

1 **Free amino acids in Antarctic aerosol: potential markers for the evolution and**  
2 **fate of marine aerosol**

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22 **Abstract**

23 To investigate the impact of marine aerosols on global climate change it is important to study their  
24 chemical composition and size distribution. Amino acids are a component of the organic nitrogen in  
25 aerosols and particles containing amino acids have been found to be efficient ice nuclei.

26 The main aim of this study was to investigate the L- and D- free amino acid composition as possible  
27 tracers of primary biological production in Antarctic aerosols from three different areas: two  
28 continental bases, Mario Zucchelli Station (MZS) on the coast of the Ross Sea, Concordia Station at  
29 Dome C on the Antarctic Plateau, and the Southern Ocean near the Antarctic continent. Studying  
30 the size distribution of amino acids in aerosols allowed us to characterize this component of the  
31 water-soluble organic carbon (WSOC) in marine aerosols near their source and after long-range  
32 transport. The presence of only free L- amino acids in our samples is indicative of the prevalence of  
33 phytoplanktonic material. Sampling at these three points allowed us to study the reactivity of these  
34 compounds during long-range transport.

35 The mean total amino acid concentration detected at MZS was  $11 \text{ pmol m}^{-3}$ , a higher percentage of  
36 amino acids were found in the fine fraction. The aerosol samples collected at Dome C had the  
37 lowest amino acid values ( $0.7$  and  $0.8 \text{ pmol m}^{-3}$ ) and the coarse particles were found to have higher  
38 concentrations of amino acids compared to the coastal site. The amino acid composition in the  
39 aerosol collected at Dome C had also changed compared to the coastal site, suggesting that physical  
40 and chemical transformations had occurred during long range transport.

41 During the sampling cruise on the R/V *Italica* on the Southern Ocean, high concentrations of amino  
42 acids were found in the total suspended particles, this we attribute to the presence of intact  
43 biological material (as microorganisms or plant material) in the sample.

44

## 45 **1. Introduction**

46 The organic composition of marine aerosols is particularly interesting as it contributes a substantial  
47 portion of the world-wide aerosol mass, especially in the submicron size fraction (Bigg, 2007). The  
48 study of marine aerosols is of interest as anything that can change their size, composition or  
49 concentration in the atmosphere may have an impact on the Earth's climate, since as noted by  
50 O'Dowd et al., (2004) "Marine aerosol contributes significantly to the global aerosol load and  
51 consequently has an important impact on both the Earth's albedo and climate". This is because, the  
52 sheer extent of the ocean means that marine aerosol is one of the most important natural aerosol  
53 sources on a global scale (O'Dowd and De Leeuw, 2007, Rinaldi et al. 2010). Several studies  
54 (Facchini et al., 2008a,b; Rinaldi et al., 2010) have demonstrated that the organic chemical  
55 composition of marine aerosols depends on a combination of different factors, such as primary  
56 emission via bubble bursting and the subsequent transformation into secondary aerosol. During the  
57 primary emission *via* bubble bursting processes, the presence of phytoplankton can further alter the  
58 organic chemical composition and physical properties of marine aerosols (Kuznetsova et al., 2005).  
59 The organic fraction of marine aerosols contains water-soluble organic compounds (WSOC), which  
60 include numerous species of organic acids, amines, carbonyl compounds and amino acids (Saxena  
61 and Hildemann, 1996). Amino acids are ubiquitous compounds, and are an active component of the  
62 organic nitrogen content of aerosols because some of them have been shown to enhance the ice  
63 nucleating ability of atmospheric particles (Szyrmer and Zawadzki, 1997). Recently Kristensson et  
64 al., (2010) investigated the ability of some amino acids (e.g. glycine or leucine) to act as cloud  
65 condensation nuclei (CCN), they found that particles containing amino acids at "atmospherically  
66 relevant mixture ratios" are good CCN. These compounds can also serve as a source of nutrients for  
67 marine ecosystems due to their high bioavailability (Zhang et al., 2002).

68 A large number of studies have confirmed the presence of amino acids in the condensed phase of  
69 aerosols (Gorzelska and Galloway, 1990; Spitzzy, 1990; Milne and Zika, 1993; Saxena and

70 Hildemann, 1996; Zhang et al., 2002; Zhang and Anastasio, 2003; Mandalakis et al., 2010;  
71 Mandalakis et al., 2011; Ge et al., 2011 and its references), in rainwater (Mopper and Zika, 1987;  
72 Mace et al., 2003a,b), fog (Zhang and Anastasio, 2001), and in dew water (Scheller, 2001). They  
73 can be present as dissolved combined amino acids (proteins and peptides) (Kuznetsova et al., 2005;  
74 Ge et al., 2011), dissolved free amino acids from the hydrolysis of the combined amino  
75 acids (Mopper and Zika 1987; Milne and Zika, 1993), and particulate amino acids (from solid  
76 microorganisms and debris particles inside the liquid aerosol phase) (Kuznetsova et al., 2005).

77 Several emission sources can affect not only the total concentration of dissolved free amino acids in  
78 the atmosphere, but also the amino acid composition of the aerosol. Amino acids have been  
79 detected in volcanic emissions (Mukhin et al., 1978; Scalabrin et al., 2012), biomass burning has  
80 also been suggested as a possible source of amino acids as part of the WSOC content (Mace et al.,  
81 2003a; Chan et al., 2005). The different amino acids found in continental particles are thought to  
82 have been originally produced by plants, pollens and algae, as well as fungi and bacterial spores  
83 (Milne and Zika, 1993; Scheller, 2001; Zhang and Anastasio, 2003; Mace et al., 2003a) and can be  
84 found in high concentrations in soil and desert dust. The continental contribution was evaluated by  
85 Mace et al. (2003b), who found that biogenic amino acids were present in the fine particles and that  
86 coarse particles contained amino acids from mainly anthropogenic sources. The anthropogenic  
87 sources currently identified are tobacco smoke (Ge et al., 2011), incinerators, waste collection  
88 centers and sewage treatment plants (Leach et al., 1999). Zhang and Anastasio (2002) identified  
89 livestock farming as the main source of amino acid ornithine in Californian aerosols. Matsumoto  
90 and Uematsu (2005) describe how long-range transport influences the concentration of amino acids  
91 in the North Pacific Ocean, while an evident marine source was verified by Weydan and Preston  
92 (2008) in the South Atlantic Ocean. Several studies investigated the free dissolved amino acids in  
93 marine aerosols (Gorzelska and Galloway, 1990; McCarthy et al., 1998; Mace et al., 2003;  
94 Matsumoto and Uematsu, 2005; Kuznetsova et al., 2005; Weydan and Preston; 2008; Mandalakis et  
95 al., 2011) but few studies have been conducted in the polar regions. Schmale et al. (2013) conducted

96 a complete study on the characterization of Sub-Antarctic marine aerosols and they identified  
97 hatching penguins as a source of amino acids in the aerosol of Bird Island in the Southern Atlantic  
98 Ocean. To our knowledge, this paper is the first to investigate the different compositions and  
99 particle-size distributions of amino acids in Antarctic aerosols.

100 Chirality is an important feature of amino acids and the homochirality of life on Earth occurs  
101 because L-amino acids are the only enantiomers used during the biosynthesis of proteins and  
102 peptides (Cronin and Pizzarello, 1997). The principal biochemical source of D-amino acids are  
103 peptidoglycans, the main structural components of bacterial cell walls (Voet and Voet, 1999).  
104 Chiral information can be useful in revealing the primary and secondary origins of aerosol  
105 components as demonstrated by several recent studies (Kuznetsova et al, 2005; Wedyan and  
106 Preston, 2008; Nozière et al., 2011; González et al., 2011; González et al., 2014). Amino acid  
107 enantiomeric ratios can be powerful markers for characterizing nitrogenous materials (McCarthy et  
108 al., 1998). Kuznetsova et al. (2005) indicated that the relative enrichment in L-amino acids may  
109 result from planktonic particles that concentrate at the sea surface while D-enantiomers come  
110 predominantly from bacteria (Wedyan and Preston, 2008). Therefore the presence of free D-isomers  
111 is indicative of a larger proportion of bacteria in aerosols (Wedyan and Preston, 2008).

112 The aims of this study are to investigate the occurrence and concentration levels of dissolved free L-  
113 and D-amino acids in the Antarctic aerosols, to determine how these compounds produced from the  
114 seawater surface are distributed in size-segregated aerosols, and to study their compositional and  
115 distribution changes after long-range atmospheric transport.

