

Why Does Prussian Blue Fade? Understanding the Role(s) of the Substrate

Claire Gervais,¹ Marie-Angélique Languille,² Solenn Reguer,³ Martine Gillet,⁴
Sébastien Pelletier,² Chantal Garnier,⁴ Edward P. Vicenzi,¹ and Loïc Bertrand^{2,3}

¹*Smithsonian Museum Conservation Institute, 4210 Silver Hill Road, Suitland, Maryland 20746, United States**

²*IPANEMA, USR 3461 CNRS/MCC, Synchrotron SOLEIL,
BP48 Saint-Aubin, F-91192 Gif-sur-Yvette, France*

³*Synchrotron SOLEIL, BP48 Saint-Aubin, F-91192 Gif-sur-Yvette, France*

⁴*Muséum national d'histoire naturelle, Centre national de la recherche scientifique,
Ministère de la Culture et de la Communication) CRCC, USR3224,
CP21, 36 rue Geoffroy-Saint-Hilaire, 75005 Paris, France*

Prussian blue (PB) and its analogues are widely studied because of their interesting and promising magnetic and optical properties. The pigment Prussian blue, found in different types of artworks (paintings, watercolors and photographs) is also studied in the frame of heritage science, where its capricious fading behavior under light or anoxia treatment poses problematic conservation issues. PB fading is due to the reduction of iron(III) into iron(II) and depends significantly on the artefact. This paper focuses on the roles of the substrate in affecting the PB structure and modifying the redox process. In particular, X-ray absorption experiments at the Fe K-edge of unfaded and faded PB-paper samples show that changes in the PB structure can happen already by simple contact with the substrate, prior to the fading treatment. Spectrophotometric measurements on series of model PB-paper samples further demonstrate the multiple influences of the substrate and show that not only its chemical composition but also its role as a dispersion and textured medium are significantly altering the fading behavior of PB. A potential roadmap is proposed to rationally investigate the complex fading process of Prussian blue in a substrate.

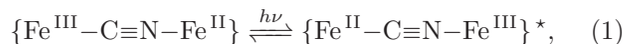
INTRODUCTION

Prussian blue (PB, iron(III) hexacyanoferrate(II)) is a pigment widely used in Europe in the 18th and 19th centuries. Among famous objects containing Prussian blue are paintings from Watteau (1684–1721), Van Gogh (1853–1890) or Picasso (1881–1973), watercolors from Hokusai (1760–1849), but also precious testimonies of our history such as the world’s second official postage stamp “Two Penny Blue”, or one of the first type of color photograph, the trichromies of Louis Ducos du Hauron [1]. Artefacts composed of Prussian blue pose several challenges for conservation efforts, among which are their sensitivity to light or to anoxic treatment [2–4]. Under intense illumination or in a low-oxygen environment, PB undergoes a discoloration. Keeping PB in the dark in ambient air reverses fading. However, it is still unclear whether the pigment fully recovers its original color or if an irreversible change in the structure and chemical state of PB occurs and to which extent [1]. Furthermore, the lightfastness of PB seems to be highly variable and contradictory fading behaviors have been observed in the literature [2, 3, 5, 6]. This variability is still not understood and hampers conservators to predict the light sensitivity of PB-containing artefacts and thus propose adequate conservation recommendations. We here investigate the physico-chemical process at the origin of PB fading in view of rationalizing the fading diversity observed in PB artefacts. For that purpose, we use model samples and focus on the substrate in which PB is embedded.

Prussian blue’s structure exists in two forms, namely

a cation-free form $\text{Fe}_4[\text{Fe}(\text{CN})_6]_3 \cdot 12\text{--}14\text{H}_2\text{O}$ (PBI) and a cation-containing form $\text{KFe}[\text{Fe}(\text{CN})_6] \cdot 3\text{--}5\text{H}_2\text{O}$ (PBS). The crystal structure of PBI can be described as a rock salt structure where $\text{Fe}^{\text{II}}(\text{CN})_6$ and Fe^{III} alternate at the vertices of the cubes [7, 8]. For charge balance, approximately 25% of the $\text{Fe}^{\text{II}}(\text{CN})_6$ sites are vacant and filled with water molecules located at empty nitrogen sites and coordinated to Fe^{III} [8]. The amount of water molecules varies with the synthesis and environmental conditions. The crystal structure of PBS remains elusive. Based on powder diffraction measurements, Keggin et al. described it as the same cubic structure than PBI, but without $\text{Fe}^{\text{II}}(\text{CN})_6$ vacancy and with K^+ cations located in the octahedral interstitial sites [9]. This structure is still nowadays used for schematically describing powder PBS obtained by wet chemistry [4].

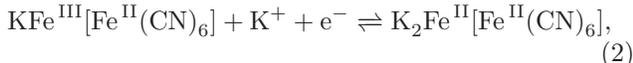
Prussian blue’s color is due to intervalence charge transfer between low-spin Fe^{II} and high-spin Fe^{III} centers, leading to an absorption around 640 nm [10],



where \star denotes an excited state. Thus, color depends on the amount of $\text{Fe}^{\text{II}}\text{--C}\equiv\text{N}\text{--Fe}^{\text{III}}$ units and will be influenced by any disturbance within this pattern, such as the amount of vacancies [11] or the presence of coordinated water within vacancies [12].

Prussian blue’s discoloration is due to the reduction of Fe^{III} into Fe^{II} ions, leading to the formation of $\text{Fe}^{\text{II}}\text{--C}\equiv\text{N}\text{--Fe}^{\text{II}}$ units, where charge transfer of Equation 1 does not happen anymore. This redox process has been particularly investigated for electrochemically syn-

thesized PB (ESPB) films due to their interesting applications in the field of biosensor and ion detection [13–15]. It can schematically be summarized to:



although the process is more complex, with different participations of the Fe^{III} sites, whether their are at the vicinity of Fe^{II} vacancies or fully surrounded by $\text{Fe}^{\text{II}}(\text{CN})_6$ clusters [15]. What stands out is that the redox properties of ESPB depend on the overall ability of cations to enter and exit the structure upon reduction and oxidation, respectively [16].

Light fading of Prussian blue in cultural artefacts is a process identical to Equation 2. Under UV-visible light, Prussian blue artefacts undergo a reduction of Fe^{III} leading to progressive discoloration [4, 17, 18]. The reaction reverses in the dark in ambient atmosphere, with slow re-oxidation of the bleached Prussian blue and recovering of the blue color. The main difference with ESPB lies however in the source of electrons and cations necessary for Equation 2 to take place. In ESPB, electrons are provided by artificial voltage and cations by the electrolytic solution. In PB artefacts, the identification of the electron-donor and the source of cations remains an open question. As the lightfastness of pure PB powder is considered to be excellent by the manufacturers and has been confirmed by ourselves on in-house synthesized PB, the source of cations and electrons has to be external to the pure powder and does not come from an internal rearrangement of the structure.

The presence of a substrate needs therefore to be considered as such an external reservoir. Recent studies on iron ink–paper [19], dyes–paper [20] and Prussian blue–paper systems have indeed shown that this correlation cannot be neglected [17, 18]. Typically, the substrate may participate to the PB reduction process in three different and non self-excluding ways:

(i) The main molecular components from the *substrate itself* may be electron-donor and thus participate to the reduction process of Prussian blue. This could be the case for paper and plant textiles, typical substrates of PB artefacts. For instance, as suggested in earlier studies [18], the chemical composition of paper (cellulose vs lignocellulosic) might have an impact on the kinetics of PB fading.

