



Atmospheric water-soluble organic nitrogen (WSON) over marine environments: a global perspective

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Abstract. To obtain a comprehensive picture of the spatial distribution of water-soluble organic nitrogen (WSON) in marine aerosols, samples were collected during research cruises in the tropical and southern Atlantic Ocean and also in the southern Indian Ocean (Amsterdam Island) for a 1-year period (2005). Samples were analyzed for both organic and inorganic forms of nitrogen, and the factors controlling their levels were examined. Fine-mode WSON was found to play a significant role in the remote marine atmosphere with enhanced biogenic activity, with concentrations of WSON ($11.3 \pm 3.3 \text{ nmol N m}^{-3}$) accounting for about 84 % of the total dissolved nitrogen (TDN). Such concentrations are similar to those observed in the polluted marine atmosphere of the eastern Mediterranean ($11.6 \pm 14.0 \text{ nmol N m}^{-3}$). Anthropogenic activities were found to be an important source of atmospheric WSON as evidenced by the levels in the Northern Hemisphere (NH) being 10 times higher than in the remote Southern Hemisphere (SH). Furthermore, the higher contribution of fine-mode WSON to TDN (51 %) in the SH, compared to the NH (13 %), underlines the important role of organic nitrogen in remote marine areas. Finally, there was a strong association of WSON with dust in coarse-mode aerosols in the NH.

1 Introduction

The ocean–atmosphere biogeochemical interactions are important in regulating atmospheric composition, marine ecosystem functioning, and the Earth's climate. Oceans are a major natural source of marine atmospheric particles (Bigg et al., 2003), with levels that depend on the meteorology, the seasonality, and the marine biodiversity of aquatic ecosystems (Bigg et al., 2003; Gantt and Meskhidze, 2013). Marine aerosols consist of salts, organic matter and living organisms originating mainly from plankton, bacteria, or other microorganisms (Monahan et al., 1983), while their organic/inorganic fraction depends strongly on the biological activity of the ocean (O'Dowd et al., 2004; Altieri et al., 2014).

Primary emissions of marine aerosols from the oceans to the atmosphere occur by bubble-bursting and wave-breaking production mechanisms and are estimated to be 5900 Tg yr^{-1} on a global scale (Bigg et al., 2003). It has been postulated that secondary aerosol formation from organic precursors emitted from the ocean, such as isoprene and monoterpenes (Yassaa et al., 2008; Arnold et al., 2009), dimethyl sulfide (Charlson et al., 1987), and aliphatic amines (Facchini et al., 2008), can significantly influence the cloud condensation nuclei (CCN) abundance and thereby the cloud albedo in the marine atmosphere (Meskhidze and Nenes, 2006).

The amount and type of organic material in seawater varies with location and time, with a large fraction remaining uncharacterized. However, an important fraction (1–3 %) of dissolved organic matter (DOM) in the surface waters comprises N-containing organic compounds (Benner, 2002). These compounds could be a potential primary source of atmospheric organic nitrogen, since peptides and free amino acids have been detected in the marine atmosphere (Kuznetsova et al., 2005; Wedyan et al., 2007). N-containing compounds in reduced forms, such as methylamines, or in oxidized forms, such as organonitrates, have been proposed as contributing to the organic matter of marine aerosols (Facchini et al., 2008; Galloway et al., 2009). During phytoplankton blooms, when biological activity is at its maximum, the organic fraction dominates in the marine aerosols mass, contributing up to 63 % of the submicron aerosol mass (O'Dowd et al., 2004).

The chemical composition of organic nitrogen in the atmosphere is complex, reflecting a wide range of direct sources, both biogenic and anthropogenic, and biological/photochemical transformations during transport. It is difficult to quantify, whatever its form (gas, particle or solution), because of its chemical and biological complexities, and no studies have yet achieved a full description of its chemical composition (Cape et al., 2011, and references therein). However, Altieri et al. (2012) found 2281 N-containing elemental formulas in marine rainwater, with the compound class, containing carbon, hydrogen, oxygen, and nitrogen (CHON+), having the largest number of formulas in the marine rainwater, as it did also in continental rainwater samples.

The importance of WSON (water-soluble organic nitrogen) in marine biogeochemical cycles and its critical role in the atmosphere has been established. However, both the origin and chemical composition of WSON in marine aerosols are largely uncertain (Kanakidou et al., 2012). The observations in marine regions are particularly scarce and limit the robust evaluation of marine organic nitrogen compared to that over continental areas. The present study provides new data for WSON, acquired during oceanographic cruises in the Atlantic Ocean as well as long-term sampling in the Indian Ocean, contributing to the global picture of WSON distribution. In the collected samples, both organic and inorganic forms of nitrogen have been analyzed and the factors controlling their levels have been examined.