116 Due to their long distance from anthropogenic and continental emission sources, polar regions are  
117 excellent natural laboratories for conducting studies on the behavior, evolution and fate of marine  
118 aerosols. In Antarctica, long-range atmospheric transport of anthropogenic pollutants is minimal  
119 because the continent is surrounded by the Southern Ocean. This means that natural sources are the  
120 main contributors to atmospheric aerosols (Bargagli, 2008, Bourcier et al., 2010). Our aim is to  
121 study concentrations of airborne amino acids, which may be related to aerosol growth in Antarctica

122 in some circumstances. Our investigation was carried out over three different Antarctic summer  
123 campaigns, including two consecutive field campaigns (2011-2012 and 2012-2013) on the Antarctic  
124 plateau at the Italian-French base of Concordia Station (DC). One sampling period (2010-2011) was  
125 carried out at the Italian coastal base MZS and finally, aerosols were sampled from the R/V *Italica*  
126 on the Southern Ocean, between Antarctica and New Zealand (2012).

## 127 **2. Experimental section**

### 128 **2.1 Sample collection**

129 Aerosol sampling was carried out over three different Antarctic expeditions during the austral  
130 summer period, in the framework of the “Progetto Nazionale di Ricerche in Antartide” (PNRA).  
131 The sampling sites are shown in Fig. 1, obtained using Google Earth maps.

132 During the first expedition one sampling campaign collected five aerosol samples from the Italian  
133 base MZS from 29<sup>th</sup> November 2010 to 18<sup>th</sup> January 2011. The sampling site was at the Faraglione  
134 Camp (74° 42' S – 164° 06' E), about 3 km south of MZS in Victoria Land. The site is a promontory  
135 at 57 m asl. It was chosen because it is located in a valley that is physically separated from the main  
136 station area by a hill, to reduce as much as possible eventual pollution from the research station.

137 During the second expedition four aerosol samples were collected from the 19<sup>th</sup> December 2011 to  
138 28<sup>th</sup> January 2012 at the Italian-French base Concordia Station located at Dome C (DC) on the East  
139 Antarctic plateau (75° 06' S – 123° 20' E), and seven other samples retrieved from the Ross Sea  
140 (Antarctica) on the R/V *Italica* during the oceanographic sampling campaign from 13 January to 19  
141 February 2012 (Fig 1).

142 In the third expedition, five aerosol samples were obtained from 07<sup>th</sup> December 2012 to 26<sup>th</sup> January  
143 2013 at Dome C. The sampling site at Dome C during both expeditions was located about 1 km  
144 south-west of the Concordia Station buildings, upwind of the dominant wind direction (from the  
145 south-west). Aerosol samples from the terrestrial bases (MZS and DC) were collected using a TE-

146 6070, PM10 high-volume air sampler (average flow  $1.21 \text{ m}^3 \text{ min}^{-1}$ ) equipped with a Model TE-235  
147 five-stage high-volume cascade impactor (Tisch Environmental Inc.) fitted with a high-volume  
148 back-up filter (quartz fiber filter Media 8" x 10") and a 5.625" x 5.375" slotted quartz fiber filter for  
149 collecting particle size fractions in the following ranges:  $10.0 - 7.2 \mu\text{m}$ ,  $7.2 - 3.0 \mu\text{m}$ ,  $3.0 - 1.5 \mu\text{m}$ ,  
150  $1.5 - 0.95 \mu\text{m}$ ,  $0.95 - 0.49 \mu\text{m}$ ,  $< 0.49 \mu\text{m}$ . The sampling period for each sample was 10 days, for a  
151 total air volume of  $\sim 15,000 \text{ m}^3$  per sample.

152 During the oceanographic cruise, airborne aerosols were collected onto circular quartz fiber filters  
153 (SKC Inc., Eighty Four, To-13 model) using a TE 5000 High Volume Air Sampler (Tisch  
154 Environmental Inc.) to determine the TSP (total suspended particulate) fraction, defined as particles  
155 with a diameter  $> 1 \mu\text{m}$ . To avoid contamination from the ship's exhaust, air samples were  
156 automatically taken under wind sector control. The sampler was located at the bow and sampling  
157 only took place when the wind came from between  $-135^\circ$  to  $135^\circ$  relative to the bow and ship  
158 direction and when the relative wind speed was  $> 1 \text{ m s}^{-1}$ . The sample collection was set to five days,  
159 but the actual sampling time varied, subject to wind sector and speed control as well as cruise  
160 events. Due to these events the actual aerosol sampling volumes varied from between 511 and 2156  
161  $\text{m}^3$ . The sea voyage track chart is reported in Fig. 1.

162 All filters were pre-combusted (4 h at  $400^\circ\text{C}$  in a muffle furnace), to avoid contamination they were  
163 wrapped in two aluminum foils, after sampling they were re-wrapped in clean double aluminum foil  
164 and were stored at  $-20^\circ\text{C}$  prior to analysis. Field blank samples were collected by loading, carrying  
165 and installing the filter holder into the instrument with the air pump closed.

## 166 **2.2 Sample processing**

167 To avoid contamination from laboratory air particles and from the operator, samples were handled  
168 under a clean laminar flow bench (class 100). The pre-analytical and sample extraction protocol has  
169 been previously described in detail by Zangrando et al.(2013) for other compounds. The same

170 protocol is summarized below and was applied to the identification of amino acids in Antarctic  
171 samples.

172 Each quartz fiber filter was cut in half using stainless steel scissors that were previously washed  
173 with methanol. Filters were broken into small pieces using clean tweezers, and were placed into  
174 50mL conical flasks. Slotted quartz fiber filters from the cascade impactor and circular quartz fiber  
175 filters from the TSP samplers were treated in the same way. They were spiked with 100  $\mu\text{L}$  of  $^{13}\text{C}$   
176 isotopically-labelled amino acid standard solutions (with concentrations ranging between 2 and 3  
177  $\mu\text{g mL}^{-1}$ ), they were then ultrasonically extracted twice for 15 minutes in an ice bath with 5 mL and  
178 then 2 mL of ultrapure water. The extracts were combined and filtered through a 0.45  $\mu\text{m}$  PTFE  
179 filter in order to remove particulate and filter traces before instrumental analysis.

180 The larger high volume back-up filters were spiked with 400  $\mu\text{L}$  of internal standard solution and  
181 were extracted with 25 mL then 5 mL of ultrapure water in an ultrasonic ice bath as described  
182 above.

### 183 **2.3 Instrumental analysis**

184 The enantiomeric determination of free L- and D-amino acids by HPLC-MS/MS has been described  
185 in detail by Barbaro et al. (2014). This instrumental method has been applied to the aqueous extracts  
186 of the aerosol samples collected during this study.

187 An Agilent 1100 Series HPLC Systems (Waldbronn, Germany; with a binary pump, vacuum  
188 degasser, autosampler) was coupled with an API 4000 Triple Quadrupole Mass Spectrometer  
189 (Applied Biosystem/MSD SCIEX, Concord, Ontario, Canada) using a TurboV electrospray source  
190 that operated in positive mode by multiple reaction monitoring (MRM).

191 Chromatographic separation was performed using a 2.1x 250 mm CHIROBIOTIC TAG column  
192 (Advanced Separation Technologies Inc, USA) with a two mobile eluents. Eluent A is ultrapure  
193 water with 0.1% v/v formic acid and eluent B is ultra pure methanol with 0.1% v/v formic acid.



194 A binary gradient elution program was followed at a flow rate of 0.2 mL min<sup>-1</sup>: 0-15 min, an  
195 isocratic step with 30% of eluent B; 15-20 min, a gradient from 30 to 100% B; 20-25 min an  
196 isocratic washing step with 100% of eluent B; 27-30 min, re-equilibration to 30% eluent B. The  
197 injection volume was 10 µL.

198 In this work the amino acids were quantified using the isotope dilution method where an  
199 isotopically labeled standard was available. For other amino acids, where a labeled standard was  
200 unavailable, an internal standard was used to quantify the analytes. A detailed description of which  
201 analytes are quantified with which method can be found in Barbaro et al. (2014). In both cases, the  
202 results were corrected for daily instrumental sensitivity variations by evaluating the instrumental  
203 response factors.

204 Reagents and materials used for this study are reported in the Supplement.

## 205 **2.4 Quality control**

206 The entire analytical procedure was validated by estimation of trueness, repeatability and efficiency  
207 (yield%) of the sample treatment process as described by Bliesner (2006). To ensure that it was fit  
208 for purpose for the enantiomeric determination of amino acids in Antarctic aerosol, the validation  
209 was carried out by spiking five cleaned quartz filters (for each type of filter) with 100 µL of a  
210 solution containing all the native L and D amino acids (with concentrations ranging between 2 and  
211 4 µg mL<sup>-1</sup>) and 100 µL of a solution containing all the isotopically-labeled<sup>13</sup>C amino acids  
212 (concentrations ranging between 2 and 3 µg mL<sup>-1</sup>). The filters were subsequently extracted as  
213 described above in section 2.2 “Sample processing”.

