In situ site-selective transition metal K-edge XAS: a powerful probe of the transformation of mixed-valence compounds†

Amélie Bordage,*a Virgile Trannoy,a Olivier Proux,b,c Hugo Vitoux,d Robinson Moulina and Anne Bleuzena

We present herein the first in situ site-selective XAS experiment performed on a proof-of-principle transformation of a mixed-valence compound: the calcination of the $K_{0.1}Co_{2+}^{III}[Co_{II}^{III}(CN)_6]_{2.7} \cdot 20H_2O$ Prussian Blue analogue (containing $Co^{2+}$ and $Co^{3+}$ ions in two different $O_h$ sites) into $Co_{3}O_4$ (containing $Co^{2+}$ ions in a $T_d$ site and $Co^{3+}$ in an $O_h$ site). By recording the Co K-edge X-ray absorption spectra using a spectrometer aligned at the maximum of the $K_{\beta_{1,3}}$ emission line, the evolution of each species was singly monitored from 20 °C up to the oxide formation. The experimental spectrum of the $Co^{2+}(T_d)$ and $Co^{3+}(O_h)$ species in $Co_{3}O_4$ is reported for the first time. Our results demonstrate the possibilities offered by site-selective XAS for the investigation of chemical transformations and the study of materials under working conditions whenever the chemical element of interest is present in several states and/or sites.

Introduction

X-ray absorption spectroscopy (XAS) is a well-known powerful tool to characterize the electronic and crystallographic structure of an element in a compound. In the hard X-ray range, XAS is now routinely performed under extreme conditions (pressure, temperature, and irradiation); in situ XAS1–7 is also a mature technique widely used to study chemical processes. However, the classical detection modes (transmission, total fluorescence and total electron yield) average over all species of the absorbing atom in the sample, which hinders the investigation of compounds with the absorbing atom present under different oxidation/spin states and/or in different sites. Site-selective XAS has recently brought new opportunities since it can overcome this limitation8–10 by taking advantage of the high sensitivity of the $K_{\beta_{1,3}}$ emission line of one element to the spin and oxidation states as well as to covalency.11–16 The spectra are recorded using a high-resolution Rowland-circle spectrometer with analyzer crystals aligned at the maximum of the $K_{\beta_{1,3}}$ emission line of a given site. The analyzer selects this particular fluorescence line for each incident energy and reflects it on a detector. This technique has been successfully used for instance to discriminate the Co ions from the core and the shell of nanoparticles17,18 or to investigate in detail the active site in [FeFe] hydrogenase.19,20 Herein, we demonstrate for the first time that this demanding XAS technique can be used for in situ investigations. This opens new perspectives to unravel the mechanisms of chemical processes (redox processes, homogeneous or heterogeneous catalysis, ...) and to study materials under working conditions (batteries, catalysts, ..., which are everyday challenges for chemists.

As a proof-of-principle study, we chose to follow the calcination in air of the monometallic Prussian Blue analogue (PBA) of formula $K_{0.1}Co_{2+}^{III}[Co_{II}^{III}(CN)_6]_{2.7} \cdot 20H_2O$ (called Co-PBA) into $Co_{3}O_4$.21 PBAs are well-known for their face-centred cubic structure made of cyanide bridges linking two transition metals in the three directions of space.22,23 In Co-PBA, $Co^{2+}$ high-spin ions (Fig. 1, orange balls) and $Co^{3+}$ low-spin ions (Fig. 1, green balls) are both present in an octahedral ($O_h$) site but surrounded by different neighbours: $Co^{2+}$ is linked to an average of four cyanide bridges (at the N side) and two water molecules, while $Co^{3+}$ is linked to six cyanide bridges (at the C side). The $Co_{3}O_4$ oxide crystallizes in a direct spinel structure with the $Co^{3+}$ ions in the $O_h$ site and the $Co^{2+}$ ones in the tetrahedral ($T_d$) site; both Co ions are linked to $O^{2−}$ ions. In the case of monometallic PBAs, reference compounds are available for each single site. For instance, in the case of Co-PBA, the $Co^{2+}$ site is analogous to the Co site in $Co_{2+}^{III}[Fe_{II}^{III}(CN)_6]_{2.7} \cdot 20H_2O$ PBA, while the $Co^{3+}$ site resembles that in $Fe_{II}^{III}[Co_{II}^{III}(CN)_6]_{2.7} \cdot 20H_2O$ PBA. However, in the case of the $Co_{3}O_4$ spinel or the transient states, such reference compounds...
are unavailable. The calcination of Co-PBA into Co$_3$O$_4$ is therefore an ideal case to test the opportunities offered by in situ site-selective XAS to follow complex chemical processes since at least four different sites are involved in the transformation (Co$^{3+}$–CN (Oh), Co$^{2+}$–NC (Oh), Co$^{2+}$–O (Oh), Co$^{2+}$–O (Td)) and the calcination completely redistributes the Co species. In this contribution we concentrate on the XANES (X-ray Absorption Near-Edge Structure) part of the X-ray absorption spectrum.

