



1 **Aerosol organic nitrogen across the global marine boundary**
2 **layer: distribution patterns and controlling factors**

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25 **Abstract**

26 Organic nitrogen (ON) is an important yet poorly constrained component of aerosol
27 total nitrogen (TN), particularly over remote oceans. We quantified aerosol ON in 92
28 total suspended particulate samples collected across approximately 160° of latitude in
29 the marine atmospheric boundary layer (MABL) during Chinese Antarctic and Arctic
30 expeditions (2019–2024), using a newly developed method that simultaneously
31 determines ON and inorganic nitrogen. A significant latitudinal gradient was observed,
32 with significantly higher ON concentrations in the Northern Hemisphere (83.3 ± 141.4
33 ng m^{-3}) than in the Southern Hemisphere ($15.4 \pm 12.4 \text{ ng mm}^{-3}$). Regionally, coastal
34 East Asia recorded the highest ON levels ($164.6 \pm 179.1 \text{ ng mm}^{-3}$) but a lower ON/TN
35 ratio (21.1%), indicating strong terrestrial and anthropogenic influence. In contrast,
36 the Arctic Ocean had lower ON concentrations ($19.1 \pm 19.0 \text{ ng mm}^{-3}$) but the highest
37 ON/TN ratio (38.6%), suggesting dominant marine biogenic sources. The Southern
38 Ocean showed the lowest ON concentration ($12.0 \pm 7.1 \text{ ng m}^{-3}$) yet a relatively high
39 ON/TN ratio (27.8%), also pointing to oceanic origins. Near Antarctica, samples
40 influenced by sea-ice air masses displayed markedly elevated ON and ON/TN ratios.
41 These increases were strongly correlated with sea ice concentration and chlorophyll-a
42 exposure, indicating enhanced biogenic emissions from sea-ice-associated ecosystems.
43 This study offers the first direct ON measurements along a global MABL transect,
44 revealing distinct latitudinal and regional patterns, and emphasizing the combined
45 roles of continental inputs and marine sources. It also identifies sea-ice dynamics as a
46 key factor influencing ON in Antarctic regions, providing crucial data for improving
47 atmospheric and climate models.

48



49 **Key points.**

- 50 1. Aerosol organic nitrogen (ON) in the global marine boundary layer was quantified
51 along a transect from the Antarctic to the Arctic ($\sim 160^\circ$ latitude) for the first time.
52 2. A strong latitudinal gradient in ON was observed, revealing distinct hemispheric
53 and regional patterns.
54 3. Near Antarctic, ON concentrations and ON/TN ratios were distinctly elevated in
55 sea-ice-influenced air masses, highlighting the role of sea-ice dynamics.

56 **1. Introduction**

57 Marine atmospheric boundary layer (MABL) aerosol particles contain significant
58 amounts of organic nitrogen (ON) and inorganic nitrogen (IN), both recognized as
59 major components of atmospheric particulate matter (Li et al., 2023). ON may
60 account for roughly 20–80% of total reactive nitrogen deposition to the surface ocean,
61 implying a potentially large, yet uncertain, role in marine nitrogen cycling and climate
62 (Altieri et al., 2016, 2021). However, ON remains poorly constrained due to analytical
63 limitations (Baker et al., 2017). Previous studies focused on the water-soluble fraction
64 of aerosol ON (WSON) inferred indirectly by subtraction IN from total nitrogen (TN)
65 ($\text{ON} = \text{TN} - \text{IN}$), while the water-insoluble organic nitrogen (WION) fraction has
66 been largely unquantified (Cornell, 1999; Mace et al., 2003). The subtraction
67 approach is prone to errors and artifacts, especially when TN and IN concentrations
68 are similar, leading to underestimation and large uncertainties in ON burdens and
69 fluxes.

70 Aerosol ON arises from diverse sources. Marine pathways include primary
71 emissions via sea spray enriched with organic matter from the sea surface microlayer
72 and secondary formation from marine precursors (e.g., alkylamines) reacting with
73 acidic species (Facchini et al., 2008; Miyazaki et al., 2011a). Continental pathways
74 include long-range transport of organic emissions from fossil fuel combustion,
75 biomass burning, soils, and vegetation (Cape et al., 2011; Jickells et al., 2013; Luo et
76 al., 2018). Primary marine emissions inject large amounts of particulate matter
77 annually, carrying organic carbon and nitrogen from plankton, bacteria, and surface



78 films (Violaki et al., 2015a). Observations have shown that sea spray can carry
79 substantial ON and that WION can dominate ocean-influenced aerosol ON (Miyazaki
80 et al., 2011a).

81 ON affects climate and biogeochemistry by supplying bioavailable nitrogen,
82 modifying cloud condensation nuclei and ice-nucleating particle populations, and
83 contributing to aerosol light absorption. Hygroscopic ON compounds (e.g., amino
84 acids, amines, sugars) enhance water uptake and cloud condensation nuclei (CCN)
85 activity; some proteinaceous organics act as efficient ice nuclei (Alsante et al., 2024;
86 Chan et al., 2005). Marine alkylamines can form salts with sulfuric acid, promoting
87 new particle formation and growth, thereby linking ON to aerosol number and
88 radiative forcing (Almeida et al., 2013; Brean et al., 2021). Nitrogen-containing
89 chromophores (brown nitrogen) can dominate the absorptive properties of organic
90 aerosol regionally and contribute substantially to global absorption by carbonaceous
91 aerosol (Li et al., 2025).

92 Despite this importance, ON sources over remote oceans remain debated. Some
93 studies implicate continental transport (e.g., dust, anthropogenic emissions), whereas
94 others point to direct sea spray emissions or secondary formation from marine-derived
95 alkylamines (Altieri et al., 2016; Lesworth et al., 2010; Zamora et al., 2011).
96 Correlations between ON and ocean biological proxies (e.g., chlorophyll-a) suggest in
97 situ marine production, particularly during phytoplankton blooms (Altieri et al., 2016;
98 Dall'Osto et al., 2019). Yet open-ocean and polar regions, where sea-ice dynamics
99 may create distinct ON emissions, remain sparsely observed (Matsumoto et al., 2022).
100 Around Antarctica in particular, the paucity of direct ON measurements—especially
101 of WION—limits understanding of ON sources, seasonality, and impacts on
102 high-latitude atmospheric chemistry.

