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In situ vertical profiles of aerosol extinction, mass, and composition over the southeast United States during SENEX and SEAC⁴RS: observations of a modest aerosol enhancement aloft

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Abstract

Vertical profiles of submicron aerosol over the southeastern United States (SEUS) during the summertime from in situ aircraft-based measurements were used to construct aggregate profiles of chemical, microphysical, and optical properties. Shallow cumulus convection was observed during many profiles. These conditions enhance vertical transport of trace gases and aerosol and create a cloudy transition layer on top of the sub-cloud mixed layer. The trace gas and aerosol concentrations in the transition layer were modeled as a mixture with contributions from the mixed layer below and the free troposphere above. The amount of vertical mixing, or entrainment of air from the free troposphere, was quantified using the observed mixing ratio of carbon monoxide (CO). Although the median aerosol mass, extinction, and volume decreased with altitude in the transition layer, they were ~ 10 % larger than expected from vertical mixing alone. This enhancement was likely due to secondary aerosol formation in the transition layer.

- organic aerosol (OA) were both similar in magnitude, only the enhancement of sulfate was statistically significant. The column integrated extinction, or aerosol optical depth (AOD), was calculated for each individual profile, and the transition layer enhancement of extinction typically contributed less than 10 % to the total AOD. Our measurements and analysis were motivated by two recent studies that have hypothesized an enhanced layer of secondary organic aerosol (SOA) aloft to explain the summertime
- 20 enhanced layer of secondary organic aerosof (SOA) aloft to explain the summertime enhancement of AOD (2–3 times greater than winter) over the southeastern United States. In contrast to this hypothesis, the modest enhancement we observed in the transition layer was not dominated by OA and was not a large fraction of the summertime AOD.



1 Introduction

Shallow cumulus convection is common over the southeastern United States (SEUS) during the summer. It enhances the vertical transport of trace gases and aerosol, and creates a transition layer between the mixed layer and free troposphere (Siebesma,

⁵ 1998). Due to the presence of clouds and entrainment in the transition layer, it has also been referred to as the cloud layer and the entrainment zone. The transition layer is intermittently mixed by thermal plumes that originate in the mixed layer and form cumulus clouds that release latent heat within the layer. There have been several observations of vertical transport and redistribution of trace gases by shallow cumulus convection
 (Angevine, 2005; Ching and Alkezweeny, 1986; Ching et al., 1988; Greenhut, 1986), and a few studies have investigated the vertical transport and aerosol formation during cumulus convection (Ching et al., 1988; Sorooshian et al., 2007, 2006; Wonaschuetz

et al., 2012).

Based on the seasonality of the surface aerosol-AOD relationship in the SEUS and

- the spatial similarity of biogenic emissions and enhanced AOD, Goldstein et al. (2009) and Ford and Heald (2013) have hypothesized the existence of a layer of enhanced SOA aloft in the summer which contributes to AOD but not to surface measurements of aerosol mass. Although, neither study speculates about meteorological and chemical mechanisms that would lead to its formation, aerosol production in the transition layer
- is a plausible mechanism that could produce the hypothesized layer. More generally, the vertical distribution of aerosol and aerosol formation are integral to understand the relationship between aerosol mass (PM_{2.5}) at the surface and AOD (Hoff and Christopher, 2009).

Submicrometer aerosol particles, which dominate aerosol mass, are largely secondary and composed of OA and sulfates during the summer in the SEUS (Edgerton et al., 2005; Weber et al., 2007). While the formation mechanisms of secondary particulate sulfate are well understood (e.g. Seinfield and Pandis, 1998), the formation of SOA is more complex and uncertain. Both biogenic and anthropogenic precursor emissions



are thought to be important (de Gouw and Jimenez, 2009). The relative importance of the homogenous and aqueous oxidation pathways for both sulfate and OA is also uncertain (Carlton and Turpin, 2013; Carlton et al., 2008; Eatough et al., 1994; Ervens et al., 2011; Luria and Sievering, 1991; McKeen et al., 2007). Based on the abundance of aerosol water and cumulus convection, aqueous processing is expected to be an

important aerosol formation pathway in the SEUS (He et al., 2013), and processing in cloud droplets would occur primarily in the transition layer.

