

1 **Aquazol as a binder for retouching paints. An evaluation through analytical pyrolysis and**
2 **thermal analysis**

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12 **Abstract**

13 Aquazol poly (2-ethyl-oxazoline) is a tertiary aliphatic amide, with physical and chemical properties
14 that are exploited in a variety of ways, from pharmaceutical applications to the conservation of
15 cultural heritage. In this study, we evaluated the use of Aquazol as a new binder for retouching paint
16 in the restoration of artworks.

17 Aquazol 500 admixed with various formulations of organic red pigments was used to prepare paint
18 replicas which were artificially aged and investigated by a multi-analytical approach based on
19 analytical pyrolysis coupled with gas chromatography and mass spectrometry (Py-GC/MS), and
20 thermogravimetry (TG), complemented by FTIR and LIBS spectroscopy. This is the first study on
21 the ageing phenomena of Aquazol 500 using analytical pyrolysis and thermogravimetric analysis.
22 The influence of the pigments' components on the pyrolysis behavior of Aquazol was also
23 investigated.

24 The paint replicas did not show significant modifications during artificial ageing. This thus highlights
25 the optimal properties of Aquazol 500 as a binder for retouching, in addition to its already established
26 suitability as a filler or consolidant in the restoration of artifacts. Interestingly, when Aquazol 500 is
27 used in formulations containing organic pigments, Aquazol-pigment interactions are observed,
28 strongly depending on the pigment used.

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30 **Keywords**

31 Aquazol, Retouch paint, Organic pigments, Analytical pyrolysis coupled with gas chromatography
32 and mass spectrometry (Py-GC/MS), Thermogravimetry (TG).

34 **1. Introduction**

35 The technology transfer from the polymer industry to conservators and conservation scientists is of
36 paramount importance in the conservation of artworks. The conservation of paintings is a delicate
37 field of material science, where tailored solutions based on advances in polymer science and on new
38 commercial synthetic macromolecules can be exploited to formulate innovative consolidants, binders,
39 varnishes or coatings. It is thus important to assess whether the newly-developed material has
40 improved features compared to those available in terms of stability, compatibility and reversibility.
41 This study evaluates the use of polymer poly (2-ethyl-oxazoline) as a new binder for retouching paint
42 in the restoration of paintings.

43 Poly (2-ethyl-oxazoline) or PEOX was first marketed in 1977 with the commercial name Aquazol,
44 by the Dow Chemical Company [1]. It is now produced by Polymer Chemistry Innovations in four
45 ranges of molecular weight: 5000 g/mol; 50000 g/mol; 200000 g/mol; and 500000 g/mol (Aquazol®
46 5, 50, 200 and 500 respectively) [2]. Aquazol is a tertiary aliphatic amide, with physical and chemical
47 properties, which are now being exploited for an increasing variety of ways, from pharmaceutical
48 applications to the conservation of the cultural heritage [3], [4], [5], [6], [7], [8], [9], [10], [11].

49 Aquazol's physico-chemical properties were first described in 1986 by T.T Chiu et al. [12], in terms
50 of the composition, polymerization mechanism and properties. The wide range of solubilities in
51 several polar and medium polar solvents including water is one of its key features in several
52 applications. This makes Aquazol a versatile material for use in the cultural heritage, since if
53 necessary it can be removed after application thus ensuring a good reversibility of the conservation
54 treatments.

55 Aquazol was proposed as a consolidant for the first time by R. Wolbers et al. in the early 1990s [5].
56 The properties of the polymer have subsequently been investigated in terms of pH, colour and
57 viscosity measurements, tensile strength and re-solubilization tests, FTIR spectroscopy, size-
58 exclusion chromatography (SEC) and thermogravimetry (TG), thus suggesting a good optical and
59 chemical stability of the material [12]. Stress/strain curves generated as a function of relative humidity
60 (RH) variations have shown that Aquazol films maintain a good elasticity when exposed to different
61 RH conditions. In addition, Aquazol is a green material [3] and is compatible with other polymers to
62 obtain mixtures with modulated properties. The few analytical studies described in the literature
63 include its characterization by pyrolysis coupled with gas-chromatography (Py-GC/MS) [13], which
64 have been shown to be suitable in investigating its molecular structure.

65 Conservators have tested and applied Aquazol as a weak adhesive [9], [14], [15], as a consolidant and
66 filler [8], [16], [17], [18], [19], [20], and as an ingredient in the preparation of bole for the application
67 of gilding [21]. However, to date it has only been tested on a few painted surfaces [22]. Several

68 potential applications of Aquazol as a conservation material are currently under investigation, such
69 as its use as a consolidant for unvarnished painted surfaces, in combination with TiO₂ nanoparticles
70 [23].

71 The properties described above along with the data available in the literature also suggest that
72 Aquazol is a suitable candidate for use as a binder for reversible paints for retouching during the
73 conservation of paintings. Retouching a painting is one of the most common and critical conservation
74 practices, where the reversibility of the applied retouching materials is key in guaranteeing the ethical
75 acceptability of the intervention, which entails removal and re-treatment. The most common
76 retouching paints are today based on Arabic gum as a paint binder, and thus are not suitable for water-
77 sensitive painted surfaces.

78 Here we present the results of a study on the stability of Aquazol-based paints for use as a retouching
79 paint alternative to Arabic gum-based paints. Aquazol 500 admixed with organic red pigments in
80 different formulations was used to prepare artificially aged paint layers. The paint replicas were
81 investigated by a multi-analytical approach based on analytical pyrolysis coupled with gas
82 chromatography and mass spectrometry (Py-GC/MS), and thermogravimetry (TG), complemented
83 by FTIR and LIBS spectroscopy.

84 To the best of our knowledge, this is the first ageing study of Aquazol 500 using analytical pyrolysis
85 and thermogravimetric analysis. These techniques have been successfully used to investigate
86 synthetic polymers in paint samples at a molecular level [24], [25], [26], [27], [28]. They yield
87 information on the thermal stability of the binder and the formation of cross-linking bonds within the
88 polymer during ageing [29], [30], [31], [32], [33]. They also highlight the presence of pigment/binder
89 interactions [34] and the formation of pyrolytic products related to degradation [35], [36], [37], [38],
90 [39].

91 Two different recipes for Aquazol-based retouching paint were prepared and the raw materials and
92 paint replicas were analyzed before and after artificial ageing in Solarbox, in order to assess their
93 stability over time. Paint layers were prepared with three commercial organic pigments, namely
94 Irgazin red (diketo-pyrrolo-pyrrole, DPP), Alizarin (synthetic anthraquinone-based pigment lake),
95 and Quindo pink (quinacridone), in order to investigate the influence of the pigments on the thermal
96 features of the material during curing and ageing. An analysis of the pure materials used to prepare
97 the paints was performed beforehand.

98 The influence of the pigments' components on the pyrolysis behavior of Aquazol was also
99 investigated.

100

101 **2. Materials and methods**

102 2.1. Paint samples and artificial ageing

103 Eight paint layers with two different recipes for the binding medium and three different organic
104 pigments (DPP, alizarin and quinacridone) were prepared on glass slides using Aquazol 500 as a
105 binder. The set also included layers of the binder without pigments. Proportions and amounts of
106 materials used for each paint preparation were chosen according to the instructions provided by
107 Professor Richard Wolbers [40].

108 The materials for the paint preparations were: Aquazol 500 (Nr. 63905 Kremer Pigmente), diketo-
109 pyrrolo-pyrrole (DPP, “Irgazine Ruby DPP-TR”, Nr. 23182 Kremer Pigmente, CI PR264), Alizarin
110 (“Alizarine Crimson Dark”, Nr. 3610 Kremer Pigmente, CI PR83), Quinacridone (“Quindo Pink D”,
111 Nr.23402 Kremer Pigmente, CI PV19), xanthan gum (La Saponaria, Pesaro, Italy, used in recipe A),
112 glycerol (86%, Nr.64900 Kremer Pigmente, used in recipe A), 2-phenoxyethanol (99%, Nr. 77699
113 Sigma-Aldrich).

114 The choice of the binder/pigment ratio was based on CPVC (Critical Pigment Volume Concentration)
115 parameters indicating the optimal concentration of pigment to add to the binder, on the basis of the
116 physical and morphological features of the pigment powder particles: large pigment particles need
117 less binder than the smaller ones [41].

118 DPP and alizarin, which have a similar particle size, were prepared in paints with a 1:4 ratio of
119 pigment and fluid binder (33% w/w aqueous Aquazol solution). Quinacridone, which has a smaller
120 particle size, was prepared in paint with 15% in weight of pigment and 85% of fluid binder (40%
121 aqueous Aquazol solution).