2 Sample collection and chemical analysis

2.1 Sampling sites and collection

2.1.1 Tropical North Atlantic Ocean

Within the framework of the OOMPH (Organics over the Ocean Modifying Particles in both Hemispheres) project, samples were collected during a cruise with the German re-

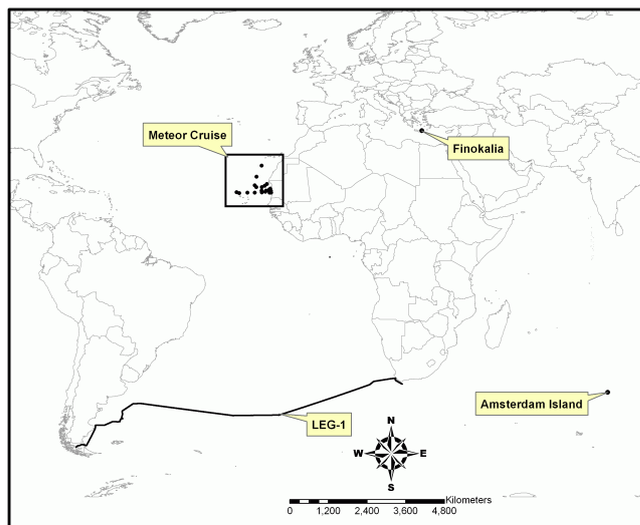


Figure 1. Sampling sites around the world during this study. Finokalia station is used for comparison purposes.

search vessel *METEOR*, which took place in the tropical North Atlantic Ocean in summer 2006 (16 July–4 August). The research cruise track, presented in Fig. 1, is characterized by frequent approaches to the African coast. During this cruise, aerosol sampling was carried out with a three-stage high-volume Sierra-type cascade impactor (Baker et al., 2007), which was operated at a flow rate of $1 \text{ m}^3 \text{ min}^{-1}$. The sampler had two impaction stages (P3 and P4 with cut-off diameters of 2.0 and $1.3 \mu\text{m}$, respectively) and one backup filter stage. During this study, fine particles were defined as the sum of P4 and the backup filter, while the P3 stage was considered to collect the coarse particles. A total of 21 daily high-volume aerosol samples were collected on quartz filters, pre-combusted at 450°C for 5 h. Samples were stored on the ship in the freezer (-20°C) until laboratory analysis. In the lab, quartz filters were extracted with 20 mL Milli-Q water for 45 min in an ultrasonic bath. Prior to analysis, the extracts were filtered through polyethersulfone (PES) membrane filters ($0.45 \mu\text{m}$ pore size diameter) to remove suspended particles. CHCl_3 was added as biocide and then the extracts were preserved at -18°C . All analyses were performed within a month after collection.

2.1.2 South Atlantic Ocean

Shipborne measurements of aerosols and trace gases were also made in the southern Atlantic Ocean as part of the same project (OOMPH). The cruise took place with the French research vessel *Marion Dufresne* during the late Southern Hemisphere summer from Cape Town (South Africa) on 19 January to Punta Arenas (Chile) until 5 February 2007 (Fig. 1). Possible local (ship)/continental contaminations were monitored by black carbon (BC) concentrations measured by an Aethalometer.

Aerosol samples were collected by a low-volume aerosol sampler placed on the prow of the research vessel. The sampler consisted of two annular glass denuders in a row, connected by a Teflon filter holder. Both denuder tubes had an overall length of 24.2 cm, a 2.2 cm inner cylinder diameter, and a 0.1 cm annulus thickness. Details on the collection efficiency of the denuder are given by Lawrence and Koutrakis (1994). The first denuder was coated with a Na_2CO_3 solution, absorbing acidic trace gases such as HNO_3 and SO_2 , and the second one was coated with a citric acid solution, absorbing NH_3 . Before use, denuders were filled with 10 mL of the coating solution, then dried using purified air, and capped with Teflon fittings.

During this cruise only fine particles were collected ($\text{PM}_{2.5}$) on Teflon filters ($n = 23$) on a 12 h basis from 21 January to 5 February 2007. Samples were stored on the ship in the refrigerator (4°C) until laboratory analysis. In the lab, the aerosol samples were extracted with 15 mL of Milli-Q water for 45 min in an ultrasonic bath and processed as the samples collected in the tropical Atlantic Ocean.

2.1.3 Amsterdam Island (Indian Ocean)

Amsterdam Island is a small island in the southern Indian Ocean ($37^\circ 48' \text{S}$, $77^\circ 34' \text{E}$), located 3400 km southeast of Madagascar and 5000 km from the coast of South Africa. Most of the year, the island experiences pristine marine conditions, especially during summer, when meteorological high-pressure conditions, accompanied by low wind speeds, prevail. For the last 20 years, atmospheric monitoring activities have been performed at this island as part of the World Meteorological Organization Global Atmospheric Watch (WMO GAW) network (Miller et al., 1993; Sciare et al., 2009).

Aerosol concentrations approaching remote marine conditions are mostly found over the oceans, especially in the Southern Hemisphere, where large expanses of open ocean and a low density of population and industry contribute to keeping human impact at a minimum. The natural aerosol over these remote ocean regions consists mainly of a mixture of sea salt particles, organics, and sulfates from the oxidation of biogenic dimethyl sulfide; some mineral dust and smoke from wildfires may also be present (Andreae, 2007).

The pristine oceanic conditions prevailing at Amsterdam Island and the very low level of local contamination from the scientific base have been documented for many atmospheric compounds such as CO, hydrocarbons, radon, black carbon, and total aerosol number concentration (Williams et al., 2001; Sciare et al., 2009).

Most atmospheric measurements were performed at 30 m above sea level and 2 km upwind of the scientific base. The aerosol samples were collected by a three-stage cascade impactor (Dekati PM_{10}) with cutoff diameters at 10, 2.5, and 1 μm , and operating at a flow rate of 30 L min^{-1} . The sampler was located on the north side of the island at

about 150 m a.s.l. A total of 42 samples were collected on Teflon filters from 29 January to 22 December 2005, with a 1-week average sampling time. In parallel, bulk aerosol samples were collected on pre-fired 47 mm diameter Whatman QMA quartz filters for BC analysis, according to the analytical protocol described in Sciare et al. (2009).