In this analysis, aircraft-based in situ measurements of aerosol chemical, physical, and optical properties are used to examine the vertical structure of aerosol in the SEUS

- ¹⁰ during shallow cumulus convection and to quantify aerosol enhancements in the transition layer and its contribution to summertime AOD. We use measurements made aboard the National Oceanic and Atmospheric Administration (NOAA) WP-3-D aircraft during the NOAA Southeast Nexus (SENEX) study in June and July of 2013 and the National Aeronautic and Space Administration (NASA) DC-8 aircraft during the Study
- ¹⁵ of Emissions and Atmospheric Composition, Clouds, and Climate Coupling by Regional Surveys (SEAC⁴RS) in August and September of 2013 to construct aggregate vertical profiles of aerosol extinction, mass, and composition as a function of altitude over the SEUS. The transition layer aerosol and trace gas concentrations are modeled as a mixture with contributions from the free troposphere and mixed layer. The in situ
- ²⁰ measurements of the extinction coefficient are used to calculate the AOD and contributions to the AOD from aerosol water, from the mixed layer, and from the transition layer.

2 Methods and measurements

In this analysis we combine data collected during two aircraft field studies that were partially conducted over the SEUS in the spring and summer of 2013. Although the SENEX study collected measurements in late spring and early summer while SEAC⁴RS collected measurements in the late summer, both studies encountered shallow cumulus



convection. Additionally, both aircraft hosted a similar set of in situ instruments which was conducive to a combined analysis.

2.1 SENEX

The SENEX study was part of the larger Southeast Atmosphere Study (SAS) in the SEUS during the June and July of 2013. The NOAA WP-3-D aircraft flew 18 research flights based out of Smyrna, Tennessee during June and July 2013 for SENEX with a payload of instruments measuring atmospheric trace gases, aerosol properties, and meteorological parameters. This analysis uses measurements of carbon monoxide (CO), carbon dioxide (CO₂), methane (CH₄), sulfur dioxide (SO₂), isoprene, aerosol composition, and aerosol optical properties (Table 1). The aerosol was sampled downstream of a low turbulence inlet (Wilson et al., 2004) and an impactor with a 1 μm aerodynamic diameter size cut. Before impaction, the sampled aerosol was initially dried by ram heating when sampling into the aircraft. The sampled aerosol was then dried further in each instrument.

15 2.2 SEAC⁴RS

SEAC⁴RS consisted of measurements aboard three aircraft based in Houston, Texas during August and September of 2013. In this analysis, we focus on in situ measurements from the NASA DC-8 aircraft, which conducted 19 research flights during SEAC⁴RS. The measurements collected onboard the DC-8 used in this analysis are summarized in Table 2. Unlike the SENEX study, there was no continuous measurement of methane during SEAC⁴RS. The aerosol extinction and black carbon instruments flown on the DC-8 during SEAC⁴RS were the same instruments used onboard the NOAA WP-3-D aircraft during SENEX. Measurements of aerosol extinction, volume, black carbon mass sampled aerosol through a shrouded diffuser inlet described

²⁵ by McNaughton et al. (2007). The aerosol extinction and volume were measured downstream of 1 μm impactor. The aerosol mass spectrometer (AMS) sampled aerosol



through a HIMIL inlet (http://www.eol.ucar.edu/homes/dcrogers/Instruments/Inlets/). In both inlets, the sampled aerosol was initially dried by ram heating and then further dried in each instrument.

2.3 Surface measurements

⁵ The Southeastern Aerosol Research and Characterization (SEARCH) Network consists of 8 continuous monitoring ground sites in Georgia and Alabama hosting several gas-phase and aerosol measurements (Edgerton et al., 2005, 2006; Hansen et al., 2003). During SENEX the NOAA WP-3-D flew over four of these sites a total of 15 times, and extinction near the surface is calculated using measurements of aerosol scattering (Radiance Research Model M903 nephelometer, Tempe, Arizona, USA) and absorption (Magee Scientific Model AE-16 Aethalometer, Berkeley, California, USA) at each SEARCH site to compare with the extinction measured onboard the NOAA WP-3-D aircraft in the mixed layer.

2.4 Aerosol water

- The enhancement of extinction due to condensation of water onto the aerosol is modeled using an empirical parameterization (shown in Eq. 1), hereafter referred to as the kappa parameterization (Brock et al., 2015). The hygroscopic growth of particle diameter is described by kappa-Kohler theory (Petters and Kreidenweis, 2007). A particle size distribution and a Mie scattering calculation would be necessary to rigorously ex-
- tend the kappa-Kohler theory to the hygroscopic enhancement of optical properties. However, Brock et al. (2015) show that if atmospheric accumulation mode size distributions typical of the SEUS are used, the functional form of kappa-Kohler theory can be applied directly to the optical extinction (Eq. 1).

$$\sigma_{\text{ext}}(\text{RH}) = \sigma_{\text{ext}}(\text{dry}) \times \left(1 + \kappa \times \left(\frac{\text{RH}}{100 - \text{RH}}\right)\right)$$

(1)

