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Abstract

This study aims at exploring the way paper samples may impact the performance of Single-Metal Deposition (SMD II), a fingerprint detection technique known for its versatility of application as well as its sensitivity regarding porous substrates. To get a broader view on how porous substrates may impact the SMD II performances, 74 North American and European papers types were collected, characterized (UV-visible and infrared spectroscopy, roughness, porosity, and surface pH), and processed as substrates bearing fingerprints. This part of the study represented a first valuable outcome by the number of samples considered. After processing with SMD II, the samples were characterized again with the techniques mentioned above, background staining and fingerprint quality were assessed and associated with a quality score. Overall, no positive nor negative trend was observed between the paper characteristics and the SMD II performance. As a consequence, it is currently still not possible to predict if a paper sample will behave well or bad with SMD II. Of all the monitored parameters, the chemical composition of the surface coating (i.e., silica or calcium carbonate) may be worth exploring further, as it has been observed that some coatings undergo partial degradation during the SMD II process. As a result, secretion residue may be damaged by the chemical solubilization of the support layer if they failed to penetrate deeper into the substrate.

Keywords	TRL level 3 Forensic Science; Chemical analysis; Porous substrate; Surface properties; Gold Nanoparticles
Manuscript category	Technology Readiness Level 3
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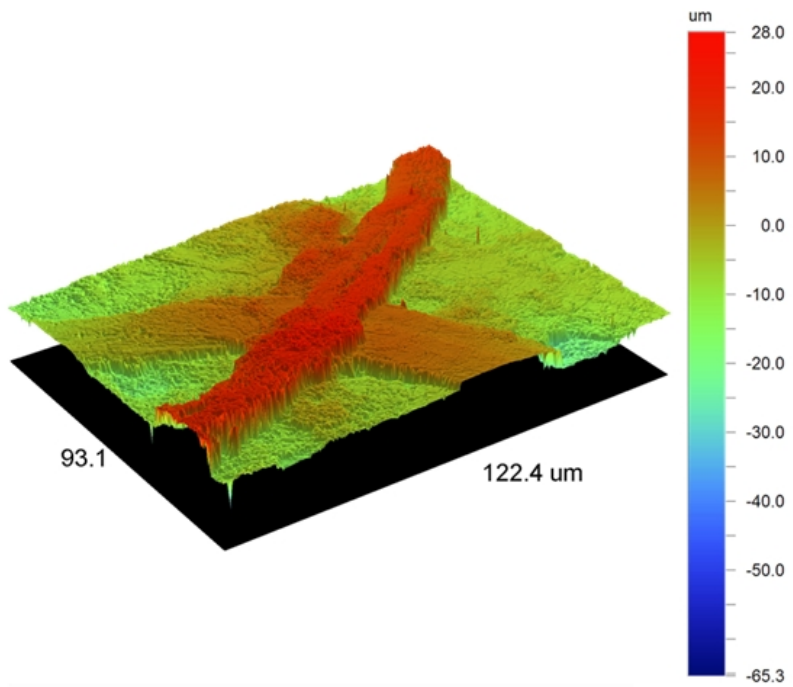
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1 **Highlights**

- 2 • Physical surface topography (roughness and porosity) as well as cellulose and lignin
3 chemical groups have no detectable influence on fingerprints detection using the SMDII
4 technique
- 5 • The only factor that may be of importance seems to be the chemical composition of
6 surface coating (silicates and carbonates).



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9 4 Paper characteristics and their influence on the ability of Single Metal Deposition to
10 5 detect fingerprints
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29 **Abstract**

30 This study aims at exploring the way paper samples may impact the performance of Single-
31 Metal Deposition (SMD II), a fingerprint detection technique known for its versatility of
32 application as well as its sensitivity regarding porous substrates. To get a broader view on
33 how porous substrates may impact the SMD II performances, 74 North American and
34 European papers types were collected, characterized (UV-visible and infrared spectroscopy,
35 roughness, porosity, and surface pH), and processed as substrates bearing fingerprints. This
36 part of the study represented a first valuable outcome by the number of samples considered.
37 After processing with SMD II, the samples were characterized again with the techniques
38 mentioned above, background staining and fingerprint quality were assessed and associated
39 with a quality score. Overall, no positive nor negative trend was observed between the paper
40 characteristics and the SMD II performance. As a consequence, it is currently still not possible
41 to predict if a paper sample will behave well or bad with SMD II. Of all the monitored
42 parameters, the chemical composition of the surface coating (*i.e.*, silica or calcium carbonate)
43 may be worth exploring further, as it has been observed that some coatings undergo partial
44 degradation during the SMD II process. As a result, secretion residue may be damaged by
45 the chemical solubilization of the support layer if they failed to penetrate deeper into the
46 substrate.

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47 **Keywords**

- 48 • Forensic Science
- 49 • Chemical analysis
- 50 • Porous substrate
- 51 • Surface properties
- 52 • Gold Nanoparticles

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57 **Highlights**

- 58 • Physical surface topography (roughness and porosity) as well as cellulose and lignin
- 59 chemical groups have no detectable influence on fingerprints detection using the SMDII
- 60 technique
- 61 • The only factor that may be of importance seems to be the chemical composition of
- 62 surface coating (silicates and carbonates).

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170 **64 1. Introduction**
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173 **65 1.1. Detection of fingerprints using single-metal deposition (SMDII)**
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175 **66** Multimetal deposition (MMD) is a fingerprint detection technique based on the use of colloidal
176
177 **67** gold. The application protocol is built along a two-step process: (i) a detection bath containing
178
179 **68** gold nanoparticles which bind to secretion residue under specific conditions, followed by (ii) a
180 **69** contrast reinforcement bath based on the selective reduction of metal on gold nanoparticles.
181
182 **70** As a result, MMD-processed fingerprints appear as dark/light-grey ridges on a relatively
183
184 **71** unstained substrate [1]. Initially named “The Universal Process” [2] for its ability to detect
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186 **72** marks on a wide range of substrates (e.g., porous, non-porous, semi-porous, adhesives), the
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188 **73** technique was proposed in 1989 [3] and has consistently been improved since, to make it
189
190 **74** more reliable and user-friendly.

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192 **75** Amongst the various improvements, it is possible to cite the optimization of the colloidal gold
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194 **76** synthesis, by Schnetz, to obtain more homogenous (in size and shape) and smaller (from 30
195
196 **77** to 14 nm) nanoparticles [4]. This led to MMD II. Another improvement of the technique
197
198 **78** consisted in replacing the silver-on-gold reinforcement step by a gold-on-gold one [5,6], that
199
200 **79** proved to produce the same quality of results, with more reliable outcomes, improved control
201
202 **80** and cheaper costs. At this stage of development, the technique was renamed Single-Metal
203
204 **81** Deposition (SMD). Finally, the colloidal gold synthesis was further optimized, as well as the
205
206 **82** application protocol to make it more end-user friendly [7,8]. The latest evolution of the
207
208 **83** technique, SMD II [8], is characterized by a modified colloidal gold synthesis and a simplified
209
210 **84** application protocol (e.g., no pH monitoring). As a result, the gold deposition process is more
211
212 **85** reliable and less pH dependent.

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214 **86** The key step of the technique (being MMD or SMD) remains the gold nanoparticles
215
216 **87** deposition onto fingerprint residue, which is not yet fully understood despite the various
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218 **88** optimization and improvement steps. This is a major limiting factor as it makes it difficult to
219
220 **89** cope with apparent unreliability when processing items or substrates. For example, the
221
222 **90** technique can give very good results on problematic substrates, such as cling films [9], but
223
224 **91** suffers from several issues on conventional substrates, such as paper [10]. Among the lack of
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226 **92** reproducibility and inconsistent detection performance observed on papers, it is possible to
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228 **93** cite: unexplained background staining that can diminish the contrast, unwanted deposition of

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94 gold nanoparticles on the substrate instead of the ridges (reversed detection), or absence of
95 detection (null result). In order to fix those issues, a better understanding of the influence
96 papers may have on the SMD performance is consequently required.

97 The main objective of this study is to monitor the effect of the composition and structure of
98 different types of paper from North American and European markets on the detection
99 efficiency of SMD II. Spectrophotometric methods as well as paper physics properties
100 (surface pH, surface profilometry, roughness and porosity) were considered to identify the
101 parameters that may influence the quality of the detected fingermarks or induce unwanted
102 background staining. Such knowledge would help designing a more robust and efficient SMD
103 formulation, so that it can be reliable independently from the types of papers. Readers
104 interested in fingermark composition and detection can refer to the most recent publications in
105 the field, such as [11].

106 **1.2. Paper composition and properties**

107 **1.2.1 Paper chemical composition**

108 Wood represents the major raw material in the manufacture of paper, aside for specialty
109 papers using cotton or linen, or low grade papers using annual plants. The main constituents
110 of wood are cellulose, hemicelluloses and lignin (Figure 1). Other components, known under
111 the general term of extractives, are present in small and variable quantities. Two types of
112 wood can be distinguished: softwoods (coniferous) and hardwoods, differing mostly by their
113 content in lignin (*i.e.*, 25-35% and 18-25%, respectively). It can be noted that the lignin
114 content in tropical hardwoods may exceed that of many softwoods. Softwoods and
115 hardwoods share a similar amount of cellulose (40-50%), and varying structures and
116 quantities of hemicellulose [12].

117 < Insert Figure 1 here >

118 **1.2.2 Cellulose**

119 Cellulose is a polysaccharide consisting of a linear chain of several hundreds to many
120 thousands of $\beta(1\rightarrow4)$ linked D-glucose units. The cellulose macromolecules are organized in
121 a unit called an elemental microfibril (10 nm in width and 5 nm in thickness), in which there

281
282 122 are about 100 cellulosic polymers connected by intra- and inter-molecular hydrogen bonds.
283
284 123 The main characteristic of this polymer is its insolubility in water, which is the result of the very
285
286 124 high molecular mass (3000 glucose units).

287 288 125 **1.2.3 Hemicelluloses**

289
290 126 Hemicelluloses differ from cellulose by the degree of polymerization (150-200), and by the
291
292 127 branching of molecular chains (Figure 1, lower right). The constitutive sugars of
293
294 128 hemicelluloses are divided into four groups: pentoses, hexoses, hexuronic acids and
295
296 129 deoxyhexoses. These units are connected by (1→4) or (1→6) links [12].

297 298 130 **1.2.4 Lignin**

299
300 131 Lignin is a thermosetting polymer with a very strong aromatic character and a molecular
301
302 132 weight that may exceed 40,000 g.mol⁻¹. The main constituting unit is the phenylpropane,
303
304 133 linked by ether-carbon or carbon-carbon bonds [12]. Lignin ensures the cohesion of the fibers
305
306 134 between each other by acting as natural glue. The complexity of lignin is such that much
307
308 135 research is still under way to define its molecular structure in a much more precise way.
309
310 136 Figure 1 shows a model of the chemical structure of softwood lignin at the top of the figure.
311
312 137 Different wood species have different lignin structure and composition.

312 313 138 **1.2.5 Surface roughness**

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315 139 Paper roughness is an important parameter for its physical characterization. It is therefore
316
317 140 essential to be able to quantify the roughness of a paper so that the given value correlates
318
319 141 with the expected use of this paper, for example printing. In our case, it would be interesting
320
321 142 to see if surface roughness can be correlated with the quality of affixing of the secretion
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323 143 residue composing the fingermarks on the different types of paper.

324 144 Roughness is defined as the average distance between the paper surface and a reference
325
326 145 plane to be defined. The roughness indices increase with the roughness of the paper [13].
327
328 146 Various parameters such as Ra, Rq, Rt and Rz are defined to quantify the roughness of a
329
330 147 paper surface (Table 1). Rq is the value we will use in our analysis.

331 148 < Insert Table 1 here >

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149 1.2.5 Porosity

150 Paper has a porous structure formed by a network of fibers. Accordingly, there is a two-phase
151 arrangement in which pores and voids between fibers form an important part of its structure
152 [14]. Paper porosity is correlated with several properties such as absorption, opacity, and ink-
153 paper interaction. The porosity is influenced by the processing conditions, the addition of
154 pigments and chemical additives. For some grades of paper, a coating is applied to the top
155 surface of the paper to change its porosity [15].

156 In forensic science, an appropriate fingerprint detection sequence is usually chosen by
157 associating the item to one of the main substrate classes: porous, non-porous or semi-porous
158 (if we exclude specific substrates such as adhesives, metals, etc.). These categories are
159 based on the apparent (empiric) porosity of the substrate, which is known to influence the
160 behavior of the secretion residue [11]. Office papers are associated with porous substrates,
161 while magazine papers are usually considered as semi-porous substrates.

162 2. Material and Methods

163 2.1. Paper collection and characterization

164 2.1.1 Paper sampling

165 74 different kinds of paper (e.g., inkjet, LaserJet, copier, envelope, newsprint, Offset, drawing,
166 artistic) from 70 to 275 g.m⁻² basis weight were used in this study (see Appendix A for
167 details). These paper samples originated from Europe (e.g., Germany, Austria, Finland,
168 France, Portugal, Sweden and Switzerland), Canada, Mexico, and United States of America.
169 The samples were characterized by a range of paper composition: sugar cane (95%),
170 cardboard, colored, recycled fibers (10, 20, 30, 50% and even 80% post-consumer fibers),
171 wheat, FSC (Forest Stewardship Council) approved, Kraft, bleached Kraft, bleached generic
172 and blended mixed FSC.

173 2.1.2 UV-Visible-NIR spectroscopy

174 UV-visible spectra were taken on a Varian/Agilent Cary 5000 UV-VIS-NIR spectrophotometer
175 equipped with a diffuse reflectance integrating sphere (350-850 nm for UV-Visible part and
176 4000-600 cm⁻¹ for IR). The choice of this method is necessary because paper is a solid,

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394 177 strongly absorbent, and highly diffusive material. The main functional groups responsible for
395
396 178 the sensitivity to light of lignin are the carbonyl and phenolic groups, quinones and various
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398 179 conjugated double bonds [12].

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400 180 Ten spectra of each paper sample were recorded, averaged and analyzed using the
401
402 181 ACD/SpecManager™ version 12.00 from ACD/Labs (*Advanced Chemistry Development*)
403
404 182 software.

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406 183 **2.1.3 Profilometry**

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408 184 3D profiles of the paper surface were recorded with a contact free optical profilometer (Veeco
409
410 185 Wyko NT1100™ instrument) using a Mirau interferometer. Phase shift interference (PSI) and
411
412 186 vertical offset interference (VSI) can be used to respectively measure smooth and rough
413
414 187 surfaces (heights that can reach up to 1mm). Those two modes were used to optimize
415
416 188 detection and measurements of the paper samples.

