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Flow climatology for physicochemical properties of dichotomous aerosol over the western North Atlantic Ocean at Bermuda

J. L. Moody¹, W. C. Keene¹, O. R. Cooper², K. J. Voss³, R. Aryal³, S. Eckhardt⁴, B. Holben⁵, J. R. Maben¹, M. A. Izaguirre⁶, and J. N. Galloway¹

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Correspondence to: J. L. Moody (moody@virginia.edu)

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¹Department of Environmental Sciences, University of Virginia, Charlottesville, VA, USA

²Cooperative Institute for Research in Environmental Sciences (CIRES), University of Colorado, and NOAA Earth System Research Laboratory, Boulder, CO, USA

³Physics Department, University of Miami, FL, USA

⁴Norwegian Institute for Air Research, Kjeller, Norway

⁵Goddard Space Flight Center, NASA, Greenbelt, MD, USA

⁶Division of Marine and Atmospheric Chemistry, University of Miami, FL, USA

Dichotomous aerosols (nominal super- and sub-um-diameter size fractions) in sectored on-shore flow were sampled daily from July 2006 through June 2009, at the Tudor Hill Atmospheric Observatory (THAO) on the western coast of Bermuda (32.27° N, 64.87° W) and analyzed for major chemical and physical properties. Flexpart retroplumes were calculated for each sampling period and aerosol properties were stratified accordingly based on transport from different regions. Transport from the northeastern United States (NEUS) was associated with significantly higher (factors of 2 to 3 based on median values) concentrations of bulk particulate non-sea-salt (nss) SO_4^{2-} , NO_3^{-} , and NH₄ and associated scattering and absorption at 530 nm, relative to transport from Africa (AFR) and the Oceanic background. These differences were driven primarily by higher values associated with the sub-um size fraction under NEUS flow. We estimate that 75 (\pm 3) % of the NEUS nss SO₄²⁻ was anthropogenic in origin, while only $25 (\pm 9) \%$ of the AFR nss SO_4^{2-} was anthropogenic. Integrating over all transport patterns, we estimate the contribution of anthropogenic sulfate has dropped 14.6 % from the early 1990's. Bulk scattering was highly correlated with bulk nss SO₄²⁻ in all flow regimes but the corresponding regression slopes varied significantly reflecting differential contributions to total scattering by associated aerosol components. Absorption by super-µm aerosol in transport from the NEUS versus AFR was similar although the super-um aerosol size fraction accounted for a relatively greater contribution to total absorption in AFR flow. Significantly greater absorption angstrom exponents (AAEs) for AFR flow reflects the wavelength dependence of absorption by mineral aerosols; lower AAEs for NEUS flow is consistent with the dominance of absorption by combustionderived aerosols. Higher AOD associated with transport from both the NEUS and AFR relative to Oceanic background flow results in a top of atmosphere direct radiative forcing on the order of -1.6 to -2.5 W m⁻², respectively, showing these aerosols drive cooling. The dominance of transport from the NEUS on an annual basis coupled with the corresponding decreases in anthropogenic nss SO₄²⁻ aerosols since the early 1990's

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implies that emission reductions in the US account for a decline in atmospheric cooling over the western North Atlantic Ocean during this period.

Introduction

Tropospheric aerosols influence Earth's climate system directly by scattering and absorbing incoming and outgoing radiation and indirectly by modulating the microphysical properties of clouds (e.g., Albrecht, 1989; Charlson and Heizenberg, 1995; Ramanathan et al., 2001; Kaufman et al., 2002; Andreae, 2009; Clarke and Kapustin, 2010). Scattering and absorption reduce radiative transfer through the atmosphere resulting in net cooling at the surface but absorption also warms the atmosphere thereby influencing its thermodynamic properties and, in heavily impacted regions, suppressing cloud formation (e.g., Koren et al., 2004). In addition, based on their size and hygroscopicity, aerosols serve as cloud condensation nuclei (CCN). For a given set of conditions, the number concentration of CCN regulates the corresponding number concentration and size distribution of cloud droplets and thus cloud brightness and evolution, surface irradiance, precipitation efficiency, the hydrologic cycle, and associated feedbacks (e.g., Andreae et al., 2004). Outstanding uncertainties regarding the overall impacts of aerosols on radiative transfer are among the major factors currently limiting our understanding of and ability to predict global climate change (IPCC, 2007).

Large, intensive field campaigns and associated modeling have contributed much towards resolving details of aerosol sources, characteristics, processing, optical properties, and influences on radiative transfer (e.g., Bates et al., 2006). A primary underlying motivation for many such campaigns is to deconvolute the climatic influences of aerosols from major anthropogenic and natural sources. Consequently, intensives have often targeted conditions within or downwind of major source regions for aerosols and precursors and, due in part to cost, most are of relatively short duration (4 to 8 weeks). Examples include investigations of pollutant plumes from North America transported over the western North Atlantic Ocean (NAO), the Tropospheric Aerosol Radiative Forc**ACPD**

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ing Observational Experiment (TARFOX) (Russell et al., 1999), the New England Air Quality Study (NEAQS) (Quinn and Bates, 2005), and the International Consortium in Transport and Transformation (ICARTT) campaign (Sierau et al., 2006), from Europe over the southeastern North Atlantic (the Aerosol Characterization Experiment (ACE) 2, Quinn and Bates, 2005), from the Indian subcontinent over the Indian Ocean (the Indian Ocean Experiment (INDOEX), Quinn and Bates, 2005), from eastern Asia over the western North Pacific (ACE-Asia) (Quinn and Bates, 2005), and from Mexico City (the Megacity Initiative: Local and Global Research Observations (MILAGRO), Molina et al., 2010), among others.

Despite the rich detail that emerges from comprehensive, short-term campaigns, such studies are not sufficient to produce the reliable aerosol climatologies needed for model development and testing (Heintzenberg et al., 2003). Intensives are typically conducted over relatively narrow ranges in time and space. Because the chemical and optical properties of plumes continue to evolve following advection downwind of study regions, such campaigns may not capture the full impact of aerosols from a given source type or region on radiative transfer nor can they directly evaluate the representativeness of results in the context of interannual variability. In addition, most investigations have been conducted during warmer months when faster photochemistry and higher temperatures sustain higher oxidant levels and faster rates of chemical transformation and secondary aerosol production. During warmer months, ammonia emitted from agricultural activities and reaction products of non-methane hydrocarbons emitted by terrestrial biota contribute significantly to the nucleation and growth of new particles whereas during colder months these sources are greatly reduced or essentially shut down. During cooler months, direct emissions of aerosols from deflation of surface soils and of aerosols and precursors from biomass burning are also generally lower, the temperature-dependent phase partitioning of semi-volatile species is shifted towards the particulate phase, and some countries (including the US) suspend regulatory controls on major aerosol precursors such as NO_x emitted from power plants. Finally, wind fields (and the associated production of marine aerosol from the surface ocean),

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cloud cover, and precipitation, all which modulate aerosol evolution, vary seasonally over most of the globe. Consequently, the physicochemical characteristics of pollutant plumes, the nature and rates at which they evolve chemically and optically, and their areal extents vary seasonally, which poses challenges for extrapolating results from short-term intensive campaigns over space and time. Remotely sensed aerosol properties from ground networks such as the Aerosol Robotic Network (AERONET) (e.g., Holben et al., 2001) and the micro pulse lidar network (MPLNET) (Welton et al., 1999) and from the growing constellation of earth-observing satellites provide useful context in this regard. However, the in situ observations necessary to constrain relationships between aerosol chemical composition, radiative effects, and climatic consequences over multi-year periods are quite limited.

Longer-term records of aerosol properties that compliment results from intensive campaigns have been generated at several observatories globally including sites in the North and South Pacific as part of the Sea-Air Exchange Experiment (SEAREX) (Prospero and Savoie, 1989) and in the NAO region as part of the Atmosphere Ocean Chemistry Experiment (AEROCE) (Arimoto et al., 1995; Savoie et al., 2002) among others. These data records are used widely by the community to parameterize and test global chemical transport and climate models. Cost and logistical considerations dictate that such long-term programs target subsets of key analytes that can be quantified reliably using measurement techniques that are both robust and require minimal servicing in the field. Similarly, temporal resolution in measurements may be sacrificed to accommodate generation of longer-term records.

The study through which results reported herein were generated was a three-year effort designed to augment and extend the data records for aerosol and precipitation characteristics at Bermuda, which are among longest-duration high-quality time series that has been generated to date (Galloway et al., 1982, 1989, 1993; Arimoto et al., 1995; Savoie et al., 2002). Bermuda is also significantly influenced over the year by airmasses containing aged emission products from North America, North Africa, and Europe as well as aged, relatively clean marine air from the central North Atlantic

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Ocean. The primary objectives of the study were (1) to extend the existing time series of aerosol and precipitation characteristics at Bermuda and evaluate long-term trends, (2) to augment the resolution of the analytical suite by characterizing dichotomous rather than bulk aerosol and by measuring additional aerosol properties including sizeresolved spectral absorption and elemental and organic carbon (EC/OC) and (3) to develop a climatology of aerosol and precipitation properties as functions of season and transport regime and the associated source regions. This paper reports the flow climatology for aerosol chemical and optical properties based on the three-year period of observation and evaluates the associated environmental implications.

Methods

Sampling site, periods, and protocols

From July 2006 through June 2009, the chemical and physical properties of ambient aerosols were characterized at the Tudor Hill Atmospheric Observatory (THAO) on the western coast of Bermuda (32.27° N, 64.87° W). The observatory includes a 23 m tall scaffolding tower and two laboratory containers configured with clean benches, deionized water, and associated electronics positioned on a steep slope approximately 25 m above mean sea level. To minimize local influences, time-integrated sampling was controlled by sensors that activated pumps only during periods of no precipitation when surface winds were off the ocean sector at speeds greater than 1 m s⁻¹. Meteorological conditions were recorded continuously from the top of the tower.