The humidified extinction coefficient $\sigma_{ext}(RH)$ is a function of the dry extinction $\sigma_{ext}(dry)$ and the hygroscopicity parameter κ . The aerosol extinction is measured in three separate constant RH channels: in dry conditions (RH less than 30%), medium RH typically 70%, and high RH greater than 80%. The hygroscopicity parameter (κ) is determined

⁵ by fitting the three measurements of extinction to Eq. (1). The resulting κ and $\sigma_{ext}(dry)$ are then used to estimate the extinction at ambient RH. Lower values of aerosol hygroscopicity generally correspond to mineral dust, aerosol with high soot fraction, or primary OA such as fresh biomass burning emissions or automotive emissions (Massoli et al., 2009; Quinn et al., 2005). High hygroscopicity usually corresponds to an oxidized, aged aerosol, large sulfate mass fractions and/or sea salt aerosol.

The calculated ambient extinction is a lower limit of the actual ambient extinction in three cases. First, if the hygroscopic growth exhibits hysteresis and the ambient RH is below the deliquesce RH, ambient particles may be on either the deliquescing (lower) or the efflorescing (upper) branch of the hysteresis curve (Santarpia et al., 2004). Our

- ¹⁵ measurements cannot distinguish between these two states, because aerosol is dried and then re-humidified in our instrument before measurement at high RH. Hence, the measured extinction at high RH and the subsequently calculated hygroscopicity parameter κ represent the hygroscopic growth of a deliquescing aerosol on the lower branch. Second, because the kappa parameterization produces an ambient extinction
- that asymptotically approaches infinity as RH approaches 100 %, we used ambient RH to calculate the ambient extinction only when RH was less than 95 %, and assumed a constant RH of 95 % when ambient RH was greater than or equal to 95 %. Therefore, the calculated ambient extinction is a lower limit of the ambient extinction when RH was greater than 95 %. Third, in the case that super-micrometer particles (which
- are not sample by instruments used in this analysis) make a significant contribution, the ambient extinction is underestimated. This is typically the case during dust events and during in-cloud sampling which were either not observed or excluded from this analysis.



2.5 Aggregation of vertical profiles

For this analysis, individual vertical profiles were selected from the research flights by inspection of the altitude time series and are generally included for all ascents and descents with an altitude difference greater than 1 km. Measurements during level flight

- ⁵ legs were not used in this analysis. Although cloud penetration was mostly avoided, aerosol data sampled during cloud penetration events were excluded due to the effects of particle shattering in the inlets. Cloud penetration events were identified using the video from the nose of the aircraft and cloud particle imaging probes mounted near the wingtips. Transects of biomass burning plumes were identified using tracers such
- ¹⁰ as the acetonitrile mixing ratio and were excluded from the analysis. Extensive aerosol parameters (mass, extinction, volume) have been corrected to standard temperature (0°C) and pressure (1 atmosphere). All calculated quantities such as ambient extinction and transition layer enhancements were determined before aggregation and then were aggregated in the same manner as the observations.
- ¹⁵ For the SENEX campaign, the vertical profiles (which were located primarily over northern Georgia and Alabama) were generally included in the flight plans for three purposes: to characterize the background boundary layer structure before and after urban and power plant plume intercepts, to characterize the vertical structure over surface measurement sites, and as enroute ascents and descents into and out of the region of
- interest. The SEAC⁴RS profiles that we use were distributed through Mississippi and Alabama and were conducted to characterize inflow and outflow near convective systems, to examine boundary layer chemistry over the SEUS, and as enroute ascents and descents. The individual vertical profiles used here include both spiral and slant ascents and descents and were typically between 5 and 15 min in duration.
- ²⁵ To reduce the effects of meteorological variability and spatial heterogeneity of emissions, we have combined the individual vertical profiles to produce aggregate profiles that are representative of the SEUS region. In this analysis we construct two types of aggregate profiles, the first of which includes all of the afternoon vertical profiles over



the SEUS and is binned according to altitude above ground level ("altitude-binned" aggregate profiles). The second type of aggregate includes only the subset of profiles during which shallow cumulus convection was present and is binned according to a normalized altitude described in Sect. 3.2 ("normalized" aggregate profiles). The profiles not included in the normalized aggregates were either collected during deeper convection and/or had a more complicated structure.

