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D. Chiriu, M. Pala, F.A. Pisu, G. Cappellini, P.C. Ricci, C.M. Carbonaro – **“Time through colors: a kinetic model of Red Vermilion darkening from Raman spectra”** Dyes and Pigments - Volume 184, January 2021, 108866

The publisher's version is available at:

<https://dx.doi.org/10.1016/j.dyepig.2020.108866>

When citing, please refer to the published version.

Time through colors: a kinetic model of Red Vermilion darkening from Raman spectra

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Abstract

Darkening of Red Vermilion (cinnabar) is a well-known phenomenon which needs, at the moment, a full comprehension of its nature. Some paintings reveal a disfiguring process of blackening degradation of Red Vermilion, albeit other relics preserve the original color during the time. The presence of halide elements like Cl⁻ ions, in addition to the light exposure, represents the most alleged cause to explain the darkening phenomenon. However, the real effect of chlorine impurities is still not completely understood. In this study, starting from a multi-technique characterization, we propose a kinetic model of mutual composition of alpha/beta cinnabar derived by the experimental darkening of pure alpha synthetic cinnabar intentionally doped with Cl⁻ ions and treated under prolonged UV light exposure. The model was further applied to ancient samples by analyzing the Raman spectra of antique cinnabar pigments belonging to manuscripts of XIII-XVII centuries.

Keywords: Vermilion darkening; kinetic model; pigment ageing; cinnabar degradation;

1. Introduction

Color technology accompanies the human history since the Neolithic period [1–4]. Among the principal pigments used for decorating artifacts or paintings, mercury sulfide HgS, known as cinnabar, assumed a prominent role as red color for many applications. In fact, cinnabar was used not only as pigment, but it found employment also as cosmetic, medicine and red ink. Several archeological sites report the use of cinnabar in ancient relics like Chinese potteries, Asian lacquer artifacts or mural paintings in Pompeii. Known also as Vermilion, it was very appreciated during the time, especially in the Middle Ages, Renaissance and Baroque, being found, among the others, in many paintings of Titian, Botticelli, Rembrandt and Rubens [2,5].

This pigment is considered generally stable, when it is pure and in absence of particular agents determining the so-called phenomenon of Vermilion darkening [2,6,7]. Indeed, there are specific impurities which may cause the effect of HgS degradation and, as evidenced in the literature, catalytic elements (halides for examples) accelerate the darkening process [2,5,8]. The mentioned degradation affected numerous artifacts during the time and involved not only paintings, but it is quite evident also in mural frescoes, obviously exposed to external agents (light, relative humidity and salt content in the atmosphere)[1,2,5,6]. Famous cases concern Egyptian and Pompeii frescoes as well as Renaissance mural paintings [2,9]. Considering the continuous studies on this phenomenon, as evidenced by recent works [2,7,10], a deep understanding of the Vermilion darkening problem can contribute to evaluate the conservation state of many above mentioned artifacts and the efforts for their preservation.

1.1 Structure of HgS

Several works report the crystal structure of the HgS system [11–13]. As reported by Ballirano et al. [1], the crystal structure of cinnabar can be identified in three different phases: red α -HgS having trigonal

75 structure (P3₁21 space group) with lattice constants a= 4.1489 Å and c= 9.4947 Å; black cubic β-HgS
76 (F43m space group) with lattice constant a= 5.8461 Å; γ-HgS (hypercinnabar) with hexagonal structure
77 and lattice constants a=7.0103 Å and c= 14.1307 Å. Due to the impurities in the composition, the α-HgS
78 phase converts to cubic metacinnabar β-HgS, in the temperature range between 373 K (HgS 99.97 %wt)
79 and 635 K (HgS 99.999 %wt) [11,14]. This transformation presents a value of enthalpy ranging from 2
80 to 8 kJ/mol. In addition, it was found a very low kinetic of the reverse transformation from β-HgS to α-
81 HgS phase at room temperature [1]. At 798 K the metacinnabar transforms to hypercinnabar.
82 As enlightened above, many works indicated the purity of HgS as the discriminant element condition to
83 prevent its darkening, thus several studies identified the production process of vermilion pigment,
84 especially used in art, as the main cause of chemical contamination [1,2,5–7,9,10].

85 1.2 Production of HgS pigment

87 The processes for obtaining vermilion involve two different techniques, sharing the characteristic that
88 both methods imply the obtaining of vermilion via conversion reaction of black β-HgS. The *dry method*
89 consists in the heating of mercury with sulphur in order to obtain the black phase β-HgS and thereafter
90 the α-HgS phase by a second heating step [9]. In this case the direct reaction leading to the formation of
91 vermilion is complex and the temperature is the crucial parameter to exceed the kinetic barrier of the
92 cubic to trigonal phase conversion. The second method provides vermilion by a *wet process*, much easier
93 than the dry one. This method starts also from β-HgS formed by the reaction of sulphur with mercury or
94 mercury salts. The complete conversion of metacinnabar to vermilion requires strong basic media and
95 moderate temperature (NaOH 4M – 60°C) [6,9].

96 1.3 Darkening phenomenon

98 Many references suggest different mechanisms ascribable to the darkening of cinnabar:

- 99 • *Phase transformation* from red hexagonal α-cinnabar into black cubic β-cinnabar (called also
100 metacinnabar): this case, formulated by Feller [15] and further studied in many other references,
101 implies that reflectance properties of red cinnabar are altered by light exposure because of the
102 production of dark phase metacinnabar [16–18]. The effect is accentuated by the original composition
103 of the pigment in addition to its production process (*wet* or *dry* synthesis [9]).
- 104 • *Formation of transition chloride compounds* (mercury chlorides or sulfochlorides) which present a
105 darker coloration with the consequent effect of total blackening of vermilion. These chlorine-based
106 compound like calomel Hg₂Cl₂, terlinguaite Hg₂ClO, eglestonite Hg₆Cl₃O(OH), corderoite Hg₃S₂Cl₂
107 and kenh suite Hg₃S₂Cl₂, are often instable and they dissociate re-forming, in particular cases of
108 relative humidity (RH) and pH, amorphous mercury sulfide [2,5].
- 109 • *Photochemical redox* to obtain metallic Hg⁰ and S⁰, after catalysis by halogens at high RH under
110 sunlight/UV exposure. The dark coloration is due to the presence of grey metallic Hg clusters into
111 the red pigment [7,19].
- 112 • *Thermodynamic effect*: dissociation of α-HgS in metallic Hg⁰ and β-HgS, due to a transition
113 temperature in the range 373-673K, as discussed before. The transition temperature can decrease
114 drastically in presence of Cl ions, in the system Hg-S-Cl-H₂O with prevalence of Hg ions. As reported
115 by Radepont et al. [6], at 300K the effect of pH leads to a simultaneous presence of Hg⁰ and α-HgS
116 with mutual concentration depending on pH. In this condition the presence of β-HgS is not excluded.
117 However, a predominant concentration of Cl ions, depending on the pH, implies the formation of Cl-
118 based compounds like calomel or corderoite.

119 By considering the above-mentioned principal mechanisms of vermilion darkening, in this work we
120 examined in depth the problem by exploiting a comparative analysis of synthetic and ancient samples.
121 To explain the phenomenon, we propose a kinetic model of mutual composition of alpha/beta cinnabar
122 derived by the experimental darkening of pure alpha synthetic cinnabar intentionally doped with Cl⁻ ions
123 and treated under prolonged UV light exposure. The analysis was carried out by means of a multi-
124 technique approach including, among the others, Raman Spectroscopy. This technique gained, in the last
125 decades, a key role in the study of chemical-physical process related to the cultural heritage [20–27]. Our
126 approach is based on different fingerprints represented by Raman spectra of above-mentioned chlorine-
127 based compounds like calomel or terlinguaite [18,28–31]. Indeed, as representatively shown in figure 1A
128 (RRUFF database - ID R060492 and R070370), the chlorine compounds present intense bands at lower
129 wavenumbers with respect to HgS phases (below 200 cm⁻¹) [32]. In addition, the intensity ratios among
130 the bands can be used to determine from one side the presence of specific compounds such as calomel
131 or terlinguaite, from the other side to discriminate between β-HgS and α-HgS phases. **This study, finally,
132 is addressed to provide important elements for understanding the problem and formulate a preliminary
133 model able to predict the degradation of the pigment as a function of the time and environment conditions.**
134

135 2. Materials and methods

136 2.1 Materials

137 2.1.1 Synthetic samples

138 To study the darkening process of vermilion in painting and inks we realized synthetic samples doped
139 with Cl⁻ with the purpose to simulate the effect of this catalyst agent. Synthetic samples were prepared
140 following a previous work of Neiman et al., which studied the role of Cl⁻ ions in relation to darkening
141 phenomenon, and interpreted the results with the formation of transition chloride compounds [33].

142 Pure red cinnabar powders, provided by “Opificio Pietre Dure” (Florence – Italy), having 65 μm
143 grain **and volume density of 8.1 g/cm³** (Natural Cinnabar Monte Amiata 100% - CAS-Nr: 1344-48-5 –
144 Red/Medium - Kremer Pigmente K10610- Pigment Red 106, C.I. 77766), were dispersed in distilled
145 water and mixed with different molar concentration of NaCl in order to obtain a set of samples with
146 variable Cl⁻ concentration. **The volume ratio between pigment and solution was 1:25.** All mixtures were
147 hold under agitation for 1 h at room temperature.

148 Four different concentrations were selected with the intent to cover a wide range of Cl⁻ ions. We
149 report the assigned nomenclature of samples connected directly to the molar concentration: 0.00M NaCl
150 called “pure”; 0.01M NaCl called “0.01M”, 0.1M NaCl called “0.1M” and 5M NaCl called “5M”.
151 Solutions were then applied to a specific support to obtain a pigment deposit available for our analysis.
152 Depending on the analysis technique different supports were exploited as reported in the following:
153

- 154 • *Raman, SEM/EDS and Pump-probe analysis*: all solutions were dropped to glass slides and dried,
155 with the final results of a solid deposit of about 100 μm thickness and covering area of about 2 cm².
- 156 • *Reflectance measurements*: Pure and 5M solutions were continuously dropped and dried upon an
157 inert polyvinyl chloride (PVC) support, tool of our measurement apparatus, until we obtained a
158 compacted homogenous powder deposit (disk with r =16 mm, thickness = 1 mm).

159
160 All synthetic samples were treated under the UV light of a LED at 365 nm (emission with Lorentzian
161 profile having full width half maximum of 10 nm), under constant power density of 10 mW/cm², for time
162 ranges between 0 and 200h.

163

164 Thermal treatments were performed keeping the glass slides with the samples in a controlled oven at
165 70°C, 135°C and 200°C for different time ranging from 0 to 10 days.

166

167 2.1.2 Ancient manuscripts from “Biblioteca Universitaria di Cagliari”

168 In order to extend the model proposed in this work to natural aging effects, we studied six ancient
169 samples, belonging to the “Biblioteca Universitaria di Cagliari” BUCA, written in the range between
170 1435 and 1511 A.D. These samples were chosen because they were the oldest sample available and the
171 most suitable to test our model. In addition, they present cinnabar inks of high purity, according to the
172 preparative procedure of cinnabar in that age [3,4,9]. They represent important texts of the library used
173 in the academia as theology, administrative and sciences references. The paper manuscripts are conserved
174 at controlled atmosphere and were subjected to little restoration to preserve the cellulose support.
175 **Parchment manuscripts are conserved at controlled atmosphere and do not present restoration**
176 **interventions.** The complete list of the samples is reported in the following:

177

- 178 • *Ruralium commodorum* (1435, **parchment support**)
- 179 • *Furs de Valencia* (1482, **paper support**)
- 180 • *Libri sententiarum* (XV century, **parchment support**)
- 181 • *De Sphaera mundi* (1482, **paper support**)
- 182 • *Kalendarium* (1485, **paper support**)
- 183 • *Opus de intellectu et de causis mirabilium effectuum* (1511, **paper support** – see figure 1B)

184 All measurements on ancient samples were performed covering at least 5 points per “card” in which
185 vermilion was applied as writing ink and color in illuminations (**see example images reported in**
186 **Supplementary Materials**). **Sampling consisted in acquiring the spectra at three different pages, namely**
187 **at the first page, in the middle and at the end of the code, to achieve a total of 25 spectra per volume.**
188 **Then, a single average spectrum was extracted as representative of the volume.** Further details about
189 codicology and sampling procedure can be found in Supplementary Materials (Figure S1).

