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## Springtime Arctic haze contributions of submicron organic particles from European and Asian combustion sources

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[1] The composition of Arctic aerosol, especially during the springtime Arctic haze, may play an important role in the radiative balance of the Arctic. The contribution of organic components to Arctic haze has only recently been investigated. Because measurements in this region are sparse, little is known about organic particle composition, sources, and concentrations. This study compares springtime measurements in the Arctic regions north of the Atlantic (ICEALOT, 2008) and Pacific (Barrow, Alaska, 2008 and 2009) oceans. The aerosol organic functional group composition from Fourier transform infrared (FTIR) spectroscopy combined with positive matrix factorization (PMF) and elemental tracer analysis indicate that mixed combustion sources account for more than 60% ( $>0.3 \mu\text{g m}^{-3}$ ) of the submicron organic mass ( $\text{OM}_1$ ) for springtime haze conditions in both regions. Correlations with typical combustion tracers (S, Zn, K, Br, V) provide evidence for the contribution of combustion sources to the Arctic  $\text{OM}_1$ . However, the two regions are influenced by different urban and industrial centers with different fuel usage. High-sulfur coal burning in northeastern Europe impacts the northern Atlantic Arctic region, while oil burning and forest fires in northeastern Asia and Alaska impact the northern Pacific Arctic region. Quadrupole and High Resolution Aerosol Mass Spectrometry measurements confirm the highly oxygenated nature of the  $\text{OM}_1$ , with an oxygenated organic aerosol (OOA) spectrum from PMF. High co-emissions of sulfate and organics from coal-burning in northeastern Europe produce significant concentrations of organosulfate functional groups that account for 10% of  $\text{OM}_1$  measured by FTIR spectroscopy during ICEALOT. These observations provide preliminary support for a heterogeneous mechanism of organosulfate formation on acidic sulfate particles.

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### 1. Background

[2] Arctic haze has been studied since it was reported by pilots over 50 years ago [Greenaway, 1950; Mitchell, 1956]. This haze is comprised mostly of sulfate aerosol, primarily from anthropogenic sources, with small amounts of ammonium, nitrate, dust, organics, and black carbon [Li and Barrie, 1993; Quinn *et al.*, 2002]. Significant anthropogenic sulfur (S) emissions exist within the Arctic region [Hole *et al.*, 2009]. On time scales of a few days to weeks, the polar

dome isolates the Arctic lower troposphere from warmer regions to the south making northern Europe and northern Russia major contributors to Arctic haze [Law and Stohl, 2007]. However, emissions from populated sub-Arctic regions are the most likely cause of the high aerosol particle concentrations in the springtime Arctic, when meridional transport from the midlatitudes to the Arctic intensifies and brings emissions from Eurasia northward [Carlson, 1981; Iversen, 1984; Iversen and Joranger, 1985]. There is more rapid transport from Europe and northern Asia to the Arctic [Stohl, 2006] than from more southern regions [Hole *et al.*, 2009] or North America. Measurements of trace metals from combustion (Mn, V) have been used to confirm the contributions of European sources to aerosol in the Arctic [Rahn, 1981]. Subsequent studies of the springtime Arctic haze have also linked contributions of sulfate aerosol to these metal tracers [Quinn *et al.*, 2009; Rahn and McCaffrey, 1980].

[3] Emissions from fuel-burning activities, including industry, electricity production, and domestic heating, have

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been identified as contributors to Arctic haze by chemical analyses of Arctic aerosol particles [Ottar *et al.*, 1986; Rahn and McCaffrey, 1979; Rahn *et al.*, 1977, 1989]. Chemical tracers for these types of anthropogenic emissions, such as sulfate, black carbon, Pb, Cd, Cu, and Zn, were found in Greenland snow [Candelone *et al.*, 1996; Davidson *et al.*, 1993; Masclet *et al.*, 2000], suggesting similar types of emission sources for these species and the constituents of Arctic haze. During the winter and early spring, fossil fuel combustion is the main source of sulfate to the Arctic since there is minimal biological activity [Ferek *et al.*, 1995]. Anthropogenic emissions from conurbations, such as Murmansk on the Kola peninsula, industrial emissions from the northern part of Russia, oil industry, and shipping all contribute to haze particles [Hole *et al.*, 2006a]. The Kola peninsula, in addition to other parts of northern Russia, has many Ni, Cu, and other non-ferrous metal producing industries, which also contribute to anthropogenic emissions [Hole *et al.*, 2006b; Weiler *et al.*, 2005].

[4] Recent evidence has shown that organic compounds comprise more than 30% of the submicron PM (PM<sub>1</sub>) in Arctic Haze [Shaw *et al.*, 2010]. Aerosol haze particles are thought to be chemically aged, very efficient at scattering solar radiation, and possibly weakly absorbing of solar radiation [Law and Stohl, 2007]. Identifying the sources of the haze and determining its chemical composition is necessary to define and mitigate its climate impact on both the shortwave and longwave radiation balance in the Arctic [Quinn *et al.*, 2007]. Here, we use springtime measurements from the International Chemistry Experiment in the Arctic Lower Troposphere (ICEALOT) and from Barrow, Alaska, to characterize and contrast the composition and identify the sources of organic components that contribute to haze from the Atlantic and Pacific sides of the Arctic.

## 2. Methods

[5] As part of the 2008–2009 International Polar Year, submicron particles were measured aboard the R/V *Knorr* during ICEALOT (19 March 2008 to 24 April 2008) over the North Atlantic and Arctic Oceans and the Greenland, Norwegian, and Barents Seas (20°–30°W and 41°–81°N). Submicron particles were collected from an isokinetic sampling inlet 18 m above sea level that extended 5 m above the aerosol measurement container. The inlet was heated to dry the aerosol to a relative humidity (RH) of 25% or less, during sampling. Sector control for the flow was used to prevent sampling emissions directly from R/V *Knorr*. From 1 March 2008 to 1 March 2010, similar measurements were made at the National Oceanic and Atmospheric Administration Earth System Research Laboratory (NOAA ESRL) observatory at Barrow, Alaska, with an analogous collection method described by Shaw *et al.* [2010] for the 2008 samples. The Barrow sampling site is at a coastal location (156.6°W, 71.5°N) in the northern Pacific side of the Arctic. Sector control is used at Barrow to avoid emissions from the local town. This stops the sample collection when the wind direction is between 0 and 130 degrees, which reduced the sampling time by 35% for the 221 collected filters in this study.

### 2.1. Filter-Based Particle Collection

[6] During ICEALOT, submicron particles were collected on 37 mm Teflon filters for 12 and 24 h periods using the previously described isokinetic inlet. At Barrow, two years of near-continuous submicron particle measurements were collected using an automated rotating filter holder that exposed 47 mm Teflon filters to ambient air drawn into the warmed inlet at 30 L min<sup>-1</sup> [Quinn *et al.*, 2002; Shaw *et al.*, 2010]. Filters at Barrow were exposed for different durations depending on season to accommodate changes in overall mass loadings: spring and winter filters were exposed for 24 h and summer filters for 96 h. After collection, the filters from both Barrow and the ICEALOT cruise were frozen, to prevent losses due to desorption or reaction, and transported back to San Diego for analysis by Fourier transform infrared (FTIR) spectroscopy. The FTIR spectra were analyzed using an automated algorithm that includes baselining, peak-fitting, and integrating at specific peak locations to quantify organic functional group mass associated with major carbon bond types based on the method outlined by Maria *et al.* [2002] and revised by Russell *et al.* [2009a]. Filters with detectable absorption at 876 cm<sup>-1</sup> were rinsed with 2 mL of hexane, which removes the organosulfate containing compound without removing bisulfate or carbonate that also absorb in this region [Blando *et al.*, 1998; Maria *et al.*, 2003]. After the rinse, there was no absorbance remaining at 876 cm<sup>-1</sup>, indicating efficient removal of organosulfate. Concentrations were calculated by dividing the mass of each functional group measured for each filter by the total volume of air sampled through the corresponding filters. Functional groups that were quantified include: hydroxyl (including alcohol, C–OH), alkane (C–CH), amine (C–NH<sub>2</sub>), carbonyl (C=O), carboxylic acid (COOH), and organosulfate (COSO<sub>3</sub>). Aromatic, alkene (C=CH), and organonitrate functional groups were below the detection limit for all filters discussed here.