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417 189 **2.1.4 Porosity measurements**

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419
420 190 Porosity measurements were carried out with a Parker Print Surf™ (PPS) device from
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422 191 Hagerty Technologies™. The flow of a fluid (air in our case) that passes through the paper
423
424 192 was measured with a pressure of 1960 kPa.

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425 426 193 **2.1.5 Surface pH measurements**

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428 194 pH measurements were carried out with a pH Pencil from HYDRION™, measuring a gradient
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430 195 of H₃O⁺ ions on the paper surface. The first step was to moisten the surface of the paper with
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432 196 distilled water, then to mark a line with the pen. After 15 minutes, the color of the line was
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434 197 compared with the shades of color (color sheet) accompanying the pen. Although this method
435
436 198 is not fully accurate, it is a good way to discriminate a wide range of surface pH otherwise
437
438 199 very difficult to measure, and as pH is the most critical parameter to control for SMD
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440 200 development, it could be planned that such an easy semi-quantitative pH tester could be
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442 201 deployed to assist practitioners.

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442 202 **2.2. Fingermark collection**

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203 Natural and sebum-rich marks from two donors were collected for this study. For the natural
204 fingermarks, the donors were asked not to wash their hands one hour prior deposition. No
205 intentional enrichment was performed before collecting the fingermarks. For the sebum-rich
206 marks, the donors rubbed their hands on their forehead before depositing the fingermarks.
207 One natural and one sebaceous-rich marks were collected in duplicate for each donor and
208 substrate. Fingermarks were left to age for one month in the dark. This aging period has been
209 chosen to avoid the processing of fresh marks (e.g., one-day-old or one-week-old marks) and
210 to focus on marks compatible with a casework timeline. Temperature and humidity were not
211 monitored, nor controlled.

212 **2.3. Fingermark detection, quality rating and background evaluation**

213 The paper samples bearing fingermarks were processed using the latest SMD II protocol [8].
214 Given that paper can modify the pH of the solution and have an adverse impact on the
215 results, the paper samples were cut so that they all weight the same mass. Each paper
216 sample was then processed in 200 ml of colloidal gold solution. Since the focus of the study is
217 to investigate the effect of the different types of paper on the SMD II performance, each paper
218 type was processed in a newly prepared bath of colloidal gold. After completion of the SMD II
219 protocol, the samples were left to dry before being scanned on an Epson Perfection V330
220 Photo™ at 1200 dpi, without any digital enhancement. Once scanned, each mark was rated
221 by three independent assessors using a scale ranging from 0 to 3 (Table 2 [16]).

222 SMD II is known to produce unwanted, uncontrolled and non-homogeneous darkening of the
223 porous substrate. In order to understand what parameters may trigger background staining,
224 the color of each paper was recorded before and after fingermark detection. Background
225 measurement was done as follows: for each paper type, an unprocessed sample was placed
226 next to a processed sample and photographed under a homogeneous lighting. Photographs
227 were taken in grey scale and the value of the color was extracted using the eyedropper tool
228 on Adobe Photoshop. Those values range from 0 (black) to 255 (white). For unprocessed
229 samples, one measurement was made in the center of the paper. Processed samples
230 required to conduct four measurements at four different locations which were then averaged,
231 to take background staining inhomogeneity into account. The obtained value was then
232 subtracted from the value of the unprocessed sample. A positive value means a darkening of
233 the substrate whereas a negative value means a lightening.

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234 < Insert Table 2 here >

235 **2.4. Statistical Analysis**

236 In order to highlight the potential correlation between the results of SMD II and the different
237 analyses performed on paper samples, a data analysis was performed. As a first step, the
238 raw results were organized using a Microsoft Office Excel spreadsheet. The analytical part
239 was performed with the data processing software "R 3.0.2". The different methods of analysis
240 considered were (i) the chi-square test where each variable extracted from the paper
241 analyses was assessed against the results of SMD II (fingerprint quality and background
242 staining) and (ii) a joint analysis of variables using principal component analysis (PCA),
243 multiple correspondence analysis (MCA) and multiple linear regression (MLR).

244 **3. Results and Discussion**

245 **3.1 Paper characterization (before SMD II)**

246 **3.1.1 UV-Visible spectroscopy**

247 Figure 2 shows the processing of the UV-Visible spectrum obtained for the sample "C03".
248 Each processed spectrum represents the variation of the log of the inverse of the reflectance
249 as a function of wavelength.

250 < Insert Figure 2 here >

251 Figure 3 shows the UV-Visible spectra of some North American (C01 and C02) and European
252 (E21 and E31) samples. The spectra show absorptions in the UV-Visible region. These
253 absorptions are due to the presence of lignin and the colored products of the paper (dyes,
254 coating pigments).

255 < Insert Figure 3 here >

256 It is possible to deconvolute the spectra to identify the electronic transitions between the
257 various occupied molecular orbitals (OMO) and unoccupied molecular orbitals (UMO), if these
258 are involved in our study.

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259 The 100 to 350 nm region has not been considered because of the intense background noise
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260 cause by the presence of even a small amount of lignin. UV-Visible analysis of SMD II-
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261 processed samples shows an increase in absorption for some of the papers and a decrease
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262 for others.
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263 The reduction in the intensity of absorption of the spectral bands can be explained by 1) the
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264 oxidation of the chromophores during the SMD II process. This has led to the displacement of
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265 the other bands at longer wavelengths, towards the red part of the spectra (bathochromic
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266 effect) and 2) the breaking the double bonds and formation of new compounds by
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267 modification of polarity. Other papers are characterized by hypochromic displacements to
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268 shorter wavelengths (towards ultraviolet).
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269 **3.1.2 IR spectroscopy**

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270 The four most important infrared regions for cellulose are the region of the bound and free OH
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271 elongations between 3660 and 3000 cm^{-1} , the region of the aliphatic CH elongations between
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272 3000 and 2800 cm^{-1} , the region of elongations C-O alcohols (or a cyclic system) of between
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273 1350 and 1000 cm^{-1} , as shown in Table 3.
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274 IR analysis showed that the many bleached papers samples designed for printing purpose
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275 contain very small amounts of lignin. We also observed that about 80% of the samples (apart
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276 from photographic papers and some colored papers) possess a carbonate coating, mainly
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277 because these papers are intended for printing.
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278 Carbonate characteristic peaks are located between 2530-2500, 1815-1770, and 1490-
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279 1370 cm^{-1} (CO_3^{2-} elongation band), 910-850 cm^{-1} (O-C-O deformation band), 885-870 cm^{-1}
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280 and 715 cm^{-1} [17]. FTIR analysis of a calcium carbonate powder allowed the identification of
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281 these bands and with the use of the ACD/SpecManager software, identification of these
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282 bands in the paper samples was possible. This allowed subsequent verification of the
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283 presence or loss of the carbonate layer after treatment with the SMDII.
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284 < Insert Table 3 here >
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285 **3.1.3 Profilometry and roughness**

617
618 286 The profilometry allowed obtaining the 3D topography of all paper samples (Figure 4). For
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620 287 sample C01, which is a glossy white paper used for inkjet printing, a uniform surface has
621
622 288 been measured, with very low Rq ($0.068 \pm 0.009 \mu\text{m}$). This makes sense given that the surface
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624 289 is coated with a layer of mineral pigments which makes the surface of this paper smoother
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626 290 and brighter.

627 291 < Insert Figure 4 here >
628

629
630 292 Figure 5 shows the value and variation of the Rq values of North American and European
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632 293 papers. The parameters Ra, Rq, Rt and Rz characterizing the roughness of each paper give
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634 294 the same variation, whether for North American or European papers. Rq corresponds to the
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636 295 quadratic mean value of the profile deviations. Rq values ranged from $0.07 \pm 0.01 \mu\text{m}$ to
637
638 296 $6.07 \pm 0.01 \mu\text{m}$ for papers from North America and from $3.1 \pm 0.6 \mu\text{m}$ to $4.5 \pm 1.1 \mu\text{m}$ for samples
639
640 297 from Europe. The standard deviations of the different measurements are high (Figure 5),
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642 298 because the samples are not uniform at the microscopic level.

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644 299 < Insert Figure 5 here >
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646 300 3.1.4 Porosity

647 301 Figure 6 shows the variation in average airflow in mL/min of North American and European
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649 302 samples. It can be noted that there is a very large variation in air flow, which makes it possible
650
651 303 to distinguish the most porous samples from less porous ones.

652 304 < Insert Figure 6 here >
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655 305 We have attempted to find a relationship between surface roughness and porosity (Figure 7).
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657 306 This analysis is presented as the variation of Rq as a function of the airflow rate. Our results
658
659 307 indicate that roughness and porosity of the paper are not correlated

660
661 308 < Insert Figure 7 here >
662

663 309 3.1.5. Surface pH

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665
666 310 The values of surface pH reported for the tested paper samples can be found in Appendix A.
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668 311 The majority of the tested papers have a neutral pH (82%), while only a few have a slightly
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670 312 acidic (10%) or basic pH (3%). This result was expected since most of writing papers are

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674 313 acid-free for document preservation purposes. Only three paper types have a pH of 4 or 5
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676 314 (C01, C20, C41).
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678 315 **3.2 Impact of SMD II on the paper properties** 679

680 316 **3.2.1 Carbonate-coated papers** 681

682
683 317 The IR spectra of carbonate-coated samples before and after the application of SMD II were
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685 318 mathematically subtracted from each other and compared with the spectrum of calcium
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687 319 carbonate. Figure 8 illustrates the results obtained for the paper sample RetroPlus50 Canada
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689 320 (C02). It appears that the loss observed on the spectrum resulting from the subtraction “after-
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691 321 before SMD II” (Figure 8 - top) could be correlated with the loss of calcium carbonate (Figure
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693 322 8 - bottom).
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696 323 < Insert Figure 8 here >
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698 324 This loss can be explained by the fact that SMD II requires to immerse the papers in an acidic
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700 325 solution (pH close to 3). IR analysis allows identifying the peaks of the different components
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702 326 and hence estimate a loss of carbonate. However, the quantitative analysis must be
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704 327 completed with chemical analysis to confirm the proportions of carbonates present. When
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706 328 assessing the % of loss by looking at the carbonate absorption band, it appears that most of
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708 329 the paper samples loose between 20 to 50% of this layer. Some papers undergo a total loss
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710 330 (100%). One hypothesis could be that the detrimental effect that SMD II has on the carbonate
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712 331 layer do have a direct impact on the detection performance. This hypothesis will be
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714 332 investigated further in this contribution.
715

716 333 **3.2.2 Photographic papers** 717

718 334 It was not easy to determine the exact composition of the photographic papers with the IR
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720 335 analyzes. Figure 9 shows the three IR spectra of the sample C01, the first spectrum at the top
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722 336 shows the result of the subtraction between the spectra recorded before and after SMD II was
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724 337 applied. What can be seen on this difference spectrum is that there is a loss of three bands
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726 338 which are at 1721, 1653 and 1423 cm^{-1} and a band amplification at 1584 cm^{-1} which is
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728 339 identifiable in the post-SMD II spectrum.
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730 340 < Insert Figure 9 here >
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730 341 The subtraction between the pre-SMD II spectrum of C01 and another sample, for example
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732 342 C04 (for which we were able to determine its cellulose and carbonate composition) shows
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734 343 that no peak corresponds to cellulose in the C01 spectrum, as shown in Figure 10. The
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736 344 photographic coating layer is probably too thick for the infrared radiation to reach the inner
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738 345 layers of the paper. All the photographic papers collected in this study are also coated with
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740 346 silicate, as shown in our FTIR spectra.

741 347 < Insert Figure 10 here >

742
743 348 Silicate in its various forms (gel, precipitate or colloidal) is the most widely used pigment for
744
745 349 the coating of photographic paper to provide a smooth and shiny surface [18]. The most
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747 350 characteristic peaks of silicate are shown in Table 4 [19,20,21]. All photographic samples
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749 351 considered in this study have a silicate coating. The positions of the spectral bands depends
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751 352 on the type of silica used, as well as on the way the coating recipe is prepared (temperature,
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753 353 solvent used, etc.).

754 354 < Insert Table 4 here >

755
756 355 Post-SMD II IR analysis showed that there was no change at the surface of the photographic
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758 356 paper, and therefore no loss of this layer. This can be explained by the fact that the silica
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760 357 layer is quite thick and that it remains stable at acidic pH (no dissolution of this layer). In
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762 358 reference [18], the author describes the factors influencing the dissolution of amorphous
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764 359 silica. Among these factors, the pH has a limited role as the dissolution is almost negligible for
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766 360 pH below 3 and above 9. Finally, the infrared analysis did not allow the determination of the
767
768 361 type (or composition) of silica used for the coating of the photographic papers considered in
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770 362 this study.

771 363 **3.2.3 Colored papers**

772 364 Some colored papers are coated with carbonate (C03, C08, C32, E30, E31, E32), while
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774 365 others contain very little or no carbonate, such as C23 and C27 (all colors), E27, E28 and
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776 366 E29. For the sample C27, IR analysis shows the presence of cellulose (Figure 11). The other
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778 367 peaks belong to the aromatic compounds present (CH aromatic elongation 3083, 3060,
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780 368 3026 cm^{-1} , elongation C=O 1730 and 1704 cm^{-1} , CC elongation of the aromatic ring at 1601,
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782 369 1493, 1452 cm^{-1} , out-of-plane deformation CH aromatic at 697 cm^{-1}).

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370 For the post-SMD II IR spectrum for sample C23 (with all colors), the intensity of certain
371 bands (1468 and 1445 and 853 cm^{-1}) decreased, while the intensity of other bands increased
372 (1731, 1155 cm^{-1}). The difference between the spectra obtained before and after SMD II
373 emphasizes this decrease but no loss of spectral bands. This is also true for sample C27.

374 < Insert Figure 11 here >

375 Several paper samples (C10, C33, C37, C40, E08, E11, E16, E18, E19, E20, E21, E22, E27
376 E28, E29 E31, E30, E32) showed spectral band losses at 3340-3350 cm^{-1} and at 2920 cm^{-1} .
377 Probably it is the $\text{CH}_2\text{-OH}$ group of the cellulose which is lost during the treatment with SMD II
378 (protonation of the alcohol in an acidic medium). The calculation of the derivative with the
379 ACD/SpecManager software made it possible to identify the IR bands corresponding to the
380 carbonate and the cellulose. The band at 711 cm^{-1} is identified in the carbonate with an
381 accuracy of $\pm 1 \text{ cm}^{-1}$. This band is used to calculate the amount of carbonate lost during the
382 processing with SMD II (this band is easily identifiable) (Figure 12).