The samples were stored in the freezer and transported frozen from Amsterdam Island to the laboratory for analysis. Filter samples were extracted overnight by soft shaking of the filter portion placed in borosilicate Erlenmeyer flasks with 15 mL of ultrapure water. Prior to analysis, the extract solution was filtered through Teflon (PTFE) filters (0.2 μm pore size diameter) to remove suspended particles.

To assess possible continental influence, radon (^{222}Rn) was monitored on a 2 h basis by measuring the decrease in the alpha radioactivity of atmospheric aerosols collected on filters. The measurement has a precision of 2 % and has been described in detail by Polian et al. (1986).

2.2 Chemical analysis

2.2.1 Anions/cations and inorganic nitrogen (IN)

A Dionex AS4A-SC column with ASRS-I suppressor in auto-suppression mode of operation was used for the analysis of anions (Cl^- , Br^- , NO_3^- , SO_4^{2-} , and $\text{C}_2\text{O}_4^{2-}$). Cations (Na^+ , NH_4^+ , K^+ , Mg^{2+} , and Ca^{2+}) were analyzed by using a CS12-SC column with a CSRS-I suppressor. The reproducibility of the measurements was better than 2 % and the detection limit was 1 ppbv for the main anions and cations (1 ppbv corresponds to 0.37 ng m^{-3} for a mean air volume of 40 m^3). Mean blank values were 5–10 ppbv for Na^+ and Ca^{2+} , and lower than 3 ppbv for the rest of the ionic species. Details on the chromatographic conditions are reported in Bardouki et al. (2003). Analysis of the aerosol filters, collected at Amsterdam Island, was performed at LSCE (Laboratoire des Sciences du Climat et de l'Environnement) by ion chromatography to determine selected anions (methanesulfonate, oxalate, chloride, sulfate, nitrate, and phosphate) and cations (sodium, ammonium, potassium, magnesium, and calcium). The method is detailed in Sciare et al. (2009).

Non-sea-salt calcium and sulfate (nss-Ca^{2+} and nss-SO_4^{2-}) for tropical North Atlantic data are estimated based on the following equations:

$$[\text{nss-Ca}^{2+}] = [\text{Ca}^{2+}] - 0.3184 \times [\text{Mg}^{2+}],$$

$$[\text{nss-SO}_4^{2-}] = [\text{SO}_4^{2-}]_{\text{total}} - [\text{ss-SO}_4^{2-}],$$

$$[\text{ss-SO}_4^{2-}] = 2.0958 \times [\text{Mg}^{2+}].$$

2.2.2 Total dissolved nitrogen (TDN)

Determination of TDN is performed using the persulfate oxidation (PO) method. As such, all nitrogenous compounds are oxidized to NO_3^- under alkaline conditions at $100\text{--}110^\circ\text{C}$. The produced NO_3^- ions are then reduced by a Cu–Cd col-

umn to nitrite (NO_2^-) and subsequently diazotized with sulfanilamide and N-(1-naphthyl)-ethylenediamine, forming a high-colored azo dye. The intensity of the produced color is measured by a spectrophotometer at 543 nm (*Standard Methods for the Examination Water and Wastewater*, 1998). The detection limit of this method is 24 ppb N, while during this study the blanks were always below the detection limit (Violaki et al., 2010).

2.2.3 WSON

Water-soluble organic nitrogen (WSON) was determined by subtracting inorganic nitrogen (NO_3^- and NH_4^+) from the total dissolved nitrogen (TDN). The determination of WSON concentrations by difference leads, in some cases, to negative concentrations, as discussed by Mace and Duce (2002) and Cornell et al. (2003). Discarding these values would have biased the statistical analysis towards the samples with higher DON concentrations. For the average and median estimations, we set them as zero in agreement with previous works (e.g., Mace et al., 2003; Violaki and Mihalopoulos, 2010).

2.2.4 DMS

For the samples collected in the southern Atlantic Ocean, dimethyl sulfide (DMS) was used as a tracer of biological activity. Stainless steel canisters filled with inert gas were first evacuated to less than 1 mbar pressure using a stand-alone turbo pump. Evacuated canisters were then closed and attached to a pump and sampling line that was previously flushed with gas for more than 5 min. Canisters were then opened to the sample air, pressurized, and emptied at least five times prior to final closure and storage. Four canisters were collected per day and analyzed for dimethyl sulfide (DMS) using GC-MS (Lee et al., 1980).

2.3 Air mass back trajectory analysis

Ten-day backward air mass trajectories were calculated with the LAGRangian ANalysis TOol (LAGRANTO) (Wernli and Davies, 1997; Stohl et al., 2005). Every 3 h, about 10 air mass back trajectories were started within a ± 30 min time range at the exact ship position. For these calculations, three-dimensional wind fields from the European Centre for Medium-Range Weather Forecasts (ECMWF) were used. The 6-hourly operational global analyses (T799L91) were complemented by intermediate 3 h forecasts interpolated onto a horizontal grid with a resolution of 1° latitude/longitude. The resolution of the LAGRANTO model was 25 km in the horizontal, 91 levels in the vertical, and the computational time step for trajectory calculations was 30 min.