103 To address these gaps, we measured aerosol ON and IN using samples collected
104 during three Chinese Antarctic research expeditions campaigns, spanning $\sim 160^\circ$ of
105 latitude from the Arctic to Antarctica. We employed a newly developed aerosol
106 nitrogen analyzer based on thermal evolution and chemiluminescence detection to
107 measure aerosol IN and ON simultaneously, eliminating subtraction-based method



108 biases and capturing both WSON and WION (Yu et al., 2021). This dataset enables
 109 evaluation of hemispheric and regional patterns, assessment of controlling factors
 110 (e.g., continental influence, marine biological activity), and explicit investigation of
 111 sea-ice-associated processes near Antarctica. The results provide observational
 112 constraints to improve representation of nitrogen cycling and atmosphere–ocean
 113 interactions in climate and atmospheric chemistry models.

114 **2. Methodology**

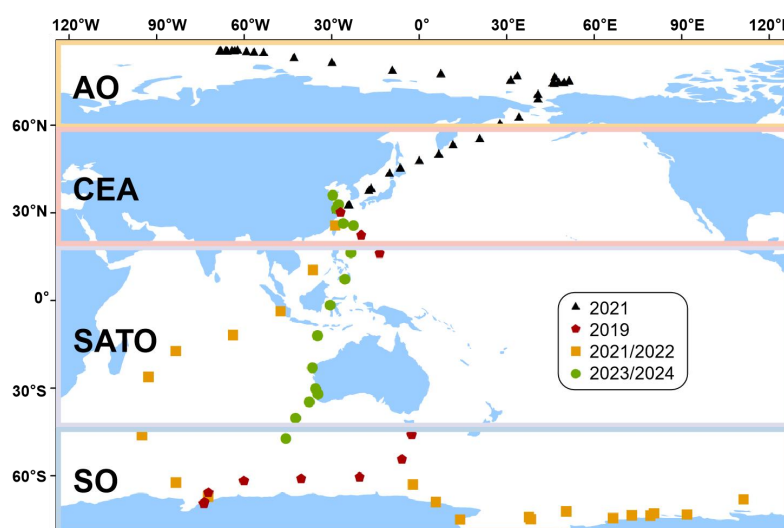
115 **2.1. Sample Collection**

116 A total of 92 total suspended particulate (TSP) samples were collected during three
 117 Chinese Antarctic research expeditions and one Arctic expedition aboard the
 118 icebreaker R/V *Xuelong*. Sampling spanned a latitudinal range of approximately 160°
 119 (86°N to 75°S), encompassing polar and mid-latitude marine regions. The Antarctic
 120 samplings were conducted in October to November in 2019 (SP2019, 14 samples),
 121 November 2021 to March 2022 (SP2021, 23 samples), and October 2023 to April
 122 2024 (SP2023, 15 samples), while the Arctic campaign occurred in July to September
 123 in 2021 (40 samples).

124 During the Antarctic campaigns, aerosols were collected using a high-volume air
 125 sampler (HVAS, TISCH Environmental, USA; flow rate: 1.2 m³ min⁻¹) equipped with
 126 pre-baked (500°C, 24 h) Whatman quartz filters (20.3 × 25.4 cm; Whatman Ltd., UK).
 127 For Arctic sampling, a DIGITEL DHA-80 sampler (flow rate: 500 L min⁻¹) with 14.2
 128 cm diameter Whatman quartz filters were employed. Each sample represented a
 129 48-hour integrated collection period, corresponding to 2–4° latitude traversed during
 130 ship transits. To minimize contamination from ship emissions, a wind sector controller
 131 restricted sampling to air masses within 120° of the ship's heading. Filters were
 132 handled using nitrile gloves and masks to avoid potential contamination.
 133 Post-sampling, filters were folded with the collection surface inward, wrapped in
 134 pre-cleaned aluminum foil, sealed in polyethylene bags, labeled with sampling time
 135 and location, and stored at -20°C. Detailed protocols followed established
 136 methodologies (Shi et al., 2021). Following expeditions, samples were transported to



137 the laboratory under frozen conditions and maintained at -20°C until analysis. The
 138 sampling location for the Antarctic and Arctic campaigns are illustrated in Fig. 1.
 139



140
 141 Figure 1. TSP Aerosol sampling locations along the cruises path from Shanghai, China to
 142 Antarctica and Arctic.

143 2.2. Chemistry Analysis for major ions, EC and OC

144 Major ions were quantified through ion chromatographic analysis of water extracts of
 145 the aerosol samples. The extraction of filters in the laboratory followed protocols
 146 comparable to those described in the previous study (Shi et al., 2021). Prior to
 147 measurement, three-quarters of each filter was sectioned into small pieces using
 148 acid-cleaned Teflon-coated scissors and transferred into high-purity Milli-Q water
 149 ($18.2\text{ M } \Omega$). The suspensions were subjected to ultrasonic treatment for 30 min,
 150 followed by continuous shaking at 120 rpm for 12 h to ensure thorough extraction of
 151 water-soluble components. The extracts were subsequently filtered through $0.22\text{ }\mu\text{m}$
 152 polytetrafluoroethylene (PTFE) membranes prior to ion analysis. The concentrations
 153 of the main ions (NO_3^- , SO_4^{2-} , Na^+ , NH_4^+ , K^+ , and Ca^{2+}) in the sample were
 154 determined by an ion chromatograph (AQ1100, RFIC, equipped with a CS12 column
 155 ($2 \times 250\text{ mm}$) for cation analysis, AS11 column ($2 \times 250\text{ mm}$) for anion analysis,