190

191 2.2 Methods

192 2.2.1 Raman Measurements

193 High resolution micro Raman scattering measurements were obtained in back scattering geometry
194 through the confocal system SOL Confotec MR750 equipped with Nikon Eclipse Ni microscope. Raman
195 spectra were gathered by using, as excitation wavelength, the 785 nm line (IO MatchBox series laser
196 diode). The system is equipped with four gratings (150, 600, 1200 and 1800 grooves/mm) which can be
197 selected depending on the measure requirements. The grating with 1800 grooves/mm was used to obtain
198 a resolution of 0.2 cm⁻¹.

199 In situ micro Raman scattering measurements were carried out in back scattering geometry with the 1064
200 nm line of an Nd:YAG laser. Measurements were performed in air at room temperature with a compact
201 spectrometer B&WTEK (Newark-USA) i-Raman Ex integrated system with a spectral resolution of 8
202 cm⁻¹. For each experimental setup, all the spectra were collected with an acquisition time of about 60 s
203 (five replicas) and power excitation between 5 and 10 mW concentrated in a spot of 0.3 mm² on the
204 surface through a Raman Video Micro-Sampling System (Nikon Eclipse for high-resolution and
205 BAC151B in the other case) equipped with a 20 × Olympus objective to select the area on the samples.

206 These conditions were selected after preliminary studies as safe conditions for the samples in order to
207 avoid the effect reported in [34]. Each measurement area represents a sampling surface of about 1 cm².

208

209 2.2.2 SEM/EDS measurements

210 SEM images were gathered by a scanning electron microscope ESEM:FEI Quanta 200 under low vacuum
211 conditions. EDS semiquantitative analyses were obtained with the help of Thermo Scientific EDS
212 UltraDry INTX-10P-A system equipped with Pathfinder. Each point of analysis was collected with an
213 acceleration voltage of 20 kV and live time of 30 s.

214

215 2.2.3 XRD measurements

216 XRD analysis was obtained by a diffractometer Rigaku Ultima IV. XRD pattern was collected using as
217 excitation the Cu-K α (40 KV, 40 mA), varying the angle θ in the range 10° - 70° with resolution of 0.1°
218 step/s. By XRD was measured the pure powder without any exposure and pure powder after UV
219 exposure. Powders were darkened superficially during the UV exposure, then shuffled and exposed again
220 to UV in order to maximize the ratio between beta and alpha phases and to reach the threshold of detection
221 limit typical for XRD technique (around 1%). The obtained pattern was analyzed by EVA database in
222 order to identify all the phases present in the starting material.

223

224 2.2.4 Pump-probe measurements

225 For transient absorption measurements a train of laser pulses obtained by a regenerative Ti:Sapphire
226 amplifier Coherent Libra-F-1K-HE-230 to produce 100-femtoseconds pulses at 800 nm with a kHz-
227 repetition rate. The train of laser pulses is divided into two parts by a beam splitter, called pump and
228 probe respectively.

229 The pump laser pulse is sent on an optical parametric amplifier (TOPAS-800-fs-UV-1) and finally
230 focused not perpendicularly on the sample, after a chopper synchronized with the frequency of the source.
231 The synchronized chopper (500 Hz) stops half of the incident pulses in alternate way so that half of the
232 obtained spectra will not be affected by the pump pulse (pump off), while the remaining will be affected
233 (pump on).

234 A white super-continuum pulse (probe) is formed by multi-frequencies generated by a sapphire plate
235 which also guarantees a sufficient stability and bandwidth flatness. The probe pulse, after passing a
236 controlled delay line, is focused and sent on the sample in the same intersection area of the pump pulse.
237 After the interaction with the sample, transmission signal is collected into the detection system
238 (UltrafastSystems HELIOS-80000-UV-VIS-NIR coupled with a CCD camera). For every single step of
239 the delay line, a single wavelength dispersed differential transmission spectrum is acquired, obtaining a
240 time spectrogram which can provide information about the temporal dynamics of the energy levels
241 depletion.

242 Transient absorption was obtained by using a 400 nm pump signal (deep blue, compatible with two
243 photons excitation from Yu et al.) and studying the interaction with the white super-continuum probe
244 signal. The pump-probe curves were obtained varying the pump power in a range between 0.200-0.600
245 mW and collecting the signal in the “short live” range (10 ps – resolution 0.02 ps) and “long live” range
246 (10 ns – resolution 0.1 ns).

247

248 2.2.5 Reflectance measurements and colorimetric parameters

249 Reflectance measurements were performed by means of UV–Vis–NIR Agilent Technologies Cary 5000
250 spectrophotometer equipped with integrating sphere module. The reflection configuration at 10°
251 measures the diffuse reflection of the sample with respect to a reference sample which is considered to
252 have a 100% reflectivity. A calibrated source Illuminant D65 was used to determine the reflectance
253 spectra and for calculating the colorimetric parameters.

254 3. Results and discussion

255 3.1 Structural analysis

257 The XRD analysis of “pure” sample (Figure S2) corresponds entirely to the phase α -HgS, down to the
258 detection threshold of 0.8-1 wt. % typical for this technique. As shown in Figure 1A, the presence of
259 specific Cl-related compounds can be easily detected, being characterized by Raman fingerprints well
260 separated from the ones of cinnabar (spectra of Cl-based compounds and metacinnabar were retrieved
261 from RRUFF database, the one of cinnabar was measured on the relic reported in Figure 1B). **As reported
262 by Frost et al.[35], cinnabar pertains to D_{3h} point group, with $2A_1'$ Raman active modes, and $3A_2'' + 5E'$
263 modes, which are both infrared and Raman active. The A_1 Raman active modes are reported at 43 and
264 256 cm^{-1} and the E' modes at 72, 88, 108, 283 and 343 cm^{-1} . Two unassigned bands were observed at
265 290 and 351 cm^{-1} . According to Frost Raman spectra of HgS and other sulphide pigments may be
266 conveniently divided into three sections namely: (a) the region centred upon 350 cm^{-1} where the
267 stretching vibrations are observed (b) the region centred upon 250 cm^{-1} ascribed to the bending vibrations
268 and (c) the region below 100 cm^{-1} assigned to the lattice modes. Among them, the bending vibration
269 region appears as the most sensitive to detect cinnabar phase variation.**

270 It is clear that to discuss vibrational features of α - and β -HgS one should consider in particular the 250-
271 280 cm^{-1} range, so that we focused our attention to this specific Regions Of Interest (ROI). Figure 1C,
272 showing the Raman spectrum of as-prepared pure sample, fully confirms the XRD results showing the
273 vibrational features of pure α -HgS phase (peaks at 253, 283 and 350 cm^{-1} , the spectrum was recorded
274 with the high-resolution system)[10]. For comparison, the figure also reports the spectrum of
275 metacinnabar, where the three main vibrations are blue-shifted of about 5-7 cm^{-1} . Beta-HgS spectrum,
276 assumed as a reference, was retrieved from RRUFF database (ID R060950) [32], the powders being from
277 natural crystals. Beside the blue-shift, the ratio between the areas of the two bands at about 250 and 280
278 cm^{-1} (A_{250}/A_{280}) undergoes a marked variation, decreasing by a factor 2 in the metacinnabar case. It is
279 worth noting that the Raman spectroscopy provides a detection threshold far exceeding the XRD one,
280 easily reaching values lower than 0.1 wt. % and down to ppm in few cases [36–38]. Based on this starting
281 achievement, we performed UV exposure on a set of samples, both pure and intentionally doped with Cl
282 at different concentration. The darkening effect is sizable, as shown by the image of the samples reported
283 in inset of figure 2A where a comparison between the *pre* and *post*-treatment is shown for the pure sample
284 case. In order to quantify the effective darkening of red color, figures 2A and 2B delineate the reflectance
285 spectra and the first derivative reflectance spectra of pure and 5M samples submitted to UV exposure.
286 The large variation in the reflectance spectra upon sample irradiation is evidenced also by first derivative
287 spectra, whose change is significative and better expresses the UV exposure effect. According to Gueli
288 et al., we calculated in table I the CIE L^*, a^*, b^* coordinates and the total color variation ΔE_{Lab} extracted
289 from reflectance spectra [39,40]. We also calculated the Chroma C^* coordinates, with hue angle h , which
290 also mark the color variation in cylindric coordinates. Chromatic variation can be observed in a color
291 reconstruction reported in figures 2C-D-E.

292 With the help of high-resolution Raman spectroscopy, we studied the spectra obtained in order to
293 find a first interpretation of the phenomenon. Figure 3 proposes the experimental spectra, obtained with
294 the high-resolution system, and the related deconvolution curves (with Lorentzian functions) to estimate

295 the phase change from cinnabar to metacinnabar. In particular, figure 3A and 3B show the spectra
296 deconvolution operated on red and partially darkened pure sample respectively. A comparison with the
297 portable system with excitation at 1064 nm and low resolution is also proposed (figure 3C and 3D).
298 Spectral resolution plays a key role for determining the above-mentioned ratio A_{250}/A_{280} correlated to the
299 transition from cinnabar to metacinnabar and a detailed section concerning the error estimation deriving
300 from the low- and high- resolution experimental systems is proposed below (*vide infra*, section 3.5). In
301 particular, the high-resolution system is able to better separate the contribution of the two bands and to
302 distinguish the presence of two extra vibrational features around 280 and 350 cm^{-1} (the fitting procedure
303 being performed with 5 and 3 Lorentzian bands in the high and low-resolution systems). However, it is
304 important to pinpoint that within the estimated uncertainty, both the systems are able to detect the
305 variation induced by light exposure (see Table SI). This is of outmost importance since the portable low-
306 resolution system is a setup that can be successfully exploited to analyze ancient samples which cannot
307 be moved out of the conservation site, as we verified in the present work on the ancient manuscript.

308 Extending our analysis to the other samples, we propose in figure Figures 4A and S3 a comparison
309 between the reference spectra of cinnabar and metacinnabar with those obtained for pure and 5M sample
310 before and after UV treatment. In the samples with low Cl concentration we do not observe any band
311 typically ascribed to chlorine-based compounds (wavenumber region below 200 cm^{-1}), and the curves
312 show a broadening of the band at 250 cm^{-1} (more accentuated in 5M sample) accompanied with a shift
313 toward low wavenumbers. Furthermore, the intensity ratio between the band at 253 cm^{-1} and the shoulder
314 at 283 cm^{-1} is completely different in samples submitted to UV exposure as compared to the pure one.
315 This trend is compatible with the transformation from cinnabar to metacinnabar phase and is reported in
316 the whole set of Cl doped samples. In addition, in the 5M sample a very weak broad band below 200 cm^{-1}
317 is detected, suggesting the formation, at this Cl concentration level, of amorphous chlorine-based
318 compound. Cl concentration appears as a crucial parameter: indeed, the Cl compound related vibrational
319 band below 200 cm^{-1} is not observed in low Cl concentration samples, confirming the role of Cl dopant
320 in elevated concentration. When we consider the darkening effect caused by UV exposure, we find that
321 Cl impurities carry out a sort of catalytic action promoting a faster darkening of the UV exposed sample:
322 whilst a complete degradation of the pigment is obtained after 148h of light exposure in the pure sample,
323 darkening is achieved in only 20 h of exposure for the 5M Cl doped sample. Further UV irradiation does
324 not affect the sample, no additional variations in Raman spectra, reflectance measurements and chromatic
325 coordinates being recorded.