For the altitude-binned profiles, we have chosen all of the available vertical profiles from both SENEX and SEAC⁴RS over Mississippi, Alabama, and Georgia in the afternoon between 12.00 p.m. and 6.30 p.m. Central Daylight Time (CDT) when we expect

- the boundary layer structure to be well developed and no residual layers left over from the previous day. The aggregate includes 74 profiles of which 41 profiles are from 6 research flights during SENEX and 33 profiles from 6 research flights during SEAC⁴RS. The locations of the profiles used in the altitude-binned aggregate are shown in Fig. 1. The data from individual vertical profiles were aggregated into 150 m vertical bins from
- the surface to 4.5 km based on the altitude above ground level. The vertical bin height of 150 m was chosen such that the slower measurements (aerosol mass and isoprene) typically contributed at least one datum to each bin for each individual profile. In each bin with data from 5 or more individual profiles, the median, interquartile range, and interdecile range were calculated. The median and percentiles were used because
- these statistics are more robust when outliers are present. The number of vertical profiles which contribute to each aggregated altitude bin varies with altitude because of the differences of the starting and ending altitudes of each individual profile (Fig. 2a). During some profiles or some portions of a profile, individual measurements of trace gases and aerosol properties did not report data (e.g. due to zeroing or calibrations).
- The normalized aggregate profiles were calculated using only those individual profiles obtained during shallow cumulus convection and were altitude-normalized as described in Sect. 3.2. Shallow cumulus convection is common globally and over the SEUS. Warren et al. (2007) have compiled a global cloud climatology based on surface observations. According to their work, the mean frequency of daytime cumulus



clouds is 49 % over Alabama during June, July, and August, and the mean cloud coverage when cumulus clouds are present is 35 %. The presence of shallow cumulus convection during individual profiles was determined by inspection of visible images from the GOES satellite and the presence of a three layer structure (mixed layer, transition

- Iayer, and free troposphere), which is expected during shallow cumulus convection. The second aggregate includes 37 of the 74 SEUS profiles from the first aggregate. The locations of profiles in the second aggregate are show in Fig. 1, and Fig. 2b shows the distributions of the mixed layer and transition layer heights determined from individual profiles which had medians of 1.2 and 2.2 km respectively. For cumulus convection
- the height of the planetary boundary layer is defined as cloud base or the mixed layer height; however, we find the term "planetary boundary layer" confusing in the context of shallow cumulus convection and have avoided using it. Determination of the mixed layer height and transition layer height is described in Sect. 3.2. For the normalized aggregate profiles, there are 25 bins assigned to each layer. Figure 2c shows the num-
- ¹⁵ ber of profiles that contribute to the second aggregate at each normalized altitude. The number of profiles varies with normalized altitude due to variability in the starting and ending altitudes of each profile and because the second aggregate is limited to portions of the individual profiles when aerosol mass, extinction, volume and CO measurements all report data. Limiting data as such facilitates quantitative comparison of the aggregate profiles. In contrast the first aggregate is constructed using all available

3 Results

data.

3.1 Altitude-binned aggregate profile

Altitude-binned aggregate profiles of dry and ambient aerosol extinction show several characteristics of note (Fig. 3). The median 532 nm dry extinction coefficient (Fig. 3a) is approximately independent of altitude below 1.5 km with median value of 50 Mm⁻¹,



and the interquartile range is 27 to $73 \,\text{Mm}^{-1}$. The interquartile (25th–75th percentiles) and interdecile (10th–90th percentiles) range are due largely to variation between individual profiles rather intra-profile point-to-point variation. Above 1.5 km the extinction coefficient decreases with altitude to a median value of $6 \,\text{Mm}^{-1}$ above 3.0 km. The

- ⁵ gradual decrease in extinction with altitude from 1.2 to 2.5 km is partly due to the variation of mixed layer and transition layer heights in the individual profiles. Figure 3b shows the RH increasing with increasing altitude below 1.2 km. Above this level the RH has a slight decreasing trend with altitude and a large interquartile range spanning 30–70%. The relative humidity of the aggregate profile could be biased low, because during SENEX the flight dates were chosen to avoid precipitation and cloud penetration.
- ¹⁰ during SENEX the flight dates were chosen to avoid precipitation and cloud penetration was mostly avoided during flights however this was not the case during SEAC⁴RS.

The median hygroscopicity parameter (κ , Fig. 3c) increases from 0.11 at the bottom of the profile to 0.18 at 3 km and is more variable above 3 km. The hygroscopic growth of the aerosol enhances the ambient extinction (Fig. 3d) throughout the profile and

significantly between 0.7 and 1.7 km. Below 1.2 km, the ambient extinction coefficient increases with altitude due to increasing RH, and above 1.2 km decreases with altitude due to a combination of decreasing RH and decreasing dry extinction. The hygroscopic growth of aerosol and the subsequent enhancement of extinction aloft could explain some of the enhancement of AOD noted by Goldstein et al. (2009) and Ford and Heald (2013).

