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# Equal abundance of summertime natural and wintertime anthropogenic Arctic organic aerosols

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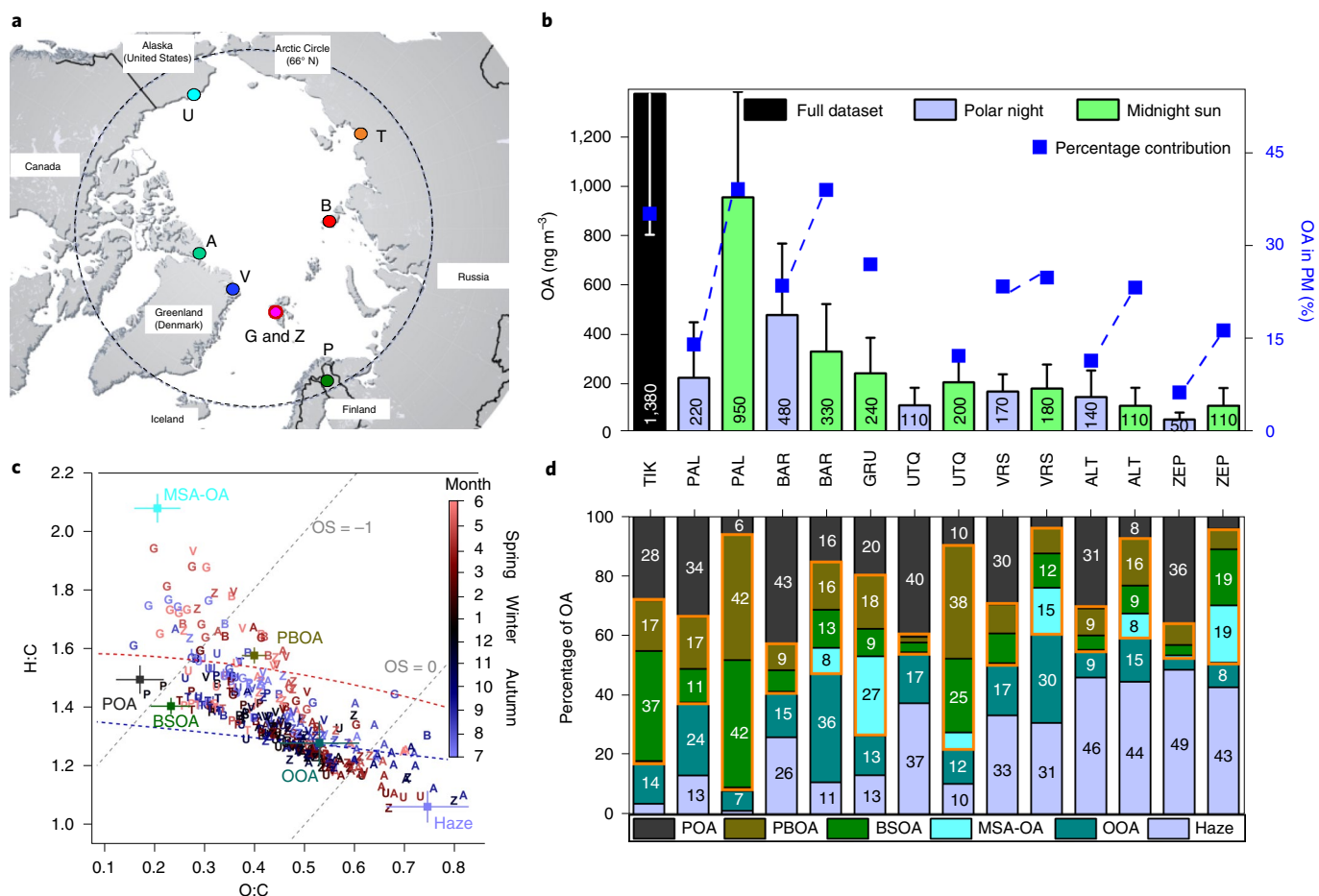
**Aerosols play an important yet uncertain role in modulating the radiation balance of the sensitive Arctic atmosphere. Organic aerosol is one of the most abundant, yet least understood, fractions of the Arctic aerosol mass. Here we use data from eight observatories that represent the entire Arctic to reveal the annual cycles in anthropogenic and biogenic sources of organic aerosol. We show that during winter, the organic aerosol in the Arctic is dominated by anthropogenic emissions, mainly from Eurasia, which consist of both direct combustion emissions and long-range transported, aged pollution. In summer, the decreasing anthropogenic pollution is replaced by natural emissions. These include marine secondary, biogenic secondary and primary biological emissions, which have the potential to be important to Arctic climate by modifying the cloud condensation nuclei properties and acting as ice-nucleating particles. Their source strength or atmospheric processing is sensitive to nutrient availability, solar radiation, temperature and snow cover. Our results provide a comprehensive understanding of the current pan-Arctic organic aerosol, which can be used to support modelling efforts that aim to quantify the climate impacts of emissions in this sensitive region.**

Organic aerosols (OAs) contribute to the Arctic aerosol mass near the surface<sup>1–4</sup> and affect the local climate through direct aerosol–radiation interactions and by altering the cloud properties<sup>5,6</sup>. OAs interact with other aerosol components<sup>7–9</sup>, for example, black carbon or elemental carbon (EC) and sulfate, and can augment or offset their radiative forcing<sup>10</sup>. The Arctic OA indirect effect is estimated to be of a similar magnitude as that of the sulfate indirect effect, and much larger than the OA direct effect<sup>11</sup>. The magnitude of these effects depends on the OA physicochemical properties, sources and formation processes<sup>12,13</sup>, which are not traceable by satellites<sup>14</sup>, and hence surface observations are indispensable<sup>15</sup>. However, OAs have received little attention in the Polar Regions<sup>16–18</sup>, mainly because of measurement challenges, and hence their complex composition and sources are poorly understood<sup>19,20</sup>. Nevertheless, an increasing abundance of natural OAs in a warming Arctic is expected<sup>21</sup> as a result of northward-expanding vegetation<sup>22</sup>,

intensifying boreal forest fires<sup>23,24</sup>, decreasing sea-ice extent<sup>25</sup> and thawing permafrost<sup>26,27</sup>. Enhanced OA emissions are also expected from increasing local anthropogenic emissions, which include oil and gas exploration, and shipping activities<sup>28,29</sup>.