326

327 3.2 Cinnabar transition and metallic Hg

328 Raman spectroscopy was successfully applied to discriminate the contribution of Cl impurities to
329 the darkening phenomenon under UV exposure, being able to display the cinnabar to metacinnabar
330 transition somehow accelerated by the presence of Cl ions. However, beside the chlorine-based
331 compounds observed in the high Cl concentration case, we could not exclude the presence of other
332 species which could also have a catalytic effect and could be not easily detected by Raman spectroscopy.
333 This is the case of metallic Hg, as already reported in the introduction [10]. To deepen this topic, we
334 performed SEM measurements and elemental EDS analysis on the 5M UV darkened sample (Figure 4B).
335 The sample image gathered by back scattered electrons is reported on a grey scale correlated to the atomic
336 weight of the elements. Table II summarizes the elemental results for the collected points indicated in
337 figure 4B. By excluding the glass slide contribution, stoichiometric ratios among the elements imply the
338 presence of HgS and NaCl starting compounds even after UV exposure. Other phases based on Cl
339 species, like calomel or terlinguaite, are excluded for stoichiometric reasons. In addition, this analysis,
340 which presents an excess of Hg not saturated with other elements, calls for the expected formation of
341 metallic Hg in the darkened area. Table II reports also which points are related to “UV” and “No UV”
342 area. In relation to these areas the ratio between atomic percentages of Hg and S (es. between points 3 or

343 4 and 2 or 5 or 6 in 5M sample) evidences a larger amount of Hg as compared to S especially in dark
344 points, suggesting the presence of metallic Hg. Although EDS analysis provide a specific elemental
345 composition of the samples, it is not able to discriminate the exact phase of possible formed compounds,
346 thus showing the need of a multi-technique approach to tackle the darkening issue. However, the
347 comparison of the compounds determined for pure and 5M samples (Table II), highlights once more the
348 role of Cl impurities in promoting HgS phase transition down to the dissociation of HgS into metallic
349 Hg.

350 The transition from cinnabar to metacinnabar phase was recently validated by ultra-fast transient
351 absorption measurements [10]. We performed spectrally resolved experiment on “pure” and “UV
352 darkened pure” samples to discriminate the presence of each phase by their different excited state
353 kinetics. Thanks to the spectral dispersion of our experiment, we observed a transient positive signal in
354 the range between 750-800 nm, and a negative signal in a broad band around 450-550 nm (Figure 5A).
355 According to Yu et al., positive signals of two-photons absorption (TPA) and excited-state absorption
356 (ESA) are associated to pure red α -HgS phase and to an intermediate state (chemical or structural phase)
357 occurring during the phase transition from α to β -HgS. Due to the lower frequency sampling of our
358 experiment we could not isolate the contribution of pure cinnabar, but we detected its presence coupled
359 to the intermediate state once a threshold power of about 0.450 mW was achieved, confirming previous
360 results. The negative ground state depletion (GSD) signals, despite a larger time scale up to ns, seem to
361 be compatible with the interpretation of Yu et al., indicating a short-lived GSD associated to β -Hgs phase
362 and a long-lived GSD ascribed to metallic Hg. Without further entering the details of the transient
363 absorption kinetics, which will be the subject of a next work, the figure shows all the elements of the
364 darkening transition, starting from TPA+ESA (α -HgS + transition phase), ESA (transition phase), short
365 GSD (β -HgS) and finally long GSD (metallic Hg).

366 The UV exposure proceeds from the surface of the cinnabar powders because of the large UV
367 absorption and the very small penetration depth (the absorption gap of HgS is 2.1 eV - [41]). To finally
368 demonstrate the transition from α -HgS to β -HgS we carried out a prolonged UV exposure experiment in
369 order to achieve a darkened sample where the amount of β -HgS reached the sensitivity level of XRD
370 measurements. The conclusive demonstration is given by the XRD pattern shown in figure 5B where the
371 main diffraction peak of α -HgS around 31 degrees shows the appearance of a shoulder towards lower
372 angles because of the formation of β -HgS [10]. These findings are in good agreement with the results
373 gathered from pure alpha and beta cinnabar (the latter produced by thermal treatment, as reported in the
374 introduction) and the ones obtained analyzing the effect of UV exposure [2,10].

375 To sum up the whole set of experimental results, we were able to highlight the transition from α -
376 HgS to β -HgS under UV exposure of samples eventually doped with Cl ions by means of a multi-
377 technique approach. Among the others, thanks to its high sensitivity and non-destructive character,
378 Raman spectroscopy is, in our opinion, the technique of election to examine in detail the kinetics of the
379 darkening process allowing to discriminate the two phases as a function of the composition of the samples
380 or the UV exposure undergone by them. Thus, a kinetic model based on the vibrational features is
381 proposed to evaluate the darkening phenomenon in laboratory samples and some artbooks to assess
382 conservation status and aging of the relics.

383 384 3.3 The model

385 As reported in the previous sections, the Raman spectra analysis and the darkening effect as a function
386 of UV exposure and composition can be rationalized considering the phase transformation from cinnabar
387 to metacinnabar as the main cause of the darkening process. The gathered results indicate that the effect

388 is promoted by the presence of chlorine ions, but high concentration of this element produces the
 389 formation of other Cl related compounds (like corderoite, terlinguaite etc.). This interpretation is
 390 confirmed by recent studies proposing, in addition, the formation of metallic Hg as a possible darkening
 391 agent [2,9,10]. The role of chlorine is commonly accepted as catalyst, in presence of light, toward the
 392 formation of transition compounds. Several studies show how a phase transformation takes place with
 393 the help of coalescence phenomena where clusters of the new phase act as further nucleation center. This
 394 condition implies an exponential speed of transformation until the complete saturation, where chlorine
 395 impurities, or metacinnabar clusters engage the reaction as catalysts. Thus, we can assume the model of
 396 auto-catalyst reaction where the reaction products act as catalyst for a new step of the reaction and we
 397 apply The Verhulst model, based on logistic function [42–44]. By assuming P as the measure of a species
 398 population and t the time, the growth logistic model is defined by the differential eq. 1:

$$399 \quad \frac{dP}{dt} = kP \left(1 - \frac{P}{K} \right) \quad (1)$$

400 where K is the asymptotic term at $t \rightarrow \infty$, the ratio P/K represents the slowdown factor of the curve and
 401 k refers to the growth rate. Equation 1 can be solved with the logistic function:

$$402 \quad P(t) = \frac{K}{1 + qe^{-kt}} \quad (2)$$

403 with

$$404 \quad q = \frac{K - P(0)}{P(0)} \quad (3)$$

405 being the term $P(0)$ the population at $t=0$.

406 In our case, the asymptotic term K is 1, since we have the complete transformation of the initial population
 407 (α -cinnabar) into the final one (β -cinnabar). Thus, the model used in this study can be simplified
 408 accordingly

$$409 \quad P(t) = \frac{1}{1 + qe^{-kt}} = \frac{1}{1 + qe^{-\frac{t}{\tau}}} \quad (4)$$

410 To apply the Verhulst model to the Raman spectra collected on synthetic samples, we need to find an
 411 observable representing cinnabar population at time t . We can focus the attention on the term $R = A_{252}/A_{283}$
 412 representing the ratio between the areas of the bands at 252 cm^{-1} and 283 cm^{-1} . As previously observed,
 413 this ratio changes when a phase transformation from cinnabar to metacinnabar is taking place. If we
 414 consider as $R(0) \propto [\alpha\text{-HgS}]$ the ratio at the time $t=0$ when only the cinnabar phase is present and $R(t) \propto$
 415 $[\alpha\text{-HgS}] + [\beta\text{-HgS}]$ the ratio due to contemporary presence of cinnabar and metacinnabar, the normalized
 416 ratio R_n expresses the fraction of $[\beta\text{-HgS}]$ with respect to the total composition according to the following
 417 relation:

$$418 \quad R_n = \frac{R(t) - R(0)}{R(t)} = \frac{[\beta\text{-HgS}]}{[\alpha\text{-HgS}] + [\beta\text{-HgS}]} \quad (5)$$

419 By equalizing the eq. 5 with eq. 4, the population $P(t)$ of $[\beta\text{-HgS}]$ can be expressed by the following
 420 relation:

$$421 \quad P(t) = R_n = \frac{R(t) - R(0)}{R(t)} = \frac{[\beta\text{-HgS}]}{[\alpha\text{-HgS}] + [\beta\text{-HgS}]} = \frac{1}{1 + qe^{-\frac{t}{\tau}}} \quad (6)$$

422 where, according to the logistic model, the following conditions hold:

$$423 \quad \lim_{t \rightarrow 0} R_n = 0 = [\alpha - \text{HgS}]$$

$$424 \quad \lim_{t \rightarrow \infty} R_n = 1 = [\beta - \text{HgS}]$$

425 The term τ of the equation is defined as the characteristic time of the reaction during the darkening
426 process. According to the literature, it depends on a large set of factors governing the pigment degradation:
427 relative humidity (RH), chlorine concentration ([Cl]), temperature (considered in terms of activation
428 energy E_{kT}), metacinnabar concentration ([β -HgS]), light exposure (both in terms of light energy E_{hv} and
429 exposure time t_{exp}). To complete the darkening model it is therefore mandatory to study the characteristic
430 time $\tau = \tau(RH, E_{kT}, E_{hv}, [Cl], [\beta - HgS], t_{exp})$ to understand the role of the different factors.

431

432 3.4 Logistic model applied to synthetic samples

433 By operating a deconvolution with Lorentzian functions on the Raman spectra, we calculated the ratio
434 $R(t)$ and $R(0)$ as a function of UV exposure time for each synthetic sample. A complete extraction
435 procedure with deconvolution functions is reported into the supplementary materials. Based on the results
436 reported in section 3.1, we pursued the analysis on the synthesized samples with the low-resolution
437 system to validate the approach for the investigation of ancient relics (considered in section 3.6). Figure
438 6 reports the experimental points obtained for any sample and the logistic fit of eq. 6. The figure
439 summarizes also the fitting parameters K , q and τ (we leave the parameter K free to verify that its value
440 is equal to 1, as previously discussed). Error bars are obtained by using the spectrometer sensitivity and
441 standard deviation associated to each determined area.

442 As perceived from the data, the characteristic time τ decreases drastically as a function of Cl
443 concentration. This insight confirms again the key role played by Chlorine during the process and the
444 observed catalytic effect of the ion. It is also worth noting that K value is equal to 1 in the whole set of
445 samples, as expected. However, albeit the value of the parameter q is quite homogeneous for the first
446 three samples, the 5M one shows a large decrease of the q value. Since q represents the percentage of β -
447 HgS at time $t=0$, we get from eq. 3 that the first three samples have the same starting concentration of
448 metacinnabar of about 1.5-2%. As for the sample 5M we have already noted that the Raman spectrum
449 displays an additional band at about 270 cm^{-1} due to the presence of chlorine-based compounds formed
450 during the reaction, as mentioned before. Thus, a deconvolution with only two Lorentzian bands at 283
451 cm^{-1} and 255 cm^{-1} did not lead to an acceptable fit and a third Lorentzian band must be considered to
452 successfully fit the data. The most suitable compound forming an additional band in the region of interest
453 for our deconvolution is calomel, which is characterized by a vibrational mode at 273 cm^{-1} [28].
454 Therefore, the ratio R_n is corrected in order to consider the contribution of this additional band:

455

$$R_n = \frac{R(t) - R(0)}{R(t)_{TOT}}$$

456 with

457

$$R(t)_{TOT} = \frac{A_{255}}{A_{283} + A_{273}}$$

458 Assuming this correction, one can validate the model considering the two extreme cases, $t \rightarrow 0$ where
459 no contribution of calomel is observed and the ratio $R_{TOT}=R$, whilst for $t \rightarrow \infty$ the products of the reaction
460 are composed by a mixture of calomel and metacinnabar. This assumption justifies a very low value of
461 q parameter (6.7) for 5M sample which corresponds to a high concentration of reactive products even at
462 $t=0$. Indeed, in figure 4E the curve presents an initial $R(0)$ parameter of 0.15, corresponding to 15% of
463 possible mixture *calomel* + *metacinnabar*. A reasonable justification of this value originates from the
464 high concentration of chlorine ions which could cause an initial transformation of cinnabar even without
465 light exposure. Figure 6E summarizes the curves obtained for each sample as a function of the UV
466 exposure time and Cl concentration. In addition, we studied the variation of the rate k expressed in eq. 1
467 as a function of Cl concentration (figure 6F). A sigmoidal fit, derived solving the logistic function with

468 k , suggests that 5M concentration is a saturation value and higher content of chlorine does not produce
469 drastic effects on the speed of darkening process.