[7] Prior to rinsing with hexane, X-ray Fluorescence (XRF; Chester LabNet, Tigard, Oregon) was performed on the FTIR analyzed filters to quantify the elemental mass concentrations of elements Na and heavier [Maria *et al.*, 2003]. For ICEALOT, S, K, Br, Ca, and Zn were measured above detection limit for 75% of the samples, Fe, Na, and Cl were above detection limit for 50% to 70%, and V and Si were above detection limit for between 35% and 40% of the samples. The remaining measured elements were above the detection limit for less than 15% of the samples and are not used in this analysis. Of the 221 filters collected from Barrow, a subset of 47 was analyzed by XRF for which Na, Cl, Fe, S, Br, K and Zn were consistently above detection limits [Shaw *et al.*, 2010]. The concentration of inorganic oxidized material (IOM – dust and fly ash) was calculated from the mass concentrations of Al, Si, Ca, Fe, and Ti assuming that each element was present in the aerosol in its most common oxide form [Bates *et al.*, 2008]. Because Al was below detection limit for all but one of the samples where V was above detection limit, the measured V was identified as non-crustal.

[8] Submicron particles (RH < 25%) were also collected with a Berner-type cascade impactor on Millipore Fluoropore filters downstream of a jet plate with a 50% aerodynamic cutoff diameter of 1 μm on the R/V *Knorr*. 1 mL of

spectral grade methanol was added to the filters followed by 5 mL of distilled, deionized water. The substrates were extracted by sonicating for 30 min. They were then analyzed by ion chromatography (IC) to quantify inorganic ions including sodium ( $\text{Na}^+$ ), chloride ( $\text{Cl}^-$ ), sulfate, nitrate, and ammonium [Quinn *et al.*, 1998]. Non-sea salt  $\text{SO}_4^{2-}$  concentrations were calculated from  $\text{Na}^+$  concentrations and the ratio of  $\text{SO}_4^{2-}$  to  $\text{Na}^+$  in seawater [Quinn *et al.*, 2000]. Sea salt concentrations were calculated from the measured concentrations of  $\text{Na}^+$  and  $\text{Cl}^-$ , in order to take into account  $\text{Cl}^-$  depletion in the atmosphere [Bates *et al.*, 2008].

## 2.2. Aerosol Mass Spectrometry

[9] A quadrupole aerosol mass spectrometer (Q-AMS) was used to measure submicron particle composition averaged over 5-min intervals aboard the R/V *Knorr*. Non-refractory ions (chemical components that vaporize at 600°C) including sulfate ( $\text{SO}_4^{2-}$ ), nitrate ( $\text{NO}_3^-$ ), and ammonium ( $\text{NH}_4^+$ ), were measured by Q-AMS, in addition to OM [Allan *et al.*, 2003; Jayne *et al.*, 2000]. A collection efficiency of 1 was used for all species described here based on the high acidity (low molar ratio of ammonium to sulfate) of measured particles and high relative humidity during measurements [Matthew *et al.*, 2008; Quinn *et al.*, 2006]. Comparisons of sulfate and ammonium concentrations measured by the Q-AMS and separately by IC of filter extracts were used to confirm the collection efficiency [Hawkins *et al.*, 2010; Russell *et al.*, 2010]. Sulfate measured by the Q-AMS was higher than that measured by IC (slope = 1.42,  $r = 0.82$ ) suggesting sufficient collection by the Q-AMS. Ammonium measured by both instruments compared well (slope = 0.99,  $r = 0.75$ ) further supporting a collection efficiency of 1. The total submicron OM ( $\text{OM}_1$ ) measured by FTIR exceeds the  $\text{OM}_1$  measured by the Q-AMS (slope = 0.91,  $r = 0.57$ ) but is within the 20% uncertainty reported for either technique and is consistent with a reduced collection efficiency of organic mass on sea salt or other non-refractory particles [Russell *et al.*, 2010, 2009b].

[10] Submicron particle composition was also measured aboard the R/V *Knorr* using a high resolution time of flight aerosol mass spectrometer (HR-ToF-AMS, Aerodyne Research Inc., Billerica, MA) [Canagaratna *et al.*, 2007; DeCarlo *et al.*, 2006]. Measurements were made every minute alternating automatically between V and W mode, with and without a thermal denuder, and in combination with particle time of flight (PToF) mode. The HR-ToF-AMS data were analyzed using SQUIRREL and PIKA software (described at <http://tinyurl.com/tofams-analysis>) to determine the concentrations of mass fragments and common aerosol species [Canagaratna *et al.*, 2007; DeCarlo *et al.*, 2006]. The speciated HR-ToF-AMS mass concentrations correlated well with those obtained from the Q-AMS. An unknown problem caused the absolute concentrations determined by the HR-ToF-AMS to be unusually and systematically low throughout this campaign, so only the Q-AMS was used for quantitative mass concentrations of the major aerosol components. The high resolution mass spectra obtained from the HR-ToF-AMS were used to investigate the normalized oxygen to carbon (O/C) ratios of the observed organic aerosol. The organic aerosol was further characterized by positive matrix factorization (PMF) of HR-ToF-AMS organic mass spectra.

## 2.3. Potential Source Contribution Function

[11] A potential source contribution function (PSCF) was used to determine the regions that have higher probabilities of being sources for the input of aerosol mass [Pekney *et al.*, 2006]. This algorithm uses the calculated 5-day isentropic Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model to generate air mass back trajectories [Draxler and Rolph, 2003] with gridded frequencies of co-occurrence with high concentration samples. Back trajectories were calculated arriving at 500 m above the sampling platform every hour for ICEALOT and every 4 h for Barrow. PSCF analysis was calculated individually on the time series of concentrations and mass fractions of organic functional groups, total  $\text{OM}_1$ , PMF derived factors, and Q-AMS mass fragments.

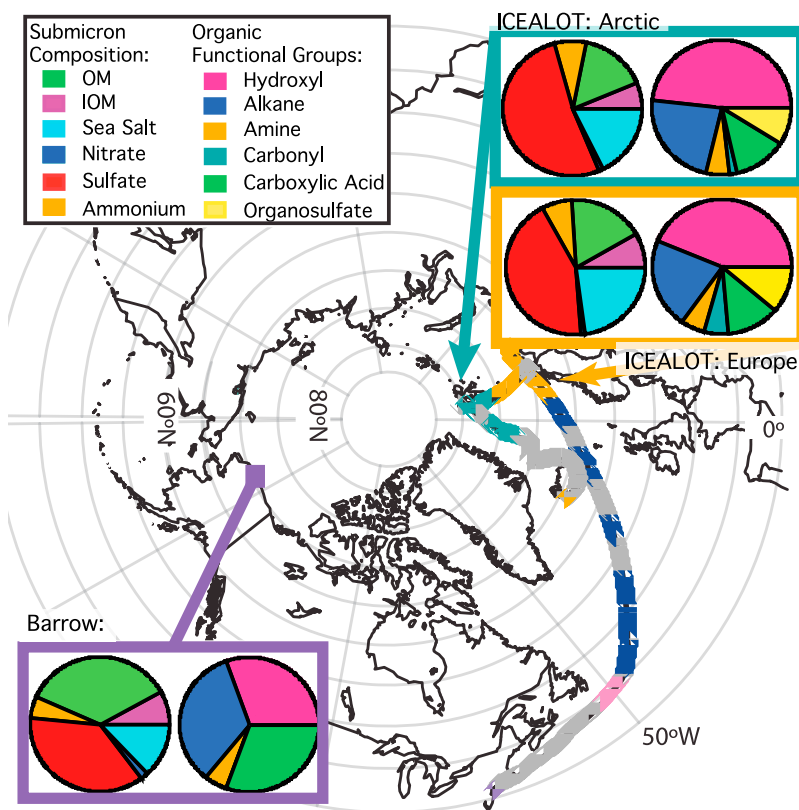
## 3. Results

[12] Five aerosol source regions were identified along the ICEALOT cruise track based on HYSPLIT 5-day isentropic back trajectories [Draxler and Rolph, 2003]. The five regions, including three continental (Long Island, North American, and European) and two marine (North Atlantic and Arctic), are shown along the ship track in Figure 1. Particles from the marine source regions had carbohydrate-like  $\text{OM}_1$  signatures mixed with sea salt and are described by Russell *et al.* [2010]. The continental regions differ from the marine regions in that they have higher  $\text{OM}_1$ ,  $\text{PM}_1$ , and combustion tracer concentrations in addition to higher radon concentrations (European source region: 540  $\text{mBq m}^{-3}$ ) relative to the marine source regions (190  $\text{mBq m}^{-3}$  for the Arctic and 340  $\text{mBq m}^{-3}$  from the North Atlantic).

