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1 **From tomato pomaces biorefinery to bio-based shape memory semi-crystalline polyester networks**

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9

10 **Abstract**

11 The development of new bio-based polyester materials is of great interest. Seed-free and pulp-free
12 peels provided by the biorefinery of industrial tomato wastes, i.e. pomaces, are rich in cutin, a
13 polyester of hydroxy fatty acids. In this study, these fatty acids were recovered after alkaline hydrolysis
14 resulting in a crude brown-colored extract, due to the co-extracted phenolic compounds. Further
15 purification drastically reduces these phenolic compounds, resulting in a pale yellow product
16 containing about 95% (9/10)-16-dihydroxy hexadecanoic acid and 5 % dicarboxylic fatty acids.
17 Polymerization of the purified extract at 150°C results in a weakly crosslinked polyester network, with
18 an estimated average number of 80 repeating units between nodes. Interestingly, the mechanical
19 behavior of this material at 25°C is determined by its semi-crystalline structure. A necking phenomenon
20 is observed during tensile tests resulting in an apparent plastic deformation of 286% and a Young
21 modulus of 72 MPa. In addition, this bio-based polyester exhibits shape memory properties with the
22 ability to be hot or cold- programmed. This work highlights the significant impact of minor compounds,
23 related to the biochemical heterogeneity of agro-industrial waste products on the polyester properties
24 and the potential of the biorefinery process to modulate the properties of biopolymers.

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26

27 **Keywords**

28 hydroxy-fatty acids, phenolic, necking, bio-sourced, cutin-like

29

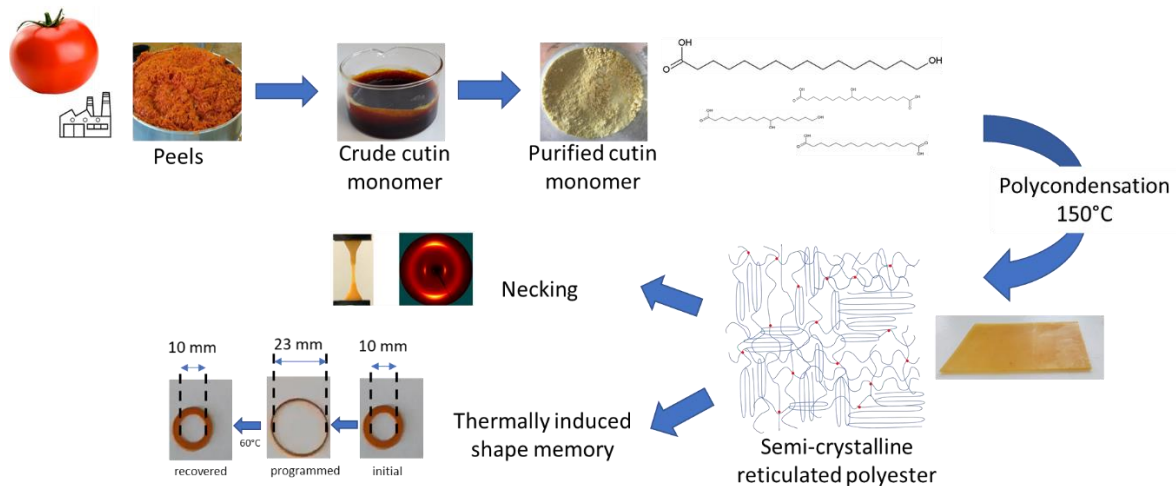
30 **Synopsis.**

31 Tuning the biorefinery of tomato pomaces enables the synthesis of a fully bio-based
32 shape memory polyester material.

33

34

35 Graphical abstract



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39 Introduction

40

41 For both economic and environmental reasons, there is a great interest in new polyesters
42 based on renewable carbon resources, without competing with food uses heredia^{1,2-4}. Intensive efforts
43 are currently focused on the valorization of agricultural and food processing wastes as biorefinery
44 feedstocks for the production of monomers⁵⁻⁷

45 Long-chain semi-crystalline polyesters with excellent mechanical properties^{8,9} can be produced from
46 plant oils, with two major drawbacks: competition with food uses, and the need to chemically convert
47 crude fatty acids into readily polymerizable monomers containing both hydroxyl and carboxylic groups
48^{3, 8, 10}. Similar chemical structures, are naturally present in plant polyester i.e. suberin and cutin
49 consisting in polyester network, biosynthesized from long-chain hydroxy fatty acids containing 16 to
50 18 carbon atoms^{11,12}. These monomers can be extracted from industrial by-products and can be used
51 for polymer synthesis^{7,13-15}. However, the natural suberin composition is complex, which makes the
52 control of the polyester functionalities more difficult. Conversely, tomato cuticle is rich in cutin
53 polyester whose monomer composition is highly homogenous (tomato cutin mainly consists of 9(10)-
54 16 hydroxy hexadecenoic acid). Tomato pomace is currently the largest and growing source of cutin-
55 rich wastes¹⁶, ranging from 6 to 9.10⁶ tons per year¹⁷⁻¹⁹. After separation from seeds and pulp, tomato
56 peels, containing more than 60% cutin, can be hydrolyzed by alkali treatment, to yield fatty acid
57 extracts containing mostly 9/10-16 dihydroxyhexadecanoic acid⁷ with typically less than 10% of
58 impurities such as α,ω -dicarboxylic fatty acids and phenolic compounds.

59 Several studies have shown that the thermal polymerization of such extracts leads to insoluble
60 polyesters^{7,13,16,20-22} with a crosslinked structure. This indicates that the role of the minor compounds
61 in the formation of the macromolecular structure should not be overlooked. Indeed, the
62 polymerization of pure 9/10-16 dihydroxyhexadecanoic acid would be expected to lead to
63 hyperbranched polymers²³ due to its ABB structure (A and B standing for acid and hydroxyl functions,
64 respectively). While dicarboxylic fatty acids undoubtedly participate in thermal polyesterification, the
65 exact role of phenolic compounds is probably more complex and difficult to control due to their high

66 reactivities ²⁴. In the present work, the amount of co-extracted phenolic compounds was greatly
67 reduced by purification, with two objectives: i) to study the influence of dicarboxylic fatty acids alone
68 on crosslinking, and ii) to get rid of the high coloring power of phenolic compounds, without resorting
69 to chemical discoloration with hydrogen peroxide, with potentially detrimental effects on the
70 polyester properties and human health, and thus on potential high value applications ^{4, 25}.

71 Serendipitously this reduction of phenolic compounds leads to semi-crystalline crosslinked
72 polyester with unusual mechanical behavior and shape memory properties. The relationships between
73 this peculiar functionality and the thermal and structural properties of this new bio-based material are
74 discussed.

75
76

77 **Materials and Methods**

78 ***Material and chemicals***

79 Industrial tomato pomaces were provided by the “Conserveries de Bergerac” (UNIPROLEDI,
80 Bergerac, France). Analytical grade solvents were obtained from Carlo Erba (Val de Reuil, France).
81 Aminopropyl silica (NH₂) gel was purchased from Macherey-Nagel. Reference chemicals (gallic acid, p-
82 coumaric acid, naringenin) were purchased from Sigma Aldrich (L’Isle d’Abeau Chesne, France).

83 **Methods**

84 ***Cutin fatty acid monomers extraction and purification***

85 The crude cutin monomer (CM) fraction was extracted from peels isolated from industrial
86 tomato pomaces according to the procedures previously described ⁷. The crude dark brown colored
87 CM extract was further purified by aminopropyl silica column chromatography according to a
88 modification of a described procedure ²⁶. The aminopropyl silica (100 g) dispersed in chloroform:2-
89 propanol 2:1 (v:v) was poured into a glass column (50 cm x 5 cm) with a fritted glass filter and a Teflon
90 tap placed on the bottom. The crude cutin monomer extract (10g) solubilized in the
91 chloroform:isopropanol mixture (50mL) was applied to the aminopropyl phase. After elution with 1L
92 of the chloroform:isopropanol mixture, the hydroxy fatty acids were eluted with 1.5L of chloroform.
93 After evaporation of the solvent under vacuum and further thorough freeze-drying, the pale yellow
94 solid was recovered and stored in a dark place at room temperature. Four purification batches were
95 pooled to perform the experiments. The depigmented hydroxy-fatty acid fraction was further referred
96 as “purified CM” fraction. Total phenolic content was determined by the Folin-Ciocalteu colorimetric
97 assay ²⁷ using gallic acid for the standard curve. It was expressed as mg of gallic acid equivalents (GAE)/g
98 cutin monomer extract. In addition, the phenolics profile was performed by a high-performance liquid
99 chromatography (HPLC)-diode array detection (DAD) method ²⁸ on an Acquity ARC system (Waters,
100 USA) with a diode array detector, using a BEH X-Bridge C18 column (250 × 4.6 mm, 5 μm) and guard
101 column (20 × 4.6 mm, 5 μm).

102 The fatty acid composition was determined as previously described by GC-FID-MS ²⁹ and
103 matrix-assisted laser desorption/ionization (MALDI)-time-of-flight (TOF) MS adapted from ³⁰. A DHB
104 (2,5-dihydroxybenzoic acid) solution at 3 mg.mL⁻¹ in H₂O/methanol (1:3) with 2.5 mM of LiCl was used
105 as the MALDI matrix. The samples were mixed with the matrix solution in a 1:9 ratio (v/v) and the
106 mixture (1 μL) was deposited on a polished steel MALDI target plate. MALDI measurements were then

107 performed on a rapifleX MALDI-TOF spectrometer (Bruker Daltonics, Bremen, Germany) equipped
108 with a Smartbeam 3D laser (355 nm, 10000 Hz) and controlled using the Flex Control 4.0 software
109 package. The mass spectrometer was operated with positive polarity in reflectron mode. Spectra were
110 acquired in the range of 180–5000 m/z. Chemical labelling of the free OH by benzyl-etherification was
111 performed as previously described³¹.

112 The monomeric content of the hydroxy fatty acid fractions was determined by size-exclusion
113 chromatography on a Waters HPLC system coupled to an evaporative light scattering detector (Sedere
114 Sedex 75 ELSD, Alfortville, France). A home-packed column (30 cm X 1 cm) of Sephadex[®] LH20 I (Sigma-
115 Aldrich) was used and elution with methanol was performed at 1 mL.min⁻¹ 25°C with methanol. PEG
116 standards were used for molecular mass calibration.