214 Tables S1, S2 and S3 report a summary of the yields, trueness and relative standard deviations  
215 (n=5) for each type of filter used in this study. Average yields of 61%, 56% and 56% were obtained  
216 from the circular, slotted and backup filters, respectively. In some cases, these values are lower than  
217 those reported in the literature (Mandalakis et al., 2010; Barbaro et al., 2011). Trueness is the most  
218 important parameter to determine during a method validation; it refers to the degree of closeness of

219 the determined value to the known "true" value. It is expressed as an error, calculated as  $(Q - T)/T$   
220  $\times 100$ , where Q is the determined value and T is the "true value".

221 For the circular filters, all D- and L-amino acids considered in this work were validated with an  
222 error percentage ranging from -13% (D-Leu/D-Ile) to +8% (L-Tyr).

223 In the backup filters, only D- and L-Hys produced unacceptable percent errors, for this reason these  
224 compounds were excluded from the quantification. The other amino acids considered in this study  
225 were quantified with an accuracy ranging from -9% (D-Met) to +9% (D-Ala, L-Thr).

226 Some amino acids (D-Ala, L-Asn, D-Asn, D-Glu, D-Phe, L-Ser, D-Ser, and D-Val) were excluded  
227 from the quantification using the slotted quartz fiber filters as very high percent errors were  
228 calculated. We believe that this behavior is probably due to the different mode of use of this  
229 sampling support: the slotted quartz fiber filters were used as impact supports while the other  
230 supports were used as filters. The other amino acids studied in this work had percent error values  
231 between -13% (D-Tyr) and +13% (D-Leu/D-Ile) and so the method was fit for purpose for their  
232 quantification.

233 The repeatability is determined as the relative standard deviation of the analytical results for the 5  
234 spiked filters. For each type of filter used in this study, the repeatability was always below 10%.

235 The method detection limit (MDL) for the analytical procedure is defined as three times the  
236 standard deviation of the average values of the field blank (n=3). Tables S1, S2 and S3 report the  
237 relative MDLs for each quantified amino acid in the three different sampling supports, the absolute  
238 mean blank values (n=3) in these tables are subtracted from the analytical results. All discussions in  
239 the following sections below are based upon blank corrected values.

240 A comparison between previously published data (Barbaro et al., 2011; Matsumoto and Uematsu,  
241 2005) and the MDLs obtained for each type of filter in this work shows that we obtained lower  
242 blank values than those previously reported.

## 243 **2.5 Back-trajectory calculation and satellite imagery**

244 Backward air trajectories arriving at MZS, Dome C and R/V *Italica* were computed using a Hybrid  
245 Single Particle Lagrangian Integrated Trajectory (HYSPLIT) transport and dispersion models  
246 (Draxler and Rolph, 2013). The meteorological data used for computing all the backward  
247 trajectories were the NCEP/NCAR Global Reanalysis Data. For MZS data, a vertical velocity  
248 model was used while an isentropic model was employed for the analysis of DC air masses, as  
249 suggested by Stohl et al (2010).

250 240 hours of back-trajectories beginning at MZS and DC were calculated for each sampling  
251 campaign period. Four runs were computed for every sampling day at six hour intervals and the  
252 resulting multiple trajectories were “mean-clustered aggregated” into 6 groups, based on the scree-  
253 plot analyses of total spatial variance.

254 A sensitivity study has been performed to verify the stability of the HYSPLIT back trajectory  
255 calculations. We calculated the back-trajectories beginning at 10 m agl (above ground level), 100  
256 m, 500 m and 1000 m at MZS and DC to evaluate how the trajectories varied with height. The  
257 results are shown in Supplementary Fig. S1-S3. It can be seen that the clusters of simulated air  
258 masses have similar trajectories although with different percentages of the total number of  
259 calculated back trajectories. For this study we used the 500 m back trajectories because we want to  
260 evaluate long range transport. This is because the mean mixed-layer height is 250–400 m agl at DC  
261 (Argentini et al., 2005) while the boundary-layer height is usually below 50 m at the Antarctic coast  
262 (Handorf et al., 1999).

263 We have also estimated the stability of the HYSPLIT model by varying the position of source at  
264 MZS as well as DC using a 121 point matrix built by adding or subtracting one degree of latitude or  
265 longitude from the real source for each sampling day. These back-trajectories calculated from the  
266 121 simulated sources have the same behavior (Supplement Fig. S4-S6), thus confirming the  
267 stability of the HYSPLIT calculations.

268 For the oceanographic cruise, trajectory matrices were performed in order to simulate the ship’s  
269 itinerary. In this case, for each 24-h sampling event, 5-day backward trajectories were computed.

270 The data related to chlorophyll were obtained *via* an Aqua/MODIS NASA satellite continually  
271 orbiting the globe (<http://neo.sci.gsfc.nasa.gov/>).

### 272 **3. Results and Discussion**

#### 273 **3.1 Free amino acid determination in the coastal area**

274 Nine L-amino acids (L-Ala, L-Asp, L-Arg, L-Glu, L-Phe, L-Pro, L-Tyr, L-Thr) and Gly had blank  
275 corrected concentrations higher than the MDLs (Supplementary Tables S2 and S3), while all D-  
276 amino acids had values below the MDLs, probably due to a negligible presence of bacteria in the  
277 aerosol source (Kuznetsova et al., 2005; Wedyan and Preston, 2008). The total concentration of  
278 amino acids, calculated as the sum of their six size distributions in all aerosol samples, has a median  
279 value of 5 pmol m<sup>-3</sup> and a mean value of 11 pmol m<sup>-3</sup>, due to the higher amino acid concentrations  
280 in the first sample (29 November-9 December), as shown in Fig. 2.

281 The mean total concentration of free amino acids determined in this study was very similar to those  
282 found in the literature for marine aerosols in remote areas. Matsumoto and Uematsu (2005) reported  
283 a mean free amino acid concentration of 10.7 pmol m<sup>-3</sup> in aerosol samples above the Pacific Ocean,  
284 while Gorzelska and Galloway (1990) and Wedyan and Preston (2008) observed means of 3 pmol  
285 m<sup>-3</sup> and 20 pmol m<sup>-3</sup> respectively in the Atlantic Ocean. Scalabrin et al. (2012) determined a mean  
286 concentration of 2.8 pmol m<sup>-3</sup> using the same aerosol sampling method reported here at an Arctic  
287 coastal station.

288 Higher mean concentrations of amino acids were found in the Mediterranean. Barbaro et al. (2011)  
289 determined a mean value of 334 pmol m<sup>-3</sup> in the Venice Lagoon (Italy); Mandalakis et al.  
290 (2010,2011) found 166 pmol m<sup>-3</sup> and 172 pmol m<sup>-3</sup> in two studies in the Eastern Mediterranean  
291 around Greece, respectively. In the Southern hemisphere, Mace et al. (2003b) performed several  
292 studies on the coast of Tasmania (Australia), and found mean free total amino acid concentrations  
293 that ranged from between 15 and 160 pmol m<sup>-3</sup>.

294 In this work, we found that the predominant compounds were Gly and Arg, which together  
295 constituted 66-85% of the total amino acid content. Gly and Arg had different proportions in the  
296 five samples, and the other compounds were present in similar proportions in all the samples, with  
297 average percentages of 9% for Glu, 7% for Ala, 5% for Thr, 4% for Asp, 2% for Val while 1% for  
298 other amino acids (Phe, Tyr and Pro). In Fig.2 it can be seen that the first sample collected between  
299 29 November and 09 December had a high proportion of Arg (74%), compared to Gly (11%). In  
300 contrast to this, in the other samples, Gly was the predominant compound, with a percentage  
301 between 48 to 56%, while Arg was present as 18% of the total.

