Multiple excitations in the K fluorescence emission of Mn, Fe and Ni compounds

P. Glatzel*, U. Bergmann†, F. M. F. de Groot** and S. P. Cramer††

*University of California at Davis, USA
†Lawrence Berkeley National Laboratory, USA
**Universiteit Utrecht, The Netherlands

Abstract. The creation and the decay of a 1s vacancy can result in the excitation of a second electron. In this paper, two different modes of 1s core hole creation as a diagnostic tool to study multi-electron excitations in the K fluorescence emission are compared. The 1s core hole excited state can be created either by photoionization or by radioactive K capture decay. In the latter case, a 1s electron reacts with a proton in the nucleus to yield a neutron and an escaping electron neutrino. We report a comparison of Kβ spectra obtained from x-ray excitation in Mn and K capture in 55-Fe in various chemical environments. The Kβ1,3 main lines of the photoexcited spectra are broader than the corresponding lines obtained after K capture and the weak satellite lines at higher energies (Kβ2,5) differ in shape. A theoretical model on the basis of different electron relaxation depending on the mode of core-hole creation is presented.

INTRODUCTION

Any electronic transition in an atom causes a readjustment of the passive electrons, i.e. the electrons that are not directly involved in the transition, to the perturbed potential. A passive electron in an orbital \( \phi_{\mu} \) before the perturbation will relax into an orbital \( \phi_{\mu'} \). For an adiabatic relaxation we have \( \mu = \mu' \), i.e. the final state orbital is described by the same set of quantum numbers. If the relaxation is non-adiabatic, the electron will occupy an orbital with different quantum numbers (\( \mu \neq \mu' \)). Those transitions can be referred to as shake or multiple electronic transitions.

In a two step picture that can be used to describe fluorescence emission, the excitation of a second electron can either occur during the creation or during the decay of a 1s vacancy. Multi-electron transitions upon inner-shell vacancy creation in 3d transition metals have been studied using photoelectron [1] and absorption spectroscopy [2, 3, 4]. In order to obtain an estimate for the fluorescence energies due to transitions from doubly ionized excited states we can use a simple Z+1 model as shown in Figure 1. For Mn we expect the KLβ lines at about 70 eV above the Kβ main lines. As an example, the KLβ lines in MnO2 are shown. The KLβ peak intensity exhibits a slow rise over several keV as predicted by the Thomas model [5]. The spectra shown in Figure 1 are not corrected for self-absorption and we therefore do not attempt for a detailed analysis. A comprehensive treatment of the KLα lines in Cu can be found in reference [6, 7].

When the core hole is filled the excitation of a second electron can occur via a radiative Auger emission (RAE) where a photon is emitted and an electron is simultaneously
FIGURE 1. Left: Term scheme for Kβ and KLβ emission in Mn. Atomic configurations are used and only the partly occupied orbitals are given. An estimate for the KL-edge and the KL fluorescence energy is given based on a simple Z+1 model. Right: Kβ main and satellite lines. For the latter, the fluorescence emissions following photoexcitation below (solid line) and above (dashed line) the KL-edge are shown.

E(KL-edge) = E(Mn K-edge)+Fe(L-edge)
E(KLβ) = E(Mn KLβ)+[E(Fe L-edge)+E(Mn L-edge)]

FIGURE 2. Left: Term scheme of KLL radiative Auger emission in a 3d metal. Right: KLL edge in Ni metal and K2NiF6. The KLL edges differ between the metallic Ni and Ni(IV) in an ionic compound, i.e. they show a chemical dependence.
FIGURE 3. Left: K capture in $^{55}$Fe and 1s photoionization in Mn. Both processes result in an ion that has a Mn (Z=25) nucleus and a hole in the 1s shell. Right: Experimental Kβ spectra for MnO and $^{55}$Fe$_2$O$_3$. The instrumental broadening is $\approx$0.8 eV for both spectra. The spectra are normalized to each other in the Kβ$_{1,3}$ peak. The energy scale is the measured fluorescence energy and the spectra were not shifted in energy relative to each other.

COMPARING DIFFERENT MODES OF 1S CORE HOLE CREATION

Kβ Main Lines

As we have seen in the previous examples, doubly excited intermediate or final states yield sufficiently shifted (i.e. can be separated experimentally) K fluorescence lines if the second vacancy occurs in the L or K shell. In contrast, valence electron shake transitions will result in overlapping spectral features between singly and doubly fluorescence final states. In order to tackle this problem one can compare K fluorescence spectra following two different modes of 1s core hole creation [10, 11]. Provided that the two modes have different valence electron shake probabilities one expects different K fluorescence lines if shake transitions have considerable probability.