Attempts to model the Arctic OA concentrations have been limited, with typical under-predictions in winter and/or spring, when haze can be omnipresent and persistent<sup>30–33</sup>. The formation of secondary organic aerosols (SOAs) from anthropogenic or natural sources in different seasons is poorly represented<sup>32,34</sup>. Of 16 models deployed in a recent AeroCom evaluation of the simulated annual aerosol optical depth in Polar Regions<sup>32</sup>, only 6 considered biogenic precursors, which can contribute to particle growth and so the size range of cloud condensation nuclei. Only in one of the models was methanesulfonic acid (MSA) considered, which resulted in an outlier Arctic aerosol optical depth in terms of both seasonal variability and year-long magnitude<sup>32</sup>. A recent study showed that natural

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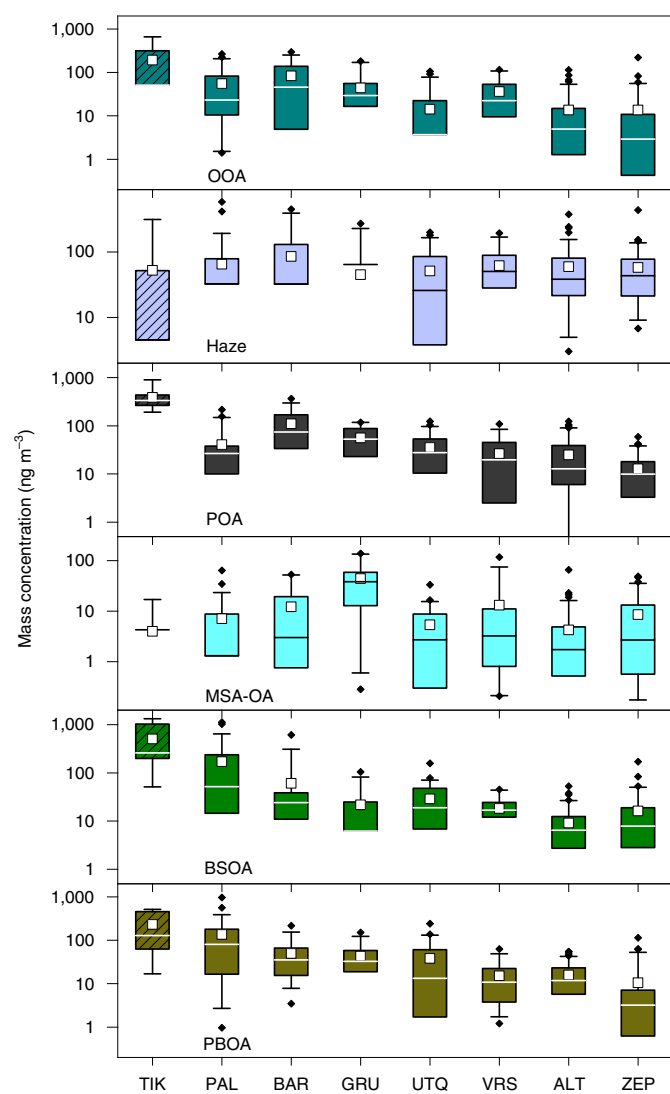
**Fig. 1 | Sites, OA factors and chemical characteristics.** **a**, Arctic political map showing the aerosol filter sampling stations (Supplementary Text 1 and Supplementary Table 1). A, Alert; B, Cape Baranov; G, Gruvebadet; P, Pallas-Matorova; T, Tiksi; U, Utqiagvik; V, Villum Research Station; Z, Zeppelin. Adapted from Hugo Ahlenius/GRID-Arendal (<https://www.grida.no/resources/8378>). **b**, Station-specific average total OA mass concentrations (whiskers are 1 standard deviation, s.d., corresponding to sample-to-sample variability) based on a statistical analysis of the water-soluble fraction to obtain factor recoveries (Supplementary Text 4) in the polar night (winter) versus midnight sun (summer) periods (Supplementary Table 2), sorted in descending order of the station annual average, and percent contribution to the particulate mass (including non-sea salt sulfate, nitrate, ammonium, EC and estimated sea salt<sup>63</sup>). The dashed blue lines connect winter and summer contributions at each station (no winter samples for G and T, and U winter samples were not analysed for ions). **c**, Van Krevelen plot as a tool for compositional differentiation among samples: atomic O:C ratio versus H:C ratio of the Arctic AMS-PMF-based factors (Supplementary Fig. 2), and individual PMF input bulk samples (colour coded by month: 1, January, through to 12, December). Red and blue dashed curves refer to the triangle reported by Ng et al.<sup>85</sup>. Grey dashed lines denote two example oxidation states (OS). Error bars correspond to 1 s.d. from a bootstrap analysis (Methods and Supplementary Text 4). **d**, Spatial distribution and seasonal variability in the average factor percentage contributions to total OA (water-soluble (Supplementary Fig. 7); entire time-series (Supplementary Fig. 10)). Factors are sorted from bottom (Haze) to top (POA) based on their onset (see Fig. 3), starting from late winter for Haze. Primary OAs, POA + PBOA (top). Orange outline, sum of natural-dominated OAs.

OAs other than MSA may account for about half of the summertime Arctic OAs<sup>33</sup>. Biological emissions, often not considered in Arctic models, may be a missing source of ice-nucleating particles<sup>2,35</sup> that is possibly further enhanced with the thawing permafrost<sup>36</sup>. The contribution of anthropogenic emissions, for example, from gas flaring<sup>37</sup>, to primary OA and SOA precursors is expected to vary substantially in space and time across the Arctic region but, unlike black carbon<sup>38</sup>, has been largely overlooked. Long-term ambient measurements of the Arctic OA composition are critically needed to identify its main sources and therefore are a first step for subsequent implementation into models. However, available studies<sup>3,39–41</sup> are single-site, short-term<sup>3,39</sup> or campaign-based<sup>39,40</sup>, and too infrequent to reveal seasonal patterns in the main sources of Arctic OAs<sup>42</sup>. Hence, Arctic aerosol source apportionment studies typically do not consider OAs<sup>43,44</sup>, or are limited to a few primary markers (for example, levoglucosan from biomass burning or EC

and radiocarbon measurements<sup>45,46</sup>. Therefore, current efforts have so far been unable to provide an understanding of the sources and formation pathways of the pan-Arctic OAs in different seasons.