470

471 3.5 Error analysis

472 An estimation of errors associated to this model can be determined considering a linear propagation of
473 relative uncertainties of each parameter present in equation (5). As evidenced in our model, the ratio R_n
474 derives from the ratios $R(t)$ and R_0 which, analogously, depend on the ratio A_{250}/A_{280} . Consequently, due
475 to the fitting procedure operated for determining the areas A_{250} and A_{280} , we estimated a variation of the
476 relative error $\Delta R_n/R_n$ in relation to experimental resolution as follows:

- 477 • $0.2 < \frac{\Delta R_n}{R_n} < 0.4$ with resolution 8 cm^{-1} ;
- 478 • $0.04 < \frac{\Delta R_n}{R_n} < 0.06$ with resolution 0.2 cm^{-1}

479 Lowest values are determined when $R_n \rightarrow 0$, while highest values when $R_n \rightarrow 1$. In figure 7A we
480 reported a comparison between the two experimental resolutions and a comparison of the relative error
481 as a function of the selected grating of the system (figure 7B). This analysis confirms once again, and
482 through the whole range of UV exposure examined, that the results obtained with the low-resolution
483 system are in very good agreement with the ones gathered with the high-resolution system, further
484 supporting the exploitation of the former in the analysis of precious relics.

485

486 3.6 Logistic model applied to ancient samples

487 The logistic model (eq. 3) was empirically applied to the set of ancient samples belonging to BUCA with
488 the intent to study if the analysis developed for synthetic samples can be exploited for natural aging.
489 Operating in this way, we did not exclude that the spectra of ancient samples can be affected by a number
490 of factors, such as impurities, natural variation, spatial variation, unknown environmental conditions
491 since the creation, beside light irradiation. However, we know the date of the ancient samples, and we
492 could test the model on these samples by assuming standard conservation conditions, such as
493 temperature, RH or light exposure, as previously done in a work dealing with paper relics [26]. Indeed,
494 the overlay procedure of results on synthetic samples and historical artworks is absolutely empirical. It
495 was a tentative approach to verify if the model studied on synthetic samples could be applied to study
496 ancient samples. We exploit the synthetic cinnabar as time zero reference for the aging curve reported in
497 figure S4, where the concentration of metacinnabar is plotted versus the age of the relics produced in the
498 1435-1511 A.D. range. **In figure S4 we also reported the experimental average spectra gathered on the
499 artworks of BUCA, to show that very small differences were recorded in those samples (as also shown
500 in Figure 8B).** The whole set of samples but the third one (*Libri Sententiarum* - indicated as red in the
501 plot), agree very well to the fit model, despite the short time range explored. As for the *Libri Sententiarum*
502 sample, it should be noted that the writing date of this copy is affected by a large uncertainty. As
503 evidenced by the curve the parameter τ associated to the characteristic time of degradation is estimated
504 in 100 years. The accuracy of the fit covers all the experimental points, or their error bars, within the
505 range of 8%. It is worth nothing that, as discussed before, the uncertainty associated to fitting curve takes
506 into account the effects of impurities, natural variation, spatial variation, unknown environmental
507 conditions since the creation (other colorants were not considered because they were not revealed by
508 Raman spectroscopy). The spread associated to experimental points in figure S4 evidences this aspect.
509 However, despite that, there is a good agreement with the fitting curve and the experimental data gathered
510 on ancient samples.

511 To extend the time range explored, we applied the model directly to other Raman spectra extracted from
512 the literature, by collecting spectra of samples belonging to XIII-XVII centuries (Figure 8B). Since we

513 had not enough information about the effective date of production from literature, but only a generic
 514 indication of the centuries, we hypothesized an error bar of 50 years, thus reporting in the figure three
 515 point for each literature sample, to account for beginning, middle and end of the correspondent century.
 516 We can observe that the model reproduces the experimental data with an 8% accuracy (as the one
 517 obtained for the BUCA samples) when considering the first half of the century as the manufacturing date
 518 of the relics. This is in good agreement with other spectra reported in the literature [25,45–48].
 519

520 3.7 Correspondence between the model applied to natural and UV-caused darkening

521 With the purpose of a complete application of the model, we can hypothesize a calibration of τ and q
 522 parameters. To achieve a correspondence between synthetic and experimentally UV-caused darkening
 523 of the cinnabar, we calculated the equivalence ratio of the two parameters as follows:

$$524 \tau_{eq} = \frac{\tau_{ancient}}{\tau_{pure}}$$

$$525 q_{eq} = \frac{q_{ancient}}{q_{pure}}$$

526 where with the subscript *ancient* we refer to the fit parameters recovered in the artbook analysis. The
 527 conversion will be obtained, in the case of sample 5M for example, by using the recalculated parameters
 528 with the formula:

$$529 \tau_{calibrated_{5M}} = \tau_{5M} \cdot \tau_{eq}$$

$$530 q_{calibrated_{5M}} = q_{5M} \cdot q_{eq}$$

531 The result is the following equation:

$$532 P(t) = R_n = \frac{1}{1 + q_{calibrated_{5M}} e^{-\frac{t}{\tau_{calibrated_{5M}}}}}$$

533 The figure 8A reports all the curves calibrated for each sample in which the double scale indicates the
 534 conversion between natural darkening (years) and UV-accelerated process (h). In the reported graph the
 535 correlation between the two process is evidenced, especially considering the role of the Cl impurities.
 536

537 3.8 Thermal treatments

538 Finally, we want to consider the role of temperature in the darkening phenomenon. With the
 539 purpose of simulating the effect of the time as a thermodynamic process, we treated the Cl-doped samples
 540 at high temperature in a home-made aging chamber. We first selected 75°C with 60% of Relative
 541 Humidity (normally used as aging standard in common processes according to ISO 5630-3) and treated
 542 the sample in a period ranging from 0 to 10 days [49]. We determined the R_n ratio (that is the
 543 metacinnabar concentration) by collecting Raman spectra at fixed time interval (12/24 h). Under these
 544 experimental conditions no evident trend of R_n value was recovered. Selecting the temperature at 200°C
 545 to drastically enhance the effect of the aging, we could not find clear indications because of a strong loss
 546 of samples weight. The mass loss was estimated of about 1% per hour for each sample. Indeed, we
 547 correlated the appearance of a volatile element to the possible formation of metallic Hg, in agreement
 548 with SEM and transient absorption measurements previously reported. By performing specific EDS
 549 analysis for these samples (see table SII in Supplementary Materials) we can evidence the loss of Hg in
 550 relation to stoichiometric ratio with S. In fact, an excess of S in the compositional analysis suggests the
 551 partially dissociation of HgS, formation of metallic Hg and its consequent evaporation [14].

552 We then selected an intermediate temperature of 135°C and the R_n trend evidenced in pure, 0.1M
 553 and 5M samples is reported in Figure S5. Despite the mass loss recorded even at this temperature (see

554 table III), we were able to measure the vibrational features over the whole aging period (10 days). As
555 shown in the figure, whilst pure and 0.1M samples present similar linear trend with very slow increasing
556 of R_n ratio, in the 5M sample the slope is larger, confirming the Cl concentration role already discussed.
557

558 **4 Conclusions**

559 This work proposes a kinetic model of red vermilion degradation by correlating the role of Cl impurities
560 in the phase changing of cinnabar to metacinnabar. The study was addressed to a set of synthetic samples
561 brought forth by the so-called *wet* method of vermilion production and doped with different chlorine
562 concentrations. For having a complete understanding of the phenomenon, we also concentrated our
563 attention to ancient manuscript samples belonging to XIII-XVII century. Our analysis conducted
564 principally through Raman spectroscopy, but also with the help of XRD and SEM/EDS, reveals a
565 transformation from the alpha-cinnabar phase to the dark beta-cinnabar with the contemporary presence
566 of metallic Hg. The latter is also confirmed by pump-probe measurements which evidenced, in transient
567 absorption measurements, a long-life ground state depletion recently assigned to metallic Hg. The
568 complete panorama of experimental results, in particular the Raman analysis, can be explained with a
569 logistic trend of the darkening effect. In fact, following the model of auto-catalyst reaction and the
570 Verhulst curve, we found the characteristic time τ for natural degradation and UV-accelerated process.
571 We estimated natural τ of 100 years for pure cinnabar and we correlated it to the UV-accelerated τ ,
572 estimated for different Cl impurities of the samples. The model addresses especially to the circumstances
573 where light exposition and high environment salinity can activate the darkening process (frescos or mural
574 painting exposed to natural agents). However, it is adaptable to all artifacts in which pure HgS or a low
575 doping of Cl is present. Finally, this work evidenced, once again, the role of no-destructive portable
576 techniques, like Raman spectroscopy able to provide important information on materials investigation
577 applied to cultural heritage.
578

579 **Acknowledgments:** All the authors acknowledge the fruitful collaboration with the staff of the Biblioteca
580 Universitaria di Cagliari (BUCA), Sardinia. Their kind collaboration and expertise helped us during all
581 the measurement sessions. The experimental measurements were performed under a convention between
582 BUCA and the Physics Department of the University of Cagliari. The authors also thank S. Grillo for
583 valuable assistance and discussions in the XRD analyses. We acknowledge the CeSAR (Centro Servizi
584 Ricerca d'Ateneo) core facility of the University of Cagliari and Dr. M. Marceddu for assistance with the
585 generation of transient absorption data.
586