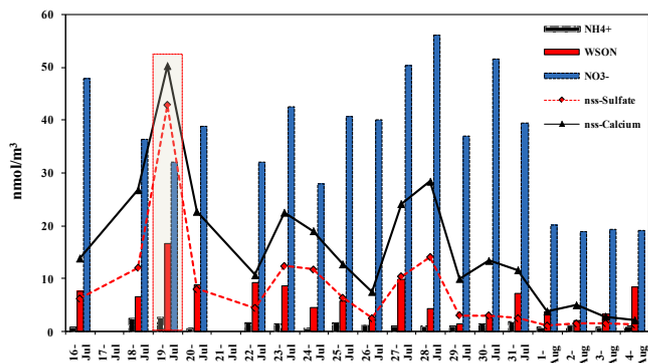


Figure 2. Temporal variation in nitrogen species and nss-SO_4^{2-} and nss-Ca^{2+} concentration in coarse particles ($D_a > 2 \mu\text{m}$) over the tropical North Atlantic atmosphere. The frame defines the air mass back trajectory with dust mixed with anthropogenic sources (19 July 2006).

3 Results and discussion

3.1 Tropical Atlantic atmosphere

During the cruise in the tropical Atlantic Ocean, the observed WSON average concentration in coarse atmospheric particles ($D_a > 2 \mu\text{m}$, $n = 18$), was $6.3 \pm 3.8 \text{ nmol N m}^{-3}$ (median $6.2 \text{ nmol N m}^{-3}$), which is in agreement with findings by Lesworth et al. (2010), who reported similar WSON concentrations ($6.1 \text{ nmol N m}^{-3}$) for coarse-mode WSON ($D_a > 1 \mu\text{m}$) in air masses originating from the Sahara. Predominance of NO_3^- was observed in coarse mode, with an average concentration of $36.1 \pm 11.6 \text{ nmol N m}^{-3}$ (median $38.0 \text{ nmol N m}^{-3}$), while the coarse-mode average concentration of NH_4^+ was $1.3 \pm 0.6 \text{ nmol N m}^{-3}$ (median $1.1 \text{ nmol N m}^{-3}$). The temporal variation in N aerosol components concentration is presented in Fig. 2. The average percentage contribution of coarse-mode nitrogen species to the TDN pool was 14, 82, and 4 % for WSON, NO_3^- , and NH_4^+ , respectively.

Almost 86 % of WSON was found in the coarse-mode atmospheric particles. Mace et al. (2003) presented the hypothesis that dust WSON might actually come mostly from adsorbed pollution, and this hypothesis has since been supported by subsequent studies (e.g., Violaki and Mihalopoulos, 2010; Zamora et al., 2011; Wang et al., 2013). No correlation was found with inorganic nitrogen (NO_3^- or NH_4^+), indicating different sources and/or their losses were at different rates (e.g., adsorption to, or chemical reaction with, larger particles). A significant correlation between WSON and nss-Ca^{2+} ($r^2 = 0.5$, $p < 0.005$, $n = 18$) confirms the continental transport of organic nitrogen by dust. WSON also correlates with nss-SO_4^{2-} ($r^2 = 0.6$, $p < 0.0005$, $n = 18$), indicating significant anthropogenic contribution to the WSON pool (see Supplement Figs. S1 and S2). During the cruise, the average concentration for coarse-

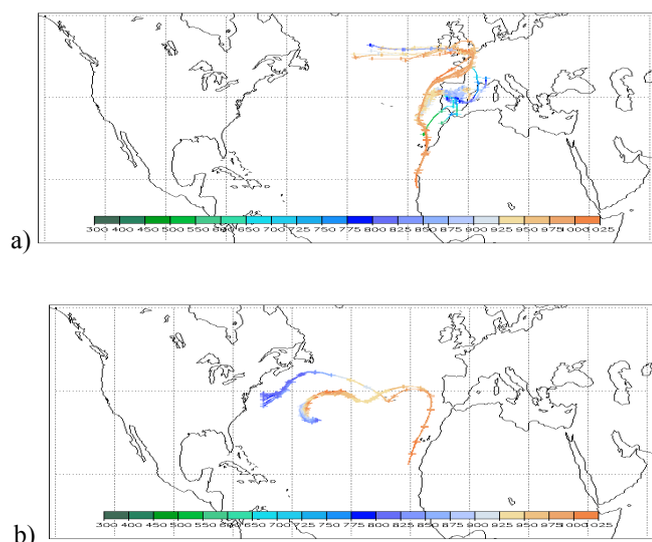


Figure 3. Five-day air mass back trajectories for 19 July (a) and 3 August 2006 (b). The color indicates the pressure level (hPa).

mode nss-Ca^{2+} and nss-SO_4^{2-} were $15.9 \pm 11.9 \text{ nmol m}^{-3}$ (median 13.0 nmol m^{-3}) and $8.1 \pm 9.7 \text{ nmol m}^{-3}$ (median 5.4 nmol m^{-3}), respectively. The highest concentrations of nss-Ca^{2+} (50.3 nmol m^{-3}), nss-SO_4^{2-} (42.8 nmol m^{-3}), and WSON (16.6 nmol m^{-3}) were observed on the same day (19 July, Fig. 2). During that day, a strong dust event took place while the ship was near the coast, resulting in air masses in which dust and anthropogenic emissions from western Europe were mixed (Fig. 3a).