### 3.1. Particle Mass

[13] Average non-sea salt  $\text{PM}_1$  concentrations from European, Long Island, and North American source regions were  $2.75 \pm 0.80$ ,  $3.40 \pm 1.10$ , and  $2.26 \pm 0.76 \mu\text{g m}^{-3}$ , respectively. The European and Arctic regions have similar relative particle compositions (Figure 1). Overall during ICEALOT, non-sea salt sulfate was the dominant anthropogenic aerosol component and had the highest measured concentration levels with an overall mean of  $1.06 \pm 0.70 \mu\text{g m}^{-3}$ , which was 53% of the total  $\text{PM}_1$  (Figure 2). The highest fractions of sulfate were measured from the European and Arctic source regions (58 and 64% of  $\text{PM}_1$ , respectively). Nitrate and ammonium levels were relatively low, accounting for less than 2% and 8% of the overall  $\text{PM}_1$ , respectively. At Barrow, the submicron particle composition had a larger fraction of  $\text{OM}_1$  (35%; see Figure 1) and a comparable fraction of sea salt (13%) and IOM (8%).

[14] During ICEALOT, the ratio of non-sea salt sulfate to non-crystal V was  $2.15 \pm 1.39$ , overall. Previously reported values at Barrow, 3 [Rahn and McCaffrey, 1980] and 2.9–4.2 [Quinn *et al.*, 2009], are within the standard deviation of this average suggesting that similar source regions were being measured, specifically European. Rahn and McCaffrey [1980] state that the ratio measured at Barrow in 1976 and 1977 is an order of magnitude greater than ratios measured at midlatitudes. This suggests that the aerosol measured during ICEALOT also has contributions of secondary aerosol, like that measured previously at Barrow. The higher ratio (2.44) measured from the European source region suggests



**Figure 1.** Track of the R/V *Knorr* during the ICEALOT cruise colored by regions of air mass origin including Long Island (purple), North America (pink), North Atlantic (dark blue), Europe (bronze), and Arctic (teal). The pies represent the submicron particle composition (left) and organic functional group composition (right) for the European and Arctic source regions during ICEALOT and Barrow (dark purple). The submicron particle composition excludes the elemental carbon (estimated on average less than 3% during ICEALOT and at Barrow).

that resulting aerosol from the European source region are more secondary and oxidized during transport.

### 3.2. Organic Mass

[15] Total  $OM_1$  measured by FTIR was calculated by summing the concentrations of the functional groups on each filter. The average  $OM_1$  concentration for ICEALOT (Figure 2) was  $0.61 \pm 0.36 \mu\text{g m}^{-3}$ . The highest concentrations (average:  $0.89 \pm 0.27 \mu\text{g m}^{-3}$ ) were observed when sampling was from the European source region. For two years at Barrow, the average springtime  $OM_1$  was  $0.41 \pm 0.36 \mu\text{g m}^{-3}$ . The organic functional group composition measured at Barrow had a larger fraction of alkane (34%) and carboxylic acid (30%) groups than the Arctic (23%, 13%) and European (21%, 12%) source regions measured during ICEALOT. In addition, organosulfate and non-acid carbonyl functional groups were measured in the Arctic and European source regions (on average 10% and 4%, respectively) and not observed at Barrow.

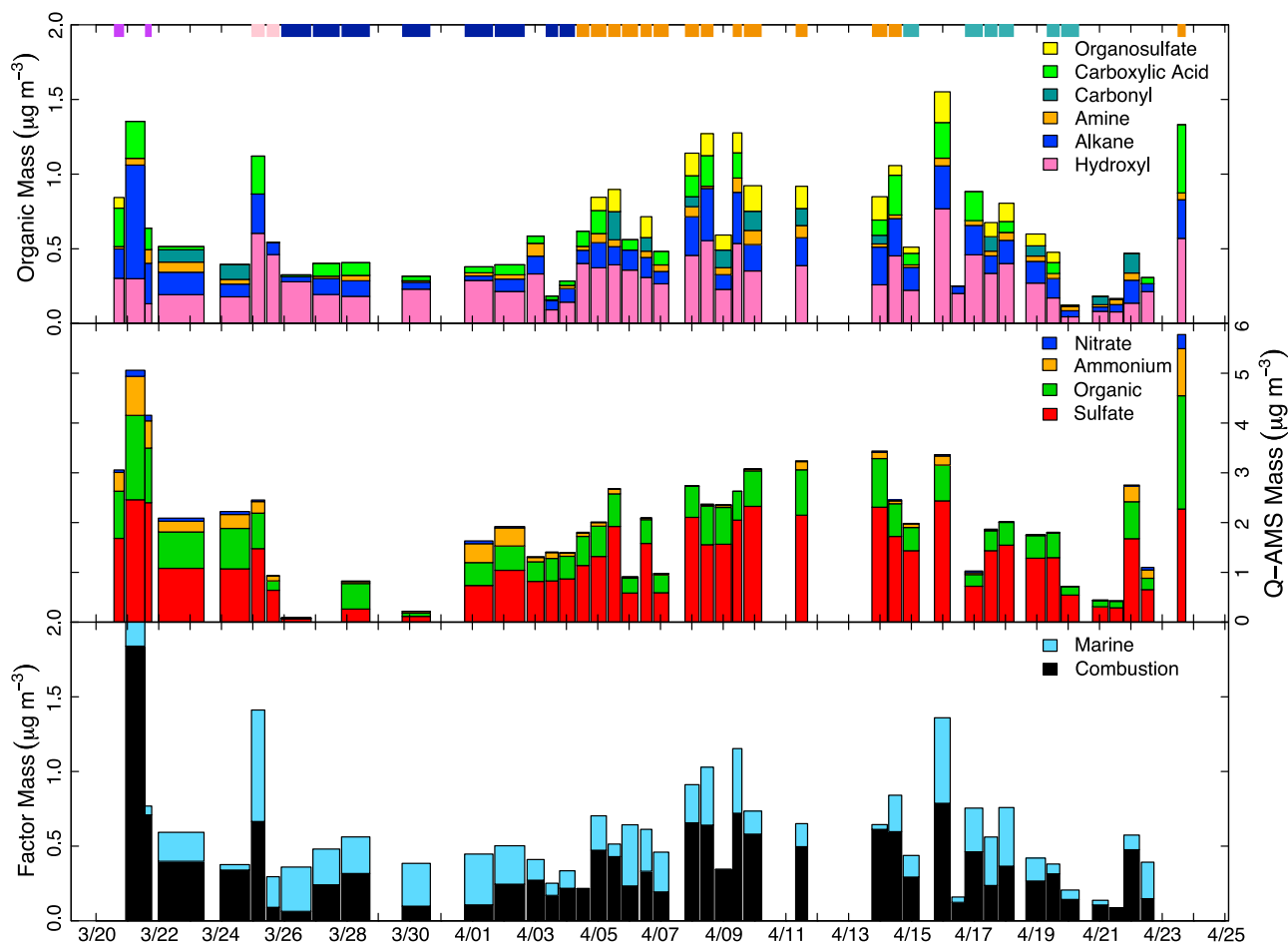
### 3.3. Ratio of Oxygen to Carbon

[16] The oxygenated fraction of organic aerosol has been used as an indicator of atmospheric processing and photochemical age [Maria *et al.*, 2004; Zhang *et al.*, 2005a]. The O/C ratio is sometimes an indicator of acid groups, which are a result of photochemical aging in the atmosphere. However,

some oxygenated groups such as hydroxyl may also increase O/C, although they can originate from primary sources such as ocean-derived marine aerosol [Russell *et al.*, 2010].