Dichotomous aerosols (nominal super- and sub-um-diameter size fractions) were sampled in parallel at a rate of 0.1 m³ min⁻¹ with two custom-designed and fabricated MSP model 130 high-flow cascade impactors configured with Liu-Pui type omnidirectional inlets (Liu et al., 1983) that were deployed at the top of the tower. The calculated passing efficiency through the inlets for 20 µm-diameter particles was 95 % and the 50% aerodynamic cut diameter between the two size factions was 0.8 µm at ambi-

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 $(\sim 15~s)$. A paired set of blank impactors was exposed and processed approximately once every two weeks. Impactors were cleaned with $18\,\mathrm{M}\Omega\,\mathrm{cm}^{-1}$ deionized water (DIW), and dried, loaded, and unloaded in a Class 100 clean bench in a laboratory container at the base of the tower. The clean bench was configured with an activated charcoal scrubber on the inlet to remove reactive trace gases from feed air. After recovery, exposed samples were folded in half, sealed in polyethylene bags, and stored (and shipped) frozen prior to analysis. Samples and corresponding blanks during each period were processed and analyzed using identical procedures.

ent RH. Relative to MSP's Micro-Orifice Uniform Deposit Impactor (MOUDI) (Marple

et al., 1991) that is in more widespread use by the research community, these hiflow impactors yield greater signal per unit deployment time while segregating aerosol

size fractions using similar nozzle technology. Air volumes were measured with sharpedged flow tubes and normalized to standard temperature and pressure (0°C and 1

atm). One impactor was configured with quartz-fiber (Pallflex 2500 QAT-UP) substrates (75 mm diameter) and back filters (90 mm diameter) for analysis of major ionic con-

stituents and organic and elemental carbon (EC/OC). The other impactor was config-

ured with Whatman 41 substrates and back filters for analysis of spectral absorption and insoluble aerosol mass. Analytical methods are described below. All data for samples were blank corrected based on analysis of impactor substrates and back filters

that were deployed at the top of the tower and through which air was pulled only briefly

Scattering coefficients at 530 nm for aerosols less than 10 µm dry diameter (total $\sigma_{s,530})$ and less than 1 µm dry diameter (sub-µm $\sigma_{s,530})$ were measured continuously at one-minute intervals by a Radiance Research 903M Integrating Nephelometer (Li et al., 1996). Scattering coefficients for the super-µm size fraction were calculated by difference (super-µm $\sigma_{s,530}$ = total $\sigma_{s,530}$ – sub-µm $\sigma_{s,530}$). Sample air was drawn though an omni directional inlet at the top of the tower and transmitted under laminar flow to the instrument at the base of the tower. The instrument was calibrated every other week with clean CO_2 (Anderson and Ogren, 1998). The sampling sequence for the

nephelometer was automatically switched among each of the two size-ranges and filtered air (to correct for Rayleigh scattering) for 20 min periods each. Data were not corrected for estimated truncation errors associated with forward scattering by large particles (Li et al., 1996). To facilitate direct comparison, scattering coefficients were averaged over periods of in-sector flow during which aerosols were sampled in parallel for characterization of ionic composition and spectral absorption.

From December 2007 through June 2009, direct solar irradiance under clear-sky conditions was measured continuously at 8 wavelengths between 340 and 1020 nm with a CIMEL Electronique Model 318A spectral radiometer deployed at the top of the tower as part of AERONET (e.g., Holben et al., 2001). Aerosol optical thickness (AOT) was calculated from spectral extinction of direct beam radiation at each wavelength based on the Beer-Bouguer Law. From March 2008 through June 2009, vertical profiles of aerosol extinction and backscatter and cloud heights were measured continuously with a micro pulse lidar deployed in one of the laboratory containers and operated as part of MPLNET (Welton et al., 1999; Reid et al., 2003).

2.2 Sample analysis

Dichotomous aerosols sampled on quartz substrates and filters were shipped to and analyzed at the University of Virginia (UVA). Half sections were extracted under sonication in 6.0 mL DIW and major ions (including CH_3COO^- , $HCOO^-$, $(COO)_2^{2-}$, $CH_3SO_3^-$, SO_4^{2-} , CI^- , Br^- , NO_3^- , NH_4^+ , Na^+ , K^+ , Mg^{2+} , and Ca^{2+}) were analyzed by IC using procedures similar to those described by (Keene et al., 2009). OC and EC in quarter sections of samples were measured in triplicate by the transmission charring compensation technique (Birch and Cary, 1996) using a Sunset Laboratory, Inc. instrument. Data were not corrected for possible absorption of gas phase organics (e.g., Novakov et al., 2000) and thus represent upper limits. The remaining quarter sections were archived frozen.

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Paired samples on Whatman 41 substrates and filters were shipped to the University of Miami (UM) for non-destructive analysis of spectral absorption ($\sigma_{a,1}$). Light absorption between 300 to 1100 nm at 10 nm intervals with a 5 nm bandwidth was measured in triplicate with an Optronics OL 740A Spectroradiometer coupled to an OL 740-70 Diffuse Reflectance Attachment (DRA) (Ball et al., 2003; Aryal et al., 2013). The DRA is an integrating sphere through which a monochromatic light beam probes the sample. The diffuse reflectance technique is comparable to the optical reflectance method employed by Reid et al. (1998) except that the latter utilizes a broadband light source and does not use an integrating sphere to capture all of the reflected light. Because on-shore flow at Bermuda is not impacted by major near-field emissions of organic aerosols or precursors from urban or industrial sources, potential artifacts associated with filter-based techniques (e.g., Lack et al., 2008) are assumed to be negligible. Following analysis of absorption, Whatman 41 substrates and filters were shipped to UVA where whole samples were extracted in three sequential 4 to 5 mL aliquots of DIW. dried, and subsequently combusted at 500 °C. The residual mass of refractory particulate material (primarily mineral aerosol) was quantified gravimetrically (Savoie et al., 1992).

Data quality and comparability

Data for aerosol samples that corresponded to in-sector sampling times of less than 10% of a day (2.4 h) exhibited poor signal-to-noise for most analytes and were of questionable representativeness; these data were excluded from the final quality-assured data set. In addition, five samples contained unusually high concentration of sea-salt constituents (30% to 380% higher than the sample with the next highest concentrations) and/or operator notes that suggested direct contamination by rainwater or splash; these samples were also excluded from the final quality-assured data set. The final data base included results for 311 individual samples.

Detection limits (DLs) for all ionic constituents, EC/OC, and non-soluble aerosol mass were estimated following Keene et al. (1989). DLs for spectral absorption were

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estimated by propagating the minimum detectable reflectance (0.5%) through the data reduction for each sample. In-sector collection times for samples included in the data base varied from 10% to 100% of the corresponding daily deployment times. In addition, signal-to-noise varied among analytes. Consequently, DLs were calculated individually for each analyte in each sample and, thus, the numbers of detectable values varied among analytes. Analytical performance was verified by intercomparison of the UVA laboratory with UM and the University of New Hampshire, among others; routine analysis of audit solutions from the National Institute of Standards and Technology, the World Meteorological Organization, and the US Environmental Protection Agency; periodic analysis of standard additions to samples; and evaluation of ion balances and diagnostic ratios of analyte concentrations. These assessments indicate that the results are unbiased and precise to quantifiable levels.

The inlet for the impactor effectively passed bulk aerosol whereas the inlet for the nephelometer was configured with a 50 % aerodynamic cut of 10 µm dry diameter. For marine aerosol, this corresponds to a cut of approximately 20 µm diameter at 80 % RH. With the exception of high-wind conditions, most marine aerosol mass and scattering is associated with particles less than 20 µm ambient diameter. Consequently, under most conditions encountered during this study, the upper end of the size distribution sampled with the impactor was approximately equal to that characterized by the nephelometer. Hereafter, the sub-10-um size fraction sampled with the nephelometer will be referred to as bulk aerosol. In addition, the impactor segragated size fractions of aerosols hydrated to ambient RH at a 50 % aerodynamic cut diameter 0.8 µm whereas the nephelometer segregated at 1.0 µm dry diameter. Consequently, at the lower end of the ambient RH distribution (<50 % RH), the cut size for the dichotomous chemical and absorption data based on impactor sampling was about 0.2 µm smaller than that for the corresponding scattering data and this difference increased with increasing ambient RH. Variability in the cut size between the two devices introduced some uncertainly in interpreting the paired dichotomous data but did not impact comparability of the summed (bulk) data. Potential influences of relative humidity and aerosol hygro**ACPD**

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Based on their thermodynamic properties (Henry's Law and dissociation constants), the equilibrium phase partitionings of NH₃, HNO₃, and HCl vary as functions of aerosol solution pH, which in marine regions typically decreases with decreasing particle size (Keene et al., 2004). In addition, larger aerosol size fractions may be undersaturated with respect to the gas phase because of their smaller surface-to-volume ratios and relatively short atmospheric lifetimes against deposition (e.g., Keene et al., 2004). Consequently, NH₄⁺, NO₃⁻, and Cl⁻ associated with bulk samples of chemically distinct marine aerosol size fractions are subject to possible artifact phase changes. Dichotomous sampling as employed for this study minimizes but does not eliminate the potential for such artifacts.

2.4 Calculations

Sea-salt and non-sea-salt constituents were differentiated following (Keene et al., 1986) for all samples for which the measured concentrations for the species of interest and the sea-salt reference species were greater than detection limits (DL). As indicated above, super- and sub-µm aerosol size fractions were sampled on 75 mm and 90 mm diameter quartz-fiber substrates and filters, respectively. Background variability in concentrations of the two mostly commonly used sea-salt reference species (Na⁺ and Mg²⁺) differed between lots for the two filter sizes such that Na⁺ offered greater resolution in calculating nss concentrations associated with the super-µm-diameter aerosol size fraction whereas Mg²⁺ offered greater resolution for those associated with the sub-µm-diameter size fraction. Super- and sub-µm-diameter nss concentrations were calculated accordingly based on the composition of surface seawater reported by Wilson (1975).

Total mineral aerosol mass was calculated as 1.3 times the measured refractory ash mass based on Trapp et al. (2010b). The Single Scattering Albedo (SSA) at 530 nm

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was calculated as the ratio of scattering to extinction ($\sigma_{s,530}/(\sigma_{s,530+},\sigma_{a,530})$). SSA's are reported for the subsets of samples for which both $\sigma_{\rm s,530}$ and $\sigma_{\rm a,530}$ were greater than DLs. Although $\sigma_{\rm s.530}$ was above DLs during virtually all periods that the nephelometer was operational, $\sigma_{\rm a.530}$ was below DLs in approximately 20 % of the corresponding samples. Because the lower end of the $\sigma_{\rm a.530}$ distribution was associated primarily to the upper end of the SSA distribution, sample statistics for SSAs are considered lower limits for ambient conditions. The Absorption Angstrom Exponent (AAE) was calculated as the negative slope for a linear regression of the absorption coefficient versus wavelength between 400 to 800 nm on a log-log plot (Bergstram et al., 2007). Only samples for which absorption was greater than DLs at all wavelengths were evaluated. For some samples, visual inspection indicated that the power-law wavelength dependence of absorption varied non-linearly consistent with relatively low correlation coefficients for the corresponding regressions. Variability in the AAE as a function of wavelength has also been reported elsewhere (see Bergstram et al., 2007). For such cases, AAEs derived over the full wavelength range are not representative diagnostics for spectral dependence over that range. We included in this analysis only AAE values for which the correlation coefficients were greater than 0.5 and, thus, reasonably linear between 400 and 800 nm.

For some evaluations, analyte concentrations associated with super- and sub-µmdiameter aerosol size fractions were summed to yield bulk-aerosol concentrations. However, Mg²⁺ (together with other sea-salt species) and NO₃ were associated primarily with the super-um size fraction (97% and 97%, respectively, based on median values for samples with detectable concentrations in both size fractions) and, consequently, corresponding concentrations of these analytes in the sub-um fraction were often below DLs (24 % and 73 %, respectively, of all samples with detectable super-um concentrations). Similarly, most CH₃SO₃⁻, nss SO₄²⁻ and NH₄⁺ was associated with the sub-µm fraction (54%, 70%, and 84%, respectively, based on median values) and, consequently, corresponding concentrations of analytes in the super-um fraction were sometimes below DLs (13 %, 2 %, and 20 %, respectively, of all samples with detectable

sub-µm concentrations). For such cases, concentrations below DLs were assumed to be 0.0 and, thus, the summed values represent lower limits for total concentrations associated with ambient bulk aerosol.