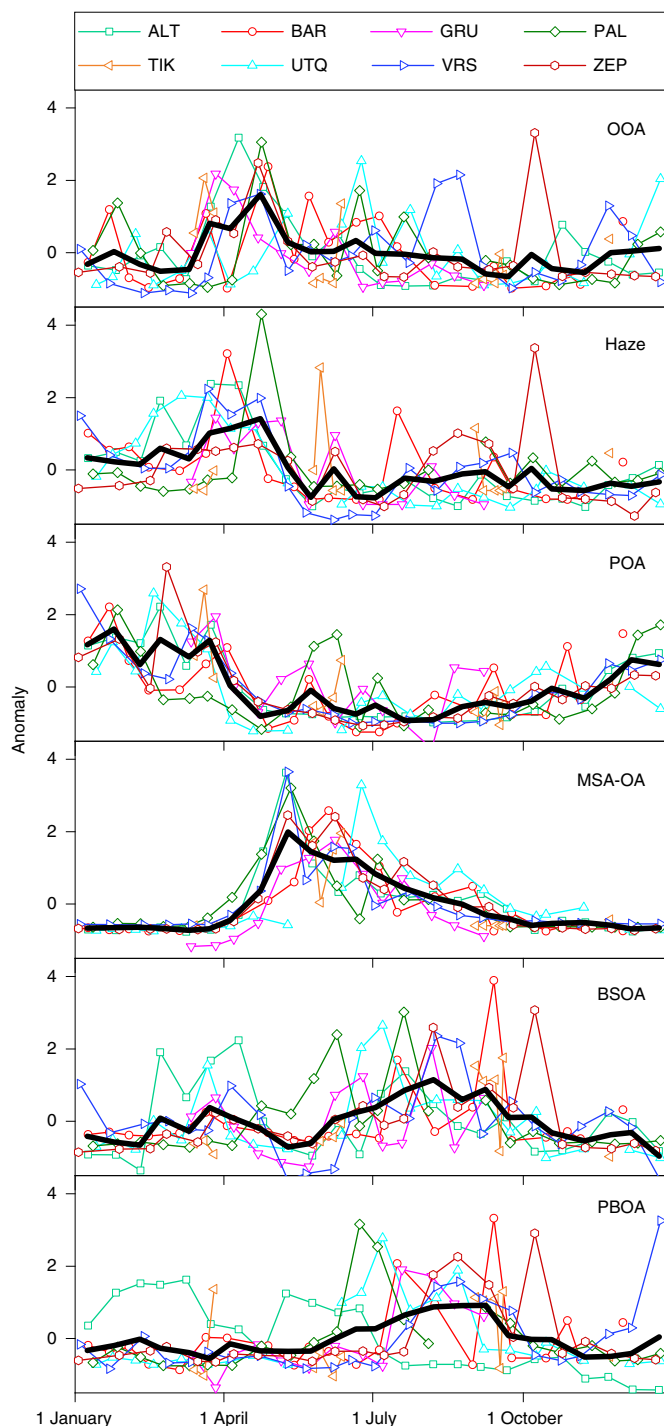
### Determination of OA sources across the Arctic

Here we fill this critical knowledge gap by determining the spatial and seasonal distribution of individual OA classes (factors) across the Arctic (Fig. 1, Methods and Supplementary Text 1). This was achieved by offline aerosol mass spectrometer (AMS) measurements of nebulized aerosol water extracts<sup>47</sup> from filter samples (Methods and Supplementary Text 2), followed by multisite positive matrix factorization (PMF) for OA source apportionment in relative terms, and subsequent quantification using externally measured water-soluble organic carbon (WSOC; see Methods). Factor identification is aided by organic marker, major ion and EC measurements (Methods and Supplementary Text 3). Source



**Fig. 2 | Station-specific yearly mass concentration of speciated Arctic OAs.** Absolute OA factor concentrations at the different stations shown as box-and-whisker plots. The stations appear in the same order as in Fig. 1b, whereas the factors appear in the same order as discussed in the main text. Note the different range of y-axis values for the different factors. The lower whiskers are missing if associated values are out of scale. Boxes for TIK are hatched to indicate incomplete (inter)annual coverage. For WSOAs, see Supplementary Fig. 8. Horizontal line and box, yearly median and interquartile range; squares and whiskers, yearly mean and range within the 5th and 95th percentiles; diamonds, outliers.

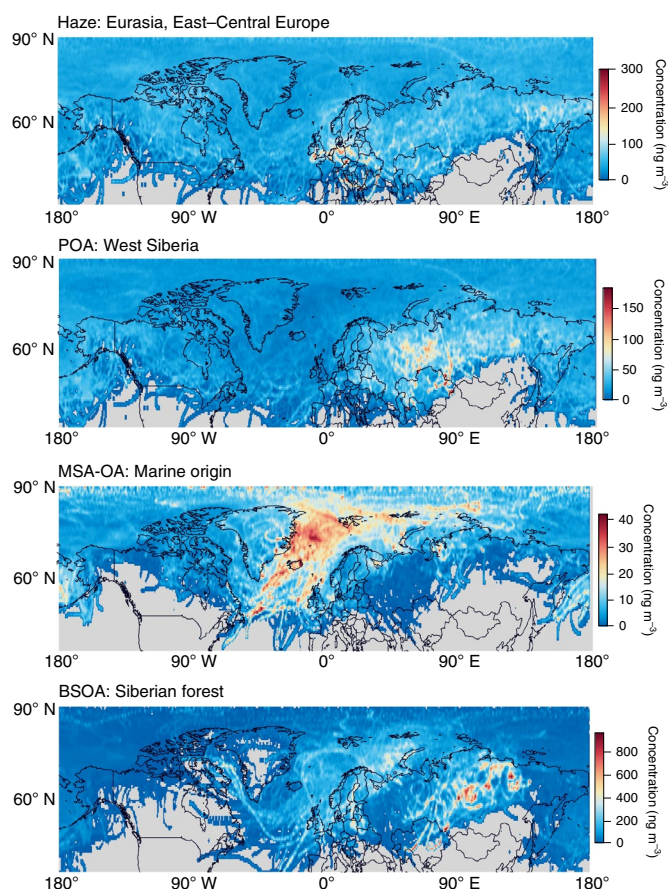
apportionment (Methods and Supplementary Text 4) results are also combined with concentration-weighted trajectory (CWT) analysis<sup>48</sup> to determine the geographical origin of the OA factors (Methods and Supplementary Text 5). We performed the analysis of 350 (bi-)weekly composite samples from filters collected at eight observatories across the Arctic (Fig. 1a) at roughly overlapping periods in 2014–2019 (Supplementary Table 1), yielding unique results of annual cycles which included the periods of winter darkness and summer daylight (Supplementary Table 2). Using this methodology, we determined the anthropogenic and natural sources that drive the mass of primary OAs and SOAs in the pan-Arctic region in both winter and summer. Our work is the first systematic, concerted and multiseason, multi-station effort with advanced analysis techniques, which goes beyond intensive observation periods and case-study work<sup>20</sup>.



**Fig. 3 | Seasonal variability of speciated pan-Arctic OAs.** Standardized annual cycles for each OA factor at the different stations. Bi-weekly averaged data from multiple years for each station are merged into a single annual cycle (the sum of individual station values in each panel equals zero). The y-axis values (anomalies) were calculated using the absolute mass concentration values as:  $(\text{value} - \text{station average}) / \text{s.d. of station}$ . The thick black lines indicate the average annual cycle of each factor over all the stations (note that here the sum of the yearly values in each panel is not equal to zero).

