Aerosol optical properties during INDOEX 1999:
Means, variability, and controlling factors

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[1] As part of the Indian Ocean Experiment (INDOEX) 1999 Intensive Field Phase, measurements of aerosol properties were made on board the R/V Ronald H. Brown in the Indian Ocean north and south of the Intertropical Convergence Zone (ITCZ) in the Arabian Sea and in the Bay of Bengal. On the basis of air mass trajectories, eight air mass source regions were identified including the southern hemisphere Atlantic; southern hemisphere Indian Ocean; northern hemisphere Indian Ocean; east Indian subcontinent where trajectories came from near Calcutta, through the southern portion of India, and then to the ship; Indian subcontinent where trajectories came from across central India to the ship; Arabia; Arabia/Indian subcontinent, a mixed region where lower-level trajectories came from Arabia and upper-level trajectories came from India; and Arabian Sea/coastal India where trajectories came from along the coast of India to the ship. Properties of the aerosol measured in the marine boundary layer included chemical composition, number size distribution, and scattering and absorption coefficients. In addition, vertical profiles of aerosol backscatter and optical depth were measured. Presented here as a function of air mass source region are the concentrations and mass fractions of the dominant aerosol chemical components, the fraction of the extinction measured at the surface due to each component, mass extinction efficiencies of the individual components, aerosol scattering and absorption coefficients, single scattering albedo, Ångström exponent, and optical depth. All results except aerosol optical depth are reported at the measurement relative humidity of 55 ± 5%. For air masses that originated from the two southern hemisphere marine regions (southern hemisphere Atlantic and Indian Ocean), sea salt dominated the extinction by sub-1 μm and sub-10 μm aerosol particles. The ratios of sub-1μm to sub-10 μm extinction were the lowest measured of all air mass source regions (mean values of 28 and 40%) due to the dominance of the aerosol mass by supermicron sea salt. In addition, aerosol optical depths were the lowest measured averaging 0.06 ± 0.03. Non-sea-salt (nss) sulfate aerosol concentrations in air masses from the northern hemisphere Indian Ocean were a factor of 6 higher than those in southern hemisphere air masses, while submicron sea-salt concentrations were comparable. Sulfate aerosol made up 40% of the sub-1μm extinction, while sea salt dominated the sub-10 μm extinction. Aerosol optical depths for this source region averaged 0.10 ± 0.03. A mean single scattering albedo near 0.89 and detectable black carbon (BC) concentrations (0.14 ± 0.05 μg m−3) indicated the transport of continentally derived aerosol to the ITCZ. The two regions influenced by low-level (500 m) airflow from Arabia had higher concentrations of submicron nss sulfate, particulate organic matter (POM), and inorganic oxidized material (IOM) than were observed in the marine regions. Concentrations of supermicron IOM were comparable to supermicron sea-salt concentrations. Nss sulfate aerosol dominated the sub-1 μm extinction and made significant contributions to the sub-10 μm extinction. Sea salt dominated the supermicron extinction. Mean BC contributions to submicron extinction were 8 and 12%. Single scattering albedo values averaged 0.93 ± 0.02 and 0.89 ± 0.02 for these two source regions. Aerosol optical depths averaged 0.19 ± 0.12 and 0.38 ± 0.07 with the higher value due to upper-level (2500 m) flow from India. Regions influenced by low-level airflow from the Indian subcontinent had the highest submicron nss sulfate, POM, BC, and IOM concentrations measured during the experiment. Supermicron sea-salt concentrations were lower than or comparable to supermicron nitrate concentrations. Sub-1 μm and sub-10

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μm extinction were dominated by nss sulfate aerosol although a burning component consisting of BC, KNO₃, and K₂SO₄ made a nearly equivalent contribution. These regions had a mean single scattering albedo of 0.85 ± 0.01, the lowest measured for any region. Mean aerosol optical depths were highest (0.3 to 0.4) for regions with low-level or upper-level airflow from the Indian subcontinent. INDEX TERMS: 0305 Atmospheric Composition and Structure: Aerosols and particles (0345, 4801); 1610 Global Change: Atmosphere (0315, 0325); 0365 Atmospheric Composition and Structure: Troposphere—composition and chemistry; KEYWORDS: aerosol chemical composition, aerosol mass extinction efficiencies, aerosol mass fractions, aerosol optical depth


1. Introduction

[2] The 1999 Indian Ocean Experiment (INDOEX) Intensive Field Phase (IFP) was conducted in February and March during the winter monsoon season. The study area included the Arabian Sea, the Bay of Bengal, and the Indian Ocean north and south of the Intertropical Convergence Zone (ITCZ) (Figure 1). In this region, during this time of year, continentally derived aerosol chemical components may be transported from the Indian subcontinent and nearby nations to the marine environment where they mix with marine aerosol components. These continental chemical components include soil dust from natural and anthropogenic processes, sulfates and fly ash from industrial processes and fossil fuel combustion, and carbonaceous species from biomass burning, biofuel burning, fossil fuel combustion, and vegetation. The marine chemical components include sea salt and biogenically derived sulfates and carbonaceous species. Hence the region is ideal for assessing the radiative significance of a number of continental and marine aerosol chemical components, one of the major goals of INDOEX.

[3] INDOEX was an international, multiplatform experiment involving two ships, three aircraft, two field stations, satellite observations, and a modeling component. The National Oceanic and Atmospheric Administration (NOAA) Research Vessel Ronald H. Brown participated in the 1999 IFP with instrumentation on board capable of characterizing both surface and column aerosol properties (in situ instrumentation.
Table 1. Air Mass Regions As a Function of Day of Year During the Experiment

<table>
<thead>
<tr>
<th>Time Period</th>
<th>Air Mass Region</th>
<th>Abbreviation</th>
<th>Characterizationb</th>
</tr>
</thead>
<tbody>
<tr>
<td>41–49.5</td>
<td>500 m: SH Atlantic</td>
<td>SHAtl</td>
<td>SHmX, SHmT</td>
</tr>
<tr>
<td></td>
<td>2500: SH Atlantic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>49.5–56.58 and</td>
<td>500 m: SH Indian Ocean</td>
<td>SHIO</td>
<td>SHmT</td>
</tr>
<tr>
<td>78.25–79.5</td>
<td>2500: SH Indian Ocean</td>
<td>none</td>
<td></td>
</tr>
<tr>
<td>63.56–65.24</td>
<td>500 m: east Indian Ocean</td>
<td>EIndSub</td>
<td>NHcT</td>
</tr>
<tr>
<td></td>
<td>2500 m: east Indian subcontinent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>65.24–66.5</td>
<td>500 m: central Indian subcontinent</td>
<td>IndSub</td>
<td>NHcT</td>
</tr>
<tr>
<td></td>
<td>2500 m: east or central Indian subcontinent</td>
<td></td>
<td></td>
</tr>
<tr>
<td>67–68 and</td>
<td>500 m: Arabia</td>
<td>Arabia/IndSub</td>
<td>none</td>
</tr>
<tr>
<td>70.6–70</td>
<td>2500 m: Indian subcontinent</td>
<td>Arabia</td>
<td>NHcX</td>
</tr>
<tr>
<td>68–70.42</td>
<td>2500 m: Arabia or Indian subcontinent</td>
<td></td>
<td>NHcX</td>
</tr>
<tr>
<td>76–78.25 and</td>
<td>500 m: NH Indian Ocean</td>
<td>NHIO</td>
<td>NHmT</td>
</tr>
<tr>
<td>79.75–81.25</td>
<td>2500 m: NH or SH Indian Ocean</td>
<td></td>
<td></td>
</tr>
<tr>
<td>81.25–82 and</td>
<td>500 m: Arabian Sea/coastal India</td>
<td>AS/CI</td>
<td>NHmT</td>
</tr>
<tr>
<td>85–88.75</td>
<td>2500 m: Arabian Sea/coastal India or Indian subcontinent</td>
<td>NHcT</td>
<td></td>
</tr>
</tbody>
</table>

aAlso shown are the 500 and 2500 m trajectory information and characterization of the air mass regions.

bSHmX, southern hemisphere marine extratropical; SHmT, southern hemisphere marine tropical; NHmT, northern hemisphere marine tropical; NHcT, northern hemisphere continental tropical; NHcX, northern hemisphere continental extratropical.

Trajectory arrival height to the ship location in meters.

for the measurement of aerosol chemical, physical, and optical properties; sunphotometers for the measurement of aerosol optical depth; and a lidar for the measurement of aerosol vertical profiles. This paper presents, as a function of aerosol source region, the measured chemical and optical properties and the relative significance of the dominant aerosol chemical components in radiative forcing.

On the basis of air mass back trajectories and aerosol number concentrations and size distributions, eight air mass source regions impacted the samples collected on the Ronald H. Brown. The geographical regions along the cruise track where air masses from the different source regions were sampled are shown in Figure 1. Also shown are the prevailing trajectories for each geographical and air mass source region. Time periods when air masses from the different source regions were encountered along the cruise track are listed in Table 1. Several of the air mass source regions overlap in space but not in time as transport patterns changed each time the ship moved through the different geographical regions. The eight air mass source regions are the southern hemisphere Atlantic (SHAtl); southern hemisphere Indian Ocean (SHIO); northern hemisphere Indian Ocean (NHIIO); east Indian subcontinent (EIndSub) where trajectories came from the northeast near Calcutta, through the southern portion of India, and then to the ship; Indian subcontinent (IndSub) where trajectories came from the northeast across central India to the ship; Arabia; Arabia/Indian subcontinent (Arabia/IndSub), a mixed region where lower-level trajectories came from Arabia and upper-level trajectories came from the northeast across India; and Arabian Sea/coastal India (AS/CI) where trajectories came from the north to northwest along the coast of India to the ship. The ship ventured into the Bay of Bengal near the end of the cruise, but instrumental problems and the limited number of samples collected made it difficult to characterize the region. Therefore it is omitted from the analysis presented here.

Presented here are the concentrations and mass fractions of the chemical components (mass fraction is defined as the component concentration divided by the total aerosol mass concentration), the fraction of measured extinction due to each component, and mass extinction efficiencies of the chemical components for the different air mass regions. The chemical components considered include non-sea-salt (nss) sulfate aerosol, sea salt, particulate organic matter (POM), black carbon (BC), and inorganic oxidized material (IOM). Also presented for the source regions are average aerosol scattering and absorption coefficients, the backscattered fraction, Ångström exponent, and aerosol optical depth.

An overdetermined data set allowed closure tests to be performed between measured and calculated parameters including aerosol mass (measured gravimetrically, derived from chemical analysis, and estimated from the number size distribution) and aerosol light extinction and single scattering albedo (derived from measured light scattering and absorption coefficients and calculated from Mie theory). Hence it was possible to test for consistency in the input data to the Mie calculations required for deriving the percent of extinction due to each chemical component and to assess the accuracy of the results of those calculations. In addition, mass extinction efficiencies were calculated with two independent methods (Mie calculations and a multiple linear regression). Results of these comparisons for each air mass source region are presented.

2. Measurements

2.1. Aerosol Sample Inlet

Sample air for the chemical and optical measurements was drawn through a 6-m sample mast. The entrance to the
mast was 18 m above sea level and forward of the ship's stack. To maintain nominally isokinetic flow and minimize the loss of supermicron particles, the inlet was rotated into the relative wind. Air entered the inlet through a 5-cm diameter hole, passed through an expansion cone, and then into the 20-cm diameter sampling mast. The flow through the mast was 1 m$^3$ min$^{-1}$.

[8] The last 1.5 m of the mast were heated to establish a stable reference relative humidity (RH) for the sample air of 55 ± 5%. A stable reference RH allows for constant instrumental size segregation in spite of variations in ambient RH and results in chemical, physical, and optical measurements which are directly comparable. In addition, measurement at a constant reference RH makes it possible, with the knowledge of appropriate growth factors [Clarke et al., 2002], for end users of the data set (process, chemical transport, and radiative transfer models) to adjust the measured parameters to a desired relative humidity. A reference RH of 55% was chosen because it has been shown to reduce impactor bounce since there is enough water associated with the hygroscopic aerosol species at this RH to make the aerosol "sticky" (P. K. Quinn, unpublished results, 1995). In addition, for the atmospheric conditions encountered during INDOEX, it was possible to maintain 55% RH without excessive heating of the aerosol. On average, the aerosol was heated 5.4°C above the ambient temperature. All results except aerosol optical depth are reported at 55 ± 5% RH.

[9] Individual 1.9 cm diameter stainless steel tubes extended into the heated portion of the mast. These were connected to the aerosol instrumentation and impactors with graphite-polyethylene conductive tubing to prevent the electrostatic loss of particles. An exception to this was the lines connected to the impactors used for collection of carbonaceous aerosol; they were constructed of stainless steel. Air was sampled only when the concentration of particles greater than 15 nm in diameter indicated the sample air was free of local contamination (i.e., there were no rapid increases in particle concentrations). The 0.18 μm refers to 0.18 μm before the cruise at 600 μm at 55% RH, and 10 μm at 55% RH. The 0.18 μm jet plate was used instead of a quartz backup filter to minimize positive artifacts due to the absorption of gas phase organics. With a jet plate, spot deposits of aerosol are collected which, when summed together, have less surface area than a quartz filter. In addition, the aluminum foils used as sampling substrates absorb less organic vapor than does a quartz filter [e.g., Turpin et al., 2000]. In addition, three samples were collected with a seven-stage impactor (D$_{50,aero}$ of 0.18, 0.31, 0.55, 1.1, 2.0, 4.1, and 10 μm) to obtain size distributions of OC and EC. Sampling periods ranged from 12 to 24 hours. Al foils were heated at the Institute for Tropospheric Chemistry (Leipzig, Germany) before the cruise at 600°C for 4 hours to remove organic contaminants. Blank levels were determined by placing substrates into a second impactor and deploying the impactor for the duration of the sampling period without drawing air through it. Foils were stored frozen until analysis.

