \delta(E_i + \hbar \omega - E_f).
$$

(2)

This is the case if one uses exclusively linear polarization and if the system is time-reversal invariant or centrosymmetric.\(^{29}\) The electric dipole and quadrupole cross-sections show an angular dependence, i.e. variations of the spectral features are expected when the sample orientation varies with respect to the incident X-ray beam. The anisotropy of the cross-sections depends on the crystal point group of the investigated compound and the complete dependence of each point group has been given by Brouder.\(^{29}\) Note that in the case of cubic crystals, the electric dipole cross-section does not show any angular dependence.

For all the compounds except pyrite, the total cross section (eqn (2)) has been calculated within DFT codes using a plane-wave basis set, pseudopotentials and periodic boundary conditions. For pyrite, the total cross section has been calculated with the FDMNES code,\(^{12}\) using a self-consistent field (SCF) electronic potential determined within the Full-potential Linearized-Augmented plane-wave framework of the Wien2k code.\(^{30}\) In each case the calculation is performed in the two following steps: (i) the determination of the SCF charge density of a supercell including the 1$s$ core-hole on the absorbing atom, through the minimization of the total energy of the system, (ii) the computation of the absorption cross section, using the continued fraction method described in ref. 31 and 32 for the plane-wave DFT codes, and using the finite difference method for the FDMNES one.

For the minerals, the calculations were carried out from the crystallographic structures given in Table 1. For the doped minerals, the atomic positions were first relaxed in order to determine the structural modifications induced by the substitution of one aluminium atom by either one chromium in the case of Cr-doped spinel, or by one vanadium atom in the case of doped-grossular (tsavorite). The results of this relaxation step are not recalled here since they have been presented in details in ref. 21 and 33 for Cr-doped spinel and tsavorite, respectively. In the case of MbCO and MbCN, there was no need to consider the whole protein in the XANES calculation, since the mean free path of the core-electron ejected during the absorption process (i.e. the so-called photoelectron) does not exceed about ten Angström. Hence, the calculations have been performed from the 48 atom cluster shown in Fig. 2, embedded in a large cubic supercell. The cluster has been orientated within the cell in such a way that the Fe–N_{p} bonds are along the $x$ and $y$ axis, and that the Fe–N_{his} along the $z$ axis. The atomic positions are those given by Della Longa et al.\(^{34}\) for MbCO and by the 2JHO protein data bank (PDB) code for MbCN.\(^{35}\) The structural parameters of the Fe environment in both proteins are given in Table 2. In the case of MbCO, since the pre-edge appears to be sensitive to the bending of the Fe–C–O chain, two values of the $\beta$ angle were considered (Fig. 2): \(14^\circ\), as in ref. 34, and a smaller one equal to \(10^\circ\).

The plane-wave DFT codes used in this study were either the PARAlllel Total Energy Code\(^{36}\) for Cr-doped spinel and the Quantum-Espresso suite of codes\(^{37}\) (including the XSpectra package\(^{38}\)) for the other compounds. Troullier–Martins\(^{38}\) norm-conserving pseudopotentials in the Kleinman–Bylander form\(^{39}\) have been used. The parameters used for their generation are listed in Table 3. Note that the pseudopotential of the absorbing atom is built with only one 1$s$ electron in its electronic configuration. The computational details of the calculations (i.e., supercell size, approximation for the exchange and correlation density functional, energy cut-off, \(k\)-point grid, broadening parameter $\gamma$ used in the continued fraction formulation of the cross-section and $S_z$ value if needed) are given in Table 4. Local and partial density of states (lp-DOS) were calculated in the case of MbCO and MbCN, using Quantum-Espresso. The lp-DOS calculations, which took into account the 1$s$ core-hole on Fe, were calculated using Löwdin projections and a Gaussian broadening of 0.07 eV. Note that in the case of MbCO, they were obtained for $\beta = 14^\circ$. In the case of pyrite, the absorption cross section calculations have been performed for a 51 atom cluster. The SCF full-potential, generated using Wien2k, has been determined by minimizing the total energy of a \(2 \times 2 \times 2\) cubic supercell, including a 1$s$ core-hole on one of the 32 Fe atoms. The other parameters of the code were set to the default values.

**Fig. 2** Local environment of Fe in MbCO, within the cluster used for XANES calculation. It contains 48 atoms, including additional hydrogen atoms introduced to ensure the neutrality of the system. The nitrogen first neighbours of iron are labelled N_p for the four belonging to the heme plane and N_{his} for the one belonging to the H93 histidine residue. The angles $\alpha$ and $\beta$ are introduced to characterize the angular geometry of the chain Fe–C–O (or Fe–C–N in the case of MbCN). This Figure has been generated by XCrystDen.$^{54}$
Table 1 Description of the crystallographic structures of the minerals under study. The point group of the absorbing atom site is indicated. In the case of Cr-doped spinel and tsavorite, the Z value is the one of the corresponding undoped minerals.

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Chem. formula</th>
<th>Cryst. syst.</th>
<th>Space group</th>
<th>Abs. atom site</th>
<th>Cell param.</th>
<th>Z</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rutile35</td>
<td>TiO₂</td>
<td>Tetragonal</td>
<td>P₄₁/mmm</td>
<td>Ti (mmmm)</td>
<td>a = 4.5937 Å 2</td>
<td></td>
</tr>
<tr>
<td>Anatase55</td>
<td>TiO₂</td>
<td>Tetragonal</td>
<td>I₄₁/amd</td>
<td>Ti (4m2)</td>
<td>a = 3.7845 Å 4</td>
<td></td>
</tr>
<tr>
<td>Pyrite56</td>
<td>Fe₂S</td>
<td>Cubic</td>
<td>P₄₃</td>
<td>Fe (3)</td>
<td>a = 5.1175 Å 4</td>
<td></td>
</tr>
<tr>
<td>Cr-doped spinel57</td>
<td>Mg₆Al₂O₄·Cr³⁺</td>
<td>Cubic</td>
<td>F₄₃m</td>
<td>Cr (3m)</td>
<td>a = 8.0806 Å 8</td>
<td></td>
</tr>
<tr>
<td>Tsavorite38</td>
<td>Ca₃Al₂(SiO₄)₃·V³⁺</td>
<td>Cubic</td>
<td>I₄₃d</td>
<td>V (3)</td>
<td>a = 11.847 Å 8</td>
<td></td>
</tr>
</tbody>
</table>

Table 2 Structural characteristics of the local surrounding of Fe in the MbCO and MbCN clusters used for XANES calculation. Distances are in Å and angles in degrees.