[17] For ICEALOT, the O/C ratio was calculated by two methods. Oxygen-containing functional groups (hydroxyl, carboxylic acid, carbonyl, and organosulfate) from FTIR measurements were summed to give the contribution of oxygen atoms to the FTIR O/C ratio and normalized by the total number of carbon atoms from each measured functional group [Hawkins *et al.*, 2010; Russell *et al.*, 2009a]. Because hydroxyl was a large fraction of the  $OM_1$  measured by FTIR, it had the largest contribution to the total O/C ratio, contributing on average 70% (0.66) of the total O/C. The O/C ratio calculated from the functional groups measured with FTIR spectroscopy was high (0.94) during ICEALOT indicating a highly oxygenated aerosol [Russell *et al.*, 2010]. The O/C from the European source region was 1.0, while the O/C in the Arctic source region was slightly lower (0.87) but still highly oxygenated.

[18] The HR-ToF-AMS O/C ratio was calculated from the individual abundances of O and C for each mass fragment using the Analytical Procedure for Elemental Separation (APES) included in the PIKA analysis software (described at <http://tinyurl.com/tofams-analysis>) [Aiken *et al.*, 2007, 2008]. In ambient environments, observed O/C values for oxidized secondary organic aerosol species range



**Figure 2.** Time series and average composition of aerosol mass. Horizontal color bar: regions of air mass origin colored according to the ship track in Figure 1. Top panel: Functional group concentrations contributing to the total  $OM_1$  measured by FTIR. Middle panel: Mass fragment contribution to total  $PM_1$  measured by Q-AMS. Bottom panel: Factors contributing to organic mass derived from PMF on FTIR spectra.

from  $\sim 0.2$  to  $\sim 0.9$  [Jimenez *et al.*, 2009]. Primary urban vehicle emissions have very low AMS O/C ratios from 0.06–0.1 [Aiken *et al.*, 2008], and an O/C ratio greater than 0.40 is considered highly oxygenated [DeCarlo *et al.*, 2008]. The calculated O/C ratio for ICEALOT correlates well ( $r = 0.91$ ) with the ratio of particulate  $CO_2$  (at nominal  $m/z$  44) to total  $OM_1$  consistent with the relationship proposed by Aiken *et al.* [2008]. The ICEALOT campaign averaged HR-ToF-AMS based O/C ratio is 0.41. Aerosol measured from the European source region has an O/C of 0.44, while the Arctic source region aerosol has a larger O/C (0.60). Taken together, these results indicate that the ICEALOT aerosol was highly oxidized. The increase in O/C in the Arctic region is likely indicative of continued processing and aging of aerosols as they are transported from sources to the Arctic.

[19] In general, the HR-ToF-AMS O/C values are systematically lower than those measured by the FTIR. The mild correlation ( $r = 0.68$ ) of the percent difference in the FTIR and HR-ToF-AMS O/C with the hydroxyl fraction of  $OM_1$  suggests that this difference in the O/C of the two methods is likely due to the lower O/C observed for hydroxyl groups by the HR-ToF-AMS [Aiken *et al.*, 2007; Hawkins *et al.*, 2010]. The HR-ToF-AMS O/C correlates

weakly with O/C calculated from only the FTIR carboxylic acid groups ( $r = 0.45$  in the European source region, but  $r = 0.27$  overall), which is consistent with the HR-ToF-AMS O/C correlation with the ratio of particulate  $CO_2$  (at nominal  $m/z$  44) to total  $OM_1$ , much of which is often attributed to acid mass fragments.

### 3.4. FTIR PMF

[20] Positive matrix factorization (PMF) was calculated for the FTIR spectra [Paatero, 1997; Paatero and Tapper, 1994] from the ICEALOT cruise and (separately) from the two-year Barrow data. A four-factor solution, which was recombined into two factors that were less correlated to each other in time, was found to best recreate the original ICEALOT time series (Figure 2), as previously discussed by Russell *et al.* [2010] (in the supporting information of that article). The two factors were determined to be a mixed combustion factor and a marine-derived factor (the latter of which is discussed in detail by Russell *et al.* [2010]). The mixed combustion factor was determined from correlations ( $0.5 < r < 1$ ) with typical combustion tracers (S, Zn, K, Br) [Qureshi *et al.*, 2006; Russell *et al.*, 2009a; Sharma and Maloo, 2005] and large fractions of alkane and carboxylic

acid functional groups consistent with combustion factors, containing both primary and secondary aerosol, from previous studies [Hawkins *et al.*, 2010; Russell *et al.*, 2011, 2009a]; see also section 4.1. The time series reconstructed from the factors of OM<sub>1</sub> correlates well with the original FTIR OM<sub>1</sub> time series ( $r = 0.86$ ). The correlation improves when the organosulfate functional group, which is not included in the wavelength range used for PMF, is removed from the FTIR OM<sub>1</sub> correlation ( $r = 0.89$ ). The slope of the line (Factor OM<sub>1</sub> = 1.01\*FTIR OM<sub>1</sub>) shows that the factors provide a good representation of the OM<sub>1</sub> and its variations.

[21] For the Barrow time series, PMF was performed on the 221 spectra resulting in a two-factor solution with FPEAK = 0. The FPEAK = 0 rotation of the PMF solution was selected since there was little sensitivity to rotations with higher or lower FPEAK values. Furthermore, the first half of the time series (from 2008) had measurements of inorganic ions by IC and trace elements by XRF on the same filters that correlated strongly with signatures for each factor. The spectra of the factors from the updated PMF analysis for the full 221 filters at Barrow are indistinguishable from those obtained from the original 118 spectra from 2008 to 2009, so we applied the same source-based definitions. The first factor was determined to be a mixed combustion factor because it exhibits strong spectral signatures in the alkane and carboxylic acid regions of the IR spectrum and correlates with combustion and industrial tracers (V, K, Fe, Zn, Br, etc.). The other factor was defined as an ocean derived factor because it exhibits strong spectral signatures of marine hydroxyl groups similar to those of ICEALOT described by Russell *et al.* [2010], correlated strongly with Na<sup>+</sup> and Cl<sup>-</sup> associated with sea salt, and corresponded to air masses that originated from open ocean in summer [Shaw *et al.*, 2010].

### 3.5. AMS PMF

[22] PMF was used for the organic portion of the HR-ToF-AMS mass spectrum calculated at unit mass resolution [Allan *et al.*, 2004b; Ulbrich *et al.*, 2009]. All analysis for HR-ToF-AMS PMF was done using PMF2 in robust mode [Paatero, 1997; Paatero and Tapper, 1994]. The PMF Evaluation Tool (PET) developed in Igor Pro (Wavemetrics, Inc., Portland, Oregon), described by Ulbrich *et al.* [2009], was used to evaluate the PMF output for HR-ToF-AMS spectra, with additional details provided in Appendix A.

[23] The three factors of the solution that best reconstructs the OM time series are: (1) an oxygenated organic aerosol (OOA) factor, with intense  $m/z$  44 signal; (2) a hydrocarbon-like organic aerosol (HOA) factor, characterized by hydrocarbon ion fragments ( $m/z$  41, 43, 55, 57, 67, 69, 71, etc.); and (3) a third factor (F3) with unique ion fragments ( $m/z$  50–55, 65, 67, 77, 79, 91). The time series and spectra of the factors are shown in Figures 3a and 3b.

[24] The OOA factor contributes 33% to the ICEALOT project average OM<sub>1</sub> and 31% in the European source region. The OOA factor spectrum correlates strongly with the reference spectra of oxalic, glyoxylic, and fulvic acids ( $r = 0.94$ ,  $r = 0.93$ ,  $r = 0.84$ , respectively) [Alfarra, 2004; Takegawa *et al.*, 2007] from the AMS spectral database (I. M. Ulbrich *et al.*, <http://cires.colorado.edu/jimenez-group/AMSSd/>) described by Ulbrich *et al.* [2009], providing an indication that this highly oxygenated factor is

representative of acid-containing organic aerosol. The OOA factor correlates mildly in time with the FTIR measured carboxylic acid functional group ( $r = 0.49$  overall,  $r = 0.71$  in the European source region) consistent with the interpretation of the OOA factor as an acid-containing, secondary, aged, organic component. The time series of the OOA factor also correlates ( $r = 0.67$ ) with sulfate, consistent with other measurements in eastern Europe [Lanz *et al.*, 2007].

