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Total electron yield (TEY) detection mode Cr K-edge XANES spectroscopy as a direct method to probe the composition of the surface of darkened chrome yellow ($\text{PbCr}_{1-x}\text{S}_x\text{O}_4$) and potassium chromate paints

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Abstract

The darkening of chromate-based yellow pigments, including chrome yellows ($\text{PbCr}_{1-x}\text{S}_x\text{O}_4$, with $0 \leq x \leq 0.8$), is a surface phenomenon leading to the formation of a thin alteration layer (thickness $< 10 \mu\text{m}$) that affects a series of iconic paintings of the late 19th-early 20th century, such are those by Vincent van Gogh.

In this paper, we systematically examined the capabilities of Cr K-edge X-ray absorption near edge structure (XANES) spectroscopy collected at the same time in X-ray fluorescence yield (XFY) and total electron yield (TEY) detection modes to selectively study the composition of the surface of darkened chromate-based yellow oil paints made up of either sulfate-rich chrome yellow type ($\text{PbCr}_{0.2}\text{S}_{0.8}\text{O}_4$) or potassium chromate.

By describing and discussing advantages and drawbacks in using either XFY or TEY detection modes in relation to XFY mode μ -XANES analysis with nano-probes from sectioned samples, we aim at assessing if: (i) TEY-XANES spectroscopy may be employed as a direct selective surface method to determine the relative abundance of different Cr-species from paint fragments; (ii) it can contribute to optimize the analytical strategy by minimizing time-consuming sample preparation; (iii) it can decrease the probability of X-ray-induced radiation damage observed when photons are concentrated in sub-micrometre probes.

1. Introduction

Darkening or fading of paint, often accompanied by loss of paint stability due to chemical changes of the pigments and/or the binding medium, affect a large number of painted (cultural heritage) objects, with serious risks for their preservation and management.

For a number of inorganic pigments, including Prussian blue ($MFe^{III}[Fe^{II}(CN)_6] \cdot xH_2O$, with $M=K^+, NH_4^+$ or Na^+), [1,2] vermilion red ($\alpha-HgS$), [3,4] orpiment ($\alpha-As_2S_3$), [5,6,7,8] cadmium yellows ($Cd_{1-x}Zn_xS$) [9,10,11,12,13,14] and reds ($CdS_{1-x}Se_x$), [15] the color changes that are observed in paintings and other artefacts are the result of redox reactions. The process is a surface phenomenon, leading to the formation of amorphous and/or crystalline secondary products that usually produce layers of limited size within the paint stratigraphy; in most cases, the altered layer at the surface has a thickness below ca. 10 μm .

Last decades have experienced a growing interest in the use of synchrotron radiation (SR)-based X-ray methods with micro-/nano-beams, in particular micro-X-ray near edge structure spectroscopy (μ -XANES) in X-ray fluorescence yield detection mode (XFY), for their capabilities to provide non-destructive spatially resolved elemental speciation information at the (sub)micrometre scale length of both amorphous and crystalline compounds across the stratigraphy of non-transparent samples. [16, 17]

Nevertheless, sample preparation in the form of cross section preparation, of which the characteristics change depending on the nature of the analyzed material and technical features of the employed method (e.g., set-up geometry, energy of the incident beam...), is often demanding and time consuming. [18] In addition, sampling usually gives access to submicrometric areas of the artwork surface, with the consequence that, in some cases, the composition of the analyzed micro-samples might be not fully representative of the studied phenomena.

Thus, it becomes relevant to explore in-depth how SR-based X-ray methods may allow to obtain direct surface elemental speciation information from larger, non-transparent and

layered fragments, while minimizing consuming and lengthy sample preparation procedures.

In this context, total electron yield (TEY) detection mode has been proved to be a suitable alternative to the indirect XAS fluorescent mode measurement because of the shallowness of the electron escape depth (of the order of tens of nm) compared with fluorescence X-rays. [19,20 21] Its use, despite limited to few case studies in the field of heritage science, has been successfully exploited for performing elemental speciation investigations in different kind of objects, including lustre decoration of Italian Renaissance pottery, [22] historical wooden buildings, [23] degraded zinc yellow and copper resinate paints. [24,25]