<table>
<thead>
<tr>
<th>MbCX</th>
<th>dₑ₋ₐₓ</th>
<th>dₑ₋ₐₓ,ₐₓ,ₐₓ</th>
<th>dₛ₋ₐₓ</th>
<th>dₐₓ₋ₐₓ</th>
<th>α</th>
<th>β</th>
<th>PDB code</th>
</tr>
</thead>
<tbody>
<tr>
<td>MbCO₃4</td>
<td>2.00</td>
<td>2.06</td>
<td>1.83</td>
<td>1.07</td>
<td>6</td>
<td>14</td>
<td>/</td>
</tr>
<tr>
<td>MbCN₃5</td>
<td>2.04</td>
<td>2.08</td>
<td>1.92</td>
<td>1.11</td>
<td>4</td>
<td>13</td>
<td>2JHO</td>
</tr>
</tbody>
</table>

Table 3 Parameterization used for the generation of the norm-conserving Troullier–Martins pseudopotentials. The core radii of the valence states are indicated in parentheses in Bohr units. The Mg pseudopotential includes non-linear core-correction.

<table>
<thead>
<tr>
<th>Atom</th>
<th>Valence states</th>
<th>Local part</th>
</tr>
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<tbody>
<tr>
<td>H</td>
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</tr>
<tr>
<td>C</td>
<td>p</td>
<td></td>
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<tr>
<td>N</td>
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<tr>
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<td>p</td>
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<tr>
<td>Mg</td>
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<tr>
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<tr>
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<tr>
<td>Ca</td>
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<tr>
<td>Ti</td>
<td>d</td>
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<tr>
<td>V</td>
<td>s</td>
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<tr>
<td>Cr</td>
<td>s</td>
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<tr>
<td>Fe</td>
<td>p</td>
<td></td>
</tr>
</tbody>
</table>

3 Results and discussion

3.1 Ti K-edge

TiO₂-rutile. The tetragonal point group of rutile is \( \frac{1}{2} \)mm (D\(_{4h}\)), which yields a dichroic behaviour of the XANES in the electric dipole approximation.\(^{29}\) The point symmetry of the Ti site is mm\(2\) (D\(_{2h}\)), thus it is centrosymmetric. Consequently, local E2 transitions and non-local E1 transitions are expected to contribute to the pre-edge features.

Note that local E1 transitions due to the atomic vibrations are not formally excluded. This point will not be investigated here.

Fig. 3 shows the comparison of the calculated polarized spectra with the experimental ones of Poumellec et al.\(^{39}\) for \( \hat{e} \) parallel and perpendicular to the 4-fold symmetry axis (i.e., the \( z \) axis of the tetragonal cell), the wave vector being along the [110] direction. The theoretical spectra are calculated either with or without the core-hole. The decomposition into E1 and E2 contributions is also shown. For both polarizations, the pre-edge region exhibits three well defined features, labelled \( A_1, A_2 \) and \( A_3 \), which are reproduced by the single-particle calculations. For \( \hat{e} \parallel z \), the theoretical E2 contribution shows two peaks, which contribute to \( A_1 \) and \( A_2 \). The E1 contribution also shows two peaks, but they contribute to \( A_2 \) and \( A_3 \). Therefore, \( A_1 \) is a pure E2 peak, originating from \( 1s \rightarrow 3d \) local transitions. Peak \( A_3 \) is a pure E1 non-local peak, originating from \( 1s \rightarrow p \) transitions, where the \( p \) empty states of the absorbing atom are hybridized with empty 3\( d \) states of the Ti second neighbours. This hybridization is achieved via the empty \( p \) orbitals of the O first neighbours. Peak \( A_2 \) originates from both kinds of transitions (local E2 and non-local E1), the E2 contribution representing around 10% of the E1 + E2 sum. For \( \hat{e} \perp z \), the E1 contribution also exhibits two peaks, explaining the origin of peaks \( A_2 \) and \( A_3 \), while the E2 contribution only displays one peak, giving the origin of peak \( A_1 \).

Now the question is why the E2 contribution is characterized by two peaks for \( \hat{e} \parallel z \) and only one for \( \hat{e} \perp z \), the wave vector \( \mathbf{k} \) being parallel to [110]. To answer this question, one should consider the crystallographic structure of rutile. The tetragonal unit cell of rutile comprises two TiO\(_6\) octahedra, which are crystallographically equivalent but are differently orientated with respect to the photon beam (see Fig. 4). When
both $\mathbf{k}$ and $\hat{\mathbf{k}}$ are between the Ti–O bonds, the E2 transition probes the local $e_g$-like orbital. This is the case for $\hat{\mathbf{k}} \parallel z$ when the site 1 is excited. On the contrary, when one of the $\mathbf{k}$ or $\hat{\mathbf{k}}$ vectors is between the Ti–O bonds and the other along a Ti–O bond, the electronic transition probes a local orbital between the bonds, which is mainly $t_{2g}$-like. This case is encountered for $\hat{\mathbf{k}} \parallel z$ when the site 2 is excited, and for $\hat{\mathbf{k}} \perp z$ when both Ti sites are excited. Therefore, when $\hat{\mathbf{k}} \parallel z$ (and $\mathbf{k} \parallel [110]$), while the $d_{xz}$ and $d_{yz}$ states of the crystal are probed, the first E2 peak corresponds to a transition from the $1s$ level to the $t_{2g}$-like orbital belonging to the Ti site 2, and the second E2 peak to a transition to the $e_g$-like orbital belonging to the Ti site 1. When $\hat{\mathbf{k}} \perp z$ (and $\mathbf{k} \parallel [110]$), while the $d_{z^2}$ states of the crystal are probed, the E2 peak corresponds to transitions from the $1s$ orbitals to $t_{2g}$-like orbitals of both Ti sites. Following similar geometrical arguments, one can observe that for the non-local E1 contributions, the $t_{2g}$- and $e_g$-like orbitals of the Ti neighbours are indirectly probed at the energy positions of $A_2$ and $A_3$ peaks, respectively. The energy position of the $t_{2g}$ and $e_g$ orbitals is different for the Ti absorbing atom and for the Ti non-excited neighbours because the $1s$ core-hole attracts the $3d$ levels of only the Ti absorbing atom (the corresponding energy shift being about 2 eV). This result is supported by the calculation, which does not take into account the core-hole effects (see green curves in Fig. 3). Indeed, the E2 contributions probing the $t_{2g}$ and $e_g$ levels of the non-excited Ti are located at the positions of $A_2$ and $A_3$, respectively. The assignment of the three Ti $K$ pre-edge features of rutile for $\hat{\mathbf{k}} \parallel z$ and $\hat{\mathbf{k}} \perp z$ is summarized in Table 5. It is in total agreement with previous cluster calculations performed using the fitting approach of the FDMNES code.