The total OA was found to be a major aerosol fraction that typically contributes 10–40% to the total particulate mass at the different stations (Fig. 1b; the dataset range is 3–65%), with higher relative contributions in the summer. Although only water-soluble





**Fig. 4 | Major source regions of long-range transported Arctic OA factors.** Merged results from the CWT-based back-trajectory (BT) analysis with ZeFir (Methods) at different Arctic stations (Supplementary Text 5 and Supplementary Fig. 11) showing long-term pan-Arctic hot spots of transported anthropogenic-dominated (Haze and POA) and natural-dominated (MSA-OA and BSOA) OA factors. The entire time series of each factor mass concentration at the different stations (time periods shown in Supplementary Table 1 and Supplementary Fig. 10) were used to create the maps (see Supplementary Text 5 for a discussion of the potential uncertainties in the source regions). The trajectories represent 5 days back in time for MSA-OA and (up to) 10 days for the other factors (Supplementary Text 5 and Supplementary Fig. 11). Colour scales indicate the water-soluble factor concentrations linked to the major source regions ('long-range' probability heat maps). The individual station results shown in Supplementary Fig. 11 were merged for each factor, except for POA, for which only six stations with winter data were considered here (no GRU and TIK), to indicate specific regions with intense gas-flaring activity during winter (for example, the Komi Republic, Khanty-Mansisk and Yamalo-Nenets autonomous districts in West Siberia). PBOA is expected to reside mainly in the coarse aerosol mode, and thus has a relatively short atmospheric lifetime (and hence more local and/or regional origins), and the formation of OOA might be linked to a prior accumulation of volatile organic compounds (thus probably not directly transported in the particle phase); therefore, the merged results for these factors are shown only in the Supplementary Information (Supplementary Fig. 11). The World Maps available with ZeFir are taken from Natural Earth Data.

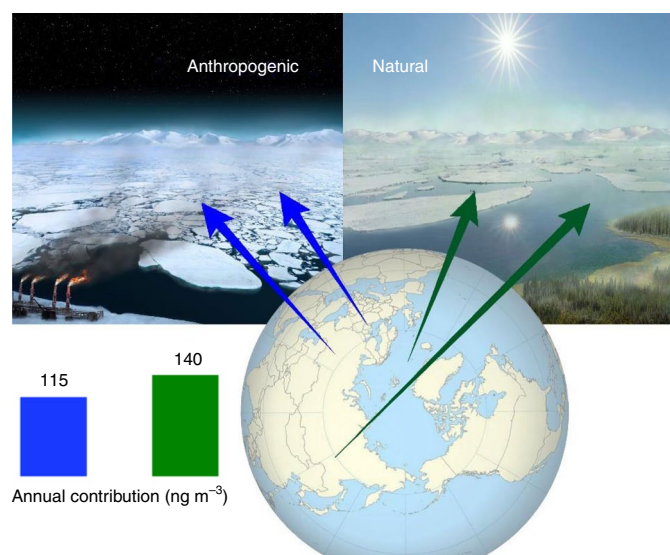
OA (WSOA) components were measured, the results represent the total OA by scaling the measurements by the water-soluble fraction of each factor (Methods and Supplementary Text 4). On average, WSOAs comprise  $82 \pm 4\%$  of the total OA. Absolute OA concentrations were generally higher at the more continental stations,

$\sim 1.4 \mu\text{g m}^{-3}$  in Tiksi (TIK),  $\sim 0.4 \mu\text{g m}^{-3}$  in Cape Baranova (BAR) and  $0.2 \mu\text{g m}^{-3}$  and  $1.0 \mu\text{g m}^{-3}$  in Pallas-Matorova (PAL) in winter and summer, respectively, whereas typical concentrations at the other stations were lower ( $0.1\text{--}0.2 \mu\text{g m}^{-3}$ ). Figure 1c shows the O:C versus H:C ratios for all the samples measured by the AMS, which highlights the large variability in the OA chemical composition across stations and seasons. We determined six major OA factors (Methods and Supplementary Text 4, Supplementary Tables 3 and 4 and Supplementary Figs. 1–10). These include factors related to anthropogenic-dominated emissions, namely, oxygenated organic aerosol (OOA), Haze and primary-anthropogenic organic aerosol (POA), and to natural-dominated emissions, namely, MSA-related organic aerosol (MSA-OA), biogenic secondary organic aerosol (BSOA) and primary biological organic aerosol (PBOA). We could not relate any factor to fresh wildfire emissions in the spring or summer, probably because of multiple reasons such as the rapid aerosol ageing during transport<sup>49</sup>, emissions remaining aloft on ascent to the middle and/or upper troposphere (if they originate from North America and East and South Asia)<sup>2,23,50</sup> and/or because this source becomes important only during short-term events. The atomic O:C versus H:C ratios of the six major OA factors (Fig. 1c) and their relative contributions across stations and seasons (Fig. 1d) are very distinct, and show the large diversity in the sources that drive the OA mass in the Arctic region. In the following, we discuss characteristic features of the factors, their spatial (Fig. 2) and seasonal (Fig. 3) variabilities, as well as their major source regions (Fig. 4).

#### Anthropogenic-dominated OA factors

The OOA factor (O:C  $\sim 0.5$ ) contains large oxygenated fragment ions (Supplementary Table 5). Its time-series shows a pan-Arctic enhancement during and after polar sunrise (Fig. 3), which potentially links this factor to oxidation products of volatile organic compounds that have accumulated during the polar night and form SOAs in the spring<sup>25,51–53</sup>. In this regard, a strong decrease in volatile organic compound concentrations was reported at PAL for April<sup>24</sup>. Although the OOA factor might be dominated by wintertime anthropogenic emissions, the contribution from natural sources during the summer cannot be excluded. The Haze factor contains more fragmented oxygenated ions (O:C  $\sim 0.75$ ; Fig. 1c), builds up in late winter, peaks in spring (Fig. 3) and correlates with sulfate (Supplementary Table 6), similar to the profile and timing of the Arctic haze phenomenon<sup>55</sup>. Except for intermittent peaks at BAR, PAL and TIK, its yearly concentration is spatially homogeneous (Fig. 2), but with a certain temporally structured variability<sup>56</sup> with the timing of the winter/spring peak occurring a few weeks later in time at most observatories as we move from west to east (Fig. 3), which indicates the effect of regional transport. This factor therefore becomes predominant in relative terms (Fig. 1d) at remote stations with lower OA loadings, that is,  $\sim 45\%$  throughout the year at Alert (ALT) and Zeppelin (ZEP) (1st and 3rd quartiles,  $Q_1\text{--}Q_3 = 38\text{--}54\%$ ). Haze organics are associated with long-range atmospheric transport from Eurasia (Fig. 4). Therefore, the transport of pollutants from distant urbanized areas to remote environments contributes substantially to the formation of pan-Arctic SOAs (OOA and Haze;  $Q_1\text{--}Q_3 = 44\text{--}56\%$  of the total OA during the polar night).