The minimum altitude of individual aircraft profiles ranged from 300–700 m above the surface. We estimate the profile of dry extinction between the surface and the minimum altitude of the profiles by combining aircraft measurements made in the mixed layer in the vicinity of surface monitoring sites using ground data from those sites. Dur-

ing the SENEX study, there were 15 overflights in the mixed layer within 10 km of four SEARCH monitoring sites. The surface aerosol extinction at each SEARCH site was calculated using the aerosol scattering coefficient measured by a nephelometer with a center wavelength of 530 nm and the aerosol absorption coefficients measured by an aethalometer at 880 nm. Because the optical absorption at this wavelength was likely



due to black carbon aerosol, we corrected the absorption coefficient to 530 nm using an Ångstrom exponent of 1 which is conventionally used for black carbon (Bergstrom et al., 2002; Lack and Langridge, 2013). Absorption typically accounted for less than 5% of the extinction. The calculated 530 nm surface extinction was not corrected to the

- ⁵ 532 nm aircraft extinction because the correction would be less than 1 %. The surface and aircraft extinction coefficients are correlated ($R^2 = 0.91$), and the slope of a linear least squares fit to the data indicates that the aircraft data are ~ 6% lower than the surface measurements (Fig. 4), which is within the combined uncertainty in the measurements. We conclude that the dry extinction is roughly independent of altitude from the surface to the top of the well-mixed layer. Crumeyrolle et al. (2014) found simi-
- lar agreement between surface and aircraft-based boundary layer measurements of ozone in the Baltimore–Washington metropolitan area.

The altitude-binned aggregate vertical profile of aerosol mass (Fig. 5a) is similar in shape to the dry extinction profile. The median mass is 13.7 μgm⁻³ at the bottom of the profile and decreases to 2.1 μgm⁻³ above 3 km. The aerosol mass is the total of all ions measured by the AMS, and these ions are typically classified based on mass to charge ratio as SO₄, NH₄, NO₃ and OA. The ions may represent simple inorganic salts such as ammonium sulfate and ammonium nitrate in the aerosol or more complex compounds with both an inorganic and organic character (i. e. organosulfates and amines). In this classification scheme, the composition (Fig. 5b) of the submicron aerosol is primarily OA, sulfate, and ammonium. The mass fraction of the inorganic components (NO₃, NH₄, and SO₄) increase with altitude up to 3 km, while the OA mass fraction decreases with altitude up to 3 km. Above 3 km, the OA fraction increases; however, at this altitude

the median aerosol mass is only $2 \mu g m^{-3}$. The increase of aerosol hygroscopicity with altitude up to 3 km corresponds with the increasing inorganic fraction of the aerosol. In particular, sulfate is typically more hygroscopic than OA, is 20% of the aerosol mass at the bottom of the profile, and is 28% of the aerosol mass at 3 km.



3.2 Normalized aggregate profiles

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The heights of the mixed and transition layers varies among individual vertical profiles and this variation obscures the transition layer in the altitude-binned aggregate profile presented in Figs. 3 and 5. For example, the layer structure is clear in a semi-rural profile measured in the vicinity of shallow cumulus convection over central Georgia on the afternoon of 16 June (Fig. 6). The mixed layer is closest to the surface, a transition layer is formed above the mixed layer, and the free troposphere is on top.

The layered structure is evident in both the physical parameters such virtual potential temperature (Θ_v) and ambient temperature as well as chemical mixing ratios such as carbon monoxide (CO) and isoprene. In the mixed layer, adiabatically conserved parameters such as virtual potential temperature (Fig. 6a) are independent of altitude.

- However, in the transition the virtual potential temperature (ng. oa) are independent of altitude. til the top of the transition layer is reached where a capping temperature inversion is present. In the mixed layer, the mixing ratio of water vapor is also independent of alti-
- tude; however, RH increases with altitude as temperature decreases (Fig. 6b). Relative humidity is high in the transition layer, and video from the nose of the aircraft confirms the presence of clouds in this layer.

The transition layer is also evident in the comparison of long-lived trace gases such as CO (Fig. 6c) with short lifetime trace gases such as isoprene (Fig. 6d). Carbon monoxide is directly emitted during combustion, produced by oxidation of hydrocarbons, lost to oxidation by OH, and typically has an atmospheric lifetime of 1–4 months which varies seasonally and regionally (Seinfeld and Pandis, 1998). In this profile, the CO mixing ratio is greater than 110 ppbv in both the mixed and transition layers and decreases to less than 100 ppbv in the free troposphere. In the mixed layer the CO

²⁵ mixing ratio is independent of altitude and decreases modestly with altitude in the transition layer. Isoprene is a short-lived trace gas that typically has an atmospheric lifetime less than 2 h (Seinfeld and Pandis, 1998) and in the summertime is emitted by vegetation common in the SEUS. In the mixed layer, the isoprene mixing ratio is greater than



1 ppbv and variable due to heterogeneous surface emissions (Fig. 6d). The isoprene mixing ratio in the transition layer is always less than 500 ppbv and typically \sim 10 % of the mixed layer value. In the free troposphere, the isoprene mixing ratio is below the detection limit of the measurement.