(ii) The substrate can act as a *carrier of impurities* that may be involved in the reduction process. This is likely to be the case for cyanotypes, an early photographic contrast-printing process based on Prussian blue, where impurities derived from the sensitizer (e.g. oxalate or citrate ions) are present in the paper substrate [4, 17]. Similarly, while Prussian blue is usually stable in paintings, its lightfastness is dramatically decreased in mixtures with lead white pigments [5]. As proposed recently, a possi-

ble explanation is the reduction of Fe^{III} at the surface of Prussian blue triggered by the near vicinity of basic lead compounds [21].

(iii) The *distribution of PB within the substrate* almost certainly plays a role regarding color and color change as well. Particle size and particle distribution are known to influence color by an interplay between absorption and scattering of light [22]. Besides, the distribution of the pigment within the substrate matrix is obviously another factor to be considered with regard to the amount of light exposure received [23, 24]. Finally, the porosity and permeability characteristics of the substrate may well play a role by controlling the access of PB to water and oxygen contents.

The goal of this paper is to probe the potential roles (i)–(iii) of the substrate on the reduction process of PB and propose potential routes for further investigation. To that aim, a methodology based on the analysis of artificially faded model PB artefacts was developed. PB–paper model samples composed of the same PBS powder synthesized in-house and artificially faded were investigated by means of X-ray Absorption Near Edge Structure spectroscopy (XANES), spectrophotometry, Scanning Electron Microscopy (SEM) and Raman spectroscopy. Two types of papers were chosen for their different stability under light and allowed to investigate the correlation between light-sensitiveness of the substrate and PB fading (effect (i)). The role of the substrate in supplying cations necessary for the reduction of PB to occur (effect (ii)) was studied by adding KCl to the PB–paper system and monitoring its fading behavior by means of spectrophotometry. The impact of PB spatial distribution within the substrate (effect (iii)) was studied by comparing the fading behavior of PB–paper systems with different PB concentrations and by probing the spatial and chemical heterogeneity of PB within the sample. Finally, the diversity and complexity of factors influencing the fading behavior of such a rather simple type of system was put into context of the fading of PB artefacts. For that purpose, a series of different PB model artefacts were prepared and their fading behavior monitored by XANES and colorimetry. A potential roadmap is proposed to study more systematically the various structural and chemical factors intrinsic to the PB–substrate systems, which might play a role on the observed fading diversity.

EXPERIMENTAL

Synthesis and sample preparation

Synthesis of Prussian blue (PBS). An equivolume of potassium hexacyanoferrate solution ($\text{K}_4\text{Fe}^{\text{II}}\text{CN}_6$, 0.2M) is added dropwise into a stirred solution of iron sulfate ($\text{Fe}^{\text{II}}(\text{SO}_4) \cdot 5\text{H}_2\text{O}$, 0.1M). The precipitate was filtered, washed with deionized water, dried at 60°C and man-

ually ground.

Paper samples (PBS–Whatman and PBS–Step3).

The two chosen papers are those used in reference [18], that is *Whatman*, composed of 100% cotton cellulose fibres and *Step3* a lignocellulosic paper (75% round wood pulp + 25% soft wood cellulose) containing fillers (kaolin at 20%) and sizing (alum, rosin) [25]. A colloidal solution of synthesized PBS was prepared (25 mg in 1 mL of deionised water) and deposited on *Whatman* and *Step3* papers by brushing (Figure 1 and 2). Weight measurements confirmed the deposition of similar amounts of PBS on both papers. To study the effect of cations in Figure 5, the colloidal solution of PBS was prepared by adding PBS in a 1 M KCl solution instead of water. For colorimetry studies (Figure 4 and 6), small strips of papers were immersed into the colloidal solutions for 5 seconds and then dried at room temperature. The initial sample (named d0) was prepared with a concentration of 25 mg PBS in 1 mL of deionised water. Dilutions were then prepared with deionised water to obtain relative concentrations of 1/2, 1/3, 1/4, 1/9, 1/20, and 1/30, respectively named d1, d2, d3, d4, d5 and d6.

Textile samples (Cotton and Silk). Cotton and silk textiles were dyed according to a modern version [2] of Napier’s recipe [26]. The tissues were soaked for 30 minutes in a solution of iron nitrate ($\text{Fe}^{\text{III}}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$, 7.65 g in 1 L water). After wringing, they were immersed in a solution of 7.56 g iron hexacyanoferrate ($\text{K}_4\text{Fe}^{\text{II}}\text{CN}_6$) and 1.9 mL of sulfuric acid (H_2SO_4 , 0.4M) in 1 L of water. This two-step immersion procedure was then repeated before soaking the clothes 2 or 3 minutes in a solution of aluminum sulphate (0.95 g in 1 L of water) in order to fix the dyeing. Finally, clothes were rinsed in water and dried at room temperature.

Ducos du Hauron samples (DDH). These samples relate to the first color prints introduced in 1868 by Louis Ducos du Hauron [1], which are obtained by superposition of three colored gelatin layers respectively in blue, red and yellow on a paper base, the blue layer being made of Prussian blue. Samples studied here refer only to the PB blue layer gelatin and are those prepared in Lavedrine *et al.* [1], according to Ducos du Hauron’s recipe from 1870.

Cyanotype samples (Cyano). Cyanotypes were prepared with Vélin paper according to Herschel recipe, which consists of an equal proportion mixing of an aqueous solution of ammonium iron(III) citrate (20 g in 100 mL of water) and iron hexacyanoferrate ($\text{K}_3\text{Fe}^{\text{III}}\text{CN}_6$, 8 g in 100 mL of water). The cyanotype is then revealed by exposing it to natural light and briefly rinsed out to stop the process.

Paint samples (Paint). Paintings consist in linen canvas covered with a ground layer composed of a 7% solution of rabbit glue and hydrated calcium sulfate. PBS was ground into linseed oil and applied on top of the ground layer.

Fading treatment. Samples were placed into a light weathering chamber equipped with a metal halide lamp (2500 W Hydrargyrum Medium-arc Iodide (HMI) lamp) and a window glass filter that removed UV radiations below 325 nm. Light aging was performed at $T=25^\circ\text{C}$ and 50%RH. We attempted to mimick alteration during 3400 museum hours (based on a 50 lux exposure) by a fading treatment of about 170 Klux at the surface of the samples and an irradiance of about 1200 W/m^2 . Three series of samples were prepared and underwent the fading treatment during five days. For each sample, a part was hidden from the light as a reference. Color was measured before and immediately after light exposure.

Characterisation methods

Scanning Electron Microscopy. SEM images were recorded on SEM-FEG able to work under controlled atmosphere (ZEISS Supra 55VP) equipped with a 30 mm^2 SDD detector (Bruker). Secondary electrons (SE-SEM) images were saved at low voltages to avoid charge effects. Higher voltages under $p(\text{N}_2)=0.1\text{ mbar}$ were used for backscattered electrons (BSE-SEM) images and EDX mappings, especially to excite the Fe– $\text{K}\alpha$ transition.

X-ray diffraction. X-ray diffraction experiments were performed on beamline CRISTAL at SOLEIL synchrotron. The wavelength of 0.7931 \AA ($E = 15634\text{ eV}$) was extracted from the undulatory beam by means of a double crystal Si(111) monochromator. The powder was introduced in a rotating quartz capillary $700\text{ }\mu\text{m}$ in diameter, mounted on a two circles diffractometer equipped with a 21 crystals multi-analyser. It took less than 30 min in continuous scanning mode to collect high angular resolution diagrams. The diagram is obtained by addition of the 21 channels data. LaB6 and NAC were both used as standards to calibrate the instrumental broadening and potential shift in location of the diffraction peaks. We tested the stability of the powder under the beam by recording several successive scans on the same area. After 10 scans, no difference in the XRD diagram was observed.