**Experimental**

**Materials**

Co-PBA ($K_{0.1}Co^{II}_4$[Co$^{III}$(CN)$_6$]$_{2.7}$·20H$_2$O) and a reference ex situ sample (obtained by the calcination of Co-PBA in air during 2 h hours at 900 °C, and called es situ) were synthesized as described in ref. 21. The two Co$_4$[Fe(CN)$_6$]$_{2.7}$·20H$_2$O (called Co$^{2+}$–N) and Rb$_2$Co$_4$[Fe(CN)$_6$]$_{3.3}$·11H$_2$O (called Co$^{2+}$–N) PBAs used as references were prepared as described in ref. 24. The other two reference compounds, [Co(OH)$_2$]$_3$[NO$_3$]$_2$ (called Co$^{3+}$–O) and K$_3$[Co(CN)$_6$] (called Co$^{3+}$–C), were supplied by Sigma-Aldrich.

**X-ray spectroscopy**

Co K-edge XANES spectra were recorded on the CRG-FAME (BM30B) beamline of the ESRF (Grenoble, France) using their five-crystal spectrometer. The energy of the incident radiation was selected using a pair of Si(220) crystals (flux of ca. 5.10$^{11}$ photons s$^{-1}$). Co K$\beta$ site-selectivity was performed by recording the XANES spectra with spherically bent Ge(444) crystals in a Rowland geometry. The detector was a Silicon Drift Detector (energy resolution: 250 eV) in order to discriminate the signal diffracted by the crystals from the scattered one and so to improve the signal-to-noise ratio. Self-absorption effects were ruled out by diluting Co-PBA in BN and we checked that no radiation damage occurred by recording fast spectra as a function of time. The combined resolution of the incident beam and the spectrometer was $\Delta E$ = 0.6 eV, measured from the pseudo-elastic peak full-width at half-maximum. Spectra were simultaneously recorded in the transmission mode. The in situ calcination was achieved using a Cyberstar gas-blower furnace. Calibration of the furnace was performed prior to measurements and the temperature stability was checked during measurements. This setup is illustrated in Fig. 2.

The X-ray emission spectrum of Co$^{2+}$–O (resp. Co$^{3+}$–C) was recorded at room temperature and was used to align the spectrometer for the +II (resp. +III) oxidation state. Data were acquired at seven temperatures from room temperature up to 400 °C. In the following text, only the spectra corresponding to the most relevant temperatures for the discussion (20 °C, 145 °C and 400 °C) are presented; for better clarity, the temperature is now specified next to the Co-PBA name (Co-PBA@temperature). For each temperature at which Co-PBA was investigated, a Co K$\beta$, X-ray emission spectrum and Co K-edge X-ray absorption spectra for the two positions of the spectrometer were recorded. The X-ray absorption spectra of Co-PBA and of the references were also recorded in transmission mode at each investigated temperature.

**Results and discussion**

**Formation of the targeted phase**

In order to attest the formation of the targeted Co$_3$O$_4$ phase during the in situ calcination process, the Co K-edge spectrum of Co-PBA was recorded. It is compared to the transmission spectrum of Co-PBA at 400 °C in Fig. 3. A very good agreement is observed both between them and the Co$_3$O$_4$ spectra reported in the literature and in particular the maximum of the white line at 7729.4 eV. This agreement attests that the targeted Co$_3$O$_4$ phase was produced during the in situ chemical transformation of Co-PBA; this is also confirmed by X-ray diffraction (see ESI†). Therefore, the information obtained from the spectra recorded at the different temperatures actually reflects the processes involved in the calcination described in ref. 21.
transition metal ions\textsuperscript{11–16} and can be performed with a high-sensitivity to the spin state of the Co K-edge \textsuperscript{23–30} 

The slight differences in the relative intensities can be explained by a lower crystallinity for the \textit{in situ} sample due to the lower temperature of calcination.