302 Scheller (2001) demonstrated that high quantities of Arg were closely linked with plant growth, but  
303 the cluster means backward trajectories (Fig. 3) calculated for our samples show that 1% of the air  
304 masses come from open-ocean areas whilst the major part (99%) principally come from the interior  
305 of the Antarctic continent, areas that are characterized by a lack of vegetation. This suggests that the  
306 local marine influence was probably the main source of amino acids in the aerosol collected at MZS  
307 and that the concentration of coastal atmospheric amino acids is probably linked to local primary  
308 production in the Ross Sea, as suggested by studies in other areas (Meskhidze and Nenes, 2006;  
309 Vignati et al., 2010; Yoon et al., 2007; Müller et al., 2009). We hypothesize that the main source of  
310 Arg in the aerosols collected at the coastal Antarctic station MZS was probably a diatom bloom as  
311 Arg is involved in their urea cycle (Bromke, 2013). The MODIS data (Fig. 4) show higher  
312 chlorophyll concentrations during the period covered by the first sampling period, while a strong  
313 decrease in the biomass production index was observed in the other sampling times. This  
314 relationship between marine primary production and Arg concentration suggests that this amino  
315 acid may have a marine biological origin and that its concentration is closely linked to algae  
316 growth.

317 Meteorological conditions play an important role in aerosol formation processes. The first sampling  
318 period (29 November-09 December) was characterized by temperatures ranging between -10°C and  
319 -1.5°C, while in the successive sampling periods, the air temperature was always above -2°C

320 (PNRA-ENEA, 2014). Studies conducted on the sea surface microlayer (Grammatika and  
321 Zimmerman, 2001; Knulst et al., 2003) established that air temperatures  $< -5^{\circ}\text{C}$  create surface slurries  
322 which may result in the expulsion of salts and particulate organic matter. Under such conditions,  
323 near-surface turbulence was increased, leading to an increase of material in the microlayer, where  
324 bubble formation and bursting actively contributed to the transport mechanisms. Leck and Bigg  
325 (2005) showed that the main occurrences of fine aerosol formation in the arctic atmosphere were  
326 observed when the ice pack is cracking forming leads that melt and refreeze. Our first sample was  
327 collected when the pack ice was melting and refreezing, and we did in fact observe the highest  
328 concentration of total amino acids in the fine aerosols during this period.

329 The hypothesis of a local marine source for the aerosols collected at the coastal station MZS was  
330 also confirmed by the distribution of the amino acids in the different particle size fractions. Fig. 2  
331 shows that 98% of the total free amino acids are generally found in the fine particles ( $< 1\mu\text{m}$ ,  
332 combined S5 and B filters). While the remaining 2% is evenly distributed over the other coarser  
333 fractions  $> 1\mu\text{m}$  (filter stages S1 to S4). Our experimental data is consistent with the observations of  
334 O'Dowd et al. (2004) and Keene et al. (2007) who showed that WSOC in sea spray submicron  
335 particles are mostly associated with the smallest size fraction (0.1-0.25  $\mu\text{m}$ ). Other authors  
336 (Facchini et al., 2008b; Modini et al., 2010) have shown that WSOC were present in all aerosol size  
337 fractions and confirm that the greatest enrichment was in the fine fraction. Our observations are in  
338 line with this literature data as amino acids are part of the WSOC family of compounds and so  
339 should have the same behavior in sea spray submicron particles.

### 340 **3.2 The determination of free amino acids at a remote continental area.**

341 Concordia Station at Dome C is an ideal site for studying the chemical composition of remote  
342 Antarctic aerosol. Several studies (Fattori et al., 2005; Jourdain et al., 2008; Becagli et al., 2012;  
343 Udisti et al., 2012) have investigated the distribution of inorganic compounds and of a few organic

344 molecules (e.g., methanesulfonic acid) in aerosol, but the free amino acid concentration and  
345 composition had not yet been studied.

346 Fig. 5 presents the concentrations of free amino acids collected during both field campaigns, and  
347 shows a similarity between the trends and compositions of the analyzed compounds between the  
348 various size fractions. Ten amino acids (L-Ala, L-Arg, L-Asp, L-Glu, L-Leu, Gly, L-Phe, L-Thr, L-  
349 Tyr, L-Val) had concentrations above MDLs (Supplementary Tables S2 and S3) in all samples  
350 collected in both field campaigns. The concentrations of D-amino acids were always below MDLs,  
351 as seen in our coastal results. It was observed that Gly, L-Asp and L-Ala together accounted for  
352 about 80% of the total amino acid content. The total mean free amino acid concentrations, as the  
353 sum of the free amino acid concentrations in all the sample stages, were  $0.8 \text{ pmol m}^{-3}$  for the 2011-  
354 2012 campaign and  $0.7 \text{ pmol m}^{-3}$  for 2012-2013 campaign (Fig. 5). To our knowledge, these mean  
355 concentrations are lower than those reported in the literature (Gorzelska and Galloway, 1990;  
356 Milne and Zika, 1993; Mace et al., 2003b; Kuznetsova et al., 2005; Matsumoto and Uematsu, 2005;  
357 Wedyan and Preston, 2008; Mandalakis et al., 2010; Barbaro et al., 2011; Mandalakis et al., 2011;  
358 Scalabrin et al., 2012), suggesting that this aerosol composition may describe the amino acid global  
359 background concentration.

360 In Fig. 5B, the sample collected from 27 December 2012 to 06 January 2013 shows an altered  
361 concentration profile, with the highest concentrations in one of the coarse fractions (S4 stage 1.5-  
362  $0.95 \mu\text{m}$ ). After evaluating the wind rose plots and activity at the base for each sample in the two  
363 summer campaigns, we believe that these samples were contaminated by human activity at  
364 Concordia station (Supplementary Fig. S7).

365

366 The mean concentrations of free amino acids in the coarse aerosol particles collected at DC for the  
367 two field campaigns were  $407$  and  $421 \text{ fmol m}^{-3}$  (see Fig. 5). At our coastal site, the mean free  
368 amino acid concentration in the coarse fraction was  $264 \text{ fmol m}^{-3}$  (Fig. 2). At DC, the free amino  
369 acid concentration in the coarse aerosol, expressed as a percent of the total free amino acids

370 concentration was found to be 13% in 2011-12 and 23% in the 2012-13 campaign. Conversely,  
371 during our 2010-2011 sampling campaign at MZS, which is located near the marine aerosol source,  
372 we found that only 2% of the total free amino acid concentration was present in the coarse fraction.

373

374 During the Antarctic summer, the surface inversion over the polar ice cap is relatively weak and  
375 aerosols produced on the ocean's surface can be transported through the upper troposphere to the  
376 Antarctic plateau where they are easily mixed down to the surface (Cunningham and Zoller, 1981).  
377 There are also transfer mechanisms from the lower stratosphere to the upper troposphere that occur  
378 near the coast of the Antarctic continent. Aerosol from different sources mixes into the upper  
379 troposphere, and this air descends uniformly over the Antarctic plateau due to surface cooling flows  
380 off the plateau causing the katabatic wind. This means that during the summer, there is a continuous  
381 flux of relatively clean air from the upper troposphere with aerosol from high altitude inputs and  
382 long range transport (Cunningham and Zoller, 1981;Stohl and Sodemann, 2010).

383 Cluster means backward trajectories analysis of all the samples collected during both summer  
384 campaigns at DC revealed a prominent marine source (Fig. 3). Fig. 3 shows that the 10-days  
385 backward trajectories came from the Southern Ocean where there are no land based man made  
386 influences.

387 Fig. 5 shows that the concentration of amino acids for the 2011-2012 summer Antarctic campaign  
388 was higher than the values reported for the 2012-2013 Antarctic campaign, and underlines that the  
389 main difference between the two campaigns is mainly in the percentages of amino acids in the  
390 coarse fraction. We suggest that the transport processes of the air masses were the main cause of  
391 these variations as the time spent inland by the air masses in the 2011-2012 summer was about 36  
392 hours (Fig. 3) whilst in 2012-2013 the time range was between 4 and 7 days (Fig. 3).

393 The analysis of the size distribution of the free amino acids (Fig 5) combined with the air mass back  
394 trajectories (Fig. 3) allowed us to suggest that the amino acids in the aerosol collected at DC can  
395 have two possible sources. The first hypothesis is that they were present in primary emitted coarse



396 mode aerosol particles, which come from phytoplanktonic sea spray coarse mode particles  
397 (Matsumoto and Uematsu, 2005), or from soil dust coarse mode particles (Mace et al., 2003).  
398 Particles and their chemical constituents can travel for many weeks in the upper troposphere  
399 without being lost, provided they are not subject to wet deposition, or that the compounds are  
400 reacting in the aerosol phase. The second hypothesis is that amino acids had a marine source and  
401 these aerosols underwent several physico-chemical transformations during long-range transport.  
402 Our results suggest that amino acids were present in the fine particles over the surface of the  
403 Southern Ocean from bubble bursting processes. The air masses subsequently passed into the upper  
404 troposphere and then over the continent where they remained for several days before descending  
405 onto the ice sheet. These fine aerosol particles could either grow during long-range transport, due to  
406 condensation of molecules from the gas phase or by collision of small and large particles  
407 (coagulation) (Petzold and Karcher, 2012;Roiger et al., 2012). However, these processes are  
408 unlikely in Antarctica due to the very clean conditions. The most likely explanation is that the fine  
409 fraction has been subjected to other processes that increased the particle size of the aerosol. The  
410 most likely remaining process is ice nucleation during long-range transport promoted by the intense  
411 cold over the plateau and presence of amino acids in the aerosol particles (Szyrmer and Zawadzki,  
412 1997). The specific reason for the increase of amino acids percentage in the coarse particles is not  
413 clear, based on the available data. In our future investigations, we will also evaluate the aerosol  
414 mass, which is probably a key parameter to measure that will help explain this increase of  
415 concentration in the coarse particles.