122 Recipe A consisted of Aquazol 500 (binder), phenoxyethanol (preservative, 2–4 drops), pigment,
123 xanthan gum and glycerol (1–2 drops), while recipe B consisted in Aquazol 500, phenoxyethanol and
124 pigment only.

125 The paint layers were investigated before and after 777 h of artificial ageing in a Solar Box 3000E
126 model (CO.FO.ME.GRA., Milan, 20 × 42 cm) equipped with a single Xenon lamp (2500 W) air
127 cooled, prepared with a glass inner filter in soda-lime at 280 nm + IR. Inside the box the irradiance
128 was uniform and perpendicular to the sample surface, with a constant irradiance of 500 W/m². During
129 the ageing protocol, the maximum temperature reached by the black body temperature probe, placed
130 next to the samples, was 45 °C.

131 2.2. Pyrolysis-gas chromatography/mass spectrometry (Py-GC/MS)

132 Samples were analyzed using a multi-shot pyrolyzer PY-3030D (Frontier Lab) coupled with a GC/MS
133 system composed of a 6890 N gas chromatograph combined with a 5973 mass selective single
134 quadrupole mass spectrometer (Agilent Technologies). The samples (~0.2 mg each) were inserted in
135 platinum sample cups. The cups were placed on top of the pyrolyzer at ambient temperature and then
136 moved into the furnace. Pyrolysis conditions were optimized as follows: pyrolysis chamber
137 temperature 700 °C, interface Py-GC 300 °C. The GC injection port operated in split mode with a
138 split ratio of 1:30. The chromatographic separation of pyrolysis products was performed with a fused
139 silica capillary column HP-5MS (J&W Agilent Technologies, 5% diphenyl-95% dimethyl-
140 polysiloxane, 30 m × 0.25 mm i.d., 0.25 µm film thickness), preceded by 2 m of deactivated fused
141 silica pre-column with an internal diameter of 0.32 mm. Chromatographic conditions were as follows:
142 40 °C for 5 min, 10 °C/min up to 310 °C, 20 min isothermal. Helium gas flow was set in constant
143 flow mode at 1.2 mL/min (purity 99.9995%). MS parameters were: electron ionization (EI, 70 eV) in
144 positive mode; ion source temperature 230 °C; scan range 50–700 m/z; interface temperature 280 °C.
145 Per-fluorotributylamine (PFTBA) was used to tune the mass spectrometer. MSD ChemStation
146 (Agilent Technologies) software was used for the data analysis, and peak assignment was based on a
147 comparison with mass spectra libraries (NIST 2.0, WILEY275), mass spectra reported in the
148 literature, and mass spectra interpretation.

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150 2.3 Attenuated total reflectance-Fourier Transform-Infrared spectrometer (ATR-FT-IR)

151 Infrared spectra were recorded using a FT-IR Agilent Technologies Spectrometer Cary 640, equipped
152 with a universal attenuated total reflectance accessory (ATRU). A Few micrograms of powdered
153 samples were analyzed with the following spectrometer parameters; resolution: 4 cm⁻¹, spectral range:
154 500-4000 cm⁻¹, number of scans: 16. Spectrum software was used to process FTIR spectra.

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156 2.4 TGA

157 A TA Instruments Thermobalance model Q5000 was used. TG measurements on ~1 mg of each
158 sample were performed at 10 °C/min scan rate, from 30 °C to 800 °C under N₂ flow (25 mL/min).

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160 2.5 LIBS

161 LIBS analyses were performed with a double pulse technique, using a Nd:YAG laser operating on
162 fundamental wavelength (1064nm). The inter-pulse delay between the two pulses was 1 µs, while
163 each laser pulse width was 15 ns. The energy released was about 60 mJ. The plasma signal was
164 collected using the double reticulum spectrometer Avantes (Stellarnet. Inc), in the range 200-900 nm
165 with an acquisition gate of 2.48 ms with a 2µs delay with respect to the first laser shot.

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2.6 PCA Analysis

Principal component analysis of the data (PCA) of the covariance matrix was performed with the software Xlstat10.0 (Addinsoft, France).

3. Results and discussion

3.1 Characterization of the pure materials

3.1.1 Aquazol 500 and additives

The TG curve of pure solid Aquazol 500, shown in Fig. 1A, shows a mass loss of about 5.9% in the temperature range 30–130 °C, typical of adsorbed water. At 434 °C (maximum of DTG curve) the polymer exhibits a main degradation step corresponding to a mass loss of about 93.5%, with a residue at 800 °C of about 0.6%. By rescaling the curve, excluding the water loss, the main degradation step corresponds to a 99.4% weight loss, indicating that the thermal degradation of Aquazol 500 occurs in a single step and that the pyrolytic process leads to volatile substances. Fig. 1B and C shows the results obtained with films made from the Aquazol-based binder prepared according to recipes A and B without pigments. The two profiles are very similar, and in both cases, when the water contribution is subtracted, a single degradation step is observed at 414 °C and 423 °C for recipes A and B, respectively. These values are slightly lower than those observed for the degradation of pure Aquazol analyzed in bulk, and correspond to a 99+% mass loss. No significant changes were observed when the TG experiments were performed on the corresponding artificially aged samples (Fig. 1D). These observations point out that reticulation and crosslinking phenomena do not occur during artificial ageing, and support the possibility of a good reversibility of Aquazol retouchings. Further solubilization experiments and cleaning tests are needed to better investigate the degree of reversibility of Aquazol paint applications.

Although glycerin and Xanthane gum are present in the recipe A formulation in a very low content, the thermal degradation of these two additives was also taken into account. Concerning glycerin, its TG curve in N₂, reported in the literature [42], exhibits a single narrow degradation step in the range 177–231 °C with a maximum in the derivative curve at 211 °C, thus corresponding to a 98% mass loss. The TG curve of Xanthane gum was acquired and shows, at quite low temperatures (50 °C), the evolution of absorbed water (corresponding to a mass loss of about 13.7%) followed by the main degradation step (48.5% of mass loss in the temperature range 200–400 °C). At 400 °C, 47.8% of sample mass is still present after which the degradation proceeds smoothly. At 900 °C, a residue of about 26.5% in mass is found. As expected considering their low amount in the recipes, the degradation of glycerin and Xanthane was not detected in the TG curves of the Aquazol recipe A paint layers.

201 The thermal degradation of the samples was investigated at a molecular level by GC/MS analysis of
202 the pyrolysis products, providing additional information on the pyrolytic behavior. The pyrolytic
203 degradation of Aquazol 500 is reported in [13], and is characterized by an initial chain scission by
204 cleavage of the C-N bond followed by two degradation pathways: one involving the cleavage of
205 another of the C-N bonds followed by a cyclization reaction producing ethyl-oxazoline, and the other
206 involving the cleavage of the C-C bond next to the nitrogen atom ($\alpha\beta$) followed by a McLafferty
207 rearrangement reaction.

208 Fig. 2A shows the pyrogram obtained in the Py-GC/MS analysis of a sample from the film of the
209 Aquazol binder prepared according to recipe B, without any pigment. Four clusters of thermal
210 degradation products produced in the flash pyrolysis at 700 °C can be identified at different retention
211 times [13], and the list of the identified pyrolysis products is reported in Table 1. The first cluster of
212 signals (1–4 min) is characterized by low molecular weight molecules, including isocyanates, nitriles,
213 amines, alkenes and ketones (LMW). The second cluster (7–14 min) is more heterogeneous and, at
214 this range of retention times, propionamide (P in the table) with related alkylated structures, and
215 monomers (M) and sesquimers with related alkylated structures are eluted. In this region, the most
216 abundant peaks correspond to the sesquimers. The dimers (D), with the related alkylated structures,
217 are eluted in retention time range of between 14 and 18 min. The last cluster, between 18 and 20 min,
218 features the presence of trimers (T) and of the related alkylated structures. Fig. 2B reports the Py-
219 GC/MS chromatographic profile of the Aquazol 500 paint layer (recipe B without pigment) after the
220 artificially ageing treatment. No significant changes in the pyrolytic profile of the polymer or in the
221 mass spectra of the pyrolysis products are observed after ageing. No peak corresponding to new
222 compounds is observed, and the relative abundance of the various classes of compounds produced
223 during pyrolysis (low molecular weight molecules, monomers, sesquimers, dimers and trimers)
224 remain almost unchanged in the artificially aged material, compared to the unaged material. The Py-
225 GC/MS profiles observed before and after artificial ageing show no significant changes for the
226 investigated Aquazol paint layers prepared according to both recipes A and B. The different pigments
227 do not induce a variation of the pyrolysis behavior after photoageing.