\section*{In situ X-ray emission spectroscopy}

A high-resolution spectrometer such as the one on FAME\textsuperscript{25} also enables us to perform X-ray emission spectroscopy (XES). This technique is well-known for its high-sensitivity to the spin state of transition metal ions\textsuperscript{23–30} and can be performed \textit{in situ}.\textsuperscript{1,2,28–30} The Co K\textsubscript{\beta,1,3} X-ray emission spectra of Co-PBA as a function of temperature are displayed in Fig. 4, as well as the spectrum of \textit{ex situ}. A shift of the maximum towards higher energy is observed upon heating and reveals a change in the Co species during the calcination process. However, it becomes clear that XES reaches a limit for the study of transformations in multisite compounds. The changes observed with the temperature are indeed (i) too small to enable a precise determination of the changes in the Co environment and (ii) averaged over the different species.

\section*{Extraction of the Co\textsuperscript{2+} and Co\textsuperscript{3+} pure contributions}

The Co K\textsubscript{\beta,1,3} X-ray emission spectrum of Co-PBA\textsubscript{20 °C} is shown in Fig. 5a, along with those of the Co\textsuperscript{2+}–O and Co\textsuperscript{3+}–C references for the two sites of Co (called simply as Co\textsuperscript{2+} and Co\textsuperscript{3+} sites in the following text) in this compound. The important feature here is the shift in energy of the K\textsubscript{\beta,1,3} line maximum from 7650.6 eV for Co\textsuperscript{2+}–O down to 7649 eV for Co\textsuperscript{3+}–C. XANES spectra were therefore successively recorded using a spectrometer aligned at these two energies. However, it is clear from Fig. 5a that experimentally, one cannot record a pure site-selective X-ray absorption spectrum: whatever the energy chosen for the spectrometer, the spectrum includes a contribution from both sites. The pure Co\textsuperscript{2+} and Co\textsuperscript{3+} contributions for each temperature must consequently be extracted from the measured spectra at the two energies of the spectrometer. We describe now this procedure, which is illustrated in Fig. 5 for the 20 °C measurement. In the following text, the term ‘K\textsubscript{\beta}-HERFD X-ray absorption spectrum’ refers to the spectrum measured using the spectrometer, while the term ‘site-selective spectrum’ refers to the ‘pure’ Co\textsuperscript{2+} and Co\textsuperscript{3+} extracted contribution. The reader should keep in mind that the pure term is used here with the understanding that the spectra it refers to are obtained after deconvolution of two experimental spectra and were not directly measured as real pure spectra.

First, from a linear combination of the normalized X-ray emission spectra of Co\textsuperscript{2+}–O and Co\textsuperscript{3+}–C wherein the Co\textsuperscript{2+} : Co\textsuperscript{3+} ratio of the integrated intensity is 4 : 2.7, we obtain the expected K\textsubscript{\beta,1,3} spectrum for a Co\textsuperscript{2+}Co\textsuperscript{3+} PBA; an excellent agreement is observed with the experimental spectrum of Co-PBA\textsubscript{20 °C} (Fig. 5a). The Co\textsuperscript{2+} fraction is then determined as the ratio of the normalized spectrum of Co\textsuperscript{2+}–O to the expected K\textsubscript{\beta,1,3} X-ray emission spectrum of Co\textsuperscript{2+}Co\textsuperscript{3+} PBA or of Co-PBA\textsubscript{20 °C} (Fig. 5b). For the two positions of the spectrometer chosen to record the K\textsubscript{\beta}-HERFD X-ray emission spectra (at 7649 eV and 7650.6 eV), the value of the Co\textsuperscript{2+} fraction ($\gamma_{7649}$ and $\gamma_{7650,6}$; Fig. 5b) is determined. They are then used to extract the pure Co\textsuperscript{2+} and Co\textsuperscript{3+} site-selective spectra (S(Co\textsuperscript{2+}) and S(Co\textsuperscript{3+})) by a linear combination of the

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig3.png}
\caption{Normalized Co K-edge XANES spectra in transmission mode of Co-PBA\textsubscript{400 °C} and \textit{ex situ}.
\label{fig3}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig4.png}
\caption{\textit{In situ} normalized Co K\textsubscript{\beta,1,3} X-ray emission spectra of Co-PBA\textsubscript{20 °C}, Co-PBA\textsubscript{145 °C} and Co-PBA\textsubscript{400 °C}, presented with the spectrum of \textit{ex situ}.
\label{fig4}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig5.png}
\caption{Extraction of the Co\textsuperscript{2+} and Co\textsuperscript{3+} site-selective XANES spectra. (a) Normalized Co K\textsubscript{\beta,1,3} emission line for the Co\textsuperscript{2+}–O and Co\textsuperscript{3+}–C references, and Co-PBA\textsubscript{20 °C}. The dotted line on the upper curve represents the linear combination of the normalized X-ray emission spectra of Co\textsuperscript{2+}–O and Co\textsuperscript{3+}–C so that the Co\textsuperscript{2+} : Co\textsuperscript{3+} ratio of the integrated intensity is 4 : 2.7 (Co\textsuperscript{2+}Co\textsuperscript{3+}). (b) Co\textsuperscript{2+} fraction at room temperature, with the values for the two positions of the spectrometer ($\gamma_{7649}$ and $\gamma_{7650,6}$). (c) Normalized Co K-edge XANES spectra of Co-PBA\textsubscript{20 °C} recorded in transmission mode (upper curve) and for the two positions of the spectrometer (s@7649, orange middle curve and s@7650,6 orange middle curve), with the extracted pure Co\textsuperscript{2+} (S(Co\textsuperscript{2+})), orange lower curve) and Co\textsuperscript{3+} (S(Co\textsuperscript{3+}); green lower curve) contributions.
\label{fig5}
\end{figure}
measured Kβ-HERFD X-ray emission spectra (s@7649 and s@7650.6) (Fig. 5c):