The fine-mode ($D_a < 2 \mu\text{m}$, $n=18$) average concentration of WSON was found to be $0.9 \pm 1.0 \text{ nmol N m}^{-3}$ (median $0.6 \text{ nmol N m}^{-3}$), while the average concentrations of NH_4^+ and NO_3^- ions were $28.0 \pm 21.6 \text{ nmol N m}^{-3}$ (median $22.9 \text{ nmol N m}^{-3}$) and $4.7 \pm 1.8 \text{ nmol N m}^{-3}$ (median $4.5 \text{ nmol N m}^{-3}$), respectively. The temporal variation in nitrogen species is presented in Fig. 4. The average percentage contributions to the TDN pool were estimated to be 5, 19, and 76% for WSON, NO_3^- , and NH_4^+ , respectively. No statistically significant correlations were found between fine-mode WSON and continental tracers such as nss-Ca^{2+} and nss-SO_4^{2-} (see Supplement Fig. S3 and S4). The average concentrations for fine-mode nss-Ca^{2+} and nss-SO_4^{2-} were $5.9 \pm 4.3 \text{ nmol m}^{-3}$ (median 4.8 nmol m^{-3}) and $32.8 \pm 15.8 \text{ nmol m}^{-3}$ (median 28.6 nmol m^{-3}), respectively. The highest concentration of WSON was observed on 3 August (Fig. 4), when the air mass back trajectory indicated marine origin, although based on nss-Ca^{2+} levels (5.6 nmol m^{-3}), influence from continental sources (e.g., dust) cannot be totally ruled out (Fig. 3b).

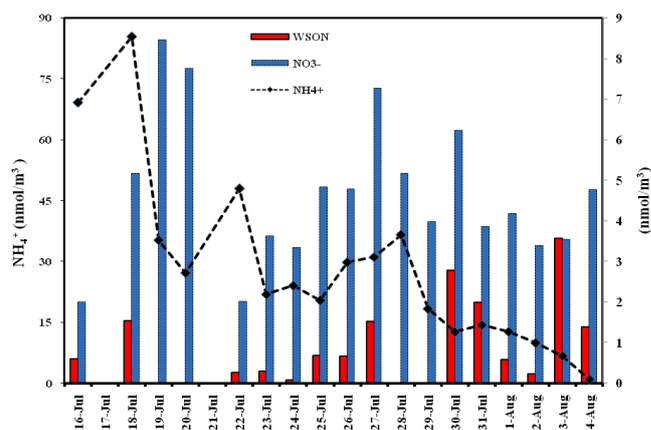


Figure 4. Temporal variation in nitrogen species concentration in fine particles ($D_a < 2 \mu\text{m}$) over the tropical North Atlantic atmosphere.

3.2 Southern Atlantic atmosphere

The cruise was performed in the temperate waters of the Atlantic Ocean during the early austral summer, when biogenic marine activity was at a maximum (Meskhidze and Nenes, 2006). Two main episodes of phytoplankton blooms were encountered during the sampling period. These are clearly seen in Fig. 5, which depicts the chlorophyll *a* map derived from SeaWiFS satellite retrievals (<http://disc.sci.gsfc.nasa.gov/giovanni>) and referred to monthly average values (January 2007). Episode A was encountered by the ship over two days (30–31 January) and episode B began on 1 February and was followed until the end of the cruise. The average concentration of WSON during the whole sampling period was estimated at $8.5 \pm 11.7 \text{ nmol N m}^{-3}$ ($n = 23$), with negligible contribution of NO_3^- ($0.2 \pm 0.5 \text{ nmol N m}^{-3}$; $n = 23$), while the NH_4^+ average concentration was estimated at $1.6 \pm 1.2 \text{ nmol N m}^{-3}$ ($n = 23$).

The highest WSON values were reported at the beginning and at the end of the cruise (Fig. 6), when the ship was near the African and the Argentinean coasts, respectively, but also when the ship crossed the second phytoplankton bloom area (episode B). Air mass back trajectories analysis indicates significant continental influence on samples collected in the beginning of episode B, which is confirmed further by the high concentration of BC ($116.9 \text{ ng C m}^{-3}$). Thus these samples are expected to result from the mixing of continental and marine sources.

Phytoplanktonic activity as a possible source of atmospheric organic nitrogenous compounds was based on DMS, since the latter could be considered an indicator of marine phytoplanktonic activity (Sciare et al., 2001). When the ship crossed the episode A area (Fig. 6), air masses had lower BC levels ($11.1 \pm 15.7 \text{ ng m}^{-3}$) compared to the average concentration observed during the sampling period ($27.2 \pm 15.7 \text{ ng m}^{-3}$). Low levels of nss-Ca^{2+}