383 < Insert Figure 12 here >

384 **3.4 Correlation between the SMD II performance and the paper characteristics**

385 Overall, 592 fingerprint samples were processed and assessed throughout the study, leading
386 to fingerprints detected with scores ranging from 0 to 3. As expected, the technique also led
387 to background noise inducing a darkening of the paper surface for most paper types. The
388 colored papers presented a lightening of the color, most certainly due to the successive water
389 baths of the SMD II protocol.

390 The quality scores that have been associated with the detected fingerprints are
391 representative of the performance of SMD II on each paper sample. To some extent, the fact
392 that the same donors provided natural marks along the study makes it possible to compare all
393 these scores and try figuring out some trends among the paper samples. Different
394 correlations with the detection quality scores were explored: vs porosity (Figure 13), vs
395 surface roughness (Figure 14), vs surface pH (Figure 15), and vs amount of carbonate lost
396 (Figure 16). It was indeed supposed that characteristics such as the paper porosity, the
397 surface roughness or the surface pH would play a direct role in the way SMD II behave
398 (detection quality and background). Also, given that a loss of calcium carbonate has been

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399 observed, it appeared interesting to see if the modification of this layer may be correlated with
400 the SMD II performance.

401 < Insert Figure 13 here >

402 < Insert Figure 14 here >

403 < Insert Figure 15 here >

404 < Insert Figure 16 here >

405 When focusing on papers presenting no apparent issues in regards to fingerprint detection
406 (quality scores close to 3), it can be seen that the porosity is rather low, with average airflow
407 ranging from 0 to 1000 mL/min, the roughness is characterized by Rq values ranging from ca.
408 3 to 4 microns, the surface pH of the paper ranges from 4 to 7 and the loss of carbonate
409 ranges from ca. 30% to ca. 90%. However, papers leading to bad quality fingerprints (quality
410 scores close to 0) also present similar characteristics; their porosity is low to average, with
411 average airflow ranging from 0 to 1500 mL/min, the roughness is characterized by Rq values
412 ranging from ca. 2.5 to 4 microns, the surface pH ranges from 6 to 7, and the loss of
413 carbonate ranges from ca. 10% to 100%.

414 Therefore, from the analysis of Figures 13 to 15, no clear trends can be identified regarding
415 the quality of fingerprints in regards to porosity, surface roughness or surface pH of the
416 papers analyzed. The same observation is made with the loss of calcium carbonate.

417 About the surface pH, it can be noted that papers with acidic surface pH (below 6) led to no
418 zero quality scores, which means that SMD II was able to detect fingerprints for each of
419 them, but with varying quality levels. Beyond that observation, there seems to be no trend
420 between surface pH and SMD II performance. 85% of the paper samples are indeed
421 characterized by surface pH between 6 and 7, with quality scores ranging from 0 to 3.
422 Contrarily to what could be expected, an acidic surface pH is consequently not necessarily
423 associated with better detection quality.

424 From the analysis of some 3D topographies of the different types of samples, it is remarkable
425 that the same type of surface does not give the same quality of revelation of the fingerprints,

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426 as it is for the C02 samples (quality score of 0.67) and C05 (quality score of 1.84). Illustrations
427 of processed paper samples bearing fingermarks are shown in Appendix B.

428 To further refine the analysis, statistical analysis was used to try and detect correlation
429 between background noise, fingermark quality and the paper variables. However, none of the
430 techniques used (Chi-square test, PCA, MCA and MLR) led to the detection of a correlation
431 between the parameters considered.

432 3.5 General discussion

433 Despite the number of different paper types collected, the various paper properties studied
434 and the large number of fingermarks processed with SMD II, no correlation between paper
435 properties and SMD II efficiency was highlighted. However, the chemical composition of the
436 surface coating is worth discussing further.

437 Regarding the experimental design, the number of donors has been voluntarily set low
438 because the study had not for aim to assess the intrinsic performance of SMD II as
439 fingermark detection technique. It rather aimed at studying the influence of paper samples on
440 its ability to detect fingermarks. Doing so requires limiting other influencing parameters, such
441 as the variability induced by donors and the age of the fingermarks. By choosing two average
442 donors, it was possible to assess how the performance of SMD II evolves when different
443 paper samples are considered. Increasing the number of donors would have not modified the
444 overall conclusions of the study and would have imply reducing the number of paper samples
445 to keep the quantity of fingermarks manageable.

446 Surface coating is made of silica or calcium carbonate. It is used to make the surface uniform
447 and improve the printing quality [22]. This coating is however soluble in acidic aqueous
448 solutions. Therefore, immersing the samples in colloidal gold will lead to its partial dissolution.
449 If the fingermark residue does not migrate deep enough in the layer of the paper [23], it will be
450 damaged. The dissolution of the fingermark may rely on two parameters: the thickness of the
451 coating layer and the depth of penetration in the paper. According to Vallette and Choudens
452 [22], and Santos et al. [24], the thickness of the coating is about 15 μm for paper of a weight
453 of 72 g/m^2 and more. It is also known that penetration depth depends on the paper type [23].
454 On coated papers, observed depth could be as deep as 30 μm . It means that even if the layer

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954 455 is entirely removed, a fraction of the fingerprint residue may remain available for detection.

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956 456 Uncoated papers contain calcium carbonate as well to improve their surface characteristics

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958 457 and whiteness. This material may also be solubilised during SMD II processing and lead to

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960 458 fingerprint degradation. Under these circumstances, the dynamic of diffusion of the secretion

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962 459 residue into the substrates is expected to play a major role. Indeed, it could be hypothesized

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964 460 that if the secretion residue has not migrated through the surface coating when the document

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966 461 is processed by SMD II, its chance of being detected would be seriously reduced. The key

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968 462 parameter to consider in a forthcoming study will be the aging time of the fingerprints, as we

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970 463 made the choice to limit our study to one-month-old fingerprints for experimental reasons.

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972 464 This observation is also compatible with another technique known to interact with the non-

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974 465 water-soluble fraction of the secretion residue, that is, physical developer (PD). Previous

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976 466 studies have shown that the performance of PD increases with the age of fingerprints [25].

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978 467 Moreover, this technique requires an acid pre-treatment to neutralize the alkali filler particles

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980 468 and to avoid an overall staining of the item. This appears compatible with the need for

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982 469 secretion residue to penetrate the substrate beyond the filler/coating layers to have a chance

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984 470 to be detected. Consequently, it may be interesting to correlate such conclusions with the

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986 471 results of the present study, based on SMD II.

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988 472 **4. Conclusions**

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990 473 This study aimed at characterizing several paper types (e.g., surface composition, surface

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992 474 pH, roughness and porosity) before and after the application of SMD II. Furthermore, we

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994 475 investigated the possibility to correlate the measured parameters with the performance of

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996 476 SMD II, in terms of ridge quality and background staining.

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998 477 At the completion of this study, we were able to show that the following parameters show no

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1000 478 correlation with the SMD II performance: paper roughness, porosity and surface pH. IR

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1002 479 analysis showed that 81% of the papers are coated with carbonates and the thickness of this

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1004 480 layer varies from one sample to another. This layer appears to be solubilized during the SMD

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1006 481 II process. Since fingerprints are originally present at the surface of this coating, further

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1008 482 investigation should be carried out considering the correlation between the calcium carbonate

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1010 483 thickness and the SMD II detection performance. One hypothesis is that secretion residue

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1012 484 may migrate below the calcium carbonate layer if it is not too thick, and be further detected by

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1014 485 SMD II despite the dissolution of the carbonate-based coating. This hypothesis is worth being

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486 further studied considering fingermarks of different ages. Moreover, it is expected that these
487 observations will be useful to physical developer as well.

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Table 1

Parameter	Description	Formula
Ra	Arithmetic mean of absolute values of deviations y	$R_a = \frac{1}{n} \sum_{i=1}^n y_i $
Rq	Quadratic mean value of the profile deviations	$R_q = \sqrt{\frac{1}{n} \sum_{i=1}^n y_i ^2}$
Rt	Maximum Profile Height	$R_t = R_p - R_v$
Rz	The mean height difference between the 5 highest peaks and the 5 lowest valleys	$R_z = \frac{1}{5} \sum_{i=1}^5 R_{pi} - R_{vi}$

Table 2

Score	Qualitative observation
0	No ridge, no fingerprint visible
1	Ridges are visible over a small area (or over the whole mark), but it is extremely difficult to retrieve level II characteristics (such as minutiae) due to extremely poor ridge details.
2	Ridges are visible on almost the whole mark; level II characteristics can be retrieved. Nevertheless, the quality is not optimal due to a low contrast, strong background staining or faint ridges.
3	Ridges are very well defined on the whole mark. Level II characteristics can easily be retrieved. The contrast is optimal with no (or extremely faint) background staining.

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Table 3

Frequency (cm ⁻¹)	Attribution
3332	O-H elongation with intramolecular
2897	CH ₂
1634	H ₂ O
1426	CH ₂ symmetrical deformation
1370	C-H deformation
1334	C-H shear (plane)
1316	CH ₂ agitation
1281	C-H deformation
1203	O-H deformation
1160	C-O and C-C elongation + CH ₂ rocking
1105	C-O and C-C elongation + CH ₂ rocking
1052	C-O elongation
1029	C-O elongation
1002	C-O and C-C elongation + CH ₂ rocking
897	Out-of-plane O-H deformation
659	Out-of-plane O-H deformation

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Table 4

Frequency (cm ⁻¹)	Attribution
3700-3200	Si-OH
3360	H ₂ O absorbed
3000-2800	Organic C-H
1733, 1653, 1634	H ₂ O absorbed
1423	CH ₂ symmetrical deformation
1870-960	Vibrational network SiO ₂
1350-500	C-H vibration
1070	Si-O-Si symmetrical elongation
900-980	Free silanol elongation
800-820	Si-O-Si symmetrical elongation

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501 **Figure captions**
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502 Figure 1 Chemical structures of the major wood components: lignin (top), cellulose (bottom
1238 left), and hemicellulose (bottom right) [12]
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504 Figure 2 Example of treatment of the UV-Visible spectrum, from % Reflectance to log of
1242 inverse reflectance allowing spectrum deconvolution, for the Staples Pastel (USA,
505 CO3). Red vertical line is λ_{max} , blue line is actual spectrum, green curves are the
1244 resulting deconvoluted bands representing electronic transitions, and black
506 spectrum represent the result of the fitted deconvolution.
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509 Figure 3 UV-Visible spectra of some of the analyzed paper samples. a) Kirkland Signature
1251 (Mexico, C01), b) RetroPlus50 (Canada, C02), c) Esquisse envelope (France,
510 E21), d) Papyrus rainbow (Europe – unspecified country, E31).
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512 Figure 4 Left half: 3D profiles of the RetroPlus50 (Canada; C02) and Staples Sustainable
1255 Earth Copy Paper (USA; C05) paper samples after they were processed with SMD
513 II. Right half: illustration of the processed samples.
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515 Figure 5 Average values of the Rq parameter (μm) for all the paper samples (see Appendix
1262 A for manufacturer details).
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517 Figure 6 Average air flow (mL/min) measured for all the paper samples (see Appendix A for
1266 manufacturer details).
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519 Figure 7 Chart illustrating the relation between the average airflow (mL/min) and the Rq
1270 values (microns) for all the paper samples. Each dot represents a paper sample.
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521 Figure 8 Top spectrum resulting from the subtraction of the IR spectra obtained before and
1273 after the application of SMD II on the paper sample RetroPlus50 Canada (C02);
522 bottom IR spectrum corresponding to calcium carbonate.
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524 Figure 9 Top spectrum obtained by subtracting the IR spectra obtained before (middle) and
1280 after (bottom) the application of SMD II (paper sample: Kirkland Signature Mexico;
525 C01).
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527 Figure 10 Difference spectra between C04 (top) and original C01 (bottom).

528 Figure 11 Infrared spectrum of the unprocessed Staples "Chemise à pochettes – 1336" paper
529 sample (C27).

530 Figure 12 Derivative calculation for the RetroPlus50 paper sample (Canada; C02). Top
531 spectra represent the paper surface with the fingermarks revealed, while bottom
532 spectra represent the opposite surface of the same paper.

533 Figure 13 Chart illustrating the relation between the airflow (mL/min) and the average quality
534 score associated with the fingermarks obtained after SMD II. Each dot represents a
535 paper sample.

536 Figure 14 Chart illustrating the relation between the Rq values (microns) and the average
537 quality score associated with the fingermarks obtained after SMD II. Each dot
538 represents a paper sample.

539 Figure 15 Chart illustrating the relation between the surface pH and the average quality score
540 associated with the fingermarks obtained after SMD II. Each dot represents a
541 paper sample.

542 Figure 16 Chart illustrating the relation between the calcium carbonate loss (estimated %)
543 and the average quality score associated with the fingermarks obtained after SMD
544 II. Each dot represents a paper sample.

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547 **Table captions**

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548 Table 1 Parameters Ra, Rq et Rz used to qualify paper surface roughness (n being the number of peaks of the profile).

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550 Table 2 Table used to assess the quality of the marks (reproduced from [16]).