⁵ To examine vertical structure in more detail, altitude-normalized aggregate profiles were calculated. Altitude normalization is commonly done by dividing the altitude by the height of the mixed layer. However, because of the more complex vertical structure encountered during shallow cumulus convection, we have defined a normalized altitude, h_{norm} , for each profile such that the top of the mixed layer, h_{ML} , is assigned a normalized altitude of 1, and the top of the transition layer, h_{TL} , is assigned a normalized altitude of 2:

$0 < h < h_{ML}$	$h_{\rm norm} = h/h_{\rm ML}$	
$h_{ML} < h < h_{TL}$	$h_{\rm norm} = 1 + (h - h_{\rm ML}) / (h_{\rm TL} - h_{\rm ML})$	(2)
$h > h_{TL}$	$h_{\rm norm} = 1 + h/h_{\rm TL}$	

For individual profiles, the mixed layer height was defined as the highest altitude at which the virtual potential temperature (Θ_v) was constant and there was a reduction in the isoprene concentration. The top of the transition layer was defined by a temperature inversion and a rapid decrease in the CO mixing ratio.

The altitude-normalized aggregate profiles of CO (Fig. 7a) and isoprene (Fig. 7b) mixing ratios demonstrate the contrast between the mixed layer and transition layer. During shallow cumulus convection, CO is transported out of the mixed layer into the transition layer due to its longer lifetime relative to isoprene. The modest decrease of CO with altitude in the mixed layer is likely due to the influence of near source emissions in some profiles. In the mixed layer the isoprene profile is variable, and the median is only modestly dependent on altitude with a median mixing ratio of 1 ppbv. However, the median isoprene mixing ratio decreases to ~ 10% of this value in the transition layer.

²⁵ The isoprene observed above the mixed layer is consistent with large eddy simulations



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performed by Kim et al. (2012) who found that cumulus clouds can transport some isoprene out of the mixed layer into the cloud layer.

4 Analysis

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4.1 Aerosol enhancements in the transition layer

⁵ During shallow cumulus convection, the air in the transition layer is a mixture of air from the mixed layer below and the free troposphere above. The concentrations of trace gases and extensive aerosol parameters C(h) in the transition layer are described in this analysis by a vertical mixing model consisting of three terms (Eq. 3): a contribution from the mixed layer, a contribution from the free troposphere, and any enhancement E(h) relative to concentration expected from the vertical mixing alone, as

$$C(h) = C_{ML}f_{m}(h) + C_{FT}(1 - f_{m}(h)) + E(h),$$

where C(h) is the aerosol or trace gas concentration, $C_{\rm ML}$ and $C_{\rm FT}$ are the aerosol or trace gas concentrations in the mixed layer and the free troposphere. Positive enhancements could be due to local production, or direct emissions to the transition layer from buoyant plumes, e.g. large biomass burning sources, and negative enhancements represent losses.

The fraction of air from the mixed layer (f_m) present in the transition layer is determined by using the CO mixing ratio as

$$f_{\rm m}(h) = \frac{{\rm CO}(h) - {\rm CO}_{\rm FT}}{{\rm CO}_{\rm ML} - {\rm CO}_{\rm FT}},$$

for which the enhancement E(h) due to local production and losses is assumed to be zero. For each profile, the mixing ratio of CO in the mixed layer CO_{ML} and the free troposphere CO_{FT} were determined using the mean between normalized altitudes of

(3)

(4)

0.5 and 0.9 for the mixed layer and 2.0 and 2.5 for the free troposphere. To investigate transition layer enhancements of chemical concentrations and aerosol extensive properties, we calculate a concentration expected from vertical mixing alone using Eq. (3) and setting E(h) to zero. The concentration expected from vertical mixing alone is cal-

- ⁵ culated for each profile and aggregated in the same manner as the observations. In the Figs. 8–10, the median concentration expected from vertical mixing alone is shown as a dashed line on top of the observed concentrations. The interquartile and interdecile ranges of observations are shown; however, they are not shown for the value expecting from vertical mixing.
- ¹⁰ One pathway of CO production is through the oxidation of VOCs, and this CO production likely accounts for a significant fraction of the CO budget during the summer in the SEUS (Hudman et al., 2008). If CO production in the transition layer is significant, the fraction of air from the mixed layer (f_m , determined using CO) would be biased high and any transition layer enhancements of other species (E(h), determined us-
- ¹⁵ ing the CO concentration and Eqs. 3 and 4) would be biased low. By comparing the observed concentration and the concentration expected from vertical mixing alone of several long lifetime species, the importance of CO production in the transition layer can be assessed.