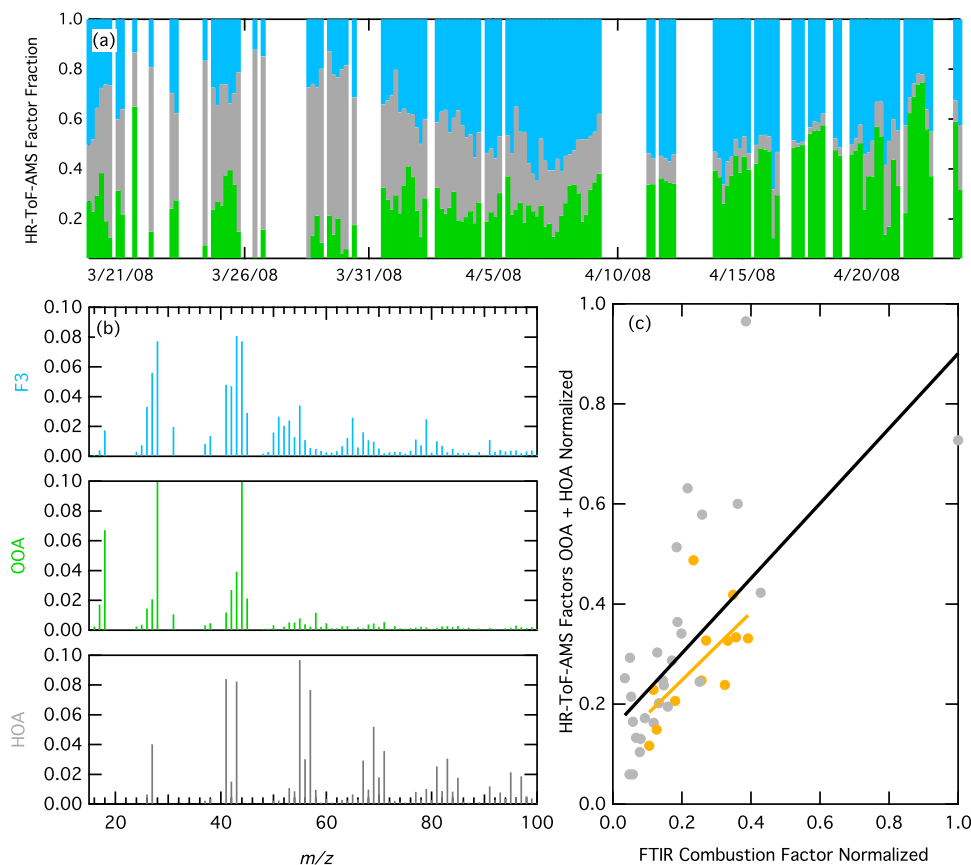
[25] The HOA factor is present mainly at the start of the cruise in the Long Island and North American source regions. HOA contributes 28% to the total OM<sub>1</sub> and 29% and 41% in the Long Island and North American source regions, respectively. The HOA factor spectrum correlates strongly with HOA spectra measured in other campaigns ( $r = 0.99$  and  $r = 0.98$ ) [Lanz *et al.*, 2009; Zhang *et al.*, 2005b] and also diesel bus exhaust ( $r = 0.98$ ) [Canagaratna *et al.*, 2004; Ulbrich *et al.*, 2009]. In addition, the HOA factor correlates ( $r = 0.68$ ) with NO<sub>x</sub> (measured on board the R/V *Knorr* using the method described by Lerner *et al.* [2009]), consistent with less processed emissions [Lanz *et al.*, 2007]. HOA is lower in the European source region (20%) where there is a longer distance between emissions and sampling, suggesting that the emissions measured in the European source region had been processed more or longer in the atmosphere.

[26] The OOA factor and the FTIR combustion factor show a correlation of  $r = 0.42$  overall and  $r = 0.74$  for the European source region, which is consistent with the large contribution of carboxylic acid functional groups to the FTIR combustion factor. The HOA factor has a moderate correlation with the FTIR combustion factor ( $r = 0.51$ ), but the correlation to the FTIR combustion factor is higher for the sum of OOA and HOA ( $r = 0.67$  overall and  $r = 0.64$  for the European source region; see Figure 3c). This suggests that the FTIR combustion factor includes both hydrogenated and oxygenated carbon mass fragments, likely including a range of recent and more processed components of combustion emissions. In the European source region, the stronger correlation of the FTIR combustion factor with OOA than with OOA + HOA suggests that the FTIR combustion factor OM<sub>1</sub> from that region is more processed.

## 4. Discussion

### 4.1. Combustion Sources of Organic Mass in Arctic Haze

[27] Emissions from combustion sources provide a large fraction of the aerosol mass in the Arctic that makes up the springtime Arctic haze. Recent measurements at Barrow and during ICEALOT show that the combustion sources typically contribute more than half of OM<sub>1</sub> and over a quarter of PM<sub>1</sub>, based on FTIR, AMS, and IC analyses. A comparison of aerosol measured during the spring in these two regions of the Arctic provides insight into the differences in emission sources and regions contributing to the springtime Arctic haze. The large fraction of oxygenated functional groups measured in the Arctic indicates a high O/C fraction close to 1 (discussed in section 3.3), which is consistent with an aged combustion source and indicates one oxygen for every carbon in an organic particle [Jimenez *et al.*, 2009; Russell *et al.*, 2010, 2009a]. Emissions from coal combustion in northeastern Europe contribute to the OM<sub>1</sub> measured



**Figure 3.** (a) Time series of HR-ToF-AMS factor fractions of total OM including HOA (dark gray), OOA (green), and F3 (light blue). (b) The mass spectra of the three distinct factors: HOA, OOA, and F3. (c) Comparison between the HR-ToF-AMS factors OOA + HOA (normalized by the maximum value) and the FTIR combustion factor (normalized by the maximum value),  $r = 0.64$  overall (black) and  $r = 0.64$  in the European source region (orange).

in the northern Atlantic region of the Arctic, while biomass burning, shipping, and industrial emissions contribute to the OM<sub>1</sub> measured in the northern Pacific region of the Arctic.

#### 4.1.1. Combustion Factor Contribution to OM<sub>1</sub>

[28] PMF of both ICEALOT and Barrow FTIR spectra provide evidence for combustion factors that constitute more than 60% of the OM<sub>1</sub> measured at both sites. The combustion factors show significant fractions of alkane, hydroxyl, and carboxylic acid functional groups. During ICEALOT, the combustion factor was 63% of the total OM<sub>1</sub> and contributed  $0.36 \pm 0.31 \mu\text{g m}^{-3}$  to the total organic mass (Figure 2). When the aerosol originated from the European source region, the combustion factor was 68% of the total OM<sub>1</sub> and contributed  $0.47 \pm 0.18 \mu\text{g m}^{-3}$ . Since OM<sub>1</sub> was approximately 33% of the overall PM<sub>1</sub> and 43% in the European air mass source region, this large contribution of combustion sources from Europe provided a substantial part of the Arctic haze aerosol concentrations in the Barents and Greenland Sea regions during the ICEALOT cruise.

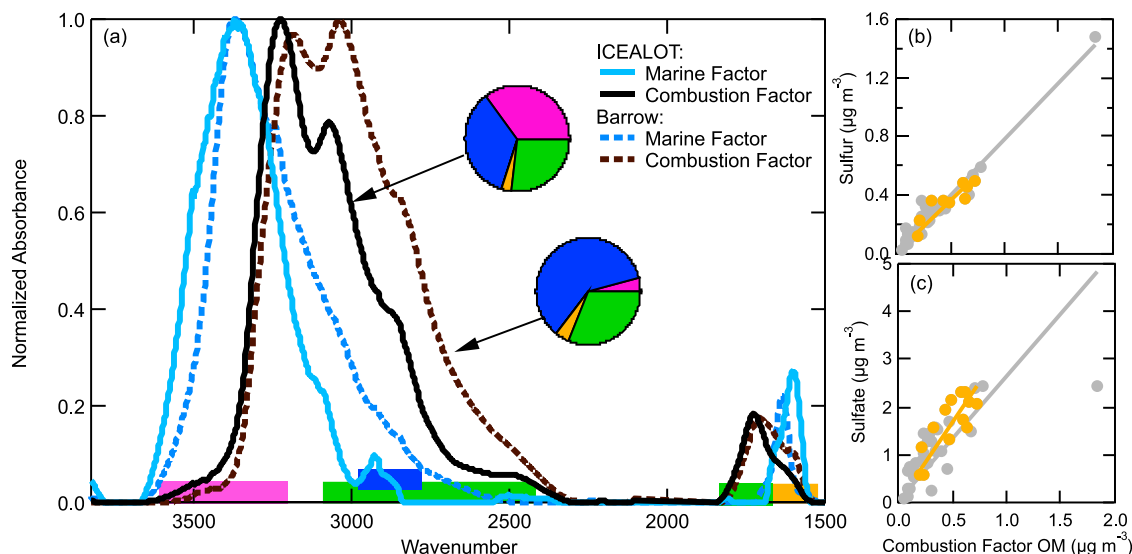
[29] Measurements taken in Barrow, Alaska, span two years and three different seasons (spring, summer and winter). The Barrow spring season was defined to represent the high OM<sub>1</sub> concentrations that prevail in March through June [Shaw *et al.*, 2010] and includes the ICEALOT sampling period. The OM<sub>1</sub> measured during spring in both 2008 and 2009 at

