Latitudinal distributions of organic nitrogen and organic carbon in marine aerosols over the western North Pacific

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Abstract

Marine aerosol samples were collected over the western North Pacific along the latitudinal transect from 44° N to 10° N in late summer 2008 for measurements of organic nitrogen (ON) and organic carbon (OC) as well as isotopic ratios of total nitrogen (TN) and total carbon (TC). Increased concentrations of methanesulfonic acid (MSA) and diethylammonium (DEA$^+$) at 40–44° N and subtropical regions (10–20° N) together with averaged satellite chlorophyll $a$ data and 5-day back trajectories suggest a significant influence of marine biological activities on aerosols in these regions. ON exhibited increased concentrations up to 260 ngN m$^{-3}$ in these marine biologically influenced aerosols. Water-insoluble organic nitrogen (WION) was found to be the most abundant nitrogen in the aerosols, accounting for 55 ± 16% of total aerosol nitrogen. In particular, the average WION/ON ratio was as high as 0.93 ± 0.07 at 40–44° N. These results suggest that marine biological sources significantly contributed to ON, a majority of which is composed of water-insoluble fractions in the study region. Analysis of the stable carbon isotopic ratios ($\delta^{13}$C) indicated that, on average, marine-derived carbon accounted for ~88 ± 12% of total carbon in the aerosols. In addition, the $\delta^{13}$C increased from −22 to −20‰ when ON/OC ratios increased from 0.15 to 0.35 in marine biologically influenced aerosols. These results clearly show that organic nitrogen is enriched in organic aerosols originated from an oceanic region with high biological productivity, indicating a preferential transfer of nitrogen-containing organic compounds from the sea surface to the marine atmosphere. Both WION concentrations and WION/water-insoluble organic carbon (WIOC) ratios showed positive correlations with local wind speeds, suggesting that sea-to-air emissions of ON via sea spray significantly contributes to marine organic aerosols over the study region.
1 Introduction

Marine aerosol, composed of primary and secondary organic and inorganic components, contributes to the Earth’s radiative forcing and indirectly to biogeochemical cycling of carbon and nitrogen and thus influences marine ecosystems. A potential oceanic organic carbon (OC) in marine aerosol has long been recognized based on significant OC concentrations observed at oceanic sites (e.g., O’Dowd et al., 2004). Recent studies have demonstrated that the chemical composition of marine organic aerosol is the complex result of different primary and secondary sources (Facchini et al., 2008a; Hawkins and Russell, 2010; Rinaldi et al., 2010). Primary emissions of biogenic organic matter via sea spray have been suggested as potential mechanisms by which phytoplankton can modulate chemical and physical properties of marine aerosols and clouds. In previous observations of marine primary organic aerosol (POA), its chemical characterizations showed hydrophobic, polysaccharide-like material (e.g., Hawkins et al., 2010) or micro-organisms such as bacteria and diatoms (Leck and Bigg, 1999; Aller et al., 2005). Moreover, the formation mechanisms of secondary organic aerosol (SOA) in the marine atmosphere have not been fully clarified. Hence, global emission of oceanic OC aerosols is still highly uncertain.

Organic nitrogen (ON) compounds are subjected to chemical transformations in the troposphere, forming products that may potentially influence the chemical and physical properties of atmospheric aerosols (e.g., Zhang et al., 2002). Despite the importance of ON in marine biogeochemical cycles and its critical role in the atmosphere, the origins and chemical composition of ON in marine aerosols are largely unknown because of very limited field research (Duce et al., 2008). In addition, the relative importance of direct emissions and secondary formation of ON is unclear. The most identified category of ON in aerosols is reduced nitrogen (N) compounds, which include amino acids, alkyl amines, urea, and N-heterocyclic compounds that originate from biomass burning (Mace et al., 2003), animal husbandry (Schade and Crutzen, 1995), and the ocean surface (Matsumoto and Uematsu, 2005). Shi et al. (2010) reported that urea
accounted for $\sim 8\%$ of water-soluble ON on average over the eastern edge of China, where air masses were frequently affected by anthropogenic sources.

Regarding aerosol water-soluble organic nitrogen (WSON) in marine atmosphere, Facchini et al. (2008b) found high concentrations of dimethylamine and diethylamine in sub-micrometer marine aerosols in the North Atlantic, pointing to the importance of alkylamines as a biological SOA tracer. They showed that alkyl-ammonium salts represented on average 11% of the marine SOA and a dominant fraction (35% on average) of aerosol WSON in the North Atlantic. Moreover, the amine concentrations measured over the North Atlantic Ocean have shown distinct seasonal variation, indicating that the production of amine is most likely influenced by the primary productivity of phytoplankton (Facchini et al., 2008a; Müller et al., 2009). Previous studies have mostly focused on the water-soluble fractions of ON, whereas the abundances, sources, and relative importance of water-insoluble organic nitrogen (WION) compared to WSON have not been previously investigated in marine aerosols. Miyazaki et al. (2010a) found that average ON concentrations were two times greater in aerosols collected in an oceanic region with higher biological productivity than in regions with lower productivity, suggesting the importance of the total aerosol ON that is linked to oceanic biological activity.