2.5 Atmospheric transport

To evaluate influences of emissions in upwind source regions, a particle dispersion model FLEXPART (Stohl et al., 1998, 2005) was run in back trajectory mode (Stohl et al., 2003; Seibert and Frank, 2004). FLEXPART is driven with ECMWF analyses and models transport using grid-resolved winds (at a resolution of 0.36 degrees, nominally 40 km at the equator) while accounting for sub-grid turbulence and deep convection. The result is an estimate of total column residence time over upwind regions, which can be displayed as a retroplume plot indicative of the column integrated contributions to sampled air from upwind regions. Starting at the tower height of 23 m above the surface and over the nominal sampling window from 15:00 UTC to 15:00 UTC each day, 40 000 particles were released from the location of the measurement site and followed backward in time for 10 days. For each day, both the potential emission sensitivity retroplume, which is proportional to the particle column residence time over a given model grid cell, and the particle footprint residence time, which indicates the contribution from within the boundary layer of a given model grid cell, were calculated (Seibert and Frank, 2004; Stohl et al., 2003).

FLEXPART trajectories were classified into groups representing transport from major source regions based on fractional footprint residence times for each 24 h period over the duration of the study (all dates in the paper refer to the sample start date). Four source-region footprints were prescribed to delineate well-separated areas that were influenced predominantly by emissions from (1) the northeastern United States (NE US FP), (2) the southeastern United States and Gulf of Mexico (GULF FP), (3) Northern Africa and the eastern tropical Atlantic (N.AF FP), and (4) the North Atlantic Ocean (NAO FP) (Fig. 1). The percent footprint residence time within each of these source regions relative to the total residence time within all four regions was then calculated

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for each 24 h period. Primary transport patterns were defined as those for which the fractional footprint residence time within one source region represented 75% or more of the total footprint residence time. Because only a subset of the days could be classified based on this criterion, the influences of combined source regions were also considered. These patterns were defined as those for which the combined fractional footprint residence times within two or three source regions represented 75% or more of the total residence time in all source regions.

The following criteria were used to classify atmospheric transport to Bermuda into five regimes based on the predominant source regions. The northeastern United States (NEUS) transport regime included days with 75% or more of the footprint residence times within the NE US FP region. The transport regime from the Gulf of Mexico and southeastern United States (East-SEUS) included days with 75% or more of the footprint residence times within the GULF FP region and days when the combined residence times within the GULF FP and the NE US FP region accounted for 75 % or more of total footprint residence times. This transport regime included contributions from sources in the eastern or southeastern United States and Gulf of Mexico region. The transport regime from North Africa and the tropical North Atlantic (AFR) included days with 75 % or more of footprint residence times within the N.AF FP region and days when the combined residence times within the N.AF FP and one other secondary footprint region accounted for 75% or more of total residence times. The Oceanic transport regime included days with 75% or more of footprint residence times within the NAO FP region, days when the combined residence times within the NAO FP and either the N.AF FP or the GULF FP regions accounted for 75% or more of the total residence times, and days for which the combined residence times within three footprint regions were required to account for 75% of the total residence time. The Oceanic transport regime was generally associated with long trajectories over the ocean and anticylonic flow around the Bermuda High (BH). Finally, the North transport regime included days when the combined residence times within the NAO FP and the NE US FP regions accounted for 75% or more of total residence times, indicative of transitions

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2.6 Statistical analysis

Many of the aerosol parameters considered in this paper exhibit data distributions with significant deviations from normal. Therefore, medians are often better measures of the central tendencies of populations, which minimizes the influence of outliers. The significance of differences between designated categories such as season or transport regime were generally evaluated using non-parametric, or rank based statistical tests. The Mann-Whitney two sample median test was used to discern differences in median values of individual aerosol chemical and physical parameters between two broad seasons, defined as the warm (April–September) and cool seasons (October–March). The significance of seasonal variability in distributions of parameters was evaluated with the Kruskal-Wallis analysis of variance test. Within each season, the significance of differences in distributions of aerosol properties among transport regimes was evaluated using the Kruskal-Wallis test.

3 Results and discussion

3.1 Chemical and optical properties

Ionic Composition. Concentrations of Na⁺, Mg²⁺, Cl⁻, and NO₃⁻ were above DLs in virtually all super- μ m size fractions (Ns = 301 to 311; Table 1). Although, as noted above, concentrations in many paired sub- μ m size fractions were less than DLs, the corresponding fractional contributions to total concentrations (super- + sub- μ m) were small (2% to 12% based on median values), which implies that concentrations less

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DLs are a minor to negligible source of negative bias for the calculated bulk concentrations. Similarly, nss SO_4^{2-} , NH_4^+ , and $CH_3SO_3^-$ were greater than DLs in most sub- μ m size fractions (Ns = 264 to 309; Table 1). However, in contrast Na⁺, Mg²⁺, Cl⁻, and NO₃, these species partitioned to a greater degree proportionately between the two 5 size fractions and, consequently, their concentrations were greater than DLs in relatively larger numbers of paired super-um size fractions (Table 1). This implies that super-µm concentrations of these analytes that were less than DLs were also relatively minor sources of negative bias for the calculated bulk concentrations. Because calculated concentrations of nss K⁺ associated with the super-um size fraction correspond to relatively small differences between substantially greater concentrations of total and sea-salt K⁺ (based on Na⁺), super-µm concentrations are poorly resolved. Only concentrations of sub- μ m nss K⁺ (N = 171), are reported and interpreted.

Concentrations of major ionic constituents measured during this study (Table 1) are consistent with previously reported measurements of aerosol composition at Bermuda (Galloway et al., 1993; Turekian et al., 2001; Savoie et al., 2002) and with the longerterm record of aerosol composition generated by the Atmosphere-Ocean Chemistry experiment (AEROCE) at THAO (J. Prospero, University of Miami, data available upon request). For example, based on the AEROCE record, the average annual concentrations of Na⁺, NO₃⁻, nss SO₄²⁻, NH₄⁺, and CH₃SO₃⁻ associated with aerosol sampled in bulk at the site between 1988 and 1997 were 160, 16.2, 20.9, 17.5, and 0.409, respectively, which agree within 2% to 14% of the average concentrations reported herein (Table 1). Significant reductions in emissions of precursor NO_x and SO₂ over the US during this period contributed to the differences in particulate NO3 and nss SO₄²⁻. A detailed evaluation of temporal trends in the ionic composition of aerosols and precipitation at Bermuda is reported in a companion paper (Moody et al., 2013). The corresponding partitioning of ionic constituents as a function of size is also consistent with reported aerosol size distributions measured at Bermuda (Turekian et al., 2001) and elsewhere in the North Atlantic marine boundary layer (MBL) (e.g., Huebert et al., 1996; Keene et al., 2004, 2009; Quinn and Bates, 2005). Differences between ambient size distributions of ionic species reflect the interplay among mixtures of aerosols from different source regions and source types (i.e., primary versus secondary, natural versus anthropogenic, marine versus continental), physical conditions (temperature, RH), thermodynamics (i.e., the pH-dependence in gas-aerosol phase partitioning), size-dependent differences in atmospheric lifetimes against deposition, and air-mass history. Taken together, the above suggests that the ionic data generated by this project are representative of ambient-aerosol composition at Bermuda.

EC/OC. In contrast to the ionic data, relatively fewer concentrations of OC and EC (and of mineral aerosol mass) were greater than DLs (Table 1), which implies that the reported concentration ranges for these analytes are not representative of the lower portions of ambient concentration distributions. In addition, the exposed portion of the impaction substrates on which super-μm aerosols were sampled was a factor of 2.8 smaller than that for the back-up filters on which sub-um aerosols were sampled. Because the analytical technique for EC/OC quantifies loadings based on a standard unit area of material punched from exposed samples, analytical resolution (signal-to-noise) for the super-μm size fraction was a factor of about 2.8 greater on average than the sub-um size fraction. Consequently, some caution and an appreciation of potential limitation are warranted when interpreting these data.

Despite the above caveats, concentrations of size-resolved OC during this study (Table 1) are within the range of those associated with size-resolved aerosol at THAO during spring (average concentrations summed over the super- and sub-um size fractions of 33.9 and 76.6 nmol C m⁻³, respectively; (Turekian et al., 2003). Although the ranges overlap, the average concentration of sub-μm OC reported herein (Table 1) is about a factor of two less than those measured in the western North Atlantic MBL under conditions of North American outflow during TARFOX (average of 488 nmol C m⁻³) and NEAQS (average of 375 nmol C m⁻³) (Quinn and Bates, 2005). The average concentration of super-μm OC (Table 1) is also somewhat less than the corresponding average reported for NEAQS (62 nmol C m⁻³). In contrast, the average concentration of sub-μm OC at Bermuda (Table 1) is greater than those measured during several campaigns in

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the eastern and tropical North Atlantic MBL (less than $\sim 80 \, \text{nmol} \, \text{C} \, \text{m}^{-3}$ (O'Dowd et al., 2004; Quinn and Bates, 2005). Similarly, the average concentration of sub- μ m EC at Bermuda (Table 1) is less than those for EC associated with North American outflow during NEAQS (36 nmol m⁻³) and TARFOX (67 nmol m⁻³) and greater than those measured in the eastern and tropical North Atlantic MBL (< 8 to 15 nmol m⁻³, respectively) (Quinn and Bates, 2005).

Mineral Aerosol. Based on a mass ratio of 9.6 for residual refractory ash to Al in the tropical North Atlantic MBL (N = 1349; $r^2 = 0.95$; D. L. Savoie, University of Miami, and R. Arimoto, New Mexico State University, Carlsbad, personal communication, 2005) and the mass ratio of 1.3 for mineral aerosol to ash (Trapp et al., 2010b), the average mineral aerosol mass for bulk aerosol with concentrations greater than DLs (Table 1) is about 25 times greater than those inferred from the average annual concentrations of crustal Al measured daily at THAO between 1988 and 1990 (0.044 to 0.056 µg Al m⁻³; (Arimoto et al., 1995). This large difference reflects the fact that residual aerosol masses for most samples during our study were less than DLs (only larger dust outbreaks were detectable). In contrast, greater sample volumes (factor of about 10), bulk (versus dichotomous) sampling, and greater analytical resolution afforded by neutron activation analysis yielded detectable Al in virtually all samples analyzed by Arimoto et al. (1995). However, the average bulk mineral mass based on our data (Table 1) was about the same as that inferred from the average for Al measured by Arimoto et al. (1995) during periods of easterly flow (1.26 µg Al m⁻³ or 15.6 µg dust m⁻³). As discussed in more detail below, the larger dust events, which accounted for the upper ends of the concentrations distributions at Bermuda during both studies, were typically associated with atmospheric transport from North Africa. Relative to Bermuda, the efficient transport of dust by the trade wind regime sustains substantially higher average and peak concentrations of mineral aerosol across the tropical North Atlantic Ocean including the MBL and free troposphere at Puerto Rico (Reid et al., 2003), the MBL at Barbados (Arimoto et al., 1995; Trapp et al., 2010a), the central MBL (Bates et al., 2001), and the free troposphere at Tenerife (Maring et al., 2000).