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229 3.1.2 Pigments

230 The thermal behavior of the commercial pigments was also investigated through TG.

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232 The irgazine ruby pigment (PR264) belongs to the diketo-pyrrolo-pyrrole (DPP) dye family with two
233 biphenyl substituents in positions 3,6. Its TG curve (Fig. 3A) is relatively simple, only showing a
234 single step mass loss at 514 °C corresponding to about 84% in weight. However, the residue at 900

235 °C is about 16% in weight, indicating that an inorganic matrix is also present, probably introduced in
236 the formulation as a filler or complexing agent. This matches the results of LIBS analysis of DPP
237 where typical spectral lines of calcium, aluminum and magnesium are present.

238 Alizarin is a derivative of anthraquinone with two hydroxyl substituents in position 1,2. Alizarin
239 based pigments, such as Garanza lake (PR83), are obtained by the precipitation of its aluminum
240 complex, usually in the form of hydrated aluminate or hydrated silicoaluminate of calcium and/or
241 magnesium. The TG curve of Kremer's alizarin shows a complex profile due to the mixture of
242 kaolinite and alizarin dye (Fig. 4A): a residue percentage of 43% at 900 °C is observed, attributed to
243 the inorganic component. This is consistent with the IR spectrum of the alizarin pigment
244 (Supplementary data, Fig. S1), which corresponds to the overlap of the spectra of pure alizarin and
245 of hydrated silicoaluminate associated with mineral kaolinite [43]. The LIBS spectra of Alizarin lake
246 (Supplementary data, Fig. S2) shows the typical lines of silicon, calcium, magnesium, aluminum
247 (probably ascribed to the presence of kaolinite): it also highlights the typical lines of titanium,
248 revealing that a titanium-based filler is also present.

249 A marked dehydration is observed in the TG in the range $T_{\text{amb}} - 135$ °C, with two distinct and
250 reproducible DTG maxima at 71° and 133 °C. These can be assigned to the release of adsorbed water
251 (at 75 °C) and of crystallization water in the kaolinite pores (at 132 °C) [44], corresponding to an
252 overall mass loss of about 13%. Increasing the temperature, the degradation pattern of alizarin lake
253 shows a weak mass loss around 312 °C, followed by the main mass loss occurring at 487 °C, preceded
254 by a shoulder at 429 °C. The overall mass loss amounts to about 45%. Over 550 °C, the mass loss
255 continues smoothly and the residue at 900 °C is about 43.1%, which is consistent with the organic
256 and inorganic composition of the pigment. The subsequent mass losses at 200–700 °C are attributed
257 to the degradation of alizarin dye and to the reorganization in the octahedral layer of kaolinite. In the
258 temperature range 450–650 °C, the dehydroxylation of kaolinite and formation of metakaolinite takes
259 place, in line with the literature.

260 Among the pigments considered within this work, only the quinacridone “Pink Quindo D” does not
261 contain any inorganic component. Its TG curve (Fig. 5A) shows a sharp single degradation step at
262 554 °C corresponding to a 97+% mass loss. The ATR-FTIR spectrum of this pigment (Supplementary
263 data, Fig. S3) matches that of pure Quinacridone from Wiley Subscription Services, Inc. (US), thus
264 confirming that no inorganic complexing agent or additive has been employed to produce this
265 pigment.

266 3.2 Characterization of the paint layers based on Aquazol 500 and pigments

267 The TG analyses of the paint replicas obtained according to both recipes A and B show very
268 interesting features.

269 The TG curves of the Aquazol 500 paint layers containing DPP (Fig. 3B and C) exhibit a single
270 degradation step at 404 °C (recipe A) and at 407 °C (recipe B), corresponding to a mass loss of about
271 87%. The residue observed at 800 °C is about 11%, in agreement with the 1:1 DPP/Aquazol mass
272 ratio. TG experiments on the same paint layers after artificial ageing (Fig. 3D) do not show any
273 significant difference compared to the unaged ones. The results indicate that a strong interaction
274 between the pigment DPP and the polymer takes place. In fact, their thermal decomposition occurs
275 in a single step at a temperature lower than those of both polymer and pigment alone. This might
276 indicate the formation of a less stable structure, due to the interactions between the amide bonds in
277 the pigment and in the polymer, leading to an alteration of the quaternary structure of the polymer.

278 The Py-GC/MS analysis of the recipes containing DPP showed the presence of specific pyrolysis
279 peaks corresponding to the thermal degradation of the organic dye. From a qualitative point of view,
280 the pyrolysis profile of Aquazol mixed with DPP is the same as pure Aquazol added with the specific
281 pyrolysis products of DPP. However, differences in the relative abundances of pyrolysis products
282 were observed, namely a relatively higher abundance of dimer and trimers relative to propionamide
283 and low molecular weight products. This might be related to the interactions between the pigment
284 and the polymer. The differences were also highlighted by the principal component analysis of the
285 pyrolysis semi-quantitative data presented in Section 3.3. No significant alteration in the pyrolysis
286 profile was observed after artificial ageing, in line with the results presented for the TG analysis.

287 Aquazol 500 paint layers containing alizarin exhibit a more complex TG curve shape (Fig. 4B and
288 C). In the temperature range below 200 °C, where the loss of water occurs, there are the typical signal
289 of alizarin- and kaolinite based-pigments with two weak maxima in the derivative curve. The main
290 degradation step occurs in the range from 384 to 394 °C, depending on the recipe used and the ageing
291 of the sample, corresponding to about a 45% mass loss (50% compared to the dehydrated sample).
292 Thus, in the paint layers containing the alizarin pigment, the main thermal degradation occurs at a
293 lower temperature than in the pure Aquazol layer. In addition, in the derivative curve it is apparent
294 that the peak related to the main weight loss is wider with respect to the pure binder.

295 These data combined suggest that a partial depolymerization has occurred with respect to the
296 unpigmented layer. At higher temperatures (temperature range 450–700 °C), the TG curves also show
297 a broad mass loss corresponding to about 22–23% (about 25% if the water loss is not considered)
298 occurring in a wide temperature range depending on the recipe used for the formulation. This second
299 mass loss is most probably due to the degradation of stable compounds, formed because of the

300 interaction between the pigment and Aquazol, and possibly characterized by a higher molecular
301 weight than the main fraction. The extent of the interaction depends on the recipe used to produce the
302 paint, and in both cases the residue at 800 °C is about 25% compared to the dehydrated sample. Both
303 the depolymerization of the binder and the formation of the stable compounds, possibly characterized
304 by a higher molecular weight compared to the unmodified Aquazol, are more pronounced after
305 artificial ageing (Fig. 4D). The main thermal degradation occurs at 384 °C for recipe A (10 °C lower
306 than in the unaged layer), while the more stable compounds start to degrade at around 450 °C and are
307 not completely degraded at 800 °C. The particular reactivity of Aquazol with alizarin lake might be
308 explained by the presence of kaolinite and metakaolinite, which are known for their catalytic
309 properties [45], [46], [47], [48]. For example, a study reported their use in catalysis, after thermal-
310 activation, in transesterification to produce biodiesel using frying oil [49]. These minerals are also
311 known for their use in the catalytic cracking reactions of alkanes and alkenes [49], [50]. Kissin
312 proposed the solid acidic catalysis of kaolinite in these reactions [50].

313 The pyrograms of the paint layers containing Alizarin, both before and after artificial ageing, present
314 significant differences in the relative abundance of pyrolysis products compared to the pyrolysis
315 profile of pure Aquazol and of Aquazol paint containing the other two pigments investigated. The
316 relative amounts of the peaks corresponding to the monomers is higher than in the pyrograms of all
317 the other aquazol paint layers which do not contain alizarin, as shown in Fig. 2C. The relative
318 abundance of the monomer (peak number 10) is in this case higher than all the other pyrolysis
319 products, resulting in a very specific chromatographic profile. The high amounts of the monomer in
320 the pyrogram is consistent with the depolymerization observed by TG. The contribution of the
321 fraction characterized by a higher molecular weight than Aquazol is predictably not visible in the
322 pyrograms, probably due to their higher thermal stability revealed in the TG experiments.

323 The Aquazol 500 paint layers containing Quinacridone (Fig. 5B and C) show TG profiles where the
324 main weight loss, due to Aquazol, occurs at very similar temperatures to those of the unpigmented
325 paint layers, corresponding to a mass loss of about 66% at 410 °C (recipe A) and 432 °C (recipe B).
326 This mass loss exactly matches the composition of the paint layer (Aquazol/pigment 2:1). The
327 secondary mass loss, probably related to the degradation of the pigment, varies from 25% (recipe A)
328 to 29% (recipe B), with a residue at 800 °C as high as 4.5% and 3.6%, respectively. In addition, the
329 secondary degradation spans from 440 °C to 530 °C in the case of the paint formula prepared
330 according to recipe A (with a single maximum in the derivative curve at 494 °C), while in the sample
331 prepared according to recipe B, the mass loss is much broader (460–660 °C with two maxima at 567
332 °C and 620 °C in the DTG curve).