117

118 ***Polyester synthesis and characterization***

119 Several films of typically 700 ± 50 µm thickness was produced by bulk poly-condensation as
120 previously described⁷. Briefly, an amount of purified CM was put in Teflon molds (5cm × 5cm or 10cm
121 x 10 cm) and baked for 24h at 150°C in an oven. Vacuum was applied during the first 120 min. to avoid
122 the formation of bubbles. This also facilitated the removal of the water molecules generated by the
123 polyesterification. Then, the water molecules vaporized by heating at high temperature were
124 trapped in a beaker containing dry silica gel. These films were characterized using the techniques
125 described below. Mechanical tests were only possible on 10 cm x 10 cm films. Shape memory was
126 qualitatively tested on all films.

127 Attenuated total reflectance Fourier-transform infrared spectroscopy (ATR-FTIR) spectra were
128 recorded on a Nicolet Magna IR 550 spectrometer (Thermo Fisher Scientific, Saint-Herblain, France)
129 equipped with a liquid nitrogen-cooled mercury-cadmium-telluride detector. The instrument was
130 continuously purged with dry air. Spectra of polyester films were obtained by attenuated total
131 reflection (ATR) using a single reflection accessory equipped with a diamond crystal at a 45° angle of
132 light incidence. All spectra (3 per polyester) were acquired in the 4000 to 700 cm⁻¹ range with a
133 resolution of 2 cm⁻¹ and accumulation of 30 scans. Spectral deconvolution was performed using Fytik
134 software³².

135 ¹³C solid-state NMR (SSNMR) was performed on a Bruker AvanceIII-400 MHz spectrometer
136 operating at 100.61 MHz for ¹³C, equipped with a double-resonance H/X CP-MAS 4-mm probe (Bruker
137 Biospin, Wissembourg, France). A cross-polarization magic-angle-spinning experiment (CP-MAS) was
138 performed. Briefly, 80 mg of the co-polyester films were packed into 4 mm zirconia rotors. The samples
139 were spun at 12,000 Hz at room temperature. The CP-MAS spectrum was acquired with a contact time
140 of 1.5 ms, a recycling delay of 8 s and an accumulation of 8192 scans. The carbonyl carbon was set to
141 176.03 ppm through external glycine calibration. Spectral deconvolutions were performed using
142 PeakFit[®] software (Systat Software, Inc., USA).

143

144 ***Rheology during polymerization***

145 Crude and purified CM samples were placed in a dynamic rotational rheometer (HAAKE MARS
146 III). A plate–plate geometry (diameter 20mm, gap 1mm) was used. The rheometer was preheated at
147 150°C before the samples were introduced and a 0.01% sinusoidal deformation was applied at 1Hz.
148 The evolution of moduli G' and G'' resulting from the thermal polymerization at 150°C was thus

149 monitored. Note that unlike the polymer synthesis conditions, vacuum could not be applied during the
150 first 120 minutes of the reaction in the rheometer. In addition, the free surface area at the edge of the
151 material between the plates is significantly smaller than that of the samples polymerized in the oven.
152 Therefore, the water molecules formed by the polyesterification reaction are expected to take more
153 time to diffuse and escape into the air surrounding the material.

154 ***Differential Scanning Calorimetry***

155 Differential Scanning Calorimetry (DSC) measurements were performed with a DSC Q100 (TA
156 Instruments, Newcastle, DE). The calorimeter was calibrated with indium ($\Delta H = 28.41 \text{ J.g}^{-1}$; melting
157 point of 156.66°C). A piece of polyester (weight about 8 mg) was loaded into a hermetically sealed
158 aluminum pan. An empty and hermetically sealed aluminum pan was used as reference. The DSC pans
159 were introduced in the calorimeter at 20°C . The scanning rate was 3°C.min^{-1} , or 5°C.min^{-1} to be in
160 similar conditions as for XRD measurements as a function of temperature. Data analysis was performed
161 using TA Universal Analysis program. The melting transition temperature (T_m) was taken at the peak
162 maximum recorded during heating. Enthalpy changes of the transitions (ΔH_m) were obtained from the
163 area under the peak. The DSC experiments were performed in triplicate, by preparing and analyzing
164 three independent pans.

165 ***X-ray diffraction***

166 X-ray diffraction (XRD) measurements as a function of temperature were performed on the
167 high brilliance SWING beam line at the SOLEIL synchrotron facility, with the monochromator set at 12
168 KeV. Using a EIGERX 4M detector at a distance of 0.519 m from the sample, diffraction patterns were
169 recorded for reciprocal spacing q . The q range allowed the characterization of the packing of the acyl
170 chains of the polyester. 1D XRD curves were obtained by circular averaging of the 2D images using the
171 Foxtrot software. A piece of polyester was loaded into thin quartz capillaries of 1.5 mm diameter (GLAS
172 W. Muller, Berlin, Germany). The capillary was inserted into a Linkam oven (Linkam Scientific
173 Instruments Ltd, Waterfield, UK). This set-up allowed synchrotron-radiation XRD as a function of
174 temperature. The XRD experiments were conducted at fixed temperature (20°C) and on heating from
175 20°C to 95°C at 5°C.min^{-1} (375 patterns; 1 frame each 0.2°C ; 0.5 sec exposure time and 0.5 sec gap
176 time). The three-dimensional plot of the XRD patterns as a function of temperature was generated
177 using R software (R Foundation for Statistical Computing, Vienna, Austria). PeakFit software (Jandel
178 Scientific, Germany) was used to determine the positions of the Bragg reflections.

179 For crystal orientation studies, two-dimensional wide-angle X-ray scattering (WAXS) diagrams
180 were recorded using a Bruker D8 X-ray diffractometer (Madison, WI) equipped with a two-dimensional
181 GADDS detector. The X-ray radiation, Cu KR1 ($\lambda = 0.154 \text{ 05 nm}$), produced in a sealed tube at 50 kV and
182 1 mA was selected and parallelized using a Gobel mirror parallel optics system and collimated to
183 produce a beam diameter of $500 \mu\text{m}$. WAXS diagrams were recorded in the transmission mode with
184 the specimens lying with their length parallel to the vertical axis of the detector. Orientation was
185 determined by azimuthal integration between 0.680 and 0.386 nm corresponding to the amorphous
186 scattering band. Background and beamstop profiles were subtracted from the sample signals; the
187 resulting signal was smoothed for more clarity. The acquisition time was 15 min.

188 ***Dynamic mechanical thermal analysis***

189 Dynamic mechanical thermal analysis (DMTA) was performed on a DMTA MKIV (Rheometric
190 Scientific, US). Rectangular specimens (20 mm × 4 mm) were cut from the films and the thickness of
191 about 0.7-1 mm was accurately measured with a micrometer. The samples were analyzed in the tensile
192 mode at the frequency of 1 Hz with a strain amplitude of 0.1 %. To keep the samples taut, a static force
193 was applied to the sample that was 10% higher than the dynamic force. A scanning rate of 3 °C·min⁻¹
194 from -50 °C to 80 °C was used. Each sample was analyzed in duplicate. For recovery measurements, a
195 very weak static force of 10⁻³N is applied to the sample in order to follow the free shrinkage of the
196 sample during its heating.

197 ***Mechanical properties***

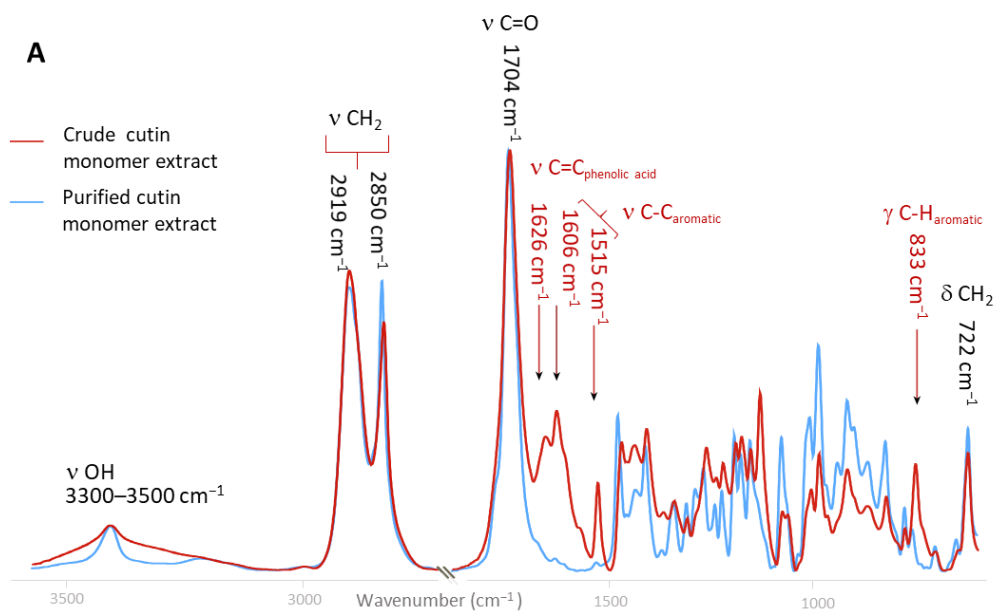
198 Tensile testing of the films was performed using the MTS Synergie 100 (MTS Systems
199 Corporation, USA). The film samples were cut into a dog-bone shape and their thickness were
200 accurately measured with a vernier caliper. The test was performed using a cross-head speed of 10
201 mm/min. The reported results, including Young's modulus, ultimate strength, and elongation at break,
202 were the average values of five specimens.