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Scattering Coefficient. Relative to the ionic data, intermittent malfunction of the nephelometer resulted in fewer days during which scattering coefficients were measured in parallel (Table 1). Total scattering at Bermuda was within the range measured at Barbados during periods of low to moderate dust concentrations (less than 40 µg m⁻³) using an identical instrument (10 to 42 Mm⁻¹; Li et al., 1996). Because scattering varies as a function of wavelength and RH, our results cannot be directly compared on a quantitative basis with those measured at different wavelengths and RHs or, in some cases, as adjusted to ambient RHs. Qualitatively, measurements of $\sigma_{s,530}$ at less than 40% RH for bulk aerosol at Bermuda (Table 1) fall within the lower ends of ranges measured at 550 nm and 55 % RH during various campaigns in the North Atlantic MBL (Quinn and Bates, 2005; Sierau et al., 2006). The fine fraction of scattering (sub-um $\sigma_{s,530}$ divided by total $\sigma_{s,530}$) based on average values at Bermuda (0.67) is within the range of averages at 550 nm and 55 % RH reported for different air-mass types sampled in the Gulf of Maine off the northeastern US coast during ICARTT (0.64 to 0.78; Sierau et al., 2006). The fine fraction of scattering at Bermuda also overlaps the range measured at 550 nm and 55 % RH over the eastern North Atlantic under conditions of European outflow during ACE-2 (average of 0.56, Quinn and Bates, 2005).

Absorption Coefficient and AAE. The average absorption coefficient at 530 nm and less than 40 % RH for bulk aerosol at Bermuda (Table 1) is a factor of 7 less than those at 550 nm and 55 % RH in North American outflow during TARFOX and NEAQS and a factor of 5 less than that at 530 nm and 55 % RH during ICARTT. However, absorption at Bermuda falls within the range of values reported for the eastern and tropical North Atlantic (Quinn and Bates, 2005). In contrast to these other campaigns which report no measureable absorption by larger aerosols in the North Atlantic MBL, super-um aerosol accounted for about 25 % of the median and average total absorption at Bermuda during our study (Table 1). We infer that the greater analytical resolution afforded by direct measurement as used here relative to calculating proportionately small contributions by subtraction (total absorption - sub-um absorption) as employed in the other studies contributed to these differences.

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The average AAE for super-µm and sub-µm size fractions (Table 1) fell within the range of AAEs for selected cases from different field campaigns in the NAO and elsewhere reported by Bergstram et al. (2007). The corresponding ranges in values were also similar to those measured during the Gulf of Mexico Atmospheric Composition and Climate Study (Bergstram et al., 2007). AAE values of about 1.0 indicate relatively little variability in absorption as a function of wavelength and are generally associated with urban pollution plumes, in which absorption is dominated by BC (also referred to as light-absorbing carbon). Higher AAEs indicate relatively greater absorption at shorter wavelengths and are generally associated with absorption by organic carbonaceous (also referred to as brown carbon) and/or mineral aerosol constituents. AAE values less than 1.0 may result from variability in the imaginary part of the index of refraction as a function of wavelength and/or random measurement uncertainty. To evaluate the sensitivity of these population statistics to the prescribed cutoff threshold of 0.5 for the correlation coefficient (discussed in Sect. 2.4), we evaluated statistics based on a higher cut off of 0.7. The resulting average and median values for the sub-um AAE increased from 1.10 to 1.23 and from 0.78 to 0.88, respectively, and the numbers of samples above the threshold decreased from 86 to 56. These differences were driven primarily by the removal of disproportionately larger numbers of samples associated with westerly flow from North America which, as discussed in more detail below, exhibited relatively lower AAE's. We return to this point below.

Single Scattering Albedo and Aerosol Optical Thickness. The single scattering albedos (SSA) for sub-µm and bulk aerosol at 530 nm and less than 40 % RH measured at Bermuda (Table 1) fall within the ranges of values measured elsewhere in the North Atlantic MBL (Quinn and Bates, 2005; Sierau et al., 2006). Mean SSA for sub-um aerosol was 1 % less than that for bulk aerosol, which is also within the range of average differences at 550 nm and 55 % RH observed elsewhere. Although the ranges overlap, the average AOT at 500 nm at Bermuda (Table 1) is lower than those for other regions in the North Atlantic, which vary from 0.18 at 525 nm in the eastern basin to 0.38 at **ACPD**

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Summary Evaluation. Based on evaluation of data quality and size-resolved composition, comparisons with past measurements at Bermuda, and comparison with measurements in surrounding regions of the NAO MBL and elsewhere, the above assessment indicates that the ionic composition and optical properties of dichotomous and bulk aerosols reported herein are reasonably representative of the western NAO MBL at Bermuda during the period of the study. Although measurements of mineral aerosol mass, OC, and EC fall within the ranges of values measured previously on Bermuda and elsewhere over the NAO, concentration distributions are truncated and do not include the lower ends of the concentration distributions for ambient aerosols. Consequently, sample statistics for these analytes are not representative of regional conditions. However, those data are reliable, provide quantitative constraints for the upper portions of concentration distributions, and, as indicated below, are useful for diagnosing relevant characteristics of individual events.

3.2 Seasonal variability

Seasonal differences in source strengths, chemical processing, and atmospheric transport all contribute to variability in aerosol chemical and physical properties. The significance of differences in distributions of aerosol properties between the warm and cool seasons and among transport regions within each season are depicted in Table 2. Two asterisks indicate 95 % confidence in rejecting the null hypothesis that the measured distributions of each variable correspond to random selection from the same population; one asterisk implies 90 % confidence in rejecting the null hypothesis. The season with the higher median concentration of each parameter that exhibits a significant difference is indicated with a "w" for warm or a "c" for cool.

We appreciate that this relatively coarse seasonal partitioning may not adequately characterize temporal variability within seasons. For example, when data are binned by month, the maximum median monthly nss SO_4^{2-} and NO_3^- concentrations (27.0

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and 30.1 nmol m⁻³, respectively) were during May and the minimum (12.5 and 9.3 nmol m⁻³, respectively) during July and September, respectively. All of these maximum and minimum monthly medians fell within the warm season. However, when data within each season were further partitioned into five transport regimes (see below), the 5 numbers of observations for each regime were relatively low, which limited the power of associated statistical tests. In addition, as described in more detail below, much of the temporal variability in aerosol characteristics within each season was driven by changes in large-scale atmospheric transport and the associated source regions for aerosols and precursors during that season. For example, the peak median values for nss SO_{A}^{2-} and NO_{3}^{-} during May were associated primarily with efficient westerly transport from North America, which is more prevalent after spring frontal passages whereas the minimum medians during summer were associated primarily with southeasterly transport around the Bermuda High, which is more prevalent during summer. The relatively coarse seasonal partitioning adopted herein represents a compromise between evaluating seasonal versus transport-related variability based on the finite data set summarized in Table 1.

Cool Season. Relative to the warm season, air sampled between October and March was, on average, colder, drier, and associated with higher wind speeds, higher pressures and more northerly transport (Table 3). Stronger winds generated higher sea-salt concentrations (Na⁺, Mg²⁺, and associated constituents). EC associated with the submicron size fraction was also relatively higher during these months.

Warm Season. Air sampled between April and September was associated with overall warmer conditions and correspondingly higher relative humidity and lower atmospheric pressure (Table 3). Concentrations of nss SO₄²⁻ and CH₃SO₃⁻ associated with bulk aerosol, nss-Ca²⁺ and nss $SO_{_{A}}^{2-}$ associated with the super- μ m size fraction and scattering by the sub-um size fraction were significantly higher during this season. In addition, the corresponding influences of aerosols on column-integrated AOD based on both downward observations of backscattering at 550 nm from the Moderate-Resolution Imaging Spectroradiomenter (MODIS) and upward observations of scatter**ACPD**

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ing at 500 nm from the AERONET CIMEL instrument were also significantly greater. However, many analytes including NO₃⁻, NH₄⁺, nss K⁺, OC, and mineral aerosol did not differ significantly between these two seasons.

Although seasonal differences accounted for some of the measured variability for 5 individual analytes, all exhibited substantial variability within each season. To investigate the nature of this residual variability, aerosol chemical and optical properties within each season were evaluated as a function of transport history. A flow climatology was constructed from FLEXPART retroplumes and footprint residence times to characterize major source regions for constituents and associated optical properties of the sampled airmasses.

Transport climatology 3.3

Transport Patterns. Figure 1a through f depicts examples of the five transport patterns defined above. Each example illustrates the column residence time retroplume in the left panel, and the contribution of footprint residence time from the designated source regions in the right panel. This flow pattern classification scheme characterizes major features of the transport climatology to Bermuda. The advance of continental air from North America to Bermuda is driven by the transient passage of low pressure systems. Cyclonic flow patterns are associated with frontal passages, and subsequent postfrontal flow. As these low pressure systems are pushed off shore by advance of a continental high, the origin of airmasses on Bermuda that arrive from the US becomes more northerly. Anticyclones that advance to a location east of Bermuda eventually result in flow becoming more southerly, with trajectories that arrive from the southeast or the southwest around the semi-permanent feature of the Bermuda High (BH). These BH conditions result in roughly equivalent and relatively short residence times over each of the prescribed source regions; most of the time, air resides over the open ocean. However, when the high is large and persistent, airmasses with a significant influence from African source regions are transported to Bermuda.

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Transport Pattern Frequency. The frequencies of flow patterns for air arriving at Bermuda vary between warm and cool seasons (Moody and Galloway, 1988). The FLEXPART model was run for every 24 h period between 1 July 2006 and 30 June 2009 and the frequencies for each flow pattern over all days during each season are depicted in Fig. 2a and b. Cool season frontal passages and post frontal transport, which favor the transport of North American air from the NEUS and East-SEUS, accounted for 49 % and 33 %, respectively, of flow during all days; transport around the BH represented 9% of the flow; and transitional flow from the North accounted for the balance (Fig. 2a). During the warm season, transport around the BH was the dominant feature with 49% of the flow associated with an African or Oceanic source region (Fig. 2b). Only 39% of the transport was classified as having a significant contribution from the US (17% NEUS, 22% East-SEUS), and the mixed influence of flow from the North occurred 12% of the time. These results reflect large-scale seasonal changes in the position and strength of the Bermuda High, and the corresponding lowered freguency of warm season cold fronts that penetrate to the latitude of Bermuda.

The frequencies of days during which (1) near-surface flow was in sector greater than 10% of the day, (2) samples were collected, and (3) the quality-assured data yielded values that were greater than analyte DLs were substantially less than 100% of all days. Therefore, as indicated previously, the number of days during which detectable values were measured varied among analytes and, consequently, the corresponding transport climatology for each analyte also varied. For example, nss SO₄²⁻ data are available for 23% of all days during the cold season and 33% of all days during the warm season. Consequently, the transport frequencies for all days (Fig. 2a and b) differ somewhat from those corresponding to the subset of days for which data are available (Table 1). The frequencies for the subsets of all days during each season for which bulk nss SO₄²⁻ was characterized are depicted in Fig. 2c and d. This flow climatology is similar to those for analytes that were present in most samples at levels greater than DLs (Table 1). Near-surface winds that were out of the marine sampling sector were most frequent under conditions of post frontal flow from the NEUS or the North.

Because these transport conditions occur more frequently in the cool season (Fig. 2a), the transport frequencies for bulk nss SO_4^{2-} from the NEUS (41%) and North (4%) during the cool season (Fig. 2c) are somewhat less than those for all days (Fig. 2a; 49% and 9%, respectively). The corresponding differences during the warm season were relatively smaller (Fig. 2b and d).

3.4 Aerosol characteristics by flow pattern

The significance of differences in distributions of aerosol properties and meteorological conditions among the five flow patterns for each season are summarized in Table 2. Distributions of selected chemical and optical properties of aerosols associated with each flow regime during each season are depicted as box and whisker plots (10th, 25th, 50th, 75th, and 90th percentiles plus individual outlier points) in Figs. 3, 4, and 5. These results illustrate clear seasonal and flow-dependent differences consistent with previous chemical climatologies for ozone (Moody et al., 1995) and the chemical composition of precipitation (Moody and Galloway, 1988) at Bermuda. Variations in mean meteorological conditions by flow pattern are shown in Table 3.