Here we present, for the first time, latitudinal distributions of both water-soluble and water-insoluble fractions of ON in marine aerosols collected over the western North Pacific in summer when marine biological activity is high. The objective of the present study is to understand the abundance and sources of the aerosol ON (for water-soluble and water-insoluble fractions) as well as the aerosol OC associated with marine biological activity. We present the latitudinal distributions of ON and OC and their isotopic ratios to better discuss the relative importance of marine biological sources of organic aerosols.
2 Experimental

2.1 Aerosol sampling

Aerosols were sampled during the R/V Hakuho-maru cruise (KH08-2) from Kushiro (42.98° N, 144.37° E) to Tokyo (35.65° N, 139.77° E), Japan, in the western North Pacific from 44° N to 10° N along 155° E between 24 August and 13 September 2008. Figure 1 presents the cruise track, together with typical 5-day back trajectories and average concentrations of satellite-derived chlorophyll $a$ during August and September 2008 obtained from SeaWiFS satellite data. The trajectories were calculated for air masses starting from the mid-points of each sampling at altitudes of 50, 100, and 200 m. For the calculation, we used the NOAA Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (http://www.arl.noaa.gov/ready/hysplit4.html, NOAA Air Resources Laboratory, Silver Spring, MD, USA). The air masses were transported mainly from the eastern-to-central Pacific due to the Pacific high-pressure system, as shown by most of the back trajectories. Moreover, the trajectories suggest that most of the air masses were transported within the marine boundary layer. The oceanic regions are suitable to study the influences of oceanic emissions on marine aerosols.

A high-volume air sampler was located at the upper deck of the ship and used to collect ambient aerosol samples. The samplings were made using quartz fiber filters (25 × 20 cm) at a sampling flow rate of 1000 L min$^{-1}$. The total volume of the samples ranged between 800 and 990 m$^3$. The samples were collected on precombusted (450°C, 3 h) quartz filters. The sampling time for each sample was approximately 12 h, starting at 08:00 LT and 20:00 LT, which are referred to as daytime and nighttime samples, respectively. Possible contamination from ship exhaust was prevented by shutting off the sampling pump during beam-side airflow and/or low wind speeds (<5 m s$^{-1}$), resulting in an effective pumping time of about 87% of the sampling period. The samples were stored individually in glass jars with a Teflon-lined screwed cap at −20°C prior to analysis. A total of 32 samples were used for the data analysis in the present study.
2.2 Chemical analysis

A total organic carbon (TOC)/total nitrogen (TN) analyzer (Model TOC-Vcs, Shimadzu) was used to determine the total aerosol carbon and nitrogen that are dissolved in water. A filter cut of 19.63 cm$^2$ was extracted with ultra-pure Milli-Q water using an ultrasonic bath (10 min × 3 times). The total extracts were then filtrated with a disc filter (Millex-GV, 0.22 µm, Millipore, Billerica, MA, USA) followed by injection of dissolved OC and TN in the extracts into the analyzer. The sample was first injected into a combustion tube in the analyzer, which is filled with an oxidation catalyst (fiber platinum on quartz) and heated to 720°C to derive CO$_2$ and nitrogen monoxide under a constant flow of ultra-pure air. The combustion gases that passed through an electrical dehumidifier for cooling and dehydration and then through a halogen scrubber were measured by a nondispersive infrared (NDIR) detector to determine the derived CO$_2$. The measured carbon is defined here as water-soluble organic carbon (WSOC). After passing through the NDIR detector the gas stream was directed to the TN unit, which provided the TN content of the liquid sample. The nitrogen content was determined by measuring nitrogen monoxide with an ozone chemiluminescence detector. Because the instrument measures TN in liquid sample, we use the term water-soluble total nitrogen (WSTN) when referring to the concentrations measured with this instrument. Field blanks for WSOC and WSTN dissolved in water were 70 ± 8 µgC L$^{-1}$ and 40 ± 5 µgN L$^{-1}$, respectively. These values represent 24% and 28% of the average WSOC and WSTN concentrations in ambient aerosols, respectively. The data presented here were corrected for blanks. The total uncertainties of the WSOC and WSTN concentrations were estimated to be 15% and 12%, respectively.

An aliquot of filter sample was analyzed for total nitrogen (TN) and total carbon (TC) using an elemental analyzer (EA) (NA 1500, Carlo Erba) (Kawamura et al., 2004). Nitrogen and stable carbon isotopic analyses were also conducted using the same EA interfaced to an isotope ratio mass spectrometer (Finnigan MAT Delta Plus) for δ$^{15}$N and δ$^{13}$C measurements (Kawamura et al., 2004). The concentrations and isotopic...
ratios reported here were corrected against the field blanks using isotope mass balance equations (Turekian et al., 2003). The blank levels of TN and TC were 22% and 10% of the measured concentrations, respectively. The nitrogen and carbon isotope corrections were 0.9‰ and 0.5‰, respectively. Some of the samples were analyzed in duplicate. The uncertainties of TN and TC measurements were within 9% whereas those of the δ\(^{15}\)N and δ\(^{13}\)C measurements were about 0.5‰.

