

1 **Amino acids, carbohydrates and lipids in the tropical oligotrophic Atlantic**  
2 **Ocean: Sea-to-air transfer and atmospheric in situ formation**

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5 **Manuela van Pinxteren<sup>1</sup>, Sebastian Zeppenfeld<sup>1</sup>, Khanneh Wadinga Fomba<sup>1</sup>, Nadja Triesch<sup>1+</sup>,**  
6 **Sanja Frka<sup>2</sup>, Hartmut Herrmann<sup>1\*</sup>**

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8 \*Corresponding author: Hartmut Herrmann ([herrmann@tropos.de](mailto:herrmann@tropos.de))  
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10 <sup>1</sup> Atmospheric Chemistry Department (ACD), Leibniz-Institute for Tropospheric Research  
11 (TROPOS), 04318 Leipzig, Germany

12 <sup>+</sup> now at: Department Safety in the Food Chain, German Federal Institute for Risk Assessment,  
13 10589 Berlin, Germany

14 <sup>2</sup> Division for Marine and Environmental Research, Ruđer Bošković Institute, 10000 Zagreb,  
15 Croatia  
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37 **Abstract**

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39 This study examines carbohydrates, amino acids, and lipids as important contributors to  
40 organic carbon (OC) in the tropical Atlantic Ocean at the Cape Verde Atmospheric Observatory  
41 (CVAO). The above compounds were measured in both surface seawater and in ambient  
42 submicron aerosol particles to investigate their sea-to-air transfer, including their enrichment  
43 in the sea surface microlayer (SML), potential atmospheric in situ formation or degradation,  
44 and their oceanic contribution to the ambient marine aerosol particles.

45 In bulk seawater and the SML, similar distributions among species were found for the lipids  
46 and carbohydrates with moderate SML enrichments (enrichment factor  $EF_{SML} = 1.3 \pm 0.2$  and  
47  $1.1 \pm 0.5$  respectively). In contrast, the amino acids exhibited a higher enrichment in the SML  
48 with an average  $EF_{SML}$  of  $2.3 \pm 0.4$  although they are less surface-active than lipids. The same  
49 compounds studied in the seawater were found on the ambient submicron aerosol particles  
50 whereas the lipids were more pronounced enriched ( $EF_{aer.} = 1.6 \times 10^5$ ) compared to the amino  
51 acids and carbohydrates ( $EF_{aer.} = 1.5 \times 10^3$  and  $1.3 \times 10^3$  respectively), likely due to their high  
52 surface activity and/or the lipophilic character. Detailed molecular analysis of the seawater  
53 and aerosol particles revealed changes in the relative abundance of the individual organic  
54 compounds. They were most pronounced for the amino acids and are likely related to an in  
55 situ atmospheric processing by biotic and/or abiotic reactions.

56 On average 49% of the OC on the aerosol particles ( $\cong 97 \text{ ng m}^{-3}$ ) could be attributed to the  
57 specific components or component groups investigated in this study. The majority (43%) was  
58 composed of lipids. Carbohydrates and amino acids made up less than 1% of the OC. This  
59 shows that carbohydrates, at least when resolved via molecular measurements of single  
60 sugars, do not comprise a very large fraction of OC on marine aerosol particles, in contrast to  
61 other studies. However, carbohydrate-like compounds are also present in the high lipid  
62 fraction (e.g., as glycolipids), but their chemical composition could not be revealed by the  
63 measurements performed here.

64 Previously determined OC components at the CVAO, specifically amines, oxalic acid, and  
65 carbonyls, comprised an OC fraction of around 6%.

66 Since the identified compounds constituted about 50% of the OC and belong to the rather  
67 short-lived biogenic material probably originating from the surface ocean, a pronounced  
68 coupling between ocean and atmosphere was indicated for this oligotrophic region. The  
69 remaining, non-identified OC fraction might in part contain recalcitrant OC, however, this  
70 fraction does not constitute the vast majority of OC in the aerosol particles here investigated.

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73 **Keywords:** organic carbon, lipids, amino acids, carbohydrates, sea surface microlayer, aerosol  
74 particles, Atlantic Ocean, CVAO

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## 78 **1 Introduction**

79 Marine aerosol particles, their composition, sources and connection to the upper ocean are  
80 not yet fully understood, however important as they impacts the carbon cycle and radiative  
81 properties of aerosol particles (Abbatt et al., 2019;Brooks and Thornton, 2018;Burrows et al.,  
82 2013;Gantt and Meskhidze, 2013;Pagnone et al., 2019;Patel and Rastogi, 2020). Furthermore,  
83 the function of (marine) aerosol particles as cloud condensation nuclei (CCN) and ice-  
84 nucleating particles (INP), i.e. marine aerosol-cloud interaction is still elusive. Important  
85 information about marine CCN is still missing (e.g. Bertram et al., 2018). Ocean-derived INPs  
86 have been suggested to play a dominant role in determining INP concentrations in near-  
87 surface air over remote areas such as the Southern Ocean, but their source strength in other  
88 oceanic regions and knowledge of which physicochemical properties determine INP efficiency  
89 remain largely unknown (Burrows et al., 2013;McCluskey et al., 2018a;McCluskey et al.,  
90 2018b). Marine aerosol particles, notably in the sub-micrometer range, have been shown to  
91 contain a large part of organic carbon (OC) in field experiments as well as in laboratory studies,  
92 where nascent aerosol particles are generated by artificial bubble-bursting mechanisms  
93 (Facchini et al., 2008;Keene et al., 2007;O'Dowd et al., 2004). Notably, the laboratory  
94 experiments, where sources other than the ocean (such as long-range transport) can be  
95 excluded, suggest that a certain part of the OC on the aerosol particles is transferred directly  
96 from the ocean via bubble bursting (Facchini et al., 2008;Keene et al., 2007). The mechanisms  
97 of the OC enrichment finally observed in aerosol particles are not yet fully understood but are  
98 likely due to complex interaction at the ocean surface when air bubbles rise and break. Air  
99 bubbles collect (organic) matter at their surface (the gas/water interface) when they ascend  
100 through the water column and when bursting, they produce film and jet droplets that transfer  
101 the OC to the atmosphere and form aerosol particles. At the ocean surface, the air bubbles  
102 enter the uppermost layer and the direct interface between the ocean and the atmosphere  
103 called the sea surface microlayer (SML) (Engel et al., 2017). The SML is described as a gel-like  
104 matrix that accumulates various organic and inorganic material (Cunliffe et al., 2013). The  
105 influence of the SML on the bubble bursting and the emission of OC into the atmosphere is  
106 difficult to determine and still controversial (Engel et al., 2017).

107 Based on the OC to sodium ratios in the ocean and the atmosphere, the OC in marine  
108 aerosol particles has shown to be strongly enriched compared to seawater concentrations. OC  
109 aerosol enrichment factors ( $EF_{aer.}$ ) of the order of  $10^2$  in supermicron aerosol particles and of  
110 the orders of  $10^3$  to  $10^5$  in submicron aerosol particles have been reported (Quinn et al., 2015  
111 and references therein). However, individual chemical groups, such as amino acids, can be  
112 even more enriched and  $EF_{aer.}$  as high as  $10^7$  for these particular compounds have been  
113 measured in submicron particles resulting from bubble bursting experiments within a tank  
114 study (Triesch et al., 2021c). The OC transfer from the ocean to the atmosphere is likely highly  
115 chemo-selective and a hydrophobic nature as well as surface-active properties of organic  
116 compounds probably favour their transfer from the sea to the air (Rastelli et al., 2017;Schmitt-  
117 Kopplin et al., 2012).

118 An important component to understand the OC on marine aerosol particles is the  
119 connection to oceanic bio-productivity. Several studies suggested that the marine aerosol  
120 composition is directly coupled to the productivity in the ocean, showing that at elevated  
121 chlorophyll-a (chl-a) concentrations in the seawater the OC on the aerosol particles is  
122 significantly higher when compared to the low oceanic productivity (O'Dowd et al.,  
123 2004;Facchini et al., 2008). A coupling between oceanic bio-productivity and aerosol  
124 composition is probably not straightforward. Wang et al., (2015) showed that two successive  
125 phytoplankton blooms in the tank seawater resulted in sea spray aerosols (SSA) with vastly  
126 different compositions and properties. Other studies, however, propose that the OC transfer  
127 from the ocean to aerosol particles is non-correlated to oceanic bio-productivity. Quinn et al.,  
128 (2014) suggested that the high reservoir of dissolved organic carbon (DOC) in the ocean is  
129 responsible for the organic enrichment in freshly emitted sea spray aerosol, thus dominating  
130 over any influence of recent local biological activity based on chlorophyll concentrations.  
131 Following this, Kieber et al., (2016) proposed that the major component in submicron sea-  
132 spray particles is of recalcitrant nature with a stability of months to millennia. They suggested  
133 that this persistent form of OC can very efficiently be transferred to the atmosphere via bubble  
134 bursting. Although they did not perform a detailed chemical analysis, they concluded that the  
135 recalcitrant organic matter exhibits surface-active properties. Applying natural abundance  
136 radiocarbon ( $^{14}\text{C}$ ) measurements it was recently suggested that 19 to 40% of the OC associated  
137 with freshly produced marine aerosol particles was refractory dissolved organic carbon (rDOC)  
138 (Beaupre et al., 2019).

139 In addition to the direct, or primary transfer of organic compounds from the ocean to  
140 the atmosphere, atmospheric processing changes the composition. Once released from the  
141 ocean to the atmosphere, organic matter can be acidified within seconds due to a pH change  
142 in the atmospheric particles or undergo fast photochemical oxidation (Kieber et al., 2016).  
143 Moreover, biogenic in situ formation and degradation can change the OC composition in  
144 marine aerosol particles and marine cloud water (Bianco et al., 2019;Malfatti et al.,  
145 2019;Matulova et al., 2014). Ervens and Amato (2020) provided a framework for estimating  
146 the production of secondary biological aerosol mass in clouds through microbial cell growth  
147 and multiplication. This pathway could be a significant source of biological aerosol material  
148 (Ervens and Amato, 2020;Khaled et al., 2021;Zhang et al., 2021). In other recent studies, the  
149 in situ formation of amino acids by biotic and abiotic processes in cloud water was measured  
150 and modelled (Jaber et al., 2021) and gel-like, organic particles, originally present in the ocean,  
151 were suggested to form in situ in the marine atmosphere via biotic and/or abiotic pathways  
152 (Haddrell and Thomas, 2017;Klein et al., 2016;van Pinxteren et al., 2022). Nevertheless,  
153 despite some studies, the atmospheric in situ transformation of marine organic compounds  
154 and its significance has not yet been extensively studied.