[10] Two-stage multijet cascade impactors [Berner et al., 1979] sampling air at 55 ± 5% RH were used to determine the submicron and supermicron concentrations of Cl$^-$, NO$_3^-$, SO$_4^{2-}$, methanesulfonate (MSA$^-$), Na$^+$, NH$_4^+$, K$^+$, Mg$^{2+}$, and Ca$^{2+}$. Sampling periods ranged from 4 to 6 hours. The RH of the sampled airstream was measured a few inches upstream from the impactor. The 50% aerodynamic cutoff diameters $D_{50,aero}$ were 1.1 and 10 μm. Throughout the paper, submicron refers to particles with $D_{aero} < 1.1$ μm at 55% RH, and supermicron refers to particles with $D_{aero} > 1.1$ μm at 55% RH. Sub-10 μm refers to particles with $D_{aero} < 10$ μm at 55% RH.

[11] The 12 μm grease cup at the inlet of the impactor was coated with silicone grease to prevent the bounce of larger particles onto the downstream stages. A Tedlar film placed on the impactor jet plate having a $D_{50,aero}$ of 10 μm was sprayed with silicone lubricant for the same reason. Since films placed on downstream jet plates were used for chemical analysis, they were not sprayed to avoid unnecessary contamination. Tedlar films were used as the collection substrate in the impaction stage, and a Millipore Fluoropore filter (1.0-μm pore size) was used for the backup filter. Films were cleaned in an ultrasonic bath in 10% H$_2$O$_2$ for 30 min, rinsed in distilled, deionized water, and dried in an NH$_3$- and SO$_2$-free glove box. Filters and films were wetted with 1 mL of spectral grade methanol. An additional 5 mLs of distilled deionized water were added to the solution, and the substrates were extracted by sonication for 30 min. The extracts were analyzed by ion chromatography [Quinn et al., 1998]. All handling of the substrates was done in the glove box. Blank levels were determined by loading an impactor with substrates but not drawing any air through it.

[12] Non-sea-salt sulfate concentrations were calculated from Na$^+$ concentrations and the ratio of sulfate to sodium in seawater. Sea-salt aerosol concentrations were calculated as

\[
\text{sea salt (μg m}^{-3}\text{-equiv)} = \text{Cl}^- (\text{μg m}^{-3}) + \text{Na}^+ (\text{μg m}^{-3}) \times 1.47
\]

where 1.47 is the seawater ratio of (Na$^+$ + K$^+$ + Mg$^{2+}$ + Ca$^{2+}$ + SO$_4^{2-}$ + HCO$_3^-$)/Na$^+$ [Holland, 1978]. This approach prevents the inclusion of non-sea-salt K$^+$, Mg$^{2+}$, Ca$^{2+}$, SO$_4^{2-}$, and HCO$_3^-$ in the sea-salt mass and allows for the loss of Cl$^-$ mass through Cl$^-$ depletion processes. It also assumes that all measured Na$^+$ and Cl$^-$ are derived from seawater. Results of Savoie and Prospero [1980] indicate that soil dust has a minimal contribution to measured soluble sodium concentrations.

[13] Uncertainties of the ionic chemical components at the 95% confidence level were propagated as a quadratic sum of all errors involved which assumes that all errors were random. Details of the uncertainty analysis are given by Quinn et al. [2000a]. The form of the uncertainty equations in the work of Quinn et al. [2000a] were used for all uncertainty analyses in this paper.

2.3. Regional Concentrations of Total Organic and Black Carbon

[14] Three-stage multijet cascade impactors [Berner et al., 1979] sampling air at 55 ± 5% RH were used to determine submicron and supermicron concentrations of total, organic, and black carbon [Neusüss et al., 2002]. The impactor had $D_{50,aero}$ of 0.18, 1.1 and 10 μm. Only in this case does submicron refer to 0.18 $< D_{aero} < 1.1$ μm. The 0.18 μm jet plate was used instead of a quartz backup filter to minimize positive artifacts due to the absorption of gas phase organics. With a jet plate, spot deposits of aerosol are collected which, when summed together, have less surface area than a quartz filter. In addition, the aluminum foils used as sampling substrates absorb less organic vapor than does a quartz filter [e.g., Turpin et al., 2000]. In addition, three samples were collected with a seven-stage impactor ($D_{50,aero}$ of 0.18, 0.31, 0.55, 1.1, 2.0, 4.1, and 10 μm) to obtain size distributions of OC and EC. Sampling periods ranged from 12 to 24 hours. Al foils were heated at the Institute for Tropospheric Chemistry (Leipzig, Germany) before the cruise at 600°C for 4 hours to remove organic contaminants. Blank levels were determined by placing substrates into a second impactor and deploying the impactor for the duration of the sampling period without drawing air through it. Foils were stored frozen until analysis.

[15] The samples were analyzed by a thermographic method using a commercial instrument (C-mat 5500, Ströhlein) at the Institute for Tropospheric Research (IFT) in Leipzig, Germany [Neusüss et al., 2000]. The sample was placed in a quartz tube and heated rapidly to a specific temperature. To separate organic carbon (OC) and BC, the sample was first heated to 590°C under nitrogen. The carbon compounds that
evaporate under these conditions are referred to as OC. Then the sample was heated under oxygen to 650°C, and all carbon except carbonate was oxidized. The evaporated carbon is completely oxidized to CO₂ followed by analysis with an IR detector. External standards were used to calibrate the measurements. With all thermal carbon measurements, the OC/BC split is a method dependent property. The BC concentrations represent an upper limit as no correction for charring of organic matter was made.

The mass of particulate organic matter (POM) was determined by multiplying the measured organic carbon concentration in μg C m⁻³ by a POM factor which is an estimated average of the molecular weight per carbon weight for the organic aerosol. A POM factor of 1.6 was used for all samples based on a review of published measurements of the composition of organic aerosol in urban and nonurban regions [Turpin and Lim, 2001]. The POM factor was assigned an absolute uncertainty of 0.35.

The uncertainties associated with positive and negative sampling artifacts can be substantial [Turpin et al., 1994, 2000]. An effort was made to minimize positive artifacts by collecting samples on impaction plates. Negative artifacts may have occurred as a result of the pressure drop across the impactor (9 mbar for the 1.1 μm jet plate and 530 mbar for the 0.18 μm jet plate). No attempt was made to correct for artifacts or to determine their associated uncertainties since the information to do so was not available.

The uncertainties for BC at the 95% confidence level were based on a quadrature sum of the uncertainty in the air volume sampled and 2 times the standard deviation of the blank values measured over the course of the experiment. The uncertainties for POM were based on a quadrature sum of the uncertainty in the air volume sampled, the uncertainty in the OC to POM conversion factor, and 2 times the standard deviation of the blank over the course of the experiment.

2.4. Regional Concentrations of IOM and Residual Mass

Total elemental composition (Na, Mg, Al, Si, P, Cl, K, Ca, Ti, V, Cr, Mn, Fe, Ni, Cu, Zn, Ba, As, and Pb) was determined by thin-film X-ray primary and secondary emission spectrometry [Feely et al., 1991, 1998] at NOAA Pacific Marine Environmental Laboratory (PMEL). Submicron samples were collected on Nuclepore filters (0.4 μm pore size) mounted in a Berner impactor downstream of the D_{50,aero} 1.1 μm jet plate. Bulk samples were collected on Nuclepore filters (0.4 μm pore size) in a filter pack having an upper D_{90,aero} of 10 μm. Supermicron elemental concentrations were determined by difference between the submicron and bulk samples. This method of sample collection allows for the sharp size cut of the impactor while collecting a thin film of aerosol necessary for the X-ray analysis. Sampling periods ranged from 12 to 24 hours.

Filters were acid-washed before sample collection by soaking in 4 N HNO₃ for 24 hours and then 2% HCL for 24 hours with multiple rinses with distilled deionized water between treatments. Filters were weighed before and after sample collection as described below. Blank levels were determined by loading an impactor or filter pack with a filter but not drawing any air through it.

A component composed of inorganic oxidized material (IOM) was constructed from the elemental data. The IOM most likely was composed of soil dust and/or fly ash. Elemental concentrations alone are not sufficient to discriminate between the two. Regional crustal and fly ash enrichment factors for Al, Fe, and Ti relative to Si were calculated and found to be less than 2 for all source regions; values significantly greater than one are required to determine if the element has a noncrustal (i.e., fly ash) or nonfly ash (i.e., crustal) source. To construct the IOM component, the mass concentrations of Al, Si, Ca, Fe, and Ti, the major elements in soil and fly ash, were combined. It was assumed that each element was present in the aerosol in its most common oxide form (Al₂O₃, SiO₂, CaO, K₂O, FeO, Fe₂O₃, TiO₂) [Seinfeld, 1986]. The measured elemental mass concentration was multiplied by the appropriate molar correction factor as follows [Malm et al., 1994; Perry et al., 1997]:

\[
[IOM] = 2.2[Al] + 2.49[Si] + 1.63[Ca] + 2.42[Fe] + 1.94[Ti]
\]

This equation includes a 16% correction factor to account for the presence of oxides of other elements such as K, Na, Mn, Mg, and V that are not included in the linear combination. In addition, the equation omits K from biomass burning by using Fe as a surrogate for soil K and an average K/Fe ratio of 0.6 in soil [Cahill et al., 1986]. The concentration of residual mass was calculated from the gravimetric mass minus the sum of the chemically analyzed species (ionic, POM, BC, and IOM).

Uncertainties at the 95% confidence level associated with the IOM component were propagated as a quadratic sum of all errors involved including those due to the X-ray analysis, blank approximation, and volume of air sampled. XRF analysis errors are based on 2 times the standard deviation of 15 replicate analyses of a sample filter. Blank errors are based on 2 times the standard deviation of the average of all blanks collected over the course of the experiment. Uncertainties for the residual mass were propagated as a quadratic sum of the errors associated with the gravimetric, ionic, POM, BC, and IOM, and calculated water mass.

2.5. Regional Mass Fractions

Submicron and supermicron regional mass fractions were calculated from concentrations of the measured chemical components and the XRF Nuclepore filters that were weighed before and after sample collection. The filters were weighed at PMEL with a Cahn Model 29 microbalance housed in a glove box kept at a humidity of 33 ± 2%. The resulting mass concentrations from the gravimetric analysis include the water mass that is associated with the aerosol at 33% RH. Additional water mass may also be present due to interactions between the collected aerosol and the sampling substrate. The response of particles collected on a filter to changes in RH has been shown to be different than that of individual particles or bulk solutions of similar chemical composition [McInnes et al., 1996].

The glove box was continually purged with room air that had passed through a scrubber of activated charcoal, potassium carbonate, and citric acid to remove gas phase organics, acids, and ammonia. Static charging, which can result in balance instabilities, was minimized by coating the walls of the glove box with a static dissipative polymer (Tech Spray, Inc.), placing an antistatic mat on the glove box floor, using antistatic gloves while handling the substrates, and exposing the substrates to a ²¹⁰Po source to dissipate any charge that had built up on the substrates. Before and after sample collection, substrates were stored double-bagged with the outer bag containing citric acid to prevent absorption of gas phase ammonia. More details of the weighing procedure are given by Quinn and Coffman [1998].
Uncertainties of the mass fractions at the 95% confidence level were based on a quadratic sum of the uncertainties of the chemical concentrations and the gravimetrically determined mass. Uncertainty in the latter includes errors due to weighing, storage and transport, and the volume of air sampled [Quinn et al., 2000a]. To maintain the sampling RH of 55%, the sample air was heated, on average, 5.4°C (range of heating was 2° to 10°C). This heating may have led to the volatilization of a portion of the semivolatile organics and ammonium nitrate from the substrate thereby resulting in artificially low masses. Heating by 30°C has been found to result in a mass loss of about 34% for an aerosol dominated by wood smoke [Meyer et al., 1992; Ayers et al., 1999]. We expect the volatilization losses to be considerably less here since the aerosol was heated, on average, only 5.4°C (compared to 30°C); wood smoke, which contains significant quantities of semivolatile organics, was not the primary aerosol source; and ammonium nitrate contributed to less than 1% of the aerosol mass. It was not possible to quantify volatilization losses, however, since the composition of the semivolatile organics was unknown.

### 2.6. Aerosol Scattering, Backscattering, and Absorption Coefficients

Measurements of aerosol scattering and hemispheric backscattering coefficients were made with an integrating nephelometer (Model 3563, TSI Inc.) at wavelengths of 450, 550, and 700 nm at 55 ± 5% RH. The RH was measured inside the nephelometer sensing volume. Two single-stage impactors, one with a \( D_{90,\text{aero}} \) of 1.1 μm and one with a \( D_{90,\text{aero}} \) of 10 μm, were placed upstream of the nephelometer. An automated valve switched between the two impactors every 15 min so that sampling alternated between sub-1 μm and sub-10 μm aerosol. Values measured directly by the nephelometer were corrected for an offset determined by measuring filtered air over a period of several hours [Anderson and Ogren, 1998]. In addition, they were corrected for the angular nonidealities, including truncation errors and nonlambertian response, of the nephelometer using

\[
\sigma_{\text{sp}} = \sigma_{\text{sp, Meas}} \frac{\sigma_{\text{sp, True}}}{\sigma_{\text{sp, Neph, sim}}},
\]  

where \( \sigma_{\text{sp, True}} \) is the “true” scattering coefficient determined from the measured number size distribution and chemistry and a Mie scattering model and \( \sigma_{\text{sp, Neph, sim}} \) is the nephelometer simulated scattering coefficient determined from a Mie scattering model which employs a Mie integral modified to simulate the nephelometer response [Quinn and Coffman, 1998]. (The Mie calculations are discussed in more detail below.) This correction is similar to that of Anderson and Ogren [1998] but uses the simultaneously measured size distribution rather than an assumed size distribution. Values are reported at 0°C and 1013 mbar.