\[
S(Co^{2+}) = \frac{(1 - \gamma_{7650.6}) s@7649 - (1 - \gamma_{7649}) s@7650.6}{\gamma_{7649} - \gamma_{7650.6}}
\]

This equation corresponds to a standard deconvolution of a multi-component system and was already used in previous site-selective XAS investigations.8,11,17

In order to assess the reliability of this extraction procedure, additional XANES spectra in the transmission mode were recorded for two references: Co\(^{3+}\)–C and Co\(^{2+}\)–N, where the Co ions are present in the same O\(_6\) Co\(^{3+}\) and Co\(^{2+}\) sites as those in Co-PBA@20 °C. An excellent agreement is observed between the site-selective spectra of Co-PBA@20 °C and the related references in transmission mode (Fig. 6). This indicates that the extraction procedure is reliable and it was therefore applied to all measured temperatures.

**Evolution of the Co\(^{2+}\) and Co\(^{3+}\) contributions with temperature**

In the following text, we do not discuss the pre-edge features of the site-selective spectra since the selection of the Co K\_\(_{\beta 1,3}\) emission line to record the XANES spectrum may result in the modification of some spectral features in the pre-edge.18,31,32

The pre-edge measured using HERFD indeed corresponds to a constant emission energy cut in the resonant XES (RXES) plane11,33,34 and it was shown that such a scan may show features that are not actual absorption features.35 The reliable interpretation of the HERFD pre-edge thus requires the measurement and careful analysis of the full RXES plane,11,35 which was beyond the scope of this study. However, it is to be noted that the main edge region of the spectrum (above the pre-edge features) should mainly be affected by sharpening as compared to a conventional XANES spectrum. This is due to the resolution that has the order of magnitude of the core hole lifetime. Detailed discussion about this point can be found in ref. 11 and 33–35.

The Co\(^{2+}\) and Co\(^{3+}\) site-selective XANES spectra of Co-PBA as a function of temperature and of exsitu are displayed in Fig. 7. At 20 °C, the shape of the spectra is the signature of the Co\(^{3+}\) (Fig. 6 and 7a) or Co\(^{2+}\) (Fig. 6 and 7b) ions in the two O\(_6\) sites of the PBA structure. Both Co-PBA@400 °C and exsitu consists of Co\(_{3}O_{4}\) their site-selective Co\(^{3+}\) (Fig. 7a) and Co\(^{2+}\) (Fig. 7b) spectra are the first experimental signature of the single Co\(^{3+}\)(T\(_{d}\)) and Co\(^{3+}\)(O\(_{h}\)) sites in the spinel structure. These experimental spectra of Co-PBA@400 °C and exsitu are also in quite good agreement with multiple-scattering calculations performed on Co\(_{3}O_{4}\) by Jiang and Ellis.36

The pure Co\(^{2+}\) and Co\(^{3+}\) site-selective XANES spectra display significant changes with temperature (Fig. 7). In the case of Co\(^{3+}\) (Fig. 7a), the spectra at 20 °C and 145 °C are very close, indicating that this site of the PBA is not modified and that the Co\(^{3+}\) ion remains sixfold-coordinated to CN\(^{–}\) ligands. At 400 °C the spectrum displays some modification, in particular for the B\(_2\) and C\(_2\) features. On the contrary the energy of the white line (7728.2 eV; peak A\(_3\)) remains nearly constant. In Co\(_{3}O_{4}\), the Co\(^{3+}\) cation is known to sit in the octahedral site, which is consistent with these spectral features observed at 400 °C and for exsitu. The high intensity of the B\(_2\) feature at 20 °C results from the significant multiple scattering (MS) in the Co\(^{3+}\)–CN linkages.37–40 Its strong and abrupt intensity-decrease above
OH bonds. In the case of Co2+ (Fig. 7a, upper curves) for an average of four Co2+(HS)–NC and two Co2+(HS)–OH2 bonds, two more references are displayed: Co2+–N for a Co2+–NC bond, Co2+–C for a Co2+–CN bond and Co2+–N for a Co2+–NC bond. (b) Comparison of the Co2+ and Co3+ site-selective XANES spectra of Co-PBA@400°C extracted using Co2+–C (upper curves) or Co3+–N (lower curves) as a reference for the +III oxidation state. The dotted (resp. dashed) lines are guides to compare the energy of the Co2+ (resp. Co3+) spectral features.