The POA factor is mainly composed of hydrocarbons (Supplementary Table 5) from primary anthropogenic emissions (O:C  $\sim 0.2$ ), most probably related to gas flaring. POA has an H:C ratio of  $\sim 1.3$  (Fig. 1c), lower than the H:C ratio ( $\sim 1.8\text{--}1.9$ ) of freshly emitted hydrocarbons detected at lower latitudes from traffic emissions<sup>57,58</sup>, which suggests a high contribution from unsaturated hydrocarbons. This factor is ubiquitous under dark and cold conditions, and peaks in December–March (Fig. 3) with a typical median pan-Arctic contribution to the total OA of 35% during the polar night (Fig. 1d,  $Q_1\text{--}Q_3 = 28\text{--}42\%$ ). In winter, POA correlates strongly with EC (Supplementary Fig. 12), which is reported to be of



**Fig. 5 | Conceptual overview of anthropogenic-dominated versus natural-dominated Arctic OAs.** A conceptual image of anthropogenic-dominated and natural-dominated emissions that drive the OA mass in the Arctic in winter and summer, respectively. The most important geographical source regions are indicated by the arrows. Bars show the entire-dataset average contributions of nearly equally contributing summed anthropogenic-dominated (blue) and summed natural-dominated (green) organic components. Credits: Helen Cawley for the landscape drawing; map made using Natural Earth.

fossil origin<sup>45,49</sup>. The POA:EC ratio of  $\sim 1.1$  is similar to the respective organic carbon (OC):EC ratios from oil and gas extraction emission estimates<sup>59</sup>. We found West Siberian locations (Fig. 4) as a major potential source region of the POA in winter (mainly at ALT, BAR, PAL and ZEP; Supplementary Fig. 11), similar to those previously found for surface black carbon<sup>42,60,61</sup>. The presence of POA at Gruvebadet (GRU) and TIK in the summer (Figs. 1d and Fig. 2) might indicate a more local origin at urban-type Arctic settlements.

### Natural-dominated OA factors

MSA-OA is characterized by the fragmentation pattern of MSA (for example, CH<sub>3</sub> and CH<sub>3</sub>SO; Supplementary Table 5), which is produced from marine dimethylsulfide oxidation. It correlates strongly with MSA, which comprises on average 80% of the factor, as measured by ion chromatography (Supplementary Table 7). This factor consistently originates from marine regions (Fig. 4), and increases with solar radiation (Supplementary Fig. 14). The MSA-OA absolute concentrations (Fig. 2) and clear annual cycles (Fig. 3) are consistent among the stations, with much higher concentrations in May–June, the main season of phytoplankton blooms. The range of maximum concentrations at ALT and Utqiagvik (UTQ) ( $\sim 20$ – $30$  ng m<sup>-3</sup>) is comparable to those of previous measurements of MSA from these stations by ion chromatography<sup>62,63</sup>. The highest weekly averaged MSA-OA levels, which exceed 100 ng m<sup>-3</sup>, occur at GRU, Villum Research Station (VRS) and ZEP. Although at these stations MSA-OA is typically 22% ( $Q_1$ – $Q_3$  = 12–34 %) of the total OA during the midnight-sun period (Fig. 1d), this factor is neither the only nor the predominant natural OA component.

BSOA is linked to biogenic emissions, for example, mono- and/or sesquiterpenes and isoprene<sup>64,65</sup>, from forests, tundra, lakes, wetlands and marine waters. This factor dominates the variability of moderately oxygenated CHO fragments related to isoprene and terpene SOAs or, potentially, to biogenic stress responses (Supplementary Table 5), and correlates strongly ( $R^2 = 0.81$ ; Supplementary Fig. 13)

with 3-methyl-1,2,3-butanetricarboxylic acid, a second-generation oxidation product of  $\alpha$ -pinene. BSOA exhibits a clear annual cycle with enhanced values in June–September (Fig. 3), consistent with the exponential increase of biogenic precursor emissions with temperature<sup>66</sup> (Supplementary Fig. 14). This factor dominates at PAL and TIK where it contributes, on average, 40% to the total OA in the summer (Fig. 1d,  $Q_1$ – $Q_3$  = 25–53%). At stations less affected by the boreal biome, BSOA appears in smaller but non-negligible amounts in the summer, and contributes 20%, on average, at UTQ and ZEP ( $Q_1$ – $Q_3$  = 11–28%) and  $\sim 9\%$ , on average, at other stations. At PAL, we expect a more local and/or semiregional source<sup>67,68</sup>, whereas the episodic occurrence of BSOA in both Russian stations is linked to northward air mass transport from the Siberian forest (Fig. 4).

PBOA (O:C  $\sim 0.4$ , H:C  $\sim 1.6$ ) is related to biological matter potentially from both terrestrial and marine origins, for example, fungal spores, bacteria, vegetative detritus, phytoplankton or fragments of it, and (ice) algae secretions<sup>1,69,70</sup>. It correlates with carbohydrate-related AMS fragments typically linked to primary biological compounds (Supplementary Table 5), and with arabitol and mannitol<sup>71</sup> ( $R^2 = 0.86$ ; Supplementary Fig. 13), which are sugar-alcohols present in fungal spores and other biological matter. PBOA exhibits substantial concentration increases in July–September, to reach up to  $1.0 \mu\text{g m}^{-3}$  at PAL (Fig. 2), and a distinct temporal evolution, peaking later than MSA-OA (Fig. 3). Similar to BSOA, PBOA exhibits higher relative contributions at lower latitude Arctic stations ( $\sim 70^\circ\text{N}$ ), averaging one-third of the total OA at PAL and UTQ in the summer (Fig. 1d,  $Q_1$ – $Q_3$  = 26–47%). PBOA anticorrelates with the snow depth at PAL and VRS (where such data are available) and with the 30-year climate-normal average snowfall at UTQ (Supplementary Fig. 14). These findings indicate that maximum local biological activity is associated with the lack of snow covering the ground<sup>72</sup>, which causes entrainment of PBOA into the air.

### Implications of Arctic OA spatiotemporal variability

Overall, the pan-Arctic OA composition and sources are not uniform. They are largely driven by the stations' latitude (and altitude; Supplementary Fig. 15), favourable conditions for long-range atmospheric transport and distance to anthropogenic and natural aerosol sources, as well as the presence of light and snow cover. We show that secondary Arctic OAs, separated into various distinct subtypes, typically dominate the OA mass, although the contribution of primary OAs from both natural and anthropogenic emissions is often equally important (Fig. 1d). This finding is in line with lower-latitude OA source apportionment studies<sup>12</sup>. The current and future abundance of condensable organics could be critical for the cloud condensation nuclei budget and cloud properties. In the Arctic winter, the sea-ice extent is at its maximum, low-level clouds have a pronounced warming effect and surface pollutants can become trapped close to the ground due to temperature inversion<sup>55</sup>. In this period, the land-surface OA sources are predominantly anthropogenic and linked to both primary emissions and secondary processes. The widespread Haze organics are dominated by aged anthropogenic-dominated emissions transported mainly from Eurasia, peak during polar sunrise and abruptly decrease in May. Under cold and dark conditions, all the stations are affected by OA emissions related to oil or gas extraction activities, mainly in West Siberia. The Arctic amplification of temperature increase<sup>73,74</sup> is more intense during winter<sup>75</sup>, and may be affected by aerosols altering the cloud properties. Hence, the future evolution of anthropogenic-dominated wintertime Arctic OA should be monitored from a climate perspective, especially in response to the development of effective emission control measures at lower latitudes<sup>11,29</sup>. In the summer, the decreasing anthropogenic pollution is replaced by natural OA emissions, with similar absolute concentrations. These emissions include marine SOAs from dimethylsulfide oxidation and little-explored primary biological emissions and biogenic SOAs. These collectively contribute to an increased