Sources of uncertainties associated with the use of the integrating nephelometer include photon counting during measurement, zeroing, and calibration; literature values of calibration gas scattering coefficients; variations in gas density within the nephelometer, and the angular correction applied in equation (3). These uncertainties were estimated using the method of Anderson et al. [1999]. Additional uncertainties include variations in measured scattering due to RH changes within the nephelometer sensing volume and inlet losses of large particles [Quinn and Coffman, 1998]. For a 30 min averaging time and a wavelength of 550 nm, a quadrature sum of errors yielded absolute uncertainties of 4.1 and 20 Mm\(^{-1}\) corresponding to low and high values of \( \sigma_{\text{sp}} \) equal to 24 and 110 Mm\(^{-1}\), respectively. Absolute uncertainties for \( \sigma_{\text{bop}} \) equal to 3.0 and 13 Mm\(^{-1}\) were 0.32 and 1.3 Mm\(^{-1}\), respectively.

The absorption coefficient for sub-1 μm aerosol, \( \sigma_{\text{ap}} \) was measured at 55 ± 5% RH by monitoring the change in transmission through a filter with a Particle Soot Absorption Photometer (PSAP, Radiance Research). Measured values were corrected for a scattering artifact, the deposit spot size, the PSAP flow rate, and the manufacturer’s calibration as per Bond et al. [1999]. Values are reported at 550 nm, 0°C, and 1013 mbar. Sources of uncertainty in the PSAP measurement include noise, drift, correction for the manufacturer’s calibration, and correction for the scattering artifact [Anderson et al., 1999]. A quadrature sum of these errors yields absolute uncertainties of 0.38 and 2.8 Mm\(^{-1}\) for \( \sigma_{\text{ap}} \) equal to 0.68 and 13 Mm\(^{-1}\), respectively, for a 30 min averaging time.

### 2.7. Number Size Distribution

Size distributions from 20 nm to 5 μm were measured with the combination of a differential mobility particle sizer (DMPS, University of Vienna (Reischle) medium column) and an aerodynamic particle sizer (APS) (TSI 3300) [Bates et al., 2002]. The DMPS was operated at 55 ± 5% RH. The APS was operated dry; diameters were shifted to 55% RH using the mass of water calculated to be associated with the aerosol at that RH (see section 3.1 for details of water calculation). Filtered mobility distributions from the DMPSs were converted to number size distributions. The APS diameters were converted to geometric diameters [Quinn and Coffman, 1998]. APS diameters were converted to geometric diameters by dividing by the square root of the particle density determined from the size-resolved chemical measurements. See section 3.1 for a description of the density calculations. An interactive routine was used to fit lognormal curves to the different modes of the number size distribution [Quinn and Coffman, 1998]. Surface area fit parameters for the total aerosol based on the number size distribution measured at 55 ± 5% RH are given in Table 2.

Uncertainties in the measured number size distribution (and mass concentrations derived from the number size distribution) result from instrumental errors of particle sizing and counting due to flow instabilities in the DMPS and APS.
The amount of observed drift in the shear and excess flows led to a ± 20% uncertainty in the number concentration [Bates et al., 2002]. Additional factors affecting the accuracy of the conversion of the number concentration to mass concentration include errors in the measured chemical composition and calculated density. Overall uncertainties in the submicron mass concentration derived from the number size distribution are ± 35% for a concentration of 3 µg m⁻³ [Quinn and Coffman, 1998]. Uncertainties for supermicron mass concentrations are ± 25% for a concentration of 20 µg m⁻³.

2.8. Aerosol Optical Depth

[33] A five-channel (380, 440, 500, 675, and 870 nm) handheld Microtops sunphotometer (Solar Light Co.) was used to derive aerosol optical depth τₐ at 500 nm for the SHAtl region. It was also used to derive Ångström exponents α at 440 and 675 nm for all of the source regions. The instrument has built-in pressure and temperature sensors and was operated with a Global Positioning System (GPS) connection to obtain position and time of the measurements. A Matlab routine used by the NASA SIMBIOS program and Brookhaven National Laboratory was used to convert the raw signal voltages to aerosol optical depths. Included in the conversion is a correction for Rayleigh scattering [Penndorf, 1957], ozone optical depth, and an air mass that accounts for the Earth's curvature [Kasten and Young, 1989]. Ozone column amounts used to calculate the ozone optical depth were based on TOMS level 3 data for the locations and time periods of the cruise. The instrument was calibrated using a Langley plot approach [Shaw, 1983] by the manufacturer prior to the cruise and again at Mauna Loa 5 months after the cruise. Calibration constants for the five wavelengths differed by less than 0.9% between the two calibrations, which corresponds approximately to 0.01 in optical depth.

[34] In addition, aerosol optical depth at 500 nm and Ångström exponents at 415 and 862 nm were derived from a fast-rotating shadowband radiometer (FFSR, Yankee Environmental Systems, Inc.). The FFSR, being an automated instrument, was able to provide more data points throughout the cruise. Unfortunately, FFSR measurements were not available during the SHAtl region. A Brookhaven National Laboratory marine version was used. In the marine version the electrical response time is decreased to about 1 ms [Reynolds et al., 2001] to accommodate for the motion of the ship. The instrument calibration coefficients for the motion of the ship. The instrument calibration coefficients for the motion of the ship. The instrument calibration coefficients for the motion of the ship.

2.9. Ancillary Parameters

[35] Also measured were meteorological parameters including surface temperature, RH, wind speed and direction, as well as vertical profiles of these parameters from radiosondes. Air mass back trajectories were calculated for three arrival altitudes (500, 2500, and 5500 m) for the ship's position at 6 hour intervals. Trajectories were calculated with the hybrid single-particle Lagrangian integrated model HY-SPLIT 4 based on the FNL global wind field [Draxler, 1992; http://www.noaa.gov/ready-bin/fnl.pl]

3. Model Calculations

3.1. Calculation of Aerosol Water Mass, Density, and Refractive Index

[36] The gravimetric analysis was performed at 33 ± 3% RH. Hence the measured mass on the filter substrates included the amount of water associated with the aerosol at that RH. The impactors, nephelometer, and PSAP sampled aerosol at 55 ± 5% RH. The chemical thermodynamic equilibrium model AeRho [Quinn et al., 1998; Quinn and Coffman, 1998] was used to estimate the water mass associated with the inorganic ions at 33 and 55% RH so that the gravimetric and chemically analyzed mass could be adjusted to the RH of the optical measurements.

[37] No information was available on the chemical composition or hygroscopicity of the organic mass. Without this information it is difficult to estimate the water mass associated with the organic matter or to estimate the error imposed by this assumption on calculated extinction. There is no consensus on the direction or magnitude of the contribution by organics to total aerosol water. Empirical evidence exists indicating that
organics enhance water absorption by aerosols at high RH under nonurban conditions and diminish water absorption under urban conditions [Saxena et al., 1995]. Dick et al., [2000] showed that organic-associated water content is considerably less than that of sulfate compounds, on a volume basis, for high RH but comparable or greater for low RH. Model estimates suggest that the contribution of secondary organic aerosols (SOA) to total aerosol water is less than 5% for cases when SOA make up 22% of the total particle mass less than 2.5 μm in diameter [Ansari and Pandis, 2000]. The suppression of light scattering at high RH (> 70%) has been attributed to the presence of organic aerosols [McInnes et al., 1998]. Given this state of knowledge and the lack of information on the chemical composition of the organic matter, no attempt was made to calculate its associated water mass or to estimate the error due to the assumption that organic-associated water mass was negligible. The trace element components estimated from the XRF analysis also were assumed to be hygroscopic.

[38] Using the aerosol chemical composition measured with seven-stage multijet cascade impactors [Berner et al., 1979] (D<sub>aero</sub> of 0.18, 0.31, 0.55, 1.1, 2.0, 4.1, and 10 μm), AeRho also was used to calculate the refractive index at 550 nm and density of the mix of all aerosol components (for the calculation of total extinction) and for the individual chemical components (for the calculation of extinction fractions and mass extinction efficiencies). Details of the AeRho calculations are given below.

[39] For the purpose of reconciling all the various in situ measurements, AeRho is a static model. It is designed to take the measured ionic composition of the aerosol and the constant sampling RH and to determine the molecular composition of the ionic chemical species within the aerosol. The molecular composition then is used to calculate the water mass associated with the aerosol and the aerosol refractive index and density. The model is not used to describe a dynamic system in which changes in the concentration of gas phase species affect the aerosol molecular composition. Therefore the model does not include interactions between the gas and aqueous phases. In addition, because of the constant sampling RH, it is not necessary to take into account changes in particle size with changes in RH.

[40] For the calculation of total extinction the aerosol was assumed to be an internal mixture containing all measured chemical components. The chemical reactions allowed to occur are shown in Table 3. The ionic molalities for each of the input species are determined initially by assuming that the activity of water is equal to the instrumental RH. Then, using the ZSR method [Zdanovskii, 1936; Robinson and Stokes, 1965], a further approximation of the water content of the aerosol is made. Aqueous phase concentrations are activity corrected using the method of Bromley [1973] which allows for the prediction of activity coefficients of strong electrolytes in multielectrolyte solutions based on binary solution activity coefficients [Pittilis and Seinfeld, 1987]. The pure-solution binary activity coefficients are calculated using the method of Pitts and Mayorga [1973]. The ionic species are partitioned between the solid and aqueous phases with the solids precipitating in the most thermodynamically favorable order. The crystallization RH used for each solid phase species is listed in Table 3. The remaining aqueous ionic species are converted to aqueous compounds in accordance with the thermodynamic equilibrium constants. Finally, thermodynamic equilibrium with respect to water is tested for, and the water activity is iterated until equilibrium is established.

[41] Polynomial fits based on data of Tang and Munkelwitz [1991, 1994] for metastable particles are used to estimate densities of individual inorganic soluble species. Data from Bray [1970] are used to estimate the density of H<sub>2</sub>SO<sub>4</sub>. The density of OC was assumed to be 1.4 g cm<sup>-3</sup> [Turpin and Lim, 2001] and that of BC 2 g cm<sup>-3</sup> [Seinfeld and Pandis, 1998]. For the purposes of the extinction calculations, the IOM and residual mass were combined into one component. The density of the component was set equal to that of a mixture of 45% montmorillonite (2.5 g cm<sup>-3</sup>) and 55% kaolinite (2.6 g cm<sup>-3</sup>) as these two minerals are common components of soil dust [Sokolik and Toon, 1999]. A volume-weighted average was taken of the density of the individual species to estimate the density of the aerosol mix in each impactor size bin. Average submicron and supermicron densities for each source region for the mix of aerosol chemical components are reported in Table 4.

[42] The method of partial molar refractions [Stelson, 1990] was used to calculate the real portion of the refractive index as a function of size at 550 nm. Values of the partial molar refractions of all chemical species except IOM were taken from [Stelson, 1990]. The complex refractive index at 550 nm was obtained by volume averaging the refractive index of the scattering and absorbing components. A refractive index of 1.5 was used for the IOM component based on the kaolinite-montmorillonite mixture at 550 nm [Sokolik and Toon, 1999].