Cool Season. The average temperature and average relative humidity differed significantly among flow patterns with relatively warmer conditions associated with Oceanic flow (transport under a Bermuda High) and relatively cooler conditions with transport from the North. Higher relative humidities were associated with Oceanic transport and flow from Africa over the tropical and subtropical NAO whereas lower relative humidities were consistently associated with transport from the NEUS (Table 3), which was dominated by post-cold frontal conditions. Relative to other flow regimes, median concentrations of nss SO₄⁻², NO₃, NH₄⁺ associated with bulk aerosol and EC associated with sub-μm aerosol were higher in flow with North American origin whereas Oceanic air contained relatively higher concentrations of CH₃SO₃ (Fig. 3). Super-μm nss Ca²⁺ was relatively higher in transport from Africa (not shown) and correlated significantly with mineral aerosol mass. Despite differences in composition, corresponding differences in the aerosol scattering and absorption during this season were not significant

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at 95 % confidence levels (Table 2). However, cool season flow was dominated by transport from North America and, consequently, the numbers of samples associated with other flow patterns were relatively few thereby limiting statistical power in evaluating the significance to variability as a function of transport regime.

Warm Season. During the warm season, flow from the NEUS was less frequent (Fig. 2) but, as in the cool season, it occurred under conditions that were cooler and drier relative to other flow regimes. The warmest and wettest airmasses arrived under the strong ridges of high pressure associated with African (AFR) transport. Absolute and relative variability in bulk nss SO₄⁻², NO₃⁻, and NH₄⁺ among flow regimes was greater during the warm season and the highest median concentrations were associated with NEUS and East-SEUS outflow from North America (Fig. 3). Relative to the cool season, CH₃SO₃ concentrations were significantly higher overall during the warm season but did not vary significantly among flow regimes. Aerosol scattering and absorption varied significantly across transport regimes (Table 2), with the greatest absolute differences associated with the sub-um size fraction (Fig. 4). Sub um EC concentrations during the warm season were lower overall than those during the cool season and varied significantly among transport regimes (Table 2, Fig. 3d). The highest median values for sub-µm EC, scattering and absorption at 530 nm were all associated with the NEUS and East-SEUS flow regimes. The higher median sub-um SSA associated with East-SEUS relative to NEUS flow indicates greater contributions of scattering to total extinction by sub-um aerosols associated with that regime (Fig. 5). In contrast, transport from the NEUS and AFR delivered aerosols that exhibit relatively greater absorption (Fig. 4). In particular, extreme events with sub-µm absorption greater than 2 Mm⁻¹ and sub-µm SSA less than 0.9 were sampled under both of these flow patterns. However, the corresponding AAEs indicate that the spectral dependence of absorption by aerosol populations from the NEUS and AFR source regions differed (Fig. 5). Absorption by mineral aerosols associated primarily with African flow increases with decreasing wavelength resulting in a relatively greater AAEs whereas absorption by combustion-derived aerosols associated primarily with the NEUS absorb more uni-

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Based on median values, super-µm OC was most concentrated in flow from the NEUS. However, the upper ends of the distributions for both the East-SEUS and North extended to relatively higher concentrations (Fig. 3f). Some of these higher concentrations were associated with identifiable biomass burning events in the eastern US and Canada, respectively. A specific example is discussed in more detail under case studies below.

It is evident from the above that emissions over continental regions surrounding the North Atlantic basin significantly influenced the chemical and optical properties of air transported to Bermuda from those regions. In contrast, Oceanic flow was associated with the lowest or near-lowest median values and the narrowest ranges in bulk nss SO_4^{2-} , NO_3^{-} , NH_4^{+} , super- μ m OC, and absorption (Figs. 3, 4). These relationships suggest that the chemical and optical characteristic of aerosols associated with the Oceanic flow regime can serve as regional backgrounds for the western North Atlantic Ocean against which the contributions of continental influences can be evaluated. We recognize that the Oceanic flow regime is influenced to some extent by emissions over surrounding continents and, consequently, the associated aerosol characteristics are viewed as upper limits for actual background conditions.

3.5 Correlations and scattering efficiency

Influences of chemical composition on the corresponding optical properties of aerosols over the western NAO were evaluated as a function of transport regime using bivariate correlations based on linear least squares regressions. Considering all data together, sectored bulk scattering at 530 nm during the warm season was significantly correlated with the paired bulk nss SO_4^{2-} concentration (r = 0.84, n = 81), indicating that overall, 71 % of the variability in total scattering can be explained by variability in bulk nss

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 SO_4^{2-} . However, the corresponding slopes for paired data associated with individual flow regimes differed significantly (Fig. 6). Table 4 includes the regression results for events characterized as transport from the NEUS, transport from AFR, transport associated with three outliers (identified as the yellow crosses in Fig. 6), and transport from AFR including the three outliers. The three outliers correspond to individual events that were transported from the NEUS but contained high concentrations of mineral aerosol. Based on their composition and radiative characteristics, these events were similar to dust events associated with transport from Africa. They are discussed in more detail in Sect. 3.7. Regression lines for the NEUS (r = .91, n = 18) and AFR + outliers (r = .79, n = 27) flow patterns exhibit the greatest divergence and bracket most of the paired data associated with the other flow regimes. Several other outliers (identified

with squares) are also discussed in more detail in Sect. 3.7.

Aerosol components other than nss SO_4^{2-} (including primary marine aerosol, crustal dust, and organic matter) also scatter incident solar radiation (e.g., Li et al., 2006). As discussed previously, many individual measurements of mineral aerosol and OC were less than DLs and, consequently, their associated contributions to scattering cannot be apportioned directly. However, the significant correlations evident in Fig. 6 (molar units) and Table 4 (same data in mass units) indicate that the relative mixture of most light-scattering aerosol components within each transport regime typically co-varied with nss SO_4^{2-} and, consequently, bulk nss SO_4^{2-} represents a reasonable proxy and a useful predictive tool for scattering by the mixed aerosol populations within each flow regime during the warm season. The lower limit for individual measurements of scattering coefficients associated with all transport regimes (about 5 Mm $^{-1}$) represents background scattering by particles such as mechanically produced marine aerosols that are uncorrelated with nss SO_4^{2-} .

Scattering efficiencies are conventionally reported in units of m 2 gm $^{-1}$. Slopes for regressions of bulk scattering (bulk- $\sigma_{s,530}$) at 530 nm versus bulk nss SO $_4^{2-}$ mass for the AFR and NEUS flow regimes (11.4 vs. 4.3 m 2 gm $^{-1}$) differ by a factor of 2.7 and the corresponding correlation coefficients indicate that these relationships account for

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greater than 81% and 62% of the associated variability in light scattering respectively (Table 4). The greater scattering per unit mass in AFR flow represents implicit contributions from other associated aerosol constituents including mineral dust and organic carbon that are transported with the SO_4^{2-} . The fine fraction of scattering also differs between these regimes. Based on median values, sub- μ m aerosol accounts for 75% of total scattering in NEUS flow but only 55% in the AFR flow. These differences are driven in part by the larger mass median diameter of mineral aerosol at Bermuda (2.0 to 2.4 μ m, Arimoto et al., 1997) relative to pollutant aerosol from North America (less than 1 μ m).

Considering only scattering associated with the sub- μ m size fraction (for the purpose of comparison to the literature), we still find a factor of two greater mass scattering efficiency (MSE) per unit nss SO_4^{2-} mass in AFR+outliers relative to NEUS flow (9.5 versus $4.7\,\mathrm{m}^2\,\mathrm{gm}^{-1}$), again, indicative of the combined influences of sub- μ m nss SO_4^{2-} and associated scattering aerosol components. However, the lower correlation coefficients indicate that these relationships for sub- μ m aerosol explain less of the observed variability in light scattering for NEUS flow and AFR flow relative to those for bulk aerosol (Table 4).

Previous studies have found that median MSEs at 550 nm and 55 % RH based on the total mass of all sub-µm aerosol components range from 2.97 m² gm⁻¹ for continental air containing pollutants and dust during ACE-Asia to 4.73 m² gm⁻¹ in air that had been over the ocean for greater than 5 days during the INDOEX campaign (see Table 6., Bates et al. 2006). MSEs at 550 nm and 55 % RH based on total sub-µm mass over the WNAO during the ICARTT campaign ranged from 2.5 to 5.0 m² gm⁻¹, and at lower wavelengths from 3.0 to 7.0 m² gm⁻¹ (Bates et al., 2006). Li et al. (1996) employed a MSE for nss SO₄⁻² over the North Atlantic of 2.8 m² g⁻¹, based on observations near the Azores (Hegg et al., 1994), a location that is generally influenced to a lesser degree by dust and North American emissions than Bermuda. MSEs based on nss SO₄²⁻ mass and dry aerosol are not directly comparable quantitatively with those based on total mass at 55 % RH as reported by Bates et al. (2006). However, based

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on the reported aerosol composition, the two data sets do reveal a consistent picture of relatively greater scattering per unit nss SO_4^{2-} mass in marine air that is strongly influenced by mineral aerosol relative to polluted continental emissions. The higher MSE relative to nss SO₄²⁻ in African flow compared to NEUS flow reflects contributions to scattering by mineral aerosol. The outliers, representing dust events associated with transport from the NEUS, are consistent with this interpretation, and will be discussed further in Sect. 3.7.

3.6 Anthropogenic versus biogenic nss SO₄⁻²

Based on a detailed evaluation of ionic and elemental tracers associated with aerosols measured at Bermuda from 1989 to 1991, Savoie et al. (2002) derived a mass ratio of 18.8 ± 2.2 for the ratio of nss SO_4^{2-} to $CH_3SO_3^-$ produced from the atmospheric oxidation of biogenic (CH₃)₂S emitted from the ocean surface. This ratio did not vary seasonally and was statistically indistinguishable with comparable ratios derived independently for the tropical NAO at Barbados (19.6 ± 2.1) and measured in the remote tropical South Pacific Ocean at American Samoa (18.1 ± 0.9). Using the ratio derived for Bermuda, relative contributions of nss SO_4^{2-} from anthropogenic versus marine biogenic sources in the current data set were differentiated and compared with results reported by Savoie et al. (2002). The reported uncertainties correspond to those for the SO₄²⁻/CH₃SO₃⁻ ratio reported by Savoie et al. (2002) and do not reflect propagation of errors associated with the measured total nss SO_4^{2-} and $CH_3SO_3^-$ in this dataset. Consequently, the reported uncertainties represent lower limits.

Based on average values, total nss SO_{λ}^{2-} concentrations decreased by 17% between the earlier measurement period (1989 to 1991) reported by Savoie et al. (2002) and that evaluated herein (2006 to 2009) (Table 5). These decreases were driven in part by large reductions in SO₂ emissions over United States as mandated by the Clean Air Act. Between 1988 and 2009, SO₂ emissions decreased by 64% (http://www.epa.gov/ttn/chief/trends/) resulting in decreased production of particulate

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SO₄²⁻ (http://www.epa.gov/airtrends) and consequently reduced export of pollutant S over the western North Atlantic Ocean. Similarly, recent analyses of fine particle mass and EC from national parks and other remote locations show a 25% decrease between 1990 and 2004 (Murphy et al., 2011). A more complete comparison of the long term trends in Bermuda aerosol and precipitation composition are reported by Moody et al. (2013).