156 Thermo Scientific, USA), and the eluents of cation and anion were 18.00 mM
 157 methylsulfonic acid (MSA) and potassium hydroxide (KOH), respectively. During
 158 sample analysis, the relative deviation of repeated assays ($n = 5$) of all ions is usually
 159 less than 5%. We used the following formula to calculate non-sea salt SO_4^{2-}
 160 (nssSO_4^{2-}), non-sea salt Ca^{2+} (nssCa^{2+}) and non-sea salt K^+ (nssK^+):

$$[\text{nssSO}_4^{2-}] = [\text{total SO}_4^{2-}] - 0.253 \times [\text{Na}^+] \quad (1)$$

$$[\text{nssCa}^{2+}] = [\text{total Ca}^{2+}] - 0.038 \times [\text{Na}^+] \quad (2)$$

$$[\text{nssK}^+] = [\text{total K}^+] - 0.037 \times [\text{Na}^+] \quad (3)$$

161 where 0.252, 0.037, and 0.038 in the above expressions are the ratios of $\text{SO}_4^{2-}/\text{Na}^+$ (Xu
 162 et al., 2013), $\text{Ca}^{2+}/\text{Na}^+$ (Anonymous, 1997), and K^+/Na^+ (Keene et al., 1986) in the sea
 163 water, respectively.

164 OC and EC concentrations were determined using a Thermal/Optical Carbon
 165 Analyzer (DRI, Model 2001, Atmoslytic Inc., USA) following the IMPROVE
 166 protocol as implemented by Wu et al., (2024). OC and EC measurements were
 167 conducted for aerosol filters collected during the 2021 Arctic and 2019 Antarctic
 168 cruises.

169 2.3. ON measurement

170 Aerosol ON and IN were simultaneously measured using the recently developed
 171 Aerosol Nitrogen Analyzer system, which enables sensitive quantification directly
 172 from filter samples without pretreatment. Detailed descriptions of the method are
 173 provided in Yu et al (2021). In brief, the analyzer integrates a thermal aerosol carbon
 174 analyzer and a chemiluminescence NO_x analyzer. Aerosol samples collected on quartz
 175 fiber filters were thermally evolved under a programmed 6-step temperature protocol
 176 (150, 180, 300, 400, 500, and 800 °C) in a 1% O_2 /99% He carrier gas. The evolved
 177 materials were catalytically oxidized to CO_2 and nitrogen oxides (NO_y), with the C
 178 signal monitored via methanator-FID detection and the N signal recorded through
 179 chemiluminescence after converting NO_y to NO. The C signal assists in
 180 differentiating IN and ON components, as ON aerosols produce both C and N signals
 181 while the IN fraction only yields an N signal. The programmed thermal evolution



182 facilitates separation of aerosol IN and ON due to their distinct thermal characteristics.
 183 Quantification of IN and ON is achieved through multivariate curve resolution (MCR)
 184 data treatment of the C and N thermograms using USEPA PMF (version 5.0).

185 **2.4. Backward Trajectory Analysis**

186 To study air mass origins, air mass backward trajectories have been calculated using
 187 the Hybrid Single-Particle Lagrangian Integrated Trajectories (HY-SPLIT) model with
 188 meteorological fields from the National Oceanic and Atmospheric Administration
 189 (NOAA) air resources laboratory GDAS database. Five-day backward trajectories
 190 were calculated in order to reveal the history of the air masses arriving at the sampling
 191 site (Stein et al., 2015). Each trajectory originated at the vessel's real-time position
 192 with an arrival height of 20 m, capturing boundary layer transport while minimizing
 193 local ship influence.

194 **2.5. Probability Source Contribution Function analysis**

195 Potential Source Contribution Function (PSCF) analysis was implemented to identify
 196 source regions of ON observed during the sampling period (Ashbaugh et al., 1985). A
 197 higher PSCF value indicates a greater potential source contribution to the receptor site.
 198 In our study, the PSCF domain was established within a grid cell encompassing all
 199 backward trajectories. The cruises were discretized into 1° latitude × 1° longitude grid
 200 cells. The PSCF value for cell ij was calculated as:

$$201 \quad \text{PSCF}_{ij} = \frac{\sum m_{ij}}{\sum n_{ij}} \quad (4)$$

202 where, m_{ij} = total trajectory endpoints within cell ij; n_{ij} = subset of endpoints
 203 associated with aerosol component concentrations exceeding the 75th percentile of
 204 cruise measurements. To mitigate uncertainty in cells with sparse trajectory density, a
 205 latitude-dependent weighting function (W) was applied:

$$206 \quad W = \begin{cases} 1.0 & \text{when } n_{ij} > N2 \\ 0.8 & \text{when } N1 < n_{ij} < N2 \\ 0 & \text{when } n_{ij} < N1 \end{cases} \quad (5)$$

207 where n_{ij} is the number of trajectories passing for each cell in the study period and
 208 $N1 = 60 \cdot \cos(\text{latitude})$, and $N2 = 300 \cdot \cos(\text{latitude})$. The cosine factor is used to



209 account for the changing grid cell size with varying latitude.

210 **2.6. Air-mass exposure to chlorophyll a**

211 The Air-mass Exposure to Chlorophyll a (Chl-a) index (AEC) serves as a quantitative
 212 metric to assess the influence of marine biogenic emissions on a target region through
 213 air mass transport (Blazina et al., 2017; Choi et al., 2019). This approach is grounded
 214 in the well-established correlation between ocean surface phytoplankton biomass and
 215 marine biogenic emissions, particularly dimethyl sulfide (DMS), where Chl-*a*
 216 concentration acts as a robust proxy for phytoplankton abundance (Siegel et al., 2013).
 217 The AEC index estimates the integrated exposure of an air mass to oceanic DMS
 218 source regions along its trajectory by accounting for both spatial distribution of Chl-a
 219 and atmospheric vertical mixing dynamics (Zhou et al., 2023).