333 The two different formulations seem to produce opposite effects: the presence of glycerin and
334 Xanthan gum (recipe A) lowers the Quinacridone decomposition temperature by about 60 °C. On the
335 other hand, in the formulation without additives (recipe B) the thermogravimetric profile suggests
336 that beside the unaltered pigment (mass loss occurring at 567 °C, very close to that of the pure
337 pigment), there are other compounds with a higher thermal stability within the paint layer, possibly
338 due to the aggregation of aquazol and/or the pigment during drying [50].

339 The Py-GC/MS analysis of the paint layers containing Quinacridone shows the presence of the
340 specific pyrolysis products derived from the pigment, although in very low amounts compared to the
341 peaks due to the binder. The pyrolysis profile of the paint layer after artificial ageing did not show
342 any significant difference in composition compared to the corresponding unaged sample. Since the
343 pyrolysis profile of the binder is not significantly altered by the presence of the quinacridone pigment,
344 it is possible that the aggregates highlighted in recipe B are due to the auto-aggregation of the pigment,
345 inhibited by xanthan gum and glycerin in recipe A.

346 The strongest differences between the freshly prepared and the aged samples were highlighted by
347 FTIR analysis. The recipe B paint layer without pigment, both recipe A and B paint layers containing
348 alizarin, and recipe B paint containing quinacridone showed the appearance of new bands as a
349 consequence of ageing. Fig. 6 shows the FT-IR spectrum of the Aquazol 500 paint layer recipe B
350 without pigment, before and after artificial ageing. In the spectra of the aged material, a band appears
351 at 1734 cm⁻¹, and in addition the absorption band at 3267 cm⁻¹ has increased in intensity while the
352 band at 3480 cm⁻¹ has decreased compared to the unaged material. The bands at 1734 cm⁻¹ and at
353 3267 cm⁻¹ are characteristic of the hydroxyl group stretching and carbonyl group stretching of
354 carboxylic acids, respectively [23] (Table 2). The formation of carboxylic acids indicates that a
355 certain degree of photo-oxidation of Aquazol 500 has occurred during ageing. The low intensity of
356 the 1734 cm⁻¹ band indicates that only a small fraction of the polymer was involved in the oxidation
357 reactions, which was insufficient to produce significant variations in the pyrolysis profiles.

358

359 3.3 Principal component analysis (PCA)

360 Due to the high number of pyrolysis products highlighted in the chromatograms, the Py-GC/MS data
361 were interpreted using Automated Mass Spectral Deconvolution and Identification System (AMDIS),
362 a free software application available from NIST. This software operates the deconvolution of the
363 overlapping peaks based on their mass spectra. A dedicated library of mass spectra of the pyrolysis
364 products of Aquazol was created and used to collect the data relative to the peak area of each
365 compound. For the PCA analysis, the relative abundances of the five classes of the thermal
366 degradation products of Aquazol were used, namely low molecular weight molecules (LMW),

367 propionamide and related alkylated structures (P), monomer and related alkylated structures (M),
368 dimer and related alkylated structures (D), trimer and related alkylated structures (T). These relative
369 abundances were calculated as the sum of the integration results of the thermal degradation products
370 of each class, expressed as a percentage of the total area. The assignment of each pyrolysis product
371 to its class is indicated in Table 1.

372 A multivariate analysis of the data obtained together with the decomposition temperature of Aquazol
373 500 determined by TG experiments was performed by principal component analysis (PCA) of the
374 covariance matrix.

375 The scatter plot shows that the samples of the paint layers containing alizarin (both recipes A and B,
376 before and after ageing) are placed at negative values of PC1 and at high values of PC2, in a different
377 region of the scatterplot compared to those of all the other paint layers. The paint samples containing
378 Aquazol without any pigment and those containing quinacridone are located in the same region as
379 the scatter plot, at positive PC1 values, which is consistent with the lack of influence of quinacridone
380 on the behavior of the polymer. The samples containing DPP form a separate compact group at
381 negative values of both PC1 and PC2, characterized by a high trimeric structure content in the
382 pyrogram.

383 The most notable differences in the pyrolysis profiles were highlighted by the PCA analysis in the
384 samples containing alizarin. An examination of the loading plot suggests that the relatively higher
385 values of the relative abundance of monomers and of the related alkylated structures is the main
386 feature of the pyrolysis profile, which differentiates the samples containing alizarin from the other
387 samples. We hypothesized that this different behavior could be due to depolymerization reactions,
388 possibly catalyzed by the inorganic fraction of the alizarin pigment.

389 As expected, depolymerization and intermolecular interactions between the polymer and the
390 pigments, responsible for the differences in the thermal degradation highlighted by PCA, do not
391 produce differences in the IR spectra, since they do not entail significant changes in the functional
392 groups. However, FT-IR spectroscopy proves efficient in highlighting oxidation phenomena.

393 **4. Conclusions**

394 The application of a multi-analytical approach based on the combination of thermoanalytical, mass
395 spectrometric and spectroscopic techniques enabled us to assess the stability of Aquazol 500 as a
396 paint binder during ageing as well as the occurrence of pigment-Aquazol 500 interactions, and their
397 evolution during ageing.

398 The FT-IR analysis of paint layers prepared with Aquazol 500 without pigments highlighted the
399 occurrence of oxidation reactions during artificial ageing, leading to the appearance of a band at 1734
400 cm⁻¹. Nevertheless, Py-GC/MS profiles of aged paint layers did not show significant variations when
401 compared to the non-aged paint, indicating that only a small fraction of the polymer was oxidized, in
402 agreement with the data reported in the literature [23]. This thus confirms its potential for use as a
403 binder for retouching paints in the restoration of works of art.

404 When Aquazol 500 was used in formulations containing pigments, Aquazol -pigments interactions
405 were observed. Depending on the pigment used, the following results were obtained:

- 406 • DPP-Aquazol 500 paint layers: a very strong DPP-Aquazol interaction was observed, leading to
407 TG curves showing a single degradation step for DPP and Aquazol 500, at a temperature lower
408 than those of pure DPP and Aquazol 500. No evidence was observed of a significant modification
409 in the thermal behavior of the paint due to the presence of additives such as glycerol and xanthan
410 gum (recipe A), or due to ageing. PCA analysis of Py-GC/MS data highlighted that the pyrolytic
411 profile appears to be significantly different from a quantitative point of view from that of pure
412 Aquazol, which may be related to the interaction of the polymeric chain and the pigment. The
413 presence of amide bonds in both structures might account for their strong interaction.
- 414 • Alizarine lake-Aquazol 500 paint layers: a strong interaction was observed leading to a partial
415 depolymerization of the binder highlighted in the TG curve and confirmed by the pyrogram
416 obtained by analytical pyrolysis, with the predominant formation of the monomer. The data
417 analysis using PCA also highlighted this behavior. In addition, stable compounds are formed
418 when alizarin lake is present in the paint layer, probably due to the presence of kaolinite in the
419 pigment formulation. The presence of additives and of artificial ageing played a minor role in the
420 degradation pattern of these paints.
- 421 • Quinacridone-Aquazol 500 paint layers: the presence of Quinacridone did not influence the
422 behavior of the polymer, as assessed both by TG and analytical pyrolysis. However, quinacridone
423 behaves differently in the paint layers than as a pure pigment, mostly depending on the recipe
424 used in the paint production, forming varying stable aggregates.

425 The adopted analytical methods achieved complementary information: FT-IR spectroscopy allowed
426 us to observe the formation of oxidized functionalities even if in very low concentration, while
427 thermal analysis techniques achieved information on intermolecular interactions between the polymer
428 and the pigments.

429 Aquazol 500 is now being used to retouch paintings. This study highlights that the stability of the
430 polymer, as shown in a previous study limited to TiO₂ nanoparticles [23], is affected by the presence
431 of pigments. This is particularly the case of organic lakes containing an inorganic fraction in the
432 formulation, leading to different physico-chemical phenomena, depending on the nature of the
433 pigment.

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449

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- 588

589 **Figure Captions**

590 Fig. 1. TG curves of: a) Aquazol 500 in bulk; b) paint layer of Aquazol 500, recipe A, without
591 pigment; c) paint layer of Aquazol 500, recipe B, without pigment; d) paint layer of Aquazol 500,
592 recipe A, without pigment, after artificial ageing.

593

594 Fig. 2. Chromatograms obtained in the GC/MS analysis of: a) paint layer of Aquazol 500, recipe B,
595 without pigment; b) paint layer of Aquazol 500, recipe B, without pigment, after artificial ageing; c)
596 paint layer of Aquazol 500, recipe B, with alizarin.