In the present study, we systematically explore the capabilities of Cr K-edge XANES spectroscopy collected at the same time in X-ray fluorescence yield (XFY) and total electron yield (TEY) detection modes to perform Cr speciation investigations of the surface of a set of darkened chromate-based yellow oil paints made up of either the light-sensitive sulfate-rich chrome yellow type ($\text{PbCr}_{0.2}\text{S}_{0.8}\text{O}_4$, hereafter denoted as $\text{Cr}_{0.2}\text{Y}$) or potassium chromate. As the above-mentioned pigments, both class of compounds undergo darkening as a result of a reduction process of the original Cr^{VI} to Cr^{III} -compounds, with Cr^{V} -species arising from the interaction between the pigment and the oily medium. [26,27,28]

The technical advantages and drawbacks in using either XFY or TEY detection modes are presented and discussed in relation to XFY mode μ -XANES analysis from sectioned samples. This is done in order to assess if: (i) TEY-XANES spectroscopy using an unfocused beam may be employed as a selective method to determine the relative abundance of different Cr species directly at the degraded surface of paint fragments; (ii) it can contribute to optimize analytical strategy by limiting time-consuming sample preparation procedures; (iii) it may help to reduce the probability of radiation damage, considering the tendency of chromate-based compounds towards photo-reduction under the exposure to X-ray beams.[29, 30]

2. Materials and methods

2.1. Preparation of paint mock-ups and photochemical aging protocol

Mock-up paints were prepared on polycarbonate slices by employing either mainly orthorhombic $\text{PbCr}_{0.2}\text{S}_{0.8}\text{O}_4$ (synthesized in our laboratory) or commercial orthorhombic K_2CrO_4 (Sigma-Aldrich) in mixture with cold-pressed linseed oil (Zecchi) in a 4:1 weight ratio.

Further information on the synthesis, crystalline structure, molecular properties and chemical reactivity of both compounds were described earlier. [26-28,31,32]

All paints were left to dry in the dark at temperatures of 25-35 °C and 35-45% relative humidity (RH) (i.e., measured indoor temperature and humidity level) and analyzed either ca. 1 month (for $\text{Cr}_{0.2}\text{Y}$) or about 10 hours (for K_2CrO_4) after their preparation (samples hereinafter called “unaged” and “0 days”, respectively).

As described earlier, [26] K_2CrO_4 oil paint films naturally undergo darkening within a few days; thus, the artificial photochemical aging was performed only on the 1-month prepared $\text{Cr}_{0.2}\text{Y}$ paint. The treatment was carried out at ~20-35% RH (measured indoor humidity level) by means of a UVA-visible light ($\lambda > 300$ nm) emitted by a UV-filtered 300 W Cermax xenon lamp (see ref. [28] for the corresponding emission spectrum). The measured irradiance and temperature at the sample position were $\sim 5 \times 10^5 \mu\text{W}/\text{cm}^2$ and 25-30 °C, respectively. The paint was irradiated for 960 h to achieve a total radiant exposure value of $\sim 5 \times 10^7 \mu\text{W}/\text{cm}^2 \cdot \text{h}$.

2.2. XFY/TEY mode XANES spectroscopy at Cr K-edge

XANES investigations at Cr K-edge of paint mock-ups were performed at the beamline BM08-LISA [33] and at the scanning X-ray microscope end station of beamline ID21 of the European Synchrotron Radiation Facility (ESRF, Grenoble, France). [34,35]

Measurements were carried out under 0.5 atm He atmosphere with a Si(311) crystal at BM08-LISA and under vacuum using a Si(220) fixed exit double-crystal monochromator at ID21.

At BM08-LISA, data were collected using an average beam size of $\sim 70 \times 100 \text{ } \mu\text{m}^2$ (h×v) and a fluence rate of $\sim 10^4 \text{ ph}/(\text{s} \cdot \mu\text{m}^2)$. An appropriate sample holder, [36] equipped with a collecting anode polarized at $\sim 20 \text{ V}$ and placed at about 10 mm from the sample, allowed performing simultaneously XANES measurements in two different detection modes: (i) TEY and (ii) XFY. To guarantee an adequate conductivity of the sample during TEY analysis, a small fragment of each paint ($\sim 20\text{-}30 \text{ mm}^2$ size) was fixed with carbon adhesive tape on the sample holder. The He atmosphere was used as an electron amplifier as described in previous literature. [37] Spectra were recorded by scanning the primary energy of the incident beam across the Cr K-edge with the following steps (345 points; 3 s/pt): 5.8892-5.9692 keV, step: 10 eV; (ii) 5.9742-6.0496 keV, step: 0.25 eV; (iii) 6.0499- 6.5375 keV, step: from 3 to 9 eV. A Cr metallic foil was used as a calibrant and a spectrum was collected at the same time as the sample to ensure the energy scale stability, which resulted to be better than 0.01 eV. For each sample, a total of 2-4 spectra were acquired.