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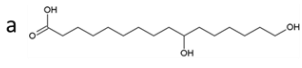
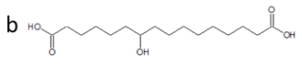
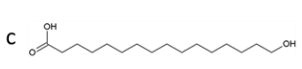
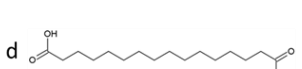
204 **Results and Discussion**

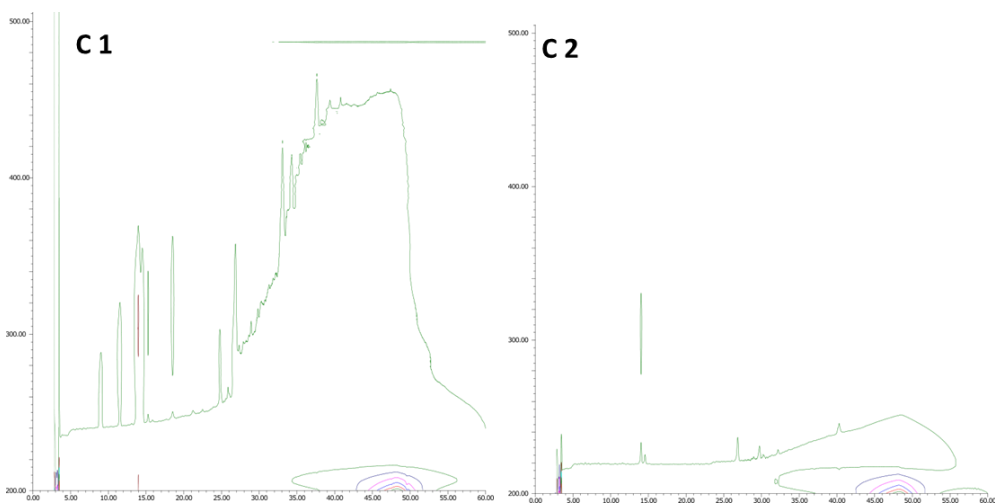
205 ***Cutin monomer purification impacts the polymerization process***

206 The purified tomato CM extracts mainly consist of hydroxy fatty acids as evidenced by FT-IR
207 spectra dominated by intense stretching vibrations of the methylene chains (asymmetric (CH₂) and
208 symmetric (CH₂) at 2919 cm⁻¹, and 2850 cm⁻¹ respectively, bending CH₂ at 722 cm⁻¹) and a sharp band
209 at 1704 cm⁻¹ assigned to carbonyl stretch of carboxylic group. A broad hydroxyl (3300–3500 cm⁻¹) band
210 was also observed. In addition, chemical analysis (Figure 1A) highlighted that the CM extracts
211 concentrated more than 90% of 9- and 10,16 dihydroxyhexadecanoic acid isomers, (hereafter designed
212 as diOHC16), as well as 1.5% of the 16-hydroxyhexadecanoic acid and 5% of the dicarboxylic fatty acids
213 (10-hydroxyhexadecanedioic acid and hexadecanedioic acid). The purification step led to the reduction
214 of several phenolic specific bands in the FT-IR spectra at 1626 cm⁻¹ (stretching of conjugated C=C in
215 aromatics), 1606 cm⁻¹ (stretching band (C-C) aromatic), 1515 cm⁻¹ (stretching of C-C aromatic
216 conjugated with C=C) and 833 cm⁻¹ (out of plane bending of (C-H) aromatic). Likewise, biochemical
217 analyses (Figure 1B) indicated a nearly 10-fold reduction from 45 mg/g (GAE) to 5 mg/g (GAE) while
218 the fatty acid composition remains almost unchanged. Regarding the lab-scale process to reduce the
219 phenolic content of the crude CM extract, some improvements can be made by replacing the non-
220 green and hazardous chloroform by the eco-friendly solvents, cyclopentyl methyl ether (CPME; De
221 Gonzalo et al, 2019) and acetone (supplemental data Figure 1). However, it should be emphasized that,
222 regardless of the solvent, this costly purification process cannot be up-scaled from an industrial
223 perspective. Indeed, the main phenolic fraction, consisting of the corresponding oxidized compounds,
224 is irreversibly bound to the aminopropyl silica gel and cannot be reused. Actually, the lab-scale process
225 was used as a proof-of-concept to highlight the effects and benefits of reducing the phenolic content
226 of the crude CM extracts of tomato pomaces.



B

Cutin monomer extract	crude	purified
a 	94 %	93 %
b 	3,7 %	4,5 %
c 	1,9 %	1,5 %
d 	0,4 %	0,5 %



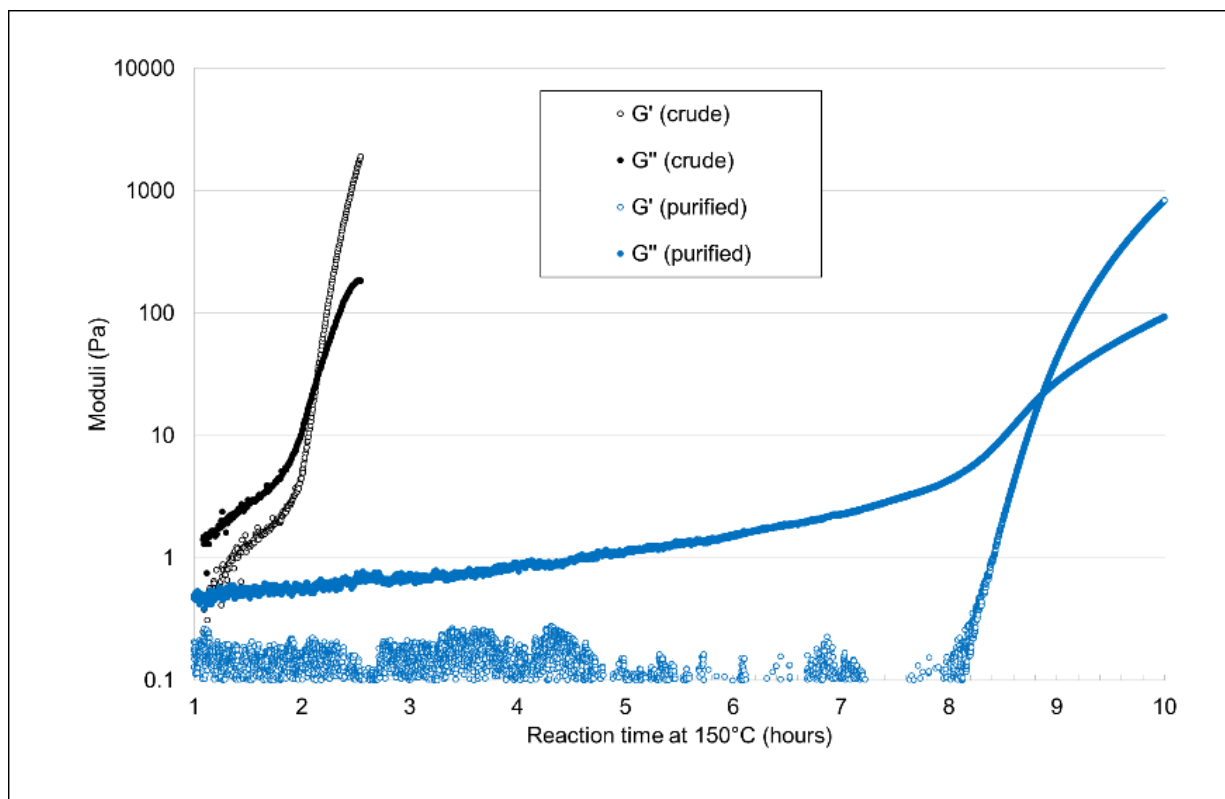
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228

229 Figure 1: (A) FT-IR characterization of the cutin monomer extracts before and after purification. (B)
 230 Lipid composition (% of fatty acids) of the cutin monomers before and after purification a: 9,10-16
 231 dihydroxyhexadecanoic acid; b: 10-hydroxyhexadecanedioic acid; c: 16-hydroxyhexadecanoic acid; d:
 232 hexadecanedioic acid. (C) Total phenolic compounds were determined by Folin Calieui analysis (inset-
 233 data expressed as mg of equivalent gallic acid per g of cutin extract) and monitored by HPLC-DAD.
 234 Contour plot of the crude cutin monomer extract (C1) and CPME-purified cutin monomer (C2).

235 Thermal polycondensation resulted in a light-brown solid material insoluble in water and
236 chloroform. Assuming that the contribution of the traces of phenolic compounds present in the
237 purified CM fraction to the polymerization can be neglected, only the fatty acid building blocks are
238 involved in the reaction. The results of the rheological tests show a strong impact of the purification
239 on network formation kinetics during polymerization. Figure 2 shows the evolution of the G' and G''
240 moduli monitored during isothermal polycondensation at 150°C for the purified and crude CM
241 fractions. Initially, both samples display a viscous liquid behavior ($G'' \gg G'$). However, gelation
242 phenomena take place during the reactions, leading to a viscoelastic solid behavior ($G' > G''$). The
243 modulus crossover time ($G' = G''$) gives a rough estimate of the gel time, which is likely to be shorter
244 during the oven polymerization of films both because of the presence of vacuum at the beginning, and
245 the larger free surface area, which favors the removal of the water molecules formed during the
246 polyesterification reaction, thus accelerating it. However, while the modulus crossover occurs at $\approx 2.2\text{h}$
247 for the crude fraction, it is observed after 9h for the purified one, which represents a 4-fold increase.
248 Such a huge difference suggests that the gelation times which are thought to be shorter during the
249 oven polymerization experiments, are still very different. Therefore, the following conclusions can be
250 drawn: i) The presence of gelation phenomena for purified CM demonstrates that the mere presence
251 of 5% dicarboxylic acids is sufficient to lead to a crosslinked polyester network by co-polymerization
252 with the main diOHC16 fatty acid component ii) The approximately 4-fold shorter gelation time
253 observed for the CM extract suggests that the presence of phenolic compounds participates in side
254 reactions inducing additional crosslinking. Therefore, it was decided to further focus on the polyester
255 obtained from the purified CM fraction.

256



257

258 Figure 2: Evolution of the moduli G' and G'' during polymerization at 150°C for the crude and purified
259 CM fractions.