Spectrophotometry. Color measurements of the samples were performed with a sphere spectrophotometer X-Rite SP64, which provides reflectance spectra (resolution 10 nm) and $L^* a^* b^*$ color parameters of the sample with L^* : lightness, a^* : red-greenness, b^* : yellow-blueness. The measured area was 4 mm in diameter. The total color difference ΔE was calculated using the CIE 1976 $L^* a^* b^*$ equation for 10° standard observer and standard illuminant D65 [10].

X-ray absorption spectroscopy. XANES spectra were collected at the DiffAbs beamline of synchrotron SOLEIL. The energy scans were measured from 7.05 keV to 7.30 keV with 0.5 eV energy resolution, using the Si(111) double crystal monochromator and a macro beam

spot size about $300\ \mu\text{m}^2$. The incident beam intensity and transmitted signal were measured using Si photodiodes, while the fluorescence signal was monitored using a silicon drift detector. Calibration of the XANES spectra was performed with reference to the inflexion point of a Fe foil at 7112 eV. XANES spectra were processed with the Athena software (Iffit 1.2.9 program suite) [27]. The pre-edge range background was removed by subtracting a Victoreen function and the absorption background was removed using a cubic spline with a normalisation range between 30 and 270 eV after the edge. All XANES spectra presented here are the average of two normalized spectra obtained successively at the same location. No beam damage could be detected (i.e. no change of the spectrum between the two successive spectra and no visible alteration of the sample).

Raman spectroscopy. Raman spectra were recorded using a Thermo Scientific DXR Raman microscope equipped with a 532 nm diode laser. The backscattered light was dispersed by a 9001/mm grating and the Raman signal was detected with a Peltier cooled charge-coupled device. The spectral resolution was in the $5.5\text{--}8.3\ \text{cm}^{-1}$ range. A $100\times$ objective (numerical aperture 0.9) was used, i.e. the estimated laser spot is of approximately 700 nm in diameter. Each spectrum was collected during 5 min, with a power of 500 mW without any observable laser-induced alteration of the samples.

RESULTS AND DISCUSSION

Characterization of PBS

Soluble Prussian blue (PBS) was synthesized by wet chemistry according to the procedure in [28]. Elemental analysis by two methods (EDX and atomic emission spectroscopy) confirmed the synthesis of soluble Prussian blue $\text{KFe}[\text{Fe}(\text{CN})_6]\cdot x\text{H}_2\text{O}$ with a ratio Fe:K of 2:1. The exact stoichiometry of water could not be precisely determined. Traces of sulfur were detected by EDX (normalized percentage of 1% w/t) and likely come from the starting product iron sulfate. Structural purity of the powder was checked by synchrotron X-ray diffraction (the signal-to-noise ratio obtained on a laboratory X-ray diffractometer was insufficient for that task). Figure 1 shows the room-temperature X-ray diffraction pattern, with its best fit (Rwp: 2.33%) by the LeBail pattern decomposition technique [29] implemented in GSAS [30]. The XRD pattern could be indexed in face-cubic symmetry (space group $\text{Fm}\bar{3}\text{m}$, $a = 10.198\ \text{\AA}$), in agreement with the literature [7, 9]. Crystallites sizes were calculated based on the Scherrer formula and estimated to be around 140 nm. Best profile refinement was achieved by taking into account microstrain effects (lattice defects), also responsible for peak broadening. Therefore, although both crystallite size and strain contributions

can be theoretically distinguished, this estimate value of 140 nm needs to be taken with caution. Scanning electron microscopy reveals a PBS powder composed of aggregates with undefined shapes and variable sizes (Figure 1, inset). Distinct nanoparticles are however visible, of a size comparable to the averaged crystallite size estimated by XRD (white circles). Both estimated crystallite and particle sizes fall into the nanoscale regime. This result is not surprising, considering the low solubility of PBS in water leading to its very fast precipitation upon synthesis. One direct consequence of this nanosize is that PBS powder has a large specific surface area and is therefore likely to be very sensitive to its environment.

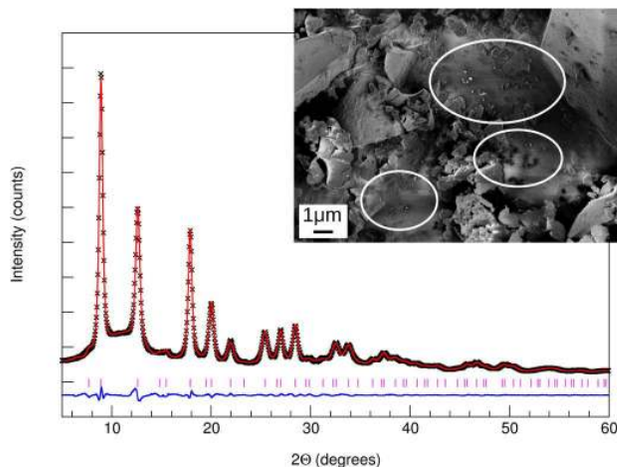


FIG. 1 Diffraction pattern of PBS at 273.0 K, $\lambda = 0.7931\ \text{\AA}$ (cross), its LeBail fit (line) and residues (vertical bars mark the Bragg reflection). Inset: SE-SEM image of PBS powder (Working distance: 2 mm; Accelerating voltage: 0.5 kV, $p=5.10^{-7}$ mbar). Circles indicate regions where distinct nano particles may be observed.

The substrate influences both the structure and redox properties of PB

In order to focus on the substrate and remove any influence due to the PB crystal structure, we investigated PBS–*Whatman* and PBS–*Step3* paper model samples, both prepared with the same PBS powder as described above. As explained in previous work [18], the two papers were chosen for their different stability under light: *Whatman* paper is a pH-neutral cellulosic paper often used as a suitable paper for conservation treatments. In contrast, *Step3* paper is a light-sensitive ligno-cellulosic paper chosen for its high acidity ($\text{pH} = 5.4$) and its ability to readily oxidize [25]. Iron speciation and color of PBS–*Whatman* and PBS–*Step3* samples were analyzed before and after fading by means of XANES at the Fe K-edge and spectrophotometry, respectively. Other

techniques revealed to be not very informative. For instance, extended X-ray absorption fine structure spectra exhibited only weak oscillations due to strong multiple scattering often found in cyano-bridged organometallic compounds [31]. Besides, X-ray diffraction data proved to be ineffective due to the presence of a large amorphous part of the cellulosic substrate and an overlap between the diffraction peaks of PB and the crystalline part of cellulose.

Structural changes of the environment of iron atoms monitored by XANES at the Fe K-edge are significantly different between the two PBS-*paper* samples (Figure 2). Unfaded PBS-*Whatman* features similar spectra than PBS powder and only a slight loss of intensity at the absorption edge is noticed after fading. On the contrary, for PBS-*Step3* systems, pronounced changes happen *both* upon deposition of the PBS powder onto the paper and after fading. Changes may be classified into four regions numbered (i)–(iv) in Figure 2.