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551 Table 3 Main infrared peaks characteristic of cellulose in the majority of papers studied
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553 Table 4 The main infrared peaks characteristic of the silicate gel [19,20,21]

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557 Competing statement

558 The authors declare that they do not have any competing interest to declare.

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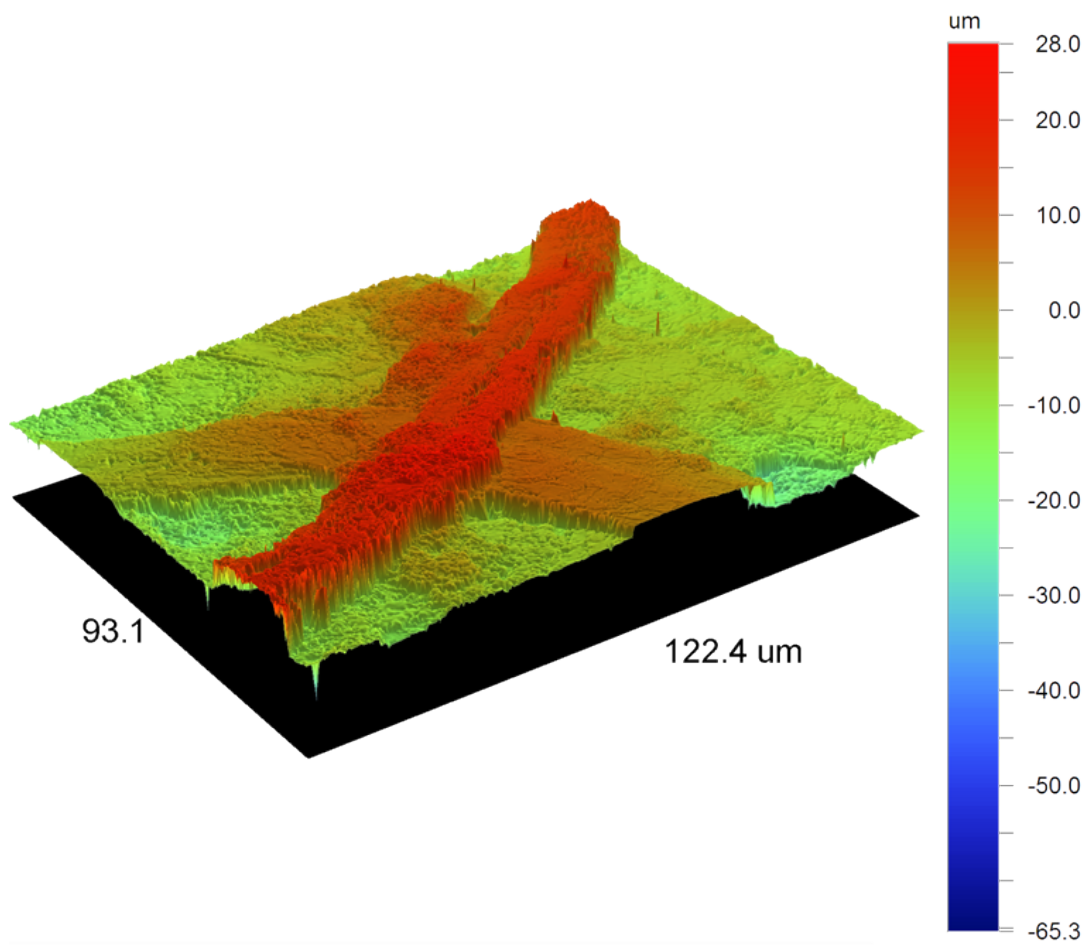
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Figures

Graphical Abstract (1328 x 531 pixels at 96 dpi)



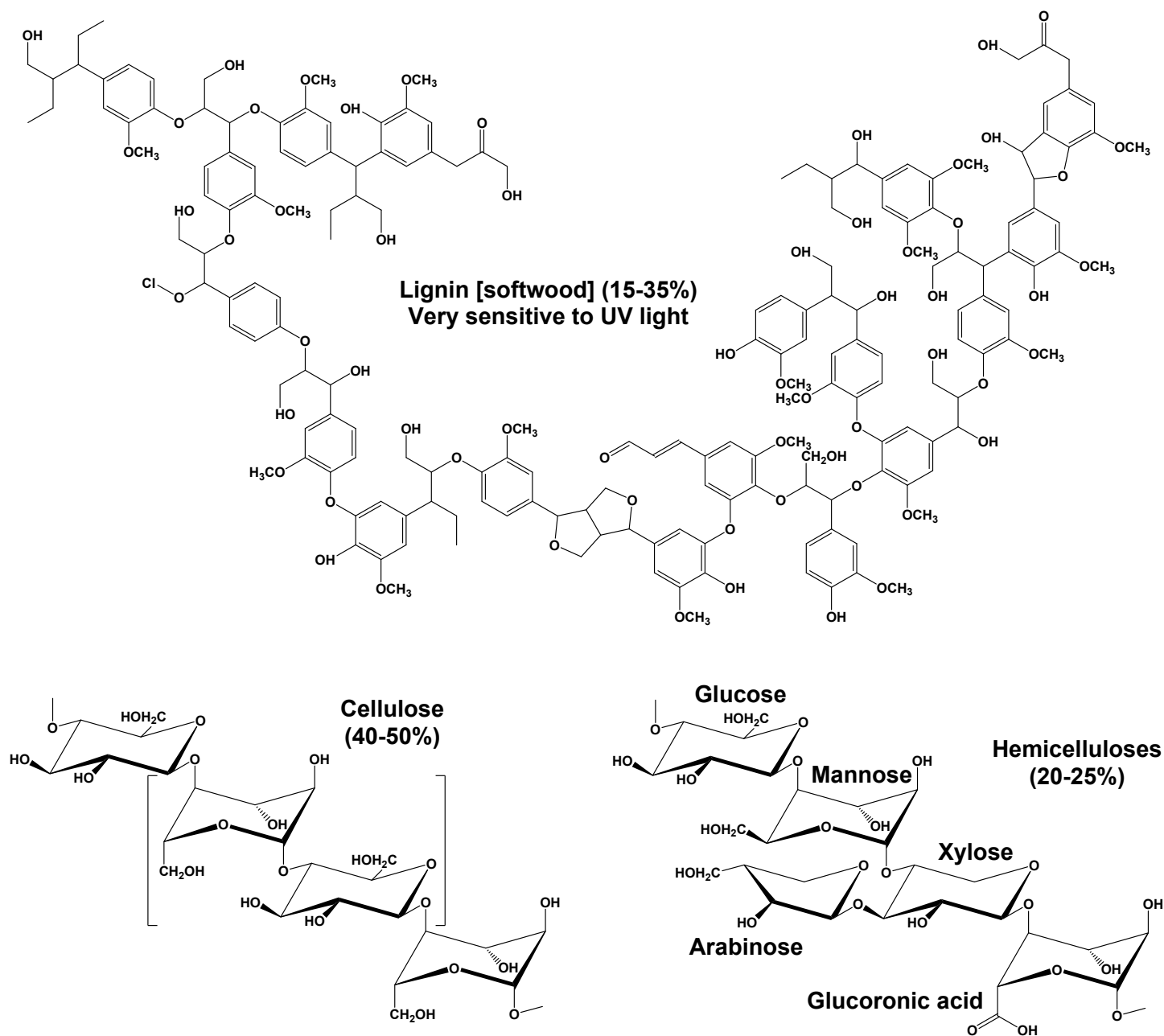


Figure 1 Chemical structures of the major wood components: lignin (top), cellulose (bottom left), and hemicellulose (bottom right) [12]

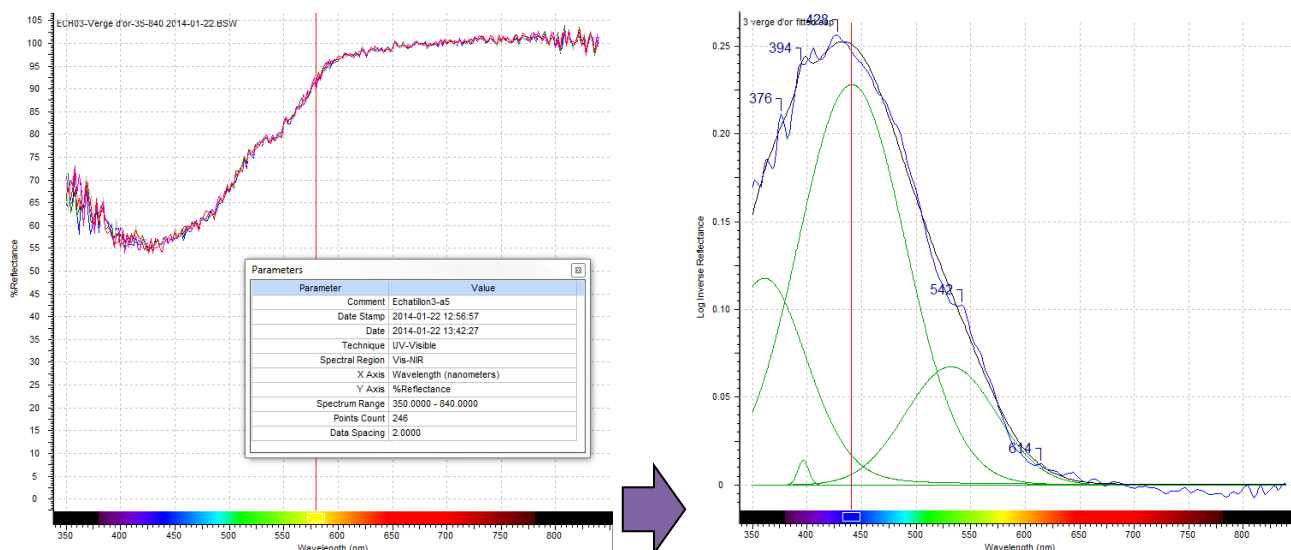


Figure 2 Example of treatment of the UV-Visible spectrum, from % Reflectance to log of inverse reflectance allowing spectrum deconvolution, for the Staples Pastel (USA, CO3). Red vertical line is λ_{max} , blue line is actual spectrum, green curves are the resulting deconvoluted bands representing electronic transitions, and black spectrum represent the result of the fitted deconvolution.

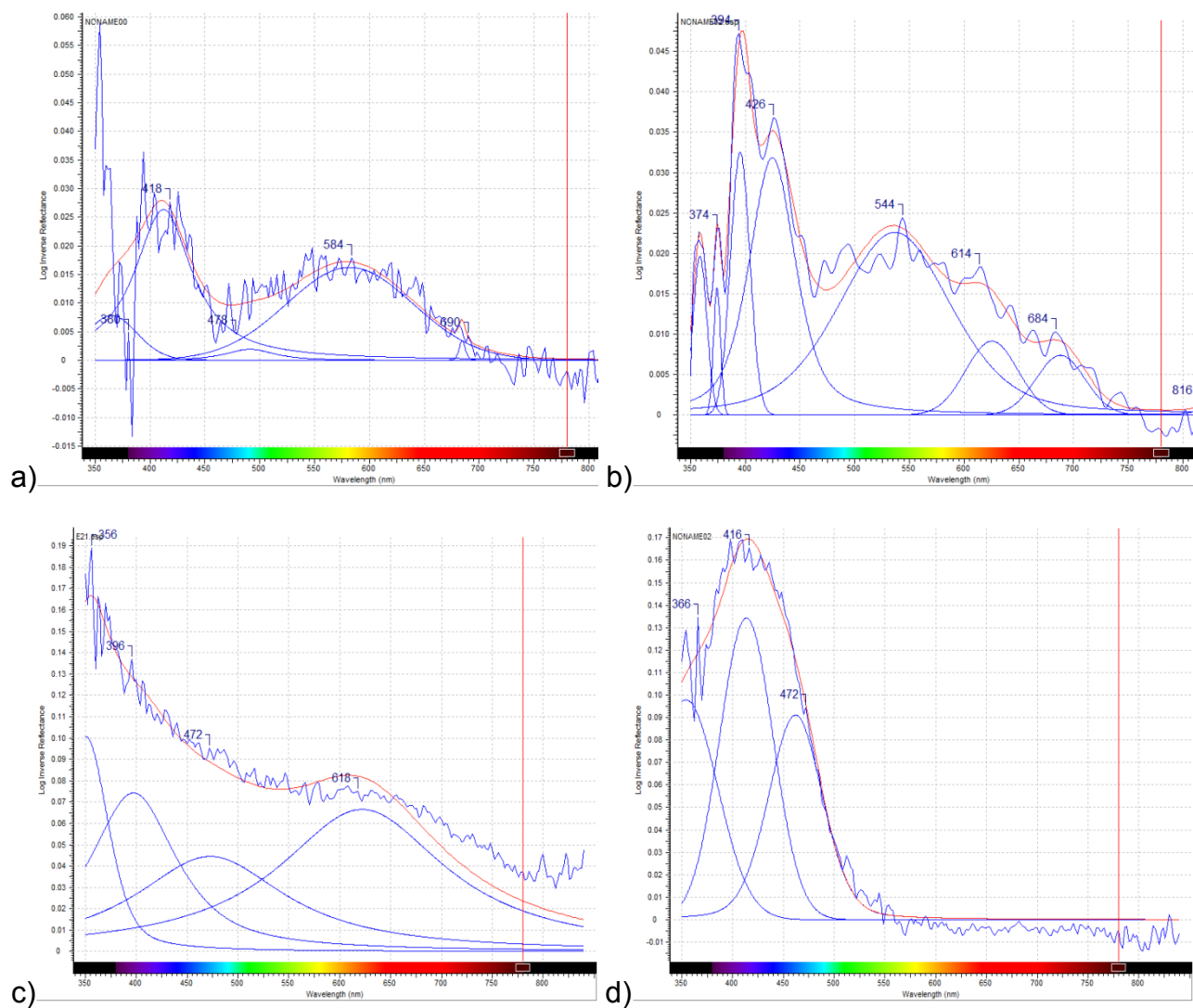


Figure 3 UV-Visible spectra of some of the analyzed paper samples. a) Kirkland Signature (Mexico, C01), b) RetroPlus50 (Canada, C02), c) Esquisse envelope (France, E21), d) Papyrus rainbow (Europe – unspecified country, E31).