597

598 Fig. 3. Structural formula of diketo-pyrrolo-pyrrole and TG curves of: a) diketo-pyrrolo-pyrrole
599 (DPP) pigment powder; b) paint layer of Aquazol 500, recipe A, containing DPP; c) paint layer of
600 Aquazol 500, recipe B, containing DPP; d) paint layer of Aquazol 500, recipe A, containing DPP,
601 after artificial ageing.

602

603 Fig. 4. Structural formula of alizarin and TG curves of: a) alizarin pigment powder; b) paint layer of
604 Aquazol 500, recipe A, containing alizarin; c) paint layer of Aquazol 500, recipe B, containing
605 alizarin; d) paint layer of Aquazol 500, recipe A, containing alizarin, after artificial ageing.

606

607 Fig. 5. Structural formula of quinacridone and TG curves of: a) Quinacridone pigment powder; b)
608 paint layer of Aquazol 500, recipe A, containing Quinacridone; c) paint layer of Aquazol 500, recipe
609 B, containing Quinacridone.

610

611 Fig. 6. ATR-FT-IR of Aquazol 500 paint layer, recipe B, without pigment: A) before artificial ageing
612 and B) after artificial ageing. Peak assignation is shown in Table 2.

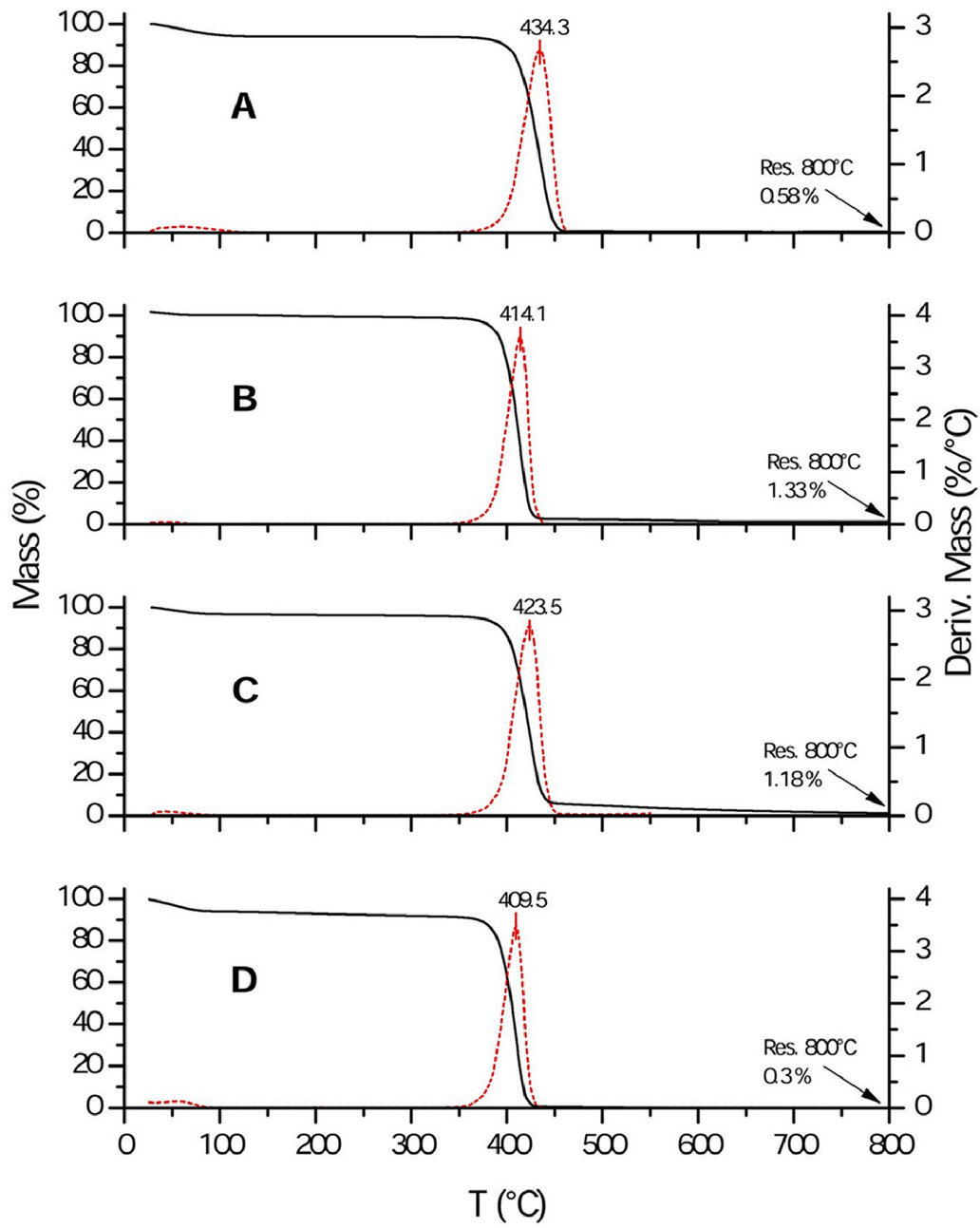
613

614 Fig. 7. PCA loading plot and scatter plot of the Py-GC/MS data (abundance of low molecular weight
615 molecules: LMW; propionamide and related alkylated structures: P; monomer and related alkylated
616 structures: M; dimer and related alkylated structures: D; trimer and related alkylated structures: T)
617 and TG data (temperature of decomposition of the polymer) of the aged and unaged samples, recipes
618 A and B, without and with the organic pigments diketo-pyrrolo-pyrrole (DPP), quinacridone (QUI)
619 and alizarin (ALI).

620

621 Fig 1

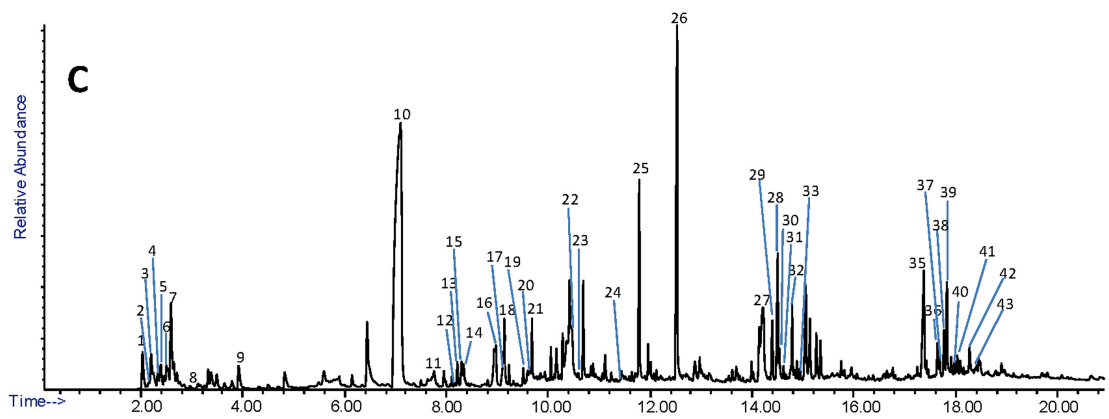
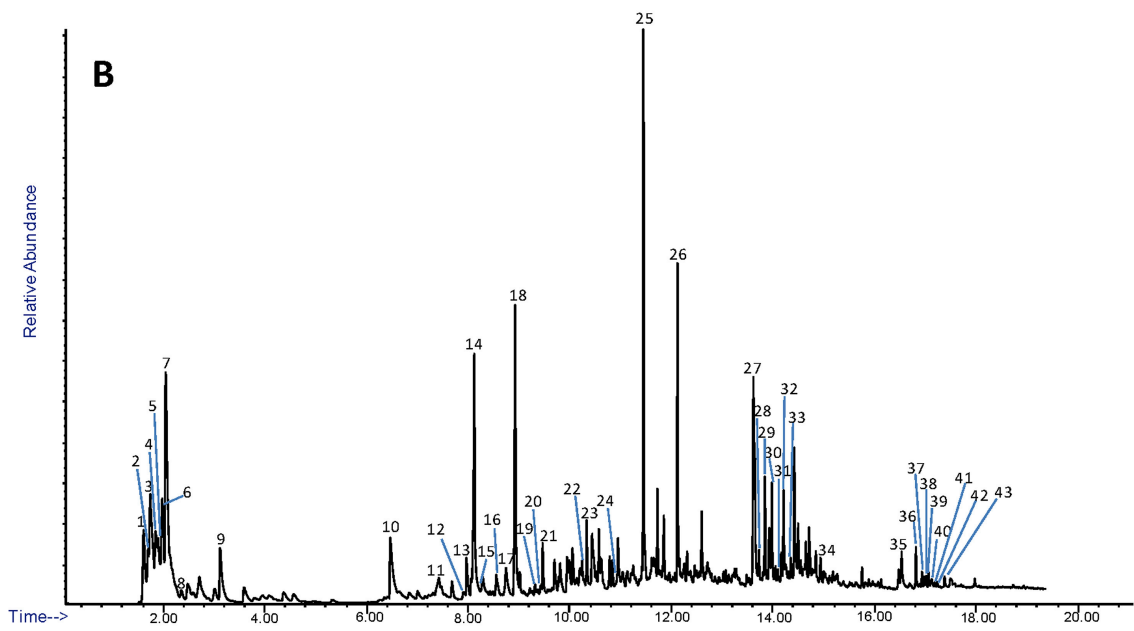
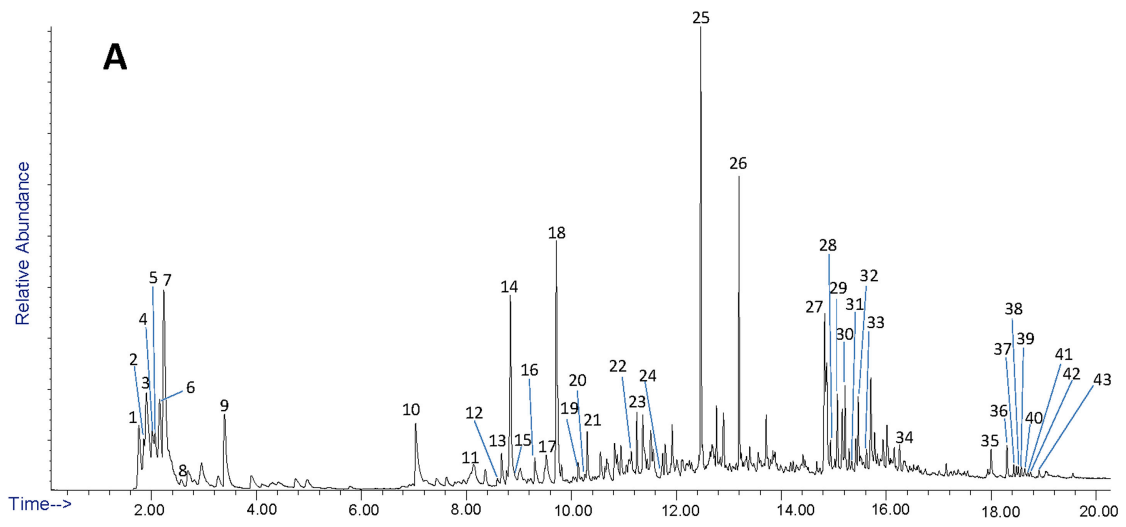
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624 Fig 2