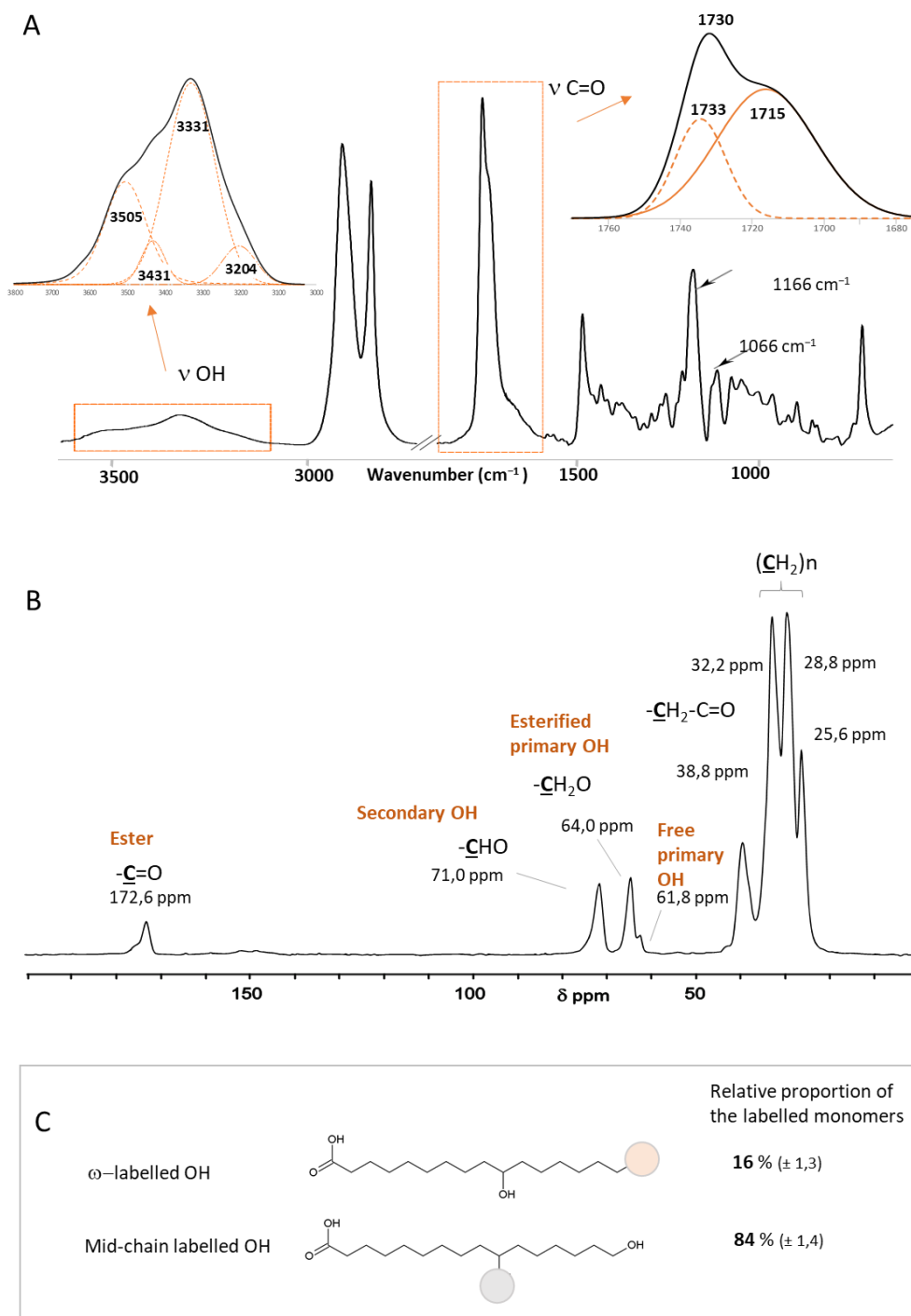
260 ***Ester and hydrogen bonds govern the polymer structure***

261 The structure of the polymer determined by ATR-FTIR and ^{13}C SSNMR (Figure 3) highlighted
262 typical polyester features. Indeed, the FTIR spectrum showed the characteristic shift of the carbonyl
263 band from 1704 cm^{-1} (Figure 1A), assigned to the carboxylic acid of the monomers, to the band around
264 1730 cm^{-1} assigned to the carbonyl ester stretching vibration ($\nu_s\text{ C=O}$) (Figure 3A)³³. The polymer also
265 displayed C-O-C ester stretching bands (1166 and 1106 cm^{-1} respectively). The $\nu_s\text{ C=O}$ band can be
266 deconvoluted into two bands at 1731 and 1715 cm^{-1} . The latter is generally assigned to carbonyl
267 involved in hydrogen bonds (H-bonds) with an hydroxyl group while the former is considered to be
268 free carbonyl group, i.e., not involved in hydrogen bonds³⁴. This band splitting has also been observed
269 in plant cuticles^{29,33}. In cuticles this lowest wavelength $\nu_s\text{ C=O}$ band is attributed to interactions of the
270 cutin polyester with cuticle-embedded polysaccharides and non-esterified hydroxy fatty acids. In this
271 study, the H-bonds can only involve the non-esterified hydroxyl group of the hydroxy-fatty acid
272 monomer units of the polyester. The vibration at 1715 cm^{-1} is broader than the band at 1733 cm^{-1}
273 indicating a relatively wide distribution of hydrogen bond distances and geometries in the polyester.
274 The H-bonds can also be observed in the large OH stretching vibrations in the $3000\text{-}3500\text{ cm}^{-1}$ region.
275 Four major peaks can be highlighted after deconvolution, i.e. at 3204 , 3331 , 3431 and 3505 cm^{-1} (Figure
276 3) in agreement with previous results on hyperbranched polyester polyol³⁵. These different
277 deconvoluted peaks can be attributed to different types of hydrogen bonding between C=O and OH
278 groups ($\text{C=O}::\text{H-O}$) and between OH groups ($\text{O-H}::\text{O-H}$)^{36,37}. The three deconvoluted peaks at 3204 ,
279 3331 , 3431 cm^{-1} are due H-bonding of different strengths, i.e., the lowest OH wavelength is related to
280 the strongest H-bonding.

281 Similarly, ^{13}C SSNMR show a broad peak at 172.6 ppm assigned to esterified carbonyl (Figure
282 3B) and a sharp peak at 64 ppm , which is assigned to primary esters²². The presence of a smaller peak
283 at 61.8 ppm indicates non-esterified primary hydroxyl group. Spectral deconvolution allowed us to
284 estimate at $72 \pm 1\%$ the proportion of esterified chains. According to the chemical structure of the
285 diOHC16 including mid-chain and ω -hydroxyl groups, both, linear and branched ester bonds, can be
286 formed in the polyesters. The esterification scheme of the polyester was further monitored by the
287 chemical labeling of the free OH groups within the diOHC16-derived polyesters³¹. After complete
288 depolymerization, the release of diOHC16 containing labeled OH groups either in ω -position or in the
289 midchain-position was compared (Figure 3C). Within these labeled monomers, the relative proportion
290 of labeled mid-chain OH/ ω -OH labeled in diOHC16 indicated that the free OH groups were mostly
291 (84%) in the mid-chain position. This indicates that the two OH groups of the ABB monomer do not
292 have the same reactivity, favoring the formation of "head to tail" linear sequences and limiting the
293 formation of side pendant chains.

294 Finally, the polymer was subjected to alkaline hydrolysis. The product obtained after alkaline
295 hydrolysis display the same fatty acid monomer composition (Supplemental Figure 2A) and its size
296 distribution superimposed on that of the purified CM fraction (Supplemental Figure 2B), indicating the
297 presence of ester linkages in the polymer. This ester structure is an advantage for the recyclability of
298 this bio-based polymer.

299 All these data indicate that the structure of the polymer is controlled by both ester and
300 hydrogen bonds. The polyester contains both linear and branched polymer chains of different masses
301 with a preferential linear polymerization scheme. Since the polyester is reticulated, the minor
302 dicarboxylic fatty acids (Figure 1B) are most likely involved in the polymer cross-linking.



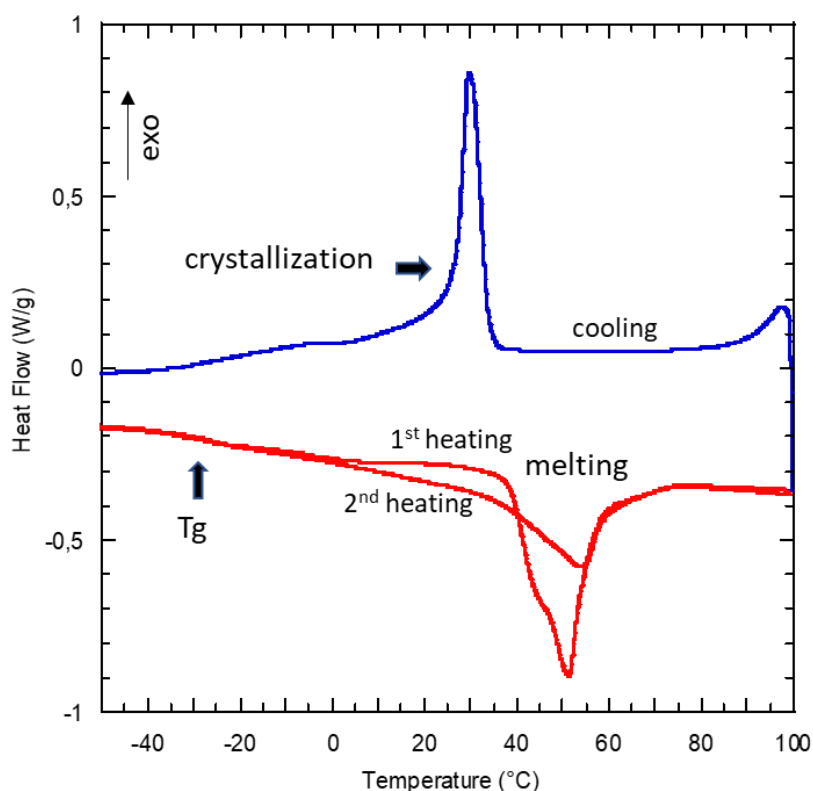
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304 Figure 3 : (A) ATR FTIR spectra of polymer produced from the purified cutin monomer. Typical
 305 polyester bands (arrows) were observed at 1730 cm^{-1} (stretching of esterified C=O), 1166 cm^{-1} (asym
 306 stretching of the C-O-C ester bound) and 1106 cm^{-1} (symetric stretching of the C-O-C ester bounds). In
 307 inset, a magnification of the OH stretching bands involved in different hydrogen bond and of the CO
 308 stretching bands at 1730 and 1715 cm^{-1} assigned respectively to ester group and ester group involved
 309 in hydrogen bond. (B) Solid state ^{13}C CP-MAS spectrum of the polyester. (C) Inset- Relative proportion
 310 of free OH groups within the polyester at the ω - and mid-chain position of the the diOHC16. Values
 311 are means of at least three replicates (\pm standard deviation).