After deposition of PBS powder on *Step3* paper, the XANES spectrum of PBS-*Step3* sample exhibits a broadening of the absorption edge (region (i)). It manifests by a lowering in intensity of the maximum of the derivatives of the XANES spectra (Figure 3) and is associated to a slight shift (below 1eV) of the edge towards lower energies. The broadening might be correlated to the presence of more irregular or disordered Fe sites [32], while the shift in energy indicates a slight reduction of Prussian blue. Both features do not seem to be enhanced by light exposure. The unfaded PBS-*Step3* sample also features a shift of ~ 2.5 eV of the position of the 1st oscillation compared to PBS powder (region (ii)). It may be indicative of an increase of the distance to neighbors associated with a geometrical distortion of the $\text{Fe}^{\text{II}}-\text{C}\equiv\text{N}-\text{Fe}^{\text{III}}$ units.

After fading, the XANES spectrum of PBS-*Step3* exhibits additionally a more intense pre-edge peak at 7108 eV (region (iii)). It could be attributed to a change in the oxidation state from Fe^{III} to Fe^{II} , a deviation from the perfect octahedral coordination state of iron clusters or an elongation/contraction of the iron-CN ligand distance [33]. Unfortunately, the pre-peak signal was too noisy to be further processed. Fading also induces a significant change of the slope in the post-peak region (range 7125–7162 eV, region (iv)). In cyanoferrates, this region is governed by very large multiple-scattering effects due to the presence of the $\text{C}\equiv\text{N}$ pattern [34]. A change in this region of the spectrum thus involves a modification of the $\text{Fe}^{\text{II}}-\text{C}\equiv\text{N}-\text{Fe}^{\text{III}}$ units, either geometrically or in terms of oxidation state. It is worth noting that the change of slope is visible on all five XANES measurements performed on the faded sample at different locations (red-orange curves), but the degree to which the slope changes varies significantly from point to point. It thus indicates that the modification

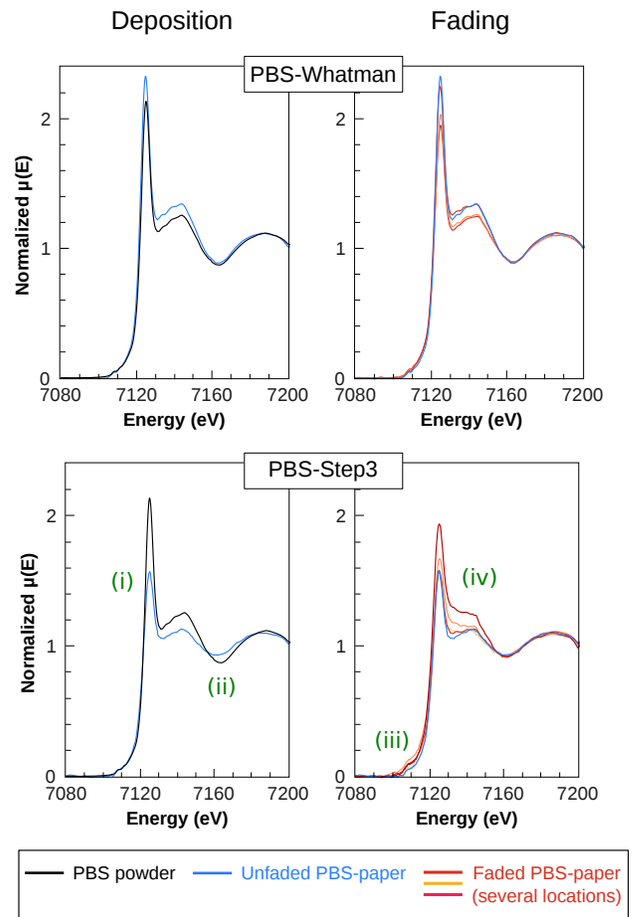


FIG. 2 XANES spectra of PBS powder (black) and PBS-*paper* samples before (blue) and after fading (red-orange). Due to heterogeneity observed in the faded PBS-*paper* samples, measurements were performed at three to five different locations of each sample (red-orange curves). For PBS-*Step3* samples, structural variations are observed both before and after light exposure and occur in four regions of the spectra denoted (i)–(iv).

in the spectral region (iv) is present all over the sample but with some spatial heterogeneity.

Reflectance curves of PBS-*paper* samples are presented in Figure 4. Before light exposure, PBS-*Whatman* and PBS-*Step3* samples already exhibit a different spectrum at the position of the maximum of reflectance (450 nm and 465 nm respectively). However, this difference between the two unfaded PBS-*paper* samples cannot be straightforwardly related with those observed on XANES spectra, as the blue-green region (400–550 nm) is dominated by interband transitions of the papers (black curves).

After fading, the change of reflectance curves differs also for PBS-*Whatman* and PBS-*Step3* samples, with ΔE values of 3.8 and 13.0, respectively. The distinction takes place in the range 400–550 nm, where reflectance

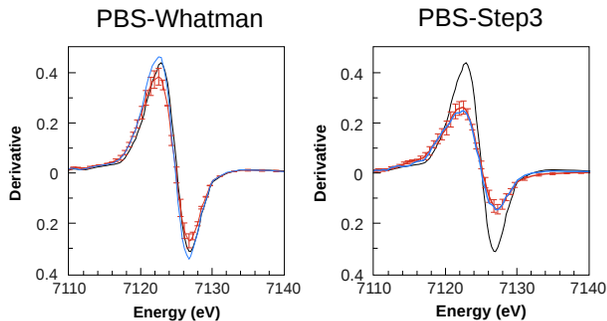


FIG. 3 Impact of the substrate on the absorption edge (region (i)). Derivative of the XANES absorption spectra illustrating the broadening of the absorption edge for PBS–*Whatman* and PBS–*Step3* samples. Powder (black) and PBS–*paper* samples before (blue) and after fading (red). Errors bars correspond to standard deviations on 4 and 5 measurements for PBS–*Whatman* and PBS–*Step3* respectively.

of both PBS–*Step3* and *Step3* paper alone are significantly decreased (red and gray curves, Figure 4, right). This may be attributed to a phenomenon of “paper yellowing”, a result of the oxidation of the lignin present in *Step3* paper, which releases chromophores absorbing in the blue-green region [35]. In the wavelength domain 550–700 nm, absorption by $\text{Fe}^{\text{II}}-\text{C}\equiv\text{N}-\text{Fe}^{\text{III}}$ occurs and gives its color to Prussian blue (Equation 1). The photoreduction of Prussian blue, leading to non-absorbing $\text{Fe}^{\text{II}}-\text{C}\equiv\text{N}-\text{Fe}^{\text{II}}$ blocks would thus impact principally this region and lead to a higher reflectance. Interestingly, the fact that both samples behave similarly in this region might thus indicate that Prussian blue reduction happens to the same extent in both samples. With respect to geometrical distortions of the crystal lattice of Prussian blue, they would lead to band broadening or shifting [36]. The fact that they are not visible does not demonstrate their absence, as the low spectral resolution (10 nm) associated to the overlap of the intervalence charge transfer in Prussian blue with paper absorption bands might hide such spectral features [24].

Clearly, all these variations upon deposition and fading need to be further investigated to understand the exact types of structural modifications taking place during the fading process. However, considering the ability of Prussian blue and its analogues to accommodate ions within the lattice [36] and the presence of a mobile and disordered water network [37], one may surmise that the geometrical distortions and other structural changes observed in PBS are likely due to a reorganisation of the water network and/or migration of ions within the structure, either in tetrahedral sites or in iron vacancies.