155 To understand the transfer processes of OC from the ocean to the aerosol particles,  
156 potential atmospheric OC in situ formation as well as the coupling of the OC on the aerosol  
157 particles to processes in the ocean, it is crucial to unravel the chemical composition of the  
158 aerosol OC content. In the present study, we investigated samples from the tropical Atlantic  
159 Ocean at the CVAO. The focus of this study was on the analysis of amino acids and

160 carbohydrates, as well as of lipid components, as these OC groups are reported as the major  
161 marine organic matter groups in the seawater and therefore likely transferred to the aerosol  
162 particles via bubble bursting (Burrows et al., 2014). We investigated these compounds on  
163 marine aerosol particles and in the ocean SML and bulk water. Specifically, we followed the  
164 concentrations and speciation of OC from the sea to the atmospheric particles, which helps to  
165 evaluate the enrichment factors of the various targeted compounds between the  
166 compartments. The results will help to gain a better understanding of the chemical  
167 composition of marine aerosol particles in this tropical location, its transfer from the ocean  
168 and in situ formation, and finally, help to elucidate the coupling of marine aerosol particles to  
169 the surface ocean in an oligotrophic region.

170

## 171 **2. Material and methods**

### 172 **2.1 Aerosol and seawater sampling during the campaign**

173

174 A field campaign (MarParCloud) was carried out at the Cape Verde Atmospheric Observatory  
175 (CVAO, 16°51'49'N, 24°52'02'E) in autumn 2017 (13.09.2017 – 13.10.2017) and the sampling  
176 sites are illustrated and explained in detail in van Pinxteren et al., (2020). The CVAO is a remote  
177 marine station in the tropical Atlantic Ocean located on the northeast coast of Sao Vicente  
178 island and described in Carpenter et al., (2010) and Fomba et al., (2014). The ocean around  
179 the Cape Verde Islands has the lowest surface chlorophyll in the North Atlantic Ocean with  
180 values below 0.2  $\mu\text{g L}^{-1}$  for the majority of each year with periodic events of slightly elevated  
181 concentrations up to 0.7  $\mu\text{g L}^{-1}$  (van Pinxteren et al., 2020 and refs therein).

182 Submicron aerosol particles were sampled on preheated 150 mm quartz fiber filters  
183 (Munktell, MK 360) at a flow rate of about 700 L  $\text{min}^{-1}$  with a high volume  $\text{PM}_{10}$  aerosol sampler  
184 (Digitel, Riemer, Germany) installed on the 30 m height tower at the coastline. The sampling  
185 times were usually set to 24 h and are listed in Tab. S4 and S5, as well as in Triesch et al.,  
186 (2021b) and in van Pinxteren et al., (2020). Regarding the aerosol sampling system, it needs  
187 to be underlined that artefact problems and overestimations due to gas phase absorption and  
188 underestimations due to re-volatilisations of the analytes from the filters cannot be accounted  
189 for and represent a certain level of uncertainty.

190 The seawater samples were taken at Bahia das Gatas, a coastal site that is upwind and  
191 about 4 km northwest of the CVAO (Fig. S1). Fishing boats were rented to go out on the open  
192 ocean and the SML was sampled with a typical glass plate SML sampling strategy (Cunliffe and  
193 Wurl, 2014). To this end, a glass plate with a sampling area of 2000  $\text{cm}^2$  was vertically  
194 immersed into the water and then slowly drawn upwards with a withdrawal rate between 5  
195 and 10  $\text{cm s}^{-1}$ . The surface film adheres to the surface of the glass and is removed using framed  
196 Teflon wipers (Stolle et al., 2010; van Pinxteren et al., 2012). Bulk seawater was collected with  
197 a specially designed device consisting of a plastic bottle mounted on a telescopic rod. The  
198 bottle was opened underwater at depth of 1 m with a specifically conceived seal-opener.

199 For the sampling of the oceanic water samples, great care was taken that all parts that  
200 were in contact with the sample (glass plate, bottles) underwent an intense cleaning with 10%

201 HCl and rinsing with ultrapure water (resistivity = 18.2 MΩ cm) prior to the campaign and in  
202 between sampling to avoid contamination and carry over problems.

203 After the sampling, all seawater and aerosol samples were stored frozen at -20°C,  
204 transported in a reefer at -20 °C to the TROPOS laboratories where they were stored frozen  
205 until analysis.

206

## 207 2.2 Chemical analysis

208

### 209 2.2.1. Seawater and aerosol analysis: general considerations

210

211 Within the seawater analysis, we measured the dissolved amino acids (DAA) and  
212 dissolved carbohydrates (DCHO) in the DOC fraction, as DOC represents by far the largest pool  
213 of organic material in the ocean (Riebesell et al., 2011). DOC is the fraction of OC that passes  
214 through a GF-F filter of 0.7 μm pore size (Wurl, 2009), however, filter pore sizes between 0.2  
215 and 1.2 μm are often applied e.g. due to requirements of the analytical system (e.g.  
216 (Romankevich, 1984; Zäncker et al., 2017). The lipid measurements (from the same samples)  
217 were taken from Triesch et al., (2021b), are included in the DOC fraction as well, being filtered  
218 via a 0.7 μm pore-sized filter (GF-F, Whatmann) and extracted in an organic solvent, and refer  
219 to dissolved lipids (DL).

220 For the aerosol particles (PM<sub>1</sub>), we focused on the water-soluble organic (WSOC)  
221 fraction of the amino acids (AA<sub>aer.</sub>) and carbohydrates (CHO<sub>aer.</sub>). The lipids, however, were  
222 extracted with an organic solvent and filtered (analogous to seawater), meaning they  
223 comprise the organic-soluble organic fraction of the aerosol particles (Lipids<sub>aer.</sub>).

224

### 225 2.2.2. Analytical methods

226

227 For the analysis of the DCHO and CHO<sub>aer.</sub> as well as and the DAA and AA<sub>aer.</sub>, molecular-  
228 resolved techniques were applied. DCHO in filtered (0.2 μm Millex syringe filters) and  
229 desalinated SML and bulk water samples as well as CHO<sub>aer.</sub>, in the filtered PM<sub>1</sub> extracts were  
230 quantified using High-Performance Anion-Exchange Chromatography coupled with Pulsed  
231 Amperometric Detection (HPAEC-PAD) (Zeppenfeld et al., 2020; Zeppenfeld et al., 2021). After  
232 an acid hydrolysis (0.8 M HCl, 100°C, 20 h) the monosaccharides fucose (Fuc), rhamnose (Rha),  
233 arabinose (Ara), galactose (Gal), glucose (Glc), xylose (Xyl), mannose (Man), galactosamine  
234 (GalN), glucosamine (GlcN), muramic acid (MurAc), galacturonic acid (GalAc) and glucuronic  
235 acid (GlcAc) were measured.

236 For the DAA analysis, seawater samples (25.5 mL) were desalinated and concentrated  
237 to a few mL as described in Triesch et al., (2021a). For the AA<sub>aer.</sub> from PM<sub>1</sub> particles, an aqueous  
238 extract of the amino acids was prepared by shaking a piece of the filter in 2 mL water. After a  
239 filtration step (filter pore size: 0.2 μm) 25 μL of ascorbic acid (20 mg mL<sup>-1</sup>, purity 99 %, Sigma-  
240 Aldrich, St. Louis, Missouri, USA) was added to a 200 μL aliquot of the desalted seawater /  
241 aqueous filter extract to avoid the oxidation of the obtained amino acids as suggested in  
242 Mandalakis et al., (2010). Following the addition of 250 μL HCl (Supra-quality,

243 ROTIPURAN®Supra 35%, Carl Roth, Karlsruhe, Germany), the hydrolysis was performed at  
244 110 °C for 20 h. After cooling to room temperature, the hydrolysed filtrate was evaporated,  
245 resolved in 500 µL milliQ-water (Millipore Elix 3 and Element A10, Merck Millipore, Darmstadt,  
246 Germany), filtered, derivatized using AccQ-Tag™ precolumn derivatization method (Waters,  
247 Eschborn, Germany), and measured by ultra-high performance liquid chromatography with  
248 electrospray ionization and Orbitrap mass spectrometry (UHPLC/ESI-Orbitrap-MS), as  
249 described in Triesch et al., (2021a). The analytes include the amino acids glycine (Gly), alanine  
250 (Ala), serine (Ser), glutamic acid (Glu), threonine (Thr), proline (Pro), tyrosine (Tyr), valine (Val),  
251 phenylalanine (Phe), aspartic acid (Asp), isoleucine (Ile), leucine (Leu), methionine (Met),  
252 glutamine (Gln) and  $\gamma$ -aminobutyric acid (GABA) (purity  $\geq$  99 %, Sigma-Aldrich, St. Louis,  
253 Missouri, USA).

254 The DL and Lipids<sub>aer.</sub> measurements were taken from Triesch et al., (2021b), where the  
255 analysis were done with a semi-molecular technique. For a better understanding of the data,  
256 a short description is given in the following: the dissolved fraction of seawater samples  
257 (obtained after filtration through pre-combusted 0.7 µm GF/F filters) and aerosol filters were  
258 extracted with dichloromethane according to specific protocols after adding 2-hexadecanone  
259 as an internal standard, as described in more detail in Triesch et al. (2021b). The extract was  
260 analysed with thin-layer chromatography (TLC). Lipid classes were separated on Chromarods  
261 SIII and calibrated with an external calibration with a mixture of standard lipids by a  
262 chromatograph flame ionisation detector (FID) Iatroscan MKVI (Iatron, Japan). The separation  
263 scheme included elution steps in the solvent systems with increasing polarity. The lipid classes  
264 included hydrocarbons (HC), fatty acid methyl esters (ME), free fatty acids (FFA), alcohols  
265 (ALC), 1,3-diacylglycerols (1,3 DG), 1,2-diacylglycerols (1,2 DG), monoacylglycerols (MG), wax  
266 esters (WE), triacylglycerols (TG), pigments (PIG), phospholipids (PP) including  
267 phosphatidylglycerols (PG), phosphatidylethanolamine (PE), phosphatidylcholines (PC), as  
268 well as glycolipids (GL) which cover sulfoquinovosyldiacylglycerols (SQDG), monogalactosyl-  
269 diacylglycerols (MGDG), digalactosyldiacylglycerols (DGDG) and sterols (ST). It needs to be  
270 underlined that, as no single lipid compound but rather lipid groups (based on varying polarity  
271 in the TLC system) were measured, the lipid results can be classified as analysis on a semi-  
272 molecular level.

273 Organic carbon (OC) on the aerosol particles (PM<sub>1</sub> samples) was measured by means  
274 of a thermal-optical method using the Sunset Laboratory Dual-Optical Carbonaceous Analyzer  
275 (Sunset Laboratory Inc., U.S.A.) from a filter piece with an area of 1.5 cm<sup>2</sup>. The EUSAAR 2  
276 temperature protocol was utilized, and a charring correction was applied (Cavalli et al., 2010).  
277 The correction value for pyrolytic carbon was determined based on measurements of a sample  
278 transmission using a 678 nm laser. Samples were thermally desorbed from the filter medium  
279 under an inert He-atmosphere followed by an oxidizing O<sub>2</sub>/He-atmosphere while applying  
280 carefully controlled heating ramps. A flame ionization detector was used to quantify methane  
281 following a catalytic methanation of CO<sub>2</sub>.