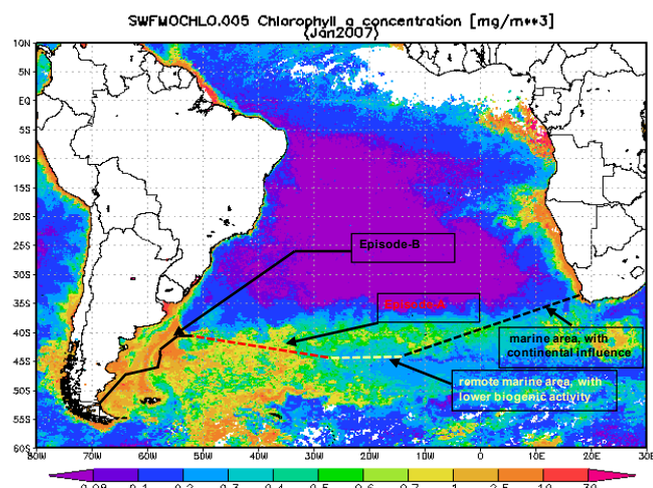


Figure 5. Two distinct phytoplankton bloom episodes are presented on a SeaWiFS chlorophyll *a* map of the South Atlantic Ocean during January 2007 along with the cruise track (<http://disc.sci.gsfc.nasa.gov/giovanni>). Episode A was encountered by the ship over 2 days (30–31 January 2007, $n = 4$) and episode B began on 1 February and was followed until the end of the cruise ($n = 5$). Remote marine conditions with low biogenic activity lasted 4 days (26–29 January 2007, $n = 5$), while marine conditions mixing with continental influence lasted 6 days (21–25 January 2007, $n = 9$).

(1.4 nmol m^{-3}) were considered a tracer of continental influence, while the measured WSON average concentration was $11.3 \pm 3.3 \text{ nmol N m}^{-3}$. These samples presented a high average contribution of WSON to TDN (84%), which could indicate an important role of marine biological activity in the biogeochemical cycle of organic nitrogen (Facchini et al., 2008). For the samples collected over the middle southern Atlantic atmosphere (26–29 January, $n = 5$), which is considered a remote marine area with low biogenic activity (Fig. 5), the average concentration of WSON was much lower ($1.3 \pm 0.8 \text{ nmol N m}^{-3}$, $n = 5$, corresponding to 51% of TDN). The important contribution of material issued from oceanic biological activity to the total ON aerosols in the marine environment has also been underlined by Miyazaki et al. (2011). During that study, average bulk ON concentrations in aerosols were measured to be 2 times higher in an oceanic region with higher biological productivity than in regions with lower productivity over the western North Pacific.

3.3 Indian Ocean (Amsterdam Island)

Amsterdam Island, located in the middle of the southern Indian Ocean, is considered a typical Southern Hemisphere marine background site, being far from major anthropogenic sources (Sciare et al., 2009). The average concentrations of WSON in coarse ($\text{PM}_{2.5-10}$) and fine ($\text{PM}_{2.5}$) particles were found to be 0.2 ± 0.4 and $0.8 \pm 1.4 \text{ nmol N m}^{-3}$, with percentage contributions to TDN of 35 and 32%, respectively. These values are in good agreement with the values reported

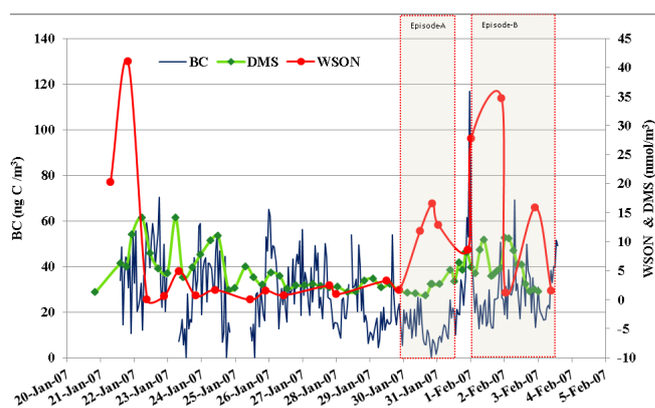


Figure 6. Temporal variation in WSON and DMS concentration in fine particles ($\text{PM}_{2.5}$) together with BC concentration during the cruise in South Atlantic atmosphere. The frames define the marine areas with intense biogenic activity (episodes A and B).

for bulk aerosols at the remote marine site of Cape Grim (average value $0.9 \text{ nmol N m}^{-3}$, $n = 2$; Mace et al., 2003b) and slightly higher compared to the values reported in Barbados ($0.5 \pm 0.3 \text{ nmol N m}^{-3}$; Zamora et al., 2011).

Concerning the inorganic nitrogen species, NH_4^+ was mainly found in fine mode, with average concentrations of 1.3 ± 1.0 and $0.3 \pm 0.1 \text{ nmol N m}^{-3}$ for fine and coarse mode, respectively, and with percentage contributions to TDN of 53 and 39%, respectively. The average concentration of NO_3^- was found to be 0.3 ± 0.2 and $0.2 \pm 0.1 \text{ nmol N m}^{-3}$ for fine and coarse mode, respectively, while the percentage contributions to TDN were 14 and 26%, respectively.

The levels of WSON, especially in fine particles (Fig. 8), have shown distinct seasonal variation with the maximum values in both modes observed during austral summer. During that season (January–March), as also confirmed by the ^{222}Rn levels, the air masses were not significantly affected by long-range transportation. As reported by Sciare et al. (2009), the meteorological regime at Amsterdam Island supports the long-range transport of air masses from southern Africa and Madagascar during winter, affecting the atmospheric composition of the area, especially with emissions from combustion processes (fossil fuel and mainly biomass burning). The simultaneous increase in average monthly concentrations of BC and radon during austral winter (Fig. 9), starting from May and maximizing in July and August (Fig. 8), confirmed the continental influence from Africa. During that season, the intense biomass burning episodes occurred in Africa, releasing fine-mode aerosols including WSON, which were subsequently transported to the marine atmosphere of Amsterdam Island (Sciare et al., 2009). At Amsterdam Island, we recorded an average BC concentration of 8.8 ng m^{-3} during the winter period and 4.1 ng m^{-3} during the summer one, which are among the lowest reported in literature. The very low levels of BC at Amsterdam Island (especially in summer) in conjunction with the very low nss-