220 For each trajectory point, Chl-a concentrations (Chl_{*a*}) were obtained from
 221 satellite remote sensing products (Aqua-MODIS, OCI algorithm; 8-day composite, 4
 222 km × 4 km resolution; <https://oceancolor.gsfc.nasa.gov/13/>) within a 20-km radius.
 223 Trajectory endpoints over Antarctica, sea-ice-covered areas, or at pressures < 850 hPa
 224 were assigned Chl-a = 0 (Zhou et al., 2023). Points without valid Chl-a data were
 225 excluded. The AEC for a single trajectory was computed as:

$$226 \quad AEC = \frac{\sum_{i=1}^{120} Chl_{a_i} \times e^{-\frac{t_i}{120}}}{n} \quad (6)$$

227 where t_i denotes time backward along the trajectory (hours), and n is the total number
 228 of valid trajectory points. The time points when the air mass passed over the continent
 229 or regions covered by sea ice were assigned a zero chlorophyll value. To ensure
 230 robustness, trajectories with $n < 90$ (75% of 120 h data points at hourly resolution)
 231 were discarded. For each sample, the final AEC value was derived from the arithmetic
 232 mean of all valid trajectories during the sampling period (Yan et al., 2024).

233 **2.7. Sea ice concentration**

234 In this study, remote sensing data are utilized to illustrate the spatiotemporal
 235 distribution of sea ice concentrations (SICs) in the Southern Ocean. The SIC of each



sample is calculated using the following formula:

$$SIC = \frac{\sum_{i=1}^{N_s} SIC_i}{N_s} \quad (7)$$

where SIC_i represents the average sea ice density at the endpoint of the specified track. N_s represents the total number of trajectory endpoints located on the sea ice area. Sea ice concentration data are from the AMSR2 dataset (Version 5.4. University of Bremen, Germany. Index of /amsr2/asi_daygrid_swath/s3125) (Melsheimer and Spreen, 2019).

3. Results

Atmospheric ON concentrations exhibited significant hemispheric differences (independent samples t-test; $p < 0.001$), with values in the Northern Hemisphere (NH: $83.3 \pm 141.4 \text{ ng m}^{-3}$, $N = 55$) being approximately five times higher than those in the Southern Hemisphere (SH: $15.4 \pm 12.4 \text{ ng m}^{-3}$, $N = 37$). The ON/TN ratios showed broadly similar magnitudes between hemispheres, with slightly higher in the NH ($30.4 \pm 13.6\%$) compared to the SH ($27.9 \pm 10.6\%$). Samples from three Antarctic cruises—SP2019 (mean = 19.4 ng m^{-3} ; range: $9.5\text{--}555.6 \text{ ng m}^{-3}$), SP2021 (mean = 20.4 ng m^{-3} ; range: $1.3\text{--}81.3 \text{ ng m}^{-3}$), and SP2023 (mean = 18.3 ng m^{-3} ; range: $1.8\text{--}457.0 \text{ ng m}^{-3}$) showed no significant variation (one-way ANOVA; $p > 0.2$), indicating that interannual variation was rather minor. A clear latitudinal gradient in ON concentrations was observed along the Antarctic-to-Arctic transect, with peak values in the $20\text{--}40^\circ \text{ N}$ zone and a gradual decline toward both polar regions (Fig. 2a). Based on spatial distribution patterns, the study transect can be divided into four regions (Fig. 1): (1) the Arctic Ocean region (AO, north of $\sim 60^\circ \text{ N}$); (2) the Coastal East Asia region (CEA, $20\text{--}60^\circ \text{ N}$); (3) the Southeast Asia-Australia Tropical Ocean region (SATO, $\sim 20^\circ \text{ N}\text{--}40^\circ \text{ S}$); and (4) the Southern Ocean region (SO, south of $\sim 40^\circ \text{ S}$).

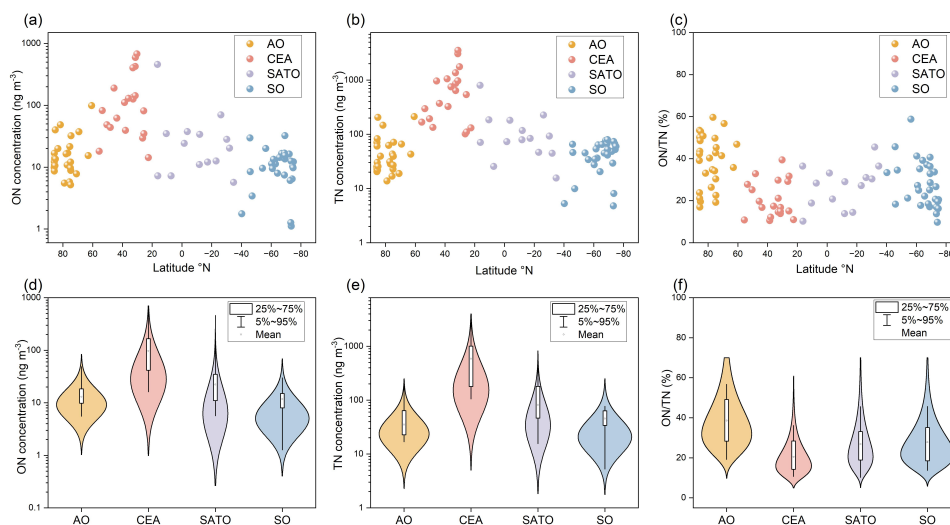
The CEA region exhibited the highest ON concentrations (mean = 164.6 ng m^{-3}) but the lowest ON/TN ratio (mean = 21.1%). In contrast, the SO region showed the lowest ON concentrations (mean = 12.0 ng m^{-3} ; range: $1.78\text{--}32.3 \text{ ng m}^{-3}$) and higher ON/TN ratios (mean = 27.8%). Notably, the AO region displayed the highest ON/TN



265 ratios (mean = 38.6%) despite relatively low ON concentrations (mean = 19.1 ng m⁻³;
266 range: 5.2–32.2 ng m⁻³). The ON/TN ratio in SATO region (26.8%) is similar to that
267 of SO but with a lower ON concentration (mean = 23.4 ng m⁻³; range: 5.7–70.1 ng
268 m⁻³), which is much lower than the CEA region, but higher than the high latitude two
269 pole regions.

