



HAL
open science

Simultaneous determination of neutral sugars, alditols and anhydrosugars using anion-exchange chromatography with pulsed amperometric detection: Application for marine and atmospheric samples

Amel Nouara, Christos Panagiotopoulos, Richard Sempere

► To cite this version:

Amel Nouara, Christos Panagiotopoulos, Richard Sempere. Simultaneous determination of neutral sugars, alditols and anhydrosugars using anion-exchange chromatography with pulsed amperometric detection: Application for marine and atmospheric samples. *Marine Chemistry*, 2019, 213, pp.24-32. 10.1016/j.marchem.2019.05.002 . hal-02140886

HAL Id: hal-02140886

<https://hal.science/hal-02140886>

Submitted on 27 May 2019

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

1 **Simultaneous determination of neutral sugars, alditols and anhydrosugars**
2 **using anion-exchange chromatography with pulsed amperometric**
3 **detection: Application for marine and atmospheric samples**

4
5
6 **Amel Nouara, Christos Panagiotopoulos*, Richard Sempéré**

7 Aix Marseille Univ., Université de Toulon, CNRS, IRD, MIO UM 110, 13288, Marseille, France

8
9 *Correspondence : Christos Panagiotopoulos

10 [e-mail : christos.panagiotopoulos@mio.osupytheas.fr](mailto:christos.panagiotopoulos@mio.osupytheas.fr)

11
12
13
14
15 **Keywords:** Anion-exchange chromatography; carbohydrates; suspended atmospheric
16 particles, marine particulate organic matter, high molecular weight dissolved organic matter

17
18
19
20 Final version

21 09th May 2019

24 **Abstract**

25 An improved high-performance anion-exchange chromatography with pulsed
26 amperometric detection (HPAEC-PAD) method is described for the simultaneous
27 determination of neutral sugars (hexoses, pentoses and deoxysugars), alditols and
28 anhydrosugars commonly found in atmospheric and marine samples. The method uses a
29 CarboPac MA1 column, at a flow rate of 0.3 mL min^{-1} , and a NaOH gradient (250-700 mM).
30 The proposed method applies a temperature gradient (from 25 to 28 °C) to the column for the
31 first 30 min of the analysis, followed by a constant temperature at 28 °C until the end of the
32 analysis. These analytical conditions allowed the separation of 15 out of 17 carbohydrates in
33 75 min with resolution factors better than 0.5 for the critical pairs levoglucosan/arabitol,
34 galactosan/arabinose, arabinose/mannose and glucose/xylose.

35 The application of this method to field samples revealed that anhydrosugars represented
36 53% of total neutral carbohydrates (TCHO) in total suspended atmospheric particles (TSP),
37 whereas they were detected for the first time in marine particulate organic matter (POM) and
38 high-molecular-weight dissolved organic matter (HMWDOM) samples accounting 2% and
39 3% of TCHO, respectively. Levoglucosan and/or galactosan were the major anhydrosugars in
40 all samples however, their concentrations are undoubtedly underestimated because hydrolysis
41 was applied to the marine samples prior to the HPAEC-PAD analysis. Despite this
42 underestimation, their presence in the marine samples clearly indicates possible terrestrial
43 input most likely via atmospheric deposition as these compounds are considered terrestrial
44 burning biomass tracers. Finally, deoxysugars were also detected for the first time in the TSP
45 sample representing 1% of TCHO, while alditols accounted for 0.4% and 0.3% of TCHO in
46 POM and HMWDOM, respectively.

47

48

49 **1. Introduction**

50 Carbohydrates are among the most abundant organic molecules on the earth because they
51 are detected in all marine and terrestrial ecosystems (Giorio et al., 2018; He et al., 2010;
52 Mopper, 1977; Mopper et al., 1980; Panagiotopoulos and Sempéré, 2005a; Repeta et al.,
53 2015; Theodosi et al., 2018). The determination of carbohydrates at the molecular level has
54 been often used to determine different biogeochemical signatures such as terrestrial inputs in
55 coastal marine environments (Cowie and Hedges, 1984; da Cunha et al., 2002;
56 Panagiotopoulos et al., 2014), organic matter sources (He et al., 2010; Panagiotopoulos et al.,
57 2012; Wicks et al., 1991), and decomposition pathways (Amon et al., 2001; Girollo et al.,
58 2003; Hedges et al., 1994; Opsahl and Benner, 1999).

59 Previous environmental studies on atmospheric suspended particulate matter (PM_{2.5-10})
60 showed that carbohydrates account for 0.2–3% and 0.7–11% of organic carbon (OC) and
61 water-soluble organic carbon (WSOC), respectively (Theodosi et al., 2018; Tsai et al., 2015;
62 Yang et al., 2005). The dominant carbohydrates reported in aerosols are glucose and sucrose
63 released by terrestrial plants (Medeiros et al., 2006; Samaké et al., 2019), levoglucosan,
64 galactosan and mannosan from biomass burning processes (Bhattarai et al., 2019; Simoneit et
65 al., 2004, 1999), as well as mannitol and arabitol from airborne fungal spores and/or various
66 vascular plants (Bauer et al., 2008; Samaké et al., 2019). Other minor carbohydrates reported
67 in aerosols include some neutral sugars (e.g., arabinose, fructose, galactose, and mannose) and
68 alditols (e.g., xylitol and sorbitol; Barbaro et al. 2015; Theodosi et al. 2018). It is worth noting
69 that fucose and rhamnose, two important deoxysugars consistently found in bacterial
70 heteropolysaccharides (Girollo et al., 2003), soil organic matter (Bock et al., 2007),
71 freshwater ecosystems (Cheng and Kaplan, 2003) as well as vascular plants (Opsahl and
72 Benner, 1999) have never been reported in atmospheric samples.

73 Monomeric constituents of carbohydrates after acid hydrolysis (e.g., fucose, rhamnose,
74 arabinose, galactose, glucose, mannose and xylose) have also been detected in the marine
75 environment. As revealed by chromatographic techniques, these monosaccharides account for
76 10–15%, <10% and <20% in particulate, dissolved and sedimentary organic carbon,
77 respectively (Panagiotopoulos and Sempéré, 2005a and references therein). These
78 monosaccharides have also been reported in marine high-molecular-weight dissolved organic
79 matter (HMWDOM) at equimolar concentrations although ¹H nuclear magnetic resonance
80 spectroscopy (NMR) showed the presence of much broader spectra of carbohydrates
81 including unhydrolyzed polysaccharides and methylated sugars (Panagiotopoulos et al.,
82 2007). Other minor carbohydrate classes also reported in marine samples include amino
83 sugars (Benner and Kaiser, 2003; Kaiser and Benner, 2009) and uronic acids (Bergamaschi et
84 al., 1999; Engel and Händel, 2011; Hung et al., 2001).

85 Surprisingly, alditols and anhydrosugars, two major carbohydrate categories consistently
86 found in aerosols and ice cores (Gambaro et al., 2008; Giorio et al., 2018), have scarcely been
87 reported in marine samples despite the growing evidence that dry or wet atmospheric
88 deposition is an important process between the atmospheric and marine compartments
89 (Pantelaki et al., 2018; Zheng et al., 2018). It is worth noting that only Panagiotopoulos et al.
90 (2013) have identified anhydrosugars (levoglucosan, mannosan, and galactosan) in surface
91 marine HMWDOM, in addition to the seven neutral sugars described above. This finding
92 raises questions about the possible sources of anhydrosugars in marine water (terrestrial vs
93 marine), which have been recently examined using compound-specific carbon isotopes
94 analysis (Nouara et al., 2019). Nevertheless, to the best of our knowledge a well-established
95 technique for the measurement of anhydrosugars and alditols does not exist for marine
96 samples. Moreover, the lack of information about deoxysugars in atmospheric samples

97 hinders our understanding of their dynamics and clearly points to the need for a universal
98 method to measure these carbohydrates in any environmental sample.

99 High-performance anion-exchange chromatography with pulsed amperometric detection
100 (HPAEC-PAD) is the most employed chromatographic technique for the analysis of
101 carbohydrates in environmental samples (Benner and Kaiser, 2003; Caseiro et al., 2007;
102 Iinuma et al., 2009; Mopper et al., 1992; Panagiotopoulos and Sempéré, 2005a; Sempéré et
103 al., 2008). However, simple carbohydrates (e.g., monosaccharides) are not ionized in the same
104 manner in alkaline media used in the HPAEC mobile phase, and therefore different
105 concentrations of NaOH must be applied to separate these monosaccharide families (e.g.,
106 alditols, neutral sugars, and uronic acids). Due to this “limitation”, most environmental
107 applications of the HPAEC-PAD technique have focused on one family of carbohydrates
108 (e.g., neutral sugars, or amino sugars) and few attempts have been made to date to separate a
109 much broader spectrum of carbohydrate categories in the same run (Engel and Händel, 2011).

110 The main objective of the present study is to establish a universal HPAEC-PAD method
111 capable of simultaneously identifying several carbohydrate families including neutral sugars,
112 alditols and anhydrosugars commonly found in terrestrial and marine ecosystems. This
113 objective was achieved after the evaluation of the performance of two different anion-
114 exchange columns and the optimization of the mobile phase concentration, column
115 temperature and flow rate. The proposed method was validated in three different
116 environmental matrices, namely, total suspended atmospheric particles (TSP), marine
117 particulate organic matter (POM), and marine HMWDOM.

118 **2. Materials and procedures**

119 *2.1. Reagents*

120 Carbohydrate standards were purchased from Sigma-Aldrich or Interchim, at their purest
121 available grade (>98%). A mixture of standard solutions at concentrations ranging from 50

122 nM to 10 μ M was prepared by dilution with ultrapure water of a standard stock solution (1
123 mM) of neutral sugars (fucose, rhamnose, arabinose, galactose, glucose, mannose, xylose,
124 fructose and ribose), alditols (xylitol, arabitol, sorbitol and mannitol), anhydrosugars
125 (levoglucosan, mannosan and galactosan), and one disaccharide (sucrose). All the prepared
126 solutions including the stock solution were stored in the dark at -25 $^{\circ}$ C until use. Despite that
127 no degradation was observed after melting and refrozen, the standard solutions (50 nM–10
128 μ M) were renewed after 2 weeks. The HCl solution (37%, Sigma-Aldrich) diluted with
129 ultrapure water to 1 M was used for sample hydrolysis, while the NaOH solution (50% w/v in
130 H₂O, low carbonate; PanReac AppliChem ITW Reagents) diluted with ultrapure water was
131 used for the mobile phase of the HPAEC-PAD chromatography. The ultrapure water used in
132 this work was produced by a Millipore Milli-Q system (Molsheim, France).

133

134 2.2. *Sampling and extraction of carbohydrates*

135 2.2.1. *TSP*

136 The aerosol sample was collected on a precombusted (450 $^{\circ}$ C, 6 h) weighed Whatman
137 quartz fiber filter (20.3 cm \times 25.4 cm, nominal retention size, 0.7 μ m) using an automatic
138 sampler (Tisch Environmental USA; flow rate 85 m³ h⁻¹). The sample was collected from
139 March 10th to March 17th of 2016, from the rooftop of the Endoume Marine Station
140 (Marseille; 43 $^{\circ}$ 16' N - 5 $^{\circ}$ 21' E). After collection, the sample was dried for 24 h in a
141 desiccator, weighed, and then stored in a freezer at -25 $^{\circ}$ C in precombusted aluminum foil
142 (450 $^{\circ}$ C, 6 h). A 1-cm² section of the filter was extracted with 6 mL ultrapure water in an
143 ultrasonic bath for 1 h and filtered through a Pasteur pipette packed with quartz wool (both
144 precombusted at 450 $^{\circ}$ C for 6 h) to remove any remaining particles, and then analyzed
145 immediately (Theodosi et al., 2018). The recovery yields of the extraction procedure were
146 estimated after the extraction of a precombusted (450 $^{\circ}$ C, 6 h) Whatman quartz fiber filter

147 spiked with a standard mixture of neutral carbohydrates spanning a concentration range from
148 100 to 1000 nM (Theodosi et al., 2018). The obtained yields of neutral sugars for all the
149 concentrations tested were between 101 and 119% ($n = 5$). The abovementioned analytical
150 procedure allowed the extraction and identification only of the free neutral sugars present in
151 the TSP sample and as such the composition of combined sugars was not determined in this
152 study.