![Fig. 3](image1.png)

**Fig. 3** Comparison between experimental (red line with circles) and calculated (solid line) Ti $K$ pre-edge spectra of rutile, for the $\mathbf{e} \parallel z$ (bottom) and $\mathbf{e} \perp z$ (top) experimental configurations. The theoretical spectra were calculated for a supercell including a core-hole (black line) or not (green lines), in order to show the effects of the presence of the $1s$ core-hole. The zero energy corresponds to the highest occupied level to the $t_{2g}$ band. The electronic eigenvalues for the Ti absorbing atom and for the Ti non-excited neighbours because the $1s$ core-hole attracts the $3d$ levels of only the Ti absorbing atom (the corresponding energy shift being about 2 eV). This result is supported by the calculation, which does not take into account the core-hole effects (see green curves in Fig. 3). Indeed, the E2 contributions probing the $t_{2g}$ and $e_g$ levels of the non-excited Ti are located at the positions of $A_2$ and $A_3$, respectively. The assignment of the three Ti $K$ pre-edge features of rutile for $\hat{\mathbf{k}} \parallel z$ and $\hat{\mathbf{k}} \perp z$ is summarized in Table 5. It is in total agreement with previous cluster calculations performed using the fitting approach of the FDMNES code.

![Fig. 4](image2.png)

**Fig. 4** Crystallographic structure of rutile, showing the orientation of the two equivalent TiO$_6$ octahedra in the unit cell with respect to the polarisation and wave vectors for the two experimental setups.

<table>
<thead>
<tr>
<th>Peak</th>
<th>Assignment for $\hat{\mathbf{k}} \parallel z$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A_1$</td>
<td>E2: 1s $\to$ 3d-$t_{2g}$ of Ti absorber</td>
</tr>
<tr>
<td>$A_2$</td>
<td>E1: 1s $\to$ $p_h$ hybrid. 3d-$t_{2g}$ of Ti neighb. + E2: 1s $\to$ 3d-$e_g$ of Ti absorber</td>
</tr>
<tr>
<td>$A_3$</td>
<td>E1: 1s $\to$ $p_h$ hybrid. 3d-$e_g$ of Ti neighb.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Peak</th>
<th>Assignment for $\hat{\mathbf{k}} \perp z$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A_1$</td>
<td>E2: 1s $\to$ 3d-$t_{2g}$ of Ti absorber</td>
</tr>
<tr>
<td>$A_2$</td>
<td>E1: 1s $\to$ $(p_x, p_y)$ hybrid. 3d-$t_{2g}$ of Ti neighb.</td>
</tr>
<tr>
<td>$A_3$</td>
<td>E1: 1s $\to$ $(p_x, p_y)$ hybrid. 3d-$e_g$ of Ti neighb.</td>
</tr>
</tbody>
</table>

Table 5 Assignment of the three $A_1$, $A_2$ and $A_3$ pre-edge features of the Ti $K$-XANES spectra of rutile for two distinct orientations of the incident photon beam polarization ($\hat{\mathbf{e}}$).
core-hole–electron interaction. The Ti K pre-edge of rutile has recently been calculated using a Bethe-Salpeter approach (BS). When the core-hole–electron interaction is included, the A1 peak appears at nearly the right position, and the overall agreement with experiment is satisfactory. Thus the BS approach provides a better description of the core-hole effects occurring in X-ray absorption. Second, the discrepancy with experiment concerns the shape of peak A2, which seems to include two components in the experimental data. According to the measurements of Le Fèvre et al., the first component would be an E2 one (1s → 3d-eg) and the second component, an E1 one (1s → p-hybridized with the t2g-shell of the Ti neighbours). In the calculations presented here, these two contributions are not resolved in energy, due again to the difficulty of single-electron approach to fully account for the core-hole–electron interaction. The last discrepancy is related to the intensity of peak A2, which is not high enough with respect to the experimental one. Atomic vibrations have been invoked to be at the origin of this discrepancy.

\( \text{TiO}_2\text{-anatase.} \) The crystallographic structure of anatase is tetragonal too, with the same point group as rutile. However the Ti site symmetry is \( 4m2(D_{2d}) \) and then is not centrosymmetric. Consequently, in addition to E2 and non-local E1 transitions, local E1 transitions are expected in the Ti K pre-edge region of anatase.

The Ti K pre-edge isotropic spectrum is compared to the experimental data of ref. 44. The results are shown in Fig. 5. The Ti K pre-edge of anatase is characterized by three peaks (A1, A2 and A3), like in rutile. However, unlike the case of rutile, the E1 part exhibits three contributions, due to the fact that local E1 transitions also occur. The E2 contribution displays two features: a 1s → 3d-\( t_2g \) well-resolved peak followed by a broad and weak 1s → 3d-\( e_g \) peak. The first E2 peak represents 22% of the total intensity of peak A1 and the second, 2% of the total intensity of peak A2. The electric quadrupole contribution is therefore weak compared to the electric dipole one, as expected in a non-centrosymmetric site. The assignment of the three pre-edge peaks is given in Table 6.