Although the average CH₃SO₃ concentration for the entire period of record from 2006 to 2009 was higher than that for 1989-1991, the ranges for annual averages overlapped (Table 5). Relative to 1989–1991, anthropogenic contributions during 2006 to 2009 were proportionately smaller (by 14.6%) and marine biogenic contributions proportionately larger (Table 5). These data were also stratified by transport pattern and by season. The results indicate that during the more recent period of observations, flow from NEUS was associated with much greater anthropogenic contributions (75%) relative to the transport from Africa (25%). Anthropogenic contributions to mean nss SO₄²⁻ concentrations associated with Oceanic flow were variable (note the percent error in Table 5) but statistically indistinguishable from 0.0%. These results imply that overall the events classified as the Oceanic flow regime represent a reasonable background for anthropogenic particulate nss SO_4^{2-} over the western NAO. Relative to the warm season, the higher frequency of transport from North America to Bermuda during the cool season coupled with the lower nss SO_4^{2-} from biogenic sources is evident in the greater fractional contribution of anthropogenic nss SO₄²⁻ from October through March (Table 5).

Case studies for events associated with different transport regimes

More detailed evaluations of specific cases yield additional insight regarding the chemical and optical properties of aerosols over the western North Atlantic Ocean as a function of transport region and the correspondence between surface observations and regional satellite total column measures of AOD and EAE. For example, the highest min-

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eral aerosol concentrations observed on Bermuda during the current study occurred in association with a large dust outbreak transported from Africa between 28 July and 3 August 2006. Figure 1c depicts one of the flexpart retroplumes (31 July to 1 August) during this period. During the first day of the event, satellite observations by MODIS revealed a high regional AOD of 0.62 at 550 nm and an EAE of 0.354, indicative of number size distributions dominated by super-um diameter particles. Mineral aerosol mass measured at the THAO was 45.6 µg m⁻³ with the majority (57%) associated with the super-um size fraction. Dusty conditions persisted for several days; on 1 August, the MODIS AOD was 0.40 at 550 nm, the EAE was 0.44, and the dust concentrations at THAO increased to a maximum of 71.0 µg m⁻³, with 61 % in the super-µm size fraction. Unfortunately, scattering data are not available for this period. However, the event was associated with high absorption by bulk aerosol (σ_{a530}) ranging from 2.4 to 2.7 Mm⁻¹ between 28 July and 1 August. Absorption also varied significantly with wavelength, yielding AAEs ranging from 2.56 to 2.72, which are consistent with the wavelength dependence of absorption by dust. Sub µm aerosols accounted for 75 % and 65 % of absorption on 28-29 July and 1-2 August, respectively. The relatively high CH₃SO₃ concentrations associated with bulk aerosol on 28-29 July and 1-2 August (1.12 and 0.88 respectively) coupled with the ratio of biogenic SO₄²⁻ to CH₃SO₃⁻ reported by Savoie et al. (2002) suggests that approximately 74% and 47%, respectively, of the associated nss SO₄²⁻ originated from (CH₃)₂S oxidation. The anthropogenic contribution to nss SO_{A}^{2-} during this period increased with time as the transport shifted from mostly African influenced to transport that had a greater contribution from the eastern United States. The relationships between bulk absorption versus bulk nss SO₄²⁻ on these days are depicted with yellow asterisks in Fig. 7.

Next we illustrate a three-day event from 18 to 20 June 2007 that was associated with high absorption and scattering coupled with flow that transitioned from the North to the East-SEUS. MODIS imagery on 19 June showed a large plume of smoke and haze near Bermuda (Fig. 8) that had been pushed out from eastern North America with a frontal passage. MODIS imagery also revealed relatively high regional AOD (daily av-

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erages of 0.19, 0.37, and 0.31 at 550 nm) and high EAEs (daily averages of 1.2, 1.1, and 1.4 at 550 nm) indicating that sub-um diameter particles dominated column extinction. The relatively high bulk σ_{a530} (1.74, 1.36, and 1.15 at 530 nm) measured at THAO did not vary significantly with wavelength yielding AAEs less than 1.0 suggesting absorption dominated by combustion-derived aerosols. Concentrations of elemental and organic carbon shifted over the three days from only super-um observations above DLs on June 18 (32.3 OC and 8.1 EC nmol C m⁻³), to those associated with the sub-um size fraction above DLs on 20 June (109.5 OC and 6.1 EC, nmol C m⁻³). Sub-µm nss K⁺ concentrations were close to the warm season average throughout the event (ranging from 0.58 to 0.62 nmol m⁻³) and bulk nss SO_4^{2-} was present at high concentrations (ranging from 84.9 to 105.6 nmol m^{-3}). The sub- μm size fraction accounted for 86 % to 90% of bulk scattering (51.2 to 58.7 Mm⁻¹) during the period. Absorption coefficients were also dominated (70 % to 80 %) by the sub-µm size fraction. The relationships between bulk scattering ($\sigma_{\rm s530}$, Fig. 6) and bulk absorption ($\sigma_{\rm a530}$, Fig. 7) versus nss SO_4^{2-} for this event are depicted in the figures with squares. Smoke and haze visible in the MODIS image (Fig. 8) coupled with transport from the eastern US and the associated high nss SO_4^{2-} concentrations suggests that the chemical and optical properties of air during this period represent a mixture of aerosols from both biomass burning and fossil-fuel combustion.

During spring 2009, we characterized three events associated with NEUS flow that were distinct outliers in the plot of scattering versus nss SO₄²⁻ (Fig. 6, yellow crosses) and Table 3. The presence of detectable mineral aerosol and the relatively large contributions of super-µm aerosol to scattering and absorption relative to nss SO₄²⁻ concentrations are more typical of dust laden air transported from Africa (such as the event described above) than NEUS pollution events. For example, on 21 April 2009 transport from the NEUS (depicted in Fig. 1a) the mineral aerosol concentration was $54.4 \, \mu g \, m^{-3}$. Half of the mineral aerosol mass was associated with the super-um size fraction and, relative to other NEUS events, much more of the scattering and absorption was driven by the super-um size fraction. All three events were associated with elevated column









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aerosol optical depth (measured by MODIS, 0.582, 0.311, and 0.222) and satellite EAEs that were small (0.43, 0.56, and 0.70), indicative of large particles. Visual inspection of the filters for 21 April indicated a yellowish tan hue to the aerosol deposit, which was distinct from the more reddish hue filter associated with high concentrations 5 of mineral dust of African origin. These events had moderate nss SO_4^{2-} concentrations and proportionately greater amounts were associated with super-um particles (average of 69%) relative to other NEUS flow. These three events are also identified in Fig. 7 with the yellow crosses, and based on their absorption characteristics, two of the days were similar to the absorption characteristics of the major dust event from the summer of 2006, discussed above. Polluted Asian dust can contribute significantly to North American background air during spring (Koch et al., 2007; Yu et al., 2012). In addition, simulations with the NAAPS model output archive, available from the Naval Research Laboratory Aerosol website (http://www.nrlmry.navy.mil/aerosol/) suggest the presence of significant Asian dust over eastern North America during this period. Based on the above, we hypothesize that these unususal events represent a mixture of polluted dust of Asian origin with polluted air from the NEUS resulting in significant scattering and absorption, relatively low SSA's of 0.92 (bulk) and 0.89 (sub-µm), and higher than average NH₄⁺.

The transport and aerosol characteristic associated with the events discussed above reveal contributions from multiple sources that often preclude simple definitive designations of source type for individual events. Nevertheless, viewed in aggregate, it is evident that physiochemical characteristics of aerosols diverge systematically among broadly different source regions.

Aerosol optical depth

Median values and distributions of satellite derived AOD at 550 nm from MODIS observations that correspond to the NEUS and AFR transport regimes differ significantly from the relatively lower values associated with the Oceanic regime during the warm season (Fig. 9a). Similar relationships are evident in the smaller dataset of AOT₅₀₀ measured

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over Bermuda by AERONET between December 2007 to June of 2009 (Fig. 9b). AOT values are systematically lower than AOD by about 0.02 to 0.06, consistent with previously reported comparisons (Smirnov et al., 2006). These differences result from several factors including the fact that satellite AOD measures backscattered radiance whereas AOT is primarily measuring forward scattered radiance. While there are no significant differences in the medians or distributions of AOD in NEUS versus AFR flow, there are differences in the MODIS EAE that reflect overall differences in the size distribution of the aerosol (EAE = 0.86 for NEUS and EAE = 0.74 for AFR). Previously it was shown that bulk nss SO₄²⁻ correlates significantly with bulk scattering. Figure 10 shows that surface measurements of bulk nss SO_4^{2-} are also correlated with column measurements of AOD; and again, there are apparent differences in the relationships for NEUS versus AFR.

The AOD data coupled with published satellite derived estimates for the radiative efficiency of aerosols per unit optical depth in North African (-35 to -55 W m⁻² δ^{-1}) and North American outflow over the ocean (-27 to -60 W m⁻² δ^{-1}) (Anderson et al., 2005) provide information with which to evaluate the potential magnitude of cooling associated with shortwave scattering and absorption by aerosols from continental sources at Bermuda, using a simplistic evaluation of the change in AOD (Table 6). Radiative efficiencies corresponding to the differences between the median AODs for the NEUS (median of 0.22) and AFR (median of 0.20) versus Oceanic regimes (median of 0.16) suggests that transport from the NEUS is associated with a top of the atmosphere (TOA) direct radiative forcing on the order of -1.6 W m⁻² (radiative cooling), and transport from AFR is associated with a TOA radiative cooling of -2.5 W m⁻², relative to the Oceanic background. Cooling associated with extreme events may be much greater. For example, based on the above approach, AOD associated with the dust event of summer 2006 discussed above (0.62) would correspond to a TOA radiative forcing of -16 to -25 W m⁻² relative to the Oceanic background. Similarly, AOD associated with the anthropogenic pollution and biomass burning event in June of 2007 (0.37) would correspond to a TOA radiative forcing of -6 to -13 W m⁻². We appreciate that this approach does not consider potential complicating influences associated with variability in the oceanic background as a function of transport regime. In this regard, wind velocity (and the associated production of marine aerosol, which scatters incident solar radiation) did not vary significantly among flow regimes during either season (Table 2). Furthermore, we do not have estimates of the relative cloud cover associated with these different flow patterns, which will significantly impact the areal influence of these radiative effects. Despite such limitations, these results do provide useful perspective for the relative impacts of aerosols from different continental sources on direct radiative forcing over the western North Atlantic Ocean.

As a comparison for these estimates, Leibensperger et al. (2012) used historical emission inventories and the GEOS-Chem model to simulate monthly mean aerosol distributions for North America. These were used with the GISS GCM to calculate the direct radiative forcing over the eastern US (east of 100° W). They report a cooling $-2.0\,\mathrm{W\,m^{-2}}$ in the period from 1970–1990, decreasing to $-1.2\,\mathrm{W\,m^{-2}}$ from 1990–2010 as a result of emission controls and declining SO₂ concentrations. The IPCC direct radiative forcing of aerosols globally is a mean cooling of $-0.5\pm0.4\,\mathrm{W\,m^{-2}}$ (Forster et al, 2007); Leibensperger et al. estimate that US sources contributed about $-0.03\,\mathrm{W\,m^{-2}}$ (or $\sim 6\,\%$) to the global direct radiative forcing by aerosols in 2010.