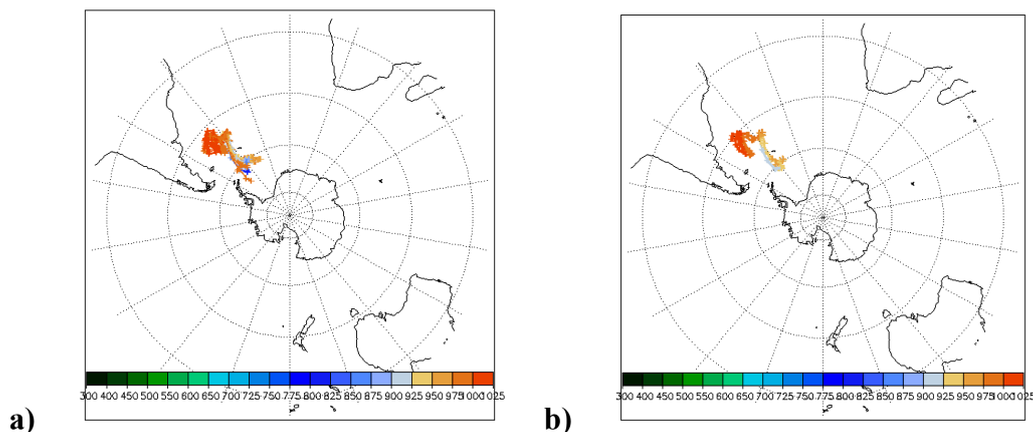


Figure 7. Five-day air mass back trajectories for 30 January 2007 (a) and 31 January 2007 (b) during episode A. The color indicates the pressure level (hPa).

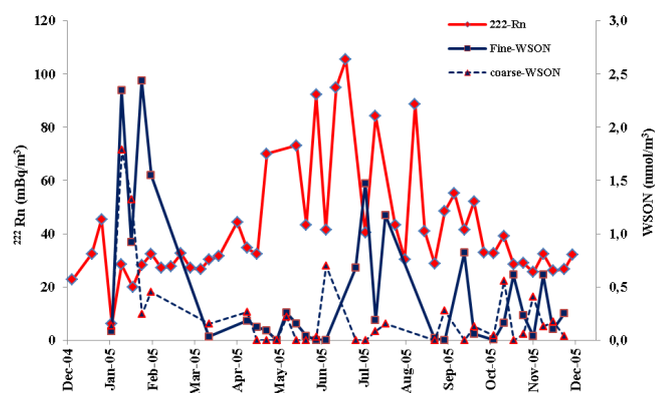


Figure 8. Seasonal variation in fine- and coarse-mode WSON with radon at Amsterdam Island. Austral summer is defined from December to April and winter from May to November.

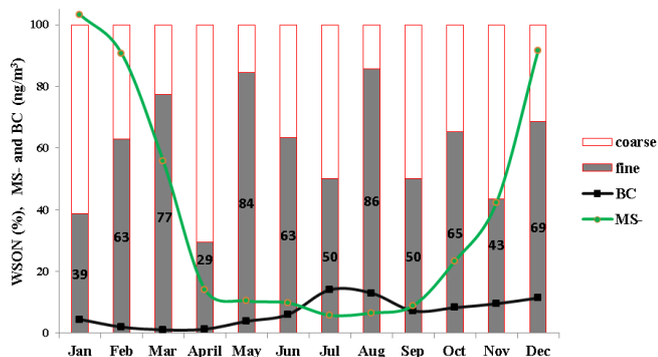


Figure 9. Average monthly concentration of bulk MS^- and BC together with the percentage contribution of WSON in fine and coarse mode at Amsterdam Island (Indian Ocean) during 2005.

Ca^{2+} values (0.3 nmol m^{-3}) highlight the very limited contribution of continental sources (combustion and/or soil) on WSON levels during that season.

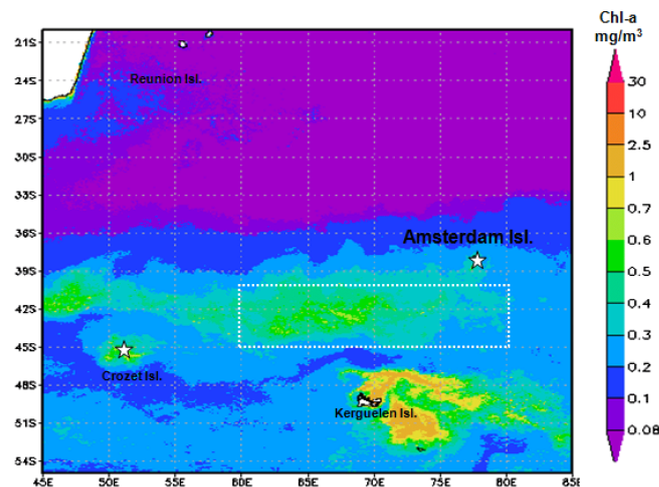


Figure 10. Chlorophyll map obtained for January (SeaWiFS data) for the Indian sector of the Austral Ocean. The white frame corresponds to the source region, which could contribute to the levels of marine organics at Amsterdam Island.