Mass concentrations of OC and elemental carbon (EC) were measured using a Sunset lab EC/OC analyzer. The equivalent OC concentration from field blanks accounted for ~18% of the average OC concentrations of the actual samples. OC data were all corrected against field blanks. TC values measured by the EC/OC analyzer (TC = OC + EC) agreed with those obtained by the EA within 5%, which was equivalent to or less than the uncertainties of each instrument.

Another filter cut of 1.54 cm\(^2\) was extracted with Milli-Q water. The total extracts (10 ml) were then filtrated using a membrane disc filter to determine major anions and cations as well as methanesulfonic acid (MSA) by a Metrohm ion chromatograph (Model 761 compact IC). Separation and quantification of diethylammonium ions (DEA\(^+\)) were also performed by IC (Miyazaki et al., 2010a).

The concentration of ON was determined by subtracting the inorganic nitrogen (IN) concentration from the TN concentration measured with the EA. Here IN is defined as the sum of nitrate (NO\(^{-}\)), nitrite (NO\(^{-}\))\(^{-}\), and ammonium (NH\(^{+}\))-nitrogen measured with the IC. Concentrations of WSON are defined as the difference between WSTN measured with the Shimadzu TOC/TN analyzer and IN (WSON = WSTN − IN). Similarly, WION was defined as WION = ON − WSON. Using the propagating errors of each parameter, the uncertainties of WSON and WION were estimated to be 18% and 16%, respectively.

### 2.3 Conversion efficiency of water-soluble organic nitrogen compounds

When interpreting the WSON and WION data, it is important to clarify the definitions of WSON and WION under our operating conditions of the analyzer. To characterize
these fractions of ON, we investigated the conversion efficiency of organic nitrogen compounds dissolved in water to nitrogen monoxide in the Shimadzu TOC/TN analyzer by using several authentic organic compounds. Briefly, authentic water-soluble nitrogen-containing organic species were dissolved in purified water and injected into the TOC/TN analyzer. Here we used several nitrogen-containing organic species according to differences in their functional groups or nitrogen positions in their chemical structure. Note that concentration levels of N in the experiments are set to be similar to those of WSON in the water extracts of ambient samples.

Table 1 summarizes the results. The conversion efficiencies of nitrogen in L-alanine, diethylamine, anthranilamide, Suwannee River fulvic acid to nitrogen monoxide were ~100%. Similarly, nitrogen in 4-imidazole carboxylic acid and imidazole-2-carboxaldehyde was almost completely converted to nitrogen monoxide in the analyzer. The result demonstrates that nitrogen in these species is almost completely converted to nitrogen monoxide within the measurement uncertainty under our operating conditions. By contrast, the conversion efficiencies of nitrogen in 4-pyrazole carboxylic acid and acetohydrazide were as low as 43 ± 14% and 11 ± 2%, respectively. Although the exact reason for the low conversion efficiencies for these compounds is unclear, both of these compounds have nitrogen atoms in adjacent positions in their molecular structure, which may have related to the lower conversion efficiency to nitrogen monoxide.

Note that conversion efficiencies of carbon in all of these compounds to CO$_2$ were ~100% in the TOC analyzer. In addition, it was also confirmed that nitrogen and carbon in these authentic organic species were converted to N$_2$ and CO$_2$ by ~100% by the EA, respectively. In summary, the laboratory experiments indicate that some classes of dissolved organic nitrogen compounds may not be quantitatively converted to nitrogen monoxide under our operating conditions of this instrument. This may lead to an uncertainty in interpreting the classification of WSON and WION in ambient aerosols.
2.4 Air mass exposure to chlorophyll $a$ along with back trajectory

The trajectory positions were calculated every 1 h along the 5-day back trajectory each. At each position along the trajectory, concentrations of chlorophyll $a$ in the ocean were taken from monthly mean global distributions (August–September 2008) obtained from SeaWiFS observations. The SeaWiFS data used for the present analysis had horizontal resolution of 0.2 degrees ($\sim$25 km). The average concentrations of chlorophyll $a$ exposure were then calculated along each trajectory to produce a time series of average air mass exposure to chlorophyll $a$ along the cruise track. The trajectories were started at two altitude levels (50 and 100 m), and chlorophyll $a$ values were averaged for each sampling point. It is noted that the chlorophyll $a$ concentrations at each grid may vary on shorter time scales. Moreover, the trajectories do not account for subgrid-scale wind and transport processes that may be active within the marine boundary layer.

3 Results and discussion

3.1 Air-mass characteristics

Figure 2 shows latitudinal distributions of the concentrations of MSA, nss-SO$_4^{2-}$, and NO$_3^-$, Five-day back-trajectory-weighted SeaWiFS chlorophyll $a$ is also shown in the figure. MSA is produced by the atmospheric oxidation of dimethylsulfide (DMS), which is released as a gas from marine microbial processes and thus can be used as an indicator of secondary aerosols of marine biogenic origin. MSA showed a distinct latitudinal gradient, with concentrations increasing at 40–44$^\circ$N. In this oceanic region, the back-trajectory-weighted chlorophyll $a$ showed a similar spatial trend to that of MSA. Back trajectories suggest that air masses sampled at 40–44$^\circ$N were transported within the marine boundary layer and frequently encountered oceanic regions with high productivity in a northeast upwind sector of the sampling locations (Fig. 1). Detailed analysis of the trajectory-weighted chlorophyll $a$ showed that aerosol samples collected
at 40–44° N must have passed over a higher chlorophyll a region approximately 12-36 h before reaching the ship. In addition, in-situ measured chlorophyll a in sea water showed substantially larger concentrations at 40–44° N (av. 0.56 mg m^{-3}) compared to those at 10–40° N (0.05–0.15 mg m^{-3}) (Ooki et al., 2010). These results suggest that the aerosols sampled at 40–44° N were largely influenced by oceanic emissions from both upwind regions with high productivity and local emissions from marine biological sources.