At ID21, the incident beam was focused with a Kirkpatrick-Baez mirror system down to a spot size of $\sim 0.6 \times 0.3 \text{ } \mu\text{m}^2$ (h×v) and, without beam attenuation, with a fluence rate of $\sim 10^{10} \text{ ph}/(\text{s} \cdot \mu\text{m}^2)$. μ -XANES spectra in XFY mode were recorded from thin sections of the aged mock-ups (thickness of 10-15 μm) by scanning the primary energy of the incident beam across the Cr K-edge (5.95–6.087 keV) with increments of 0.2 eV (640 points; 0.1 s/pt). For the calibration, the first inflection point of the first-order derivative Cr K-edge XANES spectrum of a Cr metallic foil was set at 5.9892 keV. For normalization purposes, a photodiode upstream the sample was constantly monitoring the beam intensity. μ -XRF mapping was performed *via* raster scanning of the samples with 40 ms/pixel dwell time and using a 80 mm² collimated active area silicon drift diode detector (Xflash 5100, Bruker). Cr oxidation state maps acquired from K₂CrO₄ mock-up were extracted by means of the PyMca software, [38] after setting the energy of the incident X-ray beam at two energies: (i) at 5.993 keV, for promoting the excitation of Cr^{VI}-species, and (ii) at 6.090 keV for collecting XRF

signals of all Cr species. Details about the procedure employed to extract the Cr oxidation state maps are described elsewhere. [39]

The ATHENA software package [40] was employed for the normalization and for the linear combination fitting (LCF) of the BM08-LISA and ID21 XANES spectra against a library of spectral profiles of Cr-reference powders. This procedure allowed the relative abundance of Cr^{VI}-species (expressed as %[Cr^{VI}]/[Cr_{total}]) and of reduced Cr-compounds(i.e., Cr^V and Cr^{III}) to be determined quantitatively.

The X-ray penetration depth as a function of the angle of the incident beam at the Cr K-edge energy (*cf.* Fig. 5) was obtained using the software package JGIXA. [41]. The sampling depth of the photoelectrons (predominantly originated by cascades of the Cr KLL Auger electrons at 4.8 keV) was estimated by following the procedure earlier reported in the literature. [42]

3. Results

To explore the capabilities of Cr-K edge TEY-XANES spectroscopy in performing selective surface Cr-speciation investigations, we started with the study of fragments of $\text{Cr}_{0.2}\text{Y}$ and K_2CrO_4 paint mock-ups at beamline BM08-LISA.

In line with previous studies, [26-28, 39] the photoaging treatment of $\text{Cr}_{0.2}\text{Y}$ and the natural aging of K_2CrO_4 promote a significant darkening of the paint surface (Figs. 1A-2A).

Cr-K edge XANES spectroscopy data collected at the same time in XFY and TEY detection modes from unaged- $\text{Cr}_{0.2}\text{Y}$ and 0 days- K_2CrO_4 paint mock-ups reveal that only Cr^{VI} compounds are present (Figs. 1B-2B, circle black lines; only the XFY spectra are reported as example). This is highlighted by the peculiar intense pre-edge peak at 5.993 keV, attributed to a dipole-allowed transition from the Cr 1s orbital into orbitals with Cr 3d character (t_2 symmetry). Changes in the features of the post-edge absorption region reflect the different Cr local environments in the crystal structures of $\text{Cr}_{0.2}\text{Y}$ and K_2CrO_4 . [39,43,44,45,46]

In both aged paints (Figs. 1B-2B, black and grey lines), the partial conversion of Cr^{VI} to Cr^{III} -species is pointed out by the decreasing of the pre-edge peak intensity (quantitatively proportional to the Cr^{VI} to total Cr content ratio) and by a shift of the absorption edge energy. [39,43,45,46] Nevertheless, the contribution of reduced Cr-species is higher in the spectrum recorded in TEY mode than XFY mode, as shown by a more significant decrease of the pre-edge peak intensity (see inset) and shift of the absorption edge towards lower energy along with a loss of structure of the post-absorption features. This result can be explained considering the lower sampling depth of TEY mode measurements with respect to XFY mode analysis, that, under the employed experimental conditions, allows the composition of the superficial brownish layer to be probed selectively.