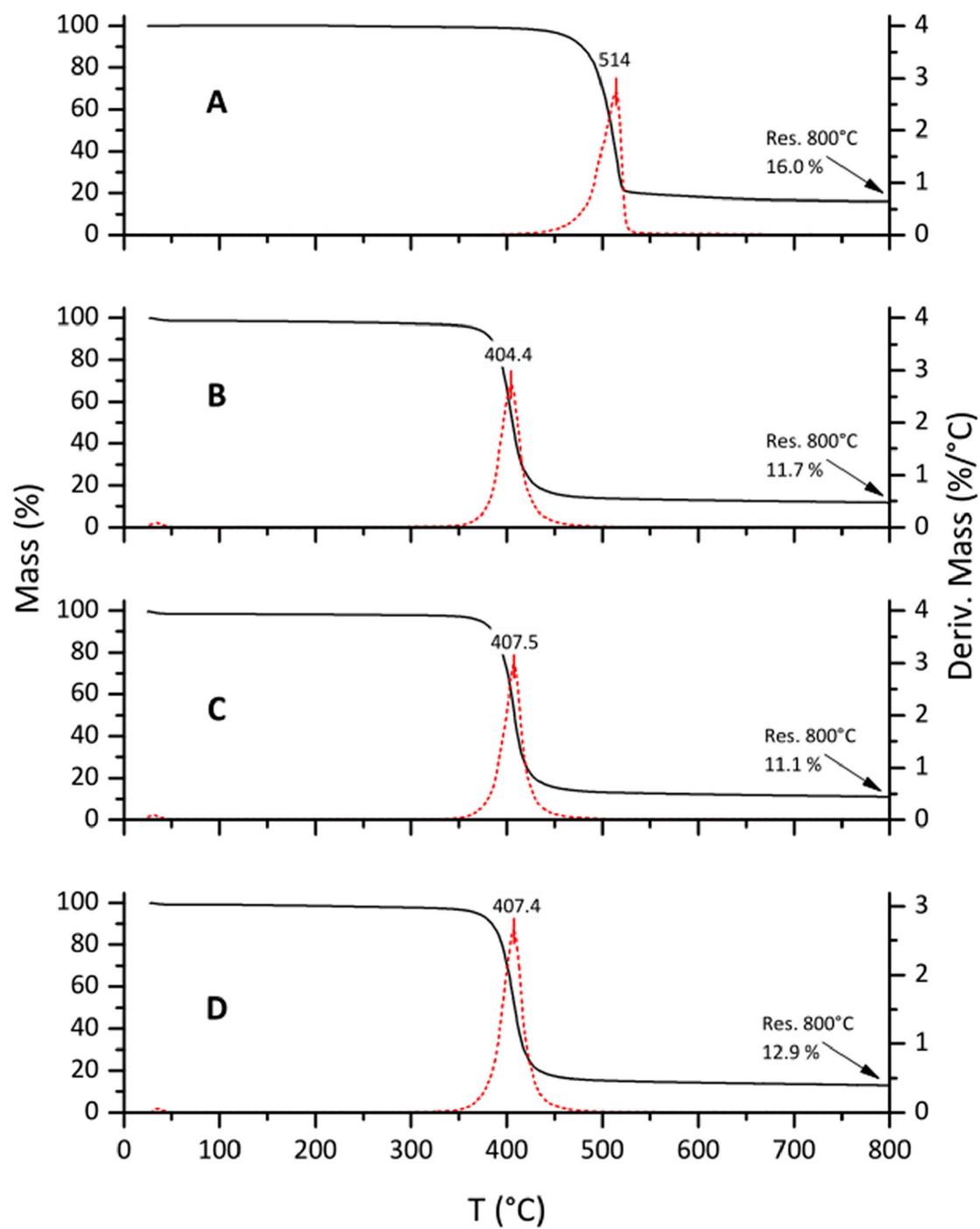
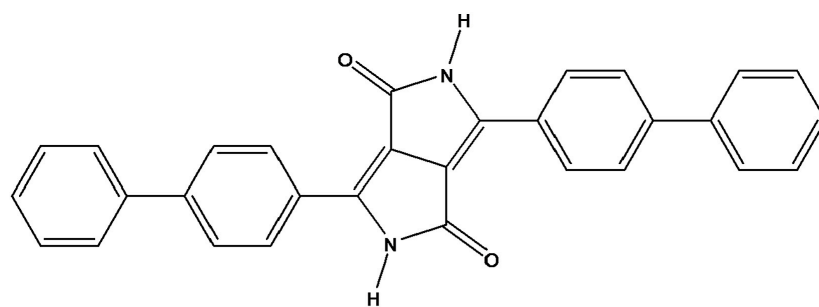
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627 Fig 3