Barrow was higher ( $0.41 \pm 0.36 \mu\text{g m}^{-3}$ ) than the average OM<sub>1</sub> measured both years ( $0.32 \pm 0.36 \mu\text{g m}^{-3}$ ). Most (80% or  $0.32 \pm 0.22 \mu\text{g m}^{-3}$ ) of the OM<sub>1</sub> measured in the spring was attributed to combustion sources based on correlations with trace metals in spring 2008 [Shaw *et al.*, 2010]. The OM<sub>1</sub> concentration and fraction of the combustion factor were significantly higher than the annual average combustion OM<sub>1</sub> ( $0.19 \pm 0.21 \mu\text{g m}^{-3}$ ), indicating that combustion sources also had a substantial impact at Barrow on the OM<sub>1</sub> fraction of springtime Arctic haze.

#### 4.1.2. Combustion Factor Composition

[30] The spectra of the combustion factors for aerosol measured in the northern Pacific and northern Atlantic regions of the Arctic are similar (Figure 4a). The differences between the two combustion factor spectra may result both from differences in their source emissions (i.e., between the mixture of combustion sources in Siberia and those in eastern Europe) and in their atmospheric oxidation processes. The ICEALOT combustion factor OM<sub>1</sub> includes large mass fractions of hydroxyl (36%), alkane (35%), and carboxylic acid (27%) functional groups, with a minor fraction (3%) of amine groups (Figure 4a). The large fraction of carboxylic acid functional groups suggests that particles aged from combustion emissions contribute to this factor [Maria *et al.*, 2004; Zhang *et al.*, 2005a]. The Barrow combustion factor





**Figure 4.** (a) Factor spectra from PMF of FTIR sample spectra for ICEALOT (solid) and Barrow (dashed). The pie charts show the average composition of the combustion factor, using the colors described in Figure 1, for both ICEALOT and Barrow. Comparisons of the ICEALOT combustion factor with (b) S ( $r = 0.97$ ,  $r = 0.9$ ) and (c) sulfate ( $r = 0.75$ ,  $r = 0.8$ ) for all air masses (gray) and the European source region (orange), respectively.

for the full 2-year study consists of alkane (60%) and carboxylic acid (31%) groups and smaller fractions of amine (5%) and hydroxyl (4%) groups (Figure 4a). The large fraction of alkane groups suggests that the combustion emissions contributing to this factor may have undergone less atmospheric processing than those measured during ICEALOT.

[31] The combustion factors retrieved from the organic FTIR spectra for both ICEALOT and Barrow have large contributions from both carboxylic acid and hydroxyl functional groups. These oxygenated groups are characteristic of the products expected from photochemical oxidation [Huebert and Charlson, 2000; Lim and Ziemann, 2005, 2009; Putaud et al., 2000] indicating that the measured particles are due in part to secondary organic aerosol. The weakness of the correlations during ICEALOT to elemental tracers (Fe and V) indicates that the variability in the combustion factor OM may lag the primary combustion products, which is expected for secondary rather than primary OM.

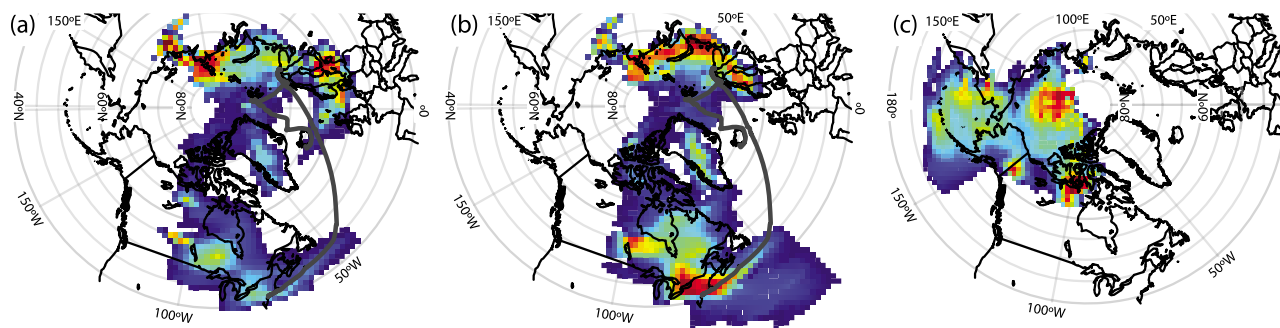
#### 4.1.3. Sources of Combustion in Northeastern Europe and Asia

[32] The combustion factor derived from measurements in the north Atlantic and adjacent regions of the Arctic Ocean during ICEALOT reflects aerosol influenced by emissions from northeastern Europe and Russia. Coal burning is one of the main sources of heating, and there are also numerous coal burning smelters in that area [Hole et al., 2006a; Tzimas et al., 2009; Weiler et al., 2005]. The Kola Peninsula, northeast of Finland, has Cu and Ni producing industry which emits high levels of  $\text{SO}_2$  and  $\text{NO}_2$  [Weiler et al., 2005]. In the Kola Peninsula, the conurbation of Murmansk is a highly populated metropolitan area and sea port, which contributes to the polluted air from that area [Law and Stohl, 2007]. The combustion factor concentration correlates strongly with S concentrations for both overall ( $r = 0.97$ ) and the European source region ( $r = 0.90$ ; see Figure 4b) indicating that coal burning is likely collocated or

co-emitted with the organic combustion sources. This strong correlation of the combustion factor OM<sub>1</sub> with sulfur likely is a consequence of S being primarily sulfate, a typical secondary aerosol product of coal combustion emissions, which also correlates strongly with  $r = 0.75$  overall (and  $r = 0.80$  in the European source region; see Figure 4c) suggesting that the sources of the organic combustion factor and those of the sulfate are collocated. Correlations with Zn (total  $r = 0.74$ ; European source region  $r = 0.57$ ) are consistent with emissions from non-ferrous metal producing industries, such as copper smelting in northeastern Europe [Polissar et al., 1998, 2001; Shaw, 1982]. The combustion factor weakly correlates with V ( $r = 0.24$  overall,  $r = 0.44$  in the European source region), suggesting that the combustion factor may also have some contributions from oil burning. The ratio of non-sea salt sulfate to non-crustal V mildly correlates ( $r = 0.55$ ) in the European source region with the combustion factor fraction of OM. In addition to the correlations with the combustion and pollution tracers (Zn, Ca, Fe, S, and V), the concentrations of these metals were 5–65% higher in the European source region than the ICEALOT average.

[33] About 50% of Europe's electricity is produced from fossil fuel combustion [Tzimas et al., 2009], and high alkane concentrations are found in petroleum combustion emissions from Europe [Blake et al., 1994, 2003]. In eastern Europe, combustion of fuels such as coal and lignite lead to high levels of  $\text{SO}_x$ ,  $\text{NO}_x$ , PM, and  $\text{CO}_2$  emissions [Ross et al., 2002]. The mixture of high  $\text{SO}_x$  emissions from coal and alkane emissions from petroleum could explain the correlation of the organic combustion factor and the sulfate observed during the European source region of the ICEALOT cruise, making emissions from northeastern Europe the main source of OM<sub>1</sub> in the Arctic haze on the northern Atlantic side.