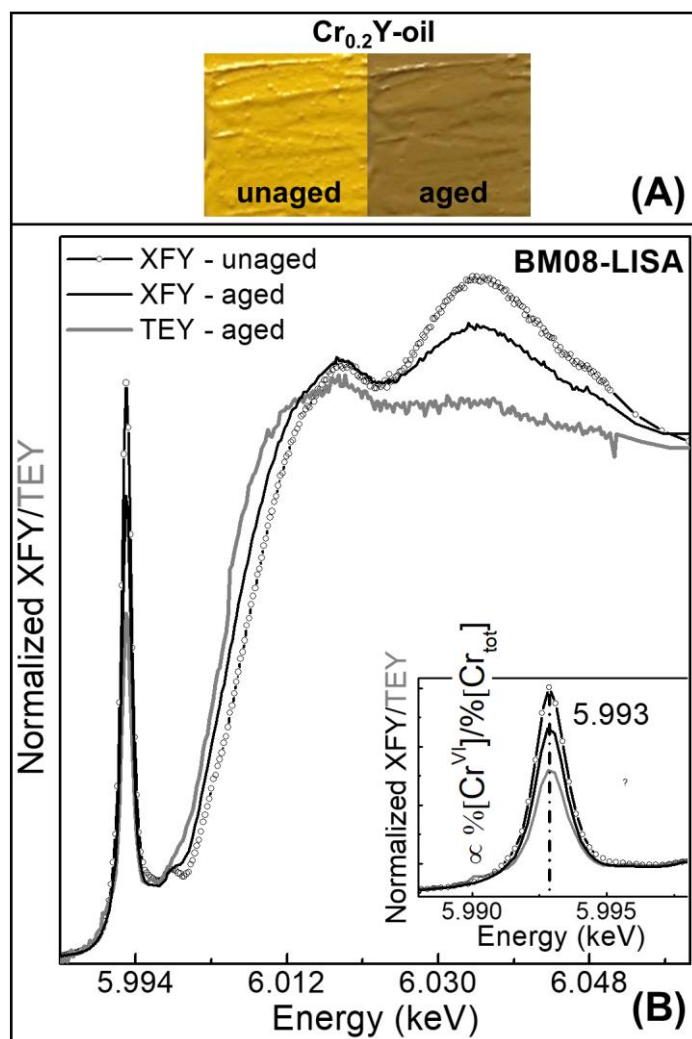


Figure 1. Photographs of sulfate rich-chrome yellow type ($\text{PbCr}_{0.2}\text{S}_{0.8}\text{O}_4$) paint mock-ups before (left) and after artificial aging with UVA-visible light. (B) Cr K-edge XANES spectra collected from the surface of unaged/aged $\text{Cr}_{0.2}\text{Y}$ paints in XFY mode (solid and circle black lines) and TEY mode (grey) at BM08-LISA.