153

154 2.2.2. *Marine POM*

155 Sinking particles (marine POM) were collected over seven-day periods from January 6th to
156 March 3rd, 2013, in the upwelling system located offshore of Lima (Peru) in the Pacific
157 Ocean ($12^{\circ} 02' S - 77^{\circ} 40' W$), using sediment traps (PPS3, Technicap) deployed in the
158 oxycline/upper oxygen minimum zone (OMZ) layer at a depth of 34 m (Bretagnon et al.,
159 2018). To avoid the bio-degradation of POM, a solution of seawater with 5% formaldehyde
160 was added to the bottom of the collection chamber. After trap recovery, the living and dead
161 swimmers were carefully removed so that only detrital particles remained in the sample.
162 These detrital particles (marine POM) were stored in the dark at 4 °C in the initial chambers
163 used in the trap. On land, the samples were filtered through 25 mm precombusted (450 °C, 6
164 h) Whatman GF/F filters, freeze-dried, and subsequently stored in the dark at 4 °C until
165 further analysis. Five portions (40–60 mg each) of each of the five samples obtained from the
166 respective collection chambers of the sediment trap were pooled together, resulting in ~265
167 mg dry POM powder. To release neutral sugars, the POM powder was hydrolyzed with 1 M
168 HCl at 100 °C for 20 h (Panagiotopoulos et al., 2014). The acid-soluble fraction recovered
169 after centrifugation (2000 rpm) was then transferred into a precombusted (450 °C, 6 h) glass
170 vial, and the acid was removed from the sample by three successive lyophilizations. The POM
171 recovery (by weight) after acid hydrolysis was 84%. A part of the final dry powder was

172 weighed (25.61 mg), redissolved in 1 mL ultrapure water and filtered through a Pasteur
173 pipette packed with quartz wool (both precombusted at 450 °C for 6 h). The filtrate was
174 diluted 500 times with ultrapure water and filtered again to remove any remaining particles
175 prior to injection into the chromatographic system.

176

177 2.2.3. *Marine HMWDOM*

178 The surface (15 m) seawater sample was collected from the Hawaii area (19° 43' N - 156°
179 03' W) in December 2003 and processed at Woods Hole Oceanographic Institution (USA) as
180 described by Panagiotopoulos et al. (2013). Briefly, the sample (8640 L) was filtered through
181 0.8 µm and 0.2 µm dual cartridge filters, and then ultrafiltered using a cross-flow
182 ultrafiltration system. The concentrated sample (40 L) that included the HMWDOM was
183 further desalted by diafiltration with ultrapure water to a final volume of 2 L and freeze dried
184 to obtain a fluffy white powder. The carbon composition of the HMWDOM collected was
185 35% by weight, with a carbon/nitrogen ratio of 16. A part of the resultant HMWDOM powder
186 (104 mg) was hydrolyzed to release the neutral sugars with 1 M HCl at 100 °C for 20 h, then
187 the hydrolysis was stopped by placing the sample in an ice bath. The HCl was removed from
188 the sample by lyophilization (3 times; final pH = 7). Then, a portion of the sample (55.83 mg)
189 was redissolved in 1 mL ultrapure water, filtered through a Pasteur pipette packed with quartz
190 wool (both precombusted at 450 °C for 6 h) and stored frozen at -20 °C. Before analysis, the
191 sample was diluted 3 times with ultrapure water.

192

193 2.3. *Chromatographic system*

194 Carbohydrate analysis was performed using a Thermo-Dionex ICS-3000 anion-exchange
195 chromatograph (HPAEC) equipped with a 250-µL injection loop and a pulsed amperometric
196 detector (PAD). The waveform used for pulsed amperometric detection was the standard

197 quadruple potential for carbohydrate analysis (Panagiotopoulos et al., 2012). The ultrapure
198 water used to prepare the mobile phase was sparged with high purity N₂ for 30 min prior to
199 use. The mobile phase consisted of ultrapure water (eluent A) and NaOH at 1 M (eluent B)
200 prepared by dilution in degassed ultrapure water. The eluents were kept continuously under
201 pure N₂ pressure to avoid exposure to atmospheric CO₂, which can cause a drastic decrease in
202 column selectivity and loss of sugar resolution. In addition, an on-line degasser (RFIC eluent
203 degasser, Thermo Fisher) was added before the analytical column to decrease interference
204 from dissolved oxygen (Cheng and Kaplan, 2001).

205 In this study, two anion-exchange analytical columns were tested for sugar separation: a
206 CarboPac MA1 (7.5 μm, 4 × 250 mm; Thermo Fisher) and a CarboPac PA1 (10 μm, 4 × 250
207 mm; Thermo Fisher), both fitted with their corresponding guard column (4 × 50 mm; Thermo
208 Fisher). The Chromeleon software (Thermo Fisher) was used for processing and data
209 acquisition. The performance of the chromatographic separation (resolution factor, R_s)
210 between the two-carbohydrate species A and B is calculated according to the following
211 formula:

$$212 \quad R_s = 2 [(t_R)_B - (t_R)_A] \times [W_A + W_B]^{-1}$$

213 Where (t_R)_A and (t_R)_B are the retention times of the two species A and B, respectively, and
214 W_A and W_B are their corresponding peak widths.

215 **3. Results and discussion**

216 *3.1. Method optimization*

217 Although previous investigations described the simultaneous discrimination of some
218 neutral sugars, alditols and anhydrosugars using CarboPac PA1 or CarboPac MA1 columns
219 (Caseiro et al., 2007; Iinuma et al., 2009), the analytical HPAEC-PAD conditions used in
220 previous studies were unable to fully resolve the 17-carbohydrate mixture assessed in the

221 present study. Application of the analytical protocol using the CarboPac MA1 column in
222 Iinuma et al. (2009) revealed that the critical monosaccharide groups that required a further
223 efficient separation were, by elution order: (a) levoglucosan, arabitol, fucose and rhamnose,
224 (b) mannosan and sorbitol, (c) mannose, arabinose and galactosan, and (d) glucose and
225 xylose. The optimization of the method therefore involved evaluating the effects of basic
226 chromatographic parameters (flow rate, mobile phase composition and column temperature)
227 to achieve optimal separation of the 17-carbohydrate mixture. Charged carbohydrates (e.g.,
228 uronic acids and phosphorylated sugars) were not considered in this study because they
229 require different mobile phase conditions compared to neutral carbohydrates using
230 CH₃COONa as the eluent (Engel and Händel, 2011; Hu et al., 2012).

231

232 *3.1.1. Column selection*

233 Although several types of anion-exchange columns are available for carbohydrate analysis
234 (CarboPac PA10, 20, 100 and 200; Ronkart et al. 2007; Meyer et al. 2008; Raessler et al.
235 2010; Raessler 2011), the most recommended are: (1) the CarboPac PA1 column for analysis
236 of neutral sugars (Cheng and Kaplan, 2001; Kerhervé et al., 2002; Panagiotopoulos et al.,
237 2012; Panagiotopoulos and Sempéré, 2005a; Theodosi et al., 2018), and (2) the CarboPac
238 MA1 column for analysis of alditols, anhydrosugars and neutral sugars (Iinuma et al., 2009;
239 Jung et al., 2014; Zhang et al., 2013).

240 The CarboPac PA1 column is well-suited for the analysis of neutral monosaccharides.
241 However, when alditols and anhydrosugars are included in the same run, a NaOH gradient is
242 required because these compounds are eluted at lower NaOH concentrations (<1 mM)
243 compared to neutral monosaccharides (>15 mM). However, the use of a NaOH gradient
244 causes disequilibrium of the baseline signal, which further results in a positive (Sullivan et al.,

245 2011) or a negative peak (Caseiro et al., 2007; Theodosi et al., 2018) that may potentially
246 mask sugar peaks that are eluted within this time window.

247 The CarboPac MA1 column was introduced much later than the CarboPac PA1 column
248 and, to date, has never been applied to marine samples. Nevertheless, its macroporous
249 stationary phase has a higher anion-exchange capacity over the CarboPac PA1 column (1450
250 $\mu\text{eq}/\text{column}$ vs 100 $\mu\text{eq}/\text{column}$), which further allows the use of gradient conditions without
251 affecting the baseline stability (Iinuma et al., 2009; Zhang et al., 2013). Moreover, because
252 the CarboPac MA1 column operates at high NaOH elution conditions (>200 mM), the pH
253 values of the mobile phase are >12.5 and as such all the hemiacetal groups of sugars are
254 deprotonated. This further increases the retention of sugars on the column and thus results in a
255 better resolution (Thermo Fisher, Technical Note 20). Finally, the higher baseline stability of
256 the CarboPac MA1 column over the CarboPac PA1 column offers an additional advantage in
257 terms of the retention time of chromatographic peaks, which does not significantly shift over
258 successive injections (Gremm and Kaplan, 1997). Due to these reasons, this study was
259 conducted using the CarboPac MA1 column and different chromatographic parameters (flow
260 rate, etc.) were appropriately adjusted for separation of different classes or sugars.

261

262 3.1.2. *Optimization of flow rate, mobile phase and column temperature*

263 According to the manufacturer, the CarboPac MA1 column operates at a recommended
264 range of flow rates of 0.2–0.5 mL min^{-1} , and the flow rate is generally fixed at 0.4 mL min^{-1}
265 for environmental applications (Iinuma et al., 2009; Jung et al., 2014; Li et al., 2016).
266 Nevertheless, the latter authors did not report co-elution problems because they did not
267 consider fucose, rhamnose, arabinose, sorbitol and xylose in their standards. To overcome the
268 co-elution problem, we first decreased the flow rate to 0.3 mL min^{-1} . This resulted in a slight

269 improvement in the separation of the above compounds, and therefore this flow rate was
270 maintained for all further improvements.

271 Previous investigations employing the CarboPac MA1 column used NaOH concentrations
272 ranging from 300 to 800 mM (Andersen and Sørensen, 2000) to elute alditols, anhydrosugars,
273 and some neutral sugars (Iinuma et al., 2009). Nonetheless, such conditions were not optimal
274 for the 17 carbohydrates considered in this study. Therefore, different NaOH gradient tests
275 were performed at a fixed column temperature of 25 °C and a flow rate of 0.3 mL min⁻¹. The
276 improved separation of the first seven eluted monosaccharides (xylitol, levoglucosan, arabitol,
277 fucose, rhamnose, sorbitol and mannosan) was achieved by applying an NaOH gradient
278 ranging from 250 mM to 350 mM in the first 30 min (Fig. 1 a, Table 1). Unfortunately, fucose
279 and rhamnose were not resolved ($R_s = 0$) under these conditions despite additional
280 adjustments of the NaOH gradient. The 250 to 350 mM gradient was then followed by a 15
281 min-NaOH gradient increasing from 350 to 450 mM that allowed an acceptable separation
282 (with resolution factors close to or higher than 1) of mannitol, galactosan, arabinose and
283 mannose.

284 Following the chromatographic separation after 45 min, the NaOH concentration was
285 increased from 450 to 700 mM and held for 55 min to elute glucose, xylose, galactose and
286 fructose. Under these elution conditions, the xylose/galactose and galactose/fructose pairs
287 were fully resolved ($R_s > 1.5$), while the glucose/xylose pair was partially resolved ($R_s = 0.3$).
288 Additional tests of decreasing or increasing the final NaOH concentration (700 mM) did not
289 improve the glucose/xylose resolution. Finally, by prolonging the 700 mM NaOH
290 concentration for an additional 20 min, the last two carbohydrates (ribose and sucrose) were
291 eluted with excellent resolution ($R_s = 8.1$). After 75 min, the column was equilibrated for 20
292 min with the initial NaOH concentration (250 mM). Regeneration and clean-up of the column

293 were not necessary, because the high NaOH concentration (700 mM) reached at the end of the
294 analysis was sufficient to restore column performance.

295 Because pairs of levoglucosan/arabitol, fucose/rhamnose, galactosan/arabinose,
296 arabinose/mannose and glucose/xylose were completely or partially overlapped under the
297 NaOH gradient conditions optimized above, we also explored the effect of temperature on the
298 resolution of these monosaccharide pairs. Earlier investigations have shown that temperature
299 plays an important role in monosaccharide separation using the CarboPac PA1 column and
300 that sub-ambient temperatures strongly affect the resolution of closely eluting
301 monosaccharide pairs, notably fucose/rhamnose and mannose/xylose (Panagiotopoulos et al.,
302 2001). The results of this study indicated that glucose co-eluted with xylose at temperatures
303 $<20\text{ }^{\circ}\text{C}$, while arabinose overlapped with mannose at $>30\text{ }^{\circ}\text{C}$. Moreover, fucose and rhamnose
304 always co-eluted under both these temperature conditions. Overall, these results suggest that
305 sub-ambient temperature conditions do not have a significant impact on the separation of the
306 abovementioned carbohydrate pairs on the CarboPac MA1 column.