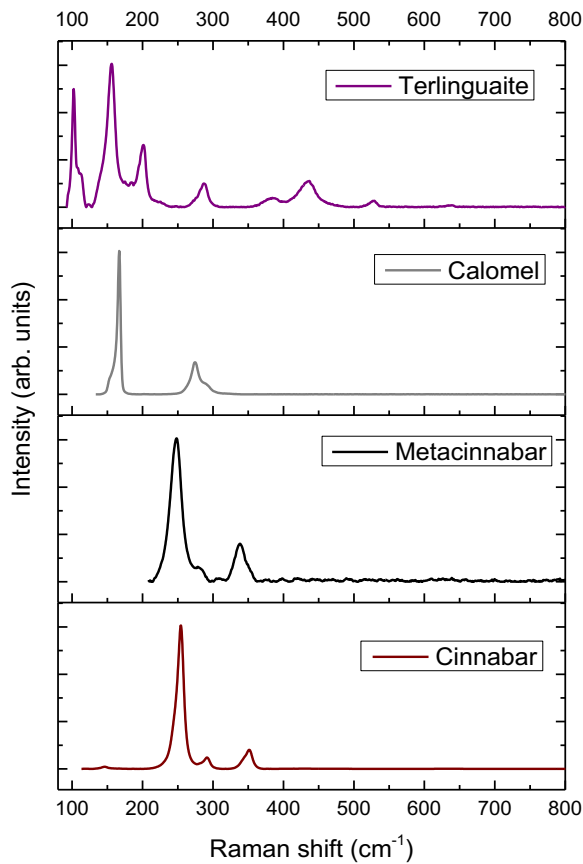
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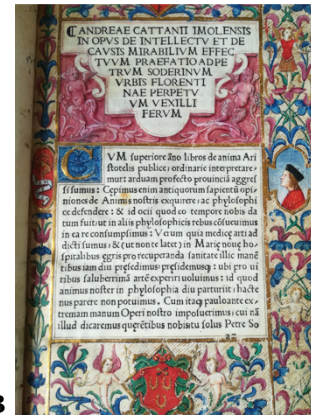
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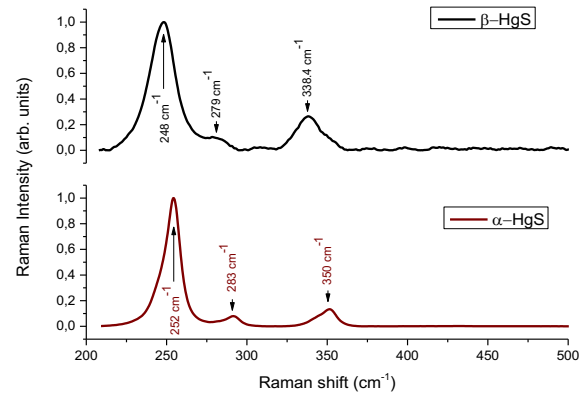
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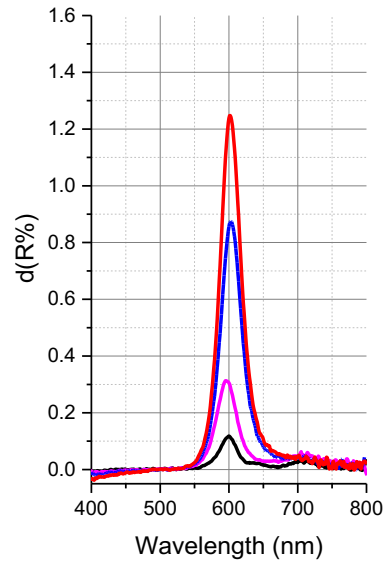
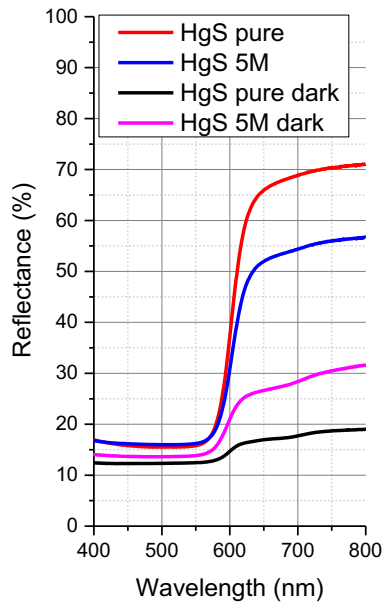


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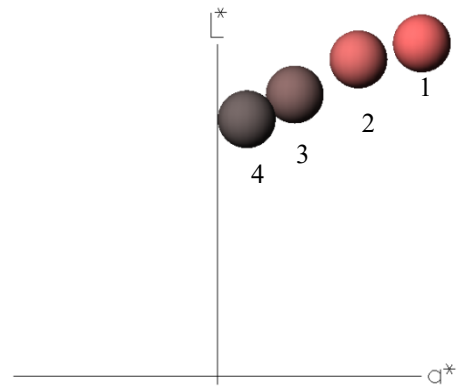
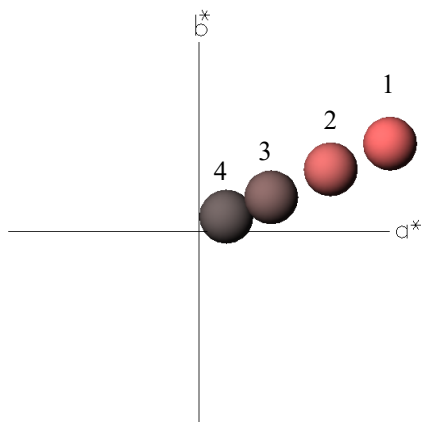
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FIGURE 1: Reference Raman spectra used for comparative analyses. Raman spectra of cinnabar phases and chlorine-based compounds related to vermilion darekening (A). Picture of representative ancient sample from Biblioteca Universitaria di Cagliari (BUCA): *Opus de intellectu et de causis mirabilium effectuum* (before 1511) (B). Comparison between β -HgS and α -HgS Raman spectra (C).



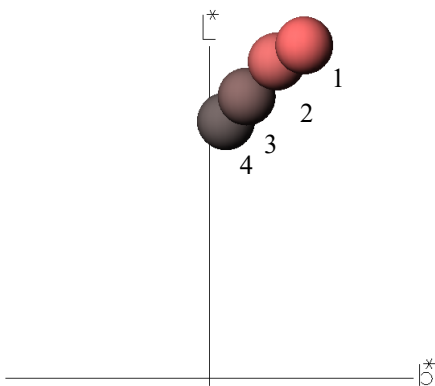
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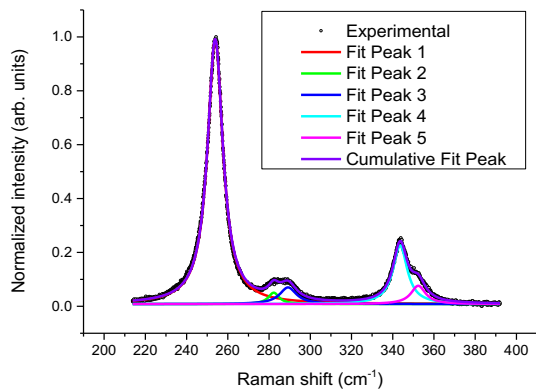
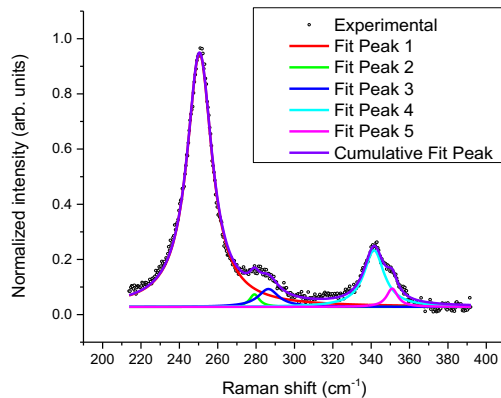
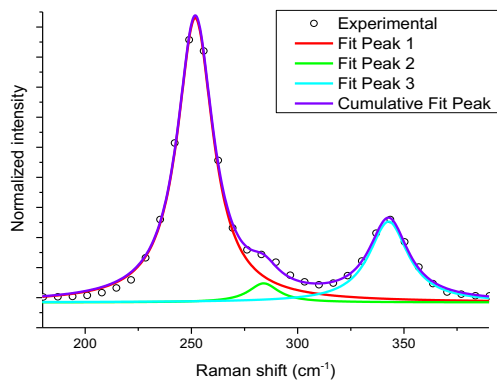
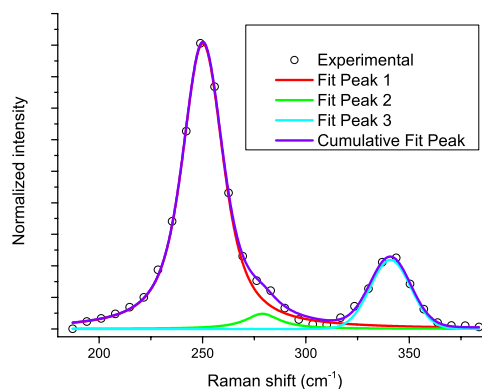
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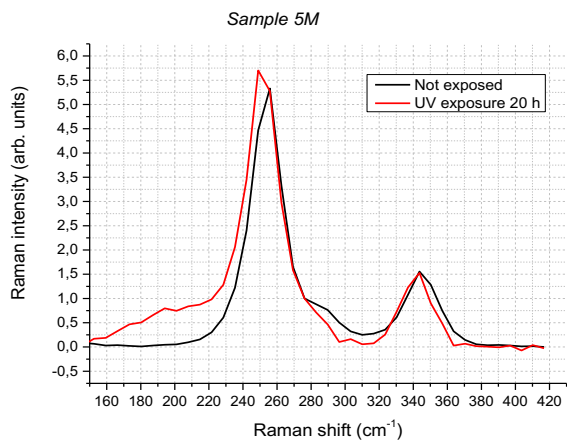
1	PURE (No UV)
2	5M (No UV)
3	5M (20 h UV exposed)
4	PURE (148 h UV exposed)

E

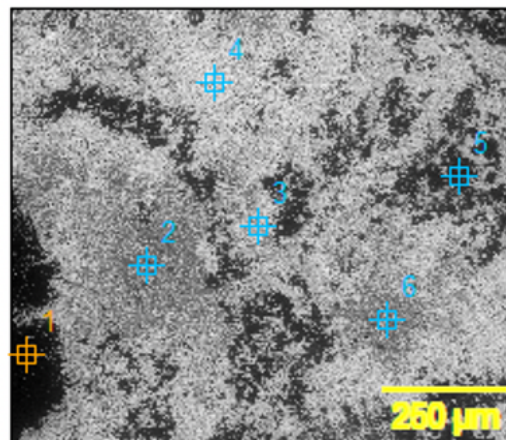
725 **FIGURE 2: Reflectance spectra and colorimetric parameters.** Reflectance spectra of sample “pure”
 726 and 5M before and after UV exposure ($\lambda=365$ nm; $10\text{mW}/\text{cm}^2$) (A); The inset show the darekend surface
 727 of “pure” sample after UV treatment. First derivative of Reflectance spectra (B); Variation of $L^*a^*b^*$
 728 coordinates calculated from reflectance spectra (C-E)

A**B****C****D**

729 **FIGURE 3: Deconvolution of Raman Spectra of “pure” sample before and after UV exposure.**
 730 Deconvolution on high resolution spectra: before (A) and after 90 h UV exposure (B); Deconvolution
 731 on low resolution spectra: before (C) and after UV exposure (D).