312

313 **Thermal transitions highlight the semi-crystalline structure and the network of the polyester**

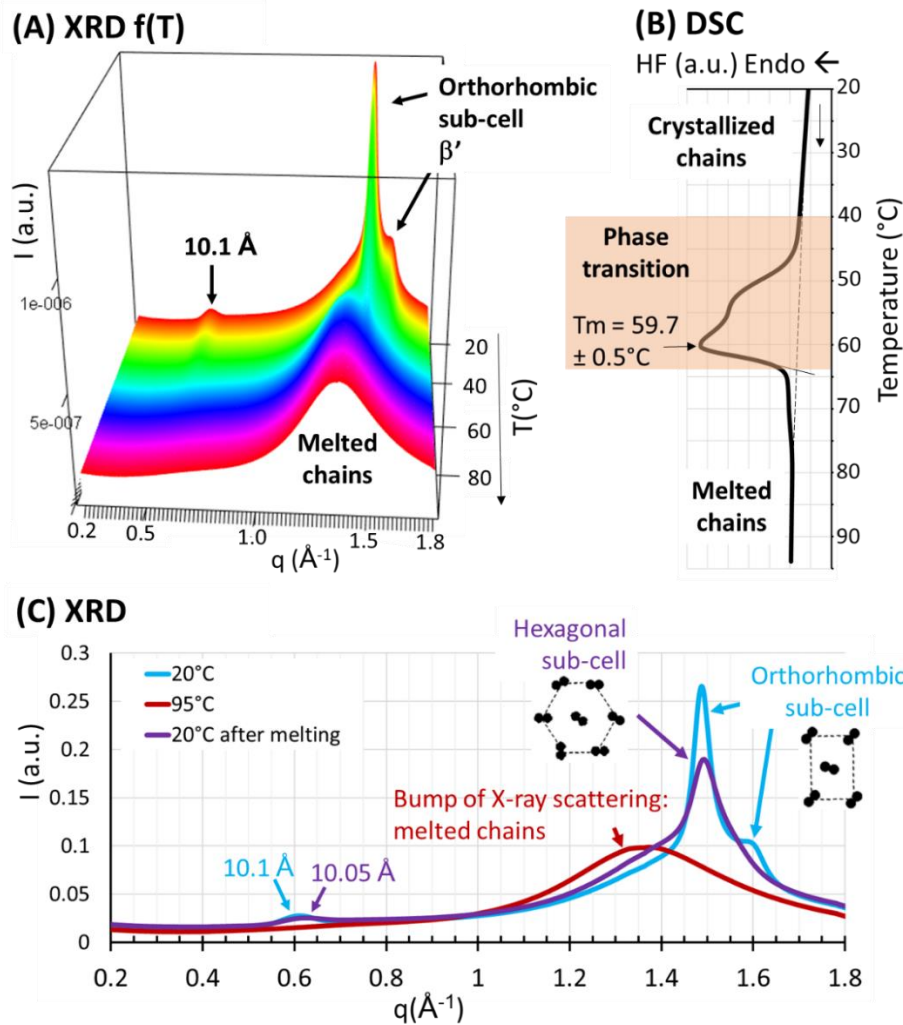
314 Figure 4 shows the thermograms obtained by DSC on a first heating of the polyester, on the
315 cooling immediately followed by a second heating. At low temperatures, the thermogram show a glass
316 transition (T_g) at -25°C , which is determined at the midpoint of the baseline fall. At higher
317 temperatures, the thermogram of the first scan shows a large melting endotherm beginning at about
318 38°C and with a maximum at 51.2°C . Above 60°C the transition appears to continue slowly and the
319 baseline recovers at about 80°C . On cooling, the DSC thermogram shows an exotherm with a maximum
320 at 30.0°C , attributed to the crystallization of the polyester chains. The second heating obtained after
321 cooling shows a broader melting endotherm that seems to start at around 10°C . These results suggest
322 an aging during the storage of the polymer which allows crystallization. This is confirmed by a melting
323 enthalpy of 61.8 J/g for the first heating and 41.1 J/g for the second heating. In order to « standardize »
324 the conditions of the study, the calorimetric, mechanical and thermomechanical experiments were
325 carried out on samples stored for 7 days at 20°C after heating above the melting temperature. The
326 thermal stability was studied by thermogravimetric analysis (supplemental Figure 3). A very low weight
327 loss (less than 0.5%) below 100°C probably corresponds to moisture evaporation. Thermal degradation
328 occurs above 300°C which is similar to other bio-sourced polyesters such as PBS. According to the
329 calorimetric results, we further investigated the structure of the crystal domains by X-ray diffraction.



330
331 Figure 4: Thermograms obtained by DSC on a first melting, cooling and second melting of the polyester.

332 The XRD patterns recorded at 20°C revealed that the polyester exhibited a crystalline phase
333 with a lateral packing of the chains in the orthorhombic sub-cell (β' polymorphic form), characterized
334 by two peaks at 1.4900 and 1.5977 \AA^{-1} (4.2 and 3.9 \AA , respectively) (figure 5A). A peak was recorded at
335 0.6208 \AA^{-1} (10.1 \AA). Since the carbon-carbon distance in an aliphatic chain is 1.27 \AA ³⁸, the chain length

336 of an acyl chain containing 16 carbon atoms, such as the diOHC16 molecules, was calculated to be
337 19.05 Å. The peak at about 10 Å was interpreted as a longitudinal organization of the chains with a
338 repeat distance corresponding to the half chain length of the acyl chains of diOHC16 in the crystallized
339 state. On heating of the polyester at 5°C.min⁻¹, the diffraction peaks corresponding to both the lateral
340 packing and the longitudinal organization of the chains, progressively decreased in intensity between
341 about 40°C and 60°C. This was interpreted as the melting of the acyl chains as a function of the increase
342 in temperature. For temperatures above 60°C, the bump of X-ray diffusion centered at 1.3694 Å⁻¹ (4.58
343 Å) was attributed to the acyl chains in their molten state, in agreement with the literature³⁸. The DSC
344 thermogram recorded at the same heating rate (figure 5B) showed a broad endotherm with T_m = 59.7
345 ± 0.5°C, and ΔH_m = 73.9 ± 1.9 J.g⁻¹ polyester (means calculated from 3 independent samples). This
346 endotherm was similar to the endotherm recorded in the temperature range from -50 to 100°C (figure
347 4). After rapid cooling of the polyester from the melt (95°C), the wide-angle XRD signal revealed the
348 formation of an hexagonal packing of the acyl chains (α polymorphic form) characterized by a single
349 peak at q = 1.4926 Å⁻¹ (d = 4.2 Å), and a longitudinal organization (d = 10.5 Å). At 20°C, the freshly
350 melted polyester was in a hexagonal packing (α polymorphic form) while the « aged » sample, after 7
351 days at room temperature was in an orthorhombic β' polymorphic form. Differences in the packing of
352 the chains as a function of the thermal history of the polyester revealed a polymorphism with a variable
353 metastability of the crystallization state (figure 5C). Regardless of the aging of the samples and the
354 polymorphic form of the crystallized acyl chains, the peak at about 10 Å corresponding to the
355 longitudinal organization of the chains was recorded at 20°C.

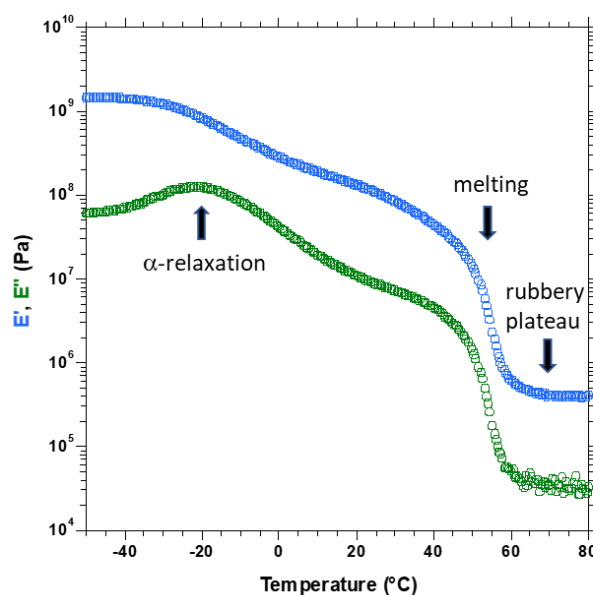


356

357 Figure 5: (A) 3-dimensional representation of XRD patterns recorded as a function of temperature on
 358 heating of the polyester from 20°C to 95°C at 5°C.min⁻¹ ; (B) Thermogram recorded on heating of the
 359 polyester from 20°C to 95°C at 5°C.min⁻¹ ; (C) XRD patterns recorded at various temperatures as
 360 indicated in the figure.

361 The characterization of dynamic properties by DMTA has been realized in order to investigate
 362 the amorphous state. Figure 6 shows thermomechanical storage (E') and loss (E'') moduli, measured
 363 by DMTA. At low temperature, the storage modulus of the material was typical of a glassy state ($E' \approx 1$
 364 GPa at -50°C). Its first drop between -40°C and 0°C, coinciding with the E'' peak centered around -20°C
 365 corresponds to the main mechanical α -relaxation associated with the calorimetric glass transition from
 366 glassy to rubbery state determined by DSC (figure 4). The storage modulus value obtained above this
 367 transition is maintained at a rather high value ($E' \approx 100$ MPa at 20°C) due to the presence of the
 368 crystalline fraction previously shown. The thermomechanical measurements showed a dramatic
 369 second drop in the moduli starting from about 45°C, due to the melting of this crystalline part of the
 370 material, leading to a plateau in the storage modulus. Its average value of $E' \approx 0.4$ MPa at 70°C, is
 371 typical of a weakly crosslinked network in the rubbery state.

372 0



373

374 Figure 6: Plot of the storage and loss modulus (E' , E'') of the polyester measured by DMTA

375

376 ***Towards a model of the polyester network***

377 In our previous work on the polymerization of the crude fatty acid ⁷, we used Flory's theory of
 378 rubber elasticity ²³ in order to estimate the crosslink density of the network from the E' modulus
 379 measured by DMTA at the rubbery plateau. The same approach can be applied to the polymer
 380 networks obtained in the present work by polymerization of the purified fatty acid. The crosslink
 381 density ν (mole.m⁻³) is given by:

$$\nu = \frac{E}{3RT}$$

382

383 Where: R is the gaz constant 8.32 J.mol⁻¹.K⁻¹, T is the temperature in Kelvin and E the rubbery state
 384 elastic modulus at small deformation.

385 The modulus E' measured by DMTA at the rubbery plateau at 70°C (above the melting temperature) is
 386 equal to 0.4 MPa, which leads to a crosslink density of 47 mol.m⁻³. This value is 16-fold lower than that
 387 of the networks obtained by polymerization of the crude fatty acid extract (728 mol.m⁻³) ⁷, confirming
 388 that when present, the phenolic compounds significantly contribute to additional crosslinking
 389 reactions. The identification of such reactions may be the subject of future studies.

390 Figure 7 gives a model representation of the network formed by the polymerization of the
 391 purified CM extract. As indicated, "A" stands for an acid function and "B" for and hydroxyl function,
 392 that can react to form ester bonds, notated "AB". For the sake of clarity, only 2 types of building blocks
 393 are used: monoacidic monomers abbreviated ABB (standing for the main diOHC16 monomer, plus the
 394 16-hydroxyhexadecanoic acid minor compounds), and dicarboxylic monomers abbreviated A-A
 395 (standing for both the 10-hydroxyhexadecanedioic acid and the hexadecanedioic acid monomers). The

396 proposed structure of the elastically active chains between crosslinks is based on the following data
397 and assumptions:

398 • The content of ABB and A-A building blocks in the chains corresponds to the composition of
399 the refined CM extract (Figure 1B), i.e. approximately 95% and 5%, respectively.