importance of natural OA in the summertime aerosol particle mass in the inner Arctic<sup>76</sup>. We found an overall nearly equal yearly abundance (Fig. 5) of summed anthropogenic-dominated OAs (average of 115 ng m<sup>-3</sup>, Q<sub>1</sub>–Q<sub>3</sub> = 20–100 ng m<sup>-3</sup>; 95th percentile, 435 ng m<sup>-3</sup>) and summed natural-dominated OAs (average of 140 ng m<sup>-3</sup>, Q<sub>1</sub>–Q<sub>3</sub> = 40–160 ng m<sup>-3</sup>; 95th percentile, 445 ng m<sup>-3</sup>). This indicates that the typically lower total aerosol volume/mass in the summer versus winter or spring<sup>77</sup> is due to species other than the total OA, which exhibits less of a seasonal cycle across the Arctic (Supplementary Fig. 16). The effects of anthropogenic and natural organics on cloud condensation nuclei and ice-nucleating particles, and potentially on the Arctic climate, have been largely unexplored. The importance of considering land coverage and biosphere–atmosphere exchanges<sup>78,79</sup> for the response of natural Arctic OAs to warming is highlighted in Supplementary Fig. 14. A small increase in temperature results in a substantial (exponential) increase of BSOA, which identifies one of the potential feedback mechanisms in the Arctic. Also, extension of the ice-free season and expansion of snow-free areas<sup>80</sup> may lead to enhanced airborne organics from biological activity and secondary marine emissions.

Currently, the Arctic system is in transition<sup>21</sup>, with long-range transported anthropogenic-dominated emissions (including sulfate) continuously decreasing due to better air quality regulations in the lower latitude regions. Meanwhile, natural emissions are expected to increase<sup>81,82</sup>, which probably enhances the magnitude and relative importance of the composition-dependent OA–cloud effect<sup>11</sup>. Our results provide the first understanding of the present-day year-long, pan-Arctic OA sources, which can be used for comparisons with past (for example, through ice-core archives<sup>83</sup>) and future measurements of these changing biogenic and anthropogenic emissions. Given practical difficulties in deploying multiple online AMS instruments for long time periods around the Arctic and the widespread availability of ambient filters, the measurement methodology and analysis techniques employed here are also applicable to emerging Arctic stations (for example, in far East Siberia<sup>84</sup>).

### Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at <https://doi.org/10.1038/s41561-021-00891-1>.

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## Methods

**Filter sampling.** Total suspended particulate matter or PM<sub>10</sub> (particulate matter with aerodynamic diameter  $d < 10 \mu\text{m}$ ) was collected on quartz fibre filters at eight stations around the Arctic (Fig. 1a). This study combined distinct locations affected by anthropogenic versus natural, marine versus terrestrial and local versus long-range transported emission sources at different seasons. PAL stands out from the other stations as it is in a sub-Arctic site influenced substantially by the boreal biome, which highlights differences with higher-latitude sites. Typically, at least one annual cycle is covered per station (Supplementary Table 1). The samples from GRU cover the spring and summer season (Supplementary Table 2) over two consecutive years, whereas the longest temporal coverage is almost four years (ALT). Although the data coverage at TIK is relatively less wide, this site experiences distinctly higher levels of pollution compared with that of other sites and includes the spring, summer and autumn periods over three consecutive years, and so covers a wide range of conditions. More details about the stations and filter sampling are provided in Supplementary Text 1, which includes a description of the conditions under which the filters were handled, transported between stations and/or labs and stored during and/or after sampling.

**AMS measurements and data analysis.** The offline AMS technique was established by Daellenbach et al.<sup>47</sup>. Briefly, punches from the quartz fibre filter samples were extracted in ultrapure water (18.2 Mohm cm, with the total organic carbon (TOC)  $< 3 \text{ ppb}$  by weight). Teflon filters were tested but not measured (Supplementary Text 2). Typical quartz-fibre-filtered water-extracted organic concentrations were  $2\text{--}3 \mu\text{g C ml}^{-1}$ . The extracts were inserted into an ultrasonic bath for 20 min at  $30^\circ\text{C}$ . Each sonicated sample was then filtered through a nylon membrane syringe (0.45  $\mu\text{m}$ ; Infocroma AG) and transferred to a 'Greiner' sample tube (50 ml). From the obtained solutions, aerosols were generated in synthetic air (80% volume N<sub>2</sub>, 20% volume O<sub>2</sub>; Carbagas) via an apex Q nebulizer (Elemental Scientific, Inc.) operated at  $60^\circ\text{C}$ , dried by a Nafion dryer and directed into a long-time-of-flight AMS. Each sample was recorded for 480 s, with a collection time for each spectrum of  $\sim 40$  s. Ultrapure water was measured for 720 s before and after each sample measurement. The technique was performed on  $\sim 370$  extracts (samples and field blanks) in total.

For the data analysis, we used Squirrel v1.59B for the  $m/z$  calibration and baseline subtraction, and Pika v1.19B for high-resolution (HR) analysis, in the Igor Pro software package 6.37. The HR peak fitting was performed in the  $m/z$  range 12–191. After the peak fitting (fragments consisted of C, O, H, N and S), the isotope ions (and the inorganic fragments) were removed, and the average water-blank signal was subtracted from the average signal of the following sample. The resulting data matrix was the organic fraction mass spectra time series (normalized fragment ion intensity) and an error matrix that included the blank variability and measurement uncertainties. The signal-to-noise ratio (was  $> 2.0$  for 390 out of 578 fitted organic fragment ions with  $m/z$  up to 133, whereas it dropped to 1.9 on average for fragments with  $m/z$  between 134 and 180. The inorganic-salt artefact on the AMS CO<sub>2</sub> (ref. <sup>86</sup>) and CO fragment ion signal was also accounted for (Supplementary Text 2), but the resulting corrections were minor.

**Auxiliary measurements.** Additional offline analyses were carried out (Supplementary Text 3); these included the measurement of EC and OC by a thermo-optical transmission method using a Sunset analyser, WSOC by a TOC analyser, major water-soluble ions (including MSA) by ion chromatography and organic markers (sugar-alcohols, sugars and organic acids) for selected samples. Several environmental parameters were retrieved as well, such as temperature, solar radiation and snow-depth data, which were averaged to match the time resolution of the filter sample composites measured by AMS (Supplementary Text 3).