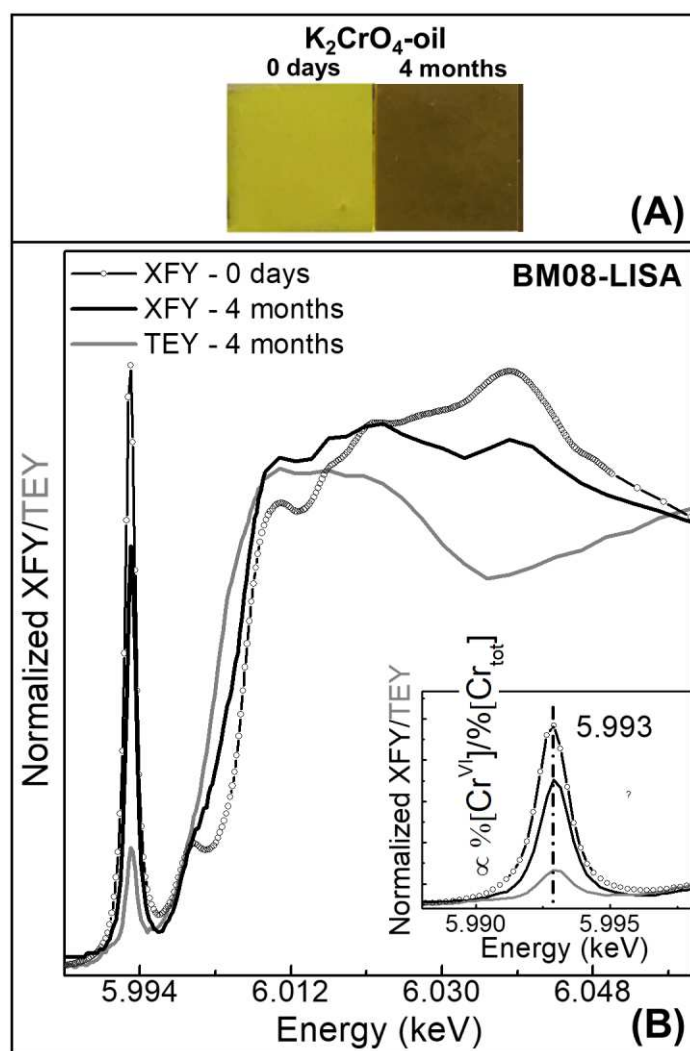


Figure 2. (A) Photographs of paint mock-ups obtained from K₂CrO₄ paint mock-ups and collected immediately (0 days) and 4 months after preparation of the sample. (B) Cr K-edge XANES spectra recorded from the surface of 4-month naturally aged K₂CrO₄ paint in XFY mode (solid and circle black lines) and TEY mode (grey) at BM08-LISA.

To better correlate the TEY-XANES results obtained from the direct analysis of the aged paint surface with the stratigraphic distribution at the submicrometric scale length of different Cr-species, we carried out Cr K-edge μ -XANES spectroscopy investigations at beamline ID21 on thin sections obtained from the photoaged $\text{Cr}_{0.2}\text{Y}$ and a 4-months naturally aged K_2CrO_4 paint.

As illustrated in the microphotographs of Figs 3A-4A, a brownish layer of about 5 μm and 20 μm thickness is clearly visible at the surface of the $\text{Cr}_{0.2}\text{Y}$ and K_2CrO_4 paints, respectively.

The XFY mode μ -XANES spectra recorded from the uppermost darkened layer of both the aged $\text{Cr}_{0.2}\text{Y}$ and K_2CrO_4 thin sections (Figs. 3B-4B; black lines “top”) show features very similar to the ones recorded in TEY modes directly at the altered surface of both paints (grey lines). In line with our previous studies, [26] the results obtained by LCF of the XFY and TEY mode (μ -)XANES spectra highlight that the abundance of reduced Cr [likely a mixture of Cr^{III} -oxides, organo- Cr^{III} compounds and/or oxochromate(V)-complexes] is ~50% and ~75-80% within the uppermost darkened layer of $\text{Cr}_{0.2}\text{Y}$ and K_2CrO_4 paints, respectively. XFY mode μ -XANES profiles reveal that Cr^{VI} -species are instead dominant in the bulk yellow paint of both samples (black lines “bulk”). In addition, Cr oxidation state maps recorded from a region of interest of the aged K_2CrO_4 paint contribute to visualize more clearly the stratigraphic distribution of different Cr-species, showing that the brownish alteration layer is mainly composed of reduced Cr-compounds, while Cr^{VI} -species are homogeneously present as the main component of the yellow paint (Fig. 4C). Overall, the stratigraphic outcomes further support the hypothesis that the discrepancies observed between the XFY- and TEY-XANES spectra recorded directly at the surface of the aged paints (Figs 1B-2B) are related to the different sampling depth of the two techniques.