Figure 8a shows the altitude normalized profile of the fraction of air from the mixed
layer (*f*_m, calculated using Eq. 4). The median is 1 in the mixed layer, 0 in the free troposphere, and decreases from 1 to 0.6 in the transition layer due to entrainment of air from the free troposphere. Figure 8b–e shows the altitude-normalized aggregate profiles and the median concentration expected from vertical mixing alone (dashed line) of methane (CH₄), carbon dioxide (CO₂), water vapor (H₂O), and black carbon aerosol mass, respectively. Both CH₄ and CO₂ have atmospheric lifetimes greater than a year and no significant production or losses in the transition layer on the time scale of atmospheric mixing. (The aggregate profile in Fig. 8b only includes data from the SENEX study because CH₄ was not measured during SEAC⁴RS.) H₂O and black carbon aerosol are also not produced in the transition layer and are not lost except in the



presence of precipitating clouds. Profiles in precipitating clouds are mostly excluded from the aggregate. Based on the agreement between the observed vertical profiles of CH₄, CO₂, H₂O, and black carbon mass (Fig. 8) and their expected concentration from vertical mixing alone, we conclude the CO production in the transition layer is not significant.

In contrast, the altitude-normalized profiles of submicrometer aerosol mass (Fig. 9a), extinction (Fig. 9b), and volume (Fig. 9c) are greater than the value expected from vertical mixing alone (dashed lines) in the transition layer. This indicates that E(h) is positive for these aerosol properties. These transition layer enhancements are quantified for individual profiles using the difference between the observed value and the value

- for individual profiles using the difference between the observed value and the value expected from vertical mixing alone. The difference is expressed as a percentage of the observed value and averaged over the transition between normalized altitudes of 1.1 and 1.9. The mean transition layer enhancements of aerosol mass, extinction, and volume were +8.6, +11.3, and +9.3% respectively. The difference in the enhancements of the enhancements of the enhancements.
- of mass, extinction, and volume may reflect actual changes in the aerosol density and extinction cross-section or could be due to imperfections in the measurements and data aggregation.

Altitude-normalized aggregate profiles of aerosol composition are shown in Fig. 10. The enhancement of each aerosol component is quantified in the same manner as aerosol mass, extinction, and volume. The observed median is greater than the value expected from vertical mixing alone by +6% for OA mass, +18% for SO₄, +25% for NH₄, and +15% for NO₃. Although enhancement of sulfate is larger than OA as a percentage, the absolute enhancement is a similar magnitude for both SO₄ and OA, ~ 0.5 μ g m⁻³.

²⁵ The transition layer enhancements can be further investigated by examining the distribution of enhancements for individual profiles (Fig. 11). For each profile, the enhancement is calculated using the absolute difference between the observed value and that expected from vertical mixing alone. The difference is averaged between normalized altitudes of 1.1 and 1.9. Because the distributions of enhancements range



from negative to positive values, the Student's *T* test is used to assess if the enhancement distributions are statistical different from zero, or no enhancement. Enhancement distributions with *p* values less than 0.05 are considered statistically significant. As expected, Fig. 11a–d shows conserved species that do not have statistically significant enhancements: CH₄, CO₂, H₂O, and black carbon mass. The enhancement distribu-

- tions of aerosol mass, extinction, and volume (Fig. 11e–g) are all statistically significant. Although both OA and inorganic aerosol components are enhanced in the transition layer, the enhancement distribution of OA is not statistically significant while the enhancement distribution of inorganic components is significant.
- Aerosol formation in the transition layer is the likely mechanism that would lead to the observed enhancement of aerosol mass, volume, and extinction. The enhancement of aerosol loading is the net result of production and loss in the transition layer; however, profiles of black carbon and total sulfur (see Sect. 4.2) suggest that the aerosol losses are small. Aerosol formation in the transition layer is a combination of aque-15 ous production (both in clouds and aerosol water), homogenous oxidation followed by
- ¹⁵ ous production (both in clouds and aerosol water), homogenous oxidation followed by condensation on existing particles, and condensation of semi-volatile species such as NH₄NO₃. The presence of clouds within the transition layer suggests a large role for aqueous production; however, our dataset does not allow us to determine the relative importance of each pathway.
- Biomass burning, common during the summer in the SEUS, emits aerosol in buoyant plumes that, if large enough, could contribute to the observed enhancement of aerosol loading in the transition layer and would not be consistent with the simple vertical mixing model used here to describe the transition layer concentrations. Profiles during which fresh biomass burning plume were sampled have been excluded from this analysis.
- ²⁵ However, the possibility remains that the aggregate profiles are contaminated by aged and diluted biomass burning plumes which have not been identified. To address this we considered biomass burning emission factors of black carbon, sulfate, and sulfur dioxide (SO₂, which is oxidized in the atmosphere to sulfate) reported by Akagi et al. (2011). The emission factors range from 0.20 to 0.91 gkg⁻¹ for black carbon mass and 0.45