The nss-SO_{4}^{2-} concentrations also showed distinct peaks at 40–44° N and 10–15° N. Indeed, Nagao et al. (1999) suggested that most nss-SO_{4}^{2-} transported from northeastern and southeastern sectors to Hahajima (142.10° E, 26.38° N) originated from DMS oxidation, based on their long-term measurements of nss-SO_{4}^{2-} and MSA. Their results support significant influences of marine biological sources on aerosols in this oceanic region. At 10–15° N, nss-SO_{4}^{2-} exhibited increased concentrations up to ~8000 ng m^{-3}. Mochida et al. (2010) reported that the plume from the eruption of Kasatochi Volcano, Aleutian Islands (52.17° N, 175.51° W), on 7 August 2008 (Martinsson et al., 2008; Rix et al., 2009) may have been associated with the enhanced nss-SO_{4}^{2-} concentration (Uematsu et al., unpublished data). The trajectories suggest that the air masses are representative of remote marine aerosols (Fig. 1).

It should be noted that the background marine aerosol composition can be influenced by aged ship emissions (e.g., Capaldo et al., 1999) as well as other anthropogenic emissions. However, an insignificant contribution of anthropogenic sources to the observed air masses is suggested by the low average concentrations of NO_{3}^{-} (116 ± 84 ng m^{-3}), which are similar to those reported for clean marine aerosols over the open ocean (<180 ng m^{-3}) (Rinaldi et al., 2009). Moreover, EC concentrations ranged from below the detection limit (<2 ngC m^{-3}) to 10 ngC m^{-3}, which is also similar to the range reported for clean marine aerosols at Mace Head in the North Atlantic (Cooke et al., 1997). The low concentrations of these anthropogenic tracers together with the back trajectories suggest that the aerosols sampled during this study were little affected by anthropogenic sources including ship emissions.
3.2 Temporal and spatial variations of organic nitrogen aerosols

Figure 3 presents time series of mass concentrations of aerosol nitrogen species/components together with MSA. Latitudinal distributions of the ON, WSON, WION, and DEA⁺ concentrations are shown in Fig. 4. During the study period, the ON concentrations ranged between 10 and 260 ngN m⁻³, with an average of 130 ± 61 ngN m⁻³. This average concentration is similar to that in PM₂.₅ aerosols (160 ngN m⁻³) observed at a forest site in the US (Lin et al., 2010). The average fraction of ON to TN was as large as 67 ± 15%. Interestingly, WION (av. 112 ± 61 ngN m⁻³) was the most abundant N in the aerosols, accounting for 55 ± 16% of TN on average. In particular, WION exhibited a maximum concentration of 260 ngN m⁻³ at 40–44° N (Fig. 4b), where marine biological influences on the observed aerosols were large. In this oceanic region, the average WION/ON ratio was 0.93 ± 0.07, which is higher than that (0.74 ± 0.12) at 10–40° N. In comparison to WION, WSON showed a smaller fraction, representing on average 12 ± 10% of TN. The fraction of WSON to TN is similar to previous findings, generally ranging from ~10:30% (Zhang et al., 2002; Mace et al., 2003; Nakamura et al., 2006). Overall NH₄⁺ and NO₃⁻ accounted for 14 ± 13% and 17 ± 11% of TN, respectively.

In general, the temporal trend of nitrogen concentrations tracked those of MSA. TN and MSA showed a linear relation with \( r^2 = 0.74 \), indicating a large influence of marine biological emission on aerosol N in the study region. In fact, Miyazaki et al. (2010a) found a bimodal size distribution of ON with two peaks at sub-micrometer and supermicrometer modes in marine aerosols over the western North Pacific in summer, where the average ON concentrations were twice more abundant in marine biologically more-influenced aerosols than in less-influenced aerosols. It should be noted that the daytime and nighttime concentrations of nitrogen species showed no distinct difference (Fig. 3). Specifically, the average ON concentration in daytime was 130 ± 59 ngN m⁻³, which is similar to that in nighttime (128 ± 66 ngN m⁻³). This result suggests that local photochemistry had an insignificant effect on the temporal variations of TN in the aerosols sampled in the present study.
As a marine biogenic organic compound of WSON, DEA$^+$ has been detected in marine aerosols (Facchini et al., 2008b; Müller et al., 2009; Sorooshian et al., 2009; Miyazaki et al., 2010a). The concentrations of aliphatic amines including DEA$^+$ measured over the northern Atlantic showed a seasonal variation, indicating that the production of amine is most likely influenced by the primary biological productivity of phytoplankton and therefore available nutrient in the ocean (Facchini et al., 2008b). Here we detected DEA$^+$ in aerosols, with concentrations ranging from a value below the detection limit ($<0.1$ ng m$^{-3}$) to 0.8 ng m$^{-3}$. The average DEA$^+$ concentration at 40–44° N ($0.52 \pm 0.19$ ng m$^{-3}$) was twice larger than that at 10–40° N ($0.23 \pm 0.11$ ng m$^{-3}$) (Table 2). The contribution of DEA$^+$ to WSON ranged between 0.1 and 9.8% with an average of 0.8%. At 40–44° N, both the DEA$^+$ concentrations and DEA$^+$/WSON ratios showed higher values compared with those at the lower latitude (10–40° N), being consistent with the large influence of marine biological activity on the observed organic aerosols.