282 Sodium was measured from filtered (0.45 µm syringe filter), aqueous extracts of the  
283 PM<sub>1</sub> samples using ion chromatography (more details in Zeppenfeld et al., 2021 and van

284 Pinxteren et al., 2022). The details on sampling and the sample treatment are summarized in  
285 Tab. S1.

286 Atmospheric concentrations were calculated from the measured analyte  
287 concentrations on the filter or in the filter extract. The masses of the analytes on the filter were  
288 extrapolated to the total filter and related to the collected air volume that flowed through the  
289 filter.

290

### 291 2.2.3 Detection limits and blank handling

292

293 Detection limits were 0.5-2.5  $\mu\text{g L}^{-1}$  (DAA) and 0.1-0.2  $\text{pg m}^{-3}$  ( $\text{AA}_{aer.}$ ) (Triesch et al.,  
294 2021a;Triesch et al., 2021c) and 0.13-0.7  $\mu\text{g L}^{-1}$  (DCHO) and 0.5-5  $\text{pg m}^{-3}$  ( $\text{CHO}_{aer.}$ ) (Zeppenfeld  
295 et al., 2021). Regarding the lipids, the method was carefully optimized for seawater analysis  
296 (Gašparović et al., 2015, 2017) and adopted for aerosol particle analytics as described in  
297 Triesch et al., (2021b). The detection limits were determined as the analyte concentrations  
298 corresponding to a signal-to-noise (S/N) ratio of 3 ranged from 0.06 to 0.33  $\mu\text{g}$  as reported  
299 previously in Penezić et al. (2022).

300 For the seawater analysis, field blanks were taken by filling high-purity water in pre-  
301 cleaned plastic bottles and handled the same as the seawater samples. The seawater blanks  
302 were in general below 10% and for the lipids always below 15% (Triesch et al. 2021b). All data  
303 are blank corrected by subtracting the field blank values from the samples.

304 Field blanks for aerosol particles were prepared using pre-baked quartz fiber filters  
305 without an active sampling and treated according to the same procedure as the field samples.  
306 The concentrations of the target analytes were calculated by external calibration. Each sample  
307 was measured twice with a relative standard deviation of typically < 10 %, and field blanks,  
308 which were for most compounds negligible and for the lipid classes always below 20% of the  
309 real aerosol particle sample, were subtracted. All presented values are corrected for the field  
310 blank.

311

### 312 2.3 Enrichment factors

313

314 The SML enrichment factor ( $\text{EF}_{\text{SML}}$ ) was calculated by dividing the concentration of the  
315 analyte in the SML with the concentration of the analyte in the bulk water after equation (1):

316

$$317 \quad \text{EF}_{\text{SML}} = \frac{c(\text{analyte})_{\text{SML}}}{c(\text{analyte})_{\text{bulk water}}} \quad (1)$$

318

319 An enrichment in the SML is indicated with  $\text{EF}_{\text{SML}} > 1$  and a depletion in the SML with  $\text{EF}_{\text{SML}} < 1$ .

320 The enrichment factor of aerosol ( $\text{EF}_{aer.}$ ) is a quantitative metric for comparing  
321 compounds in the ocean and in the atmosphere. The  $\text{EF}_{aer.}$  concept is mainly applied to closed  
322 systems (Quinn et al., 2015 and refs.therein;Rastelli et al., 2017) as degradation and formation  
323 pathways on aerosol particles including photochemical and biotic atmospheric reactions and  
324 contributions from other (non-marine) sources are excluded from this parameter.



325 Nevertheless, for comparison purposes, it is useful to apply the  $EF_{aer.}$  to open systems as well,  
 326 as shown in several studies (Russell et al., 2010;Triesch et al., 2021a;Triesch et al., 2021b;van  
 327 Pinxteren et al., 2017;Zeppenfeld et al., 2021). To this end, the concentration of the analyte  
 328 of interest in each compartment is related to the respective sodium concentration (equation  
 329 2), because sodium is regarded as a conservative sea salt tracer transferred to the atmosphere  
 330 in the process of bubble bursting (Sander et al., 2003).

$$331$$

$$332 \quad EF_{aer.} = \frac{c (analyte)_{aer.} / c (Na^+)_{aer.}}{c (analyte)_{seawater} / c (Na^+)_{seawater}} \quad (2)$$

$$333$$

334 To account for the different timescales for ocean and atmospheric processes and to  
 335 improve robustness, we applied the average concentration values of the respective compound  
 336 groups in seawater and on aerosol particles as done by Zeppenfeld et al., (2021) and van  
 337 Pinxteren et al., (2022). It should be noted that  $EF_{SML}$  and  $EF_{aer.}$  comparisons include samples  
 338 from the same campaign but not necessarily from the same exact date (details in Tab.1 and  
 339 Tab S2 - S5). However, as our focus was not on showing trends within the campaign (this was  
 340 covered by Triesch et al., 2021a,b) but rather on comparing the three different OC groups with  
 341 each other, the usage of average values is justified. To investigate the variability of the  $EF_{aer.}$ ,  
 342 the minimum ratio of  $c (analyte)_{aer.} / c (Na^+)_{aer.}$  vs. the maximum ratio of  $c (analyte)_{seawater} / c (Na^+)_{seawater}$   
 343 was applied and vice versa. As seawater concentration, the bulk water or the SML  
 344 concentration can be applied.

345 Statistical significance was evaluated using the analysis of variance (ANOVA).

346

### 347 **3. Results and Discussion**

#### 348 **3.1 SML and bulk water**

##### 349 **3.1.1 Concentration and composition of the dissolved amino acids (DAA), dissolved** 350 **carbohydrates (DCHO) and dissolved lipids (DL)**

351 Figure 1 shows the analyte concentrations in the bulk ocean water (DAA:  $80 \pm 53 \mu\text{g L}^{-1}$   
 352  $^1$ , DCHO:  $78 \pm 15 \mu\text{g L}^{-1}$ , DL:  $70 \pm 25 \mu\text{g L}^{-1}$ ) and in the SML (DAA:  $190 \pm 238 \mu\text{g L}^{-1}$ , DCHO:  $85 \pm$   
 353  $30 \mu\text{g L}^{-1}$ , DL:  $83 \pm 24 \mu\text{g L}^{-1}$ ). Hence, the average concentrations for DCHO and DL are similar  
 354 in the bulk water and in the SML (detailed values in Sect. 2.2.2, Tab. 1 and DL concentrations  
 355 can be found in Triesch et al., (2021b)). For the DAA, however, SML concentrations show a  
 356 larger variability compared to the other compounds and to bulk water. However, despite the  
 357 variability of the DAA in the SML, they are not statistically different to the other two  
 358 compound groups (DL and DCHO) as well as to the DAA in the bulk water (Anova one way,  $p >$   
 359  $0.5$  at the  $0.5$  level) that is different in the case of free amino acids (FAA). In total, no significant  
 360 difference is present between the compound groups within the SML and the ULW (Anova one  
 361 way,  $p > 0.5$  at the  $0.5$  level). Resulting from the higher SML concentrations, the average SML  
 362 enrichment factors of DAA is  $2.3 \pm 0.3$  (Tab.1) and therefore higher compared to the DCHO  
 363 ( $EF_{SML} = 1.1 \pm 0.5$ ) and DL ( $EF_{SML} = 1.3 \pm 0.2$ ). The high variability of the DAA concentrations

364 agreed well with the FAA that comprise the sum of unbound individual amino acids i.e. not  
 365 bound in a peptide or protein and were measured at this location during the MarParCloud  
 366 campaign (Triesch et al., 2021a). However, the FAA in the SML were significantly higher  
 367 compared to the FAA in the bulk water (Triesch et al., 2021a). Other studies have pointed out  
 368 highly variable amino acid concentrations as well, for example, Zänker et al., (2017) showed  
 369 FAA concentrations between 32 and 1268 nmol L<sup>-1</sup> and DAA varied between 202 and 2007  
 370 nmol L<sup>-1</sup> (for comparison: the here presented DAA values correspond on average to 1064 nmol  
 371 L<sup>-1</sup> in the bulk water and 2536 nmol L<sup>-1</sup> in the SML). High enrichments of FAA in the SML were  
 372 reported (Kuznetsova and Lee, 2002;Kuznetsova et al., 2004;Reinthaler et al., 2008;van  
 373 Pinxteren et al., 2012;Engel and Galgani, 2016) with FAA enrichments up to 300 in the SML of  
 374 the Cape Verde seawaters (Triesch et al., 2021a). A preferential enrichment of FAA over  
 375 dissolved combined amino acids as a consistent microlayer feature was proposed (Kuznetsova  
 376 et al., 2004).  
 377

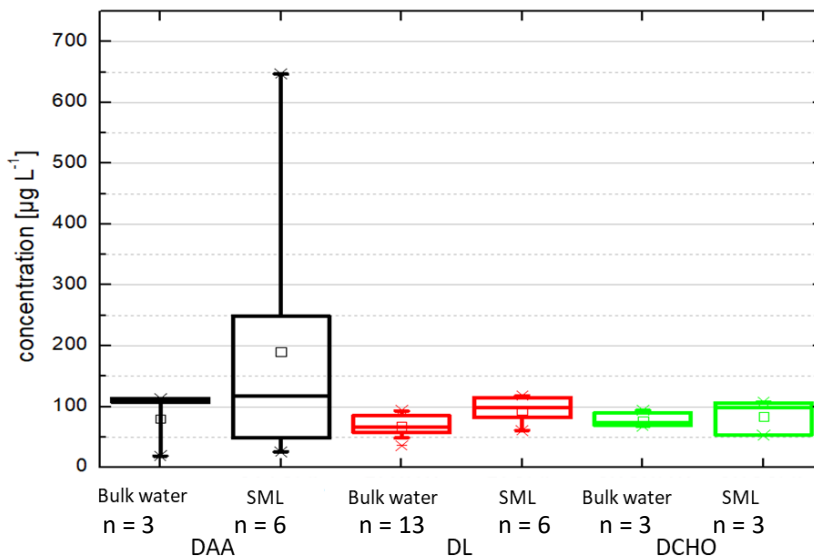


Figure 1

378  
 379 Regarding the composition of the individual DAA measured here, clear differences  
 380 between the SML and the bulk water characteristics were observed (Fig. 2, blue and orange  
 381 bars, data in Tab. S2 - 5). Besides the higher concentrations in the SML, some DAA were only  
 382 present in the SML and not in the bulk water (below detection limit). This was most  
 383 pronounced for Glu, but also evident for Tyr and Iso (detailed values in Tab. S3).