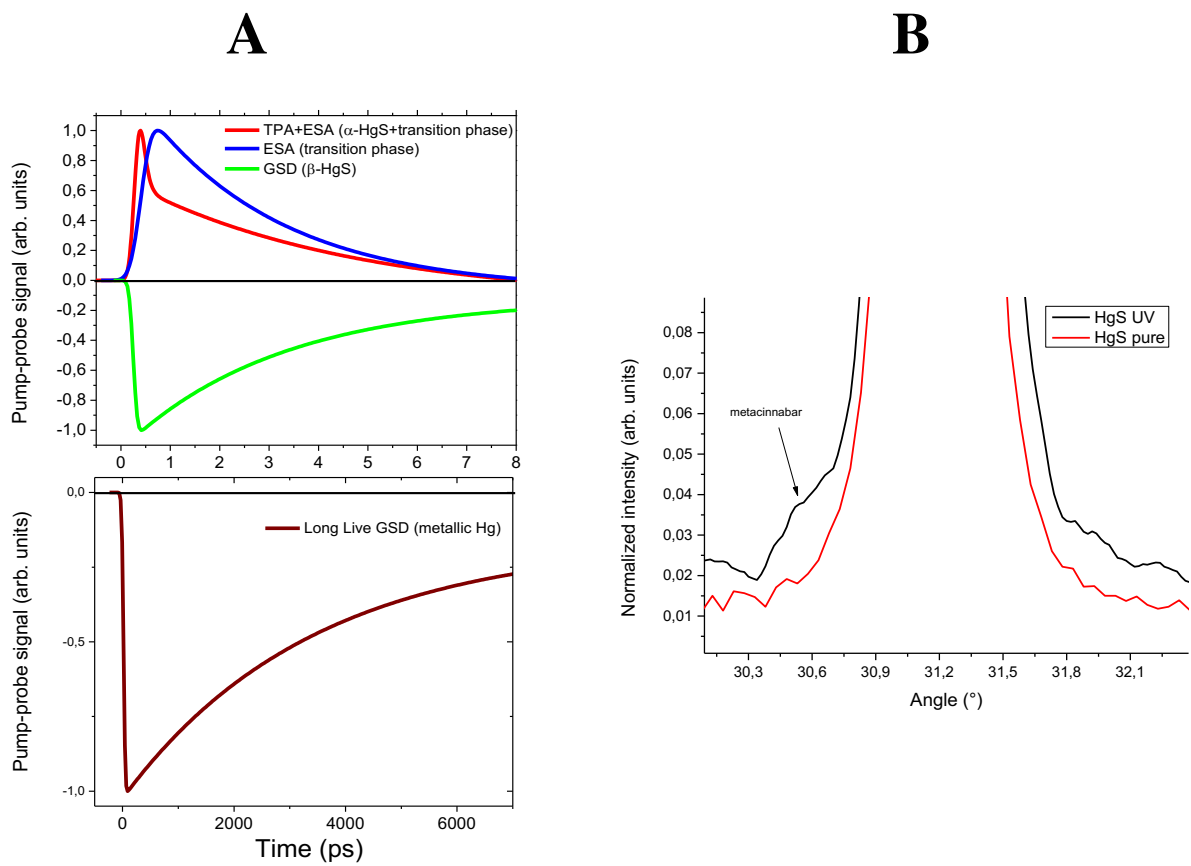
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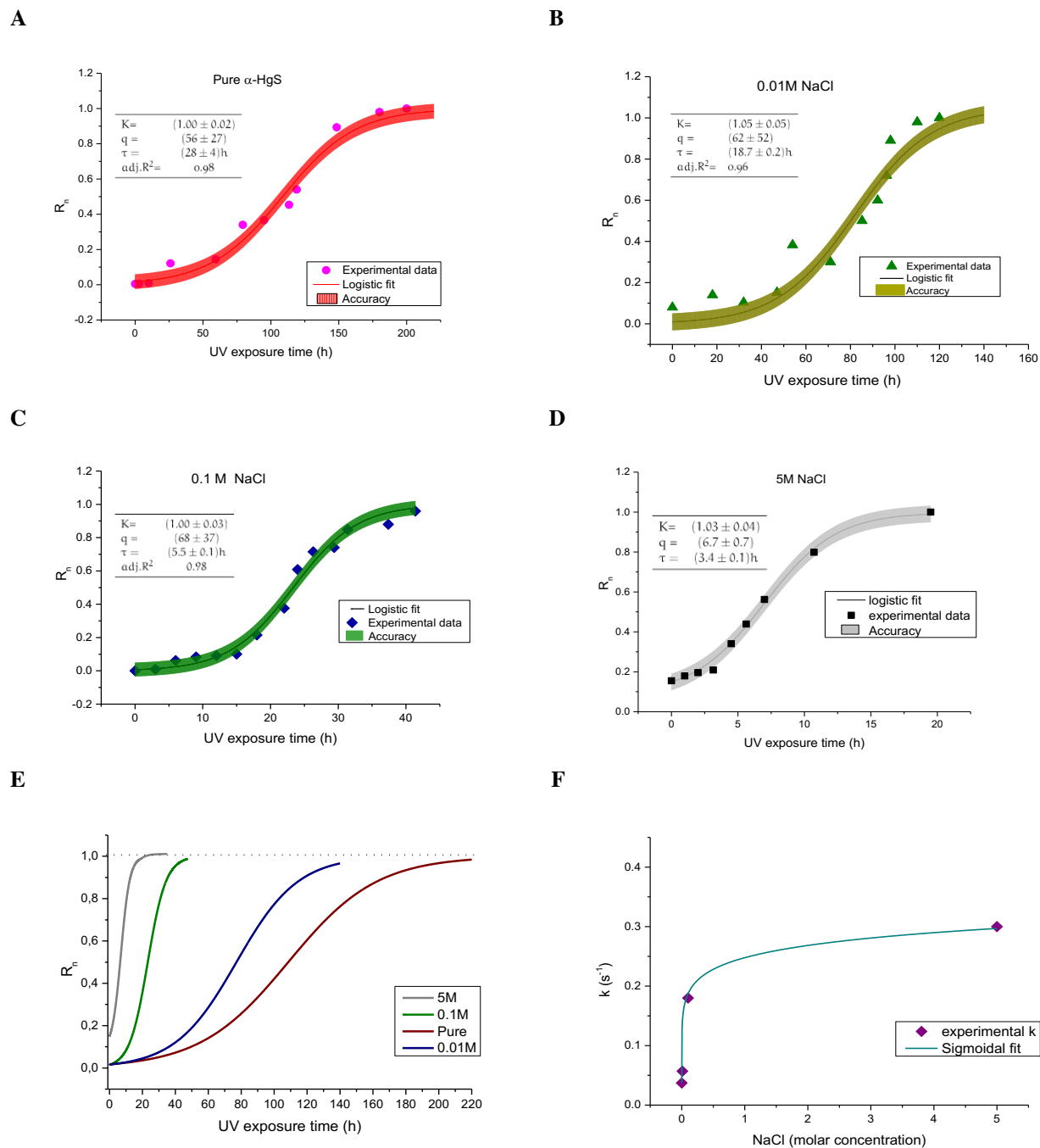
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FIGURE 4: Effect of UV exposure on experimental samples. Comparison between Raman spectra obtained before and after UV exposure on 5M sample (low resolution system) (A). SEM Back Scattered Electrons image where points 3 and 4 represents the darkened area of the pigment after UV exposure, while points 2, 5 and 6 are related to not darkened pigment. Point 1 concerns the glass slide (B).



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FIGURE 5: Pump and probe measurements of pure and 148 h UV darkened samples. Pump-probe (transient absorption) signals as a function of interpulse delays (A). XRD spectra of the degraded sample showed a broadening of the XRD peak near 30.5° related to the phase conversione. Signals normalized to the peak at 31.2° (B).



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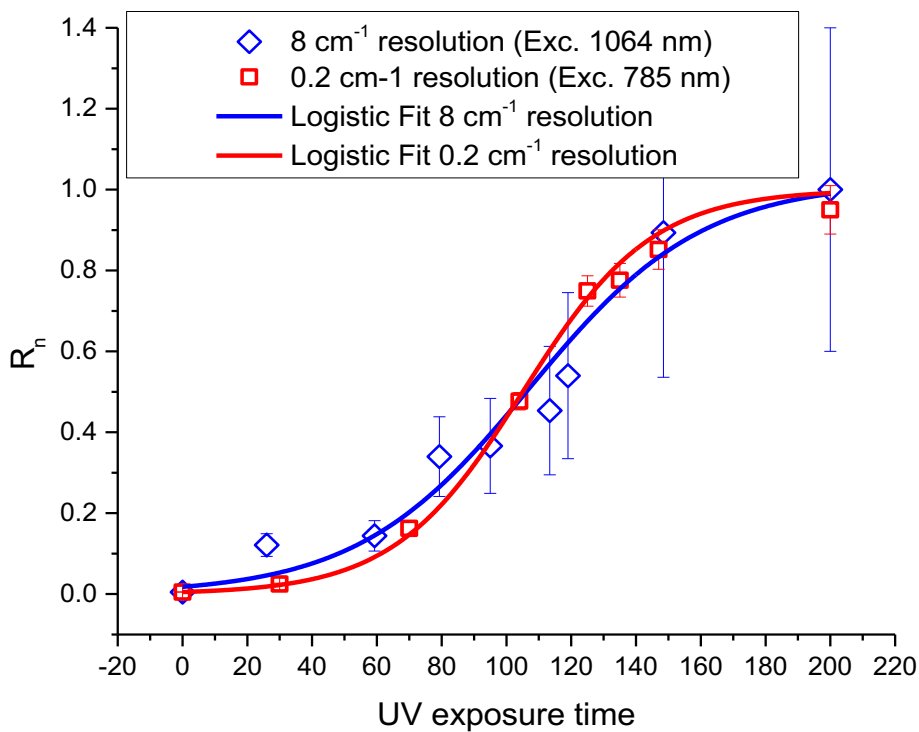
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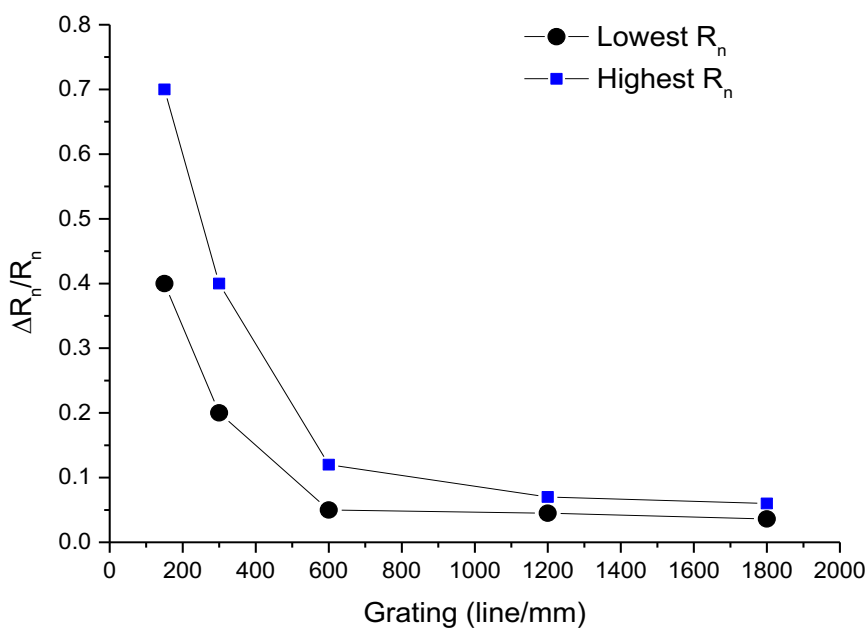
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FIGURE 6: Kinetic model follows logistic function. Experimental points and logistic model applied to the experimental samples. For each curve the characteristic time t and the parameter q is determined (A-D). Comparison among all the curves obtained evidences the role of Cl impurities in the degradation process (E). Determination of degradation rate k as a function of NaCl molar concentration (F). R_n expresses the fraction of $[\beta\text{-HgS}]$ with respect to the total composition (equation 5 and 6).



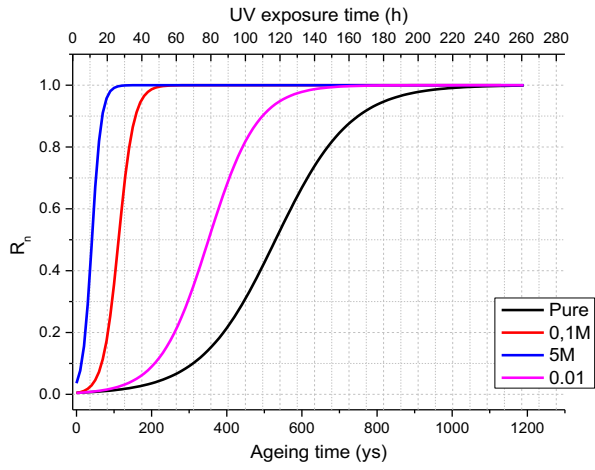
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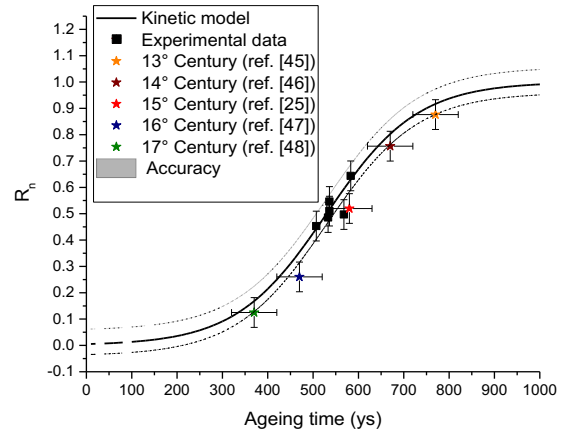
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755 **FIGURE 7: Error estimation in relation to experimental resolution.** Comparison of R_n values and
 756 error bars obtained with experimental systems with 8 cm^{-1} and 0.2 cm^{-1} resolution (A). Relative error of
 757 R_n as a function of grating characteristics.