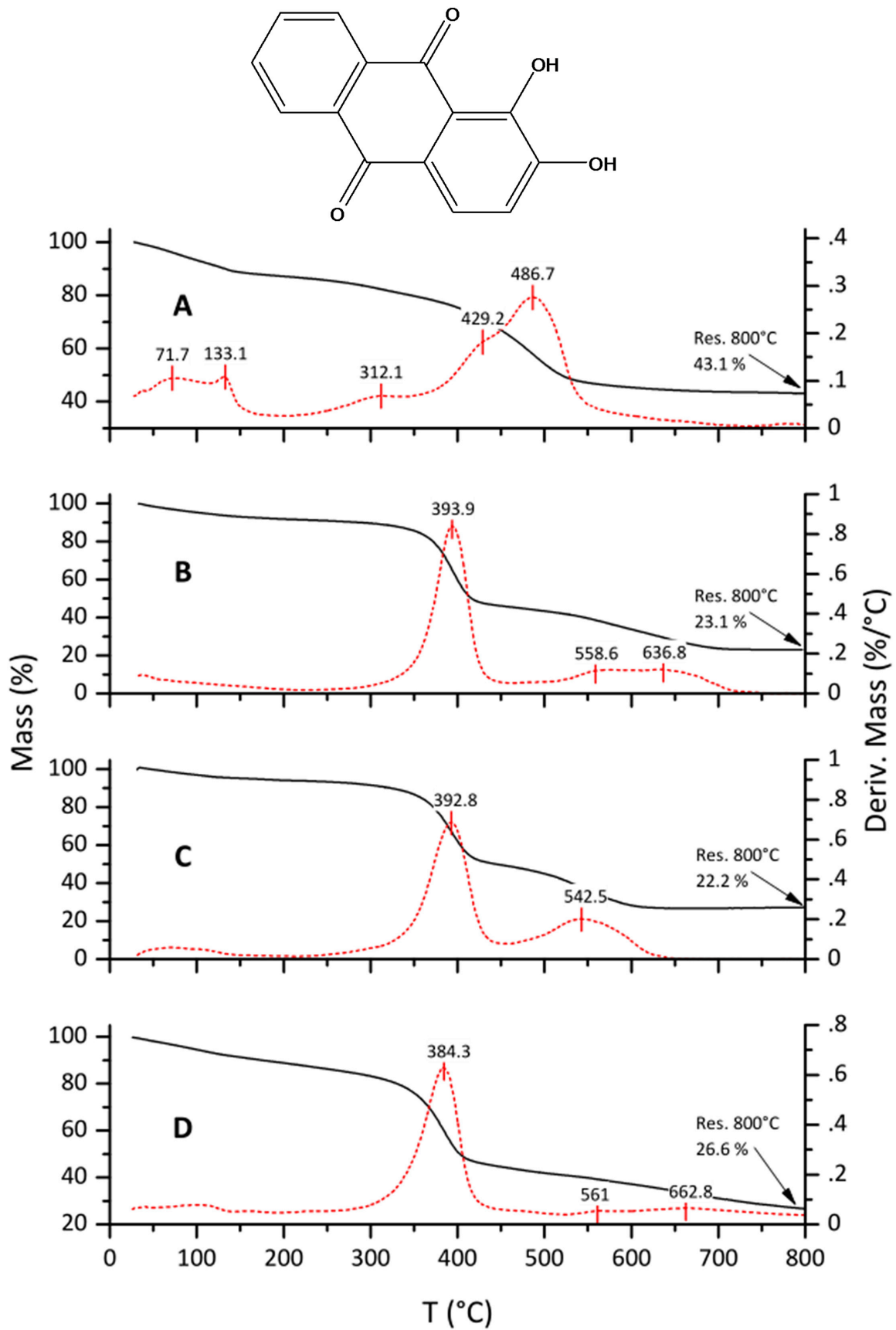
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630 Fig 4

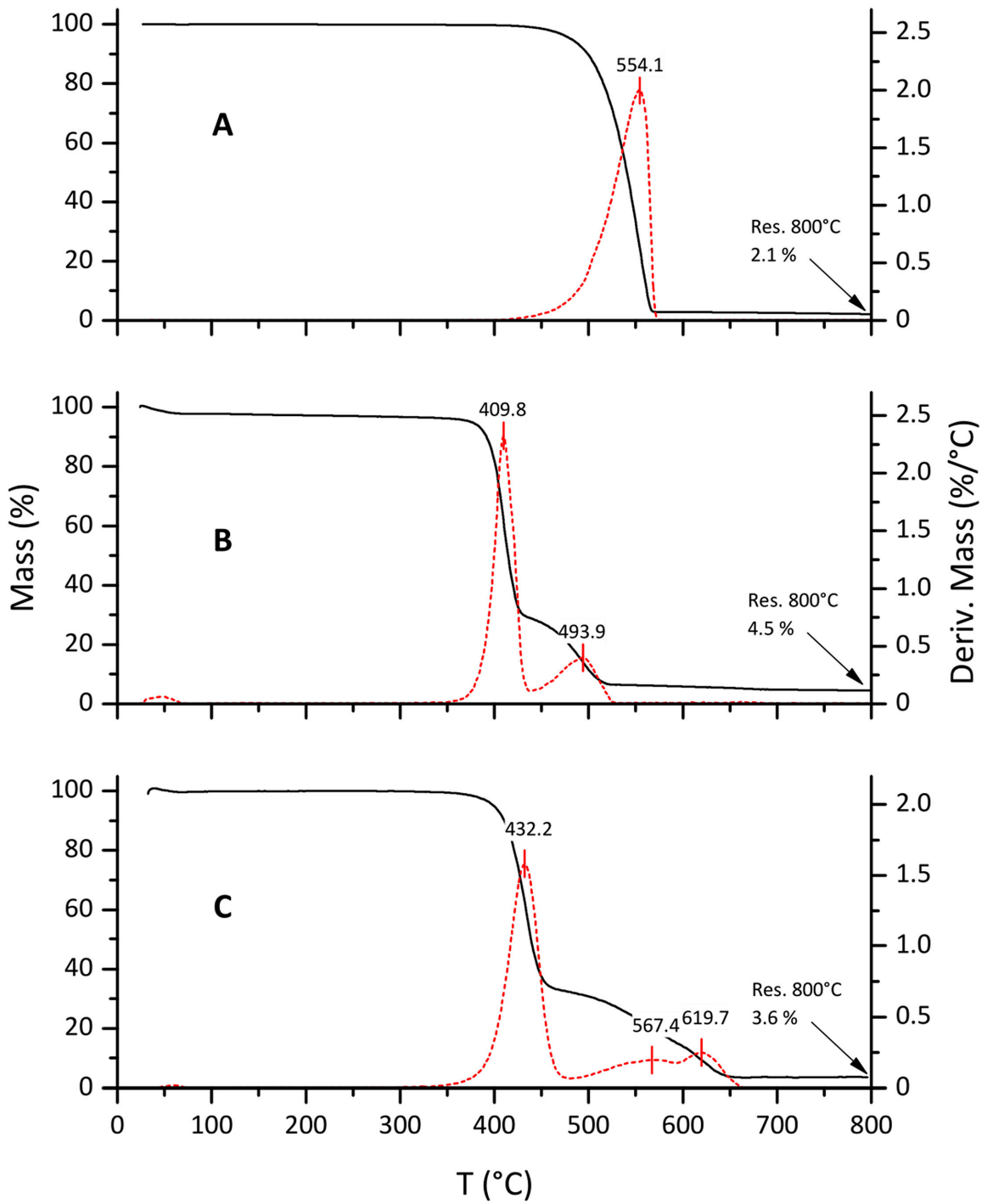
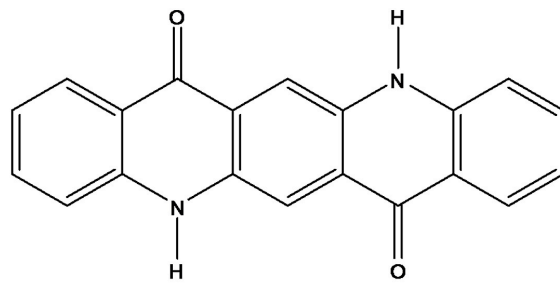
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634 Fig 5

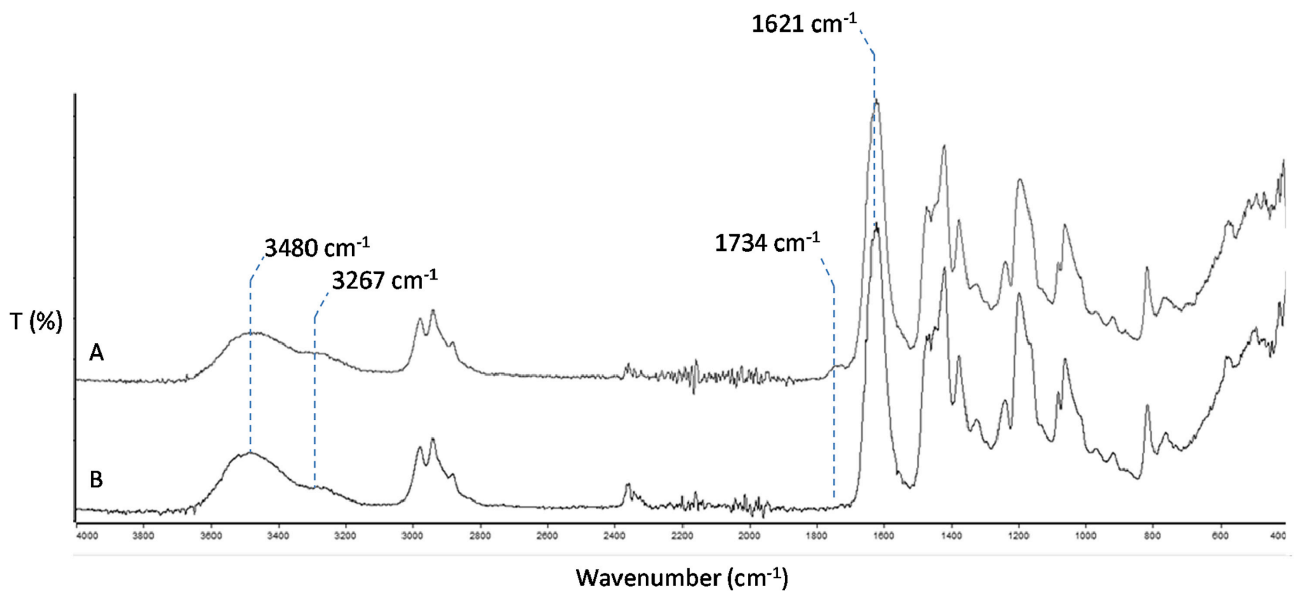
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638 **Fig 6**