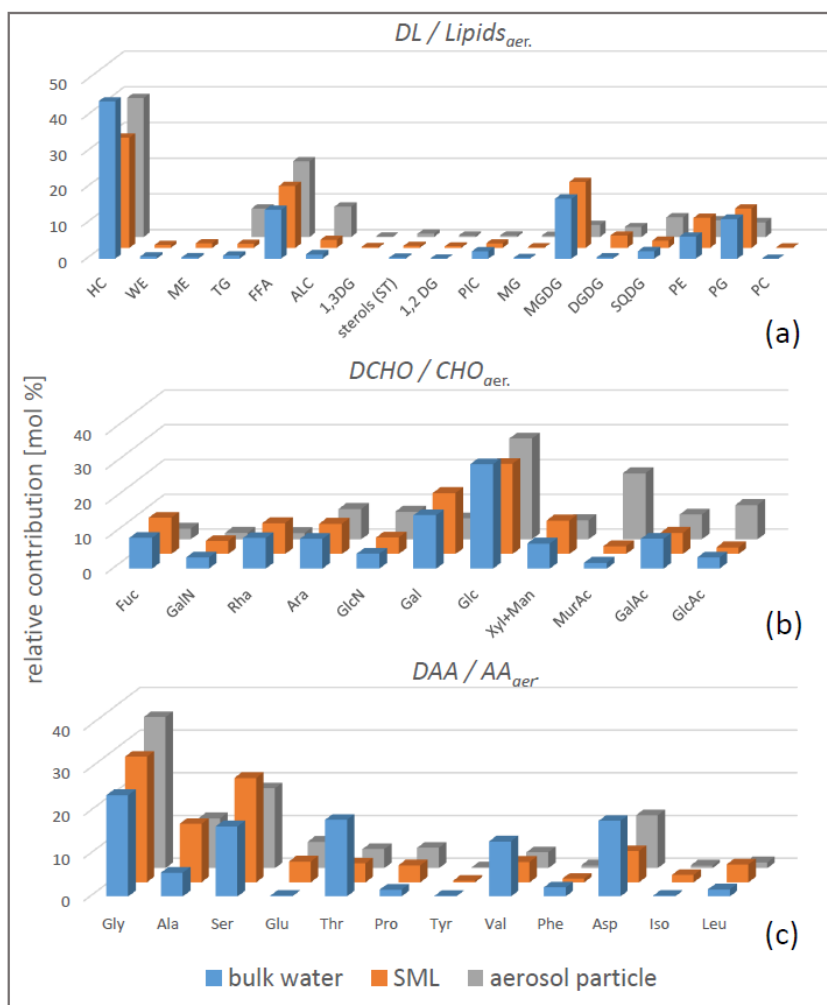


Figure 2

384

385 In contrast to the DAA, the DCHO enrichment in the SML was less pronounced with an  
 386  $EF_{SML}$  of  $1.1 \pm 0.5$  (Tab. 1), similar to SML enrichment values obtained for DCHO close to the  
 387 Peruvian upwelling regime (Zänker et al., 2017) and the Antarctic Peninsula (Zeppenfeld et al.,  
 388 2021). Regarding the relative composition, the DCHO showed a very homogeneous pattern  
 389 and were similar in the SML and the bulk water (Fig. 2, blue and orange bars).

390 The enrichment of the DL ( $EF_{SML} = 1.3 \pm 0.2$ , Tab. 1) was very similar to the DCHO  
 391 enrichment and is discussed in Triesch et al., (2021b). From the individual lipid components  
 392 and the lipolysis index, it was concluded that the lipids were degraded only to a small extent  
 393 (Triesch et al., 2021b).

394 Altogether, the high and varying concentrations and enrichments of DAA in the SML in  
 395 contrast to the DCHO and DL concentrations underline that significant changes occur for the  
 396 DAA in the SML that are less pronounced for the other two compound groups. However, it is  
 397 important to mention that the number of samples that have been analysed is different and  
 398 limited (Tab. 1 and Fig. 1) and samples from partly different dates of the campaign have been  
 399 analysed and compared.

400

### 401 3.1.2 Discussion of the SML enrichment

402

### 403 3.1.2.1 Surface vs bulk SML

404

405 The SML enrichment of DOC components is generally attributed to diffusion, turbulent  
406 mixing, as well as scavenging, and transport of surface-active matter from rising gas bubbles  
407 in the water column (Liss and Duce, 1997). Within the groups investigated here, the DL are the  
408 most hydrophobic compounds and are generally classified as highly surface-active compounds  
409 (Burrows et al., 2014). Although the surface-activity parameters (e.g. octanol-water partition  
410 coefficient, density, Topological Polar Surface Area) of the individual lipids differ among each  
411 other (values in Triesch et al., 2021b), the lipids are overall more non-polar and surface active  
412 compared to the carbohydrates and amino acids (values in Triesch et al., 2021a). Nevertheless,  
413 the enrichment of the DL in the SML was significantly lower compared to the carbohydrates  
414 and amino acids. This was obvious from a comparison of the averaged values as well as from  
415 single samples obtained at the same date (03.10.2017 and 07.10.2017, values in Tab. S2, S3,  
416 and Triesch et al., 2021b).

417 One explanation for the finding lies in the sampled SML thickness. With the glass plate  
418 technique, an SML thickness of about 100  $\mu\text{m}$  has been sampled and reported for the CVAO  
419 area (van Pinxteren et al., 2017). Hence, the 100  $\mu\text{m}$ -thick SML might be very well-mixed with  
420 regards to the soluble amino acids and carbohydrates, however the surface-active  
421 compounds, such as lipids, are potentially located on the very top and form a thin (nm-thick)  
422 monolayer. In the literature, the SML is described either as a series of sub-layers of wet and  
423 dry surfactants (Hardy, 1982) or as a gelatinous matrix (Sieburth, 1983). Independent of the  
424 model, it can be expected that a gradient along the surface likely forms with surfactants at the  
425 very top of the layer. The formation of a lipid-rich nanolayer on the very top agrees with  
426 surface-sensitive spectroscopy measurements that are able to tackle the uppermost layer and  
427 found strong indications for a nanolayer dominated by soluble surfactants (Lass and  
428 Friedrichs, 2011) and hydrophobic low molecular weight lipids (Frka et al., 2012). The  
429 nanolayer, however, cannot be probed with currently applied SML sampling methods.  
430 Therefore, the measured SML concentrations may represent a very diluted (likely highly lipid-  
431 enriched) layer. Consequently, the SML structure is even more complex, which needs to be  
432 considered, particularly when discussing lipid enrichments in the SML. Here, a combination of  
433 bulk measurements with dedicated surface probing appears highly desirable.

434

### 435 3.1.2.2 Details of SML enrichment mechanisms

436

#### 437 (i) Co-adsorption

438

439 Regarding the DAA in detail, it is interesting to note that some compounds are  
440 exclusively present in the SML, as mentioned above. They belong to hydrophilic (Glu),  
441 hydrophobic (Iso) and neutral (Tyr) fractions of amino acids, underlining that their occurrence  
442 in the SML might not be related (solely) to their physicochemical properties. Besides an air  
443 bubble-driven transfer to the surface, enrichment in the SML can be supported by co-  
444 adsorption mechanisms. Less surface-active compounds (e.g. amino acids and carbohydrates)

445 can be attached due to ionic interactions/coulomb interactions to the head groups of the air  
446 bubble-attached surfactants (e.g. lipids) that mediate their enrichment in the SML (Burrows  
447 et al., 2016;Hasenecz et al., 2019;Link et al., 2019;Schill et al., 2018). Co-adsorption can  
448 provide an explanation for the high occurrence of non-surface active, very soluble  
449 compounds, such as carbohydrates. A recent laboratory study showed different mechanisms  
450 for the co-adsorption of polysaccharides that form a second calcium-bridges sublayer  
451 underneath the monolayer whereas monosaccharides intercalate and induce reorganisation  
452 within the nanolayer (Vazques de Vasquez et al., 2022). However, in the current study, only a  
453 small SML enrichment of the DCHO and, hence, no indication for a strong co-adsorption was  
454 observed.

455

456 (ii) In-situ processing: Abiotic vs. biotic

457

458 Further explanations for the accumulation of dissolved compounds require in situ  
459 formation or degradation by SML-specific reactions that might be triggered by distinct  
460 environmental conditions in the SML. Biotic pathways and abiotic SML-specific  
461 (photo)chemical reactions may strongly impact OC cycling at the sea surface (Liss and Duce,  
462 1997). The high abundance of the amino acid Glu in the SML observed here was also reported  
463 in the FAA fraction by Triesch et al., (2021a) and can likely be explained by in situ formation.  
464 In incubated cloud water, as another marine compartment, Glu has been shown to be  
465 produced via biotic and abiotic mechanisms, e.g. via the oxidation of proline (Jaber et al., 2021  
466 and refs. therein) and such processes may be relevant in the ocean surface as well. Regarding  
467 biotic processes, it is well-known that microorganisms have complex and highly  
468 interconnected enzymatic networks and can biodegrade or biosynthesize organic compounds  
469 (KEEG pathway). Kuznetsova and Lee (2002) suggested that stressed microorganisms, rich in  
470 dissolved and combined amino acids, may be leached and released them, which in turn affects  
471 the pools of both these compounds in seawater. Although such formation mechanisms  
472 generally happen in the upper ocean, there are indications for SML-specific processes. Along  
473 a transect from upwelling regions toward oligotrophic gyres it was found that while in the bulk  
474 water a clear trend toward degradation of amino acids was observed, the production and  
475 degradation patterns of amino acids in the SML were much more complex (Reinthalder et al.,  
476 2008). This is indicative of the role of the SML in the production of labile DOC driven by coupled  
477 microbial and photochemical processes. Similarly, Kuznetsova and Lee (2001) observed that  
478 peptide turnover was always faster in the SML than in subsurface waters likely due to the  
479 greater concentrations of DOC in the SML. The authors concluded that the accumulation of  
480 organic and inorganic compounds in the SML leads to a more nutritious medium for microbial  
481 growth and consequently enzymatic hydrolytic activity compared to the bulk water.  
482 Connecting this to the results presented here, this might suggest that changes induced by  
483 abiotic and biotic processing need to be considered when regarding the SML composition.  
484 Although such reactions likely also affect lipids and carbohydrates, they seem to be most  
485 pronounced for amino acids.

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(iii) Microbial nitrogen fixation at the sea-air interface

A further mechanism contributing to the high and variable SML enrichment of the DAA at the current location might be microbial nitrogen fixation at the sea-air interface. Measurements showed that cyanobacteria are a very pronounced phytoplankton group in this region (Franklin et al., 2009; Hepach et al., 2014; Zindler et al., 2012), which was dominant during the MarParCloud campaign (van Pinxteren et al., 2020). Cyanobacteria are able to take up nitrogen from the atmosphere (Zehr, 2011). Earlier studies showed that cyanobacteria-fixed nitrogen is incorporated into amino acids (specifically glutamine (Carpenter et al., 1992)). The calculated net amino acids release from cyanobacteria colonies (*Trichodesmium thiebautii*) revealed that nitrogen fixation and the biogeochemical turnover of ambient amino acids are an important source of recently fixed (“new”) nitrogen within the oceanic surface water (Capone et al., 1994). These considerations are, however, highly speculative and demand further studies to investigate if nitrogen fixation and biosynthesis via cyanobacteria, which is often observed in subtropical and tropical oceans (Montoya et al., 2007), might establish a considerable route for amino acid formation and enrichment in the SML from the atmospheric side.