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FIGURE 8: Ageing model. Correlation between the UV exposure time (hours) and ageing time expressed in years (A). Validation of ageing model applied to literature data with an accuracy of 8% (B).

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	Pure	Pure dark	Δ	5M	5M dark	Δ
L*	55.74	43.05	-12.69	53.06	47.19	-5.87
a*	34.18	4.89	-29.29	23.58	12.92	-10.66
b*	15.84	2.75	-13.09	11.24	6.22	-5.02
ΔE_{Lab}			34.50	ΔE_{Lab}		13.70
Δ				Δ		
C*	37.67	5.61	-32.06	26.12	14.34	-11.78
h	24.86	29.35	4.49	25.49	25.71	0.22

TABLE I: Chromatic coordinates variations calculated for samples 5M and Pure

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Element	Glass slide (at %)	Sample 5M (at %)					Sample Pure (at %)	
	Point 1	Point 2 (No UV)	Point 3 (UV)	Point 4 (UV)	Point 5 (No UV)	Point 6 (No UV)	Point 1 (at %)	Point 2 (at %)
C	13.63	27.78	17.69	17.00	11.00	20.55	17.83	16.83
O	48.72	13.06	25.84	22.71	48.30	14.14	36.84	45.79
Na	7.16	22.67	7.02	5.95	7.29	20.85	4.63	6.37
Mg	1.57	0.38	1.15	0.97	1.81	0.45	1.00	1.06
Al	0.39	0.07	0.35	0.39	0.53	0.21	0.42	0.40
Si	20.31	2.55	5.61	4.55	19.99	3.26	13.32	14.82
S	1.77	5.73	14.62	17.84	3.07	8.71	10.48	5.74
Cl	1.60	21.03	5.44	4.95	1.71	21.25	0.83	0.76
K	0.43	0.07	0.04	0.06	0.41	0.08	0.20	0.29
Ca	2.30	0.29	0.47	0.38	2.19	1.41	1.60	2.29
Hg	2.12	6.36	21.75	25.21	3.70	9.09	12.85	5.63

TABLE II: EDS analysis of samples 5M and Pure.

	Identified compound – SEM/EDS	
	Dark area	Red area
Pure	Metallic Hg (3%) HgS (10%)	HgS
5M	Metallic Hg (7%) HgS (16%)	HgS and metallic Hg (1.24%)

TABLE III: Stoichiometric balance of Hg compounds from data obtained in table I

	% Mass loss per h		
	Pure	0.1M	5M
135°C	1.7%	0.1%	0.7%
200°C	2%	0.2%	0.7%

TABLE IV: Mass loss percentage due to the thermal treatment executed on the experimental samples.

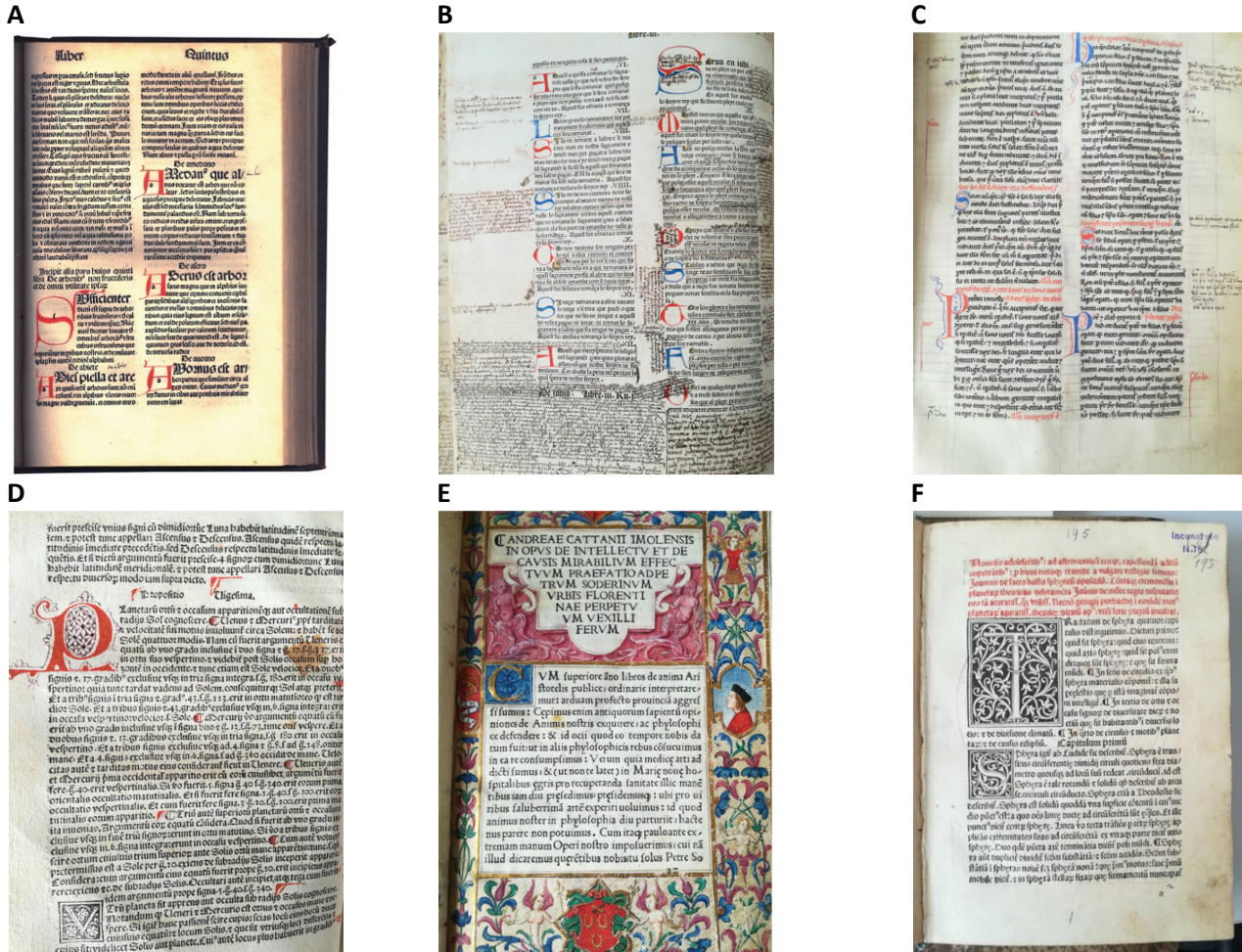
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SUPPLEMENTARY MATERIALS



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FIGURE S1: description of ancient samples from Biblioteca Universitaria di Cagliari (BUCA)

MANUSCRIPTS

Fig. S1A *Ruralium commodorum*, copy of the year 1435. The author was Pietro de' Crescenzi who was a judge from Bologna. This text written on parchment is the most famous medieval agricultural treaty. It was aimed to modernizing the technical knowledge of farmers.

Fig. S1C *Libri sententiarum*, copy written probably in the middle of the XV century. The author the text was Petrus Lombardus, who was one of the most known middle age Catholic theologian. This text written on parchment develops as a theological encyclopedia where the most authoritative opinions are exposed around the main themes of Catholic doctrine.

INCUNABULA

814 **Fig. S1B** *Furs de Valencia*, a text printed in 1482 on paper material by Lambert Palmart in Valencia.
815 Lambert Palmart from Cologne had the responsibility to direct a company of printers in Valencia during
816 the reign of James I. The text is a collection of laws governing various criminals, religious and political
817 matters that correspond to our Statutes.

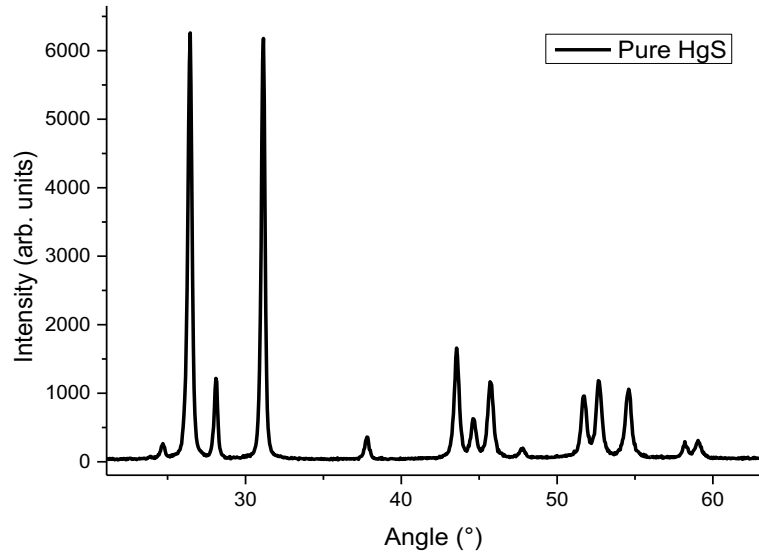
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819 **Fig. S1F** *De Sphaera mundi*, a text printed in 1482 in Venice by Erhard Ratolt on paper material. The
820 author of the original text was Ioannes Sacrobosco, a famous astronomer and mathematician from
821 Holywood (today's Halifax) who studied in Oxford and in Paris where he died around year 1256. The
822 text was the first astronomy treatise given to the press, a very useful work to start young scholars with
823 the principles of cosmology and classical astronomy.

824
825 **Fig. S1D** *Kalendarium*, a text printed in 1485 in Venice by Erhard Ratolt on paper material.
826 The author of the original text was Johannes Regiomontanus (Johann Muller: Koenisberg-1436/Rome-
827 1476)) Astronomer and mathematician.
828 He founded the first European astronomical observatory, his observations on the comet of 1472 (later
829 Halley's comet) marked the beginning of modern cometal astronomy. At the invitation of Pope Sixtus IV
830 he reformed the calendar. The *Kalendarium* was an innovative calendar that predicted solar and lunar
831 eclipses and calculated the day of the year when, from 1477 to 1531, Easter would fall. In reality the text
832 was an updated compendium of astronomical and mathematical knowledge of the time and had an overtly
833 practical and didactic purpose.

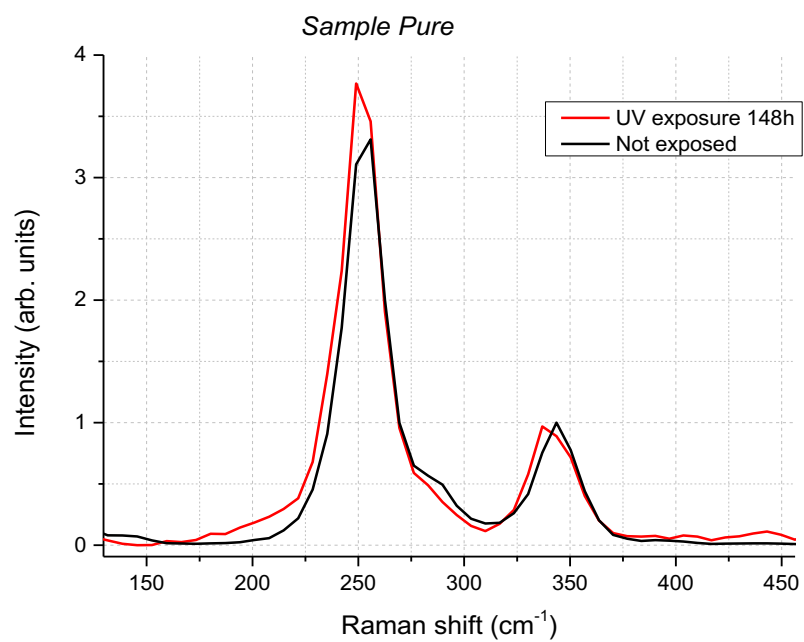
834
835 CINQUECENTINE

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837 **Fig. S1E** *Opus de intellectu et de causis mirabilium effectuum* (printed before 1511 on paper support).
838 The author of the text was Andrea Cattaneo a doctor and philosopher from Imola who was in charge of
839 courses in Firenze and Bologna. He was used to complain that his duties as a physician prevented him
840 from devoting himself to philosophy with the diligence he would have wished. His work is inspired by
841 the doctrine of Avicenna, considered the philosopher closest to Christianity.