270 Since direct measurement data of total ON in global MABL are limited, WSON
271 data were summarized for comparison (Table 1). Overall, the previous results are
272 consistent with the spatial trends of ON in our study. WSON concentrations exhibit
273 significant spatial variation, generally higher in the NH than in the SH, highlighting
274 the substantial contribution of anthropogenic sources (Violaki et al., 2015b). In
275 addition, WSON concentrations tend to be higher closer to land, while in remote
276 ocean areas, WSON levels are generally lower. The reported ratios of WSON/TN in
277 previous studies vary significantly across different investigation sites. Moreover, in
278 remote marine environments, the WSON/TN ratio is relatively high, suggesting that
279 WSON plays a substantial role in the biogeochemical cycle of nitrogen within these
280 remote regions. It is important to note that most previous studies over the remote
281 ocean measured only WSON, without accounting for the WION. As a result, the
282 ON/TN ratios in this region were likely underestimated. Based on our comparison, the
283 total ON concentration in the Southern Ocean may have been underestimated by
284 approximately 40%, hinting the significant contribution of the insoluble organic
285 fraction that has been largely overlooked in earlier datasets due to measurement
286 method limitations.

287



288

289 Figure 2. Latitudinal distributions of ON concentration, TN concentration and ON/TN ratio (a, b,
 290 c), and the statistics (d, e, f) over the the Arctic Ocean region (AO), the Coastal East Asia region
 291 (CEA), the Southeast Asia-Australia Tropical Ocean region (SATO) and the Southern Ocean
 292 region (SO), respectively.

Table 1. Measured concentrations of WSON from published reports in the marine atmospheric boundary layer.

Regions	Period	Locations	WSON (ng m ⁻³)	Ratio to TN (%)	Methods for IN	Methods for TN	Reference
Southern Atlantic	2007.01-02	35° S–45° S	119 ± 163.8	-	IC	PO	(Violaki et al., 2015b)
the Northwest Pacific Ocean	2014.2015	25–40° N, 125–150° E	43.4–564.2	11–46	IC	PO	(Luo et al., 2018)
the coast of China the East China seas	2014.2015	30–40° N, 120–130° E	96.6–7238	6–48	IC	PO	(Luo et al., 2018)
Southern Ocean	2000.11	40.41° S, 144.41° E	50.4 ± 79.8	21	IC	UV	(Mace et al., 2003)
Oahu, Hawaii	1998.7.22-8.13	21.7° N, 157.8° W	46.2	31	IC	UV	(Cornell et al., 2001)
the southern margin of the East China Sea	2005-2006	25.09° N, 121.46° E	476 ± 756	24 ± 16	IC	UV with PO	(Chen and Chen, 2010)
the western North Pacific	2008.08.24-09.13	42.98° N, 144.37° E –35.65° N, 139.77° E	130 ± 61 (10–260)	67 ± 15	IC	TOC/TN analyzer	(Miyazaki et al., 2011b)
Huaniao Island	2019	30.86° N, 122.67° E	30–2810	0.13–77	IC	TOC/TN analyzer	(Tian et al., 2023)
the northern tip of Japan	2010-2012	~45.2° N, ~141.2° E	77 ± 57	12.8 ± 15.2	IC	TOC/TN analyzer	(Matsumoto et al., 2017)
the Southern Ocean	2016-2020	38.8–69.0° S, 38.1–150.8° E	4.7	20	IC	TOC/TN analyzer	(Matsumoto et al., 2022)
the Subarctic Western North Pacific Ocean	2016.7.21-8.22	30–65° N, 130–160° W	1.62–205.8	9	IC	TOC/TN analyzer	(Jung et al., 2019)
Bermuda	2011	32.27° N, 64.87° W	105 ± 191.8	50.4 ± 18.9	nutrient analyzer	TN Analyzer	(Altieri et al., 2016)