(iv) Concluding remarks towards the SML enrichment

Although SML enrichment factors for amino acids, carbohydrates, and lipids have been reported in the available literature, they have not previously been shown in such detailed analysis for samples collected from the same site as shown here. From this study it can be concluded that the amino acids are strongly enriched in the SML compared to carbohydrates and lipids, even under the same environmental conditions. In a recent study we showed a strong enhancement of other nitrogen-containing species (aliphatic amines) in the SML at this location, while the amine concentration in the bulk water was often not detectable (van Pinxteren et al., 2019). This suggests that the pronounced SML enrichment specifically exists for nitrogen-containing organic species. In addition, the absence of a relation of the SML enrichment to physical compound parameters (e.g. hydrophobicity) suggests that enrichment processes based on physicochemical properties (e.g. surface-activity) alone do not drive SML enrichment. Rather, an SML in situ formation mechanism impacts the abundance of amino acids and likely nitrogen-containing organic species in general.

## 3.2. Aerosol particles

### 3.2.1 Concentration and composition

After evaluating the concentrations of the analytes in seawater and the SML, in the next step, their presence in the aerosol particles was investigated. The concentrations of  $AA_{aer}$ .

527 and  $\text{CHO}_{aer.}$  were  $2.4 \pm 1.1 \text{ ng m}^{-3}$  and  $1.0 \pm 1.1 \text{ ng m}^{-3}$ , respectively (Fig. 3, Tab. 1) and not  
528 significantly different ULW (Anova one way,  $p > 0.5$  at the 0.5 level).

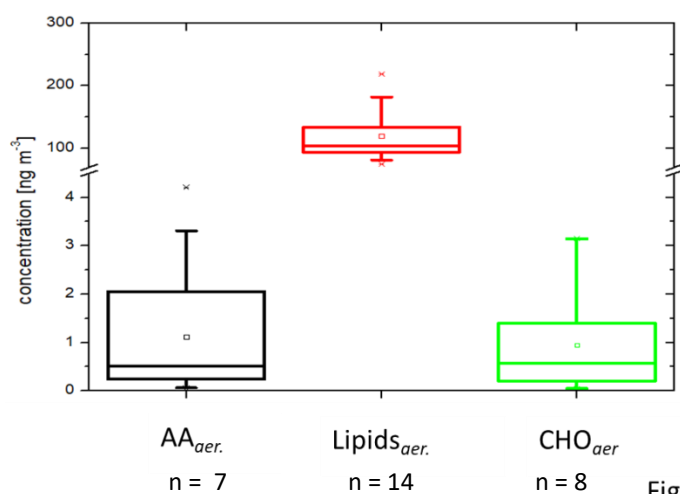


Figure 3

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530 Compared to results from the Polar Regions, the  $\text{CHO}_{aer.}$  concentrations in the tropical  
531 Atlantic Ocean analysed here are at the lower end. Leck et al., (2013) determined  
532 carbohydrates during the Arctic summer and found  $0.7 - 20 \text{ ng m}^{-3}$  in submicron particles.  
533 Zeppenfeld et al., (2021) found carbohydrate concentrations between  $0.2 - 11.3 \text{ ng m}^{-3}$  in  $\text{PM}_{10}$   
534 atmospheric particles, in the Western Antarctic Peninsula, which contributed about 3% to the  
535 OC. The same holds true for amino acids, the concentrations found here are slightly lower as  
536 reported for other marine regions, e.g. FAA in Antarctic aerosol particles were on average  $4.6$   
537  $\text{ng m}^{-3}$  in Antarctic aerosol (Barbaro et al., 2015). Triesch et al., (2021a) found FAA  
538 concentrations between  $1.5$  and  $3.0 \text{ ng m}^{-3}$  in the aerosol particles from the Cape Verdes. The  
539  $\text{AA}_{aer.}$  are generally in the same order of magnitude compared to other marine locations  
540 (Matos et al., 2016; Matsumoto and Uematsu, 2005; Renard et al., 2022; Wedyan and Preston,  
541 2008). However, concentrations of amino acid and carbohydrates in urban, rural, and forest  
542 areas are significantly higher (2-3 orders of magnitudes) compared to the values and the  
543 spectrum of compounds is strongly shifted (e.g. Dominutti et al., 2022; Samake et al., 2019; Zhu  
544 et al., 2022).

545 Comparing the data obtained here with those from analytical techniques that use  
546 functional group information (FT-IR) has shown that the latter techniques often attribute a  
547 large fraction of alcohol (hydroxyl) functional groups on the marine aerosol particles (Cravigan  
548 et al., 2020; Frossard et al., 2014; Russell et al., 2010). According to Russell et al., (2010), the  
549 primary marine signal in submicron marine aerosol over the North Atlantic and Arctic Oceans  
550 is made on average for 88% of hydroxyl groups corresponding to carbohydrate-like material.  
551 Such high fractions of carbohydrates were not found in the chromatographic analysis of  
552  $\text{CHO}_{aer.}$  presented here, nor in other studies using similar methodologies (e.g. Zeppenfeld et  
553 al., 2021). In a recent study, using thermal desorption mass spectrometry, it was suggested  
554 that carbohydrates only represented a minor fraction of the FT-IR alcohol group and another  
555 thermally stable fraction, different to carbohydrates, was the main contributor to the alcohol  
556 group (Lawler et al., (2020)). Hence, previous FT-IR measurements might have over-predicted

557 the carbohydrate fraction of marine aerosol particles and further (molecular-based) analysis  
558 should be conducted in comparison to resolve existing contradictions.

559 In contrast to the  $AA_{aer.}$  and  $CHO_{aer.}$ , the  $Lipid_{aer.}$  concentrations were  $120 \pm 43 \text{ ng m}^{-3}$  and  
560 therefore two orders of magnitude higher than the other two organic groups (Fig. 3, Tab. 1)  
561 The  $Lipid_{aer.}$  were significantly higher than the  $AA_{aer.}$  ( $p = 6.6E-13$ ) and the  $CHO_{aer.}$  ( $p = 1.6E-7$ )  
562 (Anova one way, at the 0.5 level). One possible reason for the higher  $Lipid_{aer.}$  concentration  
563 could lay in the difference in the extraction procedure (chapter 2.2) as  $AA_{aer.}$  and  $CHO_{aer.}$  were  
564 extracted in water, while the  $Lipid_{aer.}$  were extracted in an organic solvent. Previous lipid  
565 analysis on a molecular level revealed concentrations between  $0.19 - 23 \text{ ng m}^{-3}$  for ALC and  
566 between  $2.5 - 38 \text{ ng m}^{-3}$  for free fatty acids on marine aerosol particles from the western North  
567 Pacific (Kawamura et al., 2003) and a recent study found marine fatty acid concentrations  
568 between  $50$  and  $90 \text{ ng m}^{-3}$  in coastal aerosol at Qingdao (Chen et al., 2021). Mochida et al.,  
569 (2002) observed saturated fatty acids (C14 - C19) on marine aerosol particles over the  
570 northern Pacific in atmospheric concentration between  $0.8 - 24 \text{ ng m}^{-3}$ . Hence, these data are  
571 in the same order of magnitude as here measured lipid groups (ALC:  $6.3 \text{ ng m}^{-3}$ , free fatty  
572 acids:  $18.5 \text{ ng m}^{-3}$ , values in Triesch et al., 2021b). Cochran et al., (2017) showed that lipid  
573 components (long-chain fatty acids) comprised a significant fraction of up to 75% of the  
574 identified organic constituents in aerosol particles from a sea spray tank.

575 A recent study using FT-ICR-MS found that among a large number of molecular  
576 formulas, large peak numbers were assigned to proteins and lignins (phenolic  
577 macromolecules) followed by carbohydrates and lipids in aerosol originating from the Arctic  
578 ocean (vs. non-ocean aerosol) (Choi et al., 2019).

579 High lipid fractions in marine aerosol particles were also reported from NMR  
580 measurements. Measurements of nascent aerosol particles produced from North Atlantic  
581 seawater showed that the water-soluble organic aerosol fraction was purely aliphatic with  
582 hydroxylated moieties of sugars, esters, and polyols, aliphatic groups adjacent to carbonyls,  
583 amides, and acids, as well as aliphatic chains with terminal methyl-groups, typical of lipids  
584 (Facchini et al., 2008). The water-insoluble organic fraction was dominated by  
585 lipopolysaccharides, known phytoplankton exudate components. A recent study applying  
586 NMR analysis to artificially produced aerosol particles after bubbling seawater from offshore  
587 areas also showed proof of polyols and lipids (Decesari et al., 2020). NMR measurements of  
588 lipids are mainly qualitative, however, the high fraction of lipid-like components from other  
589 regions agrees well with the high  $Lipid_{aer.}$  concentrations presented here.

590 A high  $Lipid_{aer.}$  concentration as observed in the present study agrees well with the modelling  
591 results of Burrows et al., (2014), where the ocean-atmosphere transfer was calculated  
592 according to the physicochemical properties of the distinct OC groups. Lipids, as the most  
593 surface active OC group comprise the largest fraction of the aerosol fraction, although their  
594 (modelled) concentration in the seawater is lower compared to carbohydrates and amino  
595 acids (Burrows et al., 2014). In a latter model modification, where additional co-adsorption  
596 processes were included in the calculations, a more pronounced saccharidic fraction was  
597 determined on the aerosol particles from the model results (Burrows et al., 2016), that is  
598 different from the findings here, at least regarding the  $CHO_{aer.}$  measured on a molecular level.



599 However, it needs to be considered that the lipids analysed here include glycolipids (MGDG,  
600 DGDG, SQDG) which are components that have the solubility properties of a lipid but also  
601 contain one or more sugar molecules. The glycolipids comprise a non-negligible portion of the  
602 OC on the aerosol particles (values in Table S6 and S8). This underlines the complexity of  
603 attributing the OC to distinct organic groups and demonstrates that the applied analytical  
604 methods must be taken into account when comparing concentrations of substance groups.  
605 This is discussed in more detail in 3.4.1. Altogether, there seems to be a discrepancy between  
606 the measured concentrations and the modelled results underlining that the transfer of the  
607 organic compounds from the ocean to the atmosphere based on their physicochemical  
608 properties might not be the only mechanism.

609

### 610 3.2.2 Aerosol enrichment

611

#### 612 3.2.2.1 Aerosol enrichment factors

613 The finding that the Lipids<sub>aer.</sub> were much higher concentrated than the AA<sub>aer.</sub> and the  
614 CHO<sub>aer.</sub>, resulted in a very different pattern compared to the similar seawater concentrations  
615 (Fig. 1 vs. Fig. 3). To quantitatively compare the seawater and the aerosol concentration, the  
616 EF<sub>aer.</sub> was calculated (values in Tab.1).

617

618

#### ***Insert Table 1***

619

620 For the amino acids, the EF<sub>aer.</sub> was between  $9.2 \times 10^2$  (related to the SML) and  $2.1 \times 10^3$   
621 (related to the bulk water) and on average  $1.5 \times 10^3$ . For the carbohydrates, the EF<sub>aer.</sub> was  
622 between  $1.3 \times 10^3$  (related to the SML) and  $1.4 \times 10^3$  (related to the bulk water) and on average  
623  $1.3 \times 10^3$  and therefore similar to the EF<sub>aer.</sub> of the amino acids. For the lipids, however, the EF<sub>aer.</sub>  
624 was two orders of magnitude higher (EF<sub>aer.</sub> =  $1.4 \times 10^5$ , related to the SML; EF<sub>aer.</sub> =  $1.7 \times 10^5$   
625 related to the bulk water, EF<sub>aer.</sub> =  $1.6 \times 10^5$  on average, Tab. 1).