Considering the role of the substrate, results shown here allow us to draw some important conclusions concerning the structural and redox properties of Prussian

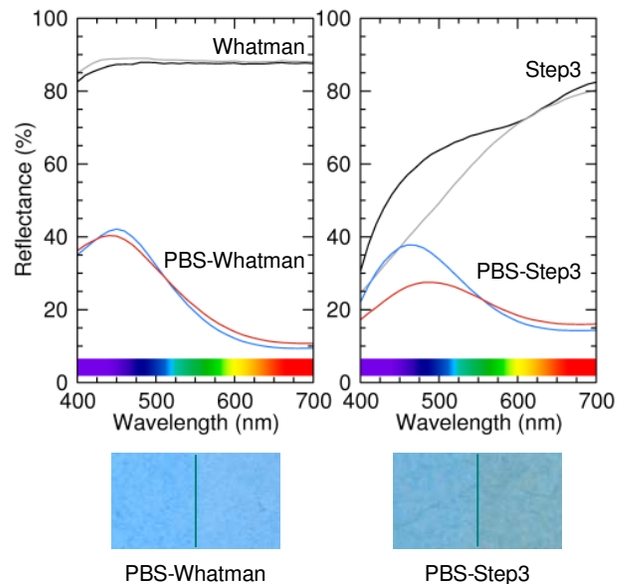


FIG. 4 Reflectance curves of PBS–*paper* samples before (blue) and after fading (red). For reference, reflectance curves of papers alone before (black) and after fading (gray) are indicated. Bottom: optical photograph of the unfaded and faded samples (respectively left and right parts of the photographs).

blue on the two papers.

First, compared to the reference PBS powder, structural changes happen already on PBS–*Step3* before fading. As observed in XANES spectra, they likely involve geometrical distortions of the Prussian blue lattice associated with a slight reduction of PB. That is, the structure of the pigment has already been modified by simple “contact” with the *Step3* paper. A reaction occurred spontaneously between the pigment and *Step3* paper that did not require visible light energy for activation. Several components in *Step3* paper could play such a catalytic role. For instance lignin is known to readily oxidize [38] and adsorption of chromium ions onto a lignocellulosic substrate resulted in spontaneous reduction of the metal and oxidation of lignin [39]. Also, alum (aluminium double salts) is responsible for the low pH of the paper (pH = 5) and might have induced a structural change of PB. To the contrary, no difference is observed between PBS–*Whatman* and PBS powder, a result showing that *Whatman* paper (composed of pure cellulose) is neutral with respect to the pigment in the frame of the experiment and does not induce spontaneous modification of its structure.

Second, changes of the PBS structure upon fading are not of the same nature on both papers, as exemplified by the variation in XANES spectra of the post-edge region occurring in PBS–*Step3* but not in PBS–*Whatman*. As just stated, the structure of PBS differs on both papers before fading (since *Step3* spontaneously modifies it) and this may be sufficient to explain the different fading

behaviors. However, UV-vis reflectance spectra clearly show a higher degradation of *Step3* paper compared to *Whatman*, a process which in turn may influence the photoreduction of Prussian blue. As comparison, acidity, such as found in *Step3*, was shown to impact the redox process of ESPB films by allowing hydrated proton transfer between the solution and Prussian blue [37]. Also, lignin present in *Step3* might be involved in the redox process by playing a role as a radical scavenger [18, 40].

In summary, these results indicate that the type of substrate has a major impact on *both* the structure and the fading process of Prussian blue and that any study of light sensitivity of PB artefacts should take it explicitly into account. While it is shown that *Step3* paper has more influence on PB fading than *Whatman*, the reasons behind are not resolved yet. As stated in the former paragraphs and in previous work [18], several hypotheses are plausible, such as the presence of lignin, acidity or the different distribution of PB within paper fibers. It seems therefore that the PBS–*Step3* system is still too complex if the goal is to rationalize, one by one, the various roles of the substrate. We have thus pursued our investigation by concentrating on the PBS–*Whatman* system, which is composed of two main components only (PBS and cellulose).

What are the roles of the substrate in the PB reduction process?

Although the relative weight of the various roles of the substrate cannot be estimated at this stage of the study, we have focused here on three properties of the substrate which seem to significantly influence PB fading: its role as a cation reservoir, as a dispersion medium and as a textured medium.

The substrate as a cation reservoir?

According to Equation 2, cations are necessary to ensure electroneutrality of the reduced PBS. The compound PBS contains potassium (and possibly sulfur coming from the synthesis). However, PBS powder alone (i.e. without substrate) does not fade. This implies that we must look at the cations brought by the substrate. In Figure 5, the influence of potassium on the fading process was tested by monitoring color changes of PBS–*Whatman* systems enriched in KCl.

The combined presence of KCl and light leads to a strong decrease of the absorption intensity in the 550–700 nm region (red curve). As *Whatman* paper enriched in KCl does not change upon fading (and KCl does not absorb in this region), this result indicates that the photoreduction process has been enhanced by ad-

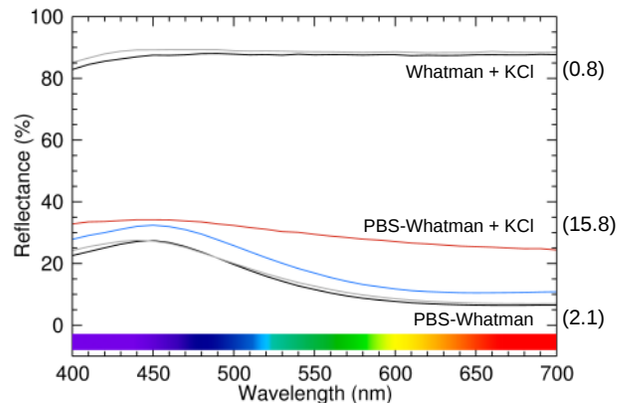


FIG. 5 Reflectance curves before (blue) and after fading (red) of PBS–*Whatman* systems enriched in KCl. Values in brackets refer to ΔE values. Control reflectances curves before and after fading of *Whatman* paper enriched in KCl and PBS–*Whatman* without KCl (sample d1) are indicated in black and gray, respectively.

dition of KCl. Several explanations can be advanced: First, a migration of K^+ cations within the PB structure could take place to maintain electroneutrality upon reduction of Fe^{III} ions. This migration phenomenon has been well established in voltametric cycles of ESPB [16]. Second, KCl is hygroscopic and its presence is likely to change the humidity content, *both* within the substrate and within the PB crystalline network. Concerning PB, water is known to influence the redox process in ESPB. Indeed, the ionic compensation following the change in ratio $Fe^{III}:Fe^{II}$ may be given by electrolytes within the solution (e.g. K^+) but may also be obtained by changes in the composition (H^+/H_3O^+) [41] or in the amount [42] of coordinated water molecules in the PB crystalline network. Concerning the substrate, an increase of moisture may alter the stability of the cellulose [43] and hence participate indirectly to the redox process. XANES at the Fe K-edge and XRD on aliquots of samples shown in Figure 5 are currently planned and should allow to better understand the influence of KCl on PB fading.

The substrate as a dispersion medium?

The influence of the concentration of PB on fading is shown Figure 6. Diluted samples exhibit a larger photoreduction of Prussian blue according to an exponential law between ΔE and relative PB concentration. This trend differs from results on the impact of water dilution on the fading of PB watercolors (PB + gum arabic) [44] but has been verified by two independent sets of measurements on our systems and is also observed for PBS–*Step3* systems. This result shows the difficulty to compare PB fading for different substrates and highlights the need to use model samples with similar concentrations, i.e. with

approximately similar percentage of reflectance in the region 550–700 nm, when comparing fading behaviors and color measurements.