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Concentrations, chemical compositions, and corresponding radiative properties of dichotomous aerosols varied significantly as a function of airmass origin. Greatest differences were typically observed during the warm season between continental outflow from the NEUS versus trans-Atlantic transport from northern Africa and the eastern tropical North Atlantic. Relative to the AFR flow regime, the NEUS regime was associated with significantly higher (factors of 2 to 3 based on median values) concentrations of bulk particulate nss SO_4^{2-} , NO_3^- , and NH_4^+ . The difference in bulk nss SO_4^{2-} was driven primarily by higher concentrations associated with the sub- μ m size fraction in

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transport off North America. Based on mean values, anthropogenic sources accounted for 75 % of particulate nss SO_4^{2-} in NEUS flow versus 25 % in AFR flow. Aerosols associated with NEUS flow also exhibited relatively greater scattering and absorption, which was driven primarily by the sub-µm size fraction. Absorption by super-µm aerosol associated with the NEUS and AFR regimes did not differ significantly although the superµm size fraction in AFR flow accounted for a greater fraction of total absorption. These differences in optical properties were driven by the nature of the aerosol populations. The significantly greater AAEs for AFR flow reflects the wavelength dependence of absorption by associated large mineral aerosols whereas the lower AAEs for NEUS flow are consistent with the dominance of smaller combustion-derived aerosols. Despite these differences in near-surface aerosol characteristics at Bermuda, downward column measurements of AOD from MODIS and upward column measurements of AOT by AERONET revealed no significant differences between these two flow regimes. The relatively greater contribution of super-um aerosol to total extinction in the AFR regime is evident in a weakly significant (p = 0.10) difference in the regional EAE estimated from MODIS for NEUS (0.86) versus AFR (0.72). Relative to the median AOD for the Oceanic regime (representative of a regional background), the higher AODs associated with aerosols transported by NEUS and AFR flow regimes relative to the Oceanic background account for a TOA direct radiative forcing on the order of -1.6 to -2.5, respectively. Given the overall predominance of transport from North America on an annual basis, it is reasonable to assume that the radiative influence of these aerosols accounts for a large fraction of atmospheric dimming over the western North Atlantic Ocean, and based on the lower contribution of anthropogenic sulfate in Bermuda aerosols relative to the early 1990's, our data provide supporting evidence that aerosol cooling has declined in recent decades.

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Table 1. Summary statistics for selected aerosol properties based on measured values > DLs.

Analyte	Nominal Size, um	Units	Median	Maximum	Minimum	Average	Standard Deviation	Ν
Na⁺	super	$nmolm^{-3}$	123	399	10	140	84	309
	sub	nmol m ⁻³	15	88	4	22	19	61
	bulk ¹	nmol m ⁻³	130	399	10	145	86	309
Mg ²⁺	super	nmol m ⁻³	11.7	44.2	0.03	13.8	9.7	301
-	sub	nmol m ⁻³	0.4	3.6	0.02	0.5	0.4	230
	bulk ¹	nmol m ⁻³	12.0	44.2	0.03	14.2	9.8	301
CI ⁻	super	nmol m ⁻³	116	445	1	135	98	311
	sub	nmol m ⁻³	2	17	0.3	3	3	99
	bulk ¹	nmol m ⁻³	116	446	1	136	98	311
NO_3^-	super	nmol m ⁻³	13.8	63.8	0.5	16.6	11.0	308
3	sub	nmol m ⁻³	0.5	6.0	0.1	1.0	1.2	82
	bulk ¹	nmol m ⁻³	14.0	68.9	1.2	16.8	11.2	308
nss SO ₄ ²⁻	super	nmol m ⁻³	4.8	56.7	<0.0	7.1	7.4	302
4	sub	nmol m ⁻³	9.7	58.9	0.9	13.3	10.4	308
	bulk ²	nmol m ⁻³	15.7	105.6	0.9	20.3	16.4	308
NH₄ ⁺	super	nmol m ⁻³	3.1	85.9	0.3	5.1	7.5	210
*	sub	nmol m ⁻³	10.3	51.3	1.3	13.0	10.2	264
	bulk ²	nmol m ⁻³	12.8	137.2	1.5	17.1	15.5	264
CH ₃ SO ₂	super	nmol m ⁻³	0.210	0.941	0.019	0.245	0.177	270
- 33	sub	nmol m ⁻³	0.213	1.61	0.001	0.264	0.202	309
	bulk ²	nmol m ⁻³	0.414	2.03	0.001	0.478	0.345	309
nss K ⁺	sub	nmol m ⁻³	0.487	5.53	0.042	0.591	0.630	171
OC	super	nmol C m ⁻³	33	399	1	49	60	157
	sub	nmol C m ⁻³	82	703	18	135	146	69
	bulk ³	$\rm nmolCm^{-3}$	130	916	23	203	202	54
EC	super	nmol C m ⁻³	5.2	48.1	0.5	7.3	8.4	33
	sub	nmol C m ⁻³	7.2	63.0	1.4	11.7	12.7	96
	bulk ³	nmol C m ⁻³	13.1	80.3	2.4	20.6	18.9	23
Mineral	super	μg m ⁻¹	5.7	43.5	1.5	8.8	7.5	57
Aerosol	sub	μg m ⁻¹	6.7	27.5	0.9	9.2	7.0	46
	bulk ³	μq m ⁻¹	15.4	71.0	5.2	20.4	15.3	31
$\sigma_{\rm s,530}^{4}$	super	Mm ⁻¹	4.6	23.1	0.3	5.8	4.1	170
8,550	sub	Mm ⁻¹	9.7	62.1	1.9	11.9	9.1	173
	bulk ³	Mm ⁻¹	16.1	68.7	2.9	17.7	10.8	170
$\sigma_{a.530}$	super	Mm ⁻¹	0.154	1.30	0.001	0.223	0.209	263
а,550	sub	Mm ⁻¹	0.478	2.70	0.000	0.556	0.436	291
	bulk ³	Mm ⁻¹	0.697	3.56	0.047	0.813	0.597	244
SSA ₅₃₀	-	0.968	0.999	0.739	0.957	0.045	114	
300	sub	_	0.958	0.998	0.575	0.947	0.051	137
	bulk ³	-	0.967	0.996	0.654	0.954	0.044	114
AAE	super	-	1.31	4.91	0.17	1.53	1.02	116
-	sub	-	0.78	3.49	0.30	1.10	0.78	86
AOT ₅₀₀	-	Mm ⁻¹	0.108	0.521	0.014	0.124	0.067	578

¹ Sub-um concentrations -DLs assumed equal to 0.0 (see text). ² Super-um concentrations -DLs assumed equal to 0.0 (see text). ³ Super-um concentrations -DLs assumed equal to 0.0 (see text). ³ Values for both size fractions > DLs. ⁴ Averaged daily over periods of in-sector flow during which aerosols were sampled for analysis of ionic composition and absorption. ⁵ Averaged daily (unsectored).

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Table 2. Results of nonparametric statistical tests on distributions of aerosol composition, radiative characteristics, and meteorological conditions during sampling, compared across season, and between flow patterns within each season.

Туре	Parameter	Size Fraction	Distributions by Season	Distributions by Transport Patterns Cool (Oct–Mar)	Distributions by Transport Patterns Warm (Apr–Sep)
Bulk	Na ⁺	bulk	**c	_	**
Aerosol	Mq ²⁺	bulk	**c	-	_
Chemical	CI¯	bulk	**c	_	_
Composition	NO ₃	bulk	_	**	**
	NH ₄	bulk	_	*	**
	CH ₃ SO ₃	bulk	**W	**	_
Size-Resolved	nss SO ₄ ²⁻	super	**W	_	**
Aerosol	•	sub	-	**	**
Chemical		bulk ²	**W	**	**
Composition	nss K ⁺	sub	_	_	*
	nss Ca2+	super	**W	*	**
	OC	super	_	_	*
		sub	_	-	_
		bulk ³	_	_	_
	EC	super	_	_	_
		sub	*c	**	**
		bulk ³	*c	_	_
	Mineral	super	_	_	*
	Aerosol	sub	_	**	_
		bulk ³	-	*	_
Aerosol Physical/	$\sigma_{\rm s,530}$	super	_	*	**
Radiative		sub	*w	-	**
Properties		bulk ³	_	-	**
	$\sigma_{\rm a,530}$	super	-	-	**
		sub	-	-	
		bulk ³	_	-	**
	SSA ₅₃₀	super	-	-	-
		sub	-	-	**
		bulk ³	-	-	-
	AAE	super	-	-	**
		sub		_	
	AOT ₅₀₀		**W	-	**
	AOD ¹		**W	*	**
Meteorological	avgT		**W	**	**
Conditions	avgRH		**W	**	**
	avgWS		**c		-
	avgWDir		**c	**	**
	avgPr		**c		**

^{**} p = .05, significant at 95 % * p = .10, significant at 90 % 1 AOD refers to the regional satellite value derived from MODIS.

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Table 3. Meteorological conditions recorded during sectored (sampling) flow periods. Seasonal average conditions and the average conditions associated with each Flexpart defined flow pattern.

Season	Parameter	Ai	Airmass Origin based on Transport Classification					
		Total	NEUS	East-SEUS	African	Oceanic	North	
Cool	avgT (°C)	19.0	18.4	19.6	20.4	21.5	16.8	
	avgRH (%)	69	64	73	82	81	66	
	avgWS (kt)	5.3	4.9	5.9	4.8	5.3	4.9	
	avgWDir (°)	260	265	254	243	244	276	
	avgPr (hPa)	1016	1016	1015	1015	1019	1018	
Warm	avgT (°C)	22.6	20.1	22.4	24.3	22.6	20.4	
	avgRH (%)	77	69	78	80	76	75	
	avgWS (kt)	4.5	4.3	4.8	4.3	4.4	4.6	
	avgWDir (°)	248	260	255	239	237	259	
	avgPr (hPa)	1014	1016	1013	1014	1016	1014	

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Table 4. Linear regression analyses of $\sigma_{s,530}$ (Mm⁻¹) against nss SO₄²⁻ (µg m⁻³), slopes are in units of m² gm⁻¹.

Model ¹	Correlation Coefficient	Intercept	(S.E.) ²	Sig. ³	Slope	(S.E.) ²	Sig. ³	Explained Variance(%)	n
Bulk- $\sigma_{\rm s,530}$ vs. Bulk	nss SO ₄ ²⁻								
NEUS	.82	5.1	1.9	.017	4.3	.49	.000	91	18
AFR	.88	1.0	1.7	.552	10.3	1.2	.000	77	24
OUTLIERS	.95	0.0^{4}	*	*	15.8	3.6	.049	90	3
AFR+OUTLIERS	.79	.80	2.6	.757	11.4	1.7	.000	63	27
Sub- $\sigma_{\rm s,530}$ vs. sub-	nss SO ₄ ²⁻								
NEUS	.84	3.7	2.1	.105	4.7	.74	.000	71	18
AFR	.60	0.1	2.5	.969	12.6	3.6	.002	36	24
OUTLIERS	.94	*	*	*	10.2	2.7	.064	87	3
AFR+OUTLIERS	.58	2.1	2.1	.322	9.5	2.7	.001	34	27

¹ Analyses are reported by transport pattern for NEUS flow and AFR flow. Three events identified as NEUS dust events have been designated OUTLIERS and are reported separately, and are also grouped with the other "dust" events in flow from AFR, and plotted as (+) in Fig. 6. ² Standard Error of the estimate (regression intercept or slope). ³ Significance level of *t* test. ⁴ Regression of outliers forced through 0.