626

#### 627 3.2.2.2 Oceanic transfer and atmospheric in situ formation

628 The overall high enrichment of OC in the aerosol particles is explained by complex, not  
629 yet - resolved interactions at the ocean surface where organic matter is enriched relative to  
630 sodium during the formation of film and jet droplets. Burrows et al., (2014) applied a  
631 conceptual model (“slab” model) where all organics partition to the surface of a “slab” of  
632 oceanic water or to both the outer and inner surfaces of a bubble film. The organic enrichment  
633 is therefore significantly higher for the thinner bubble films (bubble film thicknesses: 0.01 to  
634  $1 \mu\text{m}$ ) than for the thicker SML (typically sampled SML thicknesses: 20 to  $400 \mu\text{m}$ ). This  
635 mechanism can explain an EF<sub>aer.</sub> of OC in submicron aerosol particles of  $10^2$  to  $10^3$  compared  
636 to the SML (Burrows et al., 2014). However, EF<sub>aer.</sub> from ambient and laboratory-controlled  
637 observations show that for some compounds even higher EF<sub>aer.</sub> are obtained. In a controlled  
638 tank study, Rastelli et al., (2017) found strong enrichments for lipids (up to  $1.4 \times 10^5$ ), as well as

639 for proteins (up to  $1.2 \times 10^5$ ) and carbohydrates (up to  $1.0 \times 10^5$ , Tab. 1). A recent controlled  
640 bubble-bursting laboratory study showed that amino acids enrichments can be up to  $10^7$  in  
641 submicron SSA between 0.029 and 0.060  $\mu\text{m}$  (Triesch et al., 2021c). Similarly, Schmitt-Kopplin  
642 et al., (2012) showed that surface-active biomolecules are preferentially transferred from  
643 surface water into the atmosphere via bubble bursting. The ambient enrichment factor of the  
644 lipids ( $10^5$ ) shown here and in Triesch et al., (2021b) agreed well with laboratory-derived ones  
645 (Rastelli et al., 2017) indicating that the transfer mechanisms simulated in lab experiments  
646 agree with here performed observations in the field. Hence, the high surface activity and/or  
647 the lipophilic character of the lipid classes might explain their strong (chemo-selective)  
648 transfer to the aerosol particles. Even though the lipid composition on the aerosol particles  
649 slightly varied from the seawater concentration (Triesch et al., 2021b), their transfer is likely  
650 driven by their physicochemical properties (high surface activity and/or the lipophilic  
651 character). For the amino acids and carbohydrates, however, more complex mechanisms may  
652 determine their transfer to the atmosphere. Rastelli et al., (2017) suggested that diverse  
653 biological processes on the ocean drive the properties of proteins and carbohydrates in the  
654 ocean surface and in the atmosphere. Moreover, these compounds are known to be involved  
655 in marine gel-like particle formation, such as transparent polymer particles (TEP) and  
656 coomassie stained particles (CSP), observed in the ocean and more recently in the atmosphere  
657 (Aller et al., 2017; Kuznetsova et al., 2005; van Pinxteren et al., 2022) adding more complexity  
658 to the system. Hence not only a sea-to-air transfer but also atmospheric in situ formation and  
659 degradation might determine the concentration of the OC and notably of the amino acid and  
660 carbohydrates. This suggests that atmospheric processing plays an important role besides the  
661 physical-driven bubble bursting sea-air transfer of OC.

662

### 663 3.2.3 Limitations of the concept of an aerosol enrichment factor

664

665 When comparing OC in the ocean and the atmosphere, it needs to be considered that  
666 processes in the ocean and the atmosphere happen on different timescales. In addition, the  
667 seawater samples comprise spot samplings in the ocean while the sampling period of  $\text{PM}_{10}$   
668 aerosol particles at the CVAO covers a time span of 24h. These issues make a comparison  
669 between the ocean and atmospheric data very challenging. However, the air masses arriving  
670 at the CVAO often followed the water current (Pena-Izquierdo et al., 2012; van Pinxteren et  
671 al., 2017) and suggest a strong link between the upper ocean and the aerosol particles, as  
672 mainly winds drive the ocean currents in the upper 100 m of the ocean. Besides the ocean,  
673 Saharan dust is a strong aerosol source at the Cape Verde islands, most pronounced in the  
674 months December to February (Fomba et al., 2014). The backward trajectories during the time  
675 of the campaign (Fig. S2) as well as the mass concentrations of inorganic ions and mineral dust  
676 tracers on the aerosol particles measured during the campaign suggested a predominant  
677 marine origin with low to medium dust influences (van Pinxteren et al., 2020). Moreover, dust  
678 generally influences the supermicron particles to a larger extent than the submicron particles  
679 analysed here (Fomba et al., 2013). Hence, although different factors certainly affect the  
680 aerosol composition, it is reasonable to assume a strong oceanic contribution.

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### 3.3. Seawater and aerosol particles: Comparison of the relative composition

Regarding the organic components on the aerosol particles, the same compounds that were present in the seawater were generally present on the aerosol particles (Fig. 2, grey bars, and values in Tab. S2 - 5). However, the relative composition of distinct compounds was, at least partly, different. Regarding the carbohydrate composition, the percentages of MurAc, GlcAc, and GlcN in the aerosol particles were higher compared to the seawater. MurAc and GlcN are important constituents in the cell walls of marine microorganisms and notably, MurAc serves as a proxy for bacterial biomass (Mimura and Romano, 1985). Its high concentration might indicate an enrichment of bacteria on the aerosol particles. Zeppenfeld et al., (2021) detected similar (biogenic) carbohydrates in particles sampled in the western Antarctic peninsula and suggested that marine bacteria in atmospheric particles may metabolize a part of the oceanic carbohydrates in a selective enzymatic way analogous to the bacterial processes in seawater. Such processes might explain the changed carbohydrate composition and are likely not restricted to a specific oceanic regime, as they seem to happen in the Southern Ocean (Zeppenfeld et al., 2021) as well as in the tropical Atlantic Ocean, observed here. The elevated relative occurrence of GlcAc found here agrees well with the recent finding of a high abundance of gel-like material in aerosol particles at the CVAO, strongly enriched towards sodium compared to seawater (van Pinxteren et al., 2022) as GlcAc is one main component of marine gelatinous exopolysaccharides (Casillo et al., 2018; Krembs et al., 2002). Regarding the lipids, surfactants such as free fatty acids as well as lipophilic compounds, such as hydrocarbons, had major contributions in the seawater and on the aerosol particles, respectively. However, TG, an energy storage lipid, had a higher contribution to the aerosol particles versus the ocean water. In addition, some other, minor-contributing lipid classes were partly different within ocean and atmosphere (Triesch et al., 2021b).

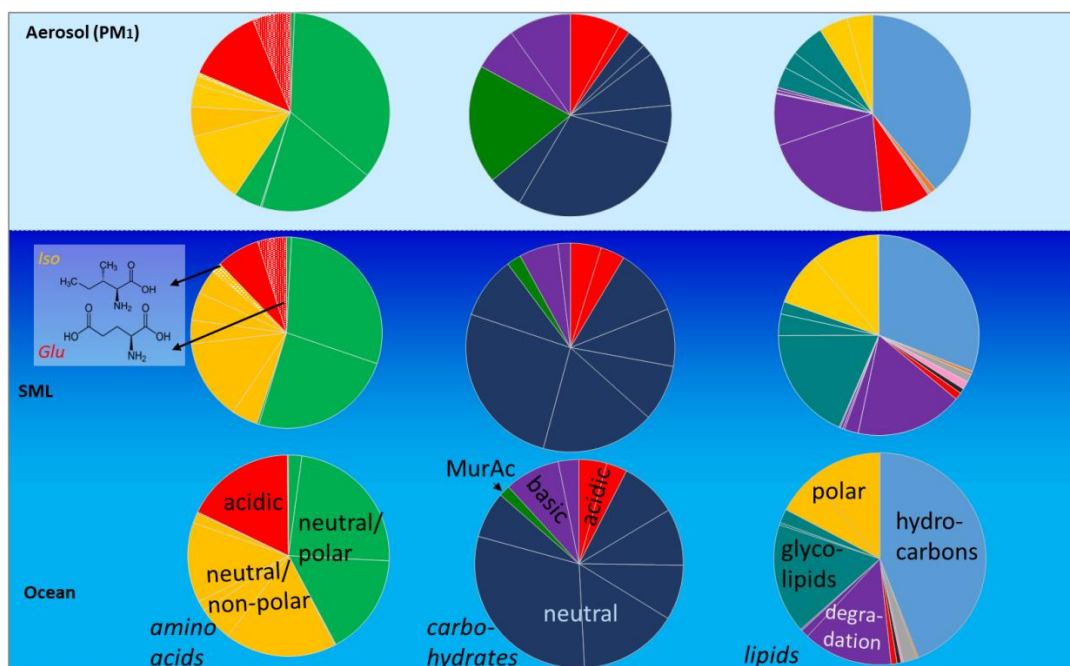


Figure 4

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The most remarkable difference in relative composition in seawater and in aerosol particles was found for amino acids, as some DAA were clearly present in the SML and in the aerosol particles but not in the bulk water (e.g. Iso and Glu, Fig. 4, individual values in Tab. S6). The amino acids generally differed a lot regarding their SML and bulk water composition. This was visible in the data set presented here for the DAA, and also reported for the FAA measured from the same campaign (Triesch et al., 2021a). Recently it was reported that the acidic amino acid Glu (in the form of FAA) is transferred to SSA to a large extent (Triesch et al., 2021c) and the results of the present study suggest that Glu might be transferred solely from the SML (and not from the bulk water) to the aerosol particles. However, besides the oceanic transfer, Glu can result from an in situ formation on the aerosol particles. Similarly to the seawater, Glu might form from biotic or abiotic reactions on the aerosol particles. From the measurements performed here, it is not possible to differentiate between a selective transfer of Glu from the SML and its biotic and abiotic in situ formation in aerosol particles. Recently, Jaber et al., (2021) and Renard et al., (2022) evaluated the atmospheric aging of the amino acids and considered biotic and abiotic (mainly oxidation) processing. Their calculations revealed different atmospheric lifetimes for the individual amino acids related to oxidation and biological processes, respectively. For example, the amino acids Ser and Ala are degraded quickly by biological processes (lifetime of a few hours) but are more stable towards oxidation (Renard et al., 2022). Such studies can help to understand the patterns of the amino acids as observed here and relate them to sources and atmospheric processing. The presence of Ser and Ala in the aerosol particles investigated here could therefore indicate that biodegradation of these compounds was not pronounced. However, additional studies are needed to better understand atmospheric biotic and abiotic processing. In addition, the transfer of individual

734 DAA exclusively from the SML shall be investigated in further research, preferably within  
735 characterized and controlled bubbling systems.