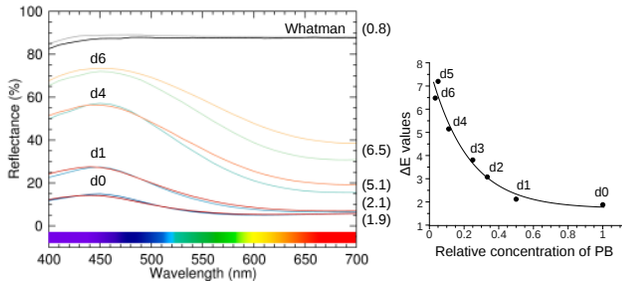


FIG. 6 Left: Reflectance curves of PBS–*Whatman* paper systems at different PB concentrations, before (blue shades) and after fading (red shades). Values in brackets refer to ΔE values. For reference, reflectance curve of *Whatman* alone before (black) and after fading (gray) is indicated. Right: Relative PB concentration vs. ΔE , showing the entire relative concentration series [d0–d6]. Data could be fitted with an exponential law of the kind $a + \exp(-x/b)$ (plain line).

The substrate as a textured medium?

Finally, even in a given sample, the distribution of Prussian blue within a textured substrate such as paper leads to spatial and compositional heterogeneities. SEM images of PBS–*Whatman* show a non-homogeneous distribution of PBS in the paper. The largest part of PBS is present in the form of a cracked crust located within, and on top of the fibers. Its characteristics are similar to the PBS powder. However, microparticles with well defined shapes are also visible, which we could not observe in PBS powder. This raises the question whether part of PBS did change upon deposition on the substrate. Indeed, the deposition step implied to put again PBS in solution. While insoluble in water, it forms sub-microparticles (colloid), that could serve as nuclei for further growth once onto the substrate. Raman spectroscopy performed on a PB particle and within a blue fiber show a difference in the 1A_g C \equiv N stretching vibration band at around 2150 cm^{-1} characteristic of PB [45]. The weak band around 2090 cm^{-1} corresponds to the E_g mode of the C \equiv N stretching vibration and does not seem to differ between the two types of PBS. Finally, the peak around 2120 cm^{-1} present in one of the spectra of PB particles could not be attributed but has been assigned to Prussian blue in reference [46]. This bimodal distribution in particle size and location has thus a direct impact on the PB structure which is heterogeneous *at this scale*. Note also the chemical heterogeneity of the cellulosic fibers, which exhibit a non-uniform luminescence on optical photographs (Figure 7). It emphasizes the

fact that a medium such as paper, itself textured and chemically heterogeneous, obviously leads to more than a simple dispersion of the pigment. It interacts chemically and topologically with it, to form ultimately a complex material with paper-pigment interfaces varying in area, chemical and structural type.

Owing to this multiple heterogeneity, it is particularly important to adapt the scale of the measurement to the property investigated, gather measurements at *all* the characteristic scales of the material [47] and be especially careful when comparing and correlating measurements obtained at different scales and information depths, such as Raman (estimated spot of 700 nm), XAS (averaging PB over a surface of $300\text{ }\mu\text{m}^2$) or colorimetry (surface of 0.5 cm^2).

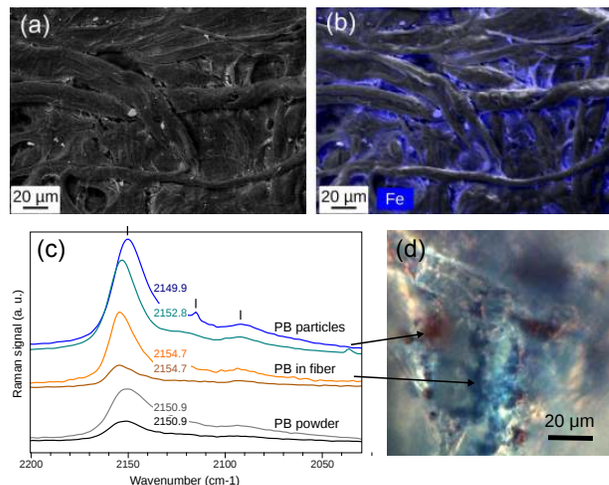


FIG. 7 Illustration of the microscale heterogeneity of Prussian blue in morphology and structure. (a) BSE-SEM image of PBS–*Whatman* (Working distance: 6.5 mm; Accelerating voltage: 12 kV and $p(\text{N}_2) = 0.1\text{ mbar}$). (b) BSE-SEM image with Fe–K α EDX mapping (300 px x 225 px). (c) Raman spectra of PBS powder and in PBS–*Whatman*. Vertical bars mark the three peaks attributed to PB and values indicate peak maxima corresponding to the 1A_g mode of the $\sigma(\text{CN})$ stretching vibration band. (d) Optical micrograph of PBS–*Whatman* sample, showing the presence of PBS particles (brown) and blue colored cellulosic fibers.

Fading behavior of model PB artefacts: a variability confirmed

Results presented so far have shown that even in “simple” PB–substrate systems, complex correlations between multiple factors influence the fading behavior of Prussian blue. This complexity should be obviously also considered for real PB artefacts, where additional chemical or spatial heterogeneities occur [47]. Table I presents a series of PB model artefacts, all subjected to the same light treatment. Colorimetry measurements confirm a

large diversity in the fading behavior, which does not depend on light or other environmental variations (e.g. temperature, humidity or anoxia) but is intrinsic to the samples.

XANES spectra at the Fe K-edge for three of these model-artefacts before and after fading further evidence the variety of both PB structures and fading behaviors (Figure 8). Even before light treatment, PB spectra have different shapes (see pre-edge and post-edge regions), confirming the influence either of the synthesis conditions on the PB structure [45] or, as demonstrated here, the modification of the structure of Prussian blue upon deposition. After fading, a shift of the absorption edge towards lower energies is observed on at least two samples, suggesting a lower average oxidation number of iron atoms, in agreement with a fading process due to a reduction of PB. As well, the extent to which XANES spectra differ after fading seems to be correlated to the amount of color change. For instance, a clearly observable shift of the edge is observed for DDH samples, together with a larger ΔE value.

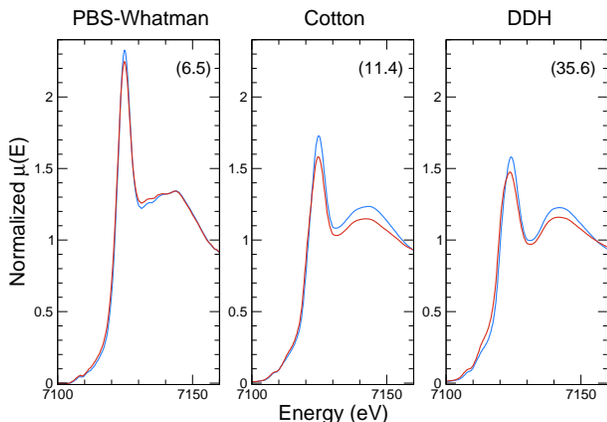


FIG. 8 Normalised XANES spectra of unfaded (blue) and faded (red) Prussian blue artefacts on paper (PBS-*Whatman*), textile (Cotton) and gelatin (DDH) samples. ΔE values are indicated in parentheses at the top of each XANES spectrum.

CONCLUSION

In order to further understand the variability of the fading behavior of Prussian blue in cultural artefacts, several factors have been shown to be particularly influential and deserve increased attention.