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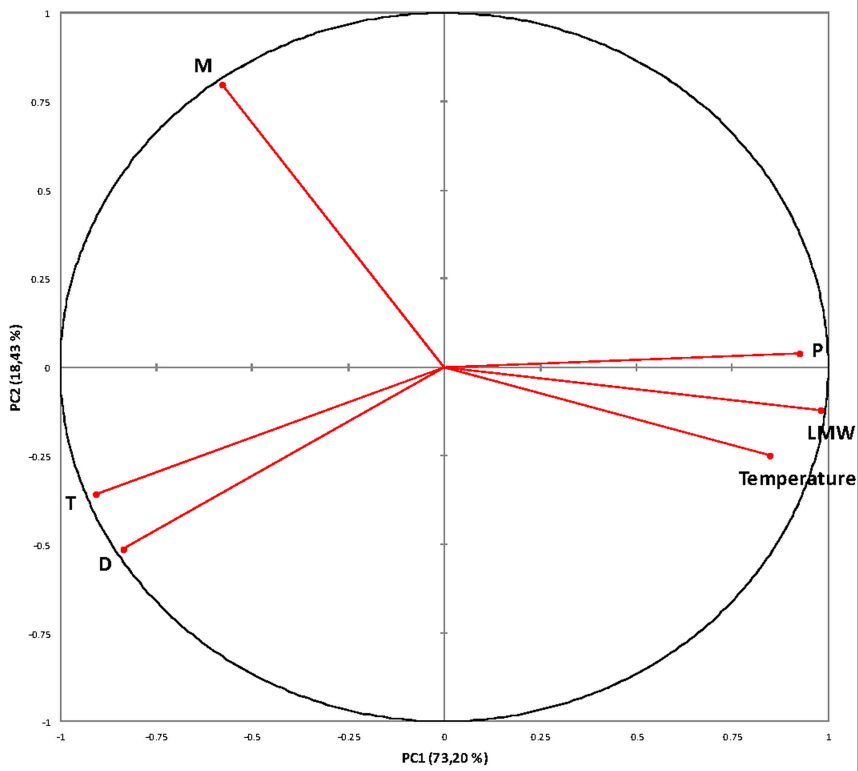
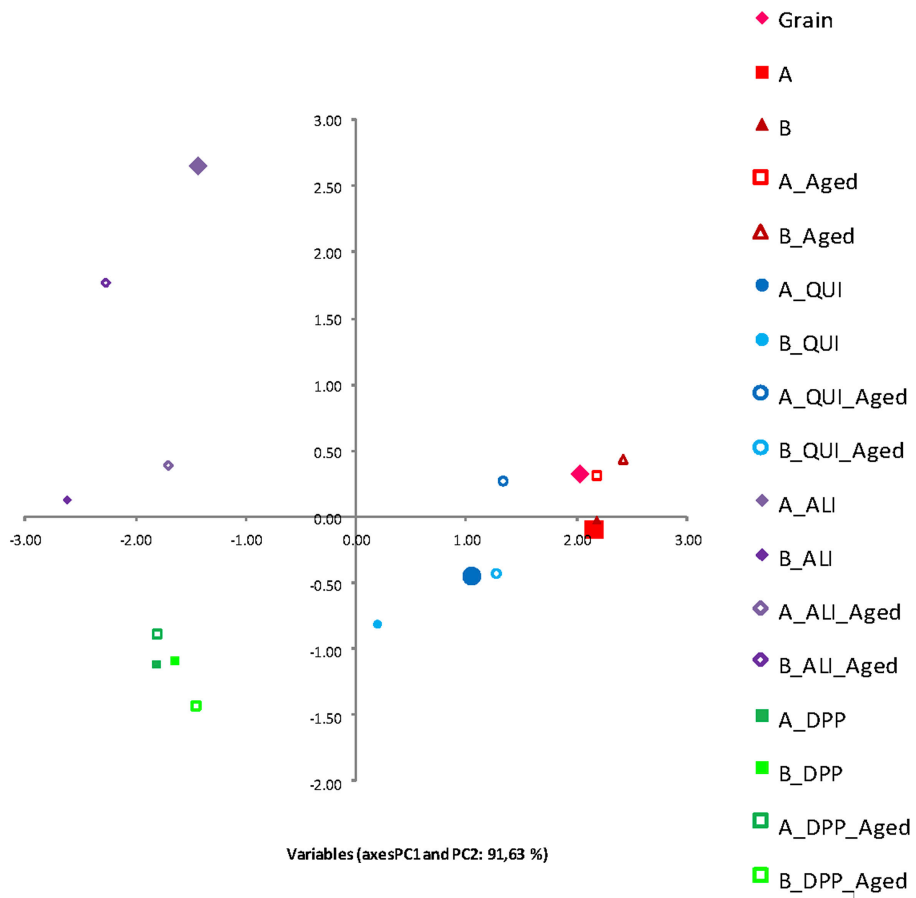


640

641

642 Fig 7

643



644

645 Table 1. Compounds identified in the Py-GC/MS chromatograms of Aquazol 500 paint layers. The
 646 pyrolytic products were classified into the following five groups: low molecular weight molecules
 647 (LMW), propionamide and related alkylated structures (P), monomer and related alkylated structures
 648 (M), dimer and related alkylated structures (D), trimer and related alkylated structures (T) [13].
 649

Peak number	m/z	Assignment	Classification
1	56	Isobutene	LMW
2	57	Isobutene-amine	LMW
3	57	Isocyanatomethane	LMW
4	67	2-butene-nitrile	LMW
5	71	Unknown	LMW
6	71	Isocyanatoethane	LMW
7	54	Propane-nitrile	LMW
8	82	2-methyl-butane-nitrile	LMW
9	86	3-pentanone	LMW
10	99	Monomer (M)	M
11	73	Propionamide (P)	P
12	107	2-ethyl-pyridine	LMW
13	108	Ethyl-pyrazine	LMW
14	111	P + CH ₂ CH ₂ CH ₃	P
15	87	P + CH ₃	P
16	101	P + CH ₂ CH ₃	P
17	101	P + CH ₂ CH ₃	P
18	113	M + CH ₃	M
19	115	M + CH ₃	M
20	125	M + CH ₂ CH ₃	M
21	127	M + CH ₂ CH ₃	M
22	127	M + CH ₂ CH ₃ isomer	M
23	125	M + CH ₂ CH ₃ isomer	M
24	155	CH ₃ CH ₂ +M + CH ₂ CH ₃	M
25	152	Sesquimer M ₂ -CH ₂ CH ₂ CH ₃	M
26	154	Sesquimer M ₂ -CH ₂ CH ₂ CH ₃	M
27	186	M ₂ -CH ₃	D
28	200	Dimer (M ₂)	D
29	198	Dimer (M ₂)	D
30	212	M ₂ +CH ₃	D
31	196	Dimer (M ₂)	D
32	226	M ₂ +CH ₂ CH ₃	D
33	224	M ₂ +CH ₂ CH ₃	D
34	254	CH ₃ CH ₂ +M ₂ +CH ₂ CH ₃	D
35	281	M ₃ -CH ₃	T
36	285	M ₃ -CH ₃	T
37	297	Trimer (M ₃)	T
38	297	Trimer (M ₃)	T
39	311	M ₃ +CH ₃	T
40	323	M ₃ +CH ₂ CH ₃	T
41	325	M ₃ +CH ₂ CH ₃	T
42	355	CH ₃ CH ₂ + M ₃ +CH ₂ CH ₃	T
43	355	CH ₃ CH ₂ + M ₃ +CH ₂ CH ₃	T

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Table 2. Characteristic absorption bands in FT-IR spectrum of Aquazol 500.

Absorption bands (cm ⁻¹)	Vibrational modes
3483	O-H (moisture, adsorbed water)
2971, 2940, 2880	C-H stretching (CH ₂ , CH ₃ groups)
1629	C=O stretching (tertiary amide)
1473	C-N amide stretching
1418, 1375, 1321	C-H bending (CH ₂ , CH ₃ groups)
1241, 1192, 1061	C-C stretching (polymer chain)

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