The agreement between theory and experiment obtained for anatase is equivalent to the one of rutile, i.e., it has two main drawbacks. First, the calculated A1 peak is at too high energy (of about 0.5 eV). Second, the A2 peak may contain two components, as suggested by Glatzel et al., which are not distinguishable in the simulation. These two components would come from the clear energy separation between the E2 and E1 contributions. These two drawbacks may have the same origin, i.e., the lack of DFT to properly model the core-hole–electron interaction. Indeed the second peak involves local and non-local transitions. The local ones are E1 and E2, thus involving the 3d-\( e_g \) shell of the Ti absorbing atom. Due to the misplacements of the 3d levels of the excited Ti, the 3d states of the absorbing Ti are located at too high an energy (i.e., the core-hole is not attractive enough).

### 3.2 Low-spin Fe\(^{2+}\) K-edge

**Carbonmonoxy-myoglobin (MbCO).** Myoglobin (Mb) is a single-chain globular protein of 153 amino acids, containing a heme, i.e., a iron-containing porphyrin. A proximal histidine (H93) group is attached directly to the iron centre, and a distal histidine group on the opposite face, not bonded to the iron. When a CO molecule is linked to Fe in the vicinity of the distal histidine, the Fe ion is in a divalent low spin state. The six coordination positions of Fe are occupied, i.e., the four nitrogen atoms in the heme plane (labelled \( \text{N}_p \)), the nitrogen atom belonging to the distal histidine (H93), and the carbon atom of the CO molecule (see Fig. 2). The protein is then called carbonmonoxymyoglobin (MbCO). MbCO crystallizes in a monoclinic lattice (space group \( P2_1 \)), with two molecules (about 1500 atoms) per unit cell.\(^{64}\) This Mb contains in the unit cell obviously are too far from each other to interact. Therefore, since the Fe site is non-centrosymmetric in an approximate 4 mm (\( C_4v \)) symmetry, two kinds of transitions are expected in the pre-edge: E2 and local E1.

Della Longa et al. have recorded angular-dependent spectra at the Fe K-edge of a single crystal of MbCO.\(^{48}\) The polarized spectra corresponding to the X-ray polarization parallel and normal to the heme plane have been deduced by neglecting the anisotropy in the heme plane.\(^{48}\) Fig. 6 (top panels) compares the polarized experimental spectra with the theoretical ones in the pre-edge region. A more complete study including the XANES features will be presented in a forthcoming publication.\(^{49}\) For both polarizations, the Fe K pre-edge region is characterized by two features A1 and A2, which are reproduced by the calculations. The E1 and E2 decomposition indicates that (i) in the case of \( \hat{\ell} \parallel \text{heme} \), peak A1 and peak A2 originate from E2 transitions and local E1 transitions, respectively. While the energy

**Table 6 Assignment of the three A1, A2 and A3 pre-edge features of the Ti K-XANES spectra of anatase.**

<table>
<thead>
<tr>
<th>Peak</th>
<th>Assignment</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1</td>
<td>E1: 1s → 3d-( t_2g ) of Ti absorber + E2: 1s → 3d-( t_2g ) of Ti absorber</td>
</tr>
<tr>
<td>A2</td>
<td>E1: 1s → p-( e_g ) of Ti absorber + E1: 1s → p hybrid. 3d-( t_2g ) of Ti neighb. + E2: 1s → 3d-( e_g ) of Ti absorber</td>
</tr>
<tr>
<td>A3</td>
<td>E1: 1s → p, hybrid. 3d-( e_g ) of Ti neighb.</td>
</tr>
</tbody>
</table>

**Fig. 5** Comparison between experimental (red line with circles) and calculated (black solid line) Ti K pre-edge isotropic spectra of anatase. The calculated spectrum is the sum of the E1 and E2 contributions, which are also shown.
separation between $A_1$ and $A_2$ is satisfactorily calculated (with an error of about 0.2 eV), the theoretical pre-edge is too high in energy with respect to the experimental one: the shift is about 1.4 eV for the $\hat{e} \parallel \text{heme}$ configuration and 0.9 eV for the $\hat{e} \perp \text{heme}$ configuration. Again, we observe that when the 3d orbitals of the absorbing atom are involved in the origin of the pre-edge features, the latter occur at a too high energy, when calculated in a single-particle DFT approach.

In order to further analyze the origin of the two features, the lp-DOS are also shown in Fig. 6 (bottom panels). The lp-DOS plots enable to visualise the reason why the E2 transitions are negligible when $\hat{e} \perp \text{heme}$ and significant when $\hat{e} \parallel \text{heme}$. Indeed, when $\hat{e} \perp \text{heme}$ (i.e. $\hat{e} \parallel [001]$ and $\mathbf{k} \parallel [110]$), only the $t_{2g}$ orbitals can be probed. Since they are mostly occupied, the resulting E2 absorption cross-section is found very weak. On the contrary, when $\hat{e} \parallel \text{heme}$ (i.e. $\hat{e} \parallel [110]$ and $\mathbf{k} \parallel [110]$), the empty $e_g$ orbitals are probed, thus leading to one peak in the E2 absorption cross-section. The lp-DOS plots also enable to reveal the various hybridizations between the orbitals, which are involved in the pre-edge. Since MbCO is a molecule, these hybridizations can be described in terms of molecular orbitals (MO). Let us first define the MO of the iron ligands, which are involved in the pre-edge. There are two axial ligands: the carbon atom of the CO molecule and the nitrogen atom of the H93 histidine group. The unoccupied MO orbitals of the CO molecule are $\pi^*$ and $\sigma^*$, partially defined by the C 2$p_x, p_y$ empty states and the C 2$p_z$ empty states, respectively. One of the antibonding $\pi^*$ MO of the histidine cycle is notably defined by the N$_{his}$-2$p_z$ empty states. In the heme plane, the four ligands are the N$_a$ atoms, and it is the $\pi^*$ MO of the porphyrin that participates in the pre-edge feature. The latter MO is partially described by the four N$_e^*$($2p_x, 2p_y$) empty states. Then the ligand MOs are hybridized with the 3d empty states of iron to form other antibonding MO, which are at the origin of the pre-edge features. In particular, it can be seen in Fig. 6 (left) that the $A_2$ peak of the $\hat{e} \perp \text{heme}$ configuration is due to E1 transitions $1^s \rightarrow 4p_{z}$ where the Fe-4$p_z$ empty states are hybridized with the MOs formed by the $t_{2g}$ $3d_{x^2-y^2}$ and $-3d_{xz}$, and the $\pi^*$ coming from CO. Consequently, the more the CO bond is tilted, i.e. the greater is the $\beta$ angle, the more efficient is