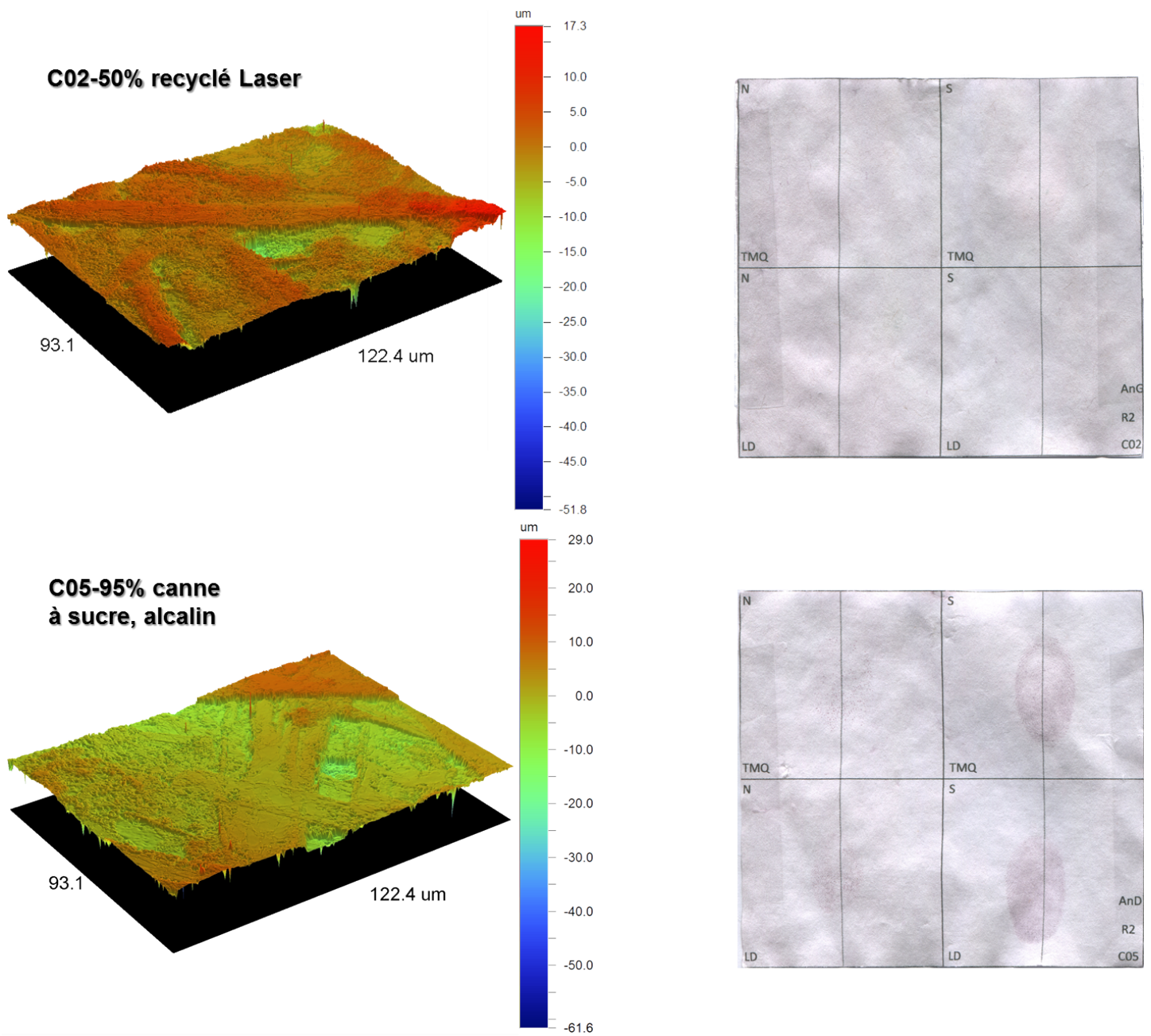


Figure 4 *Left half: 3D profiles of the RetroPlus50 (Canada; C02) and Staples Sustainable Earth Copy Paper (USA; C05) paper samples after they were processed with SMD II. Right half: illustration of the processed samples.*

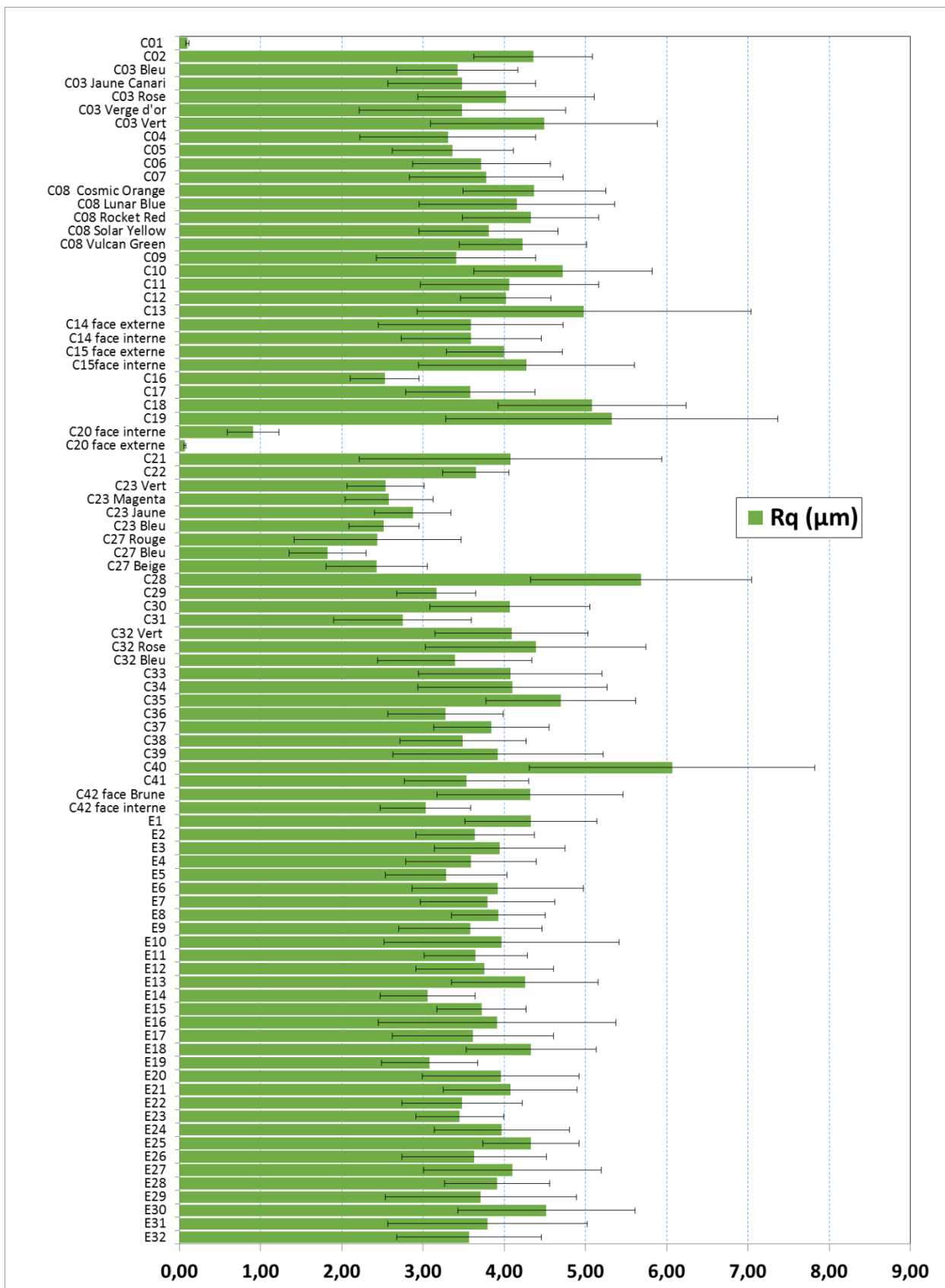


Figure 5 Average values of the Rq parameter (µm) for all the paper samples (see Appendix A for manufacturer details).

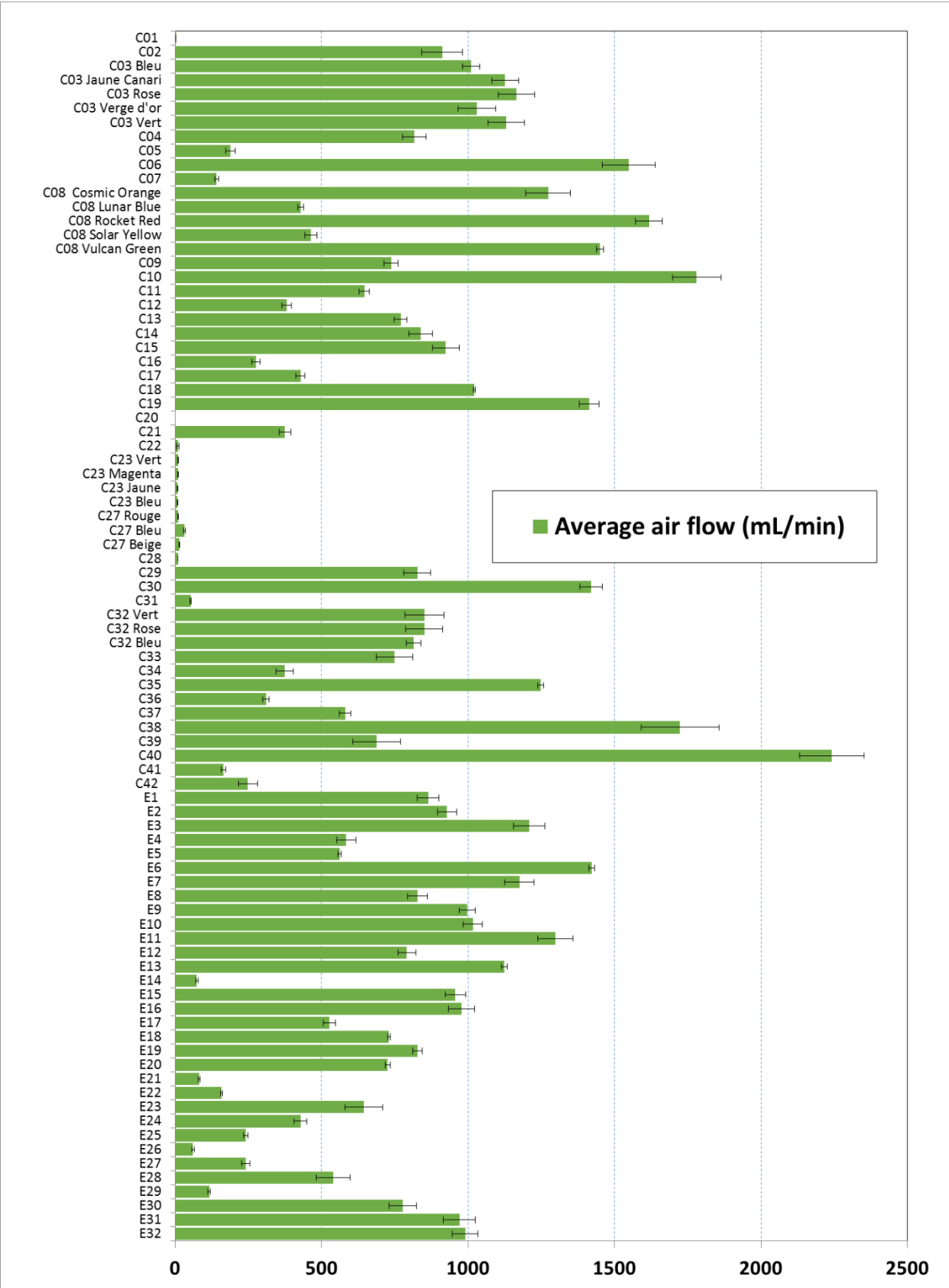


Figure 6 Average air flow (mL/min) measured for all the paper samples (see Appendix A for manufacturer details).

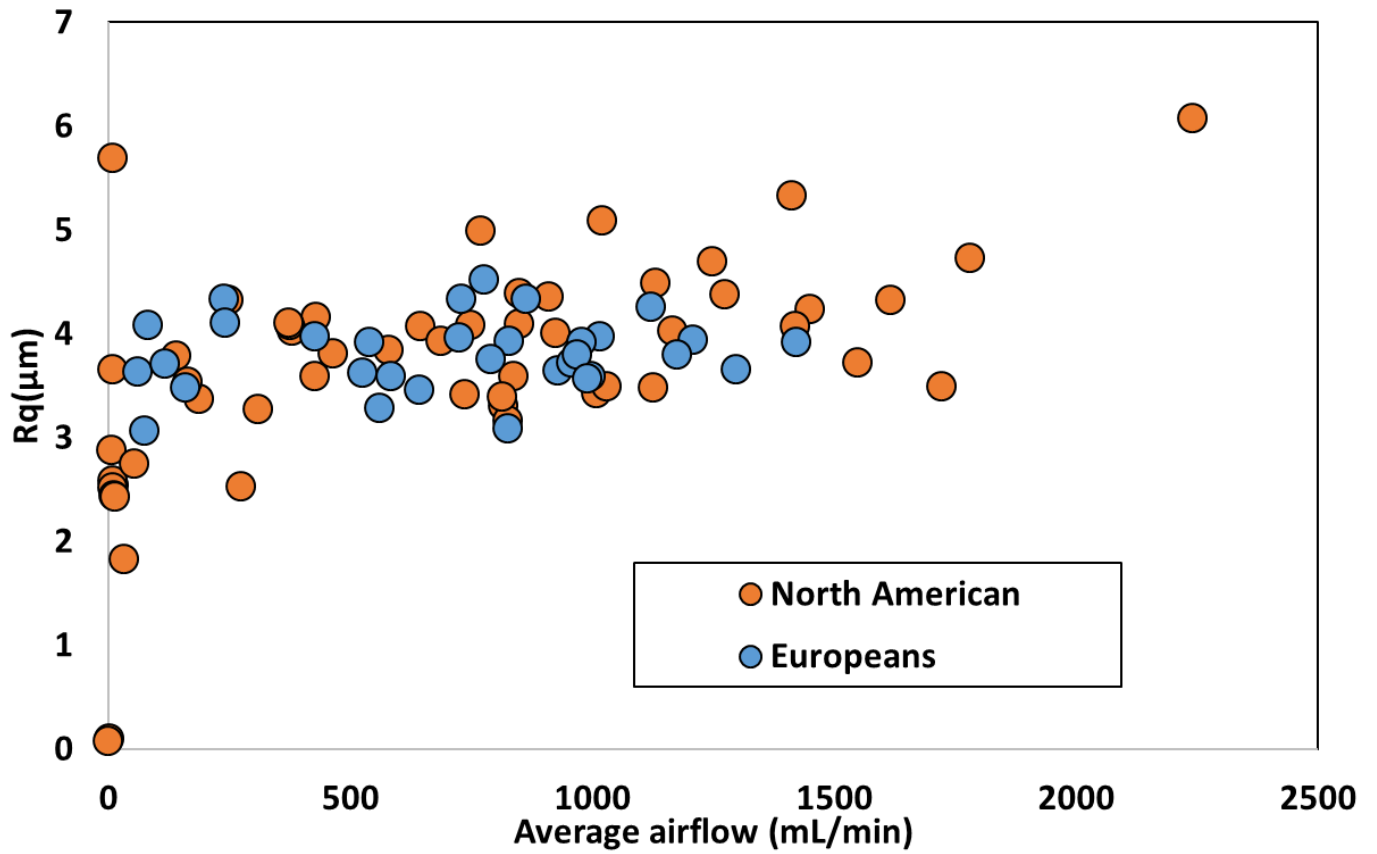


Figure 7 Chart illustrating the relation between the average airflow (mL/min) and the Rq values (microns) for all the paper samples. Each dot represents a paper sample.