Facchini et al. (2008b) suggested that DEA$^+$ is derived from an oceanic biological source and is produced by the reaction of gaseous amines with sulfuric acid or acidic sulfates. They showed that the size distributions of DEA$^+$ in marine aerosols exhibited maxima in the accumulation mode, as is also the case for SO$_4^{2-}$, MSA, and NH$_4^+$, indicating that a gas-to-particle conversion process is responsible for accumulation of ammonium salts in the fine aerosol fractions. Indeed, DEA$^+$ was detected in the submicrometer size range of biologically more-influenced aerosols obtained in a similar oceanic region in our previous study (Miyazaki et al., 2010a). Although the size distributions of DEA$^+$ and SO$_4^{2-}$ are not available in the present study, a positive correlation between DEA$^+$ and SO$_4^{2-}$ ($r^2 = 0.52$) was found in our samples, supporting the secondary formation processes of DEA$^+$ as discussed above.

Besides amine, WSON may also consist of other biologically labile components, such as amino acids (e.g., Simoneit et al., 2004) and more refractory components (e.g., humic compounds) (Cornell et al., 2003). Matsumoto and Uematsu (2005) reported that dissolved free amino acids in marine aerosols, which mostly reside in the
fine mode (aerodynamic diameter less than 2.5 µm), accounted for only <0.1% of inorganic nitrogen over the western North Pacific. Shi et al. (2010) also reported a minor contribution of free amino acids to WSON (~1%) in aerosols over the China Sea. Although detailed chemical analyses of WSON were not performed in the present study, previous studies have indicated that combined amino acids may account for a major portion of the amino compounds in aerosols (e.g., Zhang and Anastasio, 2003).

3.3 Latitudinal distributions of organic carbon aerosols

Figure 5 shows the latitudinal distributions of OC, WSOC, and WIOC. Similar to the latitudinal profiles of MSA and ON, OC and WSOC exhibited increased concentrations at 40–44°N. In the region of 40–44°N, an enhanced fraction of WSOC to OC was observed, ranging between 0.42 and 0.66 with an average of 0.53 ± 0.10. The range is similar to that (0.46–0.55) found for remote marine aerosols collected over the western North Pacific in a region farther north than the present study region (Miyazaki et al., 2010b). The WSOC and MSA concentrations showed a positive correlation in the present study ($r^2 = 0.57$), suggesting an importance of secondary production of WSOC originated from marine biological sources, similar to the case of MSA. In our previous study, WSOC and oxalic acid also showed significant correlation with MSA in the sub-micrometer size range of marine biologically influenced aerosols (Miyazaki et al., 2010b), which is in agreement with the results presented here. On the other hand, Russell et al. (2010) recently observed that the ocean-derived composition in sub-micrometer marine aerosol was dominated by carbohydrate-like material containing organic hydroxyl groups (i.e., water-soluble organics) over the North Atlantic and Arctic oceans. In our samples, however, WSOC did not show any significant correlation with Na$^+$ ($r^2 = 0.02$) or local wind speeds, suggesting that a majority of WSOC is derived from secondary production. This point will be discussed later.

The latitudinal distribution pattern of the WIOC concentration is not apparent compared to that of WSOC. The average concentration of WIOC at 40–44°N (560 ± 193 ngC m$^{-3}$) is similar to that of WSOC (645 ± 273 ngC m$^{-3}$) in the same
region (Table 2). On the other hand, the WIOC concentration at 10–40° N (av. 660 ± 274 ngC m⁻³) is similar to that at 40–44° N, and is substantially larger than the WSOC concentrations (av. 100 ± 69 ngC m⁻³). Overall, the concentrations of WIOC are larger by a factor of three than those observed in the Austral Ocean in summer when marine biological activity is high (Sciare et al., 2009). WIOC is mechanically produced through bubble-bursting processes from hydrophobic organic matter that accumulates in the micro layer of the ocean surface (Blanchard, 1964; Oppo et al., 1999). Indeed, WIOC in our samples showed a linear relation with Na⁺ (r² = 0.37), suggesting that a bubble-bursting production mechanism could explain the major fraction of WIOC in the aerosols collected. It is possible that chemical aging of water-insoluble organics (e.g., oxidation of unsaturated fatty acids, and olefins to result in water-soluble organic species) primarily emitted from the ocean surface and the subsequent production of more water-soluble organics may occur in aerosols largely influenced by marine biological productivity. This could partly explain the larger fractions of WSOC in aerosols collected at 40–44° N, which may have been transported over a higher chlorophyll a region within 12–36 h as discussed in Sect. 3.1.