**AMS-PMF analysis. Concept.** The normalized organic mass spectra from the offline AMS measurements were analysed by PMF<sup>87,88</sup> using the multilinear engine 2<sup>90</sup>. The aim was to derive source components (factors) that can be linearly combined to reproduce the observed time and chemical (mass spectral profiles) variations<sup>90</sup> in the extracted organics. The separation of OA factors was therefore based on differences in their chemical composition and temporal behaviour, regardless of their original mixing state with other particulate matter components when suspended in the atmosphere. We did not use the absolute AMS signals for the quantification of the organic fraction, as they are affected by the AMS collection and transmission efficiencies and by the nebulization efficiency. Instead, we scaled the relative factor contributions determined by PMF using the TOC-based WSOC (see 'Final AMS-PMF results'). Therefore, the total OA concentration is not affected by the collection efficiency. All extracted aerosol species are expected to be internally mixed in the nebulized particles, so the offline AMS collection efficiency should be the same for all identified OA factors<sup>91,92</sup>.

**Methodology.** The interface for the data processing was provided by the source finder toolkit<sup>93</sup> (SoFi version 6.86) for Igor Pro (WaveMetrics, Inc.). PMF attempts to solve the bilinear matrix equation,  $X_{ij} = \sum_n (G_{in}F_{nj} + E_{ij})$ , by following the weighted least-squares approach. In the case of AMS,  $i$  represents the time index,  $j$  is the fragment and  $n$  is the factor number. If  $X_{ij}$  is the matrix of the organic mass spectral data and  $s_{ij}$  the corresponding error matrix (including the blank

variability and measurement uncertainties),  $G_{in}$  is the matrix of the factor time series,  $F_{nj}$  the matrix of the factor profiles and  $E_{ij}$  the model residual matrix, then PMF determines  $G_{in}$  and  $F_{nj}$  such that the ratio of the Frobenius norm of  $E_{ij}$  over  $s_{ij}$  is minimized. The allowed  $G_{in}$  and  $F_{nj}$  are always non-negative. The PMF input matrices ( $X_{ij}$  and  $s_{ij}$ ) here included data from all the stations ( $\sim 350$  samples) and 578 HR fragments with  $m/z$  up to 133 (or 1,029 HR fragments with  $m/z$  up to 191). All fragments with a signal-to-noise ratio below 0.2 were removed from the matrices, and those with a signal-to-noise ratio below 2.0 were down-weighted according to the recommendations of Paatero and Hopke (2003)<sup>94</sup>. For factor identification<sup>90</sup>, we used a combination of criteria where applicable. These include the factor seasonal cycle, fragmentation pattern, characteristic fragments, time series correlation with external markers, time series correlation with environmental parameters and the BT analysis.

**Number of factors.** Unconstrained PMF was performed for  $n = 3\text{--}15$  factors to choose a 'base case' solution before the uncertainty analysis. Five random seed runs were performed for each  $n$  (65 runs in total). Preliminary diagnostics were then produced (Supplementary Fig. 1) to investigate the optimum  $n$  based on the explained variability of the input matrix ( $Q/Q_{\text{exp}}$  (exp, expected), scaled residuals) and the stability and/or interpretability of preliminary solutions among different runs for each  $n$ . All random seed runs provided essentially identical results (that is, the lowest  $Q/Q_{\text{exp}}$  relative s.d.) only for the 11-factor solution (the factor profiles are shown in Supplementary Fig. 2). The stability of this preliminary solution only (versus other  $n$ ) among the different random seed runs is detailed in Supplementary Table 3. The  $Q/Q_{\text{exp}}$  of this average ('base case') 11-factor solution exhibited a random pattern in both dimensions (time series and variables) of the reconstructed PMF output matrix (350 samples and 578 HR fragments up to  $m/z$  133). With regard to a partial exploration of the rotational ambiguity, the 11-factor solution was the most robust (Supplementary Text 4), as other factor solutions were less stable among the different random seed runs compared with that of  $n = 11$  for certain factors (Supplementary Table 3).

**PMF errors.** The PMF model statistical and rotational uncertainty was assessed via a bootstrapping (BS) approach (Supplementary Text 4) applied to the entire dataset (by including PMF samples from the all stations and seasons). The BS approach generated new input matrices by randomly resampling samples from the original input matrix. Each newly generated PMF input matrix had a total number of samples equal to those of the original matrices, although some of the original filter samples were represented several times, and others were not represented at all. The resulting error interval represents temporal variations of the 11-factor profiles, random errors and errors in the modelling process, such as rotational ambiguity and a mis-specified number of factors<sup>95</sup>. Note that a preliminary BS analysis was performed also for the 10-factor solution, but the results were less stable than those of the 11-factor solution.

**Retention of PMF factors related to ambient organic aerosols.** Although the optimum PMF solution contains 11 factors, only the 6 factors presented in the main text and figures were interpreted to be related to the sampled OA. The remaining factors were associated with carbonate and (field) blanks (Supplementary Text 4). The carbonate-related (inorganic) factor may originate from sea spray and/or transported dust from lower latitudes or inner-Arctic sources. It exhibited an unusually high O:C of  $\sim 1.9$  due to a very high  $f_{\text{CO}_2}$  of around 0.8 (Supplementary Fig. 2). We confirmed the presence of carbonate by HCl fumigation and offline AMS experiments for selected samples<sup>66,97</sup>. All the data presented are corrected for the carbonate (Supplementary Text 4). We separated a factor that was systematically associated with the filter substrate in terms of both the mass spectral fingerprint (factor profile) and the absolute mass concentrations at the different stations (Supplementary Text 4) and three other factors with contributions that could not be decoupled from the background signal of the filter (Supplementary Text 4 and Supplementary Fig. 4).

**Final AMS-PMF results.** The atomic ratios (O:C, H:C, N:C and S:C) and OM:OC ratios shown in Supplementary Fig. 2 were calculated for all the PMF input samples and for each AMS-PMF factor (for  $m/z$  up to 133), using the Analytical Procedure for Elemental Separation version 1.05 within Igor. The error bars in Fig. 1c correspond to 1 s.d. from the 100 BS runs. The median and interquartile range of the factor-specific ( $k$ ) fractional contribution ( $F_{c_k}$ ) time series that resulted from the BS AMS-PMF analysis were converted into absolute WSOA mass concentrations by multiplying with the AMS-based (WS)OA:(WS)OC ratios of the different PMF-input samples (on carbonate subtraction) times the corresponding TOC-analyser-based WSOC values (total dissolved OC mass without any field blank subtraction):  $\text{WSOA}_k = F_{c_k} \times \text{WSOC} \times (\text{WSOA}/\text{WSOC})$ .