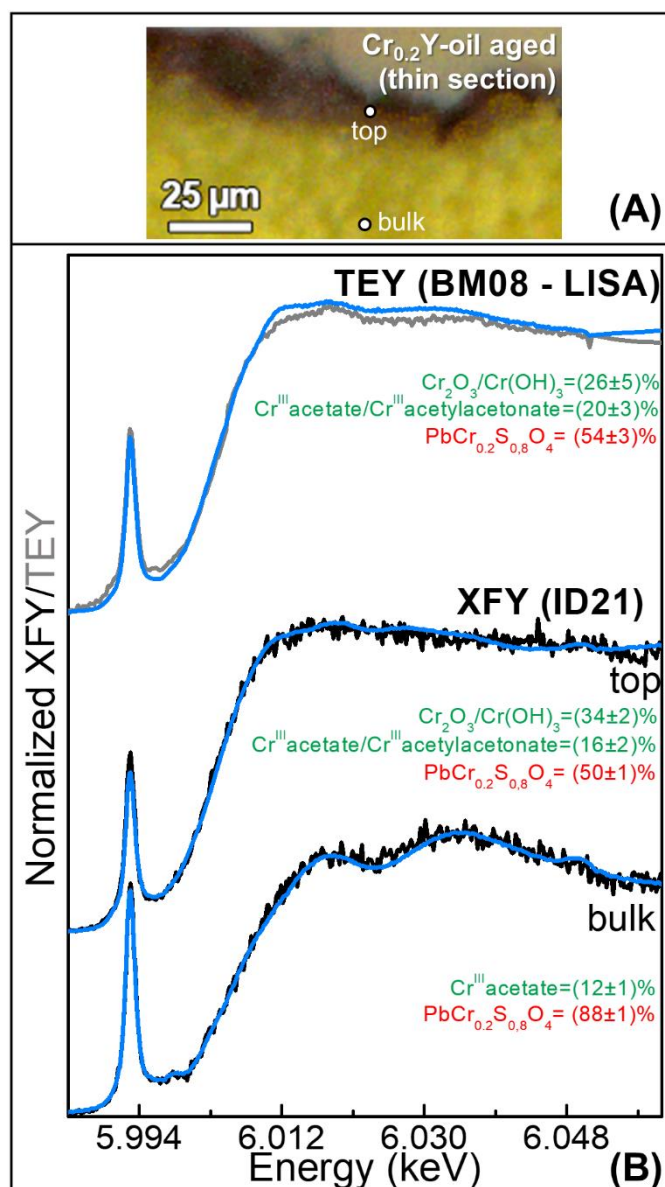


Figure 3. (A) Microphotograph of a thin section obtained from the aged $\text{Cr}_{0.2}\text{Y}$ paint. (B) Result of the linear combination fit (cyan) of the XANES spectra of $\text{PbCr}_{0.2}\text{S}_{0.8}\text{O}_4$ and different Cr^{III} -references to the ones recorded from the surface of the aged $\text{Cr}_{0.2}\text{Y}$ paint in TEY mode at BM08-LISA (grey) and from the uppermost alteration layer and the bulk paint of the thin section shown in (A) in XFY mode at ID21 (black). Spectra of (B) were recorded from the regions illustrated in (A).