### Table 3. Chemical Reactions Included in the Chemical Equilibrium Model AeRho

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Crystallization RH</th>
</tr>
</thead>
<tbody>
<tr>
<td>2H&lt;sup&gt;+&lt;/sup&gt;(aq) + SO&lt;sub&gt;4&lt;/sub&gt;&lt;sup&gt;-&lt;/sup&gt;(aq) ⇌ H&lt;sub&gt;2&lt;/sub&gt;SO&lt;sub&gt;4&lt;/sub&gt;(aq)</td>
<td>40°</td>
</tr>
<tr>
<td>H&lt;sup&gt;+&lt;/sup&gt;(aq) + HSO&lt;sub&gt;4&lt;/sub&gt;(aq) ⇌ H&lt;sub&gt;2&lt;/sub&gt;SO&lt;sub&gt;4&lt;/sub&gt;(aq)</td>
<td>40°</td>
</tr>
<tr>
<td>2NH&lt;sub&gt;4&lt;/sub&gt;(aq) + SO&lt;sub&gt;4&lt;/sub&gt;(aq) ⇌ (NH&lt;sub&gt;4&lt;/sub&gt;)&lt;sub&gt;2&lt;/sub&gt;SO&lt;sub&gt;4&lt;/sub&gt;(aq)</td>
<td>22°</td>
</tr>
<tr>
<td>NH&lt;sub&gt;4&lt;/sub&gt;(aq) + HSO&lt;sub&gt;4&lt;/sub&gt;(aq) ⇌ NH&lt;sub&gt;3&lt;/sub&gt;(aq) + H&lt;sub&gt;2&lt;/sub&gt;SO&lt;sub&gt;4&lt;/sub&gt;(aq)</td>
<td>22°</td>
</tr>
<tr>
<td>2Na&lt;sup&gt;+&lt;/sup&gt;(aq) + SO&lt;sub&gt;4&lt;/sub&gt;(aq) ⇌ Na&lt;sub&gt;2&lt;/sub&gt;SO&lt;sub&gt;4&lt;/sub&gt;(aq)</td>
<td>40°</td>
</tr>
<tr>
<td>Na&lt;sup&gt;+&lt;/sup&gt;(aq) + HSO&lt;sub&gt;4&lt;/sub&gt;(aq) ⇌ Na&lt;sub&gt;2&lt;/sub&gt;SO&lt;sub&gt;4&lt;/sub&gt;(aq)</td>
<td>40°</td>
</tr>
<tr>
<td>2K&lt;sup&gt;+&lt;/sup&gt;(aq) + SO&lt;sub&gt;4&lt;/sub&gt;(aq) ⇌ K&lt;sub&gt;2&lt;/sub&gt;SO&lt;sub&gt;4&lt;/sub&gt;(aq)</td>
<td>40°</td>
</tr>
<tr>
<td>K&lt;sup&gt;+&lt;/sup&gt;(aq) + HSO&lt;sub&gt;4&lt;/sub&gt;(aq) ⇌ KHSO&lt;sub&gt;4&lt;/sub&gt;(aq)</td>
<td>40°</td>
</tr>
<tr>
<td>H&lt;sup&gt;+&lt;/sup&gt;(aq) + NO&lt;sub&gt;3&lt;/sub&gt;(aq) ⇌ HNO&lt;sub&gt;3&lt;/sub&gt;(aq)</td>
<td>40°</td>
</tr>
<tr>
<td>NH&lt;sub&gt;4&lt;/sub&gt;(aq) + NO&lt;sub&gt;3&lt;/sub&gt;(aq) ⇌ NH&lt;sub&gt;4&lt;/sub&gt;NO&lt;sub&gt;3&lt;/sub&gt;(aq)</td>
<td>40°</td>
</tr>
<tr>
<td>Na&lt;sup&gt;+&lt;/sup&gt;(aq) + NO&lt;sub&gt;3&lt;/sub&gt;(aq) ⇌ NaNO&lt;sub&gt;3&lt;/sub&gt;(aq)</td>
<td>40°</td>
</tr>
<tr>
<td>K&lt;sup&gt;+&lt;/sup&gt;(aq) + NO&lt;sub&gt;3&lt;/sub&gt;(aq) ⇌ KNO&lt;sub&gt;3&lt;/sub&gt;(aq)</td>
<td>40°</td>
</tr>
<tr>
<td>Na&lt;sup&gt;+&lt;/sup&gt;(aq) + Cl&lt;sup&gt;-&lt;/sup&gt;(aq) ⇌ NaCl(aq)</td>
<td>45°</td>
</tr>
<tr>
<td>K&lt;sup&gt;+&lt;/sup&gt;(aq) + Cl&lt;sup&gt;-&lt;/sup&gt;(aq) ⇌ KCl(aq)</td>
<td>45°</td>
</tr>
</tbody>
</table>

*Tang and Munkelwitz [1994].
*Tang [1996].
*No data, assumed from KC1.
*Cohen et al. [1987].
*Tang et al. [1997].
each source region for the mix of aerosol chemical components are reported in Table 4.

For the calculation of the extinction due to each chemical component and mass extinction efficiencies, the method described above was followed but only individual components were considered. For components containing more than one chemical species (e.g., HSO₄⁻/NH₄HSO₄/(NH₄)₂SO₄ in the nss sulfate aerosol), a volume-weighted average was taken of the density of the individual species to estimate the density of the component in each impactor size bin. Similarly, the method of partial molar refractions was used to calculate the real portion of the refractive index as a function of size for the water soluble chemical components containing more than one chemical species.

3.2. Calculation of Extinction Due to Each Chemical Component and Mass Extinction Efficiencies

The chemical components considered in this analysis are sea salt, nss sulfate, IOM, POM, and BC. The sea-salt component includes all measured NO₃⁻ in the supermicron size range on the assumption that gas phase HNO₃ resulting from combustion processes reacts with sea salt to form NaNO₃ [Clegg and Brimblecombe, 1985]. Nss sulfate aerosol includes nss SO₄²⁻ and all measured NH₄⁺ up to an NH₄⁺ to nss SO₄²⁻ molar ratio of 2. The sea-salt and nss sulfate components also include the water calculated to be associated with these components at 55% RH. A combustion component composed of BC and submicron soluble nss K⁺, nss SO₄²⁻ and NO₃⁻ was constructed whose refractive index and density were determined by mass averaging those of the individual species. The submicron nss SO₄²⁻ added to the combustion component was found by the equilibrium calculation to be in excess of an NH₄⁺ to nss SO₄²⁻ molar ratio of 2. NO₃⁻ and nss SO₄²⁻, assumed here to be in the form of KNO₃ and K₂SO₄, have been measured in biomass burning plumes by Liu et al. [2000].

Size distributions of sea salt, nss sulfate aerosol, POM, and the combustion component were determined from the seven-stage impactor measurements coupled with the number size distributions (see below). Since only submicron and supermicron samples were collected for the IOM component, it was distributed as sea salt. As discussed below in section 4.5, mass extinction efficiencies derived with the Mie calculational method fall within the range of those derived from an independent empirical multiple linear regression method. In addition, the values are within the range of those previously reported, indicating that the assumed IOM size distribution was reasonable.

Using the output from AeRho, a volume ratio (component volume/total aerosol volume) was calculated for each component within each impactor size bin from the component mass concentration and density (both determined from AeRho) in the size bin. Component surface area ratios were then derived from the volume ratios. Extinction coefficients (σep, Mm⁻¹) at 550 nm were calculated for each component using the total aerosol surface area fit parameters (from the measured number size distributions) and the component surface area ratios. Surface area fit parameters (as opposed to number or volume fit parameters) were found to give the most accurate representation of the aerosol when attempting to fit both the accumulation and coarse modes. Surface area fit parameters for the total aerosol based on the number size distribution measured at 55 ± 5% RH are given in Table 2. This approach uses the measured chemical information but maintains the higher size resolution of the measured number size distribution. Having acquired size distributions of σep for each component, values of submicron and supermicron component σep were determined by integrating over the appropriate size range. Mass extinction efficiencies (m² g⁻¹) at 550 nm were calculated from the component σep (Mm⁻¹) and mass concentrations (µg m⁻³) for the submicron, supermicron, and sub-10 µm size ranges.

4. Results


Mean regional values of three measures of mass were compared to assess internal consistency in the impactor and number size distribution data used in the extinction calculations. Submicron and supermicron aerosol mass concentrations were determined gravimetrically, by summing the mass of the chemically analyzed species, and from the number size distribution using the density of the total aerosol mixture estimated with AeRho. The amount of water calculated to be associated with the aerosol at 55% RH was added to the
As shown in Figure 2, submicron mass concentrations from the three methods agreed for all source regions within the overall experimental uncertainty. Overall uncertainty was calculated from a quadrature sum of the uncertainties of the three methods \[\text{Quinn and Coffman, 1998}\]. This level of agreement gives confidence (for the submicron size range) to the density that was used to convert number to mass and the calculated water mass. In addition, it indicates that the chemical analyses accounted for all of the species that were present in the aerosol.

Supermicron mass concentrations from the three methods agreed for source regions where the sampled air had been over the ocean and not over land for at least 6 days prior to reaching the ship (SHAtl, SHIO, NHIO, and AS/CI). For source regions where the sampled air had been over land in the 6 days prior to reaching the ship (EIndSub, IndSub, Arabia/IndSub, and Arabia), mass derived from the number size distribution was low relative to that determined gravimetrically and through chemical analyses. One factor responsible for the disagreement could be the calculated water values used to adjust the gravimetric and chemically analyzed mass to 55% RH. For the supermicron size range the calculated water is due primarily to sea salt aerosol. If the interaction of sea salt with other chemical components decreases the water uptake and/or retention by sea salt, calculated water values will be too high. These four source regions had relatively high concentrations of non-sea-salt supermicron mass.

Extinction for submicron and supermicron aerosol was derived from the measured scattering and absorption coefficients. In addition, extinction was estimated using the measured chemical composition and number size distribution as input to Mie calculations. Measured and calculated submicron extinction agreed within the overall experimental uncertainty for all source regions (Figure 3). The most direct comparison of the nephelometer-measured and nephelometer-simulated scattering coefficients was desired. Hence the comparison used the scattering coefficients measured directly by the nephelometer without an angular correction and the nephelometer-simulated scattering coefficients.) Agreement was obtained for the supermicron size range in all source regions except EIndSub despite the lack of supermicron mass closure in the EIndSub, IndSub, Arabia/IndSub, and Arabia regions. The amount of calculated water significantly affected the gravimetric and chemically analyzed mass (the added water made up 24 to 36% of the supermicron mass for the EIndSub, IndSub, Arabia/
Table 5. Concentrations, Standard Deviations (± 1σ), and Absolute Uncertainties (95% Confidence Level, Shown in Parentheses) of the Mean of Aerosol Ionic and Carbonaceous Submicron \( (D_{\text{aero}} < 1.1 \, \mu \text{m}) \) at 55% RH Chemical Species for the Different Air Mass Regions

<table>
<thead>
<tr>
<th>Air Mass Region</th>
<th>Concentration, mg m(^{-3})</th>
<th>NH(_4^+)</th>
<th>Sea Salt</th>
<th>NO(_3^-)</th>
<th>POM(^b)</th>
<th>BC(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SH Atlantic</td>
<td>0.31 ± 0.16 (0.01)</td>
<td>0.04 ± 0.03 (0.003)</td>
<td>0.17 ± 0.10 (0.05)</td>
<td>0.003 ± 0.008 (0.001)</td>
<td>0.16 ± 0.02 (0.04)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>SH Indian Ocean</td>
<td>0.52 ± 0.18 (0.01)</td>
<td>0.09 ± 0.04 (0.003)</td>
<td>0.07 ± 0.08 (0.02)</td>
<td>&lt;0.001</td>
<td>0.07 ± 0.08 (0.04)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>NH Indian Ocean</td>
<td>1.8 ± 0.24 (0.03)</td>
<td>0.5 ± 0.12 (0.009)</td>
<td>0.09 ± 0.03 (0.03)</td>
<td>0.007 ± 0.007 (0.003)</td>
<td>0.13 ± 0.08 (0.06)</td>
<td>0.14 ± 0.05 (0.02)</td>
</tr>
<tr>
<td>East Indian subcontinent</td>
<td>13 ± 1.9 (0.32)</td>
<td>1.5 ± 0.06 (0.04)</td>
<td>0.10 ± 0.01 (0.05)</td>
<td>0.03 ± 0.02 (0.006)</td>
<td>1.4 ± 0.08 (0.19)</td>
<td>1.8 ± 0.23 (0.06)</td>
</tr>
<tr>
<td>Indian subcontinent</td>
<td>7.5 ± 2.2 (0.19)</td>
<td>2.1 ± 0.71 (0.05)</td>
<td>0.08 ± 0.02 (0.04)</td>
<td>0.02 ± 0.01 (0.006)</td>
<td>1.2 ± 0.13 (0.30)</td>
<td>1.4 ± 0.06 (0.08)</td>
</tr>
<tr>
<td>Arabia</td>
<td>1.5 ± 0.21 (0.04)</td>
<td>0.37 ± 0.04 (0.009)</td>
<td>0.13 ± 0.07 (0.04)</td>
<td>0.02 ± 0.01 (0.005)</td>
<td>0.78 ± (0.20)</td>
<td>0.08 ± 0.05 (0.05)</td>
</tr>
<tr>
<td>Arabia-Indian subcontinent</td>
<td>3.9 ± 0.93 (0.05)</td>
<td>0.96 ± 0.25 (0.01)</td>
<td>0.09 ± 0.05 (0.02)</td>
<td>0.02 ± 0.008 (0.003)</td>
<td>0.94 ± 0.88 (0.10)</td>
<td>0.54 ± 0.36 (0.02)</td>
</tr>
<tr>
<td>Arabian Sea-coastal India</td>
<td>3.7 ± 1.3 (0.06)</td>
<td>0.90 ± 0.34 (0.21)</td>
<td>0.05 ± 0.03 (0.03)</td>
<td>0.02 ± 0.007 (0.004)</td>
<td>0.42 ± 0.20 (0.09)</td>
<td>0.38 ± 0.12 (0.02)</td>
</tr>
</tbody>
</table>

\(^{a}\)Here \(\mu g \, \text{C m}^{-3}\) of OC \(\times 1.6\).

\(^{b}\)Here \(\mu g \, \text{C m}^{-3}\).

IndSub, and Arabia regions). Water does not affect as directly the calculated aerosol refractive index and density that are required for the extinction calculations.

[51] As an additional check on the extinction calculations, measured and calculated single scattering albedos were compared for the submicron size range. The comparison was not done for the supermicron size range as absorption values were below the detection limit of 0.34 Mm\(^{-3}\). As shown in Figure 3, agreement was within the overall experimental uncertainty for all source regions with detectable absorption.

[52] Agreement in extinction for submicron and supermicron size range indicates that the calculated extinction is accurate and lends confidence to the calculated extinction due to individual chemical components. Supermicron extinction fractions for those source regions where mass or extinction closure was not obtained, should be considered less certain, however.

4.2. Regional Chemical Composition and Mass Fractions

4.2.1. Sulfate Aerosol

[53] Table 5 lists the mean concentrations, standard deviations, and absolute uncertainties of the major submicron ionic chemical species for the different air mass source regions. Mean submicron nss SO\(_4^{2-}\) concentrations were lowest in the marine regions south of the ITCZ. SHAtl and SHIO nss SO\(_4^{2-}\) concentrations averaged 0.31 ± 0.16 and 0.52 ± 0.18 \(\mu g \, \text{m}^{-3}\) (average and 1σ standard deviation), respectively. These concentrations are similar to the mean values observed for different latitude bands of the central Pacific (0.13 to 0.50 \(\mu g \, \text{m}^{-3}\) [Quinn et al., 2000a]) and the southern Atlantic (0.52 ± 0.14 \(\mu g \, \text{m}^{-3}\) [Quinn et al., 2001; Andreae et al., 1995]). The NHIO region includes time periods when the ship was north of the ITCZ, within the ITCZ, and south of the ITCZ but with trajectories coming from the northern hemisphere. (The ITCZ was located between 2°S and 1°N when the ship first crossed it between 67° and 70°E on days 58 and 59. It was located between 5°S and 8°S when the ship crossed it at 73°E on days 76 to 79.) Mean nss SO\(_4^{2-}\) concentrations for this region were about a factor of 6 higher (1.8 ± 0.24 \(\mu g \, \text{m}^{-3}\)) than for regions south of the ITCZ and ranged from 1.5 to 2.2 \(\mu g \, \text{m}^{-3}\). The highest concentrations of seawater dimethylsulfide (6.3 Nm\(^{-3}\)) and atmospheric dimethylsulfide (370 pptv) (from which biogenic nss SO\(_4^{2-}\) is derived) were measured in the ITCZ suggesting that the elevated nss SO\(_4^{2-}\) for the NHIO region was, in part, biogenic in origin.