307 By narrowing the temperature range between 25 and 30 $^{\circ}\text{C}$, we found that these
308 monosaccharide pairs were better resolved (except the fucose/rhamnose pair which again co-
309 eluted), as shown by their corresponding resolution factors (Fig. 2). The results showed that
310 except for the arabinose/mannose ($R_s = 0.4$) and fucose/rhamnose pairs ($R_s = 0$), the resolution
311 factors for the levoglucosan/arabitol ($R_s = 0.7$), galactosan/arabinose ($R_s = 1.3$), and
312 glucose/xylose ($R_s = 0.6$) pairs were significantly improved at a temperature of 28 $^{\circ}\text{C}$.
313 Therefore, a temperature gradient was applied between 25 and 28 $^{\circ}\text{C}$ for the first 30 min of
314 analysis. The latter temperature was maintained for the next 45 min to distinguish the
315 galactosan/arabinose, arabinose/mannose and glucose/xylose pairs (Table 1, Table A 1; Fig. 1
316 a). After 75 min, the temperature was reset to 25 $^{\circ}\text{C}$ during column re-equilibration (20 min)
317 until the next injection.

318 *3.2. Linearity, detection limit, precision and blanks*

319 The optimized method showed a linear response for all monosaccharides ($R^2 > 0.999$) over
320 a concentration range of 50 nM to 10 μ M, which is typical for environmental monosaccharide
321 concentrations. The detection limits calculated at a signal-to-noise ratio (S/N) of three were
322 4–51 nM for neutral sugars, 2–15 nM for alditols and 8–32 nM for anhydrosugars (Table A
323 1), in agreement with previous studies using different elution conditions and anion-exchange
324 columns for monosaccharide determination (Caseiro et al., 2007; Iinuma et al., 2009;
325 Panagiotopoulos et al., 2001; Theodosi et al., 2018). The high detection limit obtained for
326 fructose (51 nM), also observed in a previous study (Iinuma et al., 2009), is probably due to
327 the low detector response toward this compound. The precision of the method was calculated
328 by performing six identical runs with the standard mixture of the 17 carbohydrates (50 nM
329 each). The relative standard deviation (RSD%) ranged from 1.8 to 16.6% for the peak area
330 and from 0.2 to 1.7% for the retention time (Table A 1). The procedural blank for the marine
331 POM consisted of a 25-mm Whatman GF/F filter (precombusted at 450 °C for 6 h),
332 hydrolyzed in the same manner as the samples. Similarly, the procedural blank for the TSP
333 consisted of a Whatman quartz fiber filter (precombusted at 450 °C for 6 h) extracted with the
334 ultrapure water. The results showed no detectable sugars in blanks (Fig. S 1).

335

336 *3.3. Comparison with the CarboPac PA1 column*

337 The final optimized MA1-HPAEC-PAD method (Table 1, Fig. 1 a) was compared with a
338 different method employing the CarboPac PA1 column (Caseiro et al., 2007; Theodosi et al.,
339 2018). The chromatogram obtained with the latter column (Fig. 1 b) showed that, at low
340 NaOH concentrations (1 mM; 0–15 min), xylitol, sorbitol, mannitol and mannosan peaks
341 were partially resolved, while the arabitol/levoglucosan and galactosan/fucose pairs
342 completely overlapped. Similar observations were made at high NaOH concentrations (19

343 mM; 15–53 min) for rhamnose/arabinose, mannose/xylose and ribose/sucrose pairs. Based on
344 the above co-elutions, these results clearly suggest that loss of molecular level information
345 may occur when a sample containing all of the abovementioned monosaccharides is analyzed
346 with the CarboPac PA1 column. Moreover, these co-elutions may further induce erroneous
347 quantification of these monosaccharides. Overall, these results indicate that the CarboPac
348 MA1 column is more appropriate than the CarboPac PA1 column for environmental
349 applications in terms of monosaccharide resolution and baseline stability.

350

351 3.4. *Application to environmental samples*

352 The method established in this study was tested on three different environmental matrices
353 including TSP, marine POM and HMWDOM. The chromatograms of these matrices are
354 presented in Fig. 3 and indicated the presence of large quantities of monosaccharides (Table
355 2).

356 3.4.1. *TSP*

357 The major carbohydrate classes identified in the TSP sample included anhydrosugars and
358 disaccharides, followed by neutral sugars and alditols. The sum of their concentration was 563
359 ng m^{-3} , representing the total neutral carbohydrate (TCHO) content recorded in the TSP
360 sample. The results of this study showed that levoglucosan (261 ng m^{-3}) and sucrose (178 ng
361 m^{-3}) were among the major neutral carbohydrates (Table 2; Fig. 3 a), which is in agreement
362 with other studies of European urban atmospheres (Yttri et al., 2007). Anhydrosugars
363 represented 53% of TCHO in the TSP sample and were dominated by levoglucosan and
364 mannosan, which accounted for 87% and 10% of the anhydrosugar pool, respectively (Fig. 4
365 a). Galactosan was also detected in the TSP sample but in lower concentrations (Table 2).

366 Neutral sugars accounted for 11% of the TCHO pool (Fig. 4 a) with fructose and glucose
367 exhibiting the highest concentrations (Fig. 4 a, Table 2). The presence of these neutral sugars
368 in the TSP sample strongly supports the influence of biogenic emission, which is typical of
369 plants growth during spring (Fu et al., 2012; Medeiros et al., 2006). Other minor neutral
370 sugars found in the TSP sample included the arabinose, mannose and xylose (Table 2).
371 Finally, alditols represented the least abundant carbohydrate class, accounting for 4% of
372 TCHO in the TSP sample (Fig. 4 a). The alditols decreased in abundance in the order of
373 xylitol, mannitol, arabitol and sorbitol (Table 2), and their presence in the sample indicates
374 inputs from fungal spores and/or microbial activities (Bauer et al., 2008; Dahlman et al.,
375 2003; Loos et al., 1994).

376 Although fucose and rhamnose co-eluted under the current HPAEC-PAD conditions, they
377 were also identified and quantified in the aerosol sample for the first time and accounted for
378 1% of TCHO and 8% of neutral sugars. Fucose and rhamnose may have multiple sources in
379 atmospheric particles including soil organic matter (Gunina and Kuzyakov, 2015; Ogner,
380 1980; Simoneit et al., 2004), vascular plants (Bianchi, 2007; Popper et al., 2004; Schädel et
381 al., 2010), microorganisms and bacteria (Cowie and Hedges, 1984; Fox et al., 1993; Petit et
382 al., 2013), while recent studies indicated that marine sources via water/air exchange may also
383 be an important contributor (Rastelli et al., 2017).

384 Nevertheless, field measurements using gas chromatography (Alves et al., 2011) or nuclear
385 magnetic resonance spectroscopy (Chalbot et al., 2013) identified only fucose or rhamnose
386 but never simultaneously reported both sugars in atmospheric samples. On the other hand,
387 studies using HPAEC-PAD have never reported fucose and rhamnose in atmospheric
388 particles, probably because of the co-elution of these deoxysugars with other
389 monosaccharides (fucose with galactosan and rhamnose with arabinose after analysis with the

390 CarboPac PA1 column; Fig. 1 b). Therefore, the presence of fucose and rhamnose in the
391 sample has never been thoroughly assessed to date.

392 The lack of data on deoxysugars in atmospheric samples may also be related to their
393 original structure. As deoxysugars are mostly found in bacterial lipopolysaccharides (Perry et
394 al., 1996; Weckesser et al., 1970; Zdrovenko et al., 2007) and/or plant structural
395 polysaccharides (Colombini et al., 2002; Grössl et al., 2005; Popper et al., 2004; Schädel et
396 al., 2010), they are either not efficiently recovered after water extraction (case of
397 lipopolysaccharides) or are recovered in the water phase but cannot be released from
398 carbohydrate polymer (case of structural polysaccharides) as no hydrolysis is applied.

399 3.4.2. *Marine POM*

400 The major carbohydrate classes identified in the marine POM sample included the neutral
401 sugars followed by anhydrosugars and alditols. The sum of their concentration was 16.2 mg
402 g⁻¹. Neutral sugars accounted for 98% of the TCHO pool (Fig. 4 b) and were dominated by
403 galactose (5.7 mg g⁻¹) and glucose (3.8 mg g⁻¹) (Table 2; Fig. 3 b). These monosaccharides
404 were followed by fructose (2.0 mg g⁻¹) and fucose/rhamnose (co-eluted; 1.8 mg g⁻¹) while
405 xylose, mannose and arabinose exhibited concentrations <1 mg g⁻¹. These results are in good
406 agreement with previously reported monosaccharide patterns for marine POM in several
407 oceanic regimes (Hernes et al., 1996; Panagiotopoulos and Sempéré, 2005b; Skoog and
408 Benner, 1997). Fructose and ribose, occasionally reported in marine POM samples due to
409 their low recovery after acid hydrolysis (Borch and Kirchman, 1997; Panagiotopoulos and
410 Sempéré, 2005a), were also identified in this sample (Table 2). This finding most likely
411 indicates the presence of storage polysaccharides (fructans) and nonstructural labile
412 compounds such as RNA or nucleotides originating from marine organisms (Cowie and
413 Hedges, 1984; Haug and Mykelstad, 1976; Hicks et al., 1994; Panagiotopoulos and Sempéré,
414 2005b). Alternatively, fructose may originate from sucrose. As sucrose was not detected in

415 the marine POM sample (the same holds for the HMWDOM sample), this most likely implies
416 its complete hydrolysis under the acid conditions used for its extraction, which results in two
417 monosaccharides (fructose and glucose).

418 The proposed analytical method allowed the identification and quantification of alditols
419 (0.4% of TCHO) and anhydrosugars (2% of TCHO), the two monosaccharide families that
420 have been largely overlooked to date in marine environmental studies (Fig. 4 b). The results
421 showed that xylitol was the major alditol type detected in the POM sample, followed by
422 sorbitol, while arabitol and mannitol were not detected in the sample (Table 2). Alditols are
423 generally reported as terrestrial tracers derived from fungal spores and/or microbial activities
424 (Dahlman et al., 2003; Vandeska et al., 1995), therefore, their presence in the marine POM
425 sample may suggest possible terrestrial input (van Pinxteren et al., 2012). Alternatively,
426 alditols may have marine origin. As carbohydrates are an important fraction of the organic
427 matter produced by marine primary production or released into the environment via
428 degradation processes, it is highly possible that alditols are also part of the carbohydrate pool.
429 Nevertheless, additional measurements using compound-specific ^{13}C analysis are warranted to
430 test this hypothesis and constrain the sources of alditols in the marine waters.

431 Compositionally, anhydrosugars were dominated by galactosan followed by levoglucosan
432 and mannosan, which represented for 63%, 25% and 13% of total anhydrosugars, respectively
433 (Fig. 4 b). The dominance of galactosan over levoglucosan has also been reported for sea
434 surface microlayer samples in Baltic sea (van Pinxteren et al., 2012) during different seasons
435 of the year however, more data are warranted in the same time from both atmospheric and
436 marine compartments to validate/confirm these findings.

437 However, the anhydrosugar concentrations found in this sample (and in the HMWDOM
438 sample) are most likely underestimated because these compounds are converted to their
439 respective hexoses after acid hydrolysis (Blanco et al., 2018). Indeed, in an additional set of

440 experiments to investigate whether the hydrolysis conditions affect the stability of
441 monosaccharides, we performed acid hydrolysis (1 M HCl, 100 °C, 20 h) on a standard
442 monosaccharide mixture containing neutral sugars, alditols and anhydrosugars. The results
443 showed that levoglucosan, mannosan and galactosan were completely converted to glucose,
444 mannose and galactose, respectively, while minimal loss was observed for neutral sugars and
445 alditols (Fig. S 2), which is consistent with previous studies (Skoog and Benner, 1997; Wang
446 et al., 2016).

447 Overall, these results revealed that the concentration of anhydrosugars is biased in
448 environmental samples if these sugars are acid-extracted (POM and HMWDOM sample).
449 Therefore, an alternative extraction method should be considered (e.g., extraction with water
450 or MeOH). Despite the possible underestimation of the amount of the anhydrosugars found in
451 the marine POM, their presence in this sample clearly indicates an external input, most likely
452 via atmospheric deposition, as shown by the isotopic signature ($\delta^{13}\text{C}$) of levoglucosan and
453 mannosan pointing to a terrestrial origin (Nouara et al., 2019).