**Recovery analysis performed using PMF.** Recovery analysis was performed using PMF, following a simplified version of the approach of D. Bhattu et al. (personal communication; see Supplementary Text 4). The lowest water solubility was  $\sim 60\text{--}80\%$  for POA, PBOA, Haze and OOA, whereas MSA-OA and BSOA can be considered fully water-soluble (Supplementary Text 4 and Supplementary Fig. 9). The AMS-PMF-based WSOA ( $x$ , the sum of six WSOA factors) versus the recovery-corrected total OA mass ( $y$ , the sum of six OA factors) time series



were correlated with an  $R^2$  of 0.99 and a slope of 1.16, which indicated that the majority of OAs are water-soluble (the remaining factors were subtracted from both the  $x$  and  $y$  variables). The reconstructed AMS-PMF-based OC mass ( $x$ , the sum of all factors) versus the measured sunset OC ( $y$ ) were correlated with an  $R^2$  of 0.87 and a slope of 1.14, which indicated a sufficient closure within the uncertainties. Although the elemental ratios shown in Supplementary Fig. 2 apply to water-extracted factors, the water solubility of Arctic OAs in this work was high (average  $\pm 1$  s.d. =  $82 \pm 4\%$ ), in agreement with previous estimates<sup>98</sup>, and does not vary significantly between locations and seasons. As a result, the data for the total OA shown in Figs. 1b,c and 2 are similar to those obtained for WSOA (Supplementary Figs. 7 and 8). The corresponding (median) total OA factor time series (Supplementary Fig. 10) were provided on conversion from WSOA values by applying median factor-specific recoveries, and were used as the result with regard to the factor time series. The median time series correlated highly with the base-case-solution time series (Supplementary Table 4).

**Source-marker AMS fragments.** We provide in Supplementary Table 5 specific fragments identified in our dataset as characteristic of specific sources, which were also identified in previous studies. These fragments were selected based on their highest contribution to these factors and the dominant contribution of these factors to these fragments.

**Uncertainties.** We briefly discuss here the PMF errors, the errors on the recovery, the errors on the relative ionization efficiency and the errors due to negative artefacts. The BS analysis indicated relative errors below 30% on average for median factor mass concentrations  $> 50 \text{ ng m}^{-3}$ , with MSA-OA clearly being the least uncertain factor (Supplementary Text 4). The interquartile range of the median time series corresponds to errors from the BS runs without contingency for errors from the recovery analysis, by applying constant WSOA-to-OA factor-specific (median) conversion factors. We used the same relative ionization efficiency for all OA fractions. Given the recently reported relative ionization efficiencies for organic compounds of 1.2–1.8 (refs. 99–101), we expect the uncertainties that result from this simplification to be on the order of 20–30%, comparable to our modelling error estimates. Although the positive filter artefact is probably captured in this work through the identification of the field blank-related factor, the negative filter artefact (loss of semivolatiles organics) could not be extensively assessed due to the current routine sampling strategies. In future analyses, it would be useful to address the negative filter artefact and other effects related to sample treatment, such as the effect of filter storage and revolatilization on water extraction, and to compare the offline results with concurrent online measurements.

**BT analysis for OA factor geographical origin assessment.** BTs show the air mass history (origins and transport paths) and thus can provide information on the geographical location of potentially advected emissions at large geographical scales. Here, BT analysis (Supplementary Text 5) was performed to assess potential source locations of individual Arctic organic components or their precursors over the entire time period covered at each station. The trajectories were calculated backward for up to ten days using the HYSPLIT4 model with meteorological data from the Global Data Assimilation System with  $1^\circ$  resolution (GDAS1; <ftp://arlftp.arl.hq.noaa.gov/pub/archives/gdas1>). The station coordinates correspond to the arriving location of each BT. We weighted the calculated BTs with the AMS-PMF-based factor time series using the CWT model to localize the air parcels responsible for high measured concentrations at the receptor site. For the CWT analysis, the Igor-based user interface 'ZeFir' was used<sup>48</sup>. Although the HYSPLIT trajectory module within ZeFir is currently limited to the use of regular GDAS meteorological field data files, it offers the option to run the model at any date and/or time (for example, an entire year). The average mass concentration of each factor in each sample was enlarged in ZeFir for 2, 7 or 14 days back in time (resolution of 6h, except for the Russian stations at 4h), depending on the average sampling time resolution at each station (for example, a 14 day enlargement for a bi-weekly sampling time resolution), and the corresponding enlarged BT results were averaged (within the software) to represent each individual sample. Air parcels rising at altitudes above 3 km a.g.l. were considered less likely to arrive at the receptor site and hence were discarded (a threshold altitude of 1.5 km did not produce substantially different results unless otherwise noted). We assessed the sensitivity of our results to the BT timescales showing that the resulting maps are robust and not too sensitive beyond certain lengths of time (Supplementary Text 5). Multisite merging is a powerful option in ZeFir to explore the bigger picture of source–receptor approaches, as combining several sites together leads to higher trajectory density values. Merged results (without normalization) are presented in Fig. 4 to indicate pan-Arctic hot spots with a greater accuracy than that of single-site results. No weighting function was applied to include the influence of less-frequent air mass trajectories from distant regions ('long-range' probability heat maps<sup>102</sup>). The main observations from individual station results are described in Supplementary Text 5.

## Data availability

The data used in this study<sup>103</sup> is openly available at Zenodo (<https://doi.org/10.5281/zenodo.5775070>). Please contact the corresponding authors when using the data.

## Code availability

Datalystica Ltd is the official distributor of SoFi Pro licenses. Origin was used for graph plotting. The open-source Igor-based package ZeFir is free to download at <https://sites.google.com/site/zefirproject/download>. The World Maps available with ZeFir are taken from Natural Earth Data.

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### Author contributions

I.E.H. and V.M. conceptualized the study. V.M. and H.L. performed the AMS measurements. V.M. performed the AMS, PMF and CWT analyses. K.D., D.B. and R.C. assisted with AMS data analysis. F.C. and K.R.D. contributed codes to the offline analysis

and SoFi. P.R. assisted with sample extractions for AMS measurements. L.H., W.Z. and S.S. provided EC, OC and ion chromatography data for ALT. O.P. coordinated the filter sampling and provided ion data from BAR and TIK. M.V. set up the filter sampling system at PAL. R.J.S. and C.E.M. organized filter collection at UTQ. C.E.M. performed ion chromatography for UTQ. A.M. and H.S. coordinated the filter sampling at VRS. J.K.N. analysed the OC and EC data from VRS. K.E.Y. and W.A. are responsible for the collection of aerosol filter samples at ZEP. J.-L.J. conducted analyses for EC, OC and dissolved organic carbon, carried out ion chromatography, high performance liquid chromatography-pulsed amperometric detection and liquid chromatography-mass spectrometry measurements on filters from the different stations that provided the concentrations for tracers, which included MSA, sugar-alcohols and organic acids. I.E.H., J.S., A.S.H.P. and U.B. supported and supervised the research. V.M., I.E.H., J.S. and U.B. interpreted the results. V.M., I.E.H. and J.S. prepared the figures and wrote the manuscript. All the co-authors commented on the manuscript.

### Competing interests

The authors declare no competing interests.

### Additional information

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