400 • The average molecular mass M_c (g.mol⁻¹) of an elastically active chain can be estimated
401 from the Flory model:

$$M_c = \frac{\rho}{\nu}$$

402

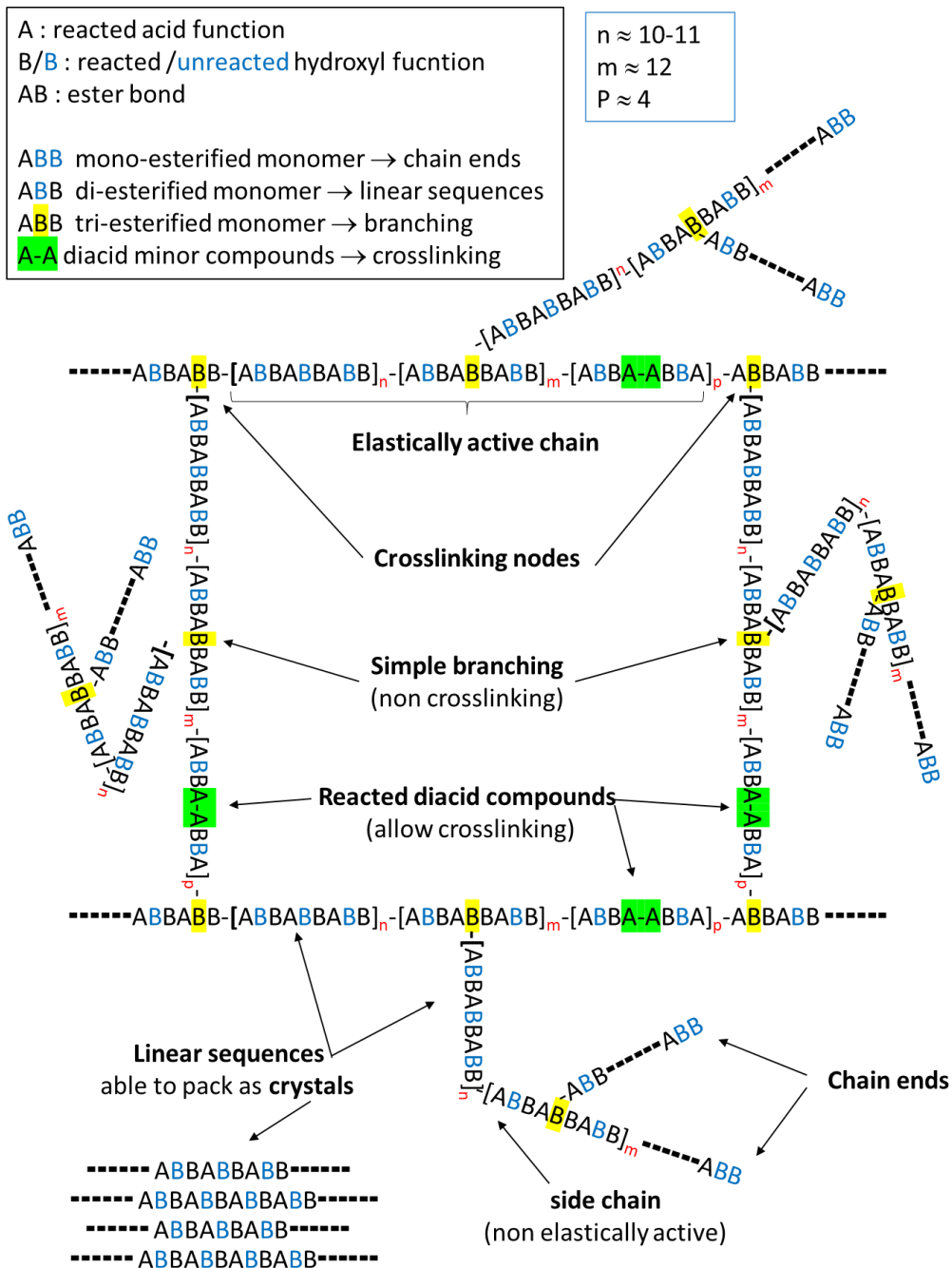
403 Assuming a density $\rho = 1 \text{ g.cm}^3$, as a first approximation, this gives $M_c \approx 21.4 \times 10^3 \text{ g.mol}^{-1}$.

404 • The number of repeating units in the chain can be estimated by dividing M_c by the mass of
405 a repeating unit: $M_0 \approx 270 \text{ g.mol}^{-1}$ (corresponding to an ABB molecule, when A and B end functions
406 are reacted). This gives ≈ 80 repeating units, i.e. ≈ 4 A-A and ≈ 76 ABB.

407 • Due to the low reactivity of the secondary α -hydroxyl groups (Figure 3C) of ABB building
408 blocks, the majority of the ester bonds are formed by reactions between acid groups and primary ω -
409 hydroxyl groups, leading to linear sequences. However, a significant fraction of the ester bonds
410 involves secondary α -hydroxyl groups, leading to side chains, and contributing to crosslinking by
411 copolymerization with A-A building blocks.

412 Indeed, this proposed model highlights a key role of the minor dicarboxylic fatty acid compounds (A-
413 A) in the formation of the three-dimensional network by connecting linear parts of branched chains.
414 Figure 7 shows that the elastically active chains resulting from these connections are delimited by
415 crosslinking nodes. Pendant chains are connected to the elastically active chain by simple branching
416 points. In this structure, long linear sequences -ABBABBABBABBABB- are assumed to be able to pack
417 as crystals (parallel packing of different elastically active chains), resulting in the semi-crystalline
418 structure evidenced by X-Ray experiments.

419



420

421 Figure 7: Model representation of the polyester network

422

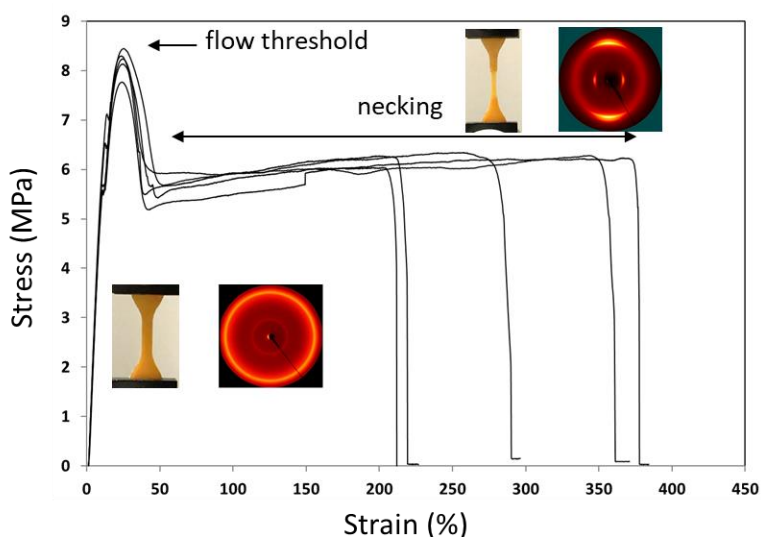
423 **Mechanical properties exhibit a plastic deformation and necking phenomena**

424 Figure 8 shows a superposition of tensile stress–strain curves for the cross-linked polyester.
 425 The average mechanical properties are a Young’s modulus of 72 MPa, a maximum strength of 8.2 MPa
 426 and an elongation at break of 286%. The superposition of the curves shows a good reproducibility in
 427 the mechanical behavior of the specimens. An elastic domain is observed until 25% strain where a
 428 maximum stress is measured at about 8 MPa. A flow threshold occurs at higher deformation
 429 accompanied by the formation of a distinct visible neck, a load drop in the stress-strain curve, and neck

430 propagation at constant stress of about 6 MPa. This corresponds to a plastic deformation until rupture.
431 The necking corresponds to a deformation propagating at a constant rate along the axis of stretching
432 ³⁹. Macroscopically the deformation is heterogeneous, the thickness of the sample decreases
433 dramatically in a localized domain. Necking is rarely reported for bio-based polymers but is well known
434 for synthetic polymers such polyethylene where it corresponds to the orientation of the polymer
435 chains ⁴⁰. In this study, the X-ray scattering anisotropy, which presents diffraction spots on equatorial
436 and polar position, is due to the orientation of the crystallized polyester particles parallel to the
437 direction of the deformation. The isotropic X-ray scattering signal recorded before stretching of the
438 polyester is due to the random arrangement of the polyester crystal particles, with the peak at 10 Å
439 and the lateral packing of the chains recorded at wide angles. DSC measurements indicate the same
440 crystal concentration before and after stretching the sample. This mechanical behavior of wide plastic
441 deformation observed in the purified polyester is completely different from the elastomeric properties
442 observed in the crude polyester ⁷. The presence of a low level of crosslinks induces crystallinity that
443 increases tensile strength. The extent of deformation is accompanied by a crystal orientation.
444 Moreover, it has been observed that the heating at 75°C of the sample elongated before breaking,
445 provokes the recovery of its initial length. It means that the deformation is thermally reversible
446 allowing shape memory properties.

447 The mechanical and thermal properties are compared with those of some common polymers
448 and biodegradable aliphatic polyesters in the supplementary table based on bibliographic data^{41 42 43}.
449 The mechanical properties of cutin polyester combined with its remarkable necking behavior during
450 deformation, are comparable to those of LDPE. In terms of glass transition, the T_g of the polyester is
451 in the same order of magnitude as that of other biodegradable polyesters such as PBS. On the other
452 hand, the melting temperature of the crystal fraction remains low at 51°C but it should be noted that,
453 at this temperature, there is no flow and the material retains its initial shape.

454

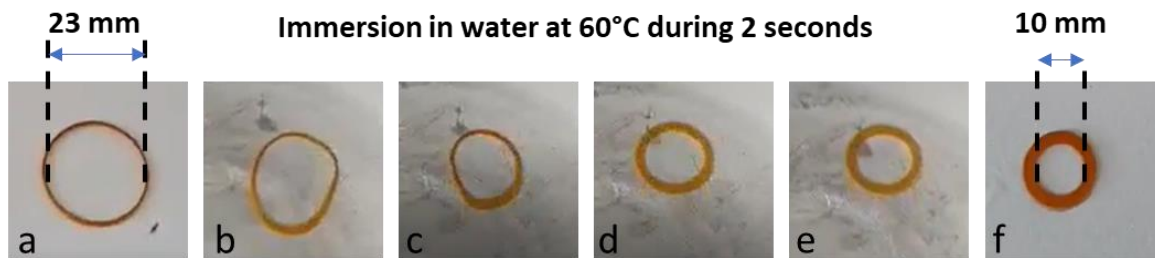


455

456 Figure 8: Stress-strain curves obtained on tensile of polyester. Inserted images show the initial
457 dogbone shape of material and necking phenomena with a juxtaposition of two-dimensional WAXS
458 diagrams.