454

455 3.4.3. *Marine HMWDOM*

456 The major carbohydrate classes identified in the HMWDOM sample included the neutral
457 sugars followed by anhydrosugars and alditols. The sum of their concentration was 766 μg
458 g^{-1} . Neutral sugars accounted for 97% of the TCHO pool in the HMWDOM (Fig. 4 c). The
459 results of this study revealed that fucose & rhamnose (300 $\mu\text{g g}^{-1}$), glucose (122 $\mu\text{g g}^{-1}$),
460 mannose (103 $\mu\text{g g}^{-1}$), galactose (83 $\mu\text{g g}^{-1}$), xylose (74 $\mu\text{g g}^{-1}$), and arabinose (34 $\mu\text{g g}^{-1}$)
461 were among the major monosaccharides, which agrees with the neutral sugar pattern reported
462 in previous studies (Aluwihare et al., 1997; Repeta et al., 2002) (Table 2; Fig. 3c). Fructose
463 and ribose were also quantified in the HMWDOM sample, and similar to POM, they most
464 likely originate from storage polysaccharide or RNA.

465 Anhydrosugars accounted for 3% of TCHO and were dominated by levoglucosan,
466 galactosan and mannosan, which represented 61%, 22% and 17% of total anhydrosugars,
467 respectively (Fig. 4 c). The lower concentration of anhydrosugars in the HMWDOM sample
468 (the same holds for the POM sample) compared to the TSP sample maybe due to the high
469 dilution occurring at sea and also to photodegradation processes by OH* radicals which take
470 place in the atmosphere for these compounds (Hoffmann et al., 2010; Lai et al., 2014). In
471 addition, as suggested for marine POM, the concentration of anhydrosugars in HMWDOM is
472 most likely biased due to the acid extraction procedure. The presence of the anhydrosugars in
473 the sample may indicate possible input of terrestrial organic matter (burning biomass tracers),
474 although additional studies using $\delta^{13}\text{C}$ are warranted to confirm this hypothesis. Finally, the
475 least abundant carbohydrate class was the alditols, which accounted 0.3% of TCHO and
476 represented by mannitol and xylitol (Fig. 4 c).

477

478 **4. Conclusions**

479 The results of this study indicated that under the optimized analytical conditions, neutral
480 sugars, alditols, and anhydrosugars commonly found in most environmental samples can be
481 analyzed with a single run in 75 min with satisfactory resolution except for the
482 fucose/rhamnose pair that completely co-eluted. These deoxysugars were identified for the
483 first time in the TSP sample by this improved HPAEC-PAD technique, which will facilitate
484 future investigations by shedding light on bacterial and/or microorganism activities on
485 atmospheric samples. However, additional deconvolution of these deoxysugars signals using
486 the traditional HPAEC-PAD method for neutral sugar determination (Panagiotopoulos et al.,
487 2001) might also be useful.

488 The proposed improved HPAEC-PAD method allowed, for the first time, the identification
489 and quantification of anhydrosugars and alditols in marine samples (marine POM and

490 HMWDOM). These two carbohydrate families have largely been overlooked in the dissolved
491 free monosaccharide pool (DFMS), and, to the best of our knowledge, they have never been
492 reported in marine DOM. Indeed, anhydrosugars and alditols can easily be extracted from
493 seawater using cation and anion-exchange resins (Kirchman et al., 2001; Mopper et al., 1992;
494 Sempéré et al., 2008; Skoog and Benner, 1997; van Pinxteren et al., 2012) in the same manner
495 as neutral sugars. However, because the subsequent PA1-HPAEC-PAD analysis has always
496 been performed at >15 mM NaOH to elute the neutral sugars; anhydrosugars and alditols
497 always co-elute at time window of 2–5 min, therefore, they have never been identified in the
498 DFMS pool.

499 On the other hand, dissolved combined monosaccharides (DCHO) or polysaccharides in
500 marine samples are generally measured as their monomeric constituents that are released after
501 acid hydrolysis. Thus, if anhydrosugars are present in the sample, they will be counted in the
502 hexose pool as the sample is hydrolyzed. Therefore, we recommend that DCHO analysis
503 always be preceded or accompanied by the analysis of DFMS to evaluate the contribution of
504 anhydrosugars – if present – in the sample. This strategy is particularly critical for samples
505 from coastal areas that receive significant terrestrial inputs, particularly in wintertime, and are
506 characterized by strong influence from wood burning.

507 The proposed HPAEC method provides an acceptable resolution for most of the
508 carbohydrates examined, and therefore its coupling with mass spectrometry may further help
509 with the structural elucidation and confirmation of the analyzed carbohydrates. Moreover, its
510 high potential may further be explored in other environmental matrices, including sediments,
511 seawater, and marine biota, which will provide a more complete profile of carbohydrates in
512 the environment. This complete profile, in turn, may be helpful to improve our understanding
513 on carbohydrate dynamics in the marine and terrestrial ecosystems and potentially better
514 evaluate the impact of biomass burning processes on marine waters.

515 **Acknowledgments**

516 This research was supported by the funding from the projects AIOLOS (Labex OT-Med;
517 ANR-11-LABX-0.061) and MANDARINE (grant No 2008-10372; Région Provence Alpes
518 Côte d'Azur). The authors thank D. Navarro for analytical assistance to conduct a part of this
519 research at the INRA laboratory of Fungal Biodiversity and Biotechnology UMR 1163
520 (13009 Marseille; France). The authors also acknowledge A. Paulmier and D.J. Repeta for
521 providing the POM sample and HMWDOM powder, respectively as well as the two
522 anonymous reviewers for valuable comments and fruitful discussions.

523 **References**

- 524 Aluwihare, L.I., Repeta, D.J., Chen, R.F., 1997. A major biopolymeric component to
525 dissolved organic carbon in surface sea water. *Nature*. <https://doi.org/10.1038/387166a0>
- 526 Alves, C.A., Vicente, A., Monteiro, C., Gonçalves, C., Evtugina, M., Pio, C., 2011.
527 Emission of trace gases and organic components in smoke particles from a wildfire in a
528 mixed-evergreen forest in Portugal. *Sci. Total Environ.* 409, 1466–75.
529 <https://doi.org/10.1016/j.scitotenv.2010.12.025>
- 530 Amon, R.M.W., Fritznar, H.P., Benner, R., 2001. Linkages among the bioreactivity, chemical
531 composition, and diagenetic state of marine dissolved organic matter. *Limnol. Ocean.* 46,
532 287–297. <https://doi.org/10.4319/lo.2001.46.2.0287>
- 533 Andersen, R., Sørensen, A., 2000. Separation and determination of alditols and sugars by
534 high-pH anion-exchange chromatography with pulsed amperometric detection. *J.*
535 *Chromatogr. A* 897, 195–204. [https://doi.org/10.1016/S0021-9673\(00\)00783-4](https://doi.org/10.1016/S0021-9673(00)00783-4)
- 536 Barbaro, E., Kirchgeorg, T., Zangrando, R., Vecchiato, M., Piazza, R., Barbante, C.,
537 Gambaro, A., 2015. Sugars in Antarctic aerosol. *Atmos. Environ.* 118, 135–144.
538 <https://doi.org/10.1016/j.atmosenv.2015.07.047>

539 Bauer, H., Claeys, M., Vermeylen, R., Schueller, E., Weinke, G., Berger, A., Puxbaum, H.,
540 2008. Arabitol and mannitol as tracers for the quantification of airborne fungal spores.
541 *Atmos. Environ.* 42, 588–593. <https://doi.org/10.1016/j.atmosenv.2007.10.013>

542 Benner, R., Kaiser, K., 2003. Abundance of amino sugars and peptidoglycan in marine
543 particulate and dissolved organic matter. *Limnol. Oceanogr.* 48, 118–128.
544 <https://doi.org/10.4319/lo.2003.48.1.0118>

545 Bergamaschi, B.A., Walters, J.S., Hedges, J.I., 1999. Distributions of uronic acids and O-
546 methyl sugars in sinking and sedimentary particles in two coastal marine environments.
547 *Geochim. Cosmochim. Acta* 63, 413–425. [https://doi.org/10.1016/S0016-](https://doi.org/10.1016/S0016-7037(99)00075-7)
548 [7037\(99\)00075-7](https://doi.org/10.1016/S0016-7037(99)00075-7)

549 Bhattarai, H., Saikawa, E., Wan, X., Zhu, H., Ram, K., Gao, S., Kang, S., Zhang, Q., Zhang,
550 Y., Wu, G., Wang, X., Kawamura, K., Fu, P., Cong, Z., 2019. Levoglucosan as a tracer
551 of biomass burning: Recent progress and perspectives. *Atmos. Res.* 220, 20–33.
552 <https://doi.org/10.1016/J.ATMOSRES.2019.01.004>

553 Bianchi, T.S., 2007. *Biogeochemistry of estuaries*. Oxford University Press, Oxford.

554 Blanco, P.H., Lad, J.B., Bridgwater, A. V., Holm, M.S., 2018. Production of Glucose from
555 the Acid Hydrolysis of Anhydrosugars. *ACS Sustain. Chem. Eng.* 6, 12872–12883.
556 <https://doi.org/10.1021/acssuschemeng.8b02202>

557 Bock, M., Glaser, B., Millar, N., 2007. Sequestration and turnover of plant and microbially
558 derived sugars in a temperate grassland soil during 7 years exposed to elevated
559 atmospheric pCO₂. *Glob. Chang. Biol.* 13, 478–490. [https://doi.org/10.1111/j.1365-](https://doi.org/10.1111/j.1365-2486.2006.01303.x)
560 [2486.2006.01303.x](https://doi.org/10.1111/j.1365-2486.2006.01303.x)

561 Borch, N.H., Kirchman, D.L., 1997. Concentration and composition of dissolved combined
562 neutral sugars (polysaccharides) in seawater determined by HPLC-PAD. *Mar. Chem.* 57,

563 85–95. [https://doi.org/10.1016/S0304-4203\(97\)00002-9](https://doi.org/10.1016/S0304-4203(97)00002-9)

564 Bretagnon, M., Paulmier, A., Garcon, V., Dewitte, B., Illig, S., Coppola, L., Campos, F.,
565 Velazco, F., Panagiotopoulos, C., Oshlies, A., Hernandez-ayon, J.M., Maske, H.,
566 Vergara, O., Montes, I., Martinez, P., Carrasco, E., Grelet, J., Desprez-De-Gesincourt,
567 Maes, C., Scouarnec, L., 2018. Modulation of the vertical particles transfer efficiency in
568 the Oxygen Minimum Zone off Peru. *Biogeosciences* 15, 1–19.
569 <https://doi.org/10.5194/bg-15-1-2018>

570 Caseiro, A., Marr, I.L., Claeys, M., Kasper-Giebl, A., Puxbaum, H., Pio, C. a., 2007.
571 Determination of saccharides in atmospheric aerosol using anion-exchange high-
572 performance liquid chromatography and pulsed-amperometric detection. *J. Chromatogr.*
573 *A* 1171, 37–45. <https://doi.org/10.1016/j.chroma.2007.09.038>

574 Chalbot, M.C.G., Gamboa da Costa, G., Kavouras, I.G., 2013. NMR Analysis of the Water-
575 Soluble Fraction of Airborne Pollen Particles. *Appl. Magn. Reson.* 44, 1347–1358.
576 <https://doi.org/10.1007/s00723-013-0492-4>

577 Cheng, X., Kaplan, L.A., 2003. Simultaneous analyses of neutral carbohydrates and amino
578 sugars in freshwaters with HPLC-PAD. *J. Chromatogr. Sci.* 41, 434–438.

579 Cheng, X., Kaplan, L.A., 2001. Improved Analysis of Dissolved Carbohydrates in Stream
580 Water with HPLC-PAD. *Anal. Chem.* 73, 458–461. <https://doi.org/10.1021/ac001059r>

581 Colombini, M.P., Ceccarini, A., Carmignani, A., 2002. Ion chromatography characterization
582 of polysaccharides in ancient wall paintings. *J. Chromatogr. A* 968, 79–88.
583 [https://doi.org/10.1016/S0021-9673\(02\)00950-0](https://doi.org/10.1016/S0021-9673(02)00950-0)

584 Cowie, G.L., Hedges, J.I., 1984. Carbohydrate sources in a coastal marine environment.
585 *Geochim. Cosmochim. Acta* 48, 2075–2087. [https://doi.org/10.1016/0016-](https://doi.org/10.1016/0016-7037(84)90388-0)
586 [7037\(84\)90388-0](https://doi.org/10.1016/0016-7037(84)90388-0)

587 da Cunha, L.C., Serve, L., Blazi, J.-L., 2002. Neutral sugars as biomarkers in the particulate
588 organic matter of a French Mediterranean river. *Org. Geochem.* 33, 953–964.
589 [https://doi.org/https://doi.org/10.1016/S0146-6380\(02\)00058-X](https://doi.org/https://doi.org/10.1016/S0146-6380(02)00058-X)

590 Dahlman, L., Persson, J., Näsholm, T., Palmqvist, K., 2003. Carbon and nitrogen distribution
591 in the green algal lichens *Hypogymnia physodes* and *Platismatia glauca* in relation to
592 nutrient supply. *Planta* 217, 41–48. <https://doi.org/10.1007/s00425-003-0977-8>

593 Elias, V.O., Simoneit, B.R.T., Cordeiro, R.C., Turcq, B., 2001. Evaluating levoglucosan as an
594 indicator of biomass burning in Carajas, Amazonia: A comparison to the charcoal
595 record. *Geochim. Cosmochim. Acta* 65, 267–272.