736

### 737 3.4 Contribution to aerosol particle OC

738

#### 739 3.4.1 Molecular and semi-molecular analysis

740 OC concentrations in marine aerosol particles during this campaign varied between  
741 0.13 and 0.31  $\mu\text{g m}^{-3}$  with an average value of 0.20  $\mu\text{g m}^{-3}$  (values in Tab. S7). This agreed well  
742 with previous OC measurements from the CVAO that were on average 0.27  $\mu\text{g m}^{-3}$  OC (van  
743 Pinxteren et al., 2017). To date, only a small percentage of OC on marine aerosol particles is  
744 characterised on a molecular level and organic biomarkers often comprise only a few percent  
745 of the OC (Chen et al., 2021). Fu et al., (2011) measured more than 140 different single organic  
746 species in marine aerosol from different oceanic areas, however the identified species  
747 composed less than 5.7% of the OC. Taking together the OC components described here  
748 (Lipids<sub>aer.</sub>, AA<sub>aer.</sub>, CHO<sub>aer.</sub>), the contribution of the identified components to the OC was  
749 calculated. Furthermore, the OC contribution of recently identified components from previous  
750 campaigns within the Cape Verde region, in detail: aliphatic amines, methane-sulfonic acid  
751 (MSA), oxalic acid and carbonyls (van Pinxteren et al., 2015) was included. The OC contribution  
752 of the single compounds and compound groups are shown in Fig. 5 (values in Tab. S8).  
753 Altogether, about 48% of the average OC could be explained by the identified components.  
754 Regarding the maximum (0.31  $\mu\text{g m}^{-3}$ ) and minimum (0.13  $\mu\text{g m}^{-3}$ ) OC concentrations within  
755 the campaign, the OC contributions of the respective compounds are between 31% (lower  
756 limit) and 74% (upper limit). The major identified OC fraction (related to the average OC) were  
757 the Lipids<sub>aer.</sub> with 43%. They were followed by the aliphatic amines (4%) that is in good  
758 agreement with a recent CVAO study, where they contributed on average with 5% to the  
759 (water-soluble) OC (van Pinxteren et al., 2019). MSA (0.9%) and oxalic acid (0.3%) were minor  
760 OC contributors. Similarly, the CHO<sub>aer.</sub> and the AA<sub>aer.</sub> made up a minor percentage with 0.3 and  
761 0.4% respectively. Regarding the lipids, it needs to be considered that the analysis performed  
762 here was not based on the detection of individual analytes but on an organic solvent  
763 extraction of the particle constituents and extract separation by solvents with different  
764 polarities applied in the TLC. The analytical method has been optimized for seawater analysis.  
765 Within atmospheric processing, additional organic compounds can form, which might contain  
766 a hydrophobic part and are potentially included in the lipid analysis performed here. However,  
767 the large similarity of the lipid groups within the seawater and the aerosol particles, as well as  
768 the agreeing concentrations of the single lipid groups (FFA, ALC) to measurements from other  
769 marine stations with molecular techniques (GC-MS) suggests that the same compound classes  
770 were present in the particles. Future analysis of the lipid fraction with mass spectrometric  
771 techniques will help to better resolve this issue.

772

773

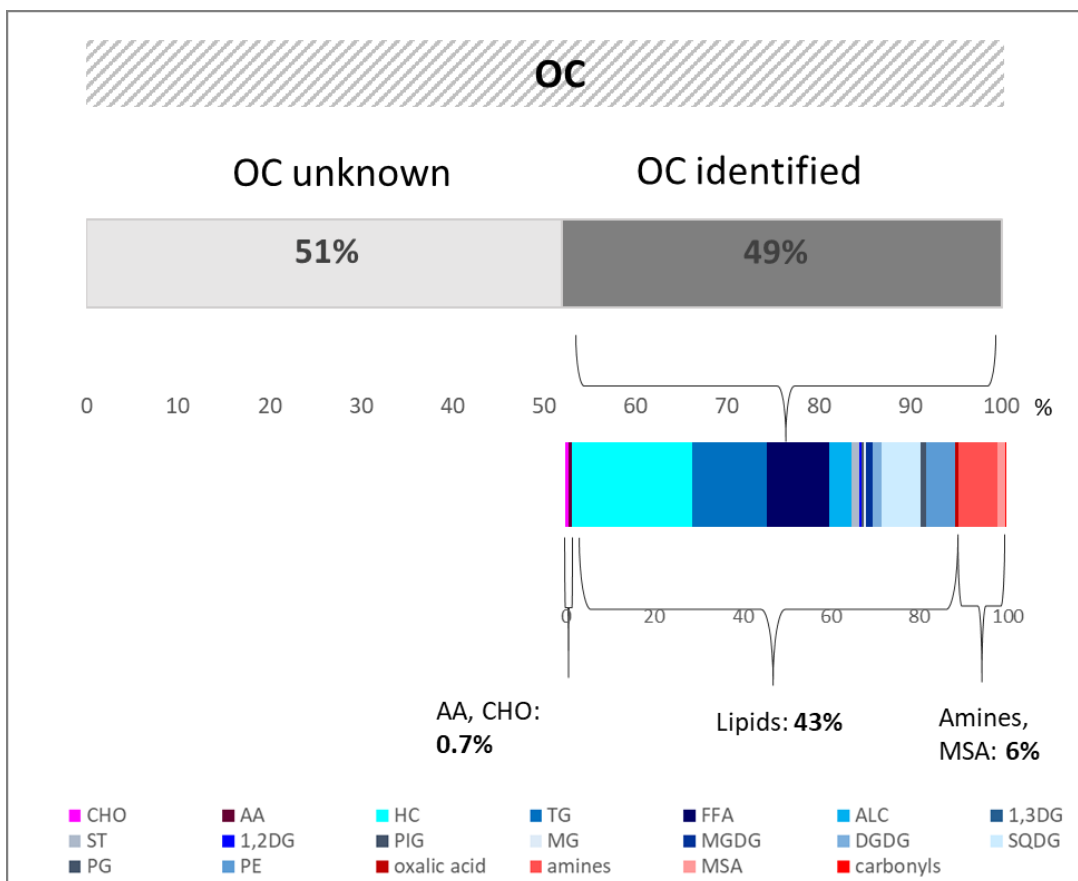


Figure 5

774

775

### 776 3.4.2 Non-identified, recalcitrant OC in aerosol particles

777

778 About 50% of the aerosol OC remained uncharacterized on a molecular and semi-  
 779 molecular level. The non-identified OC part may contain larger macromolecules that might be  
 780 composed of particulate or non-soluble forms in water (carbohydrates and proteins) or  
 781 organic solvents (lipids) that were removed in the performed analysis during the sample  
 782 preparation step. In addition, the unknown part might include component groups that belong  
 783 to the soluble carbohydrates or amino acids but are either too stable or too labile for the  
 784 sample preparation procedure (e.g. within the hydrolysis step). Moreover, other complex  
 785 molecules that cannot be captured with the methods applied here likely add to the unknown  
 786 fraction including optical active parts summarized as chromophore dissolved organic matter  
 787 (CDOM), humic-like substances (HULIS), brown carbon and water-soluble pigments. As  
 788 mentioned above, NMR analysis showed that SSA contains a large fraction of  
 789 lipopolysaccharides comprising complex, macromolecular groups of sugars, esters, carbonyls  
 790 as well as acids and lipids (Facchini et al., 2008). However, these components have, not yet  
 791 been analysed in aerosol particles using chromatographic techniques. The uncharacterized  
 792 part may also contain particulate OC compounds, such as larger aggregates of marine gels or  
 793 gel-like particles like transparent exopolymer particles (TEP). High TEP number concentrations  
 794 in aerosol particles were recently identified in the Cape Verde region (van Pinxteren et al.,

795 2022) and high mass-concentrations of TEP (e.g.  $1.2 \mu\text{g m}^{-3}$  for  $\text{PM}_{10}$ ) were identified in the  
796 western North Atlantic atmosphere (Aller et al., 2017).

797 Kieber et al., (2016) suggested that the major OC component in submicron sea-spray  
798 particles is recalcitrant and recently Beaupre et al., (2019) proposed that 19 to 40% of the OC  
799 associated with freshly produced marine aerosol particles was refractory dissolved OC. This  
800 percentage agrees with the non-identified OC part from the present study. However, we  
801 cannot identify or classify the remaining OC fraction nor attribute it with certainty to the  
802 recalcitrant OC. Further studies of the OC, in tropical as well as in other areas of the world, are  
803 needed to continue resolving OC and related transfer and formation processes. Nevertheless,  
804 the potentially recalcitrant OC fraction in the oligotrophic region does not seem to constitute  
805 the majority of OC as reported by Kieber et al., (2016).

806

### 807 3.5 Sea-to-air fluxes of the individual OC groups

808

809 The CVAO is localized in an oligotrophic region and should therefore be reasonably  
810 representative of most of the Earth's ocean surface. POA emission rates are strongly varying,  
811 however modelling studies have estimated global submicron marine POA emission rates of  
812  $10 \pm 5 \text{ Tg yr}^{-1}$  (Gantt and Meskhidze, 2013). Based on this emission flux and the contribution of  
813 the compounds to the OC fraction, we estimated the fluxes of the DAA, DL, and DCHO.  
814 Accordingly, the annual rates of emission from the ocean to the atmosphere are  $0.03 \pm 0.01$   
815  $\text{Tg yr}^{-1}$  for DCHO,  $0.04 \pm 0.02 \text{ Tg yr}^{-1}$  for DAA, and  $4.2 \pm 2.1 \text{ Tg yr}^{-1}$  for DL. The unknown OC  
816 that includes the potentially recalcitrant components, has a sea-to-air flux of  $4.8 \pm 2.4 \text{ Tg yr}^{-1}$ .  
817 However, this approach only includes the bubble-bursting-mediated transfer of the respective  
818 compounds, neglects any potential seasonal changes and neglects additional sources and  
819 formation processes. Ervens and Amato (2020) investigated the global impact of bacterial  
820 processes on carbon mass in cloud water and estimated formation rates of  $3.7 \text{ Tg C yr}^{-1}$  of  
821 secondary biological aerosol that are in the range of the POA emissions via sea spray (Gantt  
822 and Meskhidze, 2013). Hence, the emission fluxes presented here can change once such  
823 processes are quantified for these compounds.

824

## 825 4 Summary and Conclusions

826

827 A comprehensive chemical investigation of the OC in the tropical Atlantic Ocean and  
828 the atmosphere with a focus on its contribution to the OC on the marine aerosol particles in  
829 this particular region was performed.