![Fig. 6](image-url)
the hybridization. This explains why peak \( A_2 \) is more intense for \( \beta = 14^\circ \) than for \( \beta = 10^\circ \). Note that the 10° spectrum matches better with experiment. This result suggests that the 14°, resulting from the fitting procedure of the XANES spectrum performed by Della Longa et al.,\(^{34} \) is overestimated.

The assignment of the pre-edge features for both orientations of the polarization vector in terms of unoccupied MO is given in Table 7. The lp-DOS calculations presented here partially confirm and complete the tentative MO assignments of the pre-edge features given by Della Longa et al.\(^{48} \)

### FeS\(_2\)-pyrite

Pyrite crystallizes in the cubic system and iron atoms are octahedrally coordinated to sulfur atoms sitting in four equivalent sites with point symmetry \( \bar{3} (C_3) \). Thus the iron site exhibits an inversion centre, which means that local \( p-d \) hybridization is forbidden: local E1 transitions are not expected in the pre-edge region. Consequently, only E2 transitions and non local E1 transitions can occur in the pre-edge region (as in the case of rutile). Note that in the pyrite cubic cell, the four FeS\(_6\) octahedra are tilted from the cubic crystallographic axis by about 23°.

The angular dependence of the Fe K pre-edge region of pyrite has been measured in order to reveal the E2 transitions.\(^{29} \) Indeed, since the system is cubic, only the E2 transitions depend on the orientation of the crystal with respect to the X-ray beam, the E1 ones being isotropic.\(^{29} \) The X-ray linear natural dichroism (XNLD) was found to be around 0.5% of the edge jump. Experimental results are compared with single-particle calculations in Fig. 7. A good agreement is obtained between theory and both the isotropic and XNLD spectra. The pre-edge is characterized by one main peak, containing two contributions, one E1 and one E2, as expected by symmetry considerations. The isotropic E2 contribution is about 4% of the pre-edge intensity and 0.7% of the expected by symmetry considerations. The isotropic E2 peak, containing two contributions, one E1 and one E2, as XNLD spectra. The pre-edge is characterized by one main peak.

### 3.3 Cr\(^{3+} \) K pre-edge

The K pre-edge of trivalent chromium is presented through three different examples of Cr-bearing minerals, where chromium substitutes for aluminium in octahedral position: Cr-doped spinel MgAl\(_2\)O\(_4\): Cr\(^{3+} \), emerald Be\(_2\)Si\(_2\)Al\(_2\)O\(_6\): Cr\(^{3+} \) and ruby z-Al\(_2\)O\(_3\): Cr\(^{3+} \). The amount of chromium being very low in these compounds (e.g., below a few atomic-percent) the probability to have chromium atoms in neighbouring sites is also low. This enables us to exclude the contribution of non-local E1 transitions at the K pre-edge. The number of expected E2 transitions can be predicted for a \( d^3 \) configuration in octahedral environment, as can be seen from Fig. 8. The ground state of Cr\(^{3+} \) corresponds to a configuration where the three lowest \( d \)-orbitals (the \( t_{2g} \)-like) are occupied by the majority spin. In the excited state, the photoelectron can probe on the one hand the empty \( t_{2g} \) orbitals for minority spin (case B in Fig. 8), and on the other hand the empty \( e_g \) ones, for both majority and minority spins (cases A and C in Fig. 8, respectively). Therefore, three spin-polarized E2 transitions can be expected at the Cr K pre-edge. The sequence from spinel to ruby via emerald corresponds to a decreasing symmetry of the Cr-site, starting from the \( D_{4h} \) point group symmetry in Cr-spinel, to \( D_3 \) in emerald and finally \( C_3 \) in ruby. As will be shown, the degree of admixture between the empty \( p \) states

<table>
<thead>
<tr>
<th>Peak</th>
<th>Assignment for ( \hat{e} \perp \text{heme} )</th>
</tr>
</thead>
</table>
| \( A_1 \) | E1: \( 1s \rightarrow 4p \) hybrid.  
with MO \( [3d^2_s + \pi^*(H93)]^* \) and  
with MO \( [3d^2_s + \sigma^*(CO)]^* \) |
| \( A_2 \) | E1: \( 1s \rightarrow 4p \) hybrid.  
with MO \( [3d^2_s - 3d^2_g + \pi^*(CO)]^* \) |

<table>
<thead>
<tr>
<th>Peak</th>
<th>Assignment for ( \hat{e} \parallel \text{heme} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( A_1 )</td>
<td>E2: ( 1s \rightarrow \text{MO} [3d_{2g,3g} + \pi^<em>(\text{porph})]^</em> )</td>
</tr>
</tbody>
</table>
| \( A_2 \) | E1: \( 1s \rightarrow 4p \rightarrow \text{hybrid.} \)  
with MO \( [3d_{2g}, 3d_{3g} + \pi^*(\text{CO})]^* \) |

Table 7 Main molecular orbital (MO) assignment of the \( A_1 \) and \( A_2 \) pre-edge features of the Fe K-XANES spectra of MnCO for two distinct orientations of the incident photon beam polarization (\( \hat{e} \)), as deduced from single-particle lp-DOS and absorption cross-section calculations.
and the empty 3d states of the absorbing atom is consequently increasing, which affects the intensity of the local E1 transitions contributing to the pre-edge.