596 Engel, A., Händel, N., 2011. A novel protocol for determining the concentration and
597 composition of sugars in particulate and in high molecular weight dissolved organic
598 matter (HMW-DOM) in seawater. *Mar. Chem.* 127, 180–191.
599 <https://doi.org/10.1016/j.marchem.2011.09.004>

600 Fox, A., Black, G., Fox, K., Rostovtseva, S., 1993. Determination of carbohydrate profiles of
601 *Bacillus anthracis* and *Bacillus cereus* including identification of O-methyl
602 methylpentose by using gas chromatography-mass spectrometry. *J. Clin. Microbiol.* 31,
603 887–894.

604 Fu, P., Kawamura, K., Kobayashi, M., Simoneit, B.R.T., 2012. Seasonal variations of sugars
605 in atmospheric particulate matter from Gosan, Jeju Island: Significant contributions of
606 airborne pollen and Asian dust in spring. *Atmos. Environ.* 55, 234–239.
607 <https://doi.org/10.1016/j.atmosenv.2012.02.061>

608 Gambaro, A., Zangrando, R., Gabrielli, P., Barbante, C., Cescon, P., 2008. Direct
609 determination of levoglucosan at the picogram per milliliter level in antarctic ice by
610 high-performance liquid chromatography/electrospray ionization triple quadrupole mass

611 spectrometry. *Anal. Chem.* 80, 1649–1655. <https://doi.org/10.1021/ac701655x>

612 Giorio, C., Kehrwald, N., Barbante, C., Kalberer, M., King, A.C.F., Thomas, E.R., Wolff,
613 E.W., Zennaro, P., 2018. Prospects for reconstructing paleoenvironmental conditions
614 from organic compounds in polar snow and ice. *Quat. Sci. Rev.* 183, 1–22.
615 <https://doi.org/10.1016/j.quascirev.2018.01.007>

616 Giroldo, D., Henriques Vieira, A.A., Paulsen, B.S., 2003. relative increase of deoxy sugars
617 during microbial degradation of an extracellular polysaccharide released by a tropical
618 freshwater *Thalassiosira* sp. (Bacillariophyceae). *J. Phycol.* 39, 1109–1115.
619 <https://doi.org/10.1111/j.0022-3646.2003.03-006.x>

620 Gremm, T.J., Kaplan, L.A., 1997. Dissolved carbohydrates in streamwater determined by
621 HPLC and pulsed amperometric detection. *Limnol. Oceanogr.* 42, 385–393.
622 <https://doi.org/10.4319/lo.1997.42.2.0385>

623 Grössl, M., Harrison, S., Kaml, I., Kenndler, E., 2005. Characterisation of natural
624 polysaccharides (plant gums) used as binding media for artistic and historic works by
625 capillary zone electrophoresis. *J. Chromatogr. A* 1077, 80–89.
626 <https://doi.org/10.1016/j.chroma.2005.04.075>

627 Gunina, A., Kuzyakov, Y., 2015. Sugars in soil and sweets for microorganisms: Review of
628 origin, content, composition and fate. *Soil Biol. Biochem.* 90, 87–100.
629 <https://doi.org/10.1016/j.soilbio.2015.07.021>

630 Haug, A., Mykelstad, S., 1976. Polysaccharides of marine diatoms with special reference to
631 *Chaetoceros* species. *Mar. Biol.* 34, 217–222.
632 <https://doi.org/https://doi.org/10.1007/BF00388798>

633 He, B., Dai, M., Huang, W., Liu, Q., Chen, H., Xu, L., 2010. Sources and accumulation of
634 organic carbon in the Pearl River Estuary surface sediment as indicated by elemental,

635 stable carbon isotopic, and carbohydrate compositions. *Biogeosciences* 7, 3343–3362.
636 <https://doi.org/10.5194/bg-7-3343-2010>

637 Hedges, J.I., Cowie, G.L., Richey, J.E., Quay, P.D., Benner, R., Strom, M., Forsberg, B.R.,
638 1994. Origins and processing of organic matter in the Amazon River as indicated by
639 carbohydrates and amino acids. *Limnol. Oceanogr.* 39, 743–761.
640 <https://doi.org/10.4319/lo.1994.39.4.0743>

641 Hernes, P., Hedges, J., Peterson, M., Wakeham, S., Lee, C., 1996. Neutral carbohydrate
642 geochemistry of particulate material in the central equatorial Pacific. *Deep. Res. PART*
643 *II-TOPICAL Stud. Oceanogr.* 43, 1181–1204. [https://doi.org/10.1016/0967-](https://doi.org/10.1016/0967-0645(96)00012-4)
644 [0645\(96\)00012-4](https://doi.org/10.1016/0967-0645(96)00012-4)

645 Hicks, R.E., Owen, C.J., Aas, P., 1994. Deposition, resuspension, and decomposition of
646 particulate organic matter in the sediments of Lake Itasca, Minnesota, USA.
647 *Hydrobiologia* 284, 79–91. <https://doi.org/10.1007/BF00005733>

648 Hoffmann, D., Tilgner, a, Iinuma, Y., Herrmann, H., 2010. Atmospheric stability of
649 levoglucosan: a detailed laboratory and modeling study. *Environ. Sci. Technol.* 44, 694–
650 9. <https://doi.org/10.1021/es902476f>

651 Hu, Q., Tan, L., Heng, Z., Su, X., Zhang, T., Jiang, Z., Xiong, X., 2012. Quantification of
652 sugar compounds and uronic acids in enzymatic hydrolysates of lignocellulose using
653 high-performance anion exchange chromatography with pulsed amperometric detection.
654 *Energy and Fuels* 26, 2942–2947. <https://doi.org/10.1021/ef300055k>

655 Hung, C.C., Tang, D., Warnken, K.W., Santschi, P.H., 2001. Distributions of carbohydrates,
656 including uronic acids, in estuarine waters of Galveston Bay. *Mar. Chem.* 73, 305–318.
657 [https://doi.org/10.1016/S0304-4203\(00\)00114-6](https://doi.org/10.1016/S0304-4203(00)00114-6)

658 Iinuma, Y., Engling, G., Puxbaum, H., Herrmann, H., 2009. A highly resolved anion-

659 exchange chromatographic method for determination of saccharidic tracers for biomass
660 combustion and primary bio-particles in atmospheric aerosol. *Atmos. Environ.* 43, 1367–
661 1371. <https://doi.org/10.1016/j.atmosenv.2008.11.020>

662 Jung, J., Lee, S., Kim, H., Kim, D., Lee, H., Oh, S., 2014. Quantitative determination of the
663 biomass-burning contribution to atmospheric carbonaceous aerosols in Daejeon, Korea,
664 during the rice-harvest period. *Atmos. Environ.* 89, 642–650.
665 <https://doi.org/10.1016/j.atmosenv.2014.03.010>

666 Kaiser, K., Benner, R., 2009. Biochemical composition and size distribution of organic matter
667 at the Pacific and Atlantic time series stations. *Mar. Chem.* 113, 63–77.
668 <https://doi.org/10.1016/j.marchem.2008.12.004>

669 Kerhervé, P., Buscail, R., Gadel, F., Serve, L., 2002. Neutral monosaccharides in surface
670 sediments of the northwestern Mediterranean Sea. *Org. Geochem.* 33, 421–435.
671 [https://doi.org/10.1016/S0146-6380\(02\)00003-7](https://doi.org/10.1016/S0146-6380(02)00003-7)

672 Kirchman, D.L., Meon, B., Ducklow, H.W., Carlson, C.A., Hansell, D.A., Steward, G.F.,
673 2001. Glucose fluxes and concentrations of dissolved combined neutral sugars
674 (polysaccharides) in the Ross Sea and Polar Front Zone, Antarctica. *Deep Sea Res. Part*
675 *II Top. Stud. Oceanogr.* 4179–4197. [https://doi.org/10.1016/S0967-0645\(01\)00085-6](https://doi.org/10.1016/S0967-0645(01)00085-6),
676 2001

677 Lai, C., Liu, Y., Ma, J., Ma, Q., He, H., 2014. Degradation kinetics of levoglucosan initiated
678 by hydroxyl radical under different environmental conditions. *Atmos. Environ.* 91, 32–
679 39. <https://doi.org/10.1016/j.atmosenv.2014.03.054>

680 Li, X., Chen, M., Le, H.P., Wang, F., Guo, Z., Iinuma, Y., Chen, J., Herrmann, H., 2016.
681 Atmospheric outflow of PM_{2.5} saccharides from megacity Shanghai to East China Sea:
682 Impact of biological and biomass burning sources. *Atmos. Environ.* 143, 1–14.

683 <https://doi.org/10.1016/j.atmosenv.2016.08.039>

684 Loos, H., Kramer, R., Sahn, H., Sprenger, G.A., 1994. Sorbitol promotes growth of
685 *Zygomonas mobilis* in environments with high-concentrations of sugar - evidence for a
686 physiological-function of glucose-fructose oxidoreductase in osmoprotection. *J.*
687 *Bacteriol.* 176, 7688–7693.

688 Medeiros, P.M., Conte, M.H., Weber, J.C., Simoneit, B.R.T., 2006. Sugars as source
689 indicators of biogenic organic carbon in aerosols collected above the Howland
690 Experimental Forest, Maine. *Atmos. Environ.* 40, 1694–1705.
691 <https://doi.org/10.1016/j.atmosenv.2005.11.001>

692 Meyer, A., Fischer, H., Kuzyakov, Y., Fischer, K., 2008. Improved RP-HPLC and anion-
693 exchange chromatography methods for the determination of amino acids and
694 carbohydrates in soil solutions. *J. Plant Nutr. Soil Sci.* 171, 917–926.
695 <https://doi.org/10.1002/jpln.200700235>

696 Mopper, K., 1977. Sugars and uronic acids in sediment and water from the black sea and
697 north sea with emphasis on analytical techniques. *Mar. Chem.* 5, 585–603.
698 [https://doi.org/10.1016/0304-4203\(77\)90044-5](https://doi.org/10.1016/0304-4203(77)90044-5)

699 Mopper, K., Dawson, R., Liebezeit, G., Ittekkot, V., 1980. The monosaccharide spectra of
700 natural waters. *Mar. Chem.* 10, 55–66. [https://doi.org/https://doi.org/10.1016/0304-](https://doi.org/https://doi.org/10.1016/0304-4203(80)90058-4)
701 [4203\(80\)90058-4](https://doi.org/https://doi.org/10.1016/0304-4203(80)90058-4)

702 Mopper, K., Schultz, C., Chevolut, L., Germain, C., Revuelta, R., Dawson, R., 1992.
703 Determination of sugars in unconcentrated seawater and other natural waters by liquid
704 chromatography and pulsed amperometric detection. *Environ. Sci. Technol.* 26, 133–
705 138. <https://doi.org/10.1021/es00025a014>

706 Nouara, A., Panagiotopoulos, C., Balesdent, J., Violaki, K., Bard, E., Fagault, Y., Repeta,

707 D.J., Sempéré, R., 2019. Liquid chromatographic isolation of individual carbohydrates
708 from environmental matrices for stable carbon analysis and radiocarbon dating. *Anal.*
709 *Chim. Acta.* 1067, 137–146. <https://doi.org/10.1016/J.ACA.2019.03.028>

710 Ogner, G., 1980. Analysis of the carbohydrates of fulvic and humic acids as their partially
711 methylated alditol acetates. *Geoderma* 23, 1–10. [https://doi.org/10.1016/0016-](https://doi.org/10.1016/0016-7061(80)90045-2)
712 [7061\(80\)90045-2](https://doi.org/10.1016/0016-7061(80)90045-2)

713 Opsahl, S., Benner, R., 1999. Characterization of carbohydrates during early diagenesis of
714 five vascular plant tissues. *Org. Geochem.* 30, 83–94. [https://doi.org/10.1016/S0146-](https://doi.org/10.1016/S0146-6380(98)00195-8)
715 [6380\(98\)00195-8](https://doi.org/10.1016/S0146-6380(98)00195-8)