200 °C reveals the decomposition of the cyanide bridges already observed by previous thermodifferential and thermogravimetric analyses21 and is consistent with the lower MS in CoO4. The shift towards the lower energy of the B3 and C1 features at 400 °C also reflects the lengthening of the Co-to-ligand bond expected for the Co3+–CN → Co3+–O transformation,43,44 according to Natoli’s rule.43 These observations indicate that the Co3+ site mainly undergoes a change in the chemical nature of the neighbours, i.e. the abrupt replacement of the six C neighbours by the six O neighbours above 200 °C following the cyanide bridge decomposition.21 In contrast, in the case of Co2+ (Fig. 7b), a continuous evolution of the spectra is observed upon heating: at 145 °C, the relative intensity of the A2 and B3 features is almost 1 : 1, and at 400 °C it is inverted with respect to the spectrum at 20 °C. These spectral changes upon heating are consistent with the progressive transformation from Co(NC)6(OH2)2 (20 °C) to CoO4 (400 °C), probably via a Co(NC)4 transient state (145 °C). This transient state is supported by our previous thermogravimetric analyses, which showed the loss of water molecules above 90 °C.21 In addition to the determination of the steps of the transformation, this clearly different behavior of the two sites also shows that it is possible with site-selective XAS (i) to discriminate between active and inert species and (ii) to monitor the active species during the transformation.

Effect of the ligands

In this proof-of-principle in situ study, the successful extraction of the pure Co2+ and Co3+ site-selective spectra justifies a posteriori the use of only two references and consequently the recording of the spectra for only two energies (positions) of the spectrometer. It is known that for a given oxidation and spin state, the Kb1,3 lines are also sensitive to the nature of the ligand,11,12,19 as illustrated in Fig. 8, where in addition to Co2+–O (Co3+–O–OH2 bond) and Co3+–C (Co3+(LS)–CN bond), two more references are displayed: Co2+–N for an average of four Co2+(HS)–NC and two Co3+(HS)–OH2 bonds, and Co2+–N4 for an average of five Co3+(LS)–NC and one Co3+(LS)–OH2 bonds. In the case of Co2+ (Fig. 7a, upper curves), the two Kβ1,3 X-ray emission spectra are almost superimposed, as expected by the close covalency of the Co2+–O and Co3+–N bonds. On the contrary, the Kβ1,3 X-ray emission spectra of Co2+–N and Co3+–C (Fig. 8a, lower curves) display significant differences, both in the spectral shape and in the energy of the maximum. This ligand-dependence of the Kβ1,3 X-ray emission spectrum makes preferable the choice of references with a covalency of the Co-to-ligand bond as close as possible as in the chemical species under study. Nevertheless, we observed here that whatever the choice of the reference compound, the position of the features of the site-selective spectra is not modified and that only their relative intensity varies (Fig. 8b). Measurements for as many spectrometer positions as required by the different oxidation state – spin state – covalency sites possibly found in the compound under investigation should strengthen the results. However, even if references are chosen only from the known states of the material under investigation, reliable results are still obtained from the site-selective XANES spectra. This point has to be noted, since the transient (and possibly the final) species are usually unknown.

Conclusions

We present herein the first in situ site-selective XAS study, with an emphasis on the XANES part of the spectrum. We successfully managed to singly resolve the evolution of the Co2+ and Co3+ sites during the calcination of Co-PBA into CoO4. The spectra of Co3+(T2) and Co3+(O2) ions in Co3O4 were also singly recorded for the first time. Obtaining pure spectra for all the sites demonstrates again the power of site-selective XANES to investigate very complex samples such as the spinel family, which presents direct and inverse spinels as well as solid solutions. In addition, herein we showed that site-selective XANES experiments can innovatively be trustfully performed in situ, which opens new perspectives for the study of chemical transformation or materials under working conditions (catalytic systems, biological systems, materials used for data storage, battery, renewable energy, etc.).

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Notes and references