**Cr-doped spinel MgAl₂O₄ : Cr³⁺.** Spinel (MgAl₂O₄) is a cubic crystal with an *Fd³m* space group. Trivalent chromium can substitute for aluminium in the octahedral site. Upon substitution, the octahedral site retains its 3*m* (*D₃d*) symmetry. Therefore, the chromium site exhibits an inversion symmetry: this implies that local E1 transitions cannot be expected and that the *K* pre-edge features should arise from E2 transitions only. In such a site symmetry, spherical tensor analysis predicts an angular dependence (XNLD) of the electric quadrupole absorption cross-section, when the absorption is measured on a single crystal. The full angular dependence was measured and modelled at the Cr *K* pre-edge by Julin *et al.*, in order to reveal quantitative information on the crystallographic and electronic structure of Cr³⁺.¹⁹

The experimental spectra measured for two different orientations are shown in Fig. 9 (left and right). For each of them, the upper panel shows the experimental spectrum and the theoretical one, with the E1 and E2 contributions plotted separately. Both panels show that the Cr *K* pre-edge has a pure electric quadrupole character, as expected, since the E1 contribution contributes only to the edge tail without showing any features. For the two orientations (*ξ*₁ = [010], *κ*₁ = [100] and *ξ*₂ = [1 1 0], *κ*₂ = [−1 −1 0], respectively), the correct number of peaks (one and two, respectively) is well reproduced by the single-particle calculation. For the first configuration (Fig. 9, left), the experimental spectrum shows a single peak labelled *A₁*. The electric quadrupole operator, expressed as *xy*, enables to probe the empty *d* electronic density, projected on Cr, in the *xy* direction, *i.e.*, between the Cr–O bonds. The corresponding empty *d* states have almost a pure *t₂g* character, and thus they can be reached only by a photoelectron with spin ↓ (Fig. 8, case B). This prediction is consistent with the calculation. For the second configuration (Fig. 9, right), two peaks *A₁* and *A₂* are obtained in the experimental pre-edge spectrum. For this orientation, the electric quadrupole operator is expressed as *z²−r²*, which enables to probe the 3*d* electronic density both in the *xy* and *z²−r²* directions. The corresponding *d* empty states are thus respectively *t₂g* and *e₅g* ones. For the *t₂g* component, one peak is expected for spin ↓, as in the first configuration; for the *e₅g* component, two peaks are expected, one for spin ↓ and one for spin ↑ (cases A and C in Fig. 8). These predictions are in line with the theoretical spectrum, which shows that peak *A₁* arises from transitions towards *e₅g* and *t₂g* states, while peak *A₂* corresponds to transitions towards *e₅g* states. It should be mentioned that, although the symmetry of the Cr-site is *D₃d*, no splitting between the empty *t₂g* orbitals is visible, which is an indication that the trigonal distortion is very small.

![Fig. 8 Schematic mono-electronic representation of the 3d orbitals of Cr³⁺ in *O₆* symmetry for the ground state and the possible final states.](image)

Although the agreement between theory and experiment is satisfactory enough to provide a clear vision of the assignment of the 1s–3*d* transitions occurring in the pre-edge, there are two main drawbacks. First, we observe again that in the calculation, the E2 pre-edge features occur at too high energy with respect to the edge, due to the modelling of the core-hole effects. Second, the absolute and the relative intensities are not perfectly reproduced using the single-particle approach, as well as the splitting of the two peaks, which is underestimated. This has been attributed to the fact that the interelectronic repulsion on the 3*d* levels of the Cr ion cannot fully be described in the LDA framework.¹⁹

**Emerald Be₃Si₆Al₂O₁₈ : Cr³⁺ and ruby α-Al₂O₃ : Cr³⁺.** As in the case of Cr-doped spinel, the Cr *K* pre-edge spectra measured on single crystals of emerald and ruby (with their three-fold axis perpendicular to the incident beam) display two features, *A₁* and *A₂* (see respectively Fig. 10 of ref. 20 and Fig. 3 of ref. 17). Single-particle calculations have revealed that both peaks are due to local E2 1s–3*d* transitions and local E1 1s–p transitions. The latter occur since the Cr-site is not centrosymmetric. Additionally, the amount of E1 transitions is higher in ruby, since the Cr site is more distorted than in emerald.²⁰ As beryl Be₃Si₆Al₂O₁₈ and corundum α-Al₂O₃ crystallize respectively in the *P6/mcc* (*D₆h*) and *R32/c* (*D₃2h*) space groups, both E1 and E2 transitions show an angular dependence. For both compounds, the XNLD has been measured and fully interpreted using single-particle calculations. Moreover, the transitions observed in the different polarized XANES spectra are in line with the predictions made through group theory, provided the character tables of the *D₃d* and *C₃* symmetry groups is used.

### 3.4 V³⁺ *K* pre-edge

The vanadium *K* pre-edge was studied through the example of tsavorite Ca₃Al₂(SiO₄)₃·2V³⁺, a V-bearing grossular. Tsavorite is a garnet and therefore crystallizes in the cubic system with space group *Ia₃d*. Vanadium is present as a substituted minor element in the Al site, with *3* (*C₃*) point symmetry. Hence, neither local nor non-local E1 transitions are expected: only E2 transitions contribute to the pre-edge region. These E2 transitions can probe *t₂g* and *e₅g* states whatever the spin is up or down. As explained in the case of spinel, the expression of the E2 cross-section predicts an angular dependence of the pre-edge when the crystal orientation is varied with respect to the incident X-ray beam.²⁹ Polarized XANES spectra at the V *K*-edge were recorded on an oriented single-crystal by...
The recorded orientations are identical to those measured for Cr-bearing spinel.

Bordage et al. The upper panels of Fig. 10 (left and right) compare the experimental and theoretical spectra obtained for the two orientations $\epsilon_a = [010]$; $k_a = [100]$ and $\epsilon_b = [1, 1, \sqrt{2}]$; $k_b = [-1, -1, \sqrt{2}]$. For both orientations, the pre-edge exhibits three well-defined structures, which are well reproduced by the single-particle calculations, but again at too high energy. The pure electric quadrupole character of the pre-edge is well-observed, since the $E_1$ contribution presents no structure in the pre-edge region and contributes only to the edge tail. When superimposing the spectra of each orientation, one notices that the expected angular dependence of the pre-edge is satisfactorily reproduced by the calculations. Indeed, both the relative energies and intensities are correctly calculated.