716 Panagiotopoulos, C., Repeta, D.J., Johnson, C.G., 2007. Characterization of methyl sugars, 3-
717 deoxysugars and methyl deoxysugars in marine high molecular weight dissolved organic
718 matter. *Org. Geochem.* 38, 884–896. <https://doi.org/10.1016/j.orggeochem.2007.02.005>

719 Panagiotopoulos, C., Repeta, D.J., Mathieu, L., Rontani, J.F., Sempéré, R., 2013. Molecular
720 level characterization of methyl sugars in marine high molecular weight dissolved
721 organic matter. *Mar. Chem.* 154, 34–45. <https://doi.org/10.1016/j.marchem.2013.04.003>

722 Panagiotopoulos, C., Sempéré, R., 2005a. Analytical methods for the determination of sugars
723 in marine samples : A historical perspective and future directions. *Limnol. Oceanogr.*
724 *Methods* 3, 419–454. <https://doi.org/10.4319/lom.2005.3.419>

725 Panagiotopoulos, C., Sempéré, R., 2005b. The molecular distribution of combined aldoses in
726 sinking particles in various oceanic conditions. *Mar. Chem.* 95, 31–49.
727 <https://doi.org/10.1016/j.marchem.2004.07.005>

728 Panagiotopoulos, C., Sempéré, R., Jacq, V., Charrière, B., 2014. Composition and distribution
729 of dissolved carbohydrates in the Beaufort Sea Mackenzie margin (Arctic Ocean). *Mar.*
730 *Chem.* 166, 92–102. <https://doi.org/10.1016/j.marchem.2014.09.004>

731 Panagiotopoulos, C., Sempéré, R., Lafont, R., Kerhervé, P., 2001. Sub-ambient temperature
732 effects on the separation of monosaccharides by high-performance anion-exchange
733 chromatography with pulse amperometric detection - Application to marine chemistry. *J.*
734 *Chromatogr. A* 920, 13–22. [https://doi.org/10.1016/S0021-9673\(01\)00697-5](https://doi.org/10.1016/S0021-9673(01)00697-5)

735 Panagiotopoulos, C., Sempéré, R., Para, J., Raimbault, P., Rabouille, C., Charrière, B., 2012.
736 The composition and flux of particulate and dissolved carbohydrates from the Rhone
737 River into the Mediterranean Sea. *Biogeosciences* 9, 1827–1844.
738 <https://doi.org/10.5194/bg-9-1827-2012>

739 Pantelaki, I., Papatzelou, A., Balla, D., Papageorgiou, A., Voutsas, D., 2018. Characterization
740 of dissolved organic carbon in rainwater of an urban/coastal site in Mediterranean area.
741 *Sci. Total Environ.* 627, 1433–1441.
742 <https://doi.org/https://doi.org/10.1016/j.scitotenv.2018.01.339>

743 Perry, M.B., Maclean, L.L., Gmür, R., Wilson, M.E., 1996. Characterization of the O-
744 polysaccharide structure of lipopolysaccharide from *Actinobacillus*
745 *actinomycetemcomitans* serotype b. *Infect. Immun.* 64, 1215–1219.

746 Petit, E., LaTouf, W.G., Coppi, M. V., Warnick, T.A., Currie, D., Romashko, I., Deshpande,
747 S., Haas, K., Alvelo-Maurosa, J.G., Wardman, C., Schnell, D.J., Leschine, S.B.,
748 Blanchard, J.L., 2013. Involvement of a Bacterial Microcompartment in the Metabolism
749 of Fucose and Rhamnose by *Clostridium phytofermentans*. *PLoS One* 8, 1–12.
750 <https://doi.org/10.1371/journal.pone.0054337>

751 Popper, Z.A., Sadler, I.H., Fry, S.C., 2004. 3-O-methylrhamnose in lower land plant primary
752 cell walls. *Biochem. Syst. Ecol.* 32, 279–289. <https://doi.org/10.1016/j.bse.2003.07.004>

753 Raessler, M., 2011. Sample preparation and current applications of liquid chromatography for
754 the determination of non-structural carbohydrates in plants. *Trends Anal. Chem.* 30,

755 1833–1843. <https://doi.org/10.1016/j.trac.2011.06.013>

756 Raessler, M., Wissuwa, B., Breul, A., Unger, W., Grimm, T., 2010. Chromatographic analysis
757 of major non-structural carbohydrates in several wood species - An analytical approach
758 for higher accuracy of data. *Anal. Methods* 2, 532–538.
759 <https://doi.org/10.1039/b9ay00193j>

760 Rastelli, E., Corinaldesi, C., Dell’anno, A., Lo Martire, M., Greco, S., Cristina Facchini, M.,
761 Rinaldi, M., O’Dowd, C., Ceburnis, D., Danovaro, R., 2017. Transfer of labile organic
762 matter and microbes from the ocean surface to the marine aerosol: An experimental
763 approach. *Sci. Rep.* 7, 1–10. <https://doi.org/10.1038/s41598-017-10563-z>

764 Repeta, D.J., Aluwihare, L., Carlson, C., Liu, Z., Nelson, C., Stubbins, A., 2015. Introduction
765 to the special issue on the Biogeochemistry of Dissolved Organic Matter. *Mar. Chem.*
766 177, 203–204. <https://doi.org/10.1016/j.marchem.2015.10.002>

767 Repeta, D.J., Quan, T.M., Aluwihare, L.I., Accardi, A., 2002. Chemical characterization of
768 high molecular weight dissolved organic matter in fresh and marine waters. *Geochim.*
769 *Cosmochim. Acta* 66, 955–962. [https://doi.org/10.1016/S0016-7037\(01\)00830-4](https://doi.org/10.1016/S0016-7037(01)00830-4)

770 Ronkart, S.N., Blecker, C.S., Fourmanoir, H., Fougnyes, C., Deroanne, C., Herck, J.-C. Van,
771 Paquot, M., 2007. Isolation and identification of inulooligosaccharides resulting from
772 inulin hydrolysis. *Anal. Chim. Acta* 604, 81–87.
773 <https://doi.org/https://doi.org/10.1016/j.aca.2007.07.073>

774 Samaké, A., Jaffrezo, J.-L., Favez, O., Weber, S., Jacob, V., Albinet, A., Riffault, V., Perdrix,
775 E., Waked, A., Golly, B., Salameh, D., Chevrier, F., Oliveira, D.M., Besombes, J.-L.,
776 Martins, J.M.F., Conil, S., Guillaud, G., Meshba, B., Rocq, B., Robic, P.-Y., Hulin, A.,
777 Le Meur, S., Descheemaeker, M., Chretien, E., Uzu, G., 2019. Polyols and glucose
778 particulate species as tracers of primary biogenic organic aerosols at 28 french sites.

779 Atmos. Chem. Phys. 19, 3357–3374. <https://doi.org/10.5194/acp-2018-773>

780 Schädel, C., Blöchl, A., Richter, A., Hoch, G., 2010. Quantification and monosaccharide
781 composition of hemicelluloses from different plant functional types. *Plant Physiol.*
782 *Biochem.* 48, 1–8. <https://doi.org/10.1016/j.plaphy.2009.09.008>

783 Sempéré, R., Tedetti, M., Panagiotopoulos, C., Charriere, B., Van Wambeke, F., 2008.
784 Distribution and bacterial availability of dissolved neutral sugars in the South East
785 Pacific. *Biogeosciences* 5, 1165–1173. <https://doi.org/10.5194/bg-5-1165-2008>

786 Simoneit, B.R.T., Kobayashi, M., Kawamura, K., Rushdi, A.I., Rogge, W.F., Didyk, B.M.,
787 2004. Sugars - Dominant Water-Soluble Organic Compounds in Soils and
788 Characterization as Tracers in Atmospheric Particulate Matter. *Environ. Sci. Technol.*
789 38, 5939–5949. <https://doi.org/10.1021/es0403099>

790 Simoneit, B.R.T., Schauer, J.J., Nolte, C.G., Oros, D.R., Elias, V.O., Fraser, M.P., Rogge,
791 W.F., Cass, G.R., 1999. Levoglucosan, a tracer for cellulose in biomass burning and
792 atmospheric particles. *Atmos. Environ.* 33, 173–182. [https://doi.org/10.1016/S1352-](https://doi.org/10.1016/S1352-2310(98)00145-9)
793 [2310\(98\)00145-9](https://doi.org/10.1016/S1352-2310(98)00145-9)

794 Skoog, A., Benner, R., 1997. Aldoses in various size fractions of marine organic matter :
795 Implications for carbon cycling. *Limnol. Oceanogr.* 42, 1803–1813.
796 <https://doi.org/10.4319/lo.1997.42.8.1803>

797 Sullivan, A.P., Frank, N., Kenski, D.M., Collett, J.L., 2011. Application of high-performance
798 anion-exchange chromatography-pulsed amperometric detection for measuring
799 carbohydrates in routine daily filter samples collected by a national network: 2.
800 Examination of sugar alcohols/polyols, sugars, and anhydrosugars in. *J. Geophys. Res.*
801 *Atmos.* 116. <https://doi.org/10.1029/2010JD014169>

802 Thermo Fisher, technical Note 20, 2004. Analysis of Carbohydrates by High-Performance

803 Anion-Exchange Chromatography with Pulsed Amperometric Detection (HPAE-PAD).
804 Dionex.

805 Theodosi, C., Panagiotopoulos, C., Nouara, A., Zarmpas, P., Nicolaou, P., Violaki, K.,
806 Kanakidou, M., Sempéré, R., Mihalopoulos, N., 2018. Sugars in atmospheric aerosols
807 over the Eastern Mediterranean. *Prog. Oceanogr.* 163, 70–81.
808 <https://doi.org/10.1016/j.pocean.2017.09.001>

809 Tsai, Y.I., Sopajaree, K., Kuo, S.-C., Yu, S.-P., 2015. Potential PM_{2.5} impacts of festival-
810 related burning and other inputs on air quality in an urban area of southern Taiwan. *Sci.*
811 *Total Environ.* 527–528, 65–79. <https://doi.org/10.1016/j.scitotenv.2015.04.021>

812 van Pinxteren, M., Müller, C., Iinuma, Y., Stolle, C., Herrmann, H., 2012. Chemical
813 characterization of dissolved organic compounds from coastal sea surface microlayers
814 (Baltic Sea, Germany). *Environ. Sci. Technol.* 46, 10455–10462.
815 <https://doi.org/10.1021/es204492b>

816 Vandeska, E., Amartey, S., Kuzmanova, S., Jeffries, T., 1995. Effects of environmental
817 conditions on production of xylitol by *Candida boidinii*. *World J. Microbiol. Biotechnol.*
818 11, 213–218. <https://doi.org/10.1007/BF00704652>

819 Wang, Q.-C., Zhao, X., Pu, J.-H., Luan, X.-H., 2016. Influences of acidic reaction and
820 hydrolytic conditions on monosaccharide composition analysis of acidic, neutral and
821 basic polysaccharides. *Carbohydr. Polym.* 143, 296–300.
822 <https://doi.org/10.1016/j.carbpol.2016.02.023>

823 Weckesser, J., Mayer, H., Drews, G., 1970. The identification of 3-O-methyl-L-rhamnose (L-
824 acofriose) as constituent of the lipopolysaccharide of *Rhodopseudomonas capsulata*. *Eur.*
825 *J. Biochem* 16, 158–160. <https://doi.org/10.1111/j.1432-1033.1970.tb01067.x>

826 Wicks, R., Moran, M., Pittman, L., Hodson, R., 1991. Carbohydrate signatures of aquatic

827 macrophytes and their dissolved degradation products as determined by a sensitive high-
828 performance ion chromatography method. *Appl. Environ. Microbiol.* 57, 3135–3143.