*PO: the persulfate oxidation (PO) method

*UV: ultraviolet photo-oxidation

*TN analyzer: a total organic carbon (TOC) analyzer with a TN unit

*Nutrient analyzer: automated nutrient analyzer and standard colorimetric method

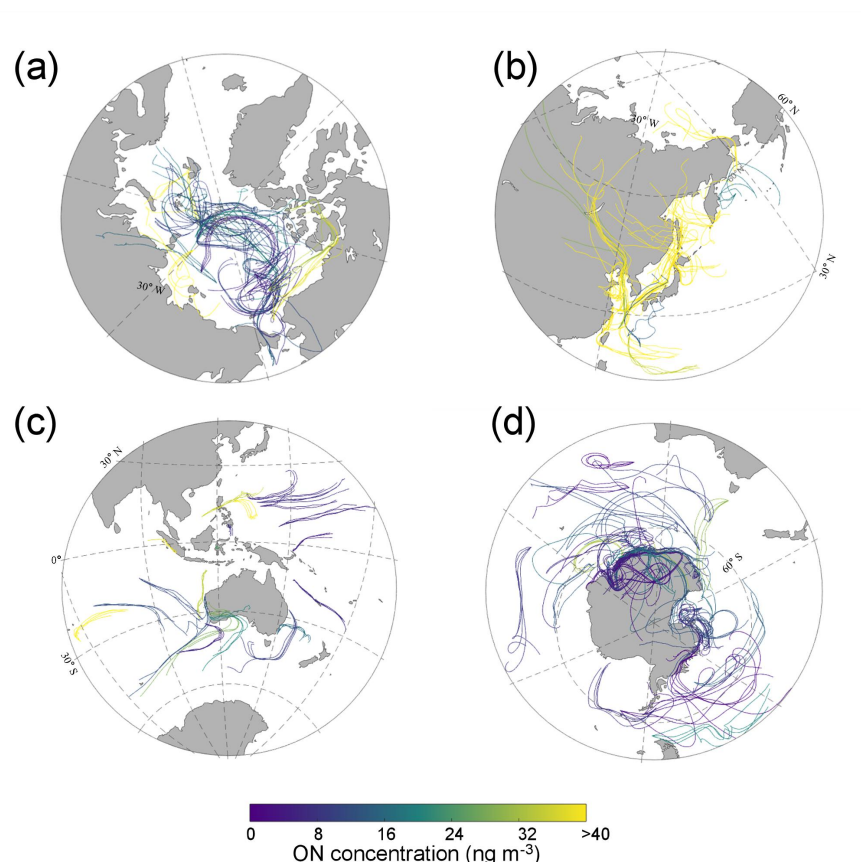


294 4. Discussion

295 4.1 Source identification of ON

296 ON in the MABL primarily originates from two main source pathways: marine
 297 emissions and long-distance continental transport. Marine sources include primary
 298 ON, predominantly associated with sea-spray particles enriched in biological material
 299 from the ocean surface microlayer, and secondary ON, which often derives from
 300 marine precursors such as alkylamines that react with acidic species to form new
 301 particles (Altieri et al., 2016; Facchini et al., 2008). Continental sources involve the
 302 long-range transport of organic emissions—including combustion byproducts, soil-
 303 and vegetation-derived compounds, and biomass burning aerosols—that can
 304 significantly influence remote ocean regions (Cape et al., 2011; Jickells et al., 2013).

305 ON concentrations in the CEA region were the highest among all study regions,
 306 with air masses spending 22.6% of their 5-day trajectories over continental areas (Fig.
 307 3b). A significant correlation between ON and crustal elements such as nssCa^{2+} ($r =$
 308 0.75 , $p < 0.01$; Fig. 4) likely suggests the influences of continental transport of
 309 particles on the ON levels in this region (Xiao et al., 2016). A significant correlation
 310 between ON and the anthropogenic tracer EC ($r = 0.81$, $p < 0.01$; Fig. 4) indicates that
 311 fossil fuel combustion and biomass burning are major ON sources (Shubhankar and
 312 Ambade, 2016; Wu and Yu, 2016). Similarly, the robust association between ON and
 313 nssK^+ ($r = 0.78$, $p < 0.01$; Fig. 4), a tracer of biomass burning, also supports
 314 contributions from agricultural and residential biomass burning (Song et al., 2018).
 315 Despite the high absolute ON concentrations, the relatively low ON/TN ratio (21.1%)
 316 likely reflects disproportionately elevated IN emissions from intensive human
 317 activities, particularly NH_3 volatilization from agriculture and vehicular NO_x
 318 emissions (Pavuluri et al., 2015). This interpretation aligns with emission inventories
 319 that identify the CEA as a global nitrogen pollution hotspot, where ON is co-emitted
 320 or formed from precursors that share common sources with EC and other
 321 combustion-related pollutants, originating from incomplete combustion and industrial
 322 processes (Deng et al., 2024).

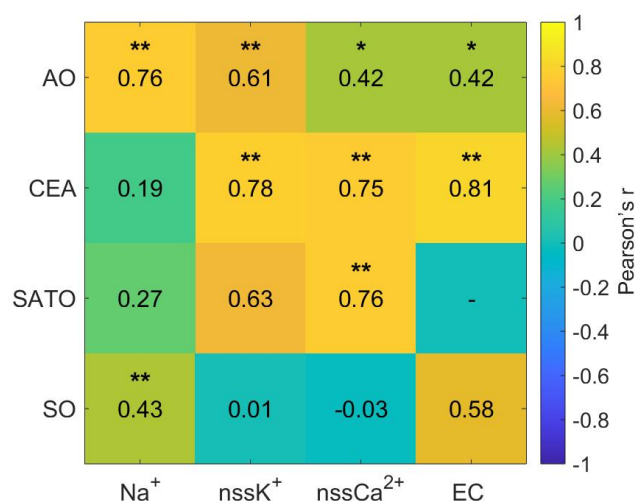


323
 324 Figure 3. 5-day air-mass backward trajectories with ON concentrations and ON/TN ratios
 325 along the Chinese Arctic/Antarctic expedition voyage over the Arctic Ocean (a), the coastal
 326 East Asia (b), the Southeast Asia-Australia Tropical Ocean (c), and the Southern Ocean (d).

327 The SATO region exhibits intermediate level of ON concentrations (mean = 23.4
 328 $\pm 18.0 \text{ ng m}^{-3}$), lower than those influenced by anthropogenic activities in CEA but
 329 higher than in polar regions. In this region, ON shows a significant positive
 330 correlation with nssCa^{2+} (Fig. 4; $r = 0.76$, $p < 0.01$), suggesting that continental inputs
 331 influence ON levels. In addition, backward trajectory analysis showed that samples
 332 affected by continental air masses have significantly higher ON concentrations than
 333 those exposed solely to marine air (Fig. 3c), suggesting the influences of continental
 334 sources. However, ON does not exhibit significant correlations with nssK^{+} or with EC
 335 ($p > 0.05$), indicating that combustion emissions may not be the primary drivers.
 336 These findings suggest that the variability of ON in the SATO region results from a
 337 mixture of marine-terrestrial interactions, primarily modulated by episodic continental



338 influences rather than continuous marine emissions. Notably, this region displays an
 339 elevated ON/TN ratio (Fig. 2), primarily due to its very low IN levels—approximately
 340 85% lower than in the CEA region—which amplifies the relative contribution of ON
 341 within TN.
 342



343
 344 Figure 4. Spatial variations in the correlation between ON and ionic species across four
 345 regions (values in cells, with “***” indicating $p < 0.01$ and “**” indicating $p < 0.05$).