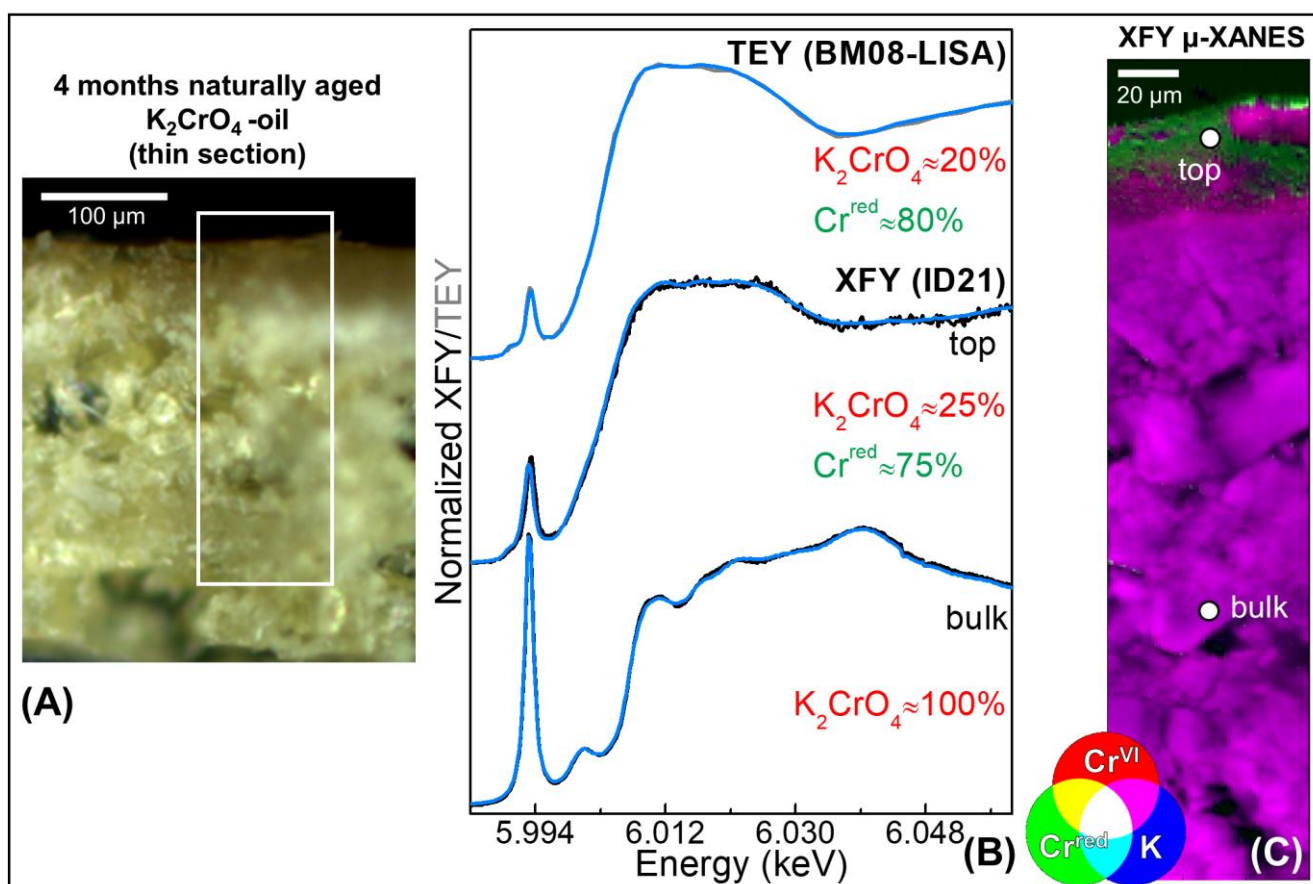


Figure 4. (A) Microphotograph of a thin section acquired from the 4-month-old K_2CrO_4 paint. (B) Result of the linear combination fit (cyan) of the Cr K-edge XANES spectra of K_2CrO_4 , $NaCr^{VO}_5(C_5H_8O)_2$ and two Cr^{III} -references [namely: $Cr(OH)_3/Cr_2O_3$ and Cr^{III} -acetate/ Cr^{III} -acetylacetonate] to the ones collected from the surface of the 4-months naturally aged K_2CrO_4 paint in TEY mode at BM08-LISA (grey line) and from the uppermost alteration layer and bulk yellow paint of the thin section in XFY mode at ID21 (black lines). (C) RGB SR μ -XRF images of $Cr^{VI}/Cr^{red}/K$ acquired from the area shown in (A). Spectra of (B) were recorded from the spots illustrated in (C).

4. Discussion

In what follows, the results described in Section 3 will be employed to discuss and compare the technical features of TEY- and XFY-XANES analysis from $\text{Cr}_{0.2}\text{Y}$ and K_2CrO_4 paint fragments (performed at BM08-LISA) and sectioned samples (carried out at ID21) with reference to penetration depth, probability of radiation damage, quality and representativeness of acquired datasets and samples preparation.

As previously observed in different research fields, such as catalysis, [47] superconducting materials, [48] and optoelectronics technology, [49] by carrying out comparative XANES experiments with combined XFY/TEY detection we find evidence that XFY measurements are less surface sensitive than TEY ones (Figs. 1B-2B). At BM08-LISA, under the experimental conditions employed (incident and XFY detection angles from the surface: 70° and 20° , respectively), the X-ray penetration depth at the Cr K-edge (5.9892 keV) (Fig. 5) is higher/comparable with respect to the thickness of the superficial alteration layer, being of the order of $\sim 5\ \mu\text{m}$ and $\sim 20\ \mu\text{m}$ in $\text{Cr}_{0.2}\text{Y}$ and K_2CrO_4 paints, respectively (Figs. 3A-4A), whereas the layer sampled by the electrons is estimated to be of $\sim 150\ \text{nm}$ in $\text{Cr}_{0.2}\text{Y}$ and $\sim 300\ \text{nm}$ in K_2CrO_4 . These calculations, along with the results obtained from thin sections at ID21 (Figs. 3-4), justify the higher relative abundance of Cr^{VI} -species observed in the XFY-XANES spectra with respect to the TEY-ones collected from the surface of paint fragments.

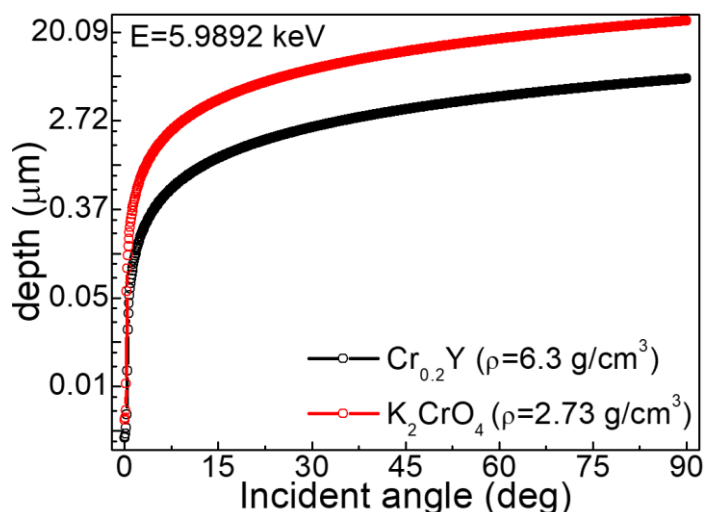


Figure 5. Plot of the X-ray penetration depth vs. incident angle at Cr K-edge energy for $\text{Cr}_{0.2}\text{Y}$ (black) and K_2CrO_4 .