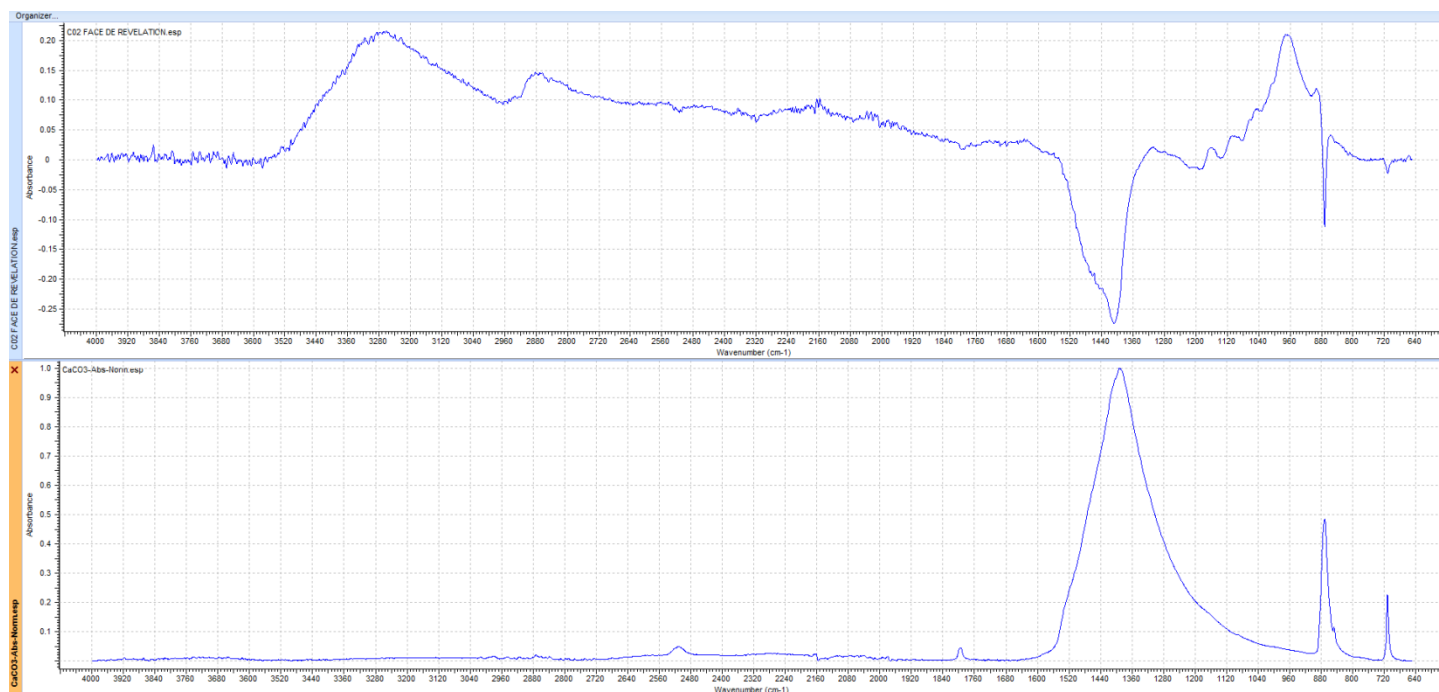


Figure 8 Top spectrum resulting from the subtraction of the IR spectra obtained before and after the application of SMD II on the paper sample RetroPlus50 Canada (C02); bottom IR spectrum corresponding to calcium carbonate.

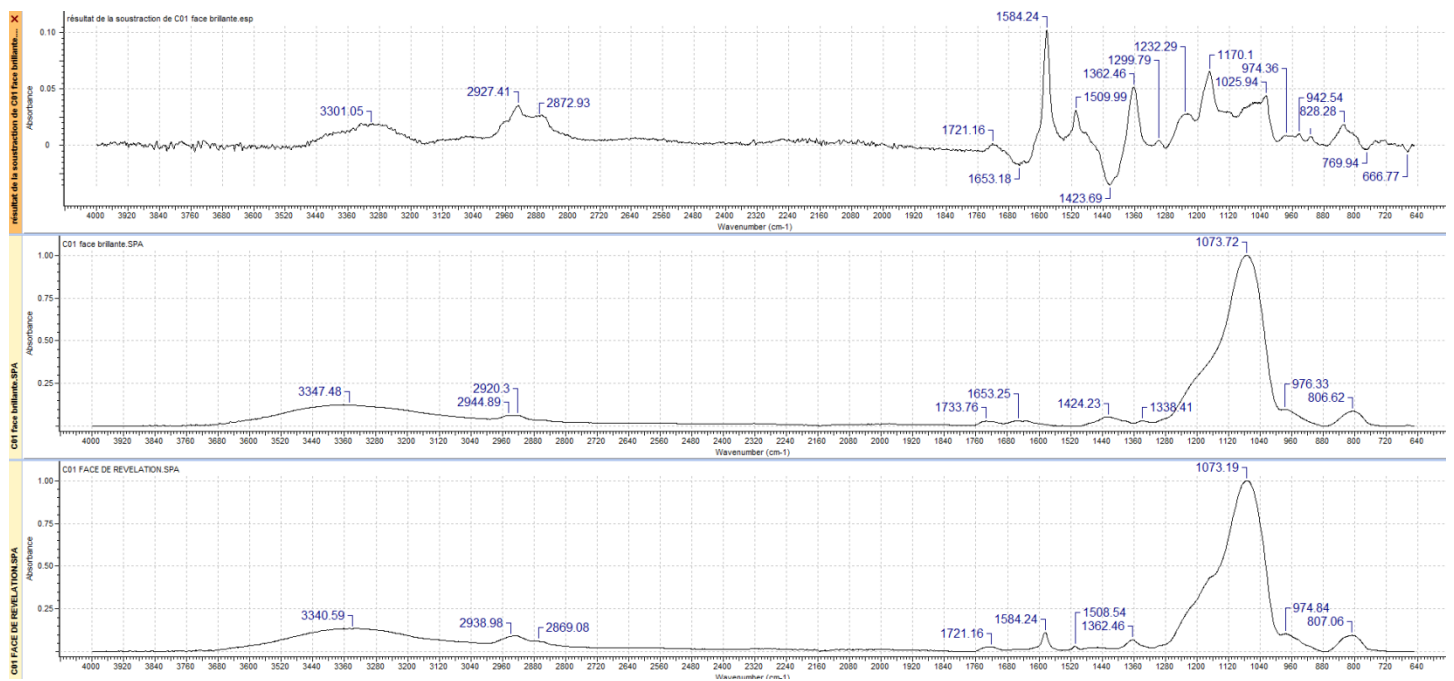


Figure 9 Top spectrum obtained by subtracting the IR spectra obtained before (middle) and after (bottom) the application of SMD II (paper sample: Kirkland Signature Mexico; C01).

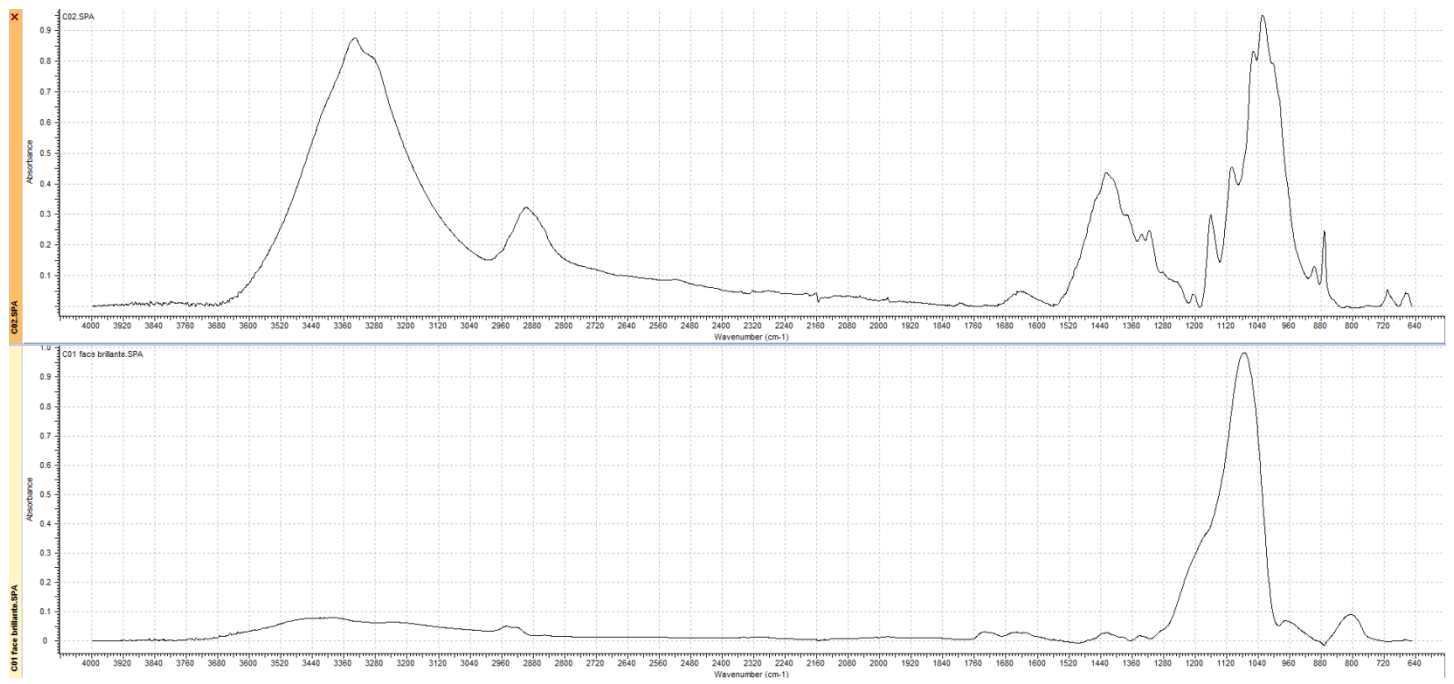


Figure 10 Difference spectra between C04 (top) and original C01 (bottom).

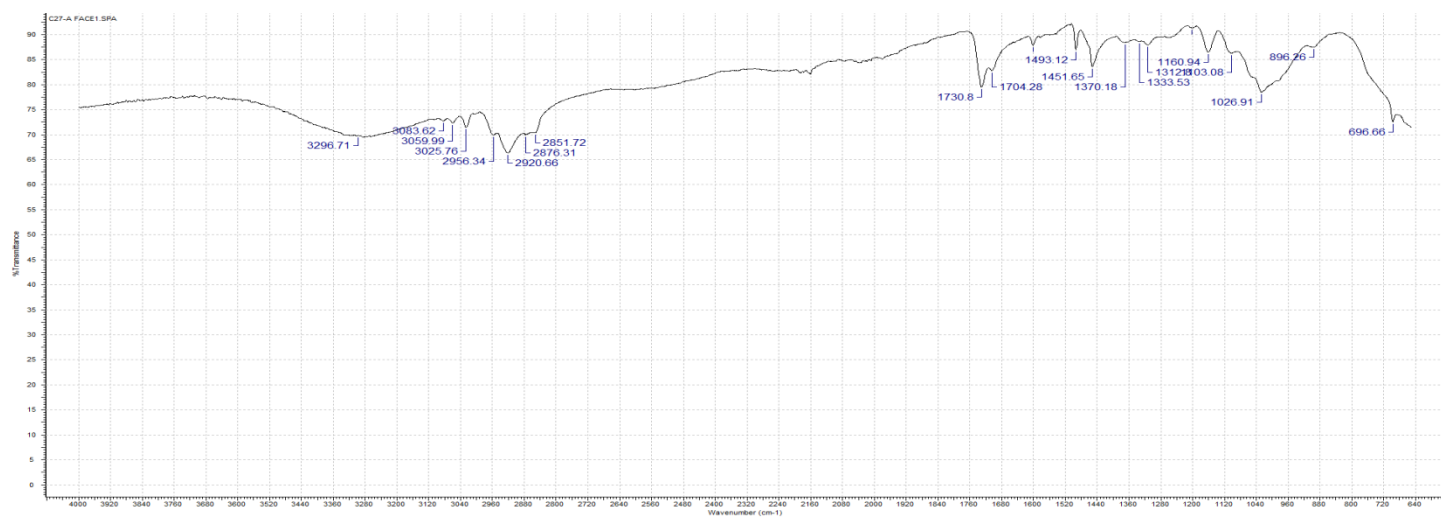


Figure 11 Infrared spectrum of the unprocessed Staples "Chemise à pochettes – 1336" paper sample (C27).

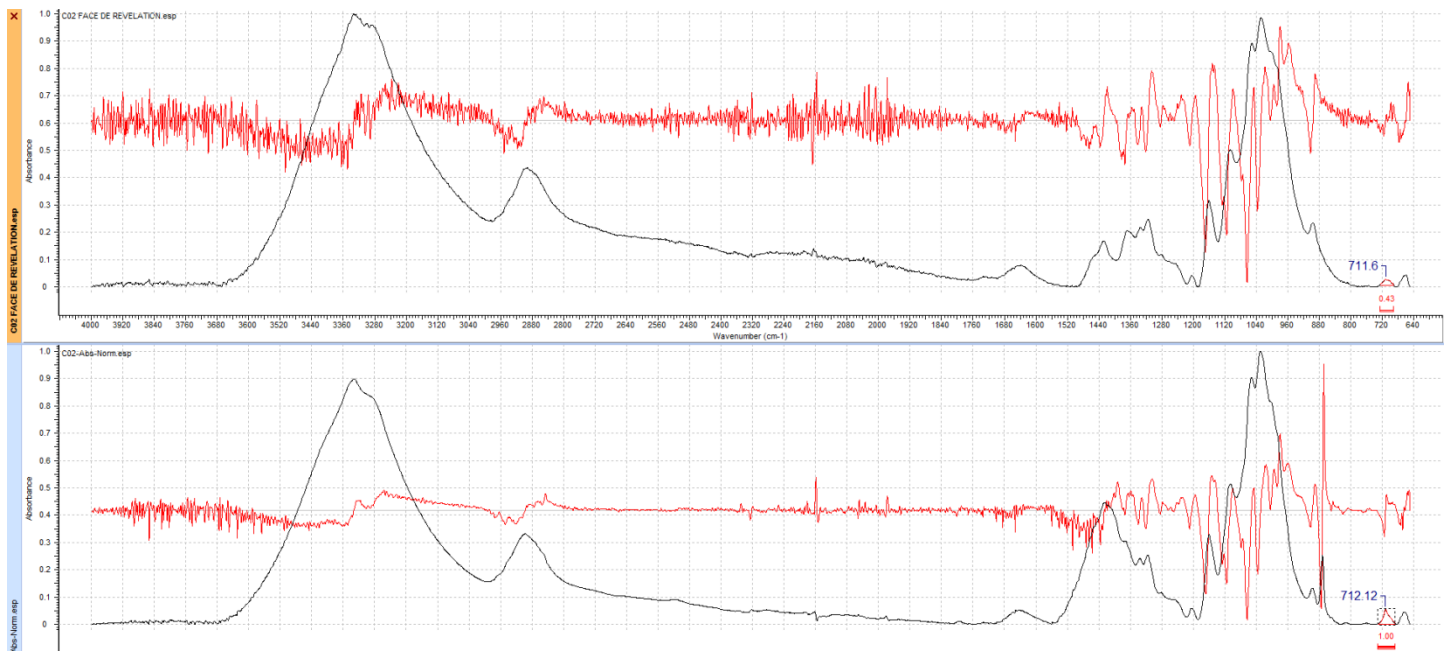


Figure 12 Derivative calculation for the RetroPlus50 paper sample (Canada; C02). Top spectra represent the paper surface with the fingermarks revealed, while bottom spectra represent the opposite surface of the same paper.