830 Regarding seawater, a similar distribution of the DL and DCHO was found with a small  
831 SML enrichment. However, the DAA, and likely the N-containing compounds in general,  
832 exhibit a high and varying enrichment in the SML (although being less surface active than  
833 lipids). Although conclusions on the detailed processes that lead to the varying DAA  
834 concentrations and the high SML enrichments cannot be resolved here and the sample  
835 number is limited, the results suggest that processes leading to changes in the organic matter  
836 composition within the upper  $100 \mu\text{m}$  oceanic layer are more pronounced for the group of

837 amino acids (and possibly for nitrogen groups in general) compared to other organic  
838 compounds groups such as lipids and carbohydrates. The SML is probably a very complex,  
839 heterogeneous, seasonality-dependent, and reactive matrix forming a lipid-rich nanolayer.  
840 The same compounds studied in the seawater were found on the ambient submicron aerosol  
841 particles and strongly enriched with respect to sea salt ( $EF_{aer.} = 10^3$  for the carbohydrates and  
842 the amino acids). To this end, the lipids were even more enriched in the submicron aerosol  
843 particles ( $EF_{aer.} = 10^5$ ) compared to the other groups. This indicates a preferred transfer of the  
844 lipids (towards the carbohydrates and the amino acids) from the ocean to the atmosphere  
845 likely driven by their physicochemical properties (high surface-activity and/or the lipophilic  
846 character). Detailed molecular analysis of the seawater and aerosol particles revealed changes  
847 in the relative composition of the single compounds. They were most pronounced for the  
848 amino acids and are likely related to an in situ atmospheric processing by biotic and/or abiotic  
849 reactions that require further investigations. A high saccharide fraction, as described in other  
850 studies, could not be found on the aerosol particles, at least when regarding the molecular-  
851 resolved carbohydrate analysis. However, saccharidic-like components (e.g. glycolipids) are  
852 also included in the lipid fraction analysed here in non-negligible concentrations. This shows  
853 that when comparing the concentrations of substance groups, the analytical methods used  
854 must be taken into account. Nevertheless, even small concentrations of carbohydrates and  
855 amino acids on marine aerosol particles can have a high impact in their microphysical  
856 properties, e.g. as ice nucleating particles, and are worth further studying.

857         Altogether, the marine aerosol particles analysis applied here shows that half of the  
858 OC can be attributed to specific components or component groups. However, the molecular-  
859 level analysed fraction explains only a small part of the OC, the  $CHO_{aer.}$  and  $AA_{aer.}$  made up less  
860 than 1%. This shows that the typical representatives of carbohydrates and amino acids within  
861 the marine OC measured here can explain only a very small fraction of the organic composition  
862 of the aerosol particles on a molecular level. Amines, MSA, oxalic acid carbonyls comprise a  
863 fraction of around 6%. Lipid analysis reveals 43% of the OC on the aerosol particles, however,  
864 the  $Lipid_{aer.}$  composition on a molecular level cannot be obtained from the measurements  
865 performed here. Altogether, about 50% of the OC remained uncharacterized on a molecular  
866 and semi-molecular level. Regarding further marine aerosol analysis, it will be important to  
867 resolve the large part of lipid compounds in more detail, as well as getting molecular-level  
868 information on the remaining, unidentified OC. This shows the need for further detailed  
869 analytical OC studies in the marine environment to resolve formation and transfer  
870 mechanisms.

871         Nevertheless, the results obtained here show that even in such an oligotrophic region,  
872 at least half of the OC on the aerosol particles consists of rather short-lived biogenic material,  
873 likely from the surface ocean, as (qualitatively) suggested by other studies (Choi et al.,  
874 2019;Schmitt-Kopplin et al., 2012). The non-resolved OC might in part be of recalcitrant  
875 nature, as indicated in other studies (Beaupre et al., 2019;Kieber et al., 2016;Lawler et al.,  
876 2020). However, the (potentially) recalcitrant OC does not constitute the majority of the OC  
877 in the oligotrophic Atlantic Ocean. Future studies should complement the data presented here  
878 with investigations of the particulate OC fraction.



879 Finally, since large parts of the open oceans are oligotrophic, the findings of this study  
880 might be relevant to the majority of the world oceans.

881

882 Data availability. The amino acid and carbohydrate data are listed in the SI. The lipid data are  
883 available through the World Data Centre PANGAEA under the following link:  
884 <https://doi.org/10.1594/PANGAEA.921832> (Triesch et al., 2020). Further data can be made  
885 available by the authors upon request.

886

887 Special issue statement.

888

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903

904 Author contributions

905 MvP led the MarParCloud campaign with support from KWF and HH. SZ performed the  
906 analytical measurements of the carbohydrates and supported the data analysis. SF was in  
907 charge of the lipid measurements. NT performed the measurements of the amino acids. MvP  
908 performed the data interpretation and wrote the manuscript with contributions from all  
909 authors.

910

911 Competing interest

912 The authors declare that they have no conflict of interest.

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1293 **Caption of Figures:**

1294 Figure 1: Box and whisker plot of the concentrations in seawater ( $\mu\text{g L}^{-1}$ ), distinguished into  
1295 SML and bulk water for the dissolved amino acids (DAA) in the SML ( $n = 6$ ) and in bulk water  
1296 ( $n = 6$ ), the dissolved carbohydrates (DCHO) in the SML ( $n = 3$ ) and in bulk water ( $n = 3$ ) and for  
1297 the dissolved lipids (DL) in the SML ( $n = 6$ ) and in bulk water ( $n = 13$ ).

1298 Each box encloses 50% of the data with the mean value represented as an open square and  
1299 the median value represented as a line. The bottom of the box marks the 25% limit of the  
1300 data, while the top marks the 75% limit. The lines extending from the top and bottom of each  
1301 box are the 5% and 95% percentiles within the data set, while the asterisks indicate the data  
1302 points lying outside of this range ("outliers").

1303

1304 Figure 2: Bar graph showing the average of the relative compositions (mol%) of dissolved lipids  
1305 (DL) and *Lipids<sub>aer.</sub>* (a), dissolved carbohydrates (DCHO) and *CHO<sub>aer.</sub>* (b) and dissolved amino  
1306 acids (DAA) and *AA<sub>aer.</sub>* (c) in the bulk water (blue bars), the SML (orange bars), and the  $\text{PM}_{10}$   
1307 aerosol particles (grey bars). Mol% were calculated from the molar masses of the respective  
1308 analytes. For the lipid groups, the molar masses of the surrogate standard (sec. 2.2.2 and Tab.  
1309 S8) are applied. The relative mol fractions are relative to the total of each type (DAA, DCHO,  
1310 and DL) analysed.

1311

1312 Figure 3: Box and whisker plot of the concentrations in the  $\text{PM}_{10}$  aerosol particles ( $\text{ng m}^{-3}$ );  $n =$   
1313 8 for *CHO<sub>aer.</sub>*,  $n = 7$  for *AA<sub>aer.</sub>*,  $n = 14$  for *Lipids<sub>aer.</sub>*

1314 Each box encloses 50% of the data with the mean value represented as an open square and  
1315 the median value represented as a line. The bottom of the box marks the 25% limit of the  
1316 data, while the top marks the 75% limit. The lines extending from the top and bottom of each  
1317 box are the 5% and 95% percentiles within the data set, while the asterisks indicate the data  
1318 points lying outside of this range ("outliers").

1319

1320 Figure 4: Scheme underlining the seawater (SML and bulk water) as well as the  $\text{PM}_{10}$  relative  
1321 compositions of DL / *Lipids<sub>aer.</sub>*, DCHO / *CHO<sub>aer.</sub>* and DAA / *AA<sub>aer.</sub>*. Assignment: amino acids:  
1322 neutral/polar: Phe, Gly, Ser, Tyr, neutral/non-polar: Thr, Ala, Pro, Val, Leu, Iso, acidic: Asp, Glu;  
1323 carbohydrates: basic: GlcN, GalN, neutral: Fuc, Rha, Ara, Gal, GIC, Xyl, Man, acidic: MurAc,  
1324 GasAc, GlAc; lipids: hydrocarbons (HC), sterols (ST), pigments (PIC), fatty acid methyl ester  
1325 (MW), membrane component: WE, metabolic reserve: TC, degradation lipids: FFA, ALC, 1,3  
1326 DG, 1,2 DG, MG, glycolipids: MGDG, DGDG, SQDG, polar lipids: PE, PG, PC.

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1328 Figure 5: Graph showing the identified and non-identified OC and the OC contribution of the  
1329 respective organic compound groups in the  $\text{PM}_{10}$  aerosol particles. The contribution of the  
1330 measured organic compounds to the total OC fraction was calculated on a carbon basis.

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1333

1334 Table 1: Average concentrations of the organic groups and enrichment factor (EF) in the SML  
 1335 ( $EF_{SML}$ ) and in the aerosol particles ( $EF_{aer.}$ ) after equations 1 and 2. EF are calculated from the  
 1336 average concentrations of the respective groups (values in Tab. 1). For  $EF_{aer.}$  the average  $Na^+$   
 1337 concentration in seawater ( $1.0E+04 \text{ mg L}^{-1}$ ) and the average  $Na^+$  concentrations in the  $PM_1$   
 1338 particles from the MarParCloud campaign ( $100 \text{ ng m}^{-3}$ , values from Triesch et al., (2021b))  
 1339 were applied. To investigate the variability of the  $EF_{aer.}$  the minimum ratio of  $c_{(analyte)aer.} / c_{(Na+)aer.}$  vs. the maximum ratio of  $c_{(analyte)seawater} / c_{(Na+)seawater}$  was applied (and vice versa) and  
 1340 the range of  $EF_{aer.}$  is given in brackets.

1341 For comparison, the last column lists the  $EF_{aer.}$  for  $PM_1$  from a chamber study (Rastelli et al.,  
 1342 2017).  
 1343

1344

DCHO							
SML ( $\mu\text{g L}^{-1}$ )	Bulk water ( $\mu\text{g L}^{-1}$ )	$PM_1$ ( $\text{ng m}^{-3}$ )	$EF_{SML}$	$EF_{aer}$ (related to the SML)	$EF_{aer}$ (related to the bulk water)	$EF_{aer}$ (average)	$EF_{aer}$ (Rastelli et al., 2017)
$85 \pm 30$	$78 \pm 15$	$1.0 \pm 1.1$	$1.1 \pm 0.5$	$1.26E+03$ ( $2.33E+02$ - $5.93E+03$ )	$1.35E+03$ ( $1.54E+02$ - $4.56E+03$ )	$1.31E+03$	$1.00E+05$
n = 3	n = 3	n = 8					
DAA							
SML	Bulk water	$PM_1$	$EF_{SML}$	$EF_{aer}$	$EF_{aer}$	$EF_{aer}$	$EF_{aer}$
190 ± 238	$80 \pm 53$	$2.4 \pm 1.1$	$2.3 \pm 0.4$	$9.23E+02$ ( $1.39E+02$ - $1.62E+04$ )	$2.07E+03$ ( $7.95E+02$ - $2.19E+04$ )	$1.50E+03$	$1.20E+05$
n = 6	n = 3	n = 7					
DL							
SML	Bulk water	$PM_1$	$EF_{SML}$	$EF_{aer}$	$EF_{aer}$	$EF_{aer}$	$EF_{aer}$
$83 \pm 24$	$70 \pm 25$	$120 \pm 43$	$1.3 \pm 0.2$	$1.42E+05$ ( $6.21E+04$ - $3.92E+05$ )	$1.71E+05$ ( $5.83E+04$ - $5.49E+05$ )	$1.57E+05$	$1.40E+05$
n = 6	n = 13	n = 14					

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