When measuring XANES in TEY mode at the surface of the sample, the beam does not need to be focussed (conversely to the micro-analysis of cross-sections) and can indeed be laterally spread over tens or hundreds of micrometers. It follows that under comparable conditions of flux and energy, the probability of radiation damage is reduced. In that respect, fluences of $\sim 10^6$ - 10^7 ph/ μm^2 were employed at BM08-LISA, that, based on our previous investigations, [29] is of a $\sim 10^4$ - 10^5 factor below the threshold value for safe analysis of chrome yellow paints. Thus, in principle, further improvements in terms of both energy spectral resolution and signal to noise ratio of the acquired spectra might be possible, by selecting smaller energy step values and using longer dwell times per points.

TEY-XANES spectroscopy, as a surface-sensitive method for obtaining solid Cr speciation information directly from the study of the surface of fragments, brings also benefits in terms of samples preparation (much faster and less challenging with respect to the preparation of thin sections) and representativeness of the analyzed material, through access to larger portion of the degraded surface.

On the other hand, within the specific context of our experiment, TEY-XANES approach has a series of drawbacks, namely:

- (i) complete loss of stratigraphic information;
- (ii) constraints related to the size and surface heterogeneity of the analyzed sample, that have to be at least of the order of few mm^2 . This implies that some historical paint fragments, often having sizes of the order of only few μm^2 , will be not analysable by TEY-XANES with X-ray macro-beams;
- (iii) impossibility to measure varnished/coated samples. The presence of a superficial organic layer with thickness greater than 5 μm will completely block the signal of the electrons arising from the paint surface underneath.

In summary, when working with samples of sizes and with a surface homogeneity that are at least of the order of few mm^2 , TEY-XANES spectroscopy can be a valuable primary

approach to assess the elemental speciation of the paint surface, to determine if the acquisition of high lateral resolution stratigraphic information is required or not and, in the case of samples showing a surface heterogeneity, to make a more accurate selection of the sampling areas from which preparing sections for elemental speciation investigations by μ -XANES spectroscopy in XFY mode.

5. Conclusions

In this paper, we have successfully used Cr K-edge TEY-XANES spectroscopy to selectively probe the composition of the surface of chrome yellow ($\text{PbCr}_{0.2}\text{S}_{0.8}\text{O}_4$) and K_2CrO_4 paints and to quantitatively determine the relative abundances of Cr^{VI} and reduced chromium compounds (i.e., Cr^{III} and Cr^{V}) associated to the alteration process of both class of compounds. Overall, the study provided accurate qualitative and semi-quantitative information on the newly formed chromium-based phases within the paint matrix, with fully consistent results by comparing the TEY-XANES results recorded directly from the darkened surface of paint fragments with respect to XFY μ -XANES investigations collected from thin sections of the same samples with submicrometric lateral resolution.

Pros and cons of XANES spectroscopy experiments involving the simultaneous combination of XFY and TEY detection were presented and discussed also in relation to μ -XANES analysis in XFY mode, so as to ultimately decrease the efforts spent on sample preparation and minimize the risks of radiation damage.

Notably, we find that TEY measurements, performed directly on paint fragments, are more surface sensitive than XFY ones. In addition, the possibility to perform TEY-XANES spectroscopy using macro-X-ray beams directly at the paint surface has the great advantages of rendering the preparation of samples both faster and easier (i.e., without need of preparing sections) and of reducing the probability of X-ray induced radiation damage.

Nevertheless, it should be noted that, in the context of our research, the TEY-XANES approach using X-ray macro-beams suffers from the following three main limitations: (i) loss of stratigraphic information; (ii) size and surface homogeneity of the analyzed sample that has to be of the order of few mm²; (iii) possibility to measure only the surface of unvarnished/coated painted fragments.

We conclude that, when working with samples that respect such requirements, TEY-XANES spectroscopy can be profitably employed as a primary approach to study the composition of sample's surface on the mm scale length, thus driving, in a subsequent step, the preparation of sectioned samples and relative measurements at the submicrometric scale length by μ -XANES spectroscopy in XFY mode. Overall, the outcomes of this paper may prompt further research into the application of such combined TEY-XANES/XFY μ -XANES methodological approach also to the study of the degradation pathways of other pigments (e.g., cadmium reds, orpiment, emerald green...) in fragments from both paint matrixes and other kind of painted objects, such as sculptures, frescos, papyri, manuscripts, and others.

Acknowledgments

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