416

417 The chemical composition of aerosols may change during long-range transport due to  
418 photochemical, chemical and ionic reactions (Milne and Zika, 1993; Nozière and Còrdova, 2008;  
419 De Haan et al., 2009). Milne and Zika (1993)verified that amino acids are destroyed *via* reactions  
420 with photochemically formed oxidants such as hydroxyl radicals, to form products such as the  
421 ammonium ion, amides and keto-acids. However, in the upper atmosphere, the chemical processes

422 take place at slower rates than in the boundary layer (Roiger et al., 2012). In aqueous-phase  
423 aerosols, glyoxal can react with amino acids, leading to scavenging processes (De Haan et al.,  
424 2009). Recent studies on organic aerosol growth mechanisms (Maria et al., 2004) underlined that  
425 oxidation processes that remove hydrophobic organic compounds, are slower in large carbonaceous  
426 aerosols.

427 From the physicochemical proprieties of amino acids, a “hydropathy” index can be made, as  
428 suggested by Pommie et al. (2004). This classifies the amino acids as hydrophilic (Asp, Hyp, Glu,  
429 Asn, Lys, Gln, Arg), hydrophobic (Ala, Val, Leu, Ile, Met, Phe) or neutral (Gly, Pro, Ser, Thr, Tyr,  
430 Hys). This helps in evaluating the contribution of each kind of amino to each class of aerosols  
431 collected over the three different field campaigns. Fig. 6 shows that the hydrophilic components  
432 were predominant in the locally produced marine aerosols released into the atmosphere near MZS,  
433 while hydrophobic compounds were dominant in the aerosols collected at the continental station  
434 (DC). The low abundance of hydrophobic amino acids in coastal aerosols was also observed by  
435 Mandalakis et al. (2011), and is probably caused by their lower tendency to dissolve in the aqueous  
436 particles contained in coastal aerosols. This classification allows us to hypothesize that a higher  
437 proportion of hydrophilic amino acids reflects a higher water content in the aerosol.

438 A comparison between the concentrations of hydrophobic Ala at the two sampling sites (MZS and  
439 DC) shows a very similar average concentration ( $70 \text{ fmol m}^{-3}$ ) in the coarse particles. This is an  
440 interesting behavior that confirms the hypothesis of limited atmospheric reactivity as proposed by  
441 Maria et al. (2004), who suggested a longer hydrophobic aerosol lifetime as a result of the slower  
442 oxidation rates. Thanks to this phenomenon, Ala significantly contributes to the amino acid content  
443 in these “remote aerosols” as it does not degrade during long range transport.

444 Fig. 6 shows that the main difference between the two campaigns is mainly in the percentage of  
445 hydrophilic and neutral amino acids present. A longer transportation time from the source to the  
446 sampling site would allow chemical transformation through photochemical reactions to take place,  
447 decreasing the concentration of hydrophilic amino acids thus modifying the composition so that the

448 more stable Gly (a neutral component) becomes the main compound (Fig. 6). In the 2012-2013  
449 summer, the time spent inland by the air masses ranged from between 4 and 7 days whilst in the  
450 2011-2012 summer it was only 36 hours.

451 Looking at the acid-base proprieties of the amino acids, some differences can be observed between  
452 two different types of aerosol. As described above, the predominant amino acid in the MZS aerosols  
453 was Arg, which contributed considerably to the percentage of basic compounds (53%). The pH  
454 neutral components represented an important percentage (40% and 68% for coastal and inland  
455 aerosols respectively). Gly is mainly present in large quantities in these aerosols because of its very  
456 low atmospheric reactivity (half life of 19 days) (McGregor and Anastasio, 2001) and its presence is  
457 usually considered an indicator of long-range aerosol transport (Milne and Zika, 1993; Barbaro et  
458 al., 2011). The acid compounds (Asp and Glu) contribution was quite different in the aerosols from  
459 the two different stations: with a low percentage in the coastal samples at MZS (7%) that was in  
460 contrast with the higher content in the aerosols from DC (33% and 26% respectively for the two  
461 consecutive field campaigns). This result can be explained by a study conducted by Fattori et al.  
462 (2005) on the DC aerosol, where high acid content was found. High concentrations of hydrochloric,  
463 nitric and sulfuric acids were found in the aerosol fine fraction, promoting numerous series of acid-  
464 base atmospheric reactions that neutralize the basic compounds. In the atmosphere, amino acids are  
465 present in very low quantities so it is thought that they do not influence the pH of aerosols.  
466 However, the pH of aerosols, can influence the chemical form of the amino acids present.

### 467 **3.3 Free amino acids during an oceanographic cruise**

468 Measurements of free amino acids were carried out on aerosol samples collected on the Southern  
469 Ocean onboard the R/V *Italica* from 13th January to 19th February 2012. Aerosols were sampled  
470 using a TSP sampler that collects particles with a diameter above 1  $\mu\text{m}$ . The first and second  
471 samples covered the track between New Zealand (from Lyttelton harbor) and MZS (Antarctica),  
472 and the sixth and last samples were collected during the return journey between Antarctica and New

473 Zealand. Samples 3, 4 and 5 were collected on the Ross Sea near the Antarctic continent (Fig. 1).  
474 Five L-amino acids (L-Asp, L-Arg, L-Glu, L-Phe, L-Pro) and Gly were present in the samples,  
475 while other L- and D-amino acids had concentrations below MDLs (Supplementary Table S1). The  
476 total concentrations of free amino acids varied between 2 and 12 pmol m<sup>-3</sup>.  
477 The first and last samples had the highest concentrations of free amino acids (Fig. 7), and their  
478 relative sampling periods were characterized by temperatures ranging between -1°C and 18°C  
479 (sample 1), in contrast, temperatures during the remaining sampling periods were always below -  
480 1°C, with a lowest value of -8°C (sample 4). Higher temperatures can facilitate metabolic processes  
481 and accelerate atmospheric chemical reactions, as well as promote bubble bursting from the sea  
482 surface. This is probably the main source of amino acids in our on-ship samples. This is also  
483 supported by the back-trajectory analysis (Supplementary Fig. S8a-g), that demonstrate only a  
484 marine influence for that period. The concentration of amino acids was strongly influenced by sea  
485 conditions during sampling. The field report (Rapporto sulla campagna Antartica, 2012), noted that  
486 during navigation from New Zealand to the ice-pack region, the winds were always above 30 knots,  
487 with maximum values of 60 knots with wave heightsof12 meters. This probably explains the higher  
488 total concentration of free amino acids in the first two samples (12 pmol m<sup>-3</sup>). Along the same track,  
489 but under calmer sea conditions (sample 7), we observed a slight reduction in the total concentration  
490 of free amino acids (8 pmol m<sup>-3</sup>).These values were very similar to those reported by Matsumoto  
491 and Uematsu (2005) in the Pacific Ocean and to those reported by Gorzelska and Galloway  
492 (1990)and Wedyan and Preston (2008) in the Atlantic Ocean. The lowest concentrations were  
493 observed in samples 2 and 6, probably due to the fact that they were collected far from Oceania and  
494 from the Antarctic coast, in an area characterized by expansive pack ice and by temperatures below  
495 -1°C, where the bubble bursting process was reduced.  
496 The samples collected near the Antarctic coast (samples 3,4 and 5) were the most interesting ones  
497 because the results could be compared with the amino acid values detected in the coastal station  
498 MZS. The mean total concentration in the samples collected on the Ross Sea was 3.5 pmol m<sup>-3</sup>,

499 about half of the values detected in our Southern Ocean samples. Such values are similar to the  
500 concentrations observed in the aerosols collected at MZS station (median 5 pmol m<sup>-3</sup>). However,  
501 this is not a true comparison: for the sampling campaign at MZS, a cascade impactor was used to  
502 collect aerosol samples with a particle-size below 10 µm, whereas the data collected during the  
503 cruise was for aerosols with a particle diameter above 1 µm. However, if we exclude data from the  
504 back-up and the fifth slotted filters, the cascade sampler covers a particle size between 0.95 µm and  
505 10 µm (stages 1 to 4), making a comparison between the two data sets more feasible. In the MZS  
506 aerosols, the median value of the amino acids concentration in the aerosols collected on stages 1 to  
507 4 was 1 pmol m<sup>-3</sup> and this concentration was lower than that measured in the cruise's aerosols (3.5  
508 pmol m<sup>-3</sup>). So we suspect that the aerosols with a diameter above 10 µm, that were collected with  
509 the TSP sampler but not the cascade impactor, could be the main source of the difference in amino  
510 acid concentration values in the samples collected on the R/V *Italica*.