[54] On the basis of trajectories arriving at the ship location at a height of 500 m, lower-level air from three of the nonmarine source regions did not pass over the Indian subcontinent prior to reaching the ship. The mean nss SO\(_4^{2-}\) concentration for the Arabia region was comparable to the NHIO region (1.5 ± 0.21 \(\mu g \, \text{m}^{-3}\)). Mean values were higher for the Arabia/IndSub (3.9 ± 0.93 \(\mu g \, \text{m}^{-3}\)) and the AS/CI (3.7 ± 1.3 \(\mu g \, \text{m}^{-3}\)) regions. The Arabia/IndSub region experienced low-level (500 m) flow from Arabia and upper-level (2500 m) flow from the Indian subcontinent. Micropulse lidar measurements show the presence of lower and upper aerosol layers for this region with mixing between the two at times [Welton et al., 2002]. A portion of the nss sulfate aerosol measured at the surface may have resulted from mixing of the upper layer Indian subcontinent air mass into the boundary layer.

[55] Mean nss SO\(_4^{2-}\) concentrations were highest for the two source regions with lower-level flow from the Indian subcontinent. The IndSub region, during which air flowed from central India to the ship, had a mean nss SO\(_4^{2-}\) concentration of 7.5 ± 2.2 \(\mu g \, \text{m}^{-3}\) with a range of 4.8 to 9.7 \(\mu g \, \text{m}^{-3}\). The EIndSub region with air flow from the Calcutta region across southern Indian or Sri Lanka to the ship had a mean concentration of 13 ± 1.9 \(\mu g \, \text{m}^{-3}\) and a range of 11 to 16 \(\mu g \, \text{m}^{-3}\). The mean concentration measured in pollution plumes at altitudes less than 500 m off the east coast of the United States during TARFOX was 11 \(\mu g \, \text{m}^{-3}\) with a range of 1 to 19 \(\mu g \, \text{m}^{-3}\) [Hegg et al., 1997]. Regional mean NH\(_4^+\) to nss SO\(_4^{2-}\) molar ratios ranged from 0.63 to 1.5 indicating that, on average, the sulfate aerosol was not completely neutralized by ammonium.

[56] Mass fractions and associated uncertainties at the 95% confidence level of the major submicron chemical species are shown in Figure 4. Mass fractions were calculated from the measured component mass concentration and the total aerosol mass concentration determined by gravimetric analysis. Nss sulfate aerosol, which includes nss SO\(_4^{2-}\), NH\(_4^+\), and H\(_2\)O at 55% RH, made up the majority of the submicron mass in all source regions except the two most remote marine regions (SHAtl and SHIO). Even in these two regions, however, the sulfate mass fraction was substantial averaging 36 ± 12% for
the SHAtl and 49 ± 15% (mean and uncertainty at the 95% confidence level) for the SHIO.

Supermicron nss SO\textsubscript{4}\textsuperscript{2-} concentrations were highest in the EIndSub (0.27 ± 0.42 μg m\textsuperscript{-3}) and IndSub (0.19 ± 0.14 μg m\textsuperscript{-3}) regions (see Table 6). In all other source regions, concentrations were less than 0.06 μg m\textsuperscript{-3}. Supermicron mean NH\textsubscript{4}\textsuperscript{+} concentrations were less than 0.03 μg m\textsuperscript{-3} for all source regions. Supermicron mass fractions of nss sulfate aerosol ranged from undetectable to 2.2% (Figure 5).

4.2.2. Sea Salt and Nitrate

Mean submicron sea-salt concentrations ranged between 0.05 and 0.17 μg m\textsuperscript{-3} for all source regions (Table 5). These concentrations are low relative to those reported for latitude bins of the central Pacific (0.11 to 0.58 μg m\textsuperscript{-3} [Quinn et al., 2000a]), for different air mass types in the northeast Atlantic (0.34 to 0.88 μg m\textsuperscript{-3} [Quinn et al., 2000b]), and for the remote Southern Ocean (1.0 ± 0.55 μg m\textsuperscript{-3} [Quinn et al., 1998]). They are comparable to concentrations reported for several different latitude bands of the Atlantic (0.08 to 0.22 μg m\textsuperscript{-3} [Quinn et al., 2001]). The low concentrations may have been due to the relatively low wind speeds during INDOEX that averaged 5.6 ± 2.8 m s\textsuperscript{-1}. For comparison, wind speeds for the Southern Ocean during ACE 1 averaged 9.5 m s\textsuperscript{-1} and for the northeast Atlantic during ACE 2 averaged 8.2 m s\textsuperscript{-1} [Quinn et al., 2000b].

Mass fractions of submicron sea-salt aerosol (including sea salt and the water associated with it at 55% RH) were high for the two remote marine source regions (49 ± 27% for the SHAtl and 23 ± 8.8% for the SHIO). They were less than 2.5% for all other source regions. The low mass fractions are in contrast to what has been observed for remote regions of the central Pacific (7 to 53% [Quinn et al., 2000a]) and marine and continentally impacted regions of the Atlantic (9 to 49% [Quinn et al., 2001]).

On a regional basis, mean supermicron sea-salt concentrations were either lower than or typical of those reported
for other ocean regions. Mean concentrations ranged from 1.4 ± 0.53 µg m⁻³ for the EIndSub region to 10 ± 3.7 µg m⁻³ for the SHAtl region (Table 6). Mean concentrations for different air mass types over the northeast Atlantic during ACE 2 ranged from 4 to 10 µg m⁻³ [Quinn et al., 2000b] and for the central Atlantic during Aerosols99 ranged from 4 to 19 µg m⁻³ [Quinn et al., 2001]. The experimental average during ACE 1 conducted in the Southern Ocean region was 9.4 ± 5.5 µg m⁻³ [Quinn et al., 1998].

[61] In the supermicron size range, the measured NO₃⁻ was assumed to be associated with sea salt through the reaction of combustion-derived HNO₃ and sea salt [Clegg and Brimblecombe, 1985]. NO₃⁻ concentrations were low relative to sea salt in the SHAtl, SHIO, and NHIO regions. They were high in the continentally impacted regions being, on average, twice the sea-salt concentration in the EIndSub region, comparable to sea salt in the IndSub region, and 17 to 44% of the sea-salt concentration in the Arabia, Arabia/IndSub, and AS/CI regions.

[62] Sea-salt aerosol dominated the supermicron mass in all source regions (Figure 5). Mean mass fractions ranged from 55 ± 4.5% in the IndSub region to 97 ± 4.3% in the SHAtl region. Sea-salt aerosol made up the majority of the sub-10 µm aerosol mass in the SHAtl (95%), SHIO (91%), NHIO (72%), and Arabia (58%) regions. It made up 23% of the sub-10 µm mass in the EIndSub region, 27% in the IndSub region, 48% in the Arabia/IndSub region, and 34% in the AS/CI region.  

4.2.3. IOM

[63] As explained in section 2.4, an inorganic oxidized material (IOM) component was constructed from the concentration of the major elements in soil and fly ash with a correction to the mass assuming that the elements were present in their normal oxidized form. This component most likely consists of fly ash with an intermittent soil dust contribution. Spectral absorption measurements of bulk aerosol collected on the Ronald H. Brown showed elevated absorption at the UV-blue wavelengths, an indicator of soil dust, only in the Arabia/IndSub and Arabia regions (D. Savoie, personal communication, 1999). During these periods, low-level trajectories were coming from Arabia to the ship.

[64] Submicron concentrations of IOM were less than 0.07 µg m⁻³ in the SHAtl, SHIO, and NHIO regions (Table 7). They were higher in the more continentally influenced source regions ranging from 0.42 µg m⁻³ in the AS/CI region to 1.7 µg m⁻³ in the IndSub region. The mean submicron mass fraction of IOM was highest in the Arabia region (21 ± 3.1%). In all other source regions the submicron mass fraction was less than 8.8%.

[65] Mean supermicron concentrations of IOM were less than 0.12 µg m⁻³ in the SHAtl and SHIO regions (Table 7). They ranged between 3.4 (NHIO) and 8 µg m⁻³ (Arabia/IndSub) for the other regions. Mean supermicron mass fractions were less than 1% in the SHAtl and SHIO regions and 14 ± 1.4% in the NHIO (Figure 5). For the more continentally influenced source regions values ranged from 23% (EIndSub) to 34% (AS/CI).

4.2.4. POM

[66] POM and BC concentrations and mass fractions are presented here to aid in the interpretation of the extinction due to these components. A more detailed discussion of these components for the INDOEX cruise can be found in the work of Neuviess et al. [2002]. Mean submicron POM concentrations were less than or equal to 0.16 µg m⁻³ for the SHAtl, SHIO, and NHIO (Table 5). Concentrations in the continentally influenced air mass source regions ranged from 0.42 ± 0.20 µg m⁻³ for the AS/CI region to 1.4 ± 0.08 µg m⁻³ for the EIndSub region. Mean submicron mass fractions for the marine source regions ranged from 2.1 ± 1.1% for NHIO to 10 ± 2.6% for SHAtl (Figure 4). Values for the continentally influenced source regions were near 5% for EIndSub, IndSub, IndSub/Arabia, and AS/CI. Highest values corresponded to the Arabia region (15 ± 4.0%).

[67] Mean supermicron POM concentrations were similar to the submicron values ranging from 0.14 to 0.56 µg m⁻³ for the marine regions and 0.25 to 1.3 µg m⁻³ for the continentally influenced regions (Table 6). The size distribution of POM depends on the processes by which it is incorporated into the aerosol particles. Its existence in the supermicron size range suggests that gas phase organics were adsorbed onto relatively large particles or it had a source in the ocean surface layer and was injected into the atmosphere in a manner similar to sea salt. Mean supermicron POM mass fractions were near 1 to 2% for the marine regions and <1 to 9% for the continentally influenced regions.
4.2.5. BC and nss K$^+$

Submicron BC concentrations were below the detection limit of 0.01 $\mu$g m$^{-3}$ in the SHAtl and SHIO regions (Table 5). They were higher in the NHIO region (mean of 0.14 ± 0.05 $\mu$g m$^{-3}$ with a range from 0.08 to 0.23 $\mu$g m$^{-3}$) indicating transport of anthropogenic aerosol from continental regions to the ITCZ region. Mean concentrations were relatively low in regions where low level air came from Arabia (0.08 ± 0.05 $\mu$g m$^{-3}$ for the Arabia region and 0.54 ± 0.36 $\mu$g m$^{-3}$ for the Arabia/IndSub region) or low-level air had not been in contact with land 6 days prior to being sampled on the ship (0.38 ± 0.12 $\mu$g m$^{-3}$ for the AS/CI region). Mean concentrations were highest for regions where low level air came from the Indian subcontinent to the ship (1.8 ± 0.23 $\mu$g m$^{-3}$ for the EIndSub region and 1.4 ± 0.06 $\mu$g m$^{-3}$ for the IndSub region). These values exceed the range measured over the northeast Atlantic during ACE 2 (0.1 to 1.1 $\mu$g m$^{-3}$ [Novakov et al., 2000]) and the range inferred for the western Atlantic during TARFOX (up to 0.9 $\mu$g m$^{-3}$ [Novakov et al., 1997]).

The submicron BC to TC mean ratio was smallest for the Arabia region (0.12 ± 0.13) and between 0.5 and 0.7 for all other source regions. Hence a relatively wide range of values were measured on the ship due to the expanse of the Indian Ocean and the variety of the air mass source regions impacting it. Mean ratios measured on three C130 flights at altitudes lower than 2 km ranged from 0.43 to 0.51 [Novakov and Ramanathan, 2001]. Submicron mean mass fractions of BC were below the detection limit for the SHAtl and SHIO regions (Figure 4), 2.7 ± 0.4% for the NHIO region, and between 1.5 (Arabia) and 7.5% (EIndSub) for the continentally influenced regions.