The lower panels of Fig. 10 (left and right) show the spin-polarized calculations of the $E_2$ cross-section performed for each experimental orientation. An assignment of the $E_2$ transitions involved in the experimental peaks, labelled $A_1$, $A_2$, and $A_3$, can be done within a single-particle view of the transitions from the $1s$ states to the empty $3d$ ones. Peak $A_1$ is thus attributed to transitions towards the $t_{2g}$ states. Peak $A_2$ arises from two contributions: transitions towards the $t_{2g}$ and $e_g^\dagger$ states. Peak $A_3$ is attributed to transitions towards the $e_g^\dagger$ states. The simple picture of the transitions involved in the V$^{3+}$ K-pre-edge in tsavorite enables to understand the spectral features observed on the experimental spectra. However, a more detailed attention must be paid to peak $A_1$ since it is situated astride the occupied and empty states. This is due to the $3d^2$ electronic configuration of V$^{3+}$: two $t_{2g}$ orbitals are occupied and one is empty. The occupied states represented in Fig. 10 (lower panels) correspond to virtual transitions towards the $t_{2g}$ orbital. The spectrum should thus display two well-separated peaks, the first in the occupied states and the second in the empty states. Nevertheless, the empty and occupied $t_{2g}$ orbitals are too close in energy for the single-particle DFT approach to reproduce their splitting into two separated components, as expected by the irreducible representations of the $C_{3i}$ vanadium site point group. Consequently, even if the agreement between experiment and calculation is quite satisfactory when the occupied states are cut, standard plane-wave DFT approach fails to properly model $3d$ incomplete spin-polarized shells. This drawback will be again illustrated with the following and last compound.

3.5 Low-spin Fe$^{3+}$ K-edge

Cyanomet-myoglobin (MbCN). The case of MbCN conjugates the difficulties of the two previous cases: the Cr K-pre-edge for the majority spin and the V K-pre-edge for the minority spin. Indeed the presence of CN in the sixth position of the Fe coordinates confers to the Mb protein a $S_z = 1/2$ spin state, with Fe in a trivalent low-spin state as schematized...
observed for peak $A$ transitions for shows that peak $A$ is due to E1 transitions for $\hat{e} \perp \text{heme}$ and to $e_g$ states, for both majority and minority spins.

The top panels of Fig. 11 (left and right) compare the experimental Fe K pre-edge polarized spectra of MbCN with cross-section calculations (for $\hat{e} \perp \text{heme}$ and $\hat{e} \parallel \text{heme}$, respectively). The spectra were recorded by Arcovito et al., applying a protocol similar to that used for MbCO. While the pre-edge spectra MbCO show two features, the experimental pre-edge spectra of MbCN are characterized by one main peak, labelled $A$, which is twice as more intense for $\hat{e} \perp \text{heme}$ than for $\hat{e} \parallel \text{heme}$. The shape and the anisotropy of the Fe K pre-edge of MbCN is well reproduced by the single-particle calculations (the fact that the calculated pre-edges are too intense with respect to experiment is essentially due to a too weak $\gamma$ broadening parameter used in the cross-section calculation). The decomposition into E1 and E2 contributions shows that peak $A$ is due to E1 transitions for $\hat{e} \perp \text{heme}$ and to E2 transitions for $\hat{e} \parallel \text{heme}$. The same decomposition has been observed for peak $A_1$ of MbCO. Since Fe lies in a nearly centrosymmetric environment within the heme plane, the absence of local E1 transitions for $\hat{e} \parallel \text{heme}$ then makes sense.

Although the expected number of pre-edge components is four (three E2 and one E1), one observes only one single broad feature in the XANES spectrum, which can be explained by the spin-polarized lp-DOS of Fe and of its neighbours (lower panels of Fig. 11). This is due to several reasons. First, because of the experimental configurations chosen, the transitions towards the partially empty $t_{2g}$ states cannot be probed. Second, as can be seen from the lp-DOS, no significant energy splitting is observed between minority and majority spins: the transitions to the empty $e_g$ states thus occur at a similar energy for both spins. Third, the local E1 transitions occur at an energy, which is close to the one of the E2 transitions (about 2.5 eV vs. 2.0 eV above the Fermi level, respectively).

These lp-DOS show that peak $A$ has the same assignment as peak $A_1$ of MbCO for both orientations. Indeed, the same MOs are involved at the energy of the calculated $A$ peak plotted in the top panels. Hence, for the $\hat{e} \perp \text{heme}$ orientation, peak $A$ is due to transitions $1s \rightarrow 4p_z$ where the $4p_z$ orbital is hybridized with the MOs $[3d_{x^2-y^2} + \sigma^*(\text{CN})]^*$ and $[3d_{z^2} + \pi^*(H93)]^*$ (the $\pi^*(\text{CN})$ and $\pi^*(H93)$ MO being displayed by the C $2p_z$ lp-DOS and the N$_{\text{his}}$ $2p_z$ lp-DOS). For the $\hat{e} \parallel \text{heme}$ orientation, peak $A$ is due to transitions $1s \rightarrow 3d_{x^2-y^2}$ where the Fe $3d_{x^2-y^2}$ orbital participates in the MO $[3d_{x^2-y^2} + \pi^*_{\text{porph}}]$ (the $\pi^*_{\text{porph}}$ MO is illustrated by the $2p_y$ and $2p_z$ partial DOS of the N atoms belonging to the heme plane).
Apart from the now well-known energy position problem of single-particle calculations, the pre-edge of MbCN appears not so critical to model (as initially expected). Nevertheless, these calculations reveal a failure similar to that observed in the case of tsavorite, i.e., the absence of energy splitting of the spin partially filled $t_{2g}$ shell. Indeed the Fermi energy, as calculated by the plane-wave DFT code, is found to be at the middle of the main peak of the spin $\uparrow$ Fe 3$d_{x^2-y^2}$ density of states (see the corresponding panels of Fig. 11). Fortunately, this problem has no incidence on the cross-section calculations because the Fe 3$d_{x^2-y^2}$ states are not probed in the orientation chosen in the experimental setup.