511 The back-trajectory analysis (Supplementary Fig. S8C-E) demonstrated that the air masses came  
512 from inland Antarctica, where no vegetation is present. The biological material present in the  
513 atmosphere with a size > 10 µm includes pollens which typically vary between 17-58 µm, fungal  
514 spores between 1-30 µm, and algal spores between 15-120 µm. Instead bacteria have a diameter  
515 between 0.25-8 µm, and viruses have diameters that are typically less than 0.3 µm (Jones and  
516 Harrison, 2004). For this reason, we propose that the biological materials influenced the  
517 concentration of the total free amino acids in the shipboard aerosols.

518 In these samples, the presence of algal spores was also confirmed by the detection of Pro at 4%  
519 (mean value) of the total concentration of amino acids. Fisher et al. (2004) measured the relevant  
520 concentration of Pro in ascospores, demonstrating that this amino acid can be used to identify the  
521 presence of spores in aerosols. In the MZS aerosols, the presence of spores could not be evaluated  
522 because the sampler did not sample the particles >10µm. This is probably the reason why the Pro  
523 concentration was always below MDLs at MZS.

524 Asp was detected in only one sample (sample 5), with a concentration of  $502 \text{ fmol m}^{-3}$ . This value is  
525 very similar to those measured in the two field campaigns on the Antarctic plateau (DC),  
526 considering only the slotted filter stages above  $1 \mu\text{m}$  ( $446$  e  $382 \text{ fmol m}^{-3}$  respectively for the 1  
527 summer field campaigns of 2011-12 and 2012-13). The back-trajectory analysis (Supplementary  
528 Fig. S8E) demonstrated that this air mass came from the plateau, where aspartic acid was a  
529 predominant component of the amino acid content.

530 In the aerosols collected during the cruise, the Arg concentration was very low because the  
531 sampling conducted on board R/V *Italica* during the summer of 2012 excluded fine particles,  
532 whereas Arg was one of the most abundant compounds observed in the coastal station found in the  
533 fine fraction.

534

#### 535 **4. Conclusions**

536 This first study on the size distribution of amino acids in Antarctica has identified possible sources  
537 of marine aerosols in this region and has characterized some chemical and physical transformations  
538 that take place during transport to the interior of the Antarctic continent.

539 Marine emissions of fine particles occurred *via* bubble bursting processes on the surface of the  
540 Southern Ocean. The mean total amino acid concentration detected at MZS was  $11 \text{ pmol m}^{-3}$ , with a  
541 higher percentage of amino acids found in the fine fraction. The aerosol samples collected at Dome  
542 C had the lowest amino acid values ( $0.7$  and  $0.8 \text{ pmol m}^{-3}$ ) and the coarse particles were found to be  
543 enriched with amino acids compared to the coastal site. Numerous chemical and photochemical  
544 events may have contributed to a decrease in the concentration in amino acids in the fine fraction,  
545 and the chemical reactions were faster for hydrophilic compounds than for hydrophobic ones, as  
546 suggested by an observed Ala enrichment.

547 The presence of only the L-enantiomers of free amino acids in Antarctic aerosols suggests that  
548 marine particles were the main sources of free amino acids in this area and that these compounds

549 can be modified when transported to the interior of the continent. Gly and Ala, are the most stable  
550 compounds, and may be used as biogenic markers of long-range marine aerosols. The back-  
551 trajectory analysis demonstrated that the differences in the transport time of air masses inside  
552 Antarctica can result in modifications to the percentage of amino acids in the coarse particles.  
553 The study of aerosols with diameters  $>10 \mu\text{m}$  indicated that bubble bursting processes can also emit  
554 microorganisms that are composed of a higher number of neutral amino acids.

555

556 **Author contributor**

557 E. Barbaro, M. Vecchiato and R. Zangrando designed the experiments, performed the HPLC-MS  
558 analyses, and elaborated the data. A. Gambaro and C. Barbante were the principal investigators of  
559 the project that supported this work. All the authors have helped in the discussion of the results and  
560 collaborated in writing the article.

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578



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748 **Figure captions**

749 Figure 1. The sampling sites: the Italian base “Mario Zucchelli Station” (MZS) (74° 42’ S – 164°  
750 06’ E), the Italian-French base “Concordia Station” (Dome C) ( 75° 06’ S – 123° 20’ E) and the  
751 track chart of the R/V *Italica*.

752 Figure 2. Amino acid size distribution in the samples collected during the summer of 2010-11 at  
753 Mario Zucchelli Station (Antarctica).

754 Figure 3. Cluster means backward trajectories analyses at 500 m aglat the coastal base “Mario  
755 Zucchelli Station” (MZS) during the summer of 2010-2011 and cluster means backward trajectories  
756 at the Italian-French base Dome C (DC) during the summers of 2011-2012 and 2012-2013.

757 Figure 4. Distribution of chlorophyll concentrations in the Ross Sea for each sampling period  
758 obtained through the Aqua/MODIS NASA satellite.

759 Figure 5. Size distributions of amino acid concentrations in the samples collected during the  
760 summer of 2011-12 (A) and during the summer of 2012-13 (B) at the Italian French base  
761 “Concordia Station” (Dome C).

762 Figure 6. Comparison between percentages of hydrophilic, neutral and hydrophobic amino acid  
763 contributions of the aerosols sampled at the Mario Zucchelli Station and at Dome C.

764 Figure 7. Amino acid distribution in the aerosols sampled on the R/V *Italica* during the  
765 oceanographic cruise on the Southern Ocean during the summer of 2012.