346 In the AO region, ON concentrations were slightly lower than in the SATO
 347 region but significantly lower than in the CEA region. In this area, ON exhibited a
 348 significant positive correlation with Na⁺, which suggest the sea salts inputs (Fig. 4; $r =$
 349 0.43, $p < 0.01$), and also showed significant correlations with nssK⁺ ($r = 0.61$, $p <$
 350 0.01). Its correlations with nssCa²⁺ ($r = 0.42$) and EC ($r = 0.42$) were weaker but still
 351 significant ($p < 0.05$). These patterns suggest that ON in the AO region may originate
 352 not only from primary sea-salt aerosols but may also be linked to biomass burning.
 353 However, the backward trajectory analysis shows no significant difference in ON
 354 concentrations between air masses influenced by continental sources and those
 355 transported solely over the ocean (Fig. S1b; independent samples t-test, $p = 0.16$),
 356 likely suggesting the limited role of terrestrial inputs in this region. Unlike the SATO
 357 region, where ON showed no correlation with AEC (Fig. S2b; $p > 0.05$), ON
 358 concentrations in the AO exhibited a strong positive correlation with the AEC (Fig.
 359 S2a; $p < 0.01$), suggesting that marine biological activity is a key driver of ON



variability in this region (Creamean et al., 2022). Collectively, these results demonstrate that the AO region is primarily governed by marine processes, with ON derived from both sea-spray organic enrichment and biogenic aerosol precursors, while terrestrial influences remain secondary (Nøjgaard et al., 2022).

In the SO region, ON concentrations were the lowest among all regions (mean = $12.0 \pm 7.1 \text{ ng m}^{-3}$), yet the ON/TN ratio was relatively high ($27.8 \pm 11.0\%$). Back-trajectory analysis indicates that air masses predominantly originated from the open ocean and Antarctic continent (Fig. 3d), with minimal anthropogenic influence. ON here exhibited a significant positive correlation with Na^+ (Fig. 4; $r = 0.43$, $p < 0.01$), but no significant relationships with nssK^+ , nssCa^{2+} or EC. This pattern suggests that primary sea-salt emissions are a major pathway for ON in the SO atmosphere (Matsumoto et al., 2022), likely through the incorporation of marine-derived organic matter into sea-spray aerosols. Meanwhile, the absence of associations with terrestrial tracers further supports the notion that ON in this remote region is controlled primarily by natural marine processes rather than continental or anthropogenic sources (Altieri et al., 2016).

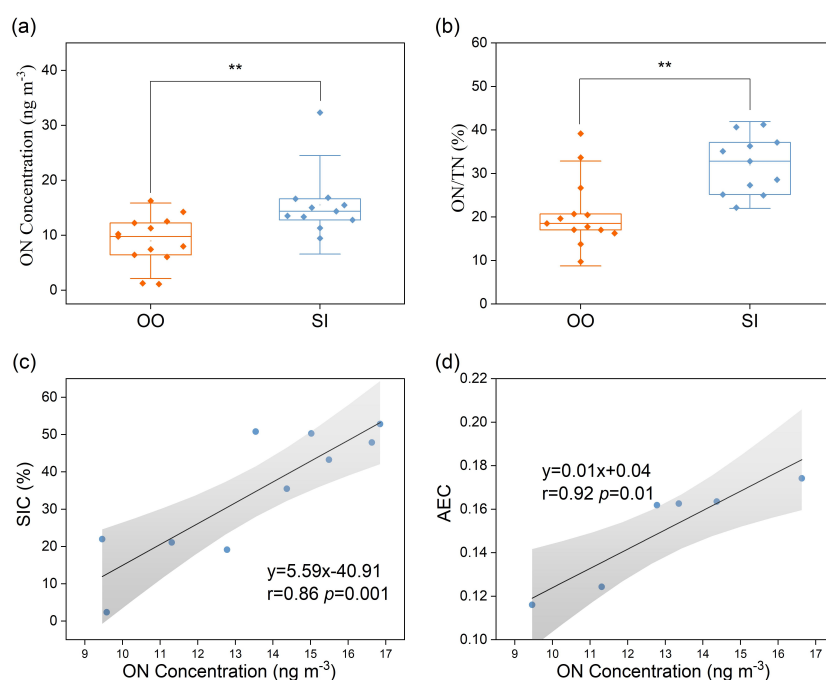
4.2 Role of sea-ice–associated biogenic processes in shaping Antarctic aerosol ON

Sea-ice and open-ocean environments create distinct conditions for the production and emission of ON. While sea ice restricts direct air–sea exchange, it hosts specialized microbial communities and accumulates organic matter within brine channels. During melt and ice-edge retreat, this organic material is released, enriching the surface microlayer and supplying precursors for aerosolization via sea spray and secondary formation (Dall’Osto et al., 2017; DeMott et al., 2016; Galgani et al., 2016; Wilson et al., 2015).

Along the Antarctic coast, we classified samples into two groups based on air-mass histories: open ocean (OO), influenced almost exclusively by open-ocean trajectories, and sea ice (SI), with air masses residing over sea ice for extended periods. SI samples exhibited significantly higher ON concentrations and ON/TN ratios than OO samples (independent-samples t-test, $p < 0.01$; Fig. 5a, b). Multiple lines of evidence point to sea-ice–linked biological processes as the driver of these enhancements: (1) Strong positive correlations of ON with sea-ice concentration (SIC; $r = 0.86$, $p < 0.01$) and with air-mass exposure to chlorophyll-a (AEC; $r = 0.91$, $p < 0.01$) in the SI group indicate that both ice cover and associated biological activity



393 elevate ON (Fig. 5c, d); (2) PSCF analysis identifies high-probability source regions
 394 (PSCF > 0.8) over sea ice and its marginal zone for SI samples (Fig. S3c), consistent
 395 with an ice-edge origin; and (3) In contrast, ON shows no significant correlation with
 396 Na^+ ($r = -0.22$, $p > 0.05$) or with IN ($p > 0.05$) for SI samples (Fig. S4a,b), suggesting
 397 that primary sea-salt emissions and purely abiotic inorganic pathways are not the
 398 dominant contributors.



399
 400 Figure 5. Comparison of measured ON concentrations (a) and ON/TN ratio (b) between SI
 401 and OO aerosol samples (“**”) indicating $p < 0.01$). And correlations between SIC (c), AEC (d)
 402 and ON concentration in SI aerosol samples (with $n = 10$ and 6 for panels c and d,
 403 respectively, owing to missing satellite data and the methodologies in Sections 2.6 and 2.7).