Aerosol nss soluble K$^+$ is a by product of biomass burning and has been found to correlate well with BC concentrations [Cachier et al., 1995]. As for BC, mean submicron nss

**Figure 5.** Regional mean supermicron mass fractions of the dominant chemical components at 55% RH. Error bars indicate the uncertainty at the 95% confidence level. The sea-salt component mass includes sea salt, nitrate, and water. The nss sulfate component mass includes nss SO$_4^{2-}$, NH$_4^+$, and water. The number of samples available for the mass fraction calculation in each region was as follows: SHAtl 7, SHIO 5, EIndSub 1, IndSub 2, Arabia/IndSub 4, Arabia 2, NHIO 3, AS/CI 3.
Table 7. Concentrations, Standard Deviations (±1σ), and Absolute Uncertainties (95% Confidence Level, Shown in Parentheses) of the Mean of Aerosol Inorganic Oxidized Matter (IOM) and Residual Mass Concentrations for the Submicron ($D_{aero} < 1.1 \mu m$ at 55% RH) and Supermicron (1.1 $< D_{aero} < 10 \mu m$ at 55% RH) Aerosol for the Different Air Mass Regimes

<table>
<thead>
<tr>
<th>Air Mass Regime</th>
<th>Submicron, $\mu g m^{-3}$</th>
<th>Supermicron, $\mu g m^{-3}$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>IOM$^{a}$</td>
<td>Residual$^{b}$</td>
</tr>
<tr>
<td>SH Atlantic</td>
<td>0.06 ± 0.05 (0.02)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>SH Indian Ocean</td>
<td>0.07 ± 0.10 (0.02)</td>
<td>0.19 ± 0.14 (0.08)</td>
</tr>
<tr>
<td>NH Indian Ocean</td>
<td>0.07 ± 0.06 (0.04)</td>
<td>0.35 ± 0.49 (0.15)</td>
</tr>
<tr>
<td>East Indian subcontinent</td>
<td>1.6 (0.23)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Indian subcontinent</td>
<td>1.7 (0.18)</td>
<td>0.57 (0.78)</td>
</tr>
<tr>
<td>Arabia</td>
<td>1.1 (0.14)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Arabia-Indian</td>
<td>0.70 ± 0.50 (0.06)</td>
<td>0.61 ± 0.58 (0.25)</td>
</tr>
<tr>
<td>Arabian Sea-coastal India</td>
<td>0.42 ± 0.34 (0.08)</td>
<td>0.93 ± 0.65 (0.28)</td>
</tr>
</tbody>
</table>

$^{a}$Based on [IOM] = 2.2[Al] + 2.49[Si] + 1.63[Ca] + 2.42[Fe] + 1.94[Ti] from Malm et al. [1994].

$^{b}$Based on gravimetric mass less the sum of the chemically analyzed mass.

K$^+$ concentrations were highest in the EIndSub (0.58 ± 0.05 $\mu g m^{-3}$) and IndSub (0.44 ± 0.14 $\mu g m^{-3}$) regions. Submicron mean mass fractions were 2.2% or less in all source regions. Mean regional ratios of nss K$^+$ to BC ranged from 0.30 (IndSub and EIndSub) to 0.52 (AS/CI) and 0.61 (Arab/IndSub) in the continental influenced regions. The mean ratio for the NHIO region was 0.66 ± 0.18. For comparison, a mean ratio of 0.52 ± 0.11 has been reported for biomass burning aerosol in Brazil [Ferek et al., 1998]. The range of values measured on the ship indicates a variable yet significant contribution of biomass burning aerosol to the Indian Ocean.

[71] Mean supermicron BC mass fractions were less than 2% in all source regions. Similarly, nss K$^+$ mean mass fractions were less than 0.5% in all source regions.

4.2.6. Residual

The concentration of residual mass was calculated from the gravimetric mass minus the sum of the chemically analyzed species (ionic, POM, BC, and IOM). Mean submicron and supermicron concentrations are reported in Table 7. Submicron mean mass fractions ranged from less than the detection limit to 15%. Supermicron mean mass fractions ranged from less than the detection limit to 0.8%. The residual mass is most likely a result of inaccuracies in the estimate of the water associated with the aerosol at 55% RH, the selection of a constant POM factor of 1.6 for the entire experiment, and the assumption of the composition of the IOM component.

4.2.7. Other Chemical Components

Other chemical components that were detected but that contributed less than one percent to the submicron mass were NO$_3^-$ (presumably associated with NH$_4^+$) and MSA$^-$. MSA$^-$ also was detected in the supermicron aerosol but made up <1% of the supermicron mass in all source regions.

4.3. Regional Optical Properties

4.3.1. Sub-1 $\mu m$ to sub-10 $\mu m$ Extinction Ratio

Sub-1 $\mu m$ and sub-10 $\mu m$ scattering and absorption coefficients were measured as described in section 2.6. Extinction coefficients at 550 nm and 55% RH were derived from the sum of the STP- and angular-corrected scattering coefficients and the absorption coefficients. Mean values and percentile information for the ratio of sub-1 $\mu m$ to sub-10 $\mu m$ extinction coefficients for the different air mass source regions are shown in Figure 6. Mean ratios were lowest in the marine regions south of the ITCZ (28 ± 7.4% for SHAtl, 40 ± 10% for SHIO, mean and 1σ standard deviation). The aerosol in these source regions was dominated by supermicron sea salt resulting in large extinction in the supermicron size range. The mean ratio was higher for the NHIO (60 ± 2.7%); supermicron mass concentrations were similar to those in the other marine regions but submicron nss sulfate aerosol concentrations also were higher. The mean ratio also was relatively low for the Arabia region (53 ± 4.3%) due to large concentrations of supermicron aerosol. On the basis of chemical analyses and spectral absorbances of filters collected in this region (D. Savoie, personal communication, 1999), the supermicron aerosol was composed of sea salt and soil dust.

[75] Mean sub-1 $\mu m$ to sub-10 $\mu m$ extinction ratios were comparable for the Arabia/IndSub (74 ± 7.7%) and AS/CI

Figure 6. Mean regional ratios of sub-1 $\mu m$ to sub-10 $\mu m$ extinction at 550 nm and 55% RH. Extinction values are based on the sum of measured STP- and angular-corrected $\sigma_{eq}$ from the nephelometer and measured STP-corrected $\sigma_{eq}$ from the PSAP.
resulting in sub-10 μm mass concentrations composed of nss sulfate aerosol, POM, and BC that those from the Indian subcontinent did. In addition, the accumulation mode surface mean diameter of the Arabia aerosol was smaller and the coarse mode surface mean diameter was larger relative to the Indian subcontinent aerosol (Table 3). Both factors result in a lower sub-10 μm extinction.

[77] Lower-level (500 m) trajectories for the AS/CI region indicated that the sampled air masses had not been over land for at least 6 days prior to reaching the ship. This region also had a relatively low mean sub-10 μm extinction of 48 ± 23 Mm⁻¹ relative to the IndSub and EIndSub regions. Mean sub-10 μm extinction for the Arabia/IndSub region was higher (60 ± 11 Mm⁻¹) due to a larger submicron mass concentration perhaps resulting from mixing of Indian subcontinent aerosol into the marine boundary layer. Highest mean extinction coefficients corresponded to the two regions with lower-level flow from the Indian subcontinent, EIndSub (130 ± 5.8 Mm⁻¹) and IndSub (89 ± 22 Mm⁻¹). These are higher than mean values for western European air measured during ACE 2 (80 ± 21 Mm⁻¹) [Quinn et al., 2000b] and within the range measured at altitudes lower than 300 m during TARFOX (24 to 200 Mm⁻¹) [Hegg et al., 1997].

[78] Sub-10 μm mean backscatter coefficients showed the same trend as the extinction coefficients being highest for the EIndSub and IndSub regions and lowest for the marine and Arabia regions. Resulting mean sub-10 μm backscattered fractions were relatively constant spanning the limited range from 0.091 ± 0.004 to 0.11 ± 0.004. Sub-1 μm mean values ranged from 0.089 ± 0.008 to 0.11 ± 0.02.

### 4.3.3. Aerosol Angström Exponents

[79] Mean regional Angström exponents α for the 450 and 700 nm nephelometer wavelength pair derived from

$$\alpha = -\frac{\log(\sigma \lambda_2/\sigma \lambda_1)}{\log \lambda_2/\lambda_1}$$

are shown for the sub-10 μm aerosol in Figure 8a. Mean values were lowest for the marine regions, particularly those in the southern hemisphere where supermicron sea salt dominated the aerosol mass. Values were near 1.0 for the regions with low-level flow from Arabia and near 1.5 for regions with low-level flow from the Indian subcontinent.

### 4.3.4. Aerosol Absorption Coefficient and Single Scattering Albedo

[80] Mean sub-1 μm σap were highest for the EIndSub (18 ± 1.3 Mm⁻¹) and IndSub (12 ± 4.4 Mm⁻¹) regions resulting from the relatively large concentrations of BC. For the majority of the sampling periods, absorption coefficients for the southern hemisphere marine regions, SHAtl and SHIO, were less than the detection limit of 0.34 Mm⁻¹. Mean values for the NHIO and Arabia regions were 2.4 ± 0.35 and 1.0 ± 0.20 Mm⁻¹, respectively. Mean values for the AS/CI and Arabia/IndSub regions were 5.0 ± 2.3 and 4.9 ± 1.8 Mm⁻¹, respectively.

[81] Single scattering albedo ωs calculated as

$$\omega_s = \frac{\sigma_{ap}}{\sigma_{ap} + \sigma_{sp}}$$

is a measure of the relative magnitude of scattering and absorption by the aerosol. Here σsp is the nephelometer-measured scattering coefficient corrected for angular nonide-
alities as per equation (3), and $\sigma_{ap}$ is the measured absorption coefficient corrected as per Bond et al. [1999]. Mean regional values of $\omega_a$ (550 nm) for sub-10 $\mu$m aerosol and 55% RH, and sub-1 $\mu$m aerosol at 55% and 75% RH are listed in Table 8. Three levels of regional mean $\omega_a$ values are apparent. For sub-1 $\mu$m aerosol at 55% RH, mean values near one were observed in the northern hemisphere marine regions. Lower values were observed in the southern hemisphere marine regions. Lower mean values were observed in the AS/CI region (0.89 $\pm$ 0.01) and in the regions with low-level flow from Arabia (0.93 $\pm$ 0.02 for Arabia and 0.89 $\pm$ 0.02 for Arabia/IndSub). Lowest mean values correspond to the regions exposed to low-level flow from the Indian subcontinent (0.85 $\pm$ 0.01 for EIndSub and 0.84 $\pm$ 0.01 for IndSub). Comparable mean values were observed in the AS/CI region (0.86 $\pm$ 0.01).

[s2] Elevated absorption at the UV-blue wavelengths suggests that the aerosol collected in the regions with low-level flow from Arabia contained soil dust. In addition, BC concentrations in these regions were low compared to the regions with low-level flow from India. The mass fraction of IOM in the Arabia and Arabia/IndSub regions was equivalent to or larger than the BC mass fraction in the EIndSub and IndSub regions. The equivalent IOM and BC mass fractions but higher $\omega_a$ in the Arabia and Arabia/IndSub regions suggests that the soil dust component had a lower mass absorption efficiency at 550 nm than did BC in the EIndSub and IndSub regions.

4.4. Regional Extinction Due to Individual Aerosol Chemical Components

[s3] The fraction of the measured extinction (scattering plus absorption) due to the major aerosol chemical components was calculated at 55% RH and 550 nm using the method described in section 3.2. The major components considered are sea-salt aerosol which includes supermicron NO$_3^-$ and water calculated to be associated with sea salt at 55% RH; nss sulfate aerosol which includes nss SO$_4^{2-}$, NH$_4^+$, and water at 55% RH; a combustion component composed of BC, KNO$_3$, and K$_2$SO$_4$; POM; and IOM. For these calculations, the IOM and residual mass were combined into one component. Results are shown for the submicron, supermicron, and sub-10 $\mu$m aerosol in Figures 9, 10, and 11. In general, the trend in extinction fractions for each of the chemical components follows the trend in their mass fractions.

4.4.1. Non-Sea-Salt Sulfate Aerosol

[s4] The contribution of nss sulfate aerosol to submicron extinction ranged from 19 to 55% (Figure 9). This range is higher than that estimated for submicron nss sulfate extinction over the Atlantic Ocean during the Aerosols99 cruise (up to 25%) [Quinn et al., 2000a]. Smallest contributions were in the southern hemisphere marine regions (SHAtl = 19 $\pm$ 6.3% and SHIO = 29 $\pm$ 14%, mean and 1$\sigma$ standard deviation) due to relatively high extinction by submicron sea-salt aerosol.

Table 8. Single Scattering Albedo $\omega_a$ at 550 nm for Sub-10 and Sub-1 $\mu$m Aerosol at the Measurement Relative Humidity of 55% and at 75%*

<table>
<thead>
<tr>
<th>Air Mass Region</th>
<th>$\omega_a$, 550 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>55% RH, sub-10 $\mu$m</td>
</tr>
<tr>
<td>SH Atlantic*</td>
<td>1.0 $\pm$ 0.02</td>
</tr>
<tr>
<td>SH Indian Ocean*</td>
<td>1.0 $\pm$ 0.02</td>
</tr>
<tr>
<td>NH Indian Ocean</td>
<td>0.95 $\pm$ 0.02</td>
</tr>
<tr>
<td>East Indian subcontinent</td>
<td>0.86 $\pm$ 0.01</td>
</tr>
<tr>
<td>Indian subcontinent</td>
<td>0.86 $\pm$ 0.02</td>
</tr>
<tr>
<td>Arabia</td>
<td>0.96 $\pm$ 0.01</td>
</tr>
<tr>
<td>Arabia/Indian subcontinent</td>
<td>0.92 $\pm$ 0.02</td>
</tr>
<tr>
<td>Arabian-Sea/coastal India</td>
<td>0.88 $\pm$ 0.02</td>
</tr>
</tbody>
</table>

*Values are reported at STP.

*Here $\sigma_{ap}$ was from the measurement RH of 55% to 75% using measured $f$(RH) curves at Kaashidoo Climate Observatory during the INDOEX 1999 intensive [Ramanathan et al., 2001].

*Upper bound; $\sigma_{ap}$ values less than the detection limit of 0.34 Mm$^{-1}$ were considered to be zero.