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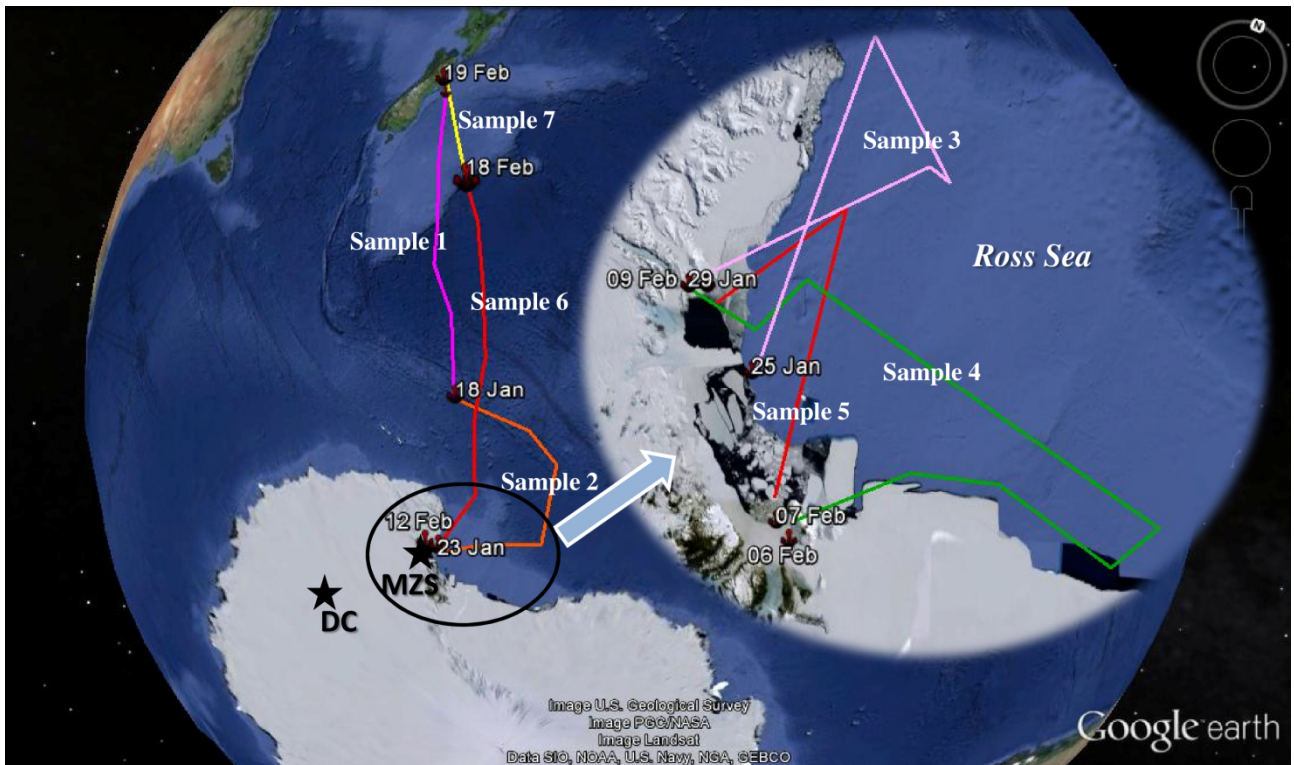
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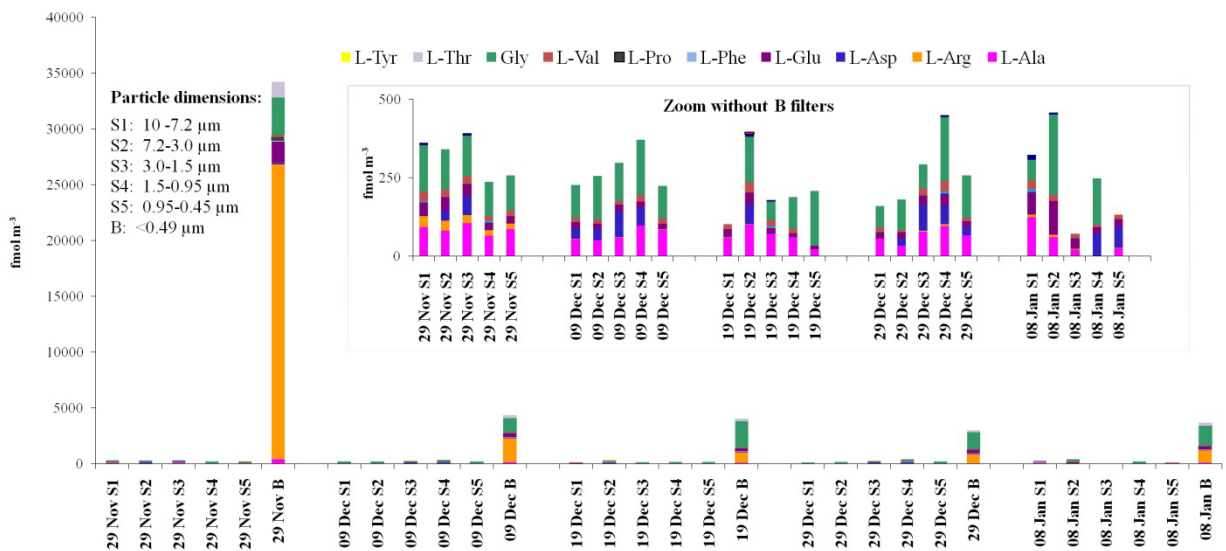
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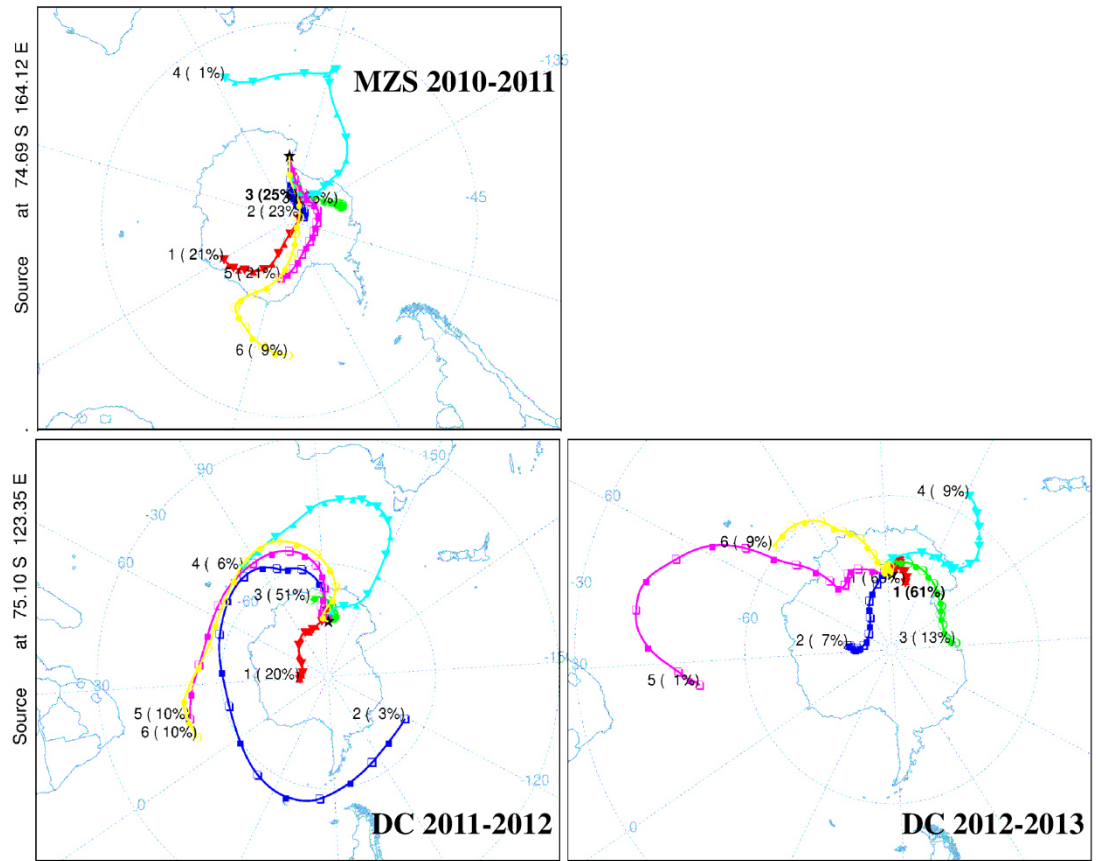
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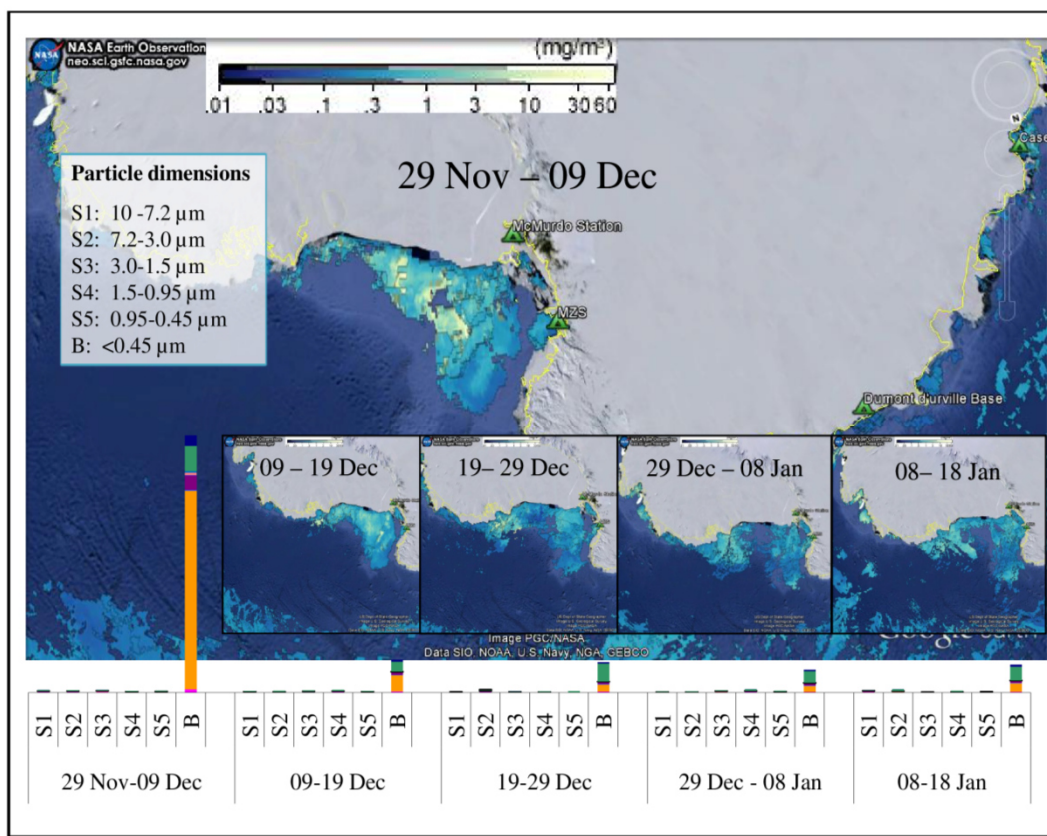