4. Conclusion

In this paper, we have presented a detailed investigation of the $K$ pre-edge of 3$d$ elements in several compounds. Through these examples, the issue of coupling closely experiment and theory to extract electronic and structural information was emphasized. In all the compounds investigated, single-particle calculations were able to reproduce satisfactorily the experimental pre-edge: the number of peaks was well reproduced, and the relative energy positions and relative intensities were in good agreement with the experimental data. This has allowed the interpretation of the experimental features within a monoelectronic view of the transitions involved in the $K$ pre-edge. The information that can be directly inferred from such an assignment are the electric dipole or electric quadrupole character of the transitions, as well as the degree of local and non-local orbital hybridization. By taking advantage of XNLD to probe specific empty orbitals, it is possible to draw a picture orientated in space. In all cases, the angular dependence of the $K$ pre-edge was indeed reproduced in a level, which was good enough to allow its interpretation. Moreover, if some $[\hat{n}, \mathbf{k}]$ orientations of the crystal with respect to the X-ray beam are not accessible experimentally, calculations are the only way to probe the corresponding empty levels. Within such a detailed analysis, the electronic structure of a transition metal

Fig. 11 Analysis of the MbCN pre-edge transitions for $\hat{\varepsilon}$ normal to the heme plane (left) and $\hat{\varepsilon}$ in the heme plane (right). Top panels: experimental and calculated pre-edge spectra (including the decomposition into E1 and E2 transitions). Bottom panels: local and partial spin-polarized DOS involved in the pre-edge region. The lp-DOS of majority spin (resp. minority spin) are displayed with positive values of states/eV/cell (resp. with negative values). The zero energy is the Fermi energy. The experimental spectra have been shifted in energy in order to make the main peak of the $\hat{\varepsilon} \perp$ heme polarized XANES experimental and theoretical spectra coincide.
ion in a given local environment can be well understood and therefore, its spectral signature well characterized.

However, single-particle calculations show some limitations. In the case of spin-polarized calculations for transition metal ions with incomplete d shells (V$^{3+}$ and LS Fe$^{3+}$), occupied and empty states are not well separated in the calculation. Such systems represent a real challenge for DFT. However, we point out that the assignment of the XANES features in terms of monoelectronic transitions still remains possible, provided that one keeps in mind that the 3d shells are incomplete. In all the compounds investigated, the calculation of K pre-edge spectra within DFT suffers from two main drawbacks, i.e., the modelling of the core-hole interaction, on one hand, and the 3d electron–electron repulsion, on the other hand.

First, for all the compounds presented in this paper, we found out that the E2 and local E1 transitions are systematically calculated at a too high energy with respect to the edge. This effect is due to the modelling of the 1s core-hole–electron interaction, which leads to an overestimation of the screening of the 1s core-hole. The relative energy positions of the pre-edge features can be improved mainly by two means. The simplest way consists of considering a core-hole with a positive charge superior to one, in order to increase artificially its attraction on the 3d empty states. Nevertheless, such calculations cannot be considered as ab initio anymore, since the value of the core-hole is a fitted parameter. An alternative, more elegant way is to consider a dynamic core-hole, instead of a static one as in the calculations presented here. This requires the Bethe–Salpeter formalism, which treats electron and hole dynamics ab initio, as well as electron–hole interactions. However, such calculations are nowadays still time-consuming, and complex systems like doped minerals and proteins are definitely challenging.

Second, the other main drawback of single-particle calculations is the modelling of electronic interactions: LDA and GGA approximations give indeed a description of these interactions in a mean-field way, which is not therefore completely satisfactory and which can be responsible for the possible differences in relative peak positions and intensities, compared to experiment. However, keeping in mind that the pre-edge features correspond to localized empty states where 3d–3d interactions are relevant, one must admit that DFT-LDA or DFT-GGA approaches already enable a good modelling of the angular dependence of the pre-edge. In the case of tsvorite, XNLD was very well-reproduced quantitatively. The less satisfactory agreement was observed for Cr-doped spinel, but it did not hamper the interpretation of the pre-edge. In certain cases, the description of 3d–3d electronic repulsion can be improved by performing LDA + U (or GGA + U) calculations. The Hubbard parameter U(3d) corresponds to the 3d Coulombian “on-site” repulsion and measures the spurious curvature of the energy functional as a function of occupation. The Hubbard parameter can be determined self-consistently using the Quantum-Espresso code, as an intrinsic linear-response property of the system. Nevertheless, DFT + U calculations cannot be performed for all systems, since the number of spins up and spins down needs to be non-zero. Only a few XANES calculations have been performed in DFT + U, i.e., Ni K-edge in NiO, Cu K-edge in CuO and La$_2$CuO$_4$ and Co K-edge in LiCoO$_2$. In these compounds, the addition of U, combined to the core-hole effects, has enabled to shift the local E2 transitions from the non-local E1 ones in the pre-edge, yielding a better agreement between calculations and experiments.

The way to take into account the many-body interactions lacking in DFT-LDA (i.e., the multi-Slater determinant nature of the electronic states) is to use the multiplet approach. For example, in the case of Cr-doped spinel, it has been shown that the angular dependence of the K pre-edge could be better modelled. However, this approach has also some drawbacks: (i) it uses a local approach, where a single transition metal ion is considered as embedded in a ligand field. Therefore, non-local E1 transitions occurring in the pre-edge cannot be calculated, (ii) because the calculation includes multiplet effects, a simple atomic picture is no longer possible to assign the transitions in terms of monoelectronic transitions, (iii) it uses some empirical parameters, which may not be available for all the systems. Hence, multiplet and single-particle methods must be considered as highly complementary. The development of approaches that go beyond DFT, such as DFT-CI (Configuration Interaction), TD-DFT and Bethe-Salpeter opens new prospects to draw a fully ab initio picture of the pre-edge structure. A nice success of TD-DFT is already illustrated by the case of K pre-edge of Fe$^{2+}$ and Fe$^{3+}$ in molecular model complexes. The application of these methods to more complex systems such as crystals requires developments, which are now under progress.

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5632 | Phys. Chem. Chem. Phys., 2010, 12, 5619–5633

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