3.4 Nitrogen-to-carbon ratios in marine aerosols

The relative contribution of organic nitrogen species to OC in aerosol mass provides insight into the origin and chemical properties of aerosol ON. The average ON/OC ratio was 0.19 ± 0.11 during the study period, which is comparable to but higher than typical ON/OC ratios (0.06–0.11) reported for oceanic dissolved materials (Hansell and Carlson, 2002). Miyazaki et al. (2010a) reported higher ON/OC ratios (0.35–0.49) in size-segregated marine aerosols obtained at farther north of our present study areas in the western Pacific. The ON/OC ratio in the present study is also higher than that for PM₂.₅ aerosols (0.04–0.09) observed at a boreal forest site (Lin et al., 2010). The average WSON/WSOC ratio was 0.23 ± 0.21, whereas the WION/WIOC ratio was 0.18 ± 0.15 in our aerosol samples. These results indicate more enrichment of organic nitrogen in marine aerosol collected in this study compared to previous studies.
Furthermore, we analyzed nitrogen isotopic ratios of TN to investigate the contributions of marine sources to the aerosol N sampled here. The $\delta^{15}N$ values of TN ranged from $-2.2\%$ to $8.9\%$ with an average of $4.9\pm2.8\%$. Although nitrogen isotopic analysis alone cannot provide conclusive evidence about atmospheric ON sources (Kelly et al., 2005), these $\delta^{15}N$ values overlap with a wide range of marine phytoplankton ($+3$ to $+12\%$) (Owens, 1987). The values are also similar to those ($1.2$–$4.6\%$) in aerosols collected in the above-mentioned oceanic region with higher biological productivity in the western North Pacific (Miyazaki et al., 2010a).

To better understand the relative contributions of marine sources to the sampled aerosols, we also analyzed stable carbon isotopic ratios. In low and mid-latitudes, $\delta^{13}C$ of OC in seawater typically ranges from $-20$ to $-22\%$ (Turekian et al., 2003), whereas the average $\delta^{13}C$ of terrestrial OC sources, such as fossil fuel combustion and biomass burning, is $-26.5\pm1\%$ (e.g., Cachier et al., 1986). The $\delta^{13}C$ values of $-22.1$ to $-20.8\%$ obtained in the present study (Table 2) are similar to those of OC in seawater. To roughly calculate the relative contribution from marine and terrestrial OC sources, we applied $\delta^{13}C$ values ranging from $-22\%$ to $-21\%$ for marine OC and those ranging from $-27\%$ to $-26\%$ for terrestrial OC following the isotopic mass balance equations given in previous studies (e.g., Turekian et al., 2003). Our calculation indicates that on average, marine sources contribute $88\% \pm 12\%$ of the aerosol carbon. The result supports that a majority of TC in the aerosols is derived from marine sources. Figure 6 shows a correlation between $\delta^{13}C$ values and ON/OC ratios classified according to the MSA concentrations. Interestingly, the $\delta^{13}C$ showed higher values (isotopically heavier) with increasing ON/OC ratios in air masses with MSA $>20$ ng m$^{-3}$. The result clearly shows an enrichment of nitrogen in organic aerosols in the oceanic region with high biological productivity and indicates a preferential transfer of nitrogen-containing compounds from the sea surface to marine atmosphere.

In general, no clear relation was found between WSOC and WSON ($r^2 = 0.04$), which suggests differences in marine biological sources and chemical properties for these two organic fractions. As discussed in the previous section, WSOC and MSA
concentrations showed positive correlation ($r^2 = 0.57$), suggesting secondary production of organic aerosols from marine biological origin. In contrast, WSON exhibited poor correlation with MSA ($r^2 = 0.05$). This result suggests that the sources and formation processes for the majority of WSON are different from those for MSA (i.e., DMS).

### 3.5 Possible formation processes of WION from marine biological sources

In this section, we discuss possible processes of WION in marine aerosols in terms of primary and secondary production. Figure 7 illustrates the WION concentrations and WION/WIOC ratios as a function of local wind speed. The WION concentrations showed a positive correlation with local wind speed, particularly at 10–40$^\circ$ N. In contrast, no relation was found between WSON and wind speed ($r^2 = 0.02$, data not shown as a figure). These results clearly suggest that significant fractions of WION were mechanically produced through bubble bursting processes as a potential mechanism. Moreover, WION/WIOC ratios tend to increase with increasing local wind speed (Fig. 7b), suggesting that nitrogen containing organic species contributes significantly to marine organic aerosols that are likely derived from bubble-bursting processes over the western Pacific region.