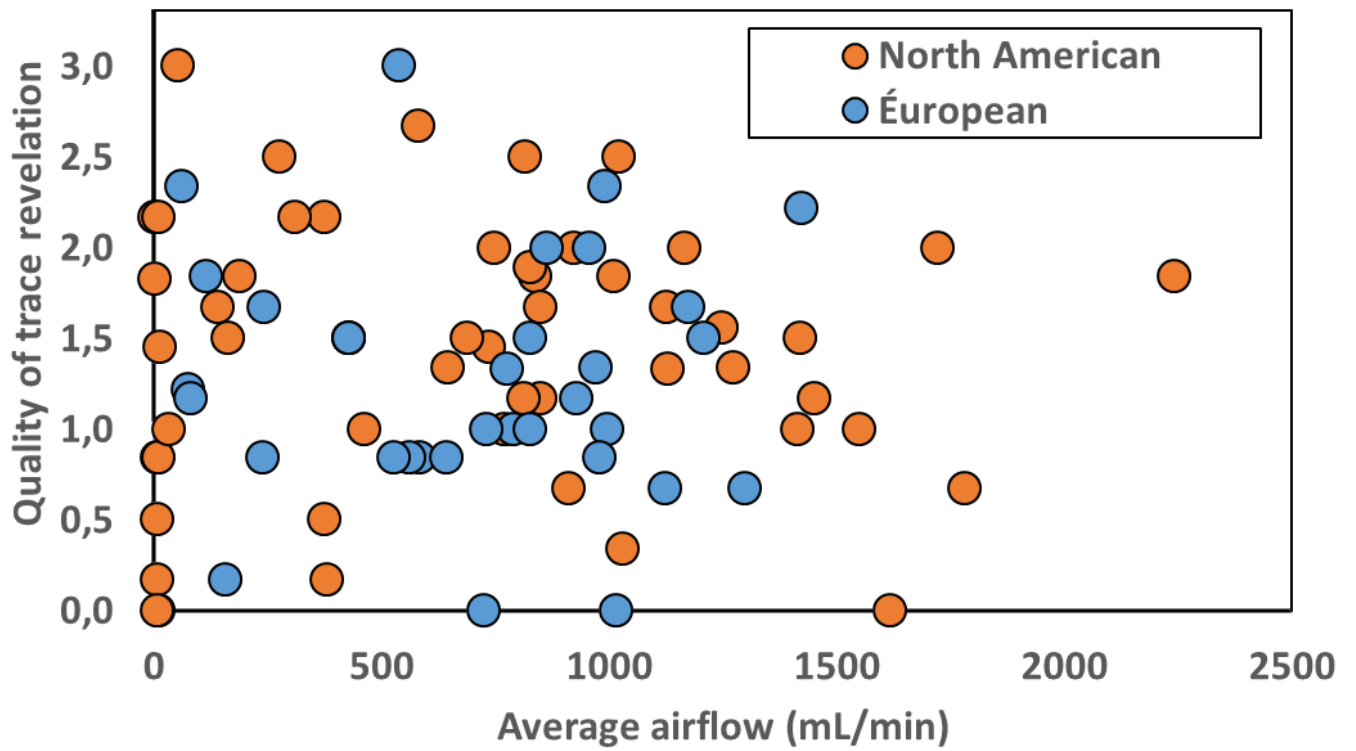


Figure 13 Chart illustrating the relation between the airflow (mL/min) and the average quality score associated with the fingermarks obtained after SMD II. Each dot represents a paper sample.

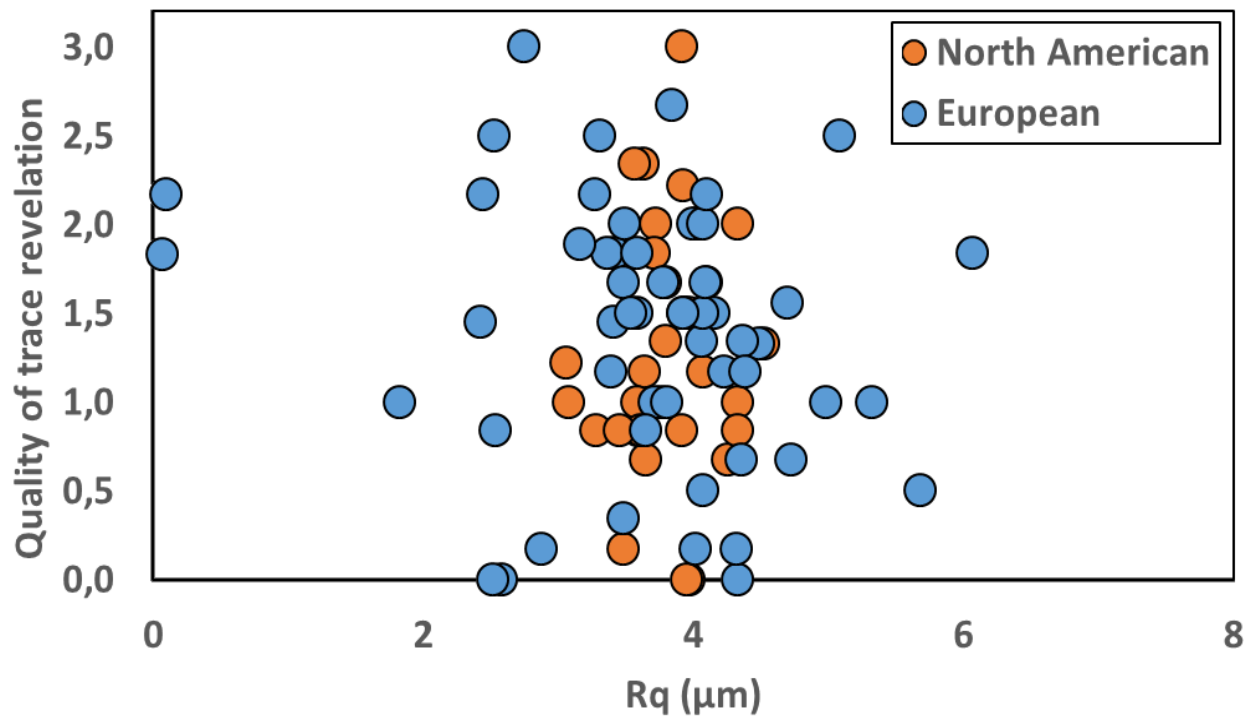


Figure 4 Chart illustrating the relation between the Rq values (microns) and the average quality score associated with the fingermarks obtained after SMD II. Each dot represents a paper sample.

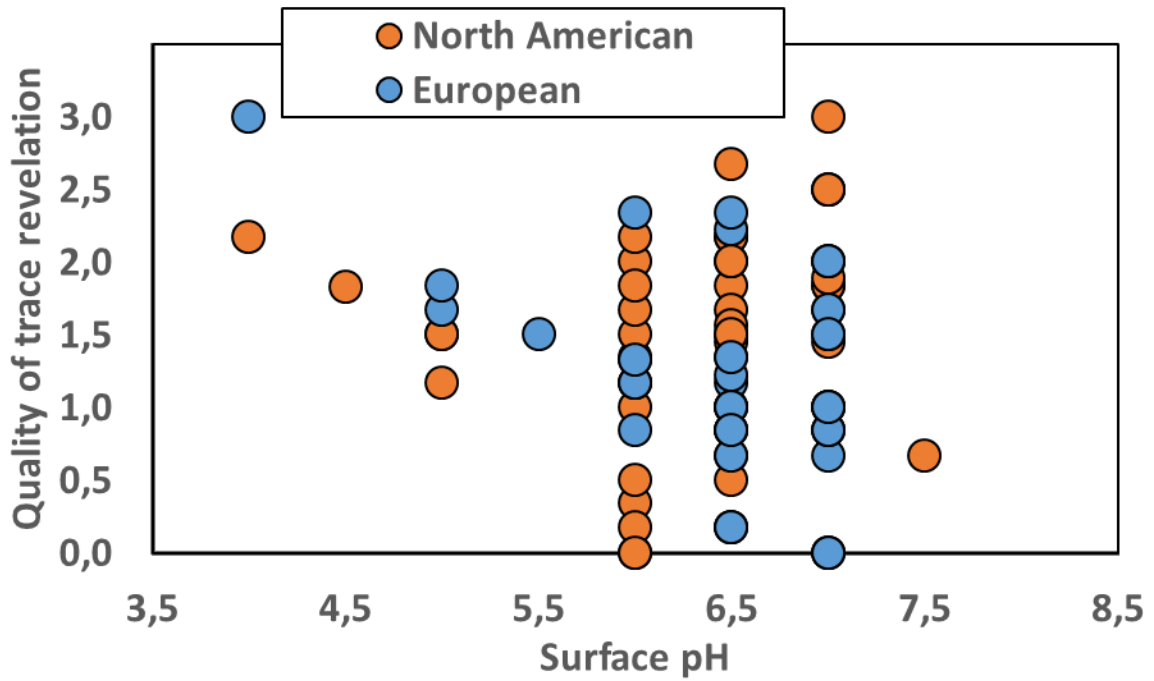


Figure 5 Chart illustrating the relation between the surface pH and the average quality score associated with the fingerprints obtained after SMD II. Each dot represents a paper sample.

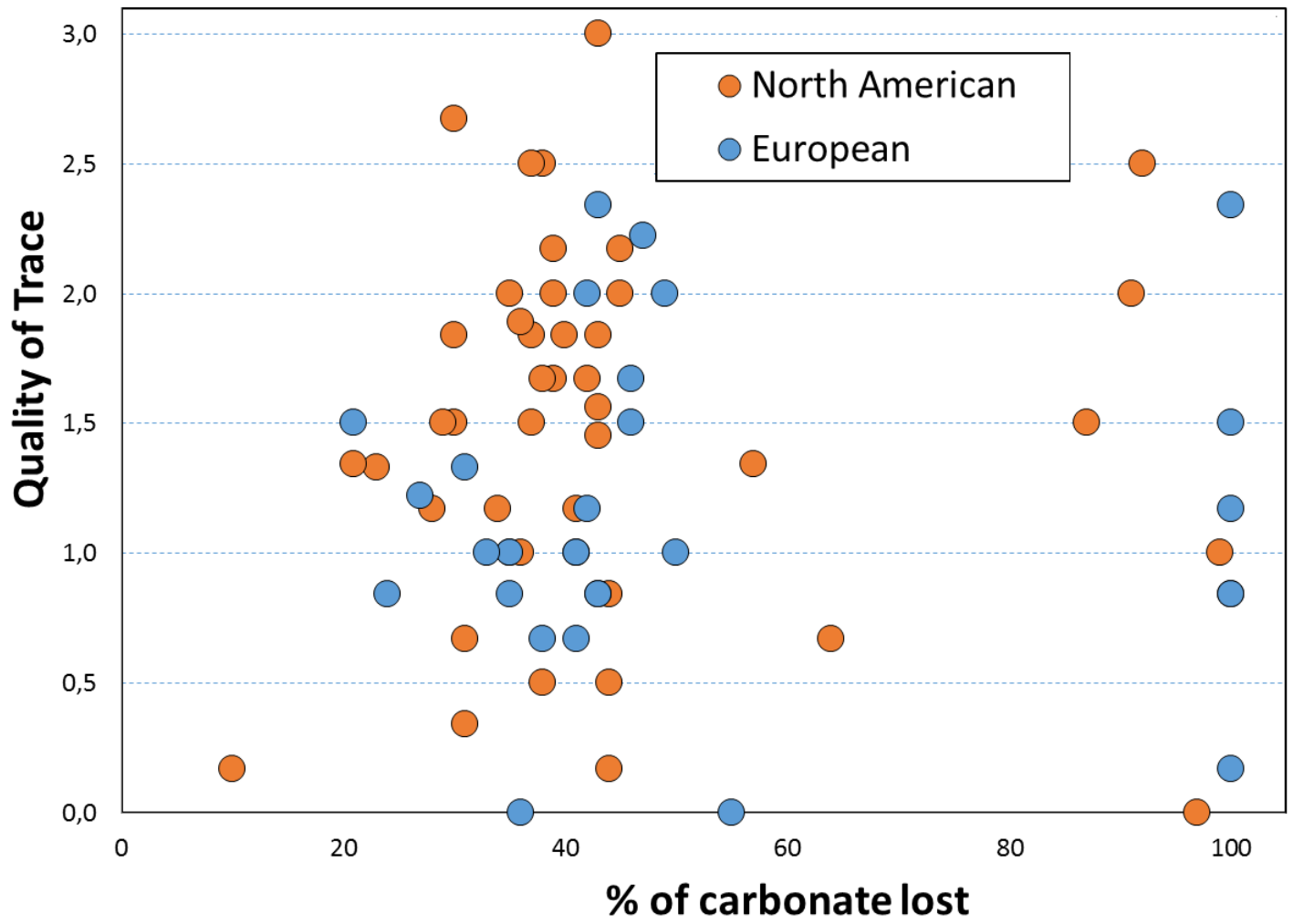


Figure 16 Chart illustrating the relation between the calcium carbonate loss (estimated %) and the average quality score associated with the fingermarks obtained after SMD II. Each dot represents a paper sample.

AUTHOR DECLARATION

We wish to confirm that there are no known conflicts of interest associated with this publication and there has been no significant financial support for this work that could have influenced its outcome.

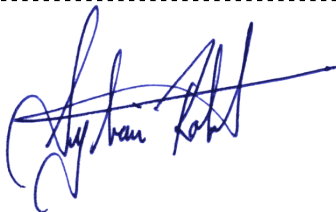

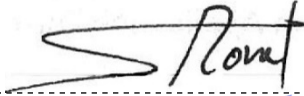

We confirm that the manuscript has been read and approved by all named authors and that there are no other persons who satisfied the criteria for authorship but are not listed. We further confirm that the order of authors listed in the manuscript has been approved by all of us.

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Frank CRISPINO	July 4, 2018	
Sébastien MORET	July 7, 2018	
Andy BÉBUE	July 12, 2018	

In case more space is needed, please attach a second page.

Appendix A: Paper samples

North American papers (C) and European papers (E). Laser (L), inkjet (I), Offset (O), copier/printer (P), utility (U). Recycled (R) with percentage when available, Forest Stewardship Council approved (FSC). Whiteness (%) and color also indicated. Basis weight units: g.m⁻². Some paper types, as well as basis weight, were not indicated by the suppliers.