829 Yang, H., Yu, J.Z., Ho, S.S.H., Xu, J., Wu, W.-S., Wan, C.H., Wang, X., Wang, X., Wang,
830 L., 2005. The chemical composition of inorganic and carbonaceous materials in PM_{2.5}
831 in Nanjing, China. *Atmos. Environ.* 39, 3735–3749.
832 <https://doi.org/10.1016/j.atmosenv.2005.03.010>

833 Yttri, K.E., Dye, C., Kiss, G., 2007. Ambient aerosol concentrations of sugars and sugar-
834 alcohols at four different sites in Norway. *Atmos. Chem. Phys. Discuss.* 7, 5769–5803.
835 <https://doi.org/10.5194/acpd-7-5769-2007>

836 Zdrovenko, G.M., Zdrovenko, E.L., Varbanets, L.D., 2007. Composition, structure, and
837 biological properties of lipopolysaccharides from different strains of *Pseudomonas*
838 *syringae* pv. *atrofaciens*. *Microbiology* 76, 683–697.
839 <https://doi.org/10.1134/S0026261707060069>

840 Zhang, Z.-S., Engling, G., Chan, C.-Y., Yang, Y.-H., Lin, M., Shi, S., He, J., Li, Y.-D., Wang,
841 X.-M., 2013. Determination of isoprene-derived secondary organic aerosol tracers (2-
842 methyltetrols) by HPAEC-PAD: Results from size-resolved aerosols in a tropical
843 rainforest. *Atmos. Environ.* 70, 468–476. <https://doi.org/10.1016/j.atmosenv.2013.01.020>

844 Zheng, L., Yang, X., Lai, S., Ren, H., Yue, S., Zhang, Y., Huang, X., Gao, Y., Sun, Y., Wang,
845 Z., Fu, P., 2018. Impacts of springtime biomass burning in the northern Southeast Asia
846 on marine organic aerosols over the Gulf of Tonkin, China. *Environ. Pollut.* 237, 285–
847 297. <https://doi.org/https://doi.org/10.1016/j.envpol.2018.01.089>

848

849

850

851 **Table and Figure captions**

852 **Table 1:** Optimal analytical conditions for the simultaneous analysis of neutral sugars,
853 alditols, anhydrosugars and sucrose

854 **Table 2.** Carbohydrate composition in atmospheric TSP, marine POM and marine
855 HMWDOM samples. The carbohydrate concentrations presented in this table are not
856 corrected for losses during the hydrolysis or the extraction procedure (see materials and
857 methods), and therefore they should be considered as minimum concentrations found in
858 the samples.

859 **Fig. 1.** HPAEC-PAD chromatograms of a standard mixture of 17 monosaccharides (1 μM
860 each) obtained with the (a) CarboPac MA1 column after optimization according to Table
861 1 (Peak identification: 1 = xylitol, 2 = levoglucosan, 3 = arabitol, 4 = fucose/ rhamnose,
862 5 = sorbitol, 6 = mannosan, 7 = mannitol, 8 = galactosan, 9 = arabinose, 10 = mannose,
863 11 = glucose, 12 = xylose, 13 = galactose, 14 = fructose, 15 = ribose, 16 = sucrose) and
864 (b) CarboPac PA1 column (Peak identification: 1 = xylitol, 2 = arabitol/levoglucosan, 3
865 = sorbitol, 4 = mannitol, 5 = mannosan, 6 = galactosan, 7 = fucose, 8 =
866 rhamnose/arabinose, 9 = galactose, 10 = glucose, 11 = mannose/xylose, 12 = fructose, 13
867 = ribose/sucrose). The dotted line and secondary axis indicate the NaOH gradient. The
868 analytical HPAEC-PAD conditions for the CarboPac PA1 column are as follows: flow
869 rate 0.7 mL min^{-1} , 0–15 min : 1 mM NaOH; 15–38 min : 19 mM NaOH (curve 5),
870 column temperature $17 \text{ }^\circ\text{C}$ and detector temperature $20 \text{ }^\circ\text{C}$ (Theodosi et al., 2018).

871 **Fig. 2.** The resolution of levoglucosan/arabitol (\blacklozenge), fucose/rhamnose (\times), galactosan/arabinose
872 (\blacksquare); arabinose/mannose (\blacktriangle), and glucose/xylose (\bullet) pairs with fixed temperatures (25,
873 28 and $30 \text{ }^\circ\text{C}$) applied during the whole HPAEC-PAD run under the optimized mobile
874 phase and flow rate conditions.

875 **Fig. 3.** HPAEC-PAD chromatograms according to Table 1 of (a) total suspended atmospheric
876 particles (TSP), (b) marine particulate organic matter (POM) and (c) marine high-
877 molecular-weight dissolved organic matter (HMWDOM). Peak identification: 1 =
878 xylitol, 2 = levoglucosan, 3 = arabitol, 4 = fucose/rhamnose, 5 = sorbitol, 6 = mannosan,
879 7 = mannitol, 8 = galactosan, 9 = arabinose, 10 = mannose, 11 = glucose, 12 = xylose,
880 13 = galactose, 14 = fructose, 15 = ribose, 16 = sucrose.

881 **Fig. 4.** Relative abundances (%) of neutral sugars, alditols, anhydrosugars and disaccharides
882 within the TCHO pool for (a) total suspended atmospheric particles (TSP), (b) marine
883 particulate organic matter (POM) and (c) marine high-molecular-weight dissolved
884 organic matter (HMWDOM). The relative abundances (%) of levoglucosan, galactosan
885 and mannosan within the anhydrosugar pool are given as well for the abovementioned
886 environmental samples.

887

888

889

890

891

892

893

894

895

896

897

898

899

900 **Table 1**901 Optimal analytical conditions for the simultaneous analysis of neutral sugars, alditols, anhydrosugars
902 and sucrose.

Parameters	Conditions	
Column	CarboPac MA1 (250 × 4 mm I.D.), Thermo Fisher	
Eluent flow rate (mL min ⁻¹)	0.3	
Detector temperature (°C)	25	
Sample loop size (μL)	250	
Injection volume (μL)	230	
Column oven temperature (°C)	25–28	(0–30 min); curve 5
	28	(30–75 min)
	25	(75–95 min)
NaOH gradient (mM)	250–350	(0–30 min) ; curve 5
	350–450	(30–45 min) ; curve 5
	450–700	(45–55 min) ; curve 5
	700	(55–75 min)
	250	(75–95 min)

903

904

905 **Table 2**906 Carbohydrate composition in atmospheric TSP, marine POM and marine HMWDOM samples. The
907 carbohydrate concentrations presented in this table are not corrected for losses during the hydrolysis or
908 the extraction procedure (see materials and methods), and therefore they should be considered as
909 minimum concentrations found in the samples.

Carbohydrate	TSP (ng m ⁻³)	POM (mg g ⁻¹)	HMWDOM (μg g ⁻¹)
Xylitol	11	0.05	0.13*
Levoglucozan	261	0.08	12
Arabitol	3.4	ND	ND
Fucose & Rhamnose	5.0	1.8	300
Sorbitol	1.9	0.02	ND
Mannosan	30	0.04	3.4
Mannitol	7.8	ND	2.3
Galactosan	9.4	0.2	4.4
Arabinose	2.1	0.1	34
Mannose	1.2	0.8	103
Glucose	15	3.8	122
Xylose	1.2	0.9	74
Galactose	ND	5.7	83
Fructose	36	2.0	11
Ribose	ND	0.7	19
Sucrose	178	ND	ND

910 ND = Not detected

911 * Xylitol concentration was outside the calibration range (50-1000 nM) but ~6 times higher than its
912 detection limit (Table A 1), thus it was quantified with an uncertainty of ~24%.