WION may be associated with organic particles of seawater origin, which include microorganisms, plankton debris, and inorganic particles containing adsorbed organic matter. They may also be composed of a large number of semi-transparent gel-like particles and proteins, all of which contain amino acids. Kuznetsova et al. (2005) used Coomassie blue dye to confirm that some of the colloidal gel-like material surrounding bacteria and viruses was proteinaceous in Mediterranean and Atlantic marine aerosol samples. More recently, Hawkins and Russell (2010) reported chemical signatures of proteins and calcareous phytoplankton as well as polysaccharides in hydrophobic marine aerosols collected in the Arctic and southeastern Pacific using Near-Edge Absorption X-ray Fine Structure (NEXAFS) spectromicroscopy. The present finding that the fraction of WION is large in marine biologically influenced aerosols is supported by these previous studies.
At 40–44° N, the relation between WION and local wind speeds, although the values of these parameters are larger, is less evident compared to that at 10–40° N (Fig. 7). On the basis of laboratory experiments on nascent sea spray aerosols, Facchini et al. (2008a) reported that sea spray organics tend to aggregate and form colloids or suspended particles, which makes the definition of water solubility a complex issue. Considering the significance of secondary production at 40–44° N as discussed above, secondary formation of ON associated with marine emissions of gaseous precursors (e.g., biogenic volatile organic compounds, DMS, and NH₃) is another possibility that could partly explain the increase of WION. As one possible process for secondary formation of nitrogen-containing organic aerosols, Galloway et al. (2009) recently reported that glyoxal uptake onto ammonium sulfate seed aerosol can irreversibly form C–N compounds. In fact, glyoxal and methylglyoxal were detected in aerosols from marine biological origin in the similar oceanic region in our previous studies (Miyazaki et al., 2010b). Fig. 8 illustrates the linear relation between WION and NH₄⁺, color-coded according to the DEA⁺ concentration. The WION concentration was positively correlated with NH₄⁺, particularly under the large influence of marine biological activity (i.e., DEA⁺ > 0.3 ng m⁻³). The positive correlation indicates that the possible formation of WION is related to ammonium salts in marine biologically influenced aerosol.

The substantial amount of WION in marine biologically influenced aerosols found in the present study may provide insights into chemical composition of WIOC, which is still mostly uncharacterized with the exception of fatty acids (Mochida et al., 2002), n-alkanes and fatty alcohols (Kawamura et al., 2003). Chemical characterizations of WION as well as WSON in aerosols from marine biological sources at a molecular level and the mechanisms of primary and secondary formation of WION remain elusive and require further study.
4 Conclusions

Latitudinal distributions of organic nitrogen (ON) and organic carbon (OC) as well as isotopic ratios of total nitrogen (TN) and total carbon (TC) were measured in marine aerosols collected in the western North Pacific in summer 2008. Increased concentrations of methanesulfonic acid (MSA) and diethylammonium (DEA\(^{+}\)) at 40–44° N and subtropical regions (10–20° N) together with averaged satellite chlorophyll \(a\) data and 5-day back trajectories suggest significant influences of marine biological activities on aerosols in these regions. ON exhibited increased concentrations (up to 260 ngN m\(^{-3}\)) in the marine biologically influenced aerosols. Water-insoluble organic nitrogen (WION) was found to be the most abundant N in the marine aerosols, accounting for 55 ± 16% of total aerosol N on average. In particular, the average WION/ON ratio was as high as 0.93 ± 0.07 at 40–44° N. Overall WSON accounted for ∼12% of total N. These results suggest that marine biological sources significantly contributed to ON, a majority of which is composed of water-insoluble fractions in the study region.

The stable carbon isotopic analysis indicated that, on average, marine-derived carbon accounted for ∼88 ± 12% of total carbon in the aerosols. Moreover, the stable carbon isotopic ratios showed higher values (from −22 to −20‰) when ON/OC ratios increased from 0.15 to 0.35 in marine biologically influenced aerosols. The results clearly indicate an enrichment of nitrogen in organic aerosols originated from an oceanic region with high biological productivity and preferential transfer of nitrogen-containing organic compounds from the sea surface to the marine atmosphere. Furthermore, both WION concentrations and WION/WIOC ratios exhibited positive correlations with local wind speeds, suggesting that sea-to-air emissions of ON via sea spray contributes significantly to marine organic aerosols over the study region.

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Japan. This research was also supported by the Environment Research and Technology Development Fund (B-0903) of the Ministry of the Environment, Japan.

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Zhang, Q. and Anastasio, C.: Free and combined amino compounds in atmospheric fine particles (PM2.5) and fog waters from Northern California, Atmos. Environ., 37, 2247–2258, 2003.
Table 1. Conversion efficiencies of selected water-soluble nitrogen-containing organic compounds to nitrogen monoxide during the measurements by the Shimadzu TOC/TN analyzer. Note that conversion efficiencies of carbon in all of these compounds to CO\textsubscript{2} were \~100\%.