Code	Origin	Company/Brand	Specification	Type	Use	Color	Basis weight	Surface pH
C01	Mexico	Kirkland	Signature - Professionnel Brillant	-	I	White	255	4
C02	Canada	ReproPlus50	-	R 50%	I and L	White	75	8
C03	USA	Staples	Pastel	R 30% - Acid free	P	Golden rofd, canary yellow, pink, blue, green	75	7
C04	Canada	Domtar	EarthChoice - Office paper	Mixed	P	White 92%	75	8
C05	USA	Staples	Sustainable Earth - Copy paper	95% sugar cane - Alkaline	P	White 92%	75	7
C06	USA	hp	Multipurpose paper	Mixed	P	White 96%	75	7
C07	Canada	Prairie Pulp & Paper Inc.	Step Forward Paper	80% corn / 20% FSC	P	White 92%	80	7
C08	USA	Neenah Paper Inc.	Astrobrights - Creative Expression	Acid free, lignin free	L, I, O, P	Solar Yellow, Vulcan Green, Cosmic Orange, Lunar Blue, Rocket Red	89	7
C09	USA	hp	LaserJet Paper	-	L, I, O, P	White 98%	90	7
C10	USA	Neenah Paper Inc.	Southworth - Parchemin	Mixed FSC	I and L	Ivory	90	7
C11	USA	Neenah Paper Inc.	Southworth - Granité	Mixed FSC	I and L	Ivory	90	7
C12	USA	Neenah Paper Inc.	Southworth - Lin	Mixed FSC	I and L	White	90	7
C13	USA	Neenah Paper Inc.	Southworth -Vergé antique	Mixed FSC	I and L	White	90	7
C14	Canada	Domtar	Business - Papier à lettre	R 20%	L, I, P	Venice	90	7
C15	Canada	Domtar	Business - Papier à lettre	R 20%	L, I, P	Windsor Marble	90	7

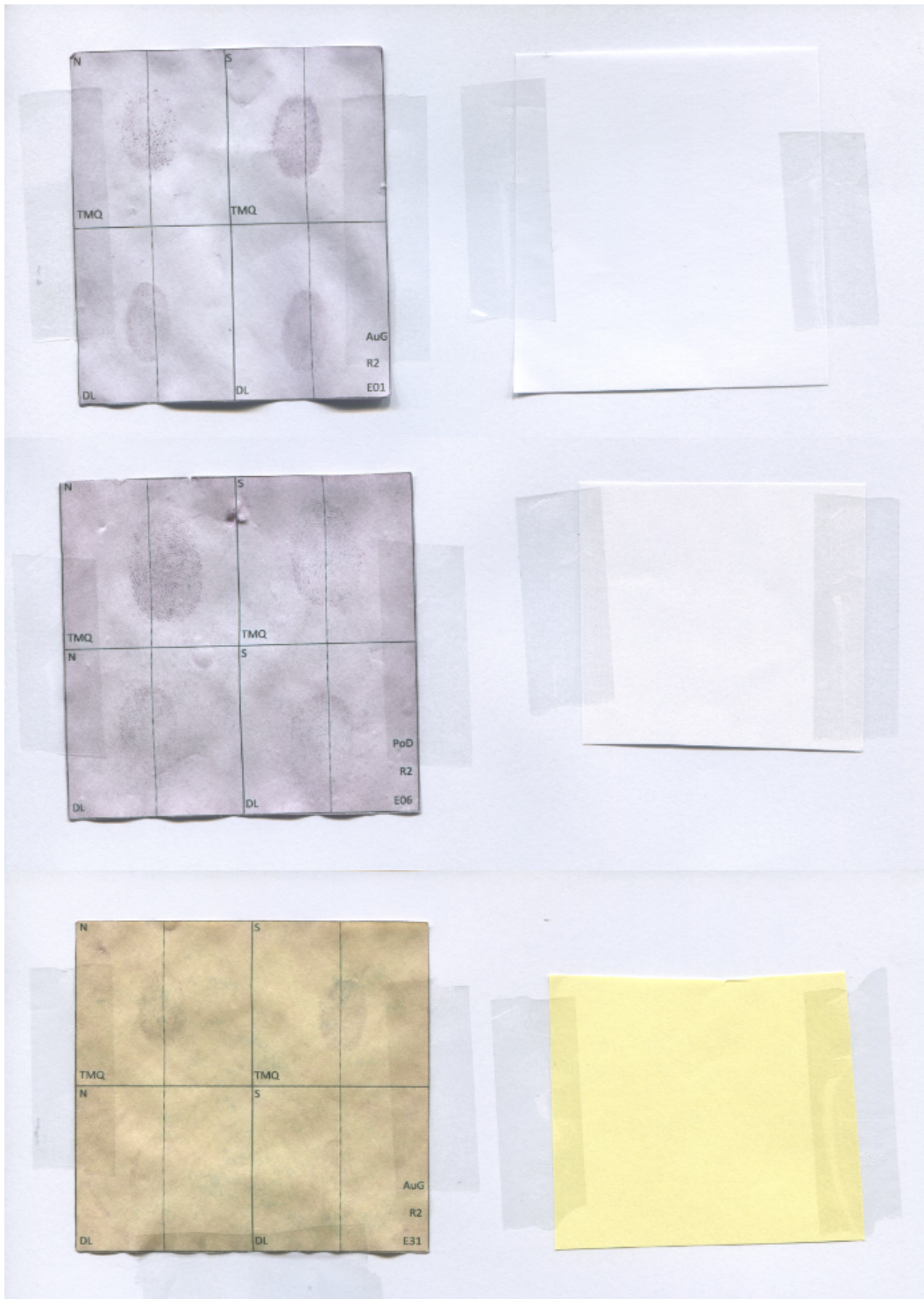
Code	Origin	Company/Brand	Specification	Type	Use	Color	Basis weight	Surface pH
C16	USA	hp	Premium Choice - Laser Paper	-	L	White 98%	120	7
C17	USA	Canon	Matte Photo Paper - MP-101	-	I	White	170	7
C18	USA	Staples	Papier à cartes	Acid and lignin free	L, I, P	Beige	250	7
C19	USA	Neenah Paper Inc.	Southworth - Parchemin (couverture)	Mixed FSC	I, L	Ivory	250	7
C20	USA	Canon	Photo paper Plus Glossy II - PP201	-	I	White	275	5
C21	USA	Staples	Tablette de papier perforé quadrillé - 22833	-	U	White	n/a	7
C22	Canada	Avery	Étiquettes CD - 5692	-	L		n/a	7
C23	Canada	Avery	Étiquettes Polyvalentes Amovibles - 6476	-	I, L	Magenta, yellow, green, blue	n/a	7
C24	USA	Staples	Protège-feuilles transparents - 10523	PVC, acid and lignin free	U	-	n/a	7
C25	Canada	Carbon Products Limited	Form-Mate	-	U	-	n/a	7
C26	Canada	Avery	Étiquettes d'adresse transparentes - 15662	-	L	-	n/a	7
C27	USA	Staples	Chemise à pochettes - 1336	Cardboard	U	-	n/a	7
C28	USA	Staples	Enveloppes d'expédition à bulles 6x9 - 17723	.	U	-	n/a	7
C29	USA	Staples	Enveloppes 5 7/8 x 9 5/8 - 594487	Mixed FSC	U	White	90	7
C30	USA	Staples	Enveloppes 5 7/8 x 9 5/9 - 194490	Mixed FSC	U	White	90	7
C31	USA	Staples	Sustainable Earth - Tablettes d'écriture	80% sugar cane residues	U	White	n/a	7
C32	Canada	3M	Post-it	R	U	Pink, green, blue	n/a	7
C33	Canada	3M	Post-it	R	U	Yellow	n/a	7
C34	USA	Staples	Étiquettes d'expédition	R Cardboard	U	Yellow	n/a	6
C35	USA	Staples	Fiches lignées - 90106	R 10% Cardboard	U	White	n/a	7
C36	USA	Staples	Tablette à réglage étroit - 66186	-	U	White	n/a	7
C37	USA	Staples	Enveloppes no1 à monnaie - 530164	FSC	U	White	105	7
C38	USA	Staples	Enveloppes no10 QuickStrip - 66396	-	U	White	75	7

Code	Origin	Company/Brand	Specification	Type	Use	Color	Basis weight	Surface pH
C39	USA	Staples	Enveloppes no10 QuickStrip - 66397	-	U	Security tint	75	7
C40	Canada	UQTR	Buvar	Bleached Kraft	U	White	n/a	6
C41	Canada	UQTR	SW,HW, Eucalyptus	-	U	White	n/a	4
C42	Canada	UQTR	Kraft couché	Kraft	P	White	n/a	7
E01	Europe	Auchan	-	-	P	White	90	7
E02	Austria	Canon Black Label zero	-	-	P	White	80	7
E03	France	Carrefour	-	-	P	White	80	7
E04	Austria	Coop	-	-	U	White	80	7
E05	Switzerland	StaplesElco switzerland Prestige	-	-	U	White	80	7
E06	France	Hp Home & Office	-	-	P	White	80	7
E07	Portugal	Inacopia Office	-	-	P	White	75	7
E08	Portugal	Inacopia Office	-	-	P	White	80	7
E09	Switzerland	Mbudget	-	-	U	White	70	7
E10	Europe	Mbudget	-	-	P	White	80	7
E11	Austria	Office	-	-	P	White	80	7
E12	Switzerland	Paetria Migos	-	-	U	White	100	7
E13	Sweden	Paetria Migos	-	-	P	White	80	7
E14	France	Paetria Premium Migos	-	-	P	White	160	7
E15	Europe	Papyrus Piano Speed	-	-	P	White	80	7
E16	Europe	Paper Team	-	-	P	White	80	7
E17	Austria	Prix Garantie Coop	-	-	U	White	70	7
E18	Germany	Sigel Office Paper Premium	-	-	P	White	80	7
E19	France	Xerox	-	-	P	White	90	7
E20	Finland	Xerox Business	-	-	P	White	80	7
E21	France	Esquisse	Envelope	R	U	Grey	80	6

Code	Origin	Company/Brand	Specification	Type	Use	Color	Basis weight	Surface pH
E22	France	La Couronne	Envelope	R	U	Brown	90	7
E23	Austria	Oecoplan coop	-	R	U	Beige	80	6
E24	France	Papeteria	-	R	P	Grey	80	6
E25	Germany	Paper Union Inapa	-	R	P	Grey	80	7
E26	Switzerland	Raygan Aligro	-	R	P	Beige	n/a	6
E27	France	Paetria	-	-	P	Pink	80	7
E28	France	Paetria	-	-	P	Yellow	80	7
E29	France	Paetria	-	-	P	Green	80	7
E30	Europe	Papyrus Rainbow	-	-	P	Pink	80	7
E31	Europe	Papyrus Rainbow	-	-	P	Yellow	80	7
E32	Europe	Papyrus Rainbow	-	-	P	Green	80	7

Appendix B: Samples of fingermarks revelations using SMDII

Four fingerprint traces were placed on each paper sample. In the upper part are traces of the first donor, in the lower part, the traces of the second donor. In the left part, the traces are natural. In the right part, the traces are loaded (passage of the fingers on the forehead and the neck to increase the quantity of secretion of the trace). Original samples are on the right, and post-revelation samples are on the left.



Appendix C: Rq pre and post-revelation of papers

Samples	Rq pre-revelation	Rq post-revelation	Difference
C01	0.097	0.968	0.870
C02	4.357	4.772	0.415
C03 Bleu	3.422	4.627	1.205
C03 Jaune Canari	3.479	5.647	2.168
C03 Rose	4.022	4.861	0.839
C03 Verge d'or	3.483	4.552	1.069
C03 Vert	4.489	5.654	1.165
C04	3.307	3.842	0.535
C05	3.366	4.901	1.535
C06	3.717	5.347	1.630
C07	3.779	4.791	1.012
C08 Cosmic Orange	4.370	5.015	0.645
C08 Lunar Blue	4.156	3.633	0.523
C08 Rocket Red	4.325	4.824	0.499
C08 Solar Yellow	3.808	4.341	0.533
C08 Vulcan Green	4.228	5.192	0.964
C09	3.407	4.186	0.779
C10	4.723	4.186	0.537
C11	4.063	5.934	1.871
C12	4.020	4.730	0.710
C13	4.983	5.884	0.901
C14	3.588	4.004	0.416
C15	4.000	4.149	0.149
C16	2.530	3.812	1.282
C17	3.582	5.879	2.297
C18	5.082	5.144	0.062
C19	5.323	6.498	1.175
C20	0.068	0.402	0.334
C21	4.076	3.507	0.569
C22	3.651	3.472	0.179
C23 Bleu	2.517	3.827	1.310
C23 Jaune	2.876	3.177	0.301
C23 Magenta	2.582	3.637	1.055
C23 Vert	2.538	2.974	0.436
C27 Beige	2.429	3.074	0.645
C27 Bleu	1.825	2.825	1.000
C27 Rouge	2.440	2.759	0.319
C28	5.685	6.245	0.560
C29	3.163	4.728	1.565
C30	4.069	6.450	2.381
C31	2.748	3.110	0.362
C32 Bleu	3.391	4.769	1.378

Samples	Rq pre-revelation	Rq post-revelation	Difference
C32 Rose	4.387	4.483	0.096
C32 Vert	4.089	4.086	0.003
C33	4.074	3.933	0.141
C34	4.098	4.694	0.596
C35	4.696	4.875	0.179
C36	3.274	3.556	0.282
C37	3.842	3.395	0.447
C38	3.492	4.269	0.777
C39	3.923	5.342	1.419
C40	6.069	6.105	0.036
C41	3.534	3.877	0.343
C42	4.317	4.971	0.654
E1	4.331	3.688	0.643
E10	3.970	4.069	0.099
E11	3.649	4.414	0.765
E12	3.756	3.845	0.089
E13	4.254	3.998	0.256
E14	3.059	4.447	1.388
E15	3.720	4.955	1.235
E16	3.911	4.332	0.421
E17	3.615	4.078	0.463
E18	4.329	4.173	0.156
E19	3.080	3.577	0.497
E2	3.641	4.003	0.362
E20	3.956	3.488	0.468
E21	4.075	4.311	0.236
E22	3.480	4.046	0.566
E23	3.451	4.959	1.508
E24	3.968	4.427	0.459
E25	4.328	3.281	1.047
E26	3.633	4.056	0.423
E27	4.099	4.036	0.063
E28	3.911	3.350	0.561
E29	3.711	4.351	0.640
E3	3.941	4.573	0.632
E30	4.519	4.160	0.359
E31	3.792	4.160	0.368
E32	3.570	3.825	0.255
E4	3.589	3.932	0.343
E5	3.284	3.274	0.010
E6	3.920	3.899	0.021
E7	3.795	3.900	0.105
E8	3.927	3.883	0.044
E9	3.584	4.536	0.952