842
843
844 For what concerns the sampling procedure, we remand to the standard codicology which analyses the
845 double pages (*recto* and *verso*) of a code for identifying the number of pages associated to each volume.
846 Each *card* is different from the other one and the images present in the page (drawings and miniatures)
847 identifies the folia univocally. In our case, the analysed pages can be recognized by referring to the main
848 detail, for example: “*card with miniature of the king of the FURS of Valencia*”. To be more precise we
849 could identify the card in the following way: paper number n counted from the first cover of the
850 document. This apart from the *Furs de Valencia* (S1B) and *De Sphaera mundi* (S1F) in which a numbering
851 is identified from the photo and that we reported in the figure.
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853
854 **FIGURE S2: XRD spectrum of experimental sample “Pure”.** Principal angles shown in the spectrum is assigned to a-HgS
855 phase confirming the Raman Analysis offered in figure 2.
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FIGURE S3: Comparison between Raman spectra obtained before and after UV exposure on Pure sample

861 **Illustrative case of extraction of the R(t) ratio from Raman spectrum**

862

863 In figure 3A and 3B we reported a deconvolution of the experimental spectra by means of Lorentzian curves. Deconvolution
864 process was operated with the help of the software ORIGIN Pro 9 by performing the Levenberg Marquardt iterative
865 algorithm. Lorentzian function was expressed by the formula:

866

$$y = y_0 + \frac{2A}{\pi} \frac{w}{4(x - x_c)^2 + w^2}$$

867

868 where y_0 is the offset in the ordinates axis, A the area of the curve, x_c the center abscissa and w the width at $y_c/2$. These four
869 parameters were free to variate in order to find the best adjusted R-Square parameter. After 100 iterations each fit operation
870 converged and a Chi-square tolerance value of $1 \cdot 10^{-15}$ was reached.

871

Table S1 reports the results for deconvolution processes of figure 3A and 3B.

872

		x_c	A	ΔA
Figure 3A (Pure 0h UV)	Peak 1	253.9	14.007	0.002
	Peak 2	282.3	0.422	0.005
	Peak 3	289.2	0.959	0.003
	Peak 4	343.6	2.875	0.004
	Peak 5	352.4	0.916	0.004
Figure 3B (Pure 90h UV)	Peak 1	250.5	23.051	0.007
	Peak 2	279.1	0.44	0.01
	Peak 3	286.4	1.32	0.02
	Peak 4	341.5	4.11	0.01
	Peak 5	350.9	0.733	0.008
Figure 3C (Pure 0h UV)	Peak 1	252.3	15.80	0.01
	Peak 2	284.0	1.3	0.3
	Peak 3	243.8	3.8	0.1
Figure 3D (Pure 90h UV)	Peak 1	250.1	25.05	0.01
	Peak 2	279.1	1.5	0.3
	Peak 3	339.6	4.2	0.1

TABLE SI: Results from deconvolution on Raman spectrum

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874

875 R_0 is calculated with the ratio determined by the Region Of Interest (ROI) at about 250cm^{-1} and 280cm^{-1} , variable in
876 relation to the darkening process and phase transformation (see Figure 1). In our case (figure 3A):

877

$$R_0 = \frac{A_{Peak1}}{A_{Peak2} + A_{Peak3}} = 10,14$$

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879 Analogously the same method can be used for calculating the ratio R(t) from peaks of figure 3B:

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$$R(t) = \frac{A_{Peak1}}{A_{Peak2} + A_{Peak3}} = 13,10$$

882

883 Following the definition of equation (5) a calculaiton of R_n is done:

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$$R_n = \frac{R(t) - R_0}{R(t)} = 0,29$$

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887 By operating a linear propagation of the relative errors from equation (5) and from the expressions of R_0 and R(t) proposed
888 before we obtain:

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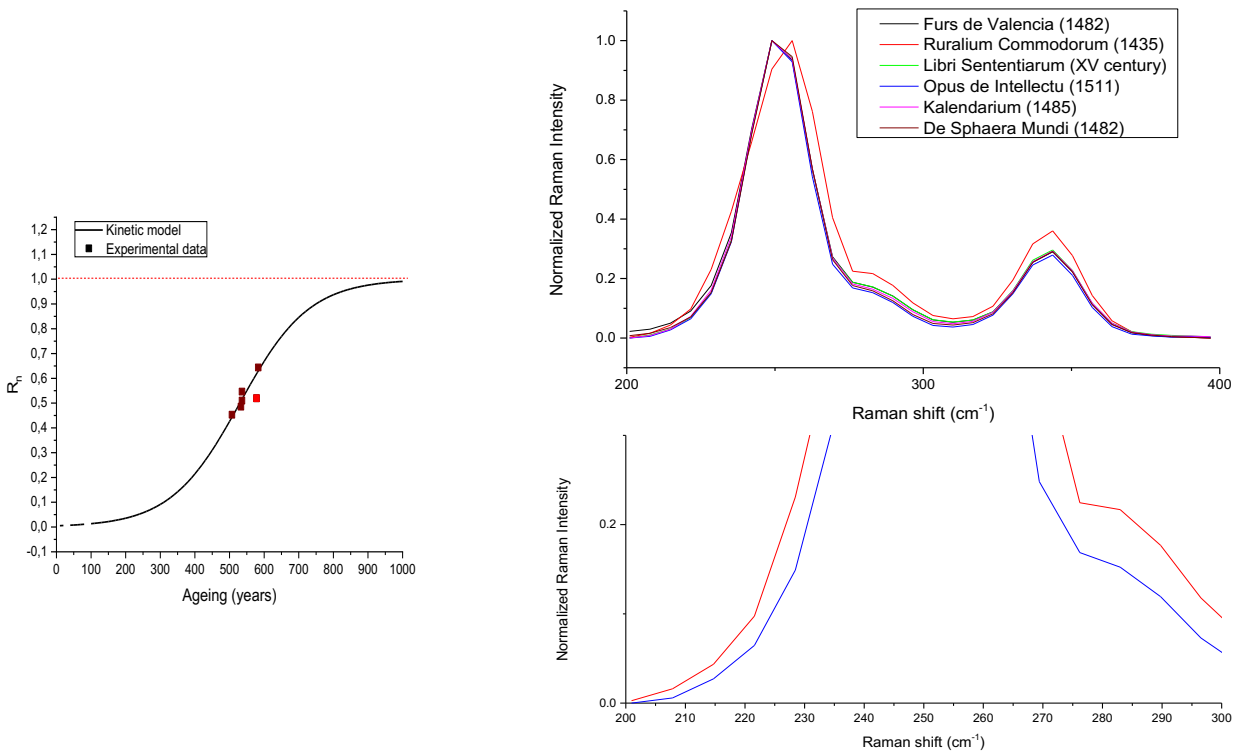
$$\frac{\Delta R_n}{R_n} = \frac{\Delta R_0}{R_0} + \frac{\Delta R(t)}{R(t)} = \frac{\Delta A_{Peak1}^{R_0}}{A_{Peak1}^{R_0}} + \frac{\Delta A_{Peak2}^{R_0}}{A_{Peak2}^{R_0}} + \frac{\Delta A_{Peak3}^{R_0}}{A_{Peak3}^{R_0}} + \frac{\Delta A_{Peak1}^{R(t)}}{A_{Peak1}^{R(t)}} + \frac{\Delta A_{Peak2}^{R(t)}}{A_{Peak2}^{R(t)}} + \frac{\Delta A_{Peak3}^{R(t)}}{A_{Peak3}^{R(t)}}$$

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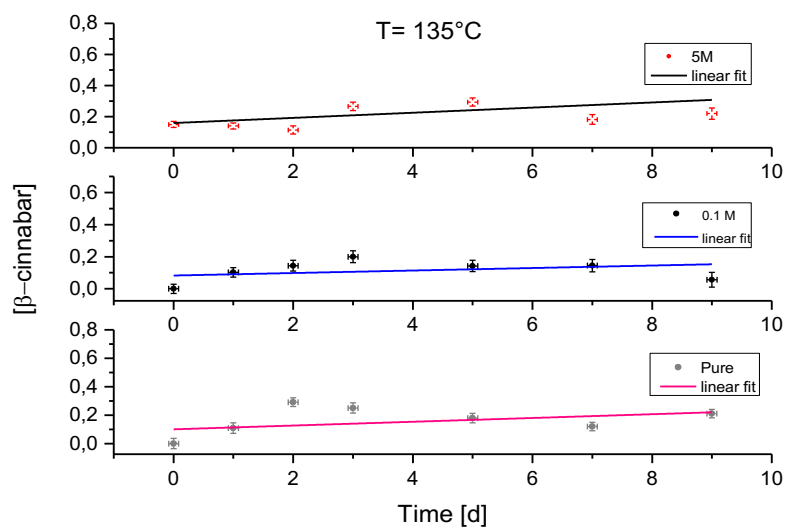
In this specific case, we obtain a relative error $\frac{\Delta R_n}{R_n} = 0,05$ and finally $R_n = (0,29 \pm 0,01)$

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FIGURE S4: Kinetic model applied to the ancient samples from Biblioteca Universitaria di Cagliari (BUCA). Left panel: R_n vs ageing time of BUCA samples. Red point is related to the sample with higher uncertainty. Right panel (up): Experimental spectra of BUCA samples in the 200-400 cm⁻¹ range. Right panel (down): Magnification in the 200-300 cm⁻¹ range for the two samples with the largest variation.



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FIGURE S5: Thermal treatment of the experimental samples. The ageing model and the b-cinnabar concentration is determined as a function of the treatment days.

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Element	5M Pt1	5M Pt2	5M Pt3	5M Pt4	5M Pt5	5M Pt6	5M Pt7	PURE Pt1	PURE Pt2	PURE Pt3	PURE Pt4
	Atom %	Atom %	Atom %	Atom %	Atom %	Atom %	Atom %	Atom %	Atom %	Atom %	Atom %
C K	4,50	3,79	5,32	4,13	4,45	8,04	9,46	4,82	8,76	0,04	0,02
O K	5,13	31,85	31,01	35,26	37,46	35,77	25,01	40,94	15,22	0,02	0,02
Na K	37,93	19,65	21,34	18,62	15,60	16,77	19,86	3,41	0,55	0,00	0,00
Mg K	0,00	1,34	1,26	1,76	0,93	1,17	0,84	0,72	0,00	0,00	0,00
Al K	0,01	0,40	0,15	0,12	0,43	0,15	0,25	0,21	0,00	0,02	0,00
Si K	0,00	8,92	6,95	6,20	9,50	2,68	1,26	7,12	0,00	0,00	0,00
S K	7,54	9,53	9,27	11,51	11,06	10,92	14,29	20,77	38,55	52,40	52,76
Cl K	37,16	15,61	16,52	14,27	11,64	16,52	18,66	1,94	3,85	0,00	0,00
K K	0,09	0,36	0,27	0,23	0,12	0,31	0,23	0,29	0,00	0,00	0,00
Ca K	0,21	0,84	0,81	0,84	1,01	0,58	0,21	0,91	0,16	0,00	0,00
Hg L	7,44	7,69	7,11	7,06	7,80	7,09	9,94	18,87	32,90	47,52	47,21
	100,00	100,00	100,00	100,00	100,00	100,00	100,00	100,00	100,00	100,00	100,00
Ratio Hg/S	0,99	0,81	0,77	0,61	0,71	0,65	0,70	0,91	0,85	0,91	0,89
Ratio Na/Cl	1,02	1,26	1,29	1,30	1,34	1,02	1,06				

TABLE SII: SEM-EDS analysis of “pure” and “5M” treated at 200°C for 24h

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