459 **Shape memory properties are consistent with a crosslinked semi-crystalline structure**

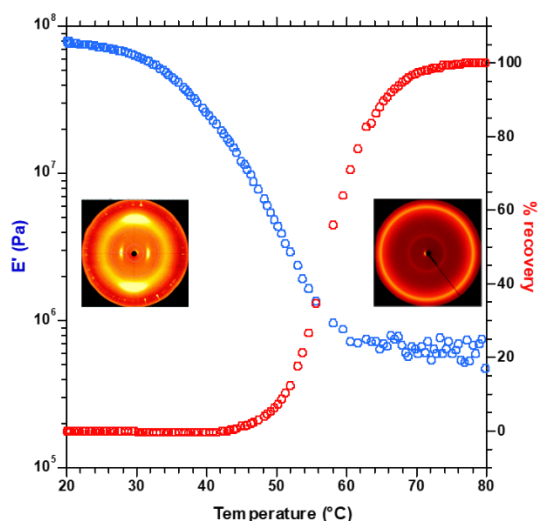
460 The polyester material exhibited shape memory properties which seems logical given the
461 slightly crosslinked semi-crystalline structure. Shape memory behavior of a polymer is the ability of a
462 material to change its shape from a temporary to a permanent shape at a given temperature. To
463 demonstrate this effect, a sample in the shape of ring, cut in the thick film (0.7 mm), was heated and
464 stretched at 75°C, above the melting temperature of the polyester, and cooled rapidly to 10°C while
465 maintaining this shape (Figure 9a). Later, the deformed sample was immersed in a water bath at 60°C
466 (Figure 9b to 9e) and the deformed sample recovered its original shape in 2 seconds (Figure 9f).



467

468 Figure 9: Shape recovery of a polyester ring immersed in water at 60°C

469 Shape recovery can be assessed by DMTA by measuring accurately the evolution of the sample
470 length simultaneously with its modulus (Figure 10). The permanent shape is a small dogbone with an
471 effective length of 3.6 mm. It was previously elongated at 75°C by 100% (7.2 mm length) and quenched
472 at room temperature, in order to obtain the temporary shape. WAXS measurements have shown quite
473 similar features as in the case of necking. The spots observed in the 2D diagram show a strong crystal
474 orientation in the temporary shape in the direction of its deformation. During heating, the sample
475 length recovery starts at 45-50°C, to reach 100% recovery at 70-75°C. Shape recovery is simultaneous
476 with the melting of the crystal previously detected by DSC in the same temperature range. At the same
477 time, the crystal orientation completely disappears as shown by WAXS after cooling. These results
478 show a clear relationship between the slightly crosslinked semi-crystalline structure of the polymer
479 and its shape memory behavior^{44, 45}. The stretching of the polymer chains at a temperature higher
480 than the melting temperature, allows their orientation, the presence of a reticulated network avoids
481 the flowing of the polymer. Freezing under stress leads to oriented crystals in the direction of
482 deformation. At room temperature the structure is maintained by crystals that allow the temporary
483 shape stability. When the material is heated above the melting temperature of crystal domains, the
484 permanent shape is spontaneously recovered driven by the reticulated elastic entropy of the extended
485 and reticulated chains. Similar shape memory properties have been observed on dogbone elongated
486 at room temperature and showing necking phenomena (result not shown). This observation makes it
487 possible to envisage a cold programming of the shape memory



488

489 Figure 10: Storage modulus E' and percentage of recovery measured by DMTA during shape recovery
 490 of polyester with a juxtaposition of two-dimensional WAXS diagrams obtained after cooling.

491 It appears clearly that this shape memory behavior is related to the presence of long elastically
 492 active chains between the nodes of the network, that are able to crystallize. Therefore, it may be
 493 affected by variations of the phenolic and diacid impurities contents: For phenolics, the potential
 494 impact may be strong. However, we cannot predict it because we currently do not know the exact
 495 nature of the additional crosslinking reactions that they induce (Future works may elucidate this point).
 496 For diacid impurities, the potential impact can be inferred from the proposed model representation of
 497 the polyester network (Figure 7): Each additional diacid molecule in the initial reaction medium can
 498 potentially lead the formation of one additional elastically active chain by “bridging” two pendent
 499 chains. In other words, assuming that all diacid molecules participate in the formation of elastically
 500 active chains implies that the crosslink density is proportional to the diacid concentration. Calculations
 501 based on this assumption (detailed in supplementary material) lead to the conclusion that for a diacid
 502 concentration range $4 < x < 6 \%$, the variations should be: $38 < \nu < 56 \text{ mol/m}^3$ for the crosslink density,
 503 and $26 > M_c > 18 \text{ Kg/mol}$ for the average mass of elastically active chains, which number of repetitive
 504 units should thus vary within the range $98 > M_c/M_0 > 65$ units. We assume that such variations are
 505 unlikely to affect the ability of the elastically active chains to crystallize, and so the presence of a shape
 506 memory behavior. However, they should affect the crystallinity index and so the tensile mechanical
 507 properties at room temperature. The rubbery storage modulus above the melting temperature will of
 508 course also be affected (in the range $0.32 < E'(70^\circ\text{C}) < 0.48 \text{ MPa}$ according to our calculations).

509

510 Conclusions

511 A fully bio-based shape memory polyester was successfully synthesized without a catalyst,
 512 from a biorefined hydroxy fatty acid extract produced from tomato pomaces. As a proof-of-concept,
 513 this work demonstrates how monomer purification can drastically affects the functional properties of
 514 the polyester, and should be considered as a key parameter in the design of a cutin biorefinery process.
 515 Nevertheless, an up-scalable phenolic and eco-friendly removal process should be further developed
 516 to take advantage of the unique shape memory and mechanical properties of the polyester from an

517 industrial and commercial perspective. Indeed, while amorphous elastomers are produced from the
518 crude hydroxy-fatty acid fraction of tomato peels, the purification step we developed, promotes the
519 production of a semi-crystalline polyester with high mechanical and shape memory properties. By
520 linking the linear parts of a branched structure dominated by the major diOHC16, the minor α , ω -
521 hexadecanedioic acids of the hydroxy fatty acid fraction as well as H-bonds, a 3D network is formed
522 that does not flow above the melting temperature. The linear parts of the chains are long enough to
523 crystallize in large parts and control the mechanical properties at room temperature. The structure
524 resulting from the combination of crystal and elastic network provides shape memory properties. In
525 the future, it seems possible to optimize the mechanical and thermomechanical properties of bio-
526 based polymers by controlling the level of minor components, first in the design of the biorefinery
527 cascade or by valorizing other bio-sourced synthons. Indeed, our findings open a new path for the
528 rational design of multifunctional polyester networks, enabling the future advancement of high-
529 performance materials to meet the expectations of a circular economy of the agro-industrial food
530 chain.