Figure 8. (a) Box plot of the Angström Exponent (450 and 700 nm wavelength pair) derived from the STP- and angular-corrected $\sigma_{ap}$ for sub-10 $\mu$m aerosol and 55% RH. Percentile information is as described in Figure 7 caption. (b) Mean and standard deviation (1$\sigma$) of the Angström Exponent (415 and 862 nm wavelength pair) derived from the measured column aerosol optical depth for the different air mass regions.
sulfate extinction in the northern hemisphere marine region (NHIO) was higher (40 ± 18%) due to lower submicron sea-salt concentrations and higher nss sulfate concentrations. The extinction fraction of sulfate aerosol was relatively low in the Arabia region (31 ± 8.2%) due to significant extinction by several components (sea salt, sulfate, IOM, and POM). The other region impacted by lower-level flow from Arabia, Arabia/IndSub, had a higher mean submicron extinction fraction of 55 ± 19% due to higher concentrations of sulfate and lower concentrations of POM and IOM. Submicron sulfate extinction fractions for the IndSub and EIndSub regions were 40 ± 14% and 42 ± 12%, respectively. Mean extinction in the AS/CI region was 38 ± 14%. Nss sulfate aerosol contributed less than 9% to supermicron extinction in all regions (Figure 10) and between 9% (SHAtl) and 46% (EIndSub) to the sub-10 μm extinction (Figure 11).

### 4.4.2. Sea-Salt Aerosol

The contribution of sea salt aerosol to submicron extinction was highest in the southern hemisphere marine regions (SHAtl = 58 ± 23%, SHIO = 56 ± 29%) and relatively high in the northern hemisphere marine region of NHIO (28 ± 11%). For the more continentally influenced regions, extinction fractions ranged from around 3 to 6% (IndSub and EIndSub) to 20% (Arabia). The sea-salt extinction fractions in the INDOEX continentally influenced regions are low compared with those reported for regions of the Atlantic impacted by continental emissions from North America (36%) and biomass burning (29%) and dust (67%) emissions from Africa [Quinn et al., 2001]. The difference is a result of higher concentrations of non-sea-salt chemical components rather than lower sea-salt concentrations.

The sea-salt aerosol contribution to supermicron extinction ranged from 51% (IndSub) to 97% (SHAtl). The dominance by sea salt (even though IOM concentrations were comparable in some regions) is a result of its propensity for water uptake with increasing RH and its reluctance to release water with decreasing RH. As the relative humidity decreases from greater than 70% to 55%, sea salt retains water to the extent that the wet particle mass is twice the drier mass [Tang et al., 1997]. Contribution of sea salt to sub-10 μm extinction ranged from 7 to 83%. This range is lower than that estimated for regions of the Atlantic (47 to 93%) [Quinn et al., 2001] indicating the importance of non-sea-salt chemical components in determining the aerosol radiative forcing in the continentally influenced INDOEX air masses.

### 4.4.3. IOM

For the purpose of the extinction calculations, the IOM and residual mass were combined. Hence the extinction fractions reported for IOM are based on both the IOM and the residual mass. The fraction of submicron extinction due to IOM was less than 1% in the SHAtl region. Mean submicron extinction due to IOM in the SHIO region was 7.4 ± 3.3% but was primarily due to residual mass. The mean extinction fraction in the NHIO region was 14 ± 7.0% and also was primarily due to residual mass. Mean submicron IOM extinction fractions were relatively low and comparable for the EIndSub.
The variability in the submicron POM extinction fractions is
relatively high in both marine and continentally-influenced air
mass regions (22 ± 7.9% in the SHAtl region and 23 ± 6.3% in
the Arabia region). Mean values also were low in both marine
and continentally influenced air mass regions (2.8 ± 0.1% in
the NHIO region and 6.8 ± 2.6% in the AS/CI region).

The mean POM submicron extinction fraction was variable
and seemingly independent of air mass type. It was
relatively high in both marine and continentally-influenced air
mass regions (22 ± 7.9% in the SHAtl region and 23 ± 6.3%
in the Arabia region). Mean values also were low in both
marine and continentally influenced air mass regions (2.8 ±
0.1% in the NHIO region and 6.8 ± 2.6% in the AS/CI region).

The variability in the submicron POM extinction fractions is
similar to what was found for marine and continentally-influ-
enced regions of the Atlantic. During the Aerosols'99 cruise,
mean extinction fractions ranged from less than 1% for air
masses influence by African dust and 18% for northern hemi-
sphere marine air masses [Quinn et al., 2002].

Mean supermicron extinction fractions were consistently
low (<6%) in all regions except the EIndSub and Ind-
Sub regions where they reached values of 23 ± 18% and 20 ±
6.4%, respectively. Mean sub-10 μm extinction fractions of
POM ranged from 1.6% (NHIO) to 15% (Arabia).

4.4.5. BC and nss K⁺ (Burning Component)

BC and nss K⁺ associated with SO₄²⁻ and NO₃⁻ were
 grouped into a burning component. Extinction due to the en-
tire component was then calculated. This component contrib-
uted less than 1% to the submicron extinction in the southern
hemisphere marine regions. Mean submicron extinction for
the NHIO was higher (15 ± 5.5%) indicating transport of
continental aerosol to the ITCZ. Mean submicron extinction
was relatively low for the two regions with lower-level flow
from Arabia (7.7 ± 1.8% for Arabia and 12 ± 2.5% for Arabia/
IndSub). It was 21 ± 7.5% for the AS/CI region and highest for
the two regions with lower-level flow from the Indian subcon-
tinent (36 ± 11% for EIndSub and 33 ± 11% for IndSub).
Contributions to supermicron extinction were negligible.

The substantial contributions of BC to extinction in the
Indian subcontinent air masses is similar to what was observed
during the 1998 INDOEX IFP. Model calculations using data
collected at the Kaashidhoo Climate Observatory estimated
that 11% of the e at ambient RH was due to soot [Satheesh et
al., 1999]. Similar results were obtained for the 1999 INDOEX
IFP [Ramanathan et al., 2001].

4.5. Mass Extinction Efficiencies of the Individual
Aerosol Chemical Components

4.5.1. Comparison of Two Methods

Mass extinction efficiencies e of individual chemical
components are defined as

\[ e_{sp,j} = \frac{\sigma_{sp,j}}{m_j}, \]

where \( \sigma_{sp,j} \) is the extinction coefficient for component \( j \) and \( m_j \)
the mass of component \( j \). An empirical and a calculational
approach were used to calculate \( e_{sp,j} \) for the sub-10 μm size
range (at 55% RH and 550 nm) to check for consistency
between the methods. A thorough discussion of the advantages
and disadvantages of each approach can be found in the work
of Charlson et al. [1999]. The empirical approach used a mul-
tiple linear regression of the mass concentration of the major
chemical components against the extinction coefficient for the
whole aerosol. The following equation was used to obtain
weighted averages of the extinction efficiencies:

\[ \sigma_{sp} = \alpha_{sp,sea} \cdot m_{sea} + \alpha_{sp,SO4} \cdot m_{SO4} + \alpha_{sp,IOM} \cdot m_{IOM} + \alpha_{sp,BC} \cdot m_{BC}, \]

where \( \sigma_{sp} \) is the sum of the measured sub-10 μm \( \sigma_{sp} \) and \( \sigma_{ep} \)
and the mass concentrations are a sum of the measured sub-
micron and supermicron concentrations for each component.
The IOM component consists of IOM and residual mass. Re-
sidual mass made up less than 4% of the sub-10 μm mass in all
source regions. The entire data set was used rather than de-
Table 9. Comparison of Mass Extinction Efficiencies Derived From Mie Calculations (Calculated) and From a Multiple Linear Regression (Empirical) at 550 nm

<table>
<thead>
<tr>
<th>Component</th>
<th>Empiricalb</th>
<th>Coefficient</th>
<th>Standard Error</th>
<th>Calculated</th>
<th>Mean</th>
<th>Range</th>
<th>Previously Reported</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sea salt</td>
<td></td>
<td>1.7</td>
<td>0.4</td>
<td></td>
<td>1.6</td>
<td>0.98–3.3</td>
<td>0.61–2.1</td>
<td></td>
</tr>
<tr>
<td>NSS SO$_4^-$ ion</td>
<td></td>
<td>2.0</td>
<td>0.78</td>
<td></td>
<td>4.9</td>
<td>2.0–6.6</td>
<td>4.2–7.5</td>
<td></td>
</tr>
<tr>
<td>IOM plus residual</td>
<td></td>
<td>0.51</td>
<td>4.3</td>
<td></td>
<td>1.5</td>
<td>0.56–2.7</td>
<td>1.07$^d$</td>
<td></td>
</tr>
<tr>
<td>POM</td>
<td></td>
<td>1.6</td>
<td>4.3</td>
<td></td>
<td>4.1</td>
<td>1.2–5.8</td>
<td>0.73–0.79$^a$</td>
<td></td>
</tr>
<tr>
<td>BC$^e$</td>
<td></td>
<td>8.5$^f$</td>
<td>1.3</td>
<td></td>
<td>8.8</td>
<td>1.6–14</td>
<td>2.6–3.6$^g$</td>
<td></td>
</tr>
</tbody>
</table>

$^a$Values are for the size range $D_{max} < 10$ μm (at 55% RH) and are means over the entire experiment. Units are in m$^2$ g$^{-1}$.  
$^b$Based on equation (6); $r^2 = 0.93$; number of samples equal to 41.  
$^c$From a regression similar to equation (6) where $y$ is the measured absorption coefficient. Only samples with measurable black carbon were included; $r^2 = 0.93$; number of samples equal to 22.  
$^d$Chiapello et al. [1999]; Sal Island, 670 nm.  
$^e$Li et al. [1996]; Barbados.  
$^f$Patterson and McMahon [1984]; Tang et al. [1982]; smoke particles in the smoldering phase.  
$^g$Mass absorption efficiency.  
$^h$Quinn et al. [1996]; Pacific Ocean.

...riving regional averages because of the small number of samples collected in some regions. The calculational approach, based on the Mie calculation described in section 3.2, was used to calculate $\alpha_{ep,j}$ for the sub-10 μm size range for comparison to the empirical approach.

[94] Mean values from the two methods are compared in Table 9. For all components except the IOM, $\alpha_{ep,j}$ calculated from the multiple linear regression falls within the range of values derived from the Mie calculation. The value of $\alpha_{ep,j}$ from the regression is about 10% below the range from the Mie calculation but has a large standard error. The agreement between the two independent methods confirms the internal consistency in the data set and indicates that the derived parameters are accurate within experimental uncertainty.

### 4.5.2. Submicron and Supermicron Component Mass Extinction Efficiencies

[95] The calculational method also was used to calculate $\alpha_{ep,j}$ at 550 nm for the submicron and supermicron size ranges for the different air mass source regions. In these calculations, sea salt and NO$_3^-$ in the supermicron size range were combined to form the sea-salt component, and nss SO$_4^-$ was combined with NH$_4^+$ to form the sulfate aerosol component. Both of these components also included the mass of water calculated to be associated with them at 55% RH. The value of $\alpha_{ep,j}$ was also calculated for the nss sulfate ion such that $\alpha_{ep,j}$ is the scattering due to sulfate aerosol (nss SO$_4^-$, NH$_4^+$, and water at 55% RH) and $m_j$ is the mass of the nss SO$_4^-$ ion. The value of $\alpha_{ep,j}$ for the nss SO$_4^-$ ion is a useful quantity as chemical transport models predict the ion concentration or column burden of sulfate rather than the sulfate aerosol concentration [e.g., Langer and Rodhe, 1991]. Mean values are reported in Table 10 for the different air mass source regions.

[96] Mean mass extinction efficiencies of submicron sea salt ranged from 5.1 to 6.0 m$^2$ g$^{-1}$. Supermicron values ranged from 1.0 to 1.2 m$^2$ g$^{-1}$. Both sets of values compare well with those estimated for latitude bands of the Atlantic during the Aerosols99 Cruise. For the Atlantic, mean submicron values ranged from 5.4 to 7.8 m$^2$ g$^{-1}$, and supermicron values ranged from 0.9 to 1.3 m$^2$ g$^{-1}$ for the aerosol at 55% RH [Quinn et al., 2002].

[97] Mean $\alpha_{ep,j}$ for submicron nss SO$_4^-$ aerosol ranged from 2.9 to 5.3 m$^2$ g$^{-1}$ and for submicron nss SO$_4^-$ from 3.5 to 6.3 m$^2$ g$^{-1}$. These values also are comparable to those reported for latitude bands of the Atlantic (2.0 to 3.8 m$^2$ g$^{-1}$ for sulfate aerosol and 2.5 to 5.8 m$^2$ g$^{-1}$ for the sulfate ion). In addition, they fall within the theoretical range of low RH sulfate scattering efficiencies predicted by Charlson et al. [1999].

[98] Mean mass extinction efficiencies for submicron IOM ranged from 3.0 to 4.3 m$^2$ g$^{-1}$. Supermicron values ranged from 0.5 to 1.4 m$^2$ g$^{-1}$. These values are similar to those estimated for African dust that had been transported to the central Atlantic (3.5 m$^2$ g$^{-1}$ for the submicron size range and 0.5 m$^2$ g$^{-1}$ for the supermicron size range) [Quinn et al., 2001].

[99] Mean submicron POM mass extinction efficiencies ranged from 4.4 to 7.6 m$^2$ g$^{-1}$ over all source regions. Supermicron values ranged from 2.8 to 5.5 m$^2$ g$^{-1}$. Because of the lack of information about the hygroscopicity of the sampled organic matter, no water was associated with the POM. By assuming that the water uptake by POM is similar to that of sulfate aerosol, Lioussse et al. [1996] calculated an increase in $\alpha_{ep,j}$ from 4 for a dry aerosol to 6.8 m$^2$ g$^{-1}$ for an aerosol at 80% RH due to an increase in particle size. The competing effect of lowering the refractive index must also be considered, however. Mean submicron mass extinction efficiencies for the burning component (BC plus submicron KNO$_3$ and K$_2$SO$_4$) ranged from 1.2 to 5.8 m$^2$ g$^{-1}$. Mean submicron mass absorption efficiencies for the burning component ranged from 9 to 15 m$^2$ g$^{-1}$.