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Table 5. Mean nss SO_4^{-2} and $CH_3SO_3^-$ concentrations in Bermuda data from 1989–1990 (Savoie et al., 2002) and current Bermuda data from 2006–2009, with estimates of the the percentage contribution of anthropogenic (and % error), based on the derived nssSO₄⁻²/CH₃SO₃ ratio of 18.8 (±2.2) (Savoie et al., 2002).

Datasat	T-1 00-2	011 00-	A t	
Dataset	Tot nssSO ₄ 2	CH ₃ SO ₃	Anthropogenic nss	Error
	Mean (µg m ⁻³)	Mean (μg m ⁻³)	SO ₄ ²⁻ (%)	(%)
Savoie '89–90*	2.35	.378	69.8	3.5
'90–91*	2.34	.353	71.6	3.3
Total	2.34	.366	70.7	3.4
Moody '06-07*	2.29	.488	60.2	4.7
'07–08*	1.83	.515	49.7	6.2
'08–09*	1.67	.332	61.6	4.4
Total	1.95	.455	56.1	5.1
Moody '06-09			By Flow F	Pattern
NEUS	2.54	.332	75.4	2.9
East-SEUS	2.12	.444	60.6	4.6
AFR	1.29	.514	25.2	8.8
Oceanic	1.20	.654	-2.1	12.0
North	2.37	.491	61.0	4.6
Moody '06-09 By Season				
Cool (Oct–Mar)	1.72	.258	71.7	3.4
Warm (Apr–Sep)	2.13	.602	46.8	6.2

^{*} Annual periods defined July-June.

Table 6. Estimates of the change in radiative forcing associated with changes in the median aerosol optical depth in transport from North America (NEUS) and North Africa (AFR) over Bermuda relative to the Oceanic background.

Transport Region	Δ Median MODIS AOD $(δ)^1$	Range of Radiative Efficiencty ² (W m ⁻² δ ⁻¹)	Median (Range) of Radiative Forcing (W m ⁻²) ³
NEUS	.037	-27 to -60	-1.6(-1.0 to -2.2)
AFR	.056	-35 to -55	-2.5 (-2.0 to -3.1)

 $^{^1}$ Median AOD for continental source region minus Median AOD for Oceanic source region, this dataset. 2 From Table 2, Anderson et al. (2005), based on regional aerosols over the ocean (Christopher and Zhang, 2002), the lower range of efficiency for the NEUS comes from using the Global Ocean estimate of $-27\,\mathrm{W\,m^{-2}\,\delta^{-1}}$ derived from the GOCART model (Yu et al., 2004). 3 TOA clear-sky 24 h averaged effect on solar radiative flux calculated from median values of change in AOD and reported range in radiative efficiency.

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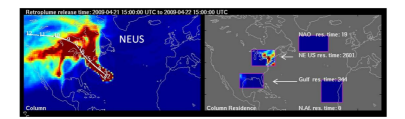


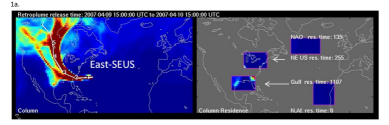
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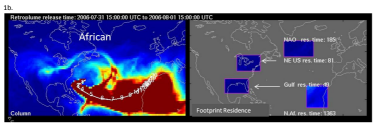
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1c.

Fig. 1. Individual examples for each of the five aggregated source region flow patterns showing the column residence time for FLEXPART retroplumes (left panels) and the corresponding residence times in the four prescribed footprint source regions (NAO, NE US, Gulf, N.Af, in right panels) that were used to assign each day to a dominant source region delivering air to Bermuda: (a) NEUS on 21 to 22 April 2009, (b) East-SEUS on 9 to 10 April 2007, (c) African (AFR) on 31 July to 1 August 2006, (d) Oceanic on 12 to 13 July 2006, and (e) North on 11 to 12 September 2007.

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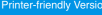
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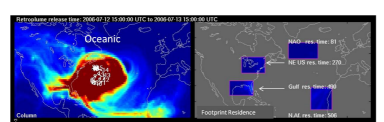
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1d.

1e.

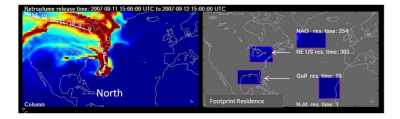


Fig. 1. Continued.

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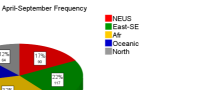
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2b.

October-March Frequency

49% 266

2a.

2c.

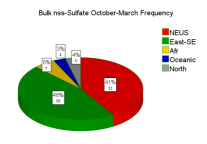
NEUS

Afr

East-SE

Oceanic

North



Bulk nss-Sulfate Frequency April-September

NEUS
East-SE
Afr
Oceanic
North

2d.

Fig. 2. Transport regime frequencies based on all days during the 3 yr duration of the project for **(a)** the cool season, October through March and **(b)** the warm season, April through September; corresponding transport frequencies based on the subset of all days for which nss SO_4^{2-} data were available during the **(c)** cool season and **(d)** warm season.

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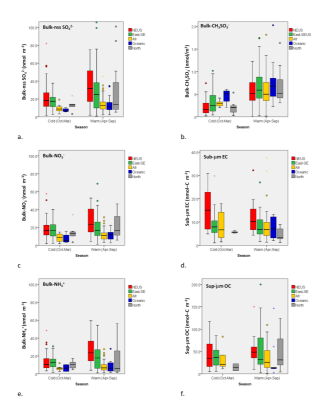


Fig. 3. Box and whisker plots depicting the 10th, 25th, 50th, 75th, and 90th percentiles and outliers (plotted individually with circles for 1.5 times the interquartile range, with asterisks for outliers 3 times the interquartile range) for (a) bulk nss SO₄²⁻, (b) bulk CH₃SO₃⁻, (c) bulk NO₃⁻, (d) sub-µm EC, (e) bulk NH₄⁺, and (f) super-µm OC associated with each of the source region flow patterns by season.

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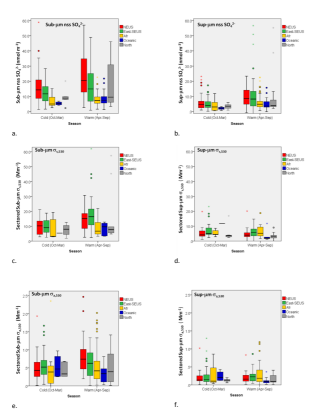


Fig. 4. Box and whisker plots (as described in Fig. 3) for **(a)** sub-μm nss SO_4^{2-} , **(b)** super-μm nss SO_4^{2-} , **(c)** sectored sub-μm $\sigma_{s,530}$, **(d)** sectored super-μm $\sigma_{s,530}$, **(e)** sectored sub-μm $\sigma_{a,530}$, and **(d)** sectored super-μm $\sigma_{a,530}$ associated with each of the source region flow patterns by season.

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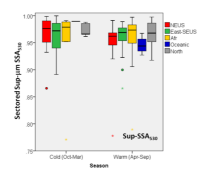
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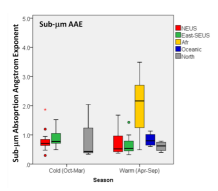
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b.



c.

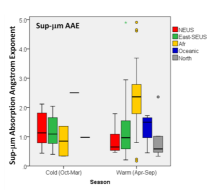


Fig. 5. Box and whisker plots (as described in Fig. 3) for **(a)** sub- μ m SSA₅₃₀, **(b)** super- μ m SSA₅₃₀, **(c)** sub- μ m AAE, and **(d)** super- μ m AAE associated with each of the source region flow patterns by season.

d.

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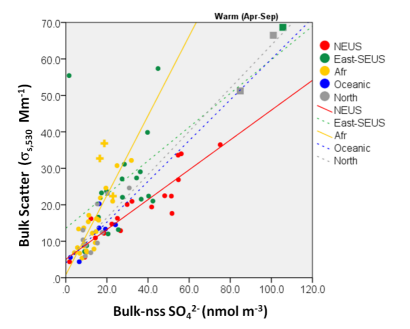


Fig. 6. Sectored bulk $\sigma_{s,530}$ versus bulk nss SO_4^{2-} associated with each of the source region flow patterns by season; color-coded lines depict the corresponding regressions.

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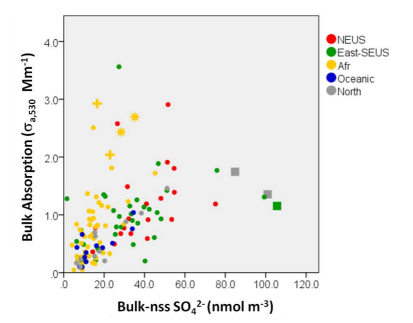


Fig. 7. Sectored bulk $\sigma_{\rm a,530}$ versus bulk nss $\rm SO_4^{2-}$ concentration associated with each of the source region flow patterns during the warm season, with specific events highlighted: asterisks mark the African dust event from 28 June and 1 August 2006; crosses mark the dust event from 21 and 23 April 2009; squares mark the polluted haze/biomass burning event from North and East-SEUS, 18, 19 and 20 June 2007.

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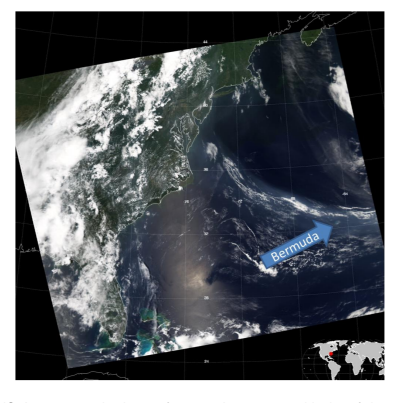


Fig. 8. MODIS Aqua true color image from 19 June 2007, midpoint of three day polluted haze/biomass burning event with very high scattering, absorption, nss SO_4^{2-} , and organic and elemental carbon, transported from North and East-SEUS, 18, 19 and 20 June 2007 as a frontal boundary advances toward Bermuda from the north.



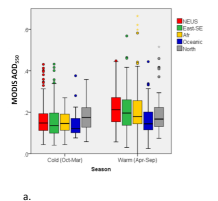
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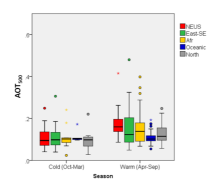
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b.

Fig. 9. Box and whisker plots (as described in Fig. 3) for (a) regional AOD measured by MODIS (Terra, at 550 nm) on all days from 2006 to 2009; and (b) Aeronet AOT (500 nm) from Bermuda January 2008 to June 2009 associated with each of the source region flow patterns by season.

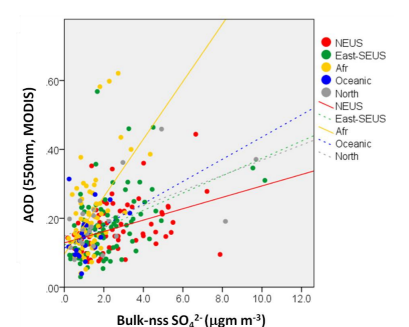


Fig. 10. MODIS regional aerosol optical depth (AOD, 550 nm) versus bulk nss SO_4^{2-} concentration by flow patterns during the warm season months of April to September; linear correlation coefficients of r = 0.51 for NEUS, r = 0.53 for AFR.

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