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Molecular structure</th>
<th>N concentrations\textsuperscript{a} (μgN L\textsuperscript{-1})</th>
<th>Conversion efficiency of N (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>L-Alanine</td>
<td><img src="image" alt="L-Alanine structure" /></td>
<td>1120</td>
<td>101 ± 2</td>
</tr>
<tr>
<td>Diethylamine</td>
<td><img src="image" alt="Diethylamine structure" /></td>
<td>1345</td>
<td>100 ± 2</td>
</tr>
<tr>
<td>Anthranilamide</td>
<td><img src="image" alt="Anthranilamide structure" /></td>
<td>1480</td>
<td>100 ± 3</td>
</tr>
<tr>
<td>Suwannee River fulvic acid\textsuperscript{b}</td>
<td><img src="image" alt="Suwannee River fulvic acid structure" /></td>
<td>730\textsuperscript{c}</td>
<td>95 ± 6</td>
</tr>
<tr>
<td>4-Imidazole carboxylic acid</td>
<td><img src="image" alt="4-Imidazole carboxylic acid structure" /></td>
<td>1000</td>
<td>99 ± 5</td>
</tr>
<tr>
<td>Imidazole-2-carboxaldehyde</td>
<td><img src="image" alt="Imidazole-2-carboxaldehyde structure" /></td>
<td>1160</td>
<td>101 ± 2</td>
</tr>
<tr>
<td>4-Pyrazole carboxylic acid</td>
<td><img src="image" alt="4-Pyrazole carboxylic acid structure" /></td>
<td>1000, 1500</td>
<td>43 ± 14</td>
</tr>
<tr>
<td>Acetoxydrazide</td>
<td><img src="image" alt="Acetoxydrazide structure" /></td>
<td>1295, 1660</td>
<td>11 ± 2</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Concentration levels of N are set to be similar to those of WSON in the water extracts of ambient samples.

\textsuperscript{b} Suwannee River fulvic acid are produced by the International Humic Substance Society (IHSS).

\textsuperscript{c} Concentrations of N is based on sample information for elemental analysis provided by the IHSS.
Table 2. Average concentrations and ratios in marine aerosols sampled in the different latitudinal regions.

<table>
<thead>
<tr>
<th></th>
<th>10–40° N</th>
<th>40–44° N</th>
</tr>
</thead>
<tbody>
<tr>
<td>MSA (ng m⁻³)</td>
<td>24 ± 16</td>
<td>129 ± 37</td>
</tr>
<tr>
<td>nssSO₄²⁻ (ng m⁻³)</td>
<td>3060 ± 2730</td>
<td>6040 ± 3400</td>
</tr>
<tr>
<td>DEA⁺ (ng m⁻³)</td>
<td>0.23 ± 0.11</td>
<td>0.52 ± 0.19</td>
</tr>
<tr>
<td>WION (ngN m⁻³)</td>
<td>91 ± 44</td>
<td>200 ± 38</td>
</tr>
<tr>
<td>WSON (ngN m⁻³)</td>
<td>16 ± 13</td>
<td>19 ± 16</td>
</tr>
<tr>
<td>WIOC (ngC m⁻³)</td>
<td>660 ± 274</td>
<td>560 ± 193</td>
</tr>
<tr>
<td>WSOC (ngC m⁻³)</td>
<td>100 ± 69</td>
<td>645 ± 273</td>
</tr>
<tr>
<td>ON/OC</td>
<td>0.15 ± 0.10</td>
<td>0.20 ± 0.09</td>
</tr>
<tr>
<td>NO₃⁻ (ng m⁻³)</td>
<td>110 ± 90</td>
<td>63 ± 62</td>
</tr>
<tr>
<td>NH₄⁺ (ng m⁻³)</td>
<td>22 ± 19</td>
<td>128 ± 40</td>
</tr>
<tr>
<td>δ¹⁵N (‰)</td>
<td>3.2 ± 5.5</td>
<td>5.1 ± 1.6</td>
</tr>
<tr>
<td>δ¹³C (‰)</td>
<td>-20.8 ± 4.5</td>
<td>-22.1 ± 1.3</td>
</tr>
</tbody>
</table>
Fig. 1. *R/V Hakuho* cruise track in the western North Pacific between 24 August and 13 September 2008, together with typical 5-day back trajectories. Also shown are monthly averaged concentrations of chlorophyll *a* for August–September 2008 derived from SeaWiFS data, available from the NASA Goddard Space Flight Center/Distributed Active Archive Centers. A white dot indicates that the satellite data is missing at a corresponding grid.
Fig. 2. Latitudinal distributions of MSA, nss-SO$_4^{2-}$, and NO$_3^-$.

Also shown are 5-day back trajectory-weighted SeaWiFS chlorophyll $a$ concentrations as square symbol.
Fig. 3. Time series of (a) the average latitudinal location of the ship, (b) concentrations of trajectory-weighted chlorophyll $a$ and (c) fractions of organic (WSON and WION) and inorganic nitrogen together with concentrations of MSA. “D” means data obtained in daytime, whereas “N” means data obtained in nighttime.
Fig. 4. Latitudinal distributions of (a) organic nitrogen (ON), (b) water-insoluble organic nitrogen (WION), (c) water-soluble organic nitrogen (WSON), and (d) diethylammonium (DEA⁺).
Fig. 5. Latitudinal distributions of (a) organic carbon (OC), (b) water-insoluble organic carbon (WIOC), (c) water-soluble organic carbon (WSOC).
Fig. 6. Scatter plots between $\delta^{13}$C and ON/OC ratios. Solid circles indicate the data with MSA $>30$ ng m$^{-3}$. 
Fig. 7. Relations of (a) WION concentrations and (b) WION/WIOC ratios with local wind speed at 40–44° N (open circles) and 10–40° N (solid circles). Values and bars for wind speed indicate averages and standard deviations, respectively, during each sampling time.
Fig. 8. A linear relation between WION and NH$_4^+$, color-coded according to the concentration of DEA$^+$. 