Fig. 9 depicts the percentage contribution of WSON in fine and coarse mode, together with bulk methanesulfonate (MS^-) and BC average monthly concentrations. During the austral summer, increased concentrations of MS^- were observed, which can be linked to a similar increase in marine productivity (Sciare et al., 2009). The closest DMS source is most probably located 1000–2000 km southwest of Amsterdam Island (Sciare et al., 2009), as indicated by the high Chl *a* levels seen by satellite at this location (Fig. 10).

The fine-mode WSON was found to correlate significantly with fine-mode MS^- ($r^2 = 0.7$, $p < 0.05$, $n = 7$; see Supplement Fig. S5) during austral summer, implying that part of WSON might have been produced secondarily from biogenic marine precursors. Significant correlation was also found between coarse-mode MS^- and coarse-mode WSON ($r^2 = 0.9$,

Table 1. Average concentration of WSON and percentage contribution to TDN for fine and coarse particles in both hemispheres. The discrimination between high and low biogenic activity in the marine atmosphere was based on the concentration levels of DMS; higher biogenic activity was considered in cases with a DMS concentration higher than 3 nmol m^{-3} .

Location	Sampling period	D_a (μm)	Fine WSON (nmol N m^{-3})	Northern Hemisphere		Coarse WSON (nmol N m^{-3})	% to TDN		
				% to TDN	D_a (μm)				
E. Mediterranean*	marine with anthropogenic influence	2005–2006	($n = 65$)	PM _{1.3}	11.6 ± 14.0	13	PM _{1.3–10}	5.5 ± 3.9	13
Tropical Ocean	Atlantic marine with dust influence	16 Jul–4 Aug 2006	($n = 18$)	< 2	0.9 ± 1	5	> 2	6.3 ± 3.8	14
Southern Hemisphere									
S. Atlantic Ocean	marine with high biogenic activity	30–31 Jan 2007	($n = 4$)	PM _{2.5}	11.3 ± 3.3	84	–	–	–
Middle S. Atlantic	remote marine with low biogenic activity	26–29 Jan 2007	($n = 5$)	PM _{2.5}	1.3 ± 0.8	51	–	–	–
Indian Ocean (Amsterdam Isl.)	remote marine	2005 ($n = 42$)		PM _{2.5}	0.8 ± 1.4	32	PM _{2.5–10}	0.2 ± 0.4	35

* Violaki and Mihalopoulos (2010)

$p < 0.0001$, $n = 9$; see Supplement Fig. S6) during the summer period, which could be explained by adsorption of MS^- produced from DMS on marine salts.

4 Comparison between different samplings sites and concluding remarks

This study investigates the concentrations of atmospheric WSON in the marine environment under different influences. A ship cruise was carried out in the tropical Atlantic during which a relatively polluted marine atmosphere was affected by strong dust outbreaks from the Sahara. The cruise in the Southern Hemisphere was planned to coincide with the large-scale summer phytoplankton bloom in order to investigate the role of marine biological activity in the sources of WSON compounds. Size-segregated aerosol sampling at Amsterdam Island in the Indian Ocean enabled the quantification of the WSON levels in a remote marine atmosphere.

Table 1 summarizes the average concentration of WSON and its percentage contribution to the TDN pool, separating fine from coarse atmospheric particles in both hemispheres. Note the different sample substrate and the different cutoffs used in the data presented in Table 1, which could have an impact on the comparison between the various sites. Coarse-mode WSON in the Northern Hemisphere (NH) was strongly associated with Saharan dust. This is confirmed by comparing the levels of coarse-mode WSON observed in the eastern Mediterranean and the tropical Atlantic Ocean, both strongly influenced by Sahara dust and notably with the same percentage contribution of WSON to TDN. Dust is considered an excellent adsorptive surface, not only for nitrogenous compounds but also for living biological organisms (Mathias-Maser et al., 1999; Prospero et al., 2005).

The concentrations of fine-mode WSON in the NH show large divergence with levels in the eastern Mediterranean which are more than 10 times higher (Violaki and Mihalopoulos, 2010) than in the tropical Atlantic Ocean. The atmospheric marine boundary layer of the eastern Mediterranean is influenced by air masses originating from eastern Europe, in particular Turkey, Ukraine, and Russia (Mihalopoulos et al., 1997; Lelieveld et al., 2002), which could enrich the atmosphere with WSON from anthropogenic activities. The important contribution of anthropogenic sources to the levels of WSON in the NH is revealed by concentration of WSON being more than 10 times higher compared with the remote marine areas of the Southern Hemisphere (middle southern Atlantic and Amsterdam Island). Nevertheless, the higher percentages of fine-mode WSON to TDN in both marine areas could indicate the substantial role of WSON in the biogeochemical cycle of nitrogen in such remote marine environments.

Particularly interesting are the increased levels of fine-mode WSON over the remote marine atmosphere when intense biogenic activity occurs (bloom area of the southern

Atlantic). Indeed, under such conditions, the observed levels of WSON ($11.3 \pm 3.3 \text{ nmol N m}^{-3}$) along with a high-percentage contribution (84 %) to TDN were almost identical to those measured over the eastern Mediterranean ($11.6 \pm 14.0 \text{ nmol N m}^{-3}$), pointing towards an important role of marine productivity as a source of WSON in remote marine regions with significant biogenic activity.

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