First of all, the structure of Prussian blue is complex and possesses interrelated features which are likely to influence the final fading behavior of the pigment. In particular, the location of cations within the structure, the water network and the ordering of vacancies determine the availability of the $\text{Fe}^{\text{II}}-\text{C}\equiv\text{N}-\text{Fe}^{\text{III}}$ units and

could play a significant role in the final capacity of the structure to photoreduce. Investigation by XANES and spectrophotometry of PBS-*paper* samples have shown that the process of PB photoreduction goes beyond a mere reduction of iron(III) and implies complex and correlated changes within the structure, such as significant geometrical distortions of the iron coordination sphere, migration of cations within the structure and/or rearrangement of the water network. The next mandatory steps to further elucidate the interrelationship between PB structure, color and light sensitivity will certainly involve detailed structural interpretation of the XANES spectra, coupled with other analytical techniques such as X-ray diffraction or Mössbauer spectroscopy.

Second, the structure of PBS and its variations are highly dependent of the substrate, which definitely contributes to the various fading behaviors observed. The substrate does indeed play a role at several levels: it can modify the structure of Prussian blue by simple contact (e.g. *Step3* paper), play an active role during its photoreduction (e.g. by providing K^+ that catalyze the redox process), or it simply contributes indirectly to the fading process, either by participating to the color change (e.g. yellowing of *Step3* paper) or by influencing the distribution of PB particules and thus their exposure to light. Owing to its various roles, further in-depth investigation of the substrate’s photodegradation [48, 49] in relation with PB photoreduction will certainly help in getting a general picture of the fading mechanism.

As summarized in Figure 9, the redox properties of PB-substrate systems are influenced by a number of internal and external factors, which are likely to be correlated one to the other. A systematic study of all these features, with a back-and-forth investigation of the entire system at both microscopic and macroscopic levels is required to further understand this complex physico-chemical process generally termed “Prussian blue fading”. This approach is here applied to Prussian blue but is also valid for many other cultural heritage materials featuring the trio pigment-substrate-environment. In a broader view, this study, like other recent papers [19, 20], confirms the importance and interest to investigate thoroughly the physico-chemical processes happening in cultural materials and the bias that could be brought by oversimplistic model based on unsupported pigments. In that respect, model samples where history, composition and environmental factors are controlled are an invaluable tool to conceptualize these processes. They help in finding the principal causes for degradation and factors which need to be particularly monitored. We hope to demonstrate the interest of developing such an approach based on models, which, coupled to the characterization of real cultural artefacts, could help conservators and restorators in choosing *knowingly* appropriate conservation treatments, on a more rational basis than immediate, visible impacts.

Sample	Syn. †	Substrate	Major components	Expected ‡ impurities / additives	ΔE values
DDH	in-	gelatin	peptides / proteins	yes	DDH
Paint	PBS	oil	fatty acids	no	Paint
Silk	in-	textile	proteins	yes (sp*)	Silk
PBS–Step3	PBS	paper	cellulose / lignin	yes (fs*)	PBS-Step3
Cotton	in-	textile	cellulose	yes (sp*)	Cotton
Cyano	in-	paper	cellulose	yes (sp*)	Cyano
PBS–Whatman	PBS	paper	cellulose	no	PBS-Whatman
PBS	PBS	–	–	no	PBS (no fading observed)

† Synthesis realized in-situ (in-) or ex-situ (powder PBS)
‡ Expected to be present beside the major components of the substrate
*sp: starting products from synthesis; fs: fillers and sizing

TABLE I Left: Basic sample description showing the diversity of the model PB artefacts investigated. Major components and impurities in the samples are given based on knowledge about the substrate and/or about sample preparation procedure. Right: Color changes (ΔE values) induced by a 5-days light exposure.

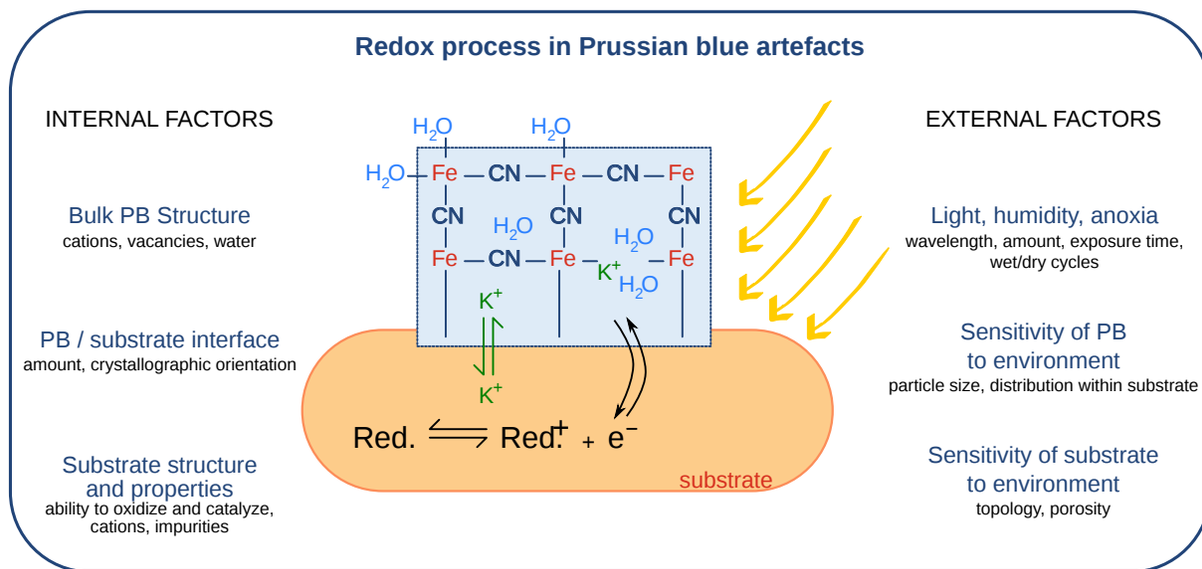


FIG. 9 Summary of internal and external factors playing a role on the redox processes in Prussian blue–substrate systems. In this paper, we have particularly emphasized on the impact of “Bulk PB structure” and “Substrate structure and properties” on the fading behavior of model PB–paper systems. The impact of “Light, humidity, anoxia” was partly investigated on similar PB–paper systems in Gervais et al [18].

ACKNOWLEDGEMENTS

K. Chaouchi and S. Blanchandin are deeply thanked for their help in preparing the samples, Dr. E. Elkaïm for his help on the CRISTAL beamline and Dr. P. Dumas for access to the Raman spectrometer. We thank also Dr. P. Ropret for the paint samples and M. Ballard for information about Prussian blue dyeing. C. Gervais would like to acknowledge the Swiss National Science Foundation for partial funding of this project through the grant n°138986, as well as Dr. T. Wüst and Prof. P. Laszlo for useful comments. We thank Dr. R.J. Koestler for suggesting us to investigate Prussian blue fading in cultural artefacts. This work has been developed as part of the IPANEMA / Smithsonian Institution agreement on science cooperation.

* Present affiliation: Bern University of the Arts, Fellerstrasse 11, CH-3027 Bern, Switzerland; claire.gervais@bfh.ch

- [1] B. Lavédrine, C. Pesme, C. Garnier, and M. Gillet, *Support/Tracé* **5**, 38 (2005).
- [2] S. Rowe, *Studies in Conservation* **49**, 259 (2004).
- [3] R. Koestler, E. Parreira, E. Santoro, and P. Noble, *Studies in Conservation* **38**, 265 (1993).
- [4] M. Ware, *Journal of Chemical Education* **85**, 612 (2008).
- [5] J. Kirby and D. Saunders, in *National Gallery Technical Bulletin*, Vol. 25, edited by A. Roy (National Gallery Company Limited, 2004) pp. 73–99.
- [6] C. Asai, *Zeitschrift für Kunsttechnologie und Konservierung* **17**, 261 (2004).