### 4.6. Aerosol Optical Depth

[100] Lidar measurements made on the Ronald H. Brown indicated the presence of a single aerosol layer in the southern hemisphere marine regions which was embedded within the marine boundary layer (MBL) and reached up to 3 km [Welton et al., 2002]. No lidar measurements were made during the time period corresponding to the NHIO region. However, both 500 and 2500 m trajectories indicate that the sampled air had been over the northern hemisphere Indian Ocean for several days prior to being sampled (Table 1). In the northern hemisphere continental grid regions, an upper layer (UL) of aerosol was present between altitudes of 1 to 3 km.
Table 10. Mean and Standard Deviation (1σ) of the Mass Extinction Efficiencies of the Major Chemical Components in Each Air Mass Region

<table>
<thead>
<tr>
<th>Region</th>
<th>Sea Salt</th>
<th>NSS Sulfate Ion</th>
<th>NSS Sulfate Aerosol</th>
<th>IOM Plus Residual</th>
<th>POM</th>
<th>BC and nss K⁺</th>
<th>Total Aerosol</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mean</td>
<td>s.d.</td>
<td>mean</td>
<td>s.d.</td>
<td>mean</td>
<td>s.d.</td>
<td>mean</td>
</tr>
<tr>
<td>SH Atlantic</td>
<td>5.1</td>
<td>1.3</td>
<td>3.5</td>
<td>1.5</td>
<td>2.9</td>
<td>0.5</td>
<td>3.0</td>
</tr>
<tr>
<td>SH Indian Ocean</td>
<td>5.3</td>
<td>1.5</td>
<td>4.1</td>
<td>2.2</td>
<td>3.4</td>
<td>0.9</td>
<td>3.5</td>
</tr>
<tr>
<td>NH Indian Ocean</td>
<td>5.7</td>
<td>1.8</td>
<td>5.1</td>
<td>2.5</td>
<td>4.0</td>
<td>1.0</td>
<td>3.6</td>
</tr>
<tr>
<td>East Indian</td>
<td>5.7</td>
<td>0.7</td>
<td>6.3</td>
<td>0.1</td>
<td>5.3</td>
<td>0.1</td>
<td>4.3</td>
</tr>
<tr>
<td>Subcontinent</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Arabian Sea</td>
<td>5.3</td>
<td>1.2</td>
<td>5.7</td>
<td>2.1</td>
<td>4.4</td>
<td>0.9</td>
<td>3.3</td>
</tr>
<tr>
<td>Arabian-Indian</td>
<td>5.3</td>
<td>1.1</td>
<td>4.2</td>
<td>1.2</td>
<td>3.2</td>
<td>0.4</td>
<td>3.1</td>
</tr>
<tr>
<td>Arabian Sea coastal</td>
<td>5.9</td>
<td>1.2</td>
<td>5.4</td>
<td>1.8</td>
<td>4.2</td>
<td>0.6</td>
<td>3.2</td>
</tr>
<tr>
<td>India</td>
<td>6.0</td>
<td>1.5</td>
<td>5.3</td>
<td>2.1</td>
<td>4.1</td>
<td>0.8</td>
<td>3.3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Region</th>
<th>Supermicron</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>mean</td>
</tr>
<tr>
<td>SH Atlantic</td>
<td>1.0</td>
</tr>
<tr>
<td>SH Indian Ocean</td>
<td>1.1</td>
</tr>
<tr>
<td>NH Indian Ocean</td>
<td>1.0</td>
</tr>
<tr>
<td>East Indian</td>
<td>1.0</td>
</tr>
<tr>
<td>Subcontinent</td>
<td></td>
</tr>
<tr>
<td>Arabian Sea</td>
<td>1.1</td>
</tr>
<tr>
<td>Arabian-Indian</td>
<td>1.1</td>
</tr>
<tr>
<td>Arabian Sea coastal</td>
<td>1.2</td>
</tr>
</tbody>
</table>

*Integral values are reported at 55% RH for the submicron (D_{aero} < 1.1 μm) and supermicron (1.1 < D_{aero} < 10 μm) size ranges. Values are based on a Mie calculation at 550 nm. Units are in m² g⁻¹.

1 Includes NO₃ and water at 55% RH.
2 Includes scattering due to nss SO₄²⁻, NH₄⁺, and water at 55% and mass due to nss SO₄²⁻ ion.
3 Includes scattering due to nss SO₄²⁻, NH₄⁺, and water at 55% and mass due to nss SO₄²⁻, NH₄⁺, and water at 55% RH.

éral well-defined column aerosol situations become apparent [Welton et al., 2002]. These situations include (1) regions where there was a single aerosol layer confined to the MBL (SHAtl and SHIO), (2) regions where the MBL aerosol and the UL aerosol originated from the same or different source regions and were well mixed in the atmospheric column (EIndSub, IndSub, and portions of Arabia/IndSub, Arabia, and AS/CI), and (3) regions where the MBL aerosol and the UL aerosol originated from the same or different source regions and were not well mixed in the atmospheric column (portions of Arab/IndSub, Arabia, and AS/CI). Relationships between surface extinction and column extinction are expected to vary for these different situations.

[102] Mean regional τ_a are compared to surface extinction coefficients derived from measured σ_{ap} and σ_{sp} in Figure 7. As for surface extinction, relatively low τ_a were measured in the southern hemisphere marine regions (0.08 ± 0.02 for SHAtl and 0.06 ± 0.03 for SHIO). The mean value was slightly higher in the northern hemisphere marine region (0.10 ± 0.03 for NHIO). Also similar to surface extinction, relatively high τ_a were observed in the EIndSub (0.31 ± 0.04) and IndSub (0.32 ± 0.03) regions. For the rest of the source regions there is a departure in the pattern of τ_a from the pattern observed in surface extinction. Mean τ_a for the Arabia/IndSub (0.38 ± 0.07), Arabia (0.19 ± 0.12), and AS/CI region (0.30 ± 0.10) regions were higher than what would be expected based on the surface extinction values.

[103] The high variability in τ_a during the Arabia and AS/CI regions was a result of changing source regions for the MBL and/or UL aerosol. During a portion of the time period of the AS/CI region, trajectory analysis shows that the MBL aerosol originated from the AS/CI region and the UL aerosol originated from the Indian subcontinent. Values of τ_a during this period range from 0.37 to 0.54. Then a transition period occurred where the transport path of the MBL aerosol to the ship moved away from the Indian coast and the UL aerosol spent less time over India before reaching the ship. Values of τ_a during this period ranged from 0.25 to 0.29 with an outlier of 0.55. On the basis of vertical profiles of RH, at the end of the period, the surface moist layer was mixed up to a height of 8 km. The 500 and 2500 m trajectories both indicated flow from the AS/CI to the ship. Values of τ_a ranged from 0.18 to 0.25. For the Arabia region there were two distinct τ_a ranges corresponding to different source regions of the UL aerosol. Values of τ_a ranged from 0.06 to 0.17 when UL aerosol originated over Arabia and from 0.29 to 0.41 when the UL aerosol originated over India.

[104] Values of surface extinction and τ_a were compared more rigorously in a linear regression of individual data points. The regression reveals two distinct groups of data (Figure 12). All of the data from the marine regions (SHAtl, SHIO, and NHIO), EIndSub, and IndSub, and a portion of the data from the Arabia/IndSub, Arabia, and AS/CI regions fall along a regression line with a coefficient of determination, r², of 0.83 (as indicated by solid symbols in Figure 12). These data fall into the first and second situations described above, i.e., either a single aerosol layer was present (SHAtl, SHIO, and NHIO) or the MBL and UL aerosols were well mixed (EIndSub,
tend to have similar vertical aerosol distributions when the distinct air mass types that contain different aerosol mixtures but a significant variation in surface and column extinction within air mass regions. Aerosol layers had similar vertical distributions even though the aerosol originating from similar source regions. When MBL and UL aerosols are well mixed, or when the MBL and UL aerosol originated from similar sources (Arabia/IndSub and AS/CI). In addition, these data include periods from the Arabia region during periods when the MBL and UL aerosol were not well mixed but both came from these regions the MBL and UL aerosol originate from similar source regions.

Also apparent in Figure 12 is the absence of a correlation between surface and column extinction. A regression of all data points results in an $r^2$ of 0.43. If only the solid symbols are included (representing data derived from periods of a single aerosol layer, when the MBL and UL aerosol were well mixed, and when the MBL and UL aerosol originated from similar sources) $r^2$ equal 0.70. To lessen differences due to the choice of wavelength pairs, the Microtops $a$ calculated for the 440/675 wavelength pair also is shown. In this case, $r^2 = 0.67$. The relationship between surface and column $a$ is not as apparent as between surface and column extinction.

UL aerosol properties, including the $a$, were measured onboard the C130 [Sheridan et al., 2002]. Comparing these data to those measured on the ship is difficult, however, since the plane and the ship were rarely in the same vicinity experiencing similar MBL and UL transport patterns. In addition, the aircraft $a$ are for the sub-1 μm aerosol, while the shipboard values in Figures 8 and 13 are for sub-10 μm aerosol (a more appropriate size range for the comparison to column $a$ and $a$). This difference in size ranges does not limit the utility of ship/aircraft comparisons since, for many of the air mass source regions, submicron aerosol dominated extinction (Figure 6). A regression of surface sub-1 μm $a$ against surface sub-10 μm $a$ over all air mass source regions results in an $r^2$ value of 0.96.

On the basis of C130 flights over the northern Indian Ocean (defined as latitudes greater than 5°N), the $a$ decreased with altitude. Mean values for 0 to 1 km, 1 to 3 km, and 3 to 5 km were $1.99 \pm 0.31, 1.83 \pm 0.52$, and $1.78 \pm 0.85$, respectively [Sheridan et al., 2002]. When the northern hemisphere regions are sorted by trajectory, the trend in $a$ with altitude is not as apparent, however. For example, in the six instances where the C130 sampled MBL air from AS/CI and UL air (1 to 4 km) from India (a transport pattern often encountered by the ship), there were flights when the mean UL level leg $a$ was greater than, equal to, or less than the MBL level leg $a$. Variability of the Angstrom Exponent within an air mass source region appears to preclude a well-defined relationship between surface and column values.

5. Conclusions

Measurements made on board the R-V Ronald H. Brown during the 1999 INDOEX IFP indicated a large degree of variability in aerosol chemical and optical properties in the marine boundary layer over the Indian Ocean during the winter monsoon season. This variability is due to the wide range of
aerosol sources that impact the region and to changing transport pathways from the Indian subcontinent and nearby nations to the marine atmosphere. The southern hemisphere Indian Ocean was relatively uninfluenced by continental sources. Aerosol chemical composition, scattering and absorption coefficients, and optical depth for this region were similar to what has been reported for remote regions of the Pacific and Atlantic Oceans. Sea salt dominated both the sub-1 µm and the sub-10 µm extinction by aerosol particles. Compared to the southern hemisphere, northern hemisphere marine regions, which include the ITCZ, had elevated black carbon and nss sulfate aerosol concentrations as well as lower single scattering albedo values indicating the transport of continental aerosol as far south as the ITCZ. Nss sulfate aerosol dominated the sub-1 µm extinction, while sea salt dominated the sub-10 µm extinction.

Simultaneous transport of low-level (500 m) and upper-level (2500 m) air masses from Arabia to the Indian Ocean resulted in higher concentrations of submicron nss sulfate aerosol, particulate organic matter, black carbon, and inorganic oxidized material (IOM). Yet marine boundary layer extinction was low (compared with regions influenced by the Indian subcontinent) due to the relatively small diameter of the accumulation mode and large diameter of the coarse mode. In addition, mean single scattering albedo values were relatively high (0.93 ± 0.02) perhaps indicating a difference in the mass absorption efficiencies of an aerosol whose primary absorbing component is dust and not BC. When low-level flow from Arabia was accompanied by upper-level flow from the Indian subcontinent, mean aerosol optical depths increased from 0.19 to 0.38. In addition, marine boundary layer BC concentrations increased by a factor of 3, and single scattering albedo decreased to 0.89 ± 0.02 presumably due to subsidence of upper troposphere air into the boundary layer.

Transport of low-level air from the Indian subcontinent resulted in the highest concentrations of nss sulfate, POM, BC, and IOM measured during the experiment. Of these air masses, transport from the eastern Indian subcontinent near Calcutta resulted in the highest concentrations of each of these chemical components. Sulfate aerosol and a burning component composed of BC, KNO₃, and K₂NO₃ dominated both the sub-1 µm and sub-10 µm extinction.

The contribution of sea salt to sub-10 µm extinction for the INDOEX continentally influence regions ranged from 7% for the eastern Indian subcontinent to 44% for air masses from Arabia. These values are lower than those reported for the Pacific and Atlantic Oceans, indicating the importance of non-sea-salt chemical components in determining the aerosol radiative forcing in the heavily impacted regions of the Indian Ocean.

The Ronald H. Brown component of the 1999 INDOEX IFP had the instrumentation required for characterizing both surface and column aerosol properties relevant to radiative forcing. Improvements could be made, however, to achieve more accurate estimates of extinction due to each chemical component. These include (1) measuring the size distribution and determining the molecular composition of IOM and (2) determining the chemical composition of the aerosol organic matter. The latter would help in determining the hygroscopicity of the organic aerosol and the origin of the organic species in the sub- and supermicron size ranges. Despite these shortcomings, the regional aerosol properties presented here provide a unique data set for comparison to parameters derived from chemical transport and radiative transfer models and satellite retrievals.

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