[34] At Barrow, the combustion factor also correlates strongly to S but, in addition, is correlated to non-dust K,



**Figure 5.** PSCF of (a) the total  $OM_1$  measured during ICEALOT, (b) the ICEALOT combustion factor, and (c) the Barrow combustion factor. The higher potential source regions are indicated by red and the lower potential source regions are indicated with blue.

characteristic of a biomass burning source, where  $[non-dust\ K] = [K] - 0.62*[AI]$  [Cachier et al., 1995; Gilardoni et al., 2009]. Alaskan and Siberian wildfires were observed in spring 2008 and 2009 [U.S. Department of Agriculture, 2010; Shaw et al., 2010; Warneke et al., 2009], which provide evidence for the influence of biomass burning in the combustion factor. V, which is characteristic of processed emissions from oil burning, is mildly correlated with the combustion factor. V is a tracer for the combustion of heavy residual oil only used in the sub-Arctic since it is too viscous to use at cold temperature and is hence a tracer of the long range transport of emissions from oil combustion [Isakson et al., 2001].

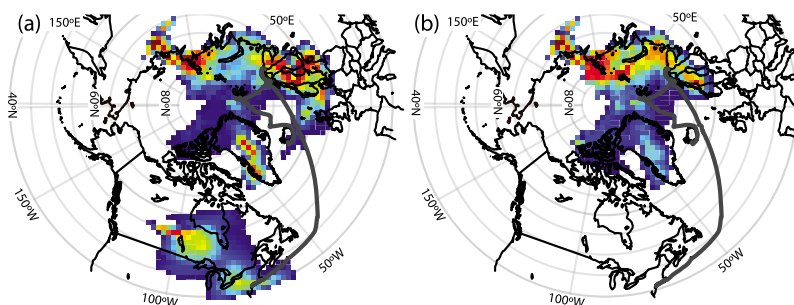
[35] To investigate the probability that these two different urban and industrial centers were the emission sources that were responsible for the observed high  $OM_1$  from combustion factors, PSCF were calculated for Barrow and ICEALOT [Pekney et al., 2006]. The region with high potential of being a source of the ICEALOT combustion factor is northeastern Europe, with the highest potential sources shown in Figure 5b as Scandinavia and the Kola Peninsula, as well as the northwestern edge of Russia. These areas of northeastern Europe have significant industrial activity, including non-ferrous metal producing plants that contribute to the organic emissions [Blake et al., 2003; Hole et al., 2006a; Mira-Salama et al., 2008; Weiler et al., 2005]. The PSCF of sulfate measured during ICEALOT shows a trend similar to the potential source of the organic combustion factor (see Figures 6a and 5b for comparison), supporting the suggestion that the organic and sulfate sources are collocated. In addition, PSCF for  $OM_1$  measured during ICEALOT also shows potential sources in northeastern Europe (Figure 5a). PSCF of the Barrow combustion factor shows potential sources from northeastern Asia, shipping lanes in the Bering Strait, and along the Canadian-Alaskan border (Figure 5c), which are similar to the potential sources observed in 2008. The differences in the potential source regions shown in Figure 5 highlight the importance of northeastern Asian sources for Arctic haze at Barrow in contrast to the northeastern European sources of the Arctic haze during ICEALOT.

[36] In summary, the functional group composition indicates that the majority of the organic aerosol in Arctic haze is produced from combustion emissions, and back trajectories from both regions suggest that the combustion sources are in northeastern Europe and Asia. The two different sources of

measured Arctic aerosol are demonstrated in the differences between the composition of their combustion factors and the potential source regions determined from PSCF.

#### 4.2. Organosulfate Formation From Combustion Emissions

[37] One interesting chemical consequence of the co-emissions of sulfate from coal burning and volatile organic compound (VOC) emissions in northeastern Europe is the production of organosulfate functional groups. In air masses sampled from both the European and Arctic source regions of the ICEALOT campaign, organosulfate groups contributed 9% and 11% of the organic mass, respectively, whereas no organosulfate groups were measured at Barrow. The organosulfate functional group fraction of  $PM_1$  has a strong correlation ( $r = 0.84$ ) with the sulfate fraction of  $PM_1$  for the European region and a moderate correlation ( $r = 0.50$ ) for the entire ICEALOT cruise (Figure 7b) and was only detected in samples with sulfate concentrations greater than  $1.25\ \mu\text{g m}^{-3}$ . This observed increase in organosulfate group fraction of  $PM_1$  with increased sulfate fraction of  $PM_1$  is consistent with organosulfate formation on sulfate particles, many of which are likely acidic in this region [Surratt et al., 2007a]. The molar ratio of ammonium to sulfate was 1.13 for the whole project and 1.22 during the periods of filter sampling. The lower average molar ratio of ammonium to sulfate (0.96) of particles measured from the European and Arctic source regions and their implied higher acidity provides conditions consistent with the acid-catalyzed particle-phase formation of organosulfate in air masses from those source regions [Inuma et al., 2009, 2007; Liggio et al., 2005; Perri et al., 2010; Surratt et al., 2007a, 2010, 2007b, 2008]. Organosulfate group fraction of  $PM_1$  has a mild negative correlation ( $r = -0.50$ ) with relative humidity for the European source region (Figure 7c). In addition, organosulfate group concentration is mildly negatively correlated ( $r = -0.64$ ) with absolute humidity in the European source region. These relationships suggest that organosulfate formation decreases with higher aerosol water concentrations, which may be due in part to the decrease in aerosol acidity with an increase in relative humidity ( $r = 0.52$  overall,  $r = 0.51$  for the European source region), consistent with the findings of a previous study in the southeastern Pacific [Hawkins et al., 2010; Liggio et al., 2005]. In summary, four conditions coincide with organosulfate group formation in the European and Arctic source regions, relative to the ICEALOT measurement



**Figure 6.** PSCF of (a) sulfate and (b) organosulfate functional group concentrations. The higher potential source regions are indicated by red and the lower potential source regions indicated with blue.

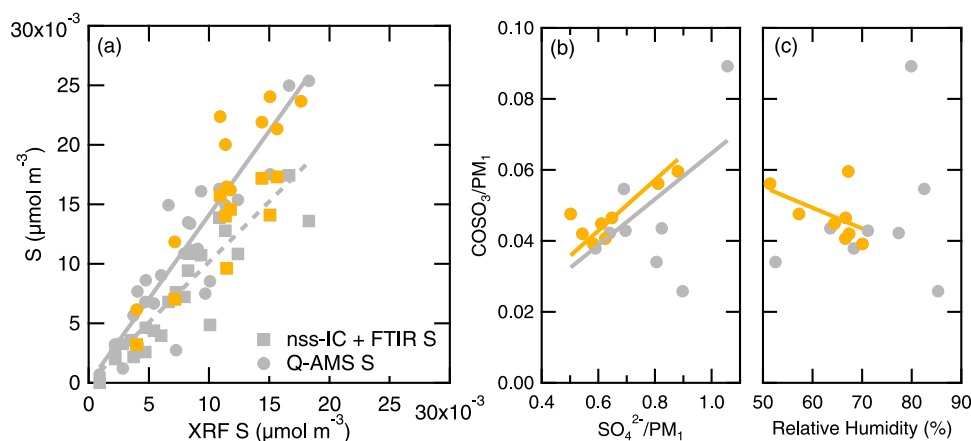
period: high organic mass ( $>0.75 \mu\text{g m}^{-3}$ ), high sulfate ( $>1.25 \mu\text{g m}^{-3}$ ), low ammonium to sulfate ratio ( $<1.0$ ), and high overall relative humidity ( $>50\%$ ). These observations are consistent with sulfate particles providing acidic aqueous conditions that enhance organosulfate formation from organic precursors. The correlation of the sulfate and the organosulfate fractions of  $\text{PM}_{10}$  also suggests both similar sources and similar processing times in the atmosphere (given the time resolution of the filter sampling, namely 12 h).

[38] The four conditions for organosulfate formation observed during ICEALOT did not all occur at Barrow, which could explain the lack of organosulfate formation there. During the spring seasons, the average sulfate concentration measured at Barrow was  $0.64 \mu\text{g m}^{-3}$ , which is less than observed during ICEALOT but consistent with the range reported by Quinn *et al.* [2002] for the spring seasons of 1998–2000. On the other hand, the average molar ratio of ammonium to sulfate (0.75) was consistent with the acidic ratio measured during ICEALOT. Average  $\text{OM}_{10}$  concentration ( $0.70 \mu\text{g m}^{-3}$ ) measured during the spring season is slightly less than was observed for periods with organosulfate functional groups during ICEALOT. The process of acid-catalyzed particle haze formation of organosulfate likely does not occur during the spring seasons at Barrow.