On the left side of Figure 3 it is shown that photoionization in Mn and radioactive electron capture decay from the K shell (K capture) lead to the formally identical fluorescence initial state with a Mn-55 nucleus and a hole in the 1s shell. We compare on the right side of Figure 3 the Kβ main lines, that arise from 3p to 1s transitions, for MnO and $^{55}$Fe$_2$O$_3$. Both compounds have a metal 3d$^5$ configuration in the ionic approximation. The striking difference between the two spectra is that the Kβ$_{1,3}$ peak appears sharper on the low energy side in the $^{55}$Fe$_2$O$_3$ than in the MnO spectrum than. A multiplet approach using atomic self-consistent field calculations can account for the Kβ$_{1,3}$ and Kβ'$^{i}$ features[12]. For a more detailed treatment the local symmetry (O$_h$ for both comounds) and orbital hybridization can be taken into account [13, 14]. It is found
FIGURE 4. Left: Term scheme for $K\beta$ emission after core hole creation via $K$ capture and photoionization, respectively. The change of effective potential upon photoionization leads to a populated shake configuration in the intermediate state. Assuming that the shake configuration does not relax into the lowest intermediate state we obtain two $K\beta$ emitting transitions. Right: Calculated $K\beta$ main lines after (a) $K$ capture and (b) photoionization. For the latter the two spectra as described in the term scheme on the left are shown and the spectra are scaled according to the calculated intensities of the $1s$ intermediate states.

that different hybridization and crystal field splittings between MnO and Fe$_2$O$_3$ cannot explain the differences observed in the experimental spectra [15].

Figure 4 shows how the dependence of the spectral features on the mode of $1s$ core hole creation can be incorporated into a theoretical model. The change of effective potential experienced by the valence electrons after $1s$ photoionization causes shake transitions and, in a simplified model, two populated intermediate states. Both states act as fluorescence initial states and give rise to $K\beta$ emission. We therefore assume that the shake configuration in the $1s$ intermediate state does not decay into the lowest $1s$ intermediate state before the decay of the $1s$ core hole, i.e. the fluorescence initial state is not fully relaxed. On the other hand, the effective potential experienced by the valence electrons hardly changes in K capture decay because a negative charge (the $1s$ electron) annihilates with a positive charge (a proton in the nucleus) [15].

The calculated spectra on the right side of Figure 4 are based on atomic multiplet calculations including crystal field splitting and ligand-to-metal charge transfer [13, 15]. Two $K\beta$ spectra are shown for MnO after photoionization. They are scaled according to the calculated population of the $1s$ excited states that are reached in a photoionization process. Only one calculated $K\beta$ spectrum is shown in the case of $K$ capture because only one $1s$ excited state is reached. Adding up the two photoionization $K\beta$ spectra
FIGURE 5. $K\beta$ satellite emission in Fe compounds after K capture (KC) and photoionization (PI). To facilitate direct comparison a common energy scale (relative fluorescence energy) is given in the bottom. The feature in Fe$_2$O$_3$ at about 0 eV relative fluorescence intensity is the $K\beta''$ or cross over peak. It was used to align the photoionization and K capture spectra relative to each other.

will yield a spectrum that is broadened on the low energy side of the $K\beta_{1,3}$ peak in comparison to the K capture spectrum and thus reproduce the experimental observation [15].

**K$\beta$ Satellite Lines**

Transitions from shells higher than 3$p$ ($K\beta$ satellite lines) can be interpreted using density functional theory [16, 17]. The $K\beta$ satellite lines in Fe$_2$O$_3$, Fe$_4$Fe(CN)$_6$ and Fe$_4$Fe(CN)$_6$ are shown in Figure 5. While the $K\beta$ main lines are dominated by splittings due to intraatomic Coulomb and spin-orbit interactions whose magnitudes depend on the nuclear charge, the $K\beta$ satellite lines are mainly shaped by ligand field effects and band formation. We therefore compared a Mn to an Fe-55 compound in the previous section and now compare identical Fe and Fe-55 compounds for the $K\beta$ satellites. The fluorescence energy scale had to be shifted for a comparison. We used the $K\beta''$ peak that has been assigned to a ligand 2$s$ to metal 1$s$ transition [18] to align the K capture and photoionization spectra.

All spectral features recorded after photoionization are broader than in the data taken after K capture. Furthermore, the intensity ratios of the two bands in the $K\beta_{2,5}$ structure of the cyanides change between the two modes of excitation. The broadening indicates that more final states are populated after photoionization as we already concluded for the $K\beta$ main lines.

On the right of Figure 5 we compare Fe$_2$O$_3$ valence band photoemission spectra (taken from reference [19]) to the $^{55}$Fe$_2$O$_3$ $K\beta$ satellite spectra. Fujimori *et al.* interpreted the valence band UPS spectra using a ligand-to-metal charge transfer model and
assigned the strong peak at low binding energies to a screened 3d$^5L$ configuration and an O 2p band and the weaker structure at about 14 eV binding energy to an unscreened 3d$^4$ configuration. The K$\beta$ satellite lines are dominated by transitions from orbitals with mainly O 2s and 2p character. These orbitals exhibit some p-character relative to the metal center and therefore serve for dipole allowed transitions to the metal 1s shell.

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