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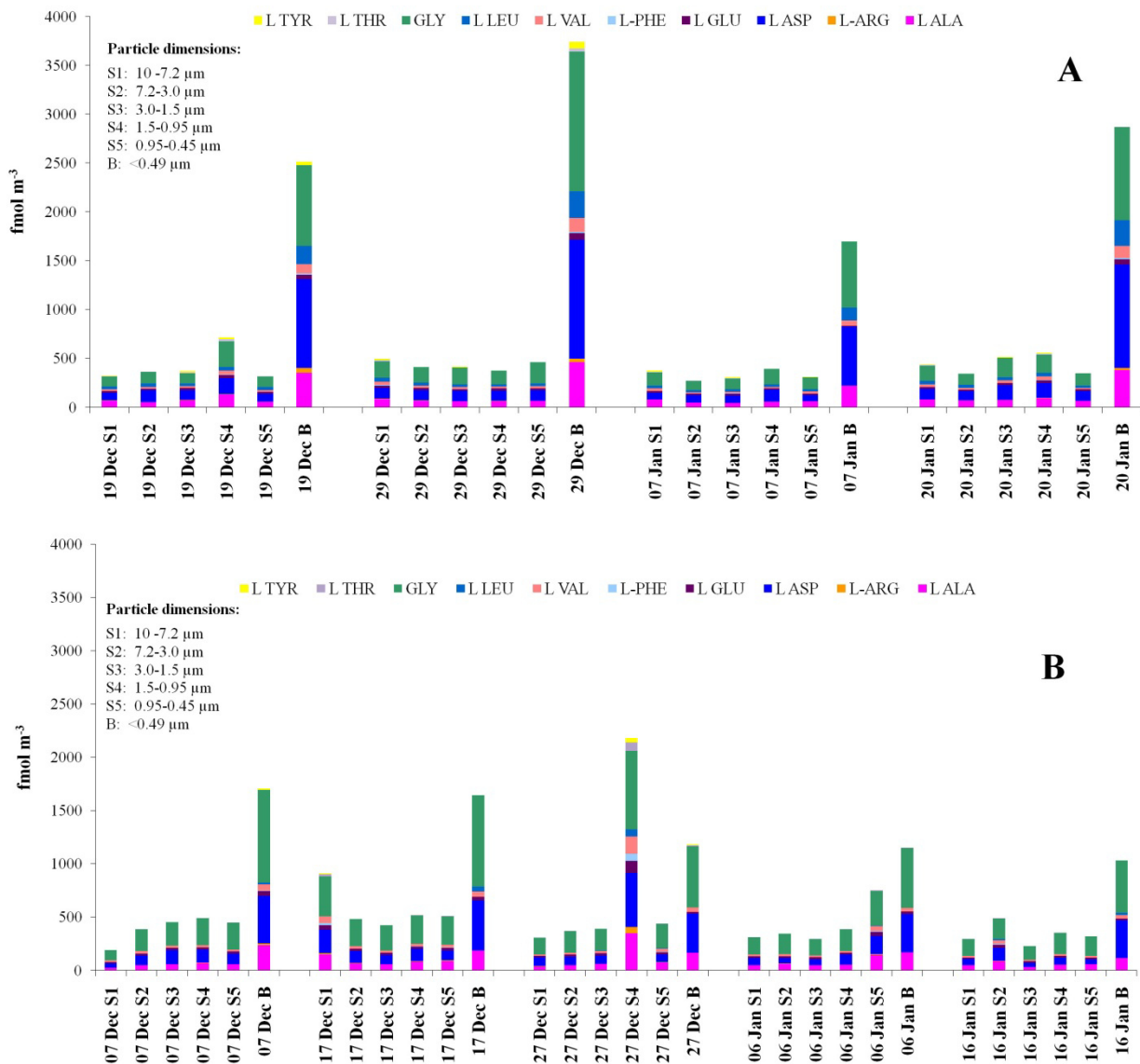
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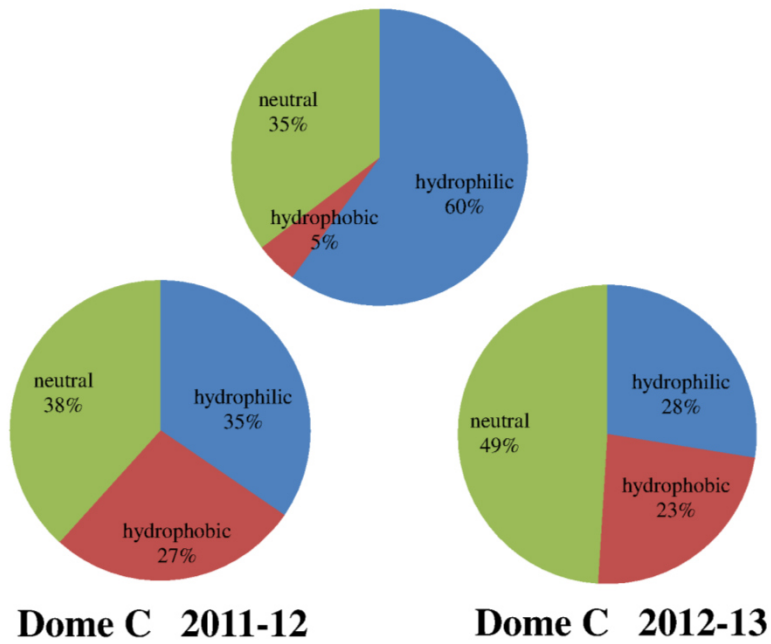
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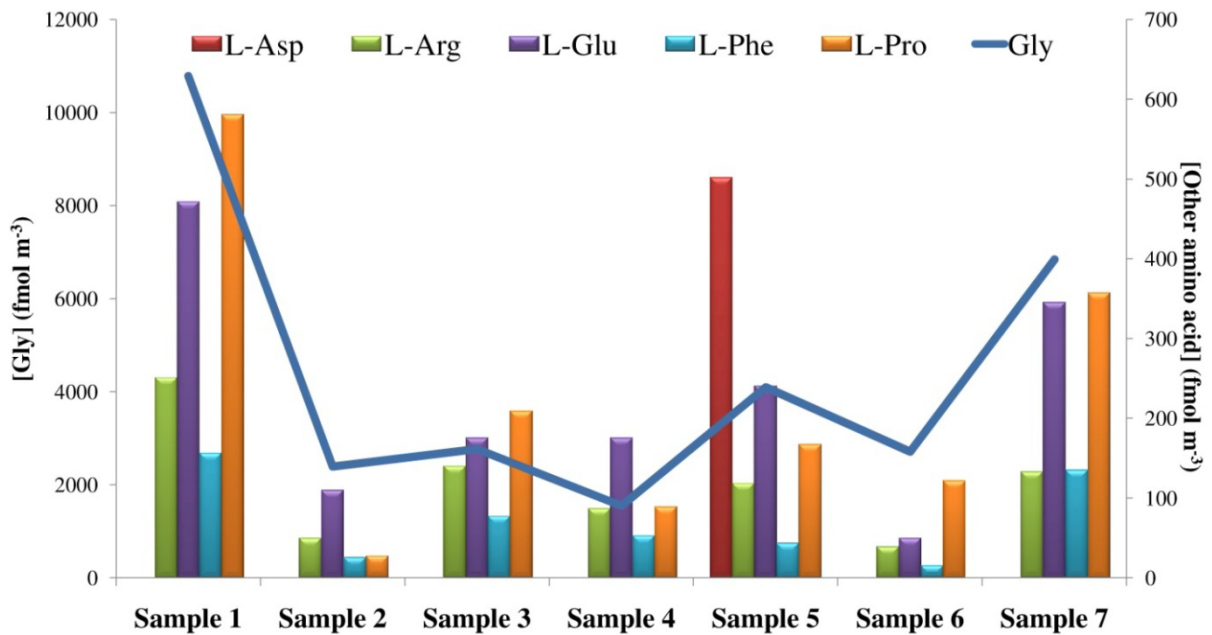
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### MZS 2010-11



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