[39] The sum of S ( $\mu\text{mol m}^{-3}$ ) from inorganic sulfate measured by IC and organosulfate groups measured by FTIR correlates well ( $r = 0.92$ ) with the total S measured by XRF (Figure 7a). Without the organosulfate group component, there is still a strong correlation ( $r = 0.92$ ) to elemental (XRF) S, but the slope of the line (0.94) suggests that there is a missing source of S. When organosulfate is added to the IC measured sulfate, the total S is better represented, with a slope of 1.02. This suggests that the majority (94%) of the S concentration measured during ICEALOT is inorganic sulfate, with the remaining 6%, on average, coming from organic sulfate. The S from Q-AMS measurements of sulfate also correlates well ( $r = 0.91$ ) with XRF measurements of S, consistent with the expectation that the Q-AMS detects both organic and inorganic sulfate [Allan *et al.*, 2004a; Farmer *et al.*, 2010].

[40] Using PSCF, the highest potential source contributions of the ICEALOT sulfate concentrations were found in northeastern Europe, similar to the locations of potential sources for the combustion factor, as is evident by comparing Figures 5b and 6a. In addition, the potential source of organosulfate groups is collocated with sulfate, as shown in Figure 6b.

[41] Other studies have shown organosulfate formation from biogenic precursors [Gómez-González *et al.*, 2008;



**Figure 7.** (a) Comparison of S measured by XRF with IC + FTIR derived S (slope = 1.02,  $r = 0.92$ ) and Q-AMS derived S ( $y = 1.41$ ,  $r = 0.91$ ) for the total project (gray) and the European source region (orange). (b) Correlation of organosulfate group fraction of  $\text{PM}_{10}$  with sulfate fraction of  $\text{PM}_{10}$  ( $r = 0.84$  for the European source region). (c) Negative correlation of organosulfate group fraction of  $\text{PM}_{10}$  and relative humidity ( $r = -0.50$  for the European source region).

*Iinuma et al.*, 2009; *Surratt et al.*, 2007a, 2010, 2008]. Because there were no large sources of terrestrial biogenic emissions measured during ICEALOT, biogenic precursors during ICEALOT were not identified as a likely source of organosulfate functional groups. While the emissions from the European source region measured during ICEALOT have been attributed to fossil fuel burning, due to the absence of biogenic volatile organic compounds (BVOCs) in the air measured at the ship, it remains possible that co-located BVOCs did contribute to OM and to subsequent reactions with sulfate in particles to form organosulfate. *Claeys et al.* [2010] found evidence that organosulfate functional groups can occur with acidic particles produced from oxidation products of dimethyl sulfide (DMS) and bacteria sources in a clean marine environment. The organosulfate measured during ICEALOT was unlikely to be the result of marine emissions since organosulfate functional groups were only observed when air masses were from European and Arctic source regions. Organosulfate group concentrations or fractions did not correlate with the marine factor, as would be expected for marine sources.

## 5. Conclusions

[42] Emissions from combustion sources greatly impact the concentrations of submicron aerosol particles in the Arctic and contribute to Arctic haze in both the Atlantic and Pacific sides of the Arctic region. Complementary measurement techniques show higher OM<sub>1</sub> in air masses sampled from the European source regions than from the marine source regions, both measured on the Atlantic side of the Arctic during ICEALOT. A large fraction (74%) of the OM<sub>1</sub> was attributed to eastern European combustion sources (industrial and other petroleum burning) based on PMF-derived factors from FTIR measurements. Measurements from Barrow, Alaska, on the Pacific side of the Arctic show the largest percent of OM<sub>1</sub> from mixed combustion sources (biomass burning and shipping) in the spring season. The organic functional group compositions indicate that mixed combustion sources account for more than 60% (>0.3 μg m<sup>-3</sup>) of the total organic mass measured in air masses influenced from both the European and Arctic source regions during ICEALOT and the spring season at Barrow.

[43] Carboxylic acid and hydroxyl groups account for more than 35% of the combustion factor organic functional groups in the Atlantic and Pacific sides of the Arctic, reflecting the products of oxidation of emissions from fuel burning. The ICEALOT combustion factor, dominated by carboxylic acid, alkane, and hydroxyl functional groups, correlates well with sulfate suggesting that their sources are collocated. The large fraction of the carboxylic acid functional group is consistent with the secondary products expected from oxidation of VOCs. Q-AMS and HR-ToF-AMS measurements confirm the highly oxygenated nature of the OM<sub>1</sub> from combustion, with an OOA spectrum similar to that of oxygenated acid groups. Aerosol measured in the Pacific side of the Arctic at Barrow has a larger fraction of alkane suggesting that the aerosol is less aged. The differences in the combustion-derived spectra from the two regions result from the mixtures of sources in each region, although more specific contributions could not be resolved in these campaigns.

[44] On the Atlantic side of the Arctic, high levels of sulfate and OM<sub>1</sub> were typically associated with back trajectories that originated in northeastern Europe. The correlations between the combustion factor for ICEALOT and S and sulfate suggest emissions from the high coal burning regions of northeastern Europe. On the other hand, the correlations between the Barrow combustion factor and V and non-dust K are characteristic of emissions from northern Asia, Siberia, and, to a lesser extent, interior regions of Alaska and Canada.

[45] The high emissions of sulfate from coal-burning in northeastern Europe produce significant concentrations of the organosulfate functional groups that account for as much as 10% of OM<sub>1</sub> measured by FTIR over the Barents and Greenland seas during ICEALOT. There is some evidence suggesting that organosulfate formation occurs heterogeneously when compounds react in acidic (aqueous) sulfate particles. Organosulfate functional groups were measured in air masses from both the European and Arctic source regions, when high concentrations (1.25–2.40 μg m<sup>-3</sup>) of sulfate were present but not at Barrow, which had lower sulfate concentrations (0.64 μg m<sup>-3</sup>, average). The potential source region of organosulfate groups is collocated with the potential source region of sulfate during ICEALOT, where there are high S emissions from abundant sources of coal combustion. The high correlation between the organosulfate fraction of PM<sub>1</sub> and sulfate fraction of PM<sub>1</sub> and the low molar ratio of ammonium to sulfate (high acidity) suggests that organosulfate functional groups are formed on sulfate particles through acid-catalyzed aqueous reactions.

[46] Our findings suggest that pan-Arctic air quality is not uniform in composition but rather regionally dependent on different transport mechanisms and source regions. Since these unique aerosol contributions produce different chemical and microphysical consequences, climate models that are able to simulate direct and indirect aerosol effects must be able to treat the specific areas differently in order to make predictions of radiative forcing in the Arctic.

## Appendix A: PMF of HR-ToF-AMS

[47] Unit mass resolution PMF analysis was performed on the HR-ToF-AMS data. The PET software and analysis methods outlined by *Ulbrich et al.* [2009] were used to perform the PMF analysis. The data and error matrices input into the PMF analysis were obtained with the SQUIRREL analysis program (described at <http://tinyurl.com/tofams-analysis>) [*Allan et al.*, 2003, 2004b]. The error matrices account for Poisson statistics in ion counting. A minimum error constraint was additionally placed on the error matrix according to *Ulbrich et al.* [2009]. Due to low signal to noise (S/N), ion fragments with masses above *m/z* 200 were removed from the analysis. The remaining ion fragments with 0.2 < S/N < 2 were downweighted by a factor of 3 [*Ulbrich et al.*, 2009]. PMF analysis results were compared for solutions with 1 through 10 factors. The largest drop in Q/Q<sub>expected</sub> occurred going from 1 to 2 factors, showing that at least two factors are needed to accurately represent the organic mass and explain the variance. The three factor solution, which had a minimum Q value and did not show any evidence of factor splitting, was chosen as the optimum solution for this analysis. The three factors together account

for 97% of the observed OM. The effect of rotation on the PMF solution was examined by changing the FPEAK parameter from  $-1$  to  $1$  with  $0.2$  intervals. Solutions with  $-0.22 < \text{FPEAK} < 0$  yielded component spectra and time series that did not contain unreasonable zeros. The chosen solution is supported by the strongest correlations with reference spectra posted at <http://cires.colorado.edu/jimenez-group/AMSsd/> [Ulbrich et al., 2009].

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