404 These observations support a mechanistic pathway whereby organic matter
 405 released from sympagic (ice-associated) communities during melt enriches the surface
 406 microlayer and is transferred to the atmosphere via sea spray as ON-rich particles
 407 (DeMott et al., 2016; Wilson et al., 2015). Concurrently, a portion of this organic
 408 nitrogen is rapidly microbially degraded to volatile alkylamines (e.g., methylamines)
 409 (Taubert et al., 2017), which then form aminium salts through acid–base reactions
 410 with marine emissions-derived acids (e.g., H_2SO_4 , MSA), contributing to both ON



411 and IN in SI conditions (Brean et al., 2021; Dawson et al., 2012; Fitzsimons et al.,
 412 2023). This process results in the formation of both organic (amine salts, contributing
 413 to ON) and inorganic nitrogen aerosol species (NH_4^+ and NO_3^-), which explains their
 414 elevated levels in the SI group samples (Fig. S5). The elevated ON/TN ratios in SI
 415 samples (31.0%) relative to OO samples (20.8%) further indicate a greater fractional
 416 contribution of ON under sea-ice influence (Fig. 5b), consistent with reported releases
 417 of organic material from the sympagic ecosystem during melt (Jang et al., 2023;
 418 Mirrielees et al., 2024; Yan et al., 2020).

419 For OO samples, PSCF hotspots ($\text{PSCF} > 0.8$) shift toward the offshore Southern
 420 Ocean (Fig. S3d), in line with trajectories dominated by open-ocean air masses. The
 421 positive association between ON and oceanic residence time ($r = 0.66$, $p < 0.01$; Fig.
 422 S6) suggests that, as sea-ice influence diminishes, ON variability becomes
 423 increasingly governed by open-ocean biological processes and long-range marine
 424 aerosol transport.

425 Overall, these results establish the ice-edge/sympagic environment as an
 426 important regulator of Antarctic aerosol ON. Sea-ice dynamics modulate both the
 427 magnitude (higher ON and ON/TN) and sources (biogenic enrichment and
 428 amine-driven secondary formation) of ON, underscoring the need to represent
 429 sea-ice-associated processes in polar atmospheric chemistry and climate models.

430 5. Conclusions and Implications

431 Taking advantage of a new analytical tool for ON and aerosol samples collected from
 432 three Antarctic and Arctic expeditions from 2019 to 2024, we quantified aerosol ON
 433 and IN in 92 TSP samples spanning 160° of latitude in the MABL. This dataset
 434 provides the first direct, subtraction-free ON measurements along a global-scale
 435 marine transect, capturing both water-soluble and water-insoluble fractions.

436 We observed a pronounced hemispheric and latitudinal gradient in ON, with
 437 substantially higher concentrations in the Northern Hemisphere ($83.3 \pm 141.4 \text{ ng m}^{-3}$)
 438 than in the Southern Hemisphere ($15.4 \pm 12.4 \text{ ng m}^{-3}$). Regionally, Coastal East Asia
 439 exhibited the highest ON ($164.6 \pm 179.1 \text{ ng m}^{-3}$) but a low ON/TN ratio (21.1%),
 440 consistent with strong terrestrial and anthropogenic influences that elevate IN. The
 441 Southeast Asia–Australia Tropical Ocean showed intermediate ON and a relatively
 442 high ON/TN ratio due to low IN. The Arctic Ocean had lower ON but the highest
 443 ON/TN ratio (38.6%), indicating prominent marine biogenic contributions. The



444 Southern Ocean showed the lowest ON ($12.0 \pm 7.0 \text{ ng m}^{-3}$) yet a relatively high
445 ON/TN ratio (27.8%), also suggestive of oceanic sources. Interannual variability
446 across the three Antarctic campaigns was minor.

447 Multiple lines of evidence, including correlations with tracers, back-trajectory
448 analysis, and PSCF, indicate that ON in CEA is dominated by continental inputs from
449 combustion and dust, whereas ON in AO and SO is primarily controlled by marine
450 processes. Along the Antarctic coast, air masses influenced by sea ice exhibited
451 significantly higher ON and ON/TN than those influenced by the open ocean, with
452 strong positive relationships to sea-ice concentration and air-mass exposure to
453 chlorophyll-a. These patterns point to sympagic and ice-edge biogenic
454 activity—through organic enrichment of sea spray and amine-driven secondary
455 formation—as key regulators of ON near Antarctica.

456 Comparison with prior WSON-only datasets suggests that earlier studies likely
457 underestimated total ON—by approximately 40% in the Southern Ocean—due to
458 omission of WION. Accounting for both soluble and insoluble phases is therefore
459 essential for constraining nitrogen deposition to the oceans and for representing ON's
460 roles in aerosol–cloud interactions and radiative effects.

461 These findings fill a critical observational gap, establish robust hemispheric and
462 regional patterns of marine aerosol ON, and provide essential constraints for
463 atmospheric chemistry and climate models. Future efforts should explicitly represent
464 ON sources, including sea-ice–associated biogenic processes and amine chemistry,
465 and expand year-round, size-resolved, and composition-resolved measurements paired
466 with isotopic and molecular tracers to refine source apportionment and evaluate model
467 parameterizations across regions and seasons.

468 **Data availability.**

469 The data on organic nitrogen concentrations in aerosol are available at National
470 Tibetan Plateau/Third Pole Environment Data Center,
471 <https://cstr.cn/18406.11.Atmos.tpd.303043>. DOI:
472 <https://doi.org/10.11888/Atmos.tpd.303043> (Sun, 2025) [Dataset].

473 **Author contribution.**

474 Ningning Sun: Data curation, Writing-original draft. Yu Xu: Methodology. Bo
475 Zhang and Ye Hu: Visualization, Software. Zhe Li: Methodology. Yilan Li: Carried
476 out data analysis. Zhenlou Chen: Review. Jian Zhen Yu: Supervision, Writing –



477 review & editing. Guitao Shi: Supervision, Writing – review & editing.

478 **Competing interests.**

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

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