- [7] H. J. Buser, D. Schwarzenbach, W. Petter, and A. Ludi, *Inorganic Chemistry* **16**, 2704 (1977).
- [8] F. Herren, P. Fischer, A. Ludi, and W. Halg, *Inorganic Chemistry* **19**, 956 (1980).
- [9] J. F. Keggin and F. D. Miles, *Nature* **137**, 577578 (1936).
- [10] K. Nassau, *The Physics and Chemistry of Color: The Fifteen Causes of Color*, Vol. 1 (John Wiley and Sons, 1983) pp. 147–148.
- [11] R. J. Mortimer and D. R. Rosseinsky, *Journal of the Chemical Society-Dalton Transactions* **9**, 2059 (1984).
- [12] V. B. Nechitayilo, V. I. Styopkin, Z. A. Tkachenko, Y. G. Goltsov, V. P. Sherstyuk, and V. V. Zhilinskaya, *Electrochimica Acta* **40**, 2501 (1995).
- [13] K. Itaya, K. Shibayama, H. Akahoshi, and S. Toshima, *Journal of Applied Physics* **53**, 804 (1982).
- [14] A. A. Karyakin, *Electroanalysis* **13**, 813 (2001).
- [15] J. Agrisuelas, P. R. Bueno, F. F. Ferreira, C. Gabrielli, J. J. Garcia-Jareno, D. Gimenez-Romero, H. Perrot, and F. Vicente, *Journal of the Electrochemical Society* **156**, P74 (2009).
- [16] B. J. Feldman and O. R. Melroy, *Journal of Electroanalytical Chemistry* **234**, 213 (1987).
- [17] M. Ware, *Cyanotype: The History, Science, and Art of Photographic Printing in Prussian Blue* (National Museum of Photography, Film, and Television, Bradford, UK, 1999).
- [18] C. Gervais, M.-A. Languille, S. Reguer, M. Gillet, S. Chagnot, F. Baudelet, E. P. Vicenzi, and L. Bertrand, *Applied Physics A*, accepted (2012).
- [19] V. Rouchon, M. Durantou, C. Burgaud, E. Pellizzi, B. Lavedrine, K. Janssens, W. de Nolf, G. Nuyts, F. Vanmeert, and K. Hellemans, *Analytical Chemistry* **83**, 2589 (2011).
- [20] J. Thomas, J. H. Townsend, S. Hackney, and M. Strlic, *Polymer Degradation and Stability* **95**, 2343 (2010).
- [21] L. Samain, G. Silversmit, J. Sanyova, B. Vekemans, H. Salomon, B. Gilbert, F. Grandjean, G. J. Long, R. P. Hermann, L. Vincze, and D. Strivaya, *Journal of Analytical Atomic Spectrometry* **26**, 930 (2011).
- [22] W. Caseri, *Journal of Materials Chemistry* **20**, 5582 (2010).
- [23] L. Matisová-Rychlá, V. Bukovský, J. Rychlý, and M. Pleteníková, *Macromolecular Symposia* **247**, 349 (2007).
- [24] H. Liang, *Applied Physics A* **106**, 309 (2011).
- [25] *The Effects of Air Pollutants on the Accelerated Ageing of Cellulose Containing Materials – Paper*, Tech. Rep. (TNO Report BU 3.94/1068/JH, Delft, The Netherlands, 1994 http://www.viks.sk/chk/res_4.95_209_233.doc).
- [26] J. Napier, *A Manual of Dyeing and Dyeing Receipts*, 3rd ed. (Charles Griffin and Co, 1875).
- [27] M. Newville, *Journal of Synchrotron Radiation* **8**, 322 (2001).
- [28] E. Reguera, J. Fernandez-Bertran, A. Dago, and C. Diaz, *Hyperfine Interactions* **73**, 295308 (1992).
- [29] A. Le Bail, H. Duroy, and J. Fourquet, *Materials Research Bulletin* **23**, 447452 (1988).
- [30] A. C. Larson and R. B. Von Dreele, *General Structure Analysis System (GSAS)*, Tech. Rep. (Los Alamos National Laboratory Report LAUR 86-748, 2000).
- [31] K. Hayakawa, K. Hatada, P. D’Angelo, S. Della Longa, C. R. Natoli, and M. Benfatto, *Journal of the American Chemical Society* **126**, 1561815623 (2004).
- [32] G. A. Waychunas, M. J. Afted, and G. E. Brown, *Physics and Chemistry of Minerals* **10**, 1 (1983).
- [33] T. Westre, P. Kennepohl, J. DeWitt, B. Hedman, K. Hodgson, and E. Solomon, *Journal of the American Chemical Society* **119**, 6297 (1997).
- [34] A. Bianconi, M. Dell’Ariccia, P. Durham, and J. Pendry, *Physical Review B* **26**, 65026508 (1982).
- [35] A. L. Dupont, C. Egasse, A. Morin, and F. Vasseur, *Carbohydrate Polymers* **68**, 1 (2007).
- [36] J. C. Wojdel and S. T. Bromley, *Journal of Physical Chemistry B* **110**, 24294 (2006).
- [37] K. Kim, I. Jureviciute, and S. Bruckenstein, *Electrochimica Acta* **46**, 41334140 (2001).
- [38] J. Rychly, L. Matisova-Rychla, V. Bukovsky, M. Pletenikova, and M. Vrska, *Macromolecular Symposia* **231**, 178 (2006).
- [39] L. Dupont, J. Bouanda, J. Ghanbaja, J. Dumonceau, and M. Aplincourt, *Journal of Colloid and Interface Science* **279**, 418 (2004).
- [40] J. A. Schmidt, C. S. Rye, and N. Gurnagul, *Polymer Degradation and Stability* **49**, 291 (1995).
- [41] N. R. de Tacconi, K. Rajeshwar, and R. O. Lezna, *Chemistry of Materials* **15**, 3046 (2003).
- [42] D. R. Rosseinsky, J. S. Tonge, J. Berthelot, and J. F. Cassidy, *Journal of the Chemical Society-Faraday Transactions I* **83**, 231 (1987).
- [43] H. D. Burgess, “Historic textile and paper materials,” (American Chemical Society, 1986) Chap. 20, pp. 363–376, gel Permeation Chromatography. Use in Estimating the Effect of Water Washing on the Long-Term Stability of Cellulosic Fibers.
- [44] A. Lerwill, J. H. Townsend, H. Liang, J. Thomas, and S. Hackney, *e-Preservation Science* **5**, 17 (2008).
- [45] S. F. A. Kettle, E. Diana, E. M. C. Marchese, E. Boccaleri, and P. L. Stanghellini, *Journal of Raman Spectroscopy*, 2006 (2011).
- [46] K. Castro, M. D. Rodriguez-Laso, L. A. Fernandez, and J. M. Madariaga, *Journal of Raman Spectroscopy* **33**, 1725 (2002).
- [47] L. Bertrand, M. Thoury, and E. Anheim, *Journal of Cultural Heritage*, <http://dx.doi.org/10.1016/j.culher.2012.09.003> (2012).
- [48] M. Strlic and J. Kolar, eds., *Ageing and stabilisation of paper* (National and University Library, Ljubljana, Slovenia, 2005).
- [49] M. Manso and M. Carvalho, *Spectrochimica Acta Part B: Atomic Spectroscopy* **64**, 482 (2009).