531

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539

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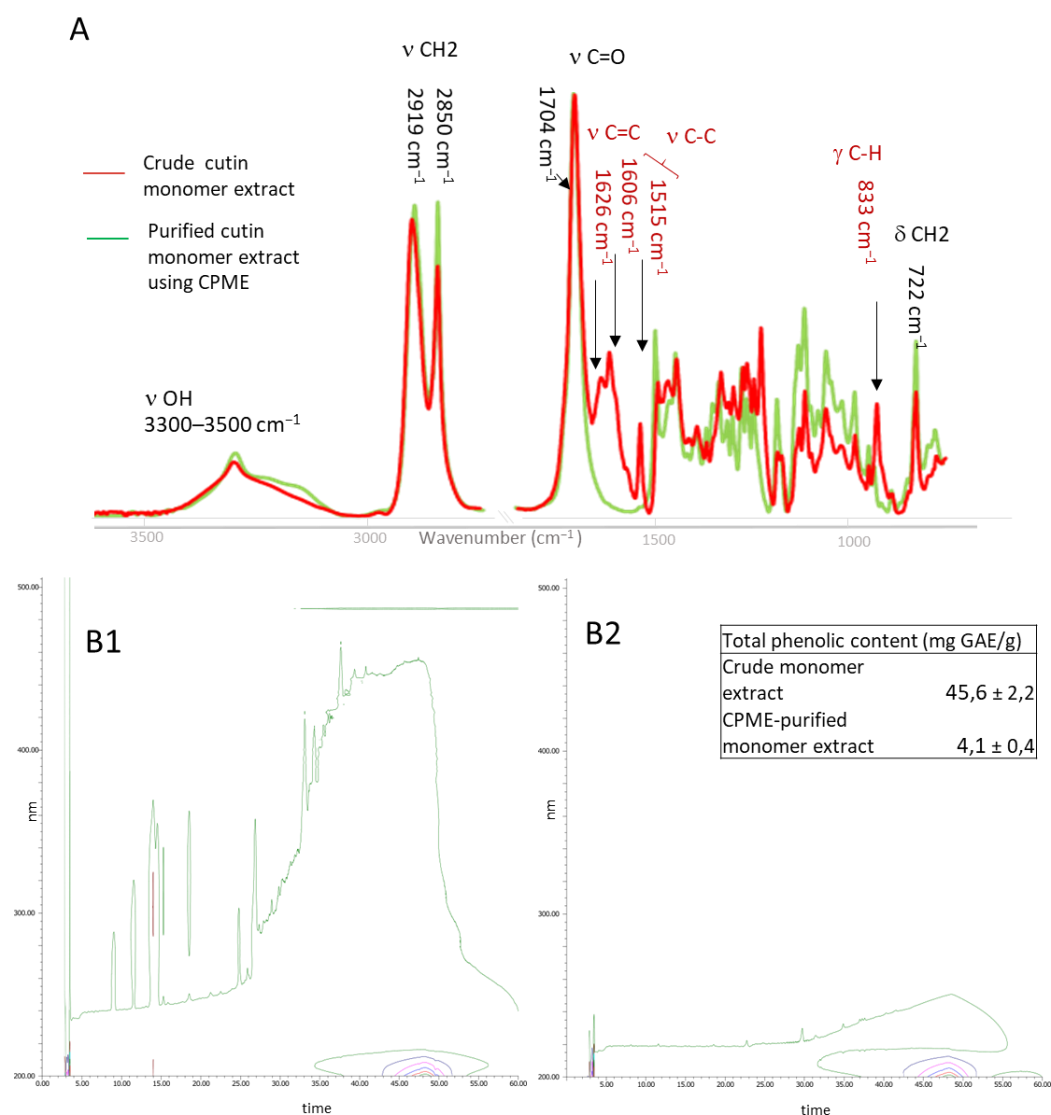
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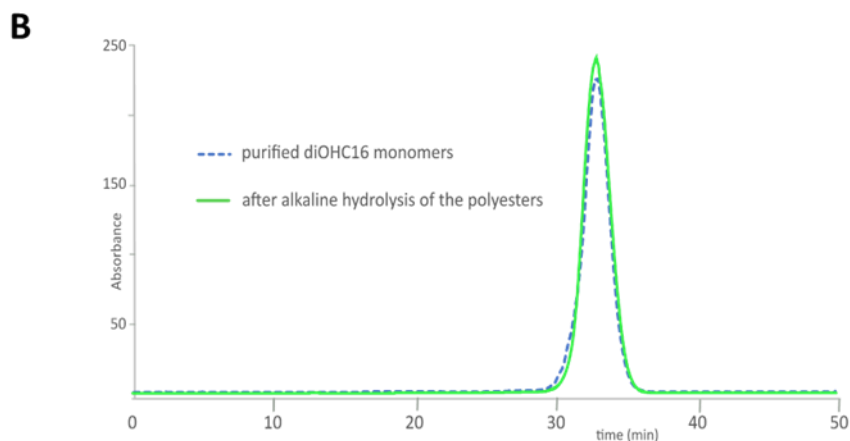
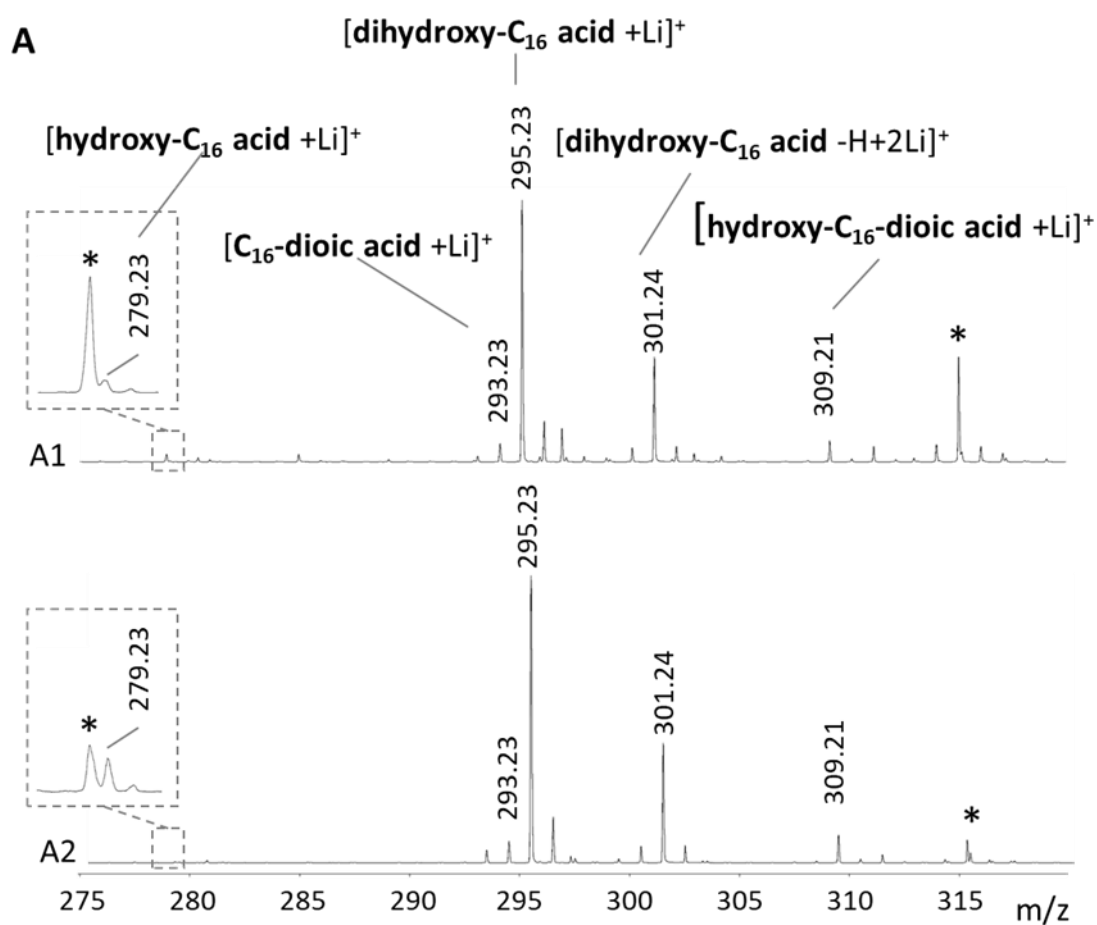
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692 Supplemental figure 1: Characterization of the cutin monomers extracts before and after purification
 693 using green solvent.

694 The same conditions of purification of the cutin monomer were used, except that the chloroform was
 695 replaced by cyclopentyl methyl ether (CPME) and a washing step of the column with Acetone was
 696 added before elution.

697 (A) FT-IR analyses. Spectra are dominated by intense stretching vibrations of the methylene chains
 698 (asymmetric (CH₂) and symmetric (CH₂) at 2919 cm⁻¹, and 2850 cm⁻¹ respectively, bending CH₂ at
 699 722 cm⁻¹) and a sharp band at 1704 cm⁻¹ assigned to carbonyl stretch of carboxylic acid. A broad
 700 hydroxyl (3300–3500 cm⁻¹) band was also observed. All these features are typical signatures of
 701 hydroxy fatty acids. The purification step led to the reduction of phenolic specific bands (red arrows)
 702 at 1626 cm⁻¹ (stretching of conjugated C=C in aromatics), 1606 cm⁻¹ (stretching band (C-C) aromatic
 703), 1515 cm⁻¹ (stretching of C-C aromatic conjugated with C=C) and 833 cm⁻¹ (out of plane bending of
 704 (C-H) aromatic).

705 Phenolic content determined by Folin Calieu analysis (inset- data expressed as mg of equivalent gallic
 706 acid per g of cutin extract) and by HPLC-DAD. Contour plot of the crude cutin monomer extract (B1)
 707 and CPME-purified cutin monomer (B2).

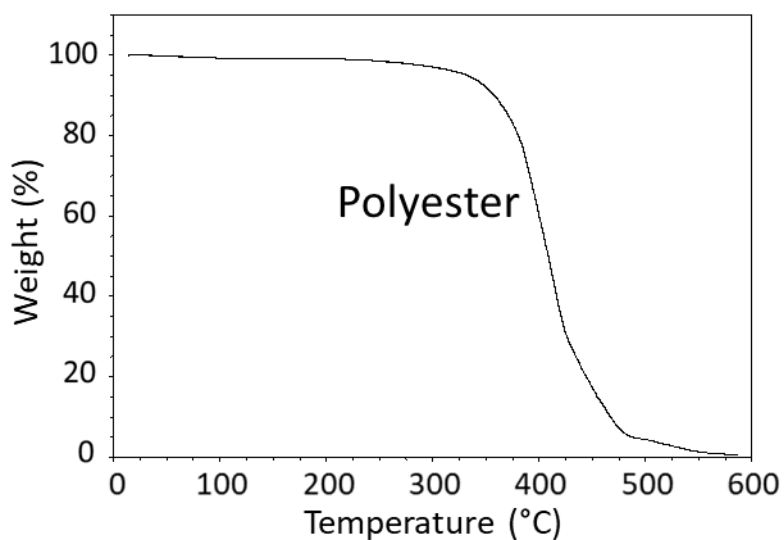


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711 Supplemental Figure 2: Alkaline hydrolysis of the polymer.

712 A. MALDI-MS(+) spectra of the purified cutin monomers used for the polyester synthesis (A1) and after
 713 alkaline hydrolysis of the polyester (A2). Annotations were deduced from exact mass measurements
 714 in the range 275-320 Da. Black star indicates an ion from the MALDI matrix. B. Size Exclusion
 715 chromatography of the purified diOHC16 monomers before polymer synthesis and after polyester
 716 alkaline hydrolysis.



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718 Supplemental Figure 3: Thermogravimetric analysis traces of the polyester

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	Tg (°C)	Tm (°C)	Tensile strenght (MPa)	Elongation at break (%)	ref
PET	70-87	243-268	48-72	20-300	41
HDPE	-125 to -90	135	22-31	100 to >1000	41
LDPE	-125 to -100	112-135	8-31	200-900	41
PP	-10	167-177	31-41	100-600	41
PCL	-60 to -65	150-162	20-42	300-1000	42
PHB	5-15	168-182	40	5-8	42
PHBV	-1	136-162	30-38	20	42
PHBHHX	-1	127	21	400	42
PBS	-32	114	34	560	43
Cutin polyester	-25	51	8.2	286	

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728 Supplemental table : Mechanical and thermal properties of common polymers and biodegradable
729 aliphatic and semi-crystalline polyester, in comparison with cutin polyester

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733 Estimation of the potential impact of diacid impurities content variations:

734 As explained in the text of the article, our calculations are based on the assumption that the crosslink
735 density is proportional to the diacid concentration. Since the current $x=5\%$ leads to a crosslink density
736 of 47 mol/m^3 a variation Δx (%) can potentially lead to a crosslink density:

737
$$v(\Delta x) = \frac{47}{5} \times (5 + \Delta x)$$

738 Therefore, for a range of variation of +/- 1% of diacid, We can expect the following values for crosslink
739 density and related properties:

Diacid %	Crosslink density v (mol/m^3)	Average mass between network nodes M_c (kg/mol)	Average number of repetitive units M_c/M_0	Rubbery modulus above melting temp. $E'(70^\circ\text{C})$ MPa
4	38	26	98	0.32
5	47	21	79	0.4
6	56	18	65	0.48

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741 The variations of the average mass and number of repeat units in elastically active chains are estimated
742 assuming a density $\rho= 1 \text{ g.cm}^3$, and a repetitive unit mass $M_0 \approx 270 \text{ g.mol}^{-1}$. The model of Flory is used
743 to calculate the corresponding rubbery storage modulus values.

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