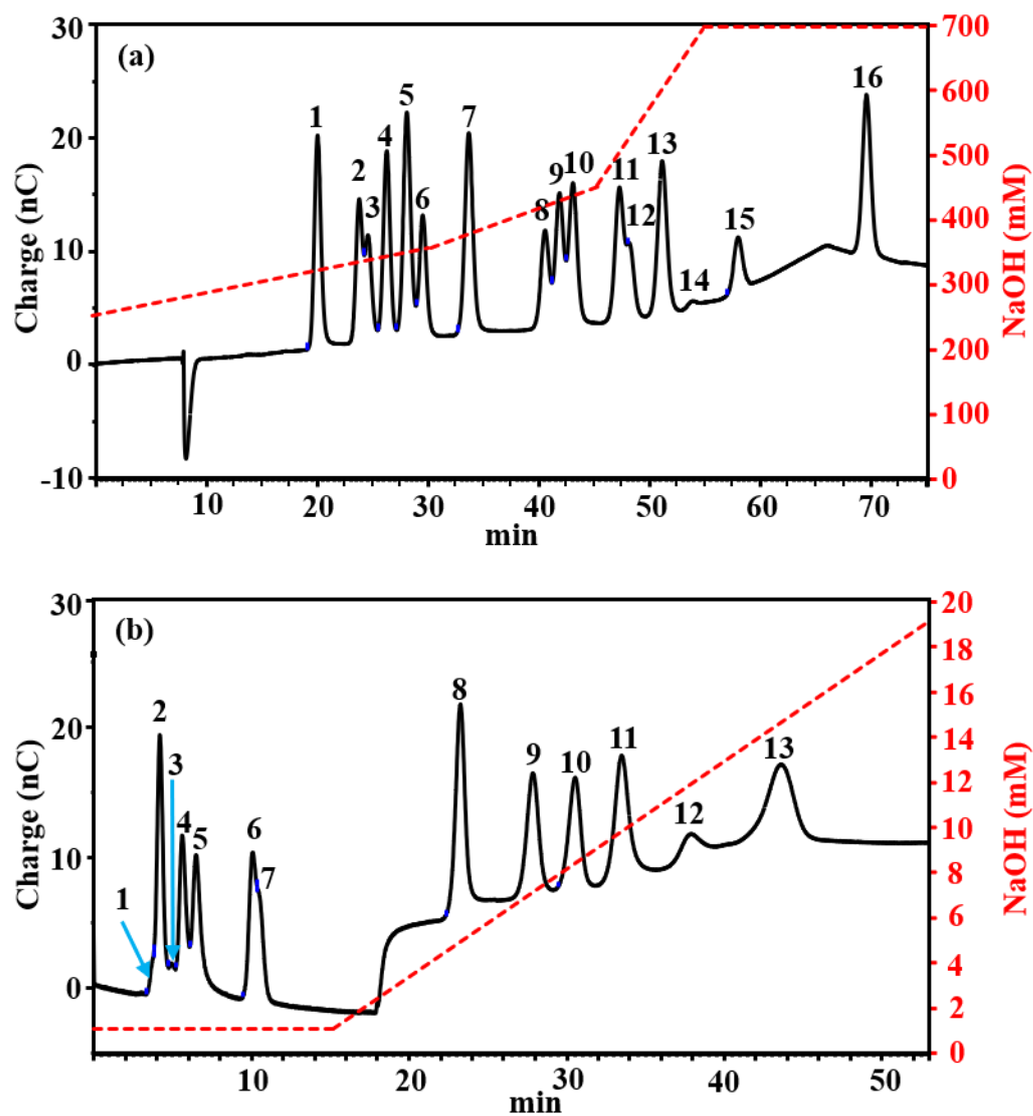


FIGURE 1

913

914

915

916

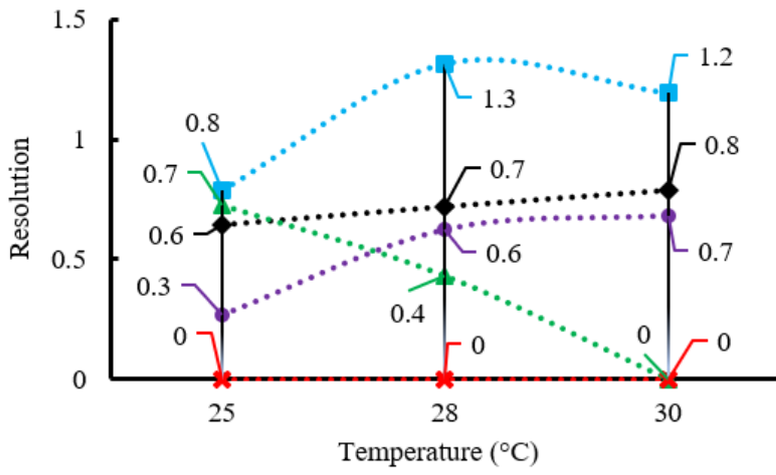
917

918

919

920

921



922

923

FIGURE 2

924

925

926

927

928

929

930

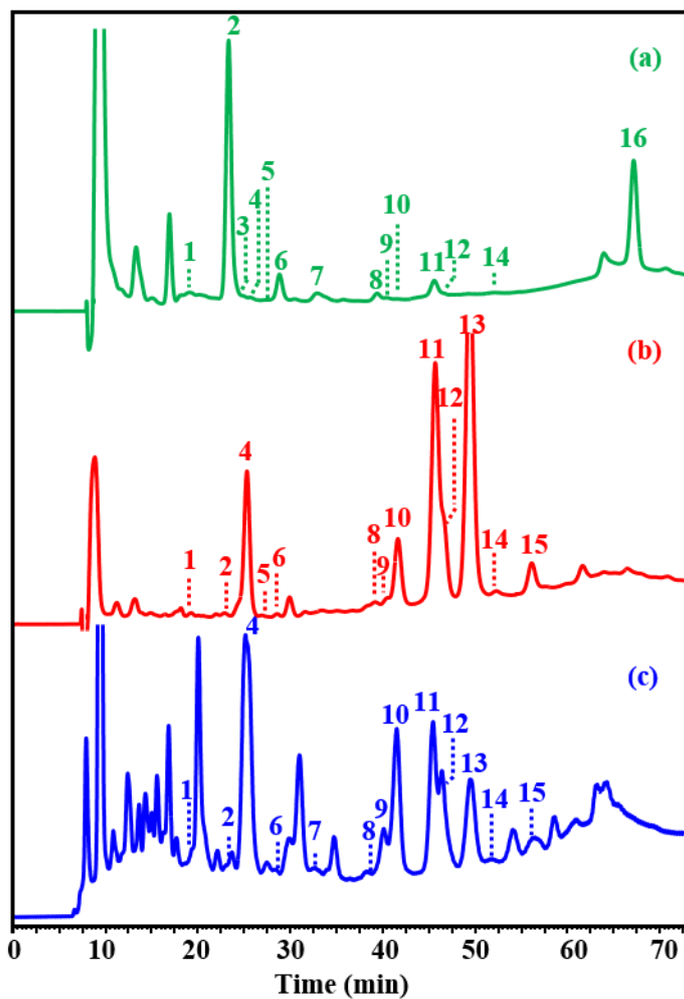
931

932

933

934

935



936

937

FIGURE 3

938

939

940

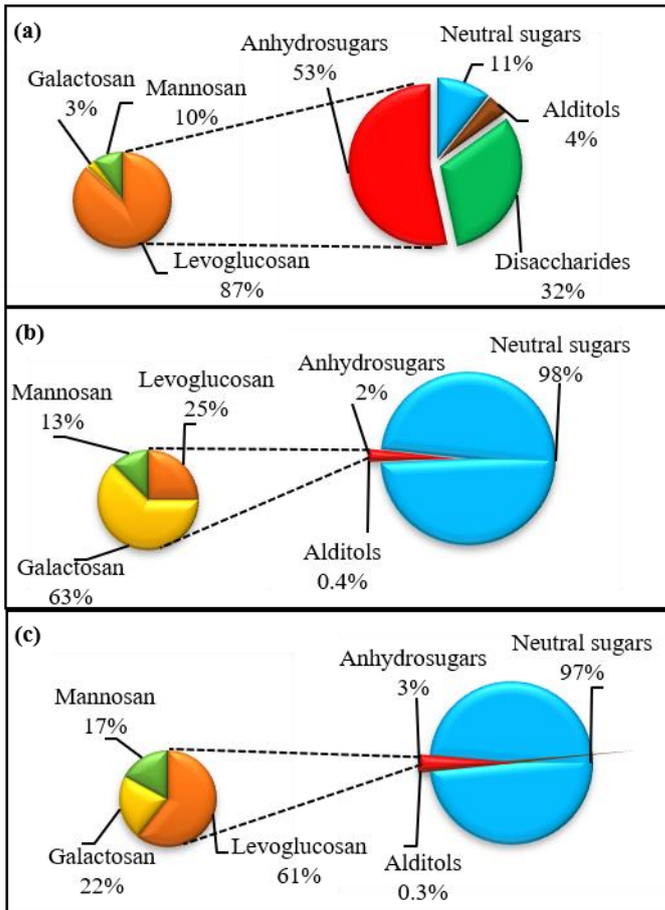
941

942

943

944

945



946

947

FIGURE 4

948

949

950

951

952

953

954

955

956

957
958
959
960
961
962
963
964
965
966
967
968
969
970
971
972
973
974
975
976
977
978

Supplemental Information for

Simultaneous determination of neutral sugars, alditols, and anhydrosugars using anion-exchange chromatography with pulsed amperometric detection: Application for marine and atmospheric samples

Amel Nouara, Christos Panagiotopoulos*, Richard Sempéré

Aix Marseille Univ., Université de Toulon, CNRS, IRD, MIO UM 110, 13288, Marseille, France

*Correspondence : Christos Panagiotopoulos

[e-mail : christos.panagiotopoulos@mio.osupytheas.fr](mailto:christos.panagiotopoulos@mio.osupytheas.fr)

09th May 2019

979 **Table of Contents**

980

981 **Table S 1.** HPAEC-PAD performance after optimization (Table 1). The average retention time (t_R),
982 and resolution factor (R_s) of the carbohydrate standards is given at 1 μ M level ($n = 3$). The relative
983 standard deviations (RSD) of peak area and retention time of the same standards is calculated at 50
984 nM level ($n = 6$), except for fructose (1 μ M level; $n = 6$). The detection limit (DL) was calculated at
985 signal to noise ratio of 3.....45

986

987 **Fig S 1.** Example of HPAEC-PAD chromatograms of procedural blanks (a) ultrapure water (b) TSP
988 blank and (c) POM blank. Blanks were prepared in triplicate for each sample (TSP and marine POM)
989 and run according to the method established in Table 1.....46

990

991 **Fig S 2.** High-performance liquid chromatography with refractive index detection (HPLC-RI)
992 chromatograms of a monosaccharide mixture (1 mM each) before hydrolysis (black solid line) and
993 after hydrolysis (red dotted line) with 1 M HCl under 100 °C for 20 h. Peak identification : 1 =
994 glucose, 2 = xylose, 3 = galactose & rhamnose, 4 = galactosan, 5 = arabinose & fucose, 6 = mannose,
995 7 = fructose, 8 = mannitol, 9 = levoglucosan, 10 = xylitol, 11 = sorbitol, 12 = mannosan. The
996 analytical HPLC-RI conditions are given in Nouara et al. (2019) and are as follows: cation exchange
997 column in Pb^{2+} form, isocratic elution with ultrapure water at 0.6 mL min^{-1} , column temperature 75
998 °C.47

999

1000

1001

1002

1003

1004

1005

1006

1007 **Table S 1.**

1008 HPAEC-PAD performance after optimization (Table 1). The average retention time (t_R), and
 1009 resolution factor (R_s) of the carbohydrate standards is given at 1 μ M level ($n = 3$). The relative
 1010 standard deviations (RSD) of peak area and retention time of the same standards is calculated at 50
 1011 nM level ($n = 6$), except for fructose (1 μ M level; $n = 6$). The detection limit (DL) was calculated at
 1012 signal to noise ratio of 3.

Carbohydrate	t_R (min)	R_s	Area RSD (%)	t_R RSD (%)	DL (nM)
Xylitol	19.9	3.1	5.6	0.2	2
Levoglucozan	23.8	0.6	4.0	0.5	8
Arabitol	24.4	1.3	6.7	0.4	13
Fucose & Rhamnose	26.0	1.3	6.5	0.3	17
Sorbitol	27.8	1.2	5.0	0.3	15
Mannosan	29.3	3.2	1.9	0.3	18
Mannitol	33.3	5.1	2.7	0.4	7
Galactosan	40.2	0.8	9.3	0.4	32
Arabinose	41.4	0.7	4.9	0.4	7
Mannose	42.6	2.4	1.8	0.3	6
Glucose	46.7	0.5	4.4	0.3	9
Xylose	47.7	1.6	6.7	0.3	24
Galactose	50.5	1.7	5.2	0.6	4
Fructose	53.1	3.0	16.6	1.7	51
Ribose	57.2	8.1	6.0	0.4	13
Sucrose	68.9	-	5.8	0.7	13

1013

1014

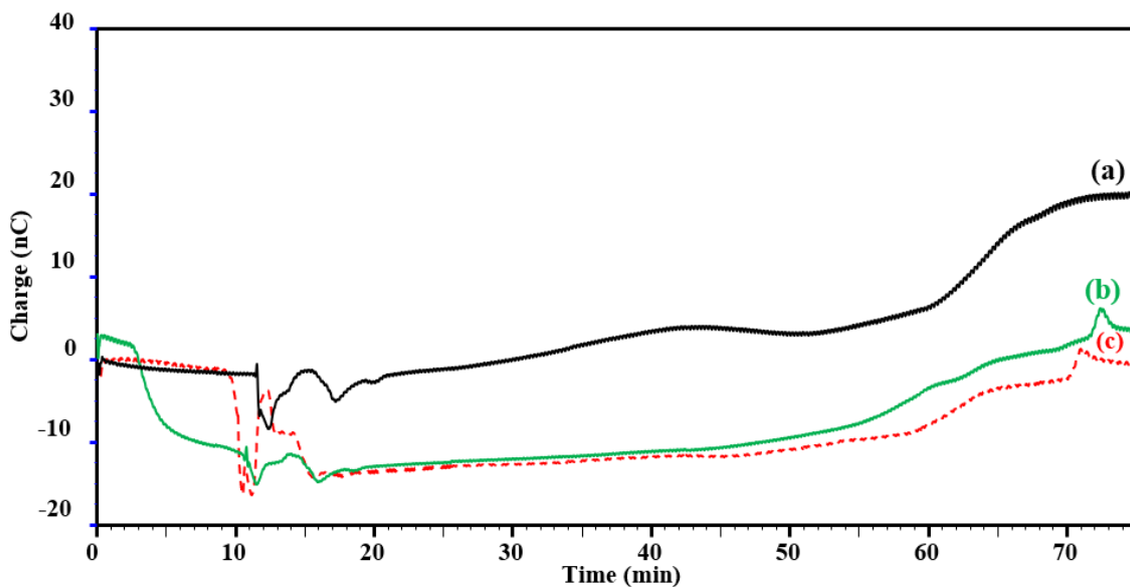
1015

1016

1017

1018

1019



1020

1021 **Fig. S 1.** Example of HPAEC-PAD chromatograms of procedural blanks (a) ultrapure water (b)
 1022 TSP blank and (c) POM blank. Blanks were prepared in triplicate for each sample (TSP and marine
 1023 POM) and run according to the method established in Table 1.

1024

1025

1026

1027

1028

1029

1030

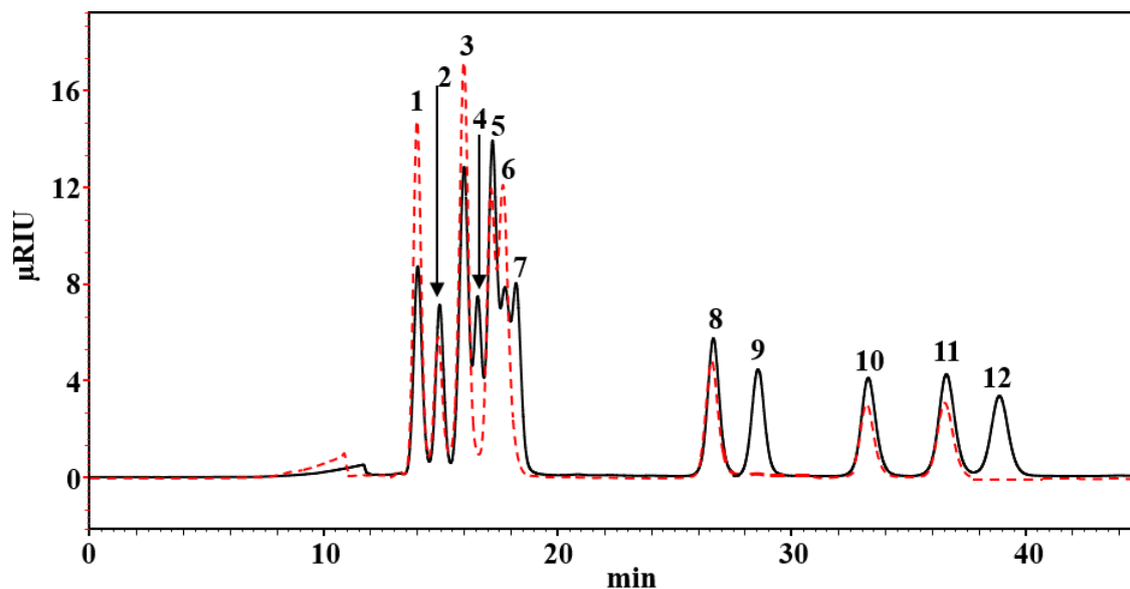
1031

1032

1033

1034

1035



1036

1037 **Fig. S 2.** High-performance liquid chromatography with refractive index detection (HPLC-RI)
1038 chromatograms of a monosaccharide mixture (1 mM each) before hydrolysis (black solid line) and
1039 after hydrolysis (red dotted line) with 1 M HCl under 100 °C for 20 h. Peak identification : 1 =
1040 glucose, 2 = xylose, 3 = galactose & rhamnose, 4 = galactosan, 5 = arabinose & fucose, 6 = mannose,
1041 7 = fructose, 8 = mannitol, 9 = levoglucosan, 10 = xylitol, 11 = sorbitol, 12 = mannosan. The
1042 analytical HPLC-RI conditions are given in Nouara et al. (2019) and are as follows: cation exchange
1043 column in Pb²⁺ form, isocratic elution with ultrapure water at 0.6 mL min⁻¹, column temperature 75
1044 °C.

1045

1046