to 0.87 gkg⁻¹ for the combination of SO₂ and sulfate. Based on these emission factors, we would expect the ratio of the combination of SO₂ and sulfate mass to black carbon mass in biomass burning plumes to range from 0.5 to 4.35. If the observed enhancement of sulfate (~ $0.5 \,\mu gm^{-3}$), were due exclusively to biomass burning, we

⁵ would expect a concomitant enhancement of black carbon (based on the ratio of emission factors for black carbon and sulfate) in the range of 100 to 1000 ng m⁻³, which is not observed in the profile of black carbon mass (Fig. 8e). Hence, we conclude the enhancement observed in the altitude-normalized aggregate profile is not due to biomass burning.

10 4.2 Sulfur budget

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Further evidence for the transition layer enhancement of particulate sulfate comes from the reduction of gas-phase SO_2 in the transition layer. Particulate sulfate is produced through gas-phase and aqueous oxidation of SO_2 (Seinfeld and Pandis, 1998). We expect that mixing in the transition layer would conserve total sulfur which we define as the sum of particulate sulfate and gas phase SO_2 . While particulate sulfate is enhanced

- in the transition layer as described in Sect. 4.1, there is also a reduction in the mixing ratio of gas phase SO₂ in the transition layer. Figure 12a–c shows altitude-normalized aggregate profiles and values expected from vertical mixing alone for particulate sulfate, gas phase SO₂, and the total sulfur. Particulate sulfate (Fig. 12a) is enhanced by approximately the same amount as the reduction of SO₂ (Fig. 12b), ~ 0.1 ppbv in
- the transition layer. Consequently, the median value of total sulfur agrees well with the value expected from vertical mixing alone.

The enhancement distributions for particulate sulfate, SO_2 , and total sulfur are shown in Fig. 12d–f. While the transition layer enhancement of particulate sulfate is significant with a *p* value of 3×10^{-5} (Fig. 12d), the reduction of SO_2 in the transition layer (Fig. 12e) is not. The lack of statistical significance in SO_2 reduction is due to positive outliers in the enhancement distribution. The enhancement distribution of total sulfur



indicates a small enhancement that is not statistically significant (Fig. 12f). We note that the conservation of sulfate and SO_2 is only apparent when mixing in the transition layer is taken into account. If biomass burning were the source of the transition layer enhancement of particulate sulfate, we expect total sulfur would be enhanced as well.

5 4.3 Aerosol optical depth

Aerosol optical depth (AOD) is typically measured remotely from space-based satellites (King et al., 1999) and by ground-based sun photometer networks (Holben et al., 2001). These remote measurements of AOD have been complemented by AOD calculated from aircraft-based in situ measurements of extinction which have the ability to quantify contributions to the AOD from individual layers and aerosol water (Crumeyrolle et al., 2014; Esteve et al., 2012). Calculated from in situ measurements, AOD is the integral of the ambient aerosol extinction coefficient (σ_{ext}), Eq. (5)

$$AOD = \int_{\text{surface}}^{\text{TOA}} \sigma_{\text{ext}}(z) dz$$

where σ_{ext} is a function of altitude *z* and the integration extends to the top of the atmosphere (TOA). The extinction coefficient in Eq. (6) is at ambient relative humidity, pressure and temperature. Several assumptions are necessary to calculate ambient extinction and subsequently AOD. First, the aircraft profiles of dry extinction, relative humidity, pressure, and temperature must be extrapolated to the surface. The dry extinction is extrapolated as a constant to the surface based on the mean extinction measured in

the lowest 200 m of each profile. Relative humidity is extrapolated to the surface using the linear trend in the lowest 200 m of each profile if the trend is positive (RH increases with increasing altitude); otherwise, it is extrapolated as a constant based on the mean RH of the lowest 200 m of each profile. Pressure and temperature are both extrapolated using the linear trend in the lowest 200 m of each profile. The second assumption

(5)

is that the contributions to AOD from aerosol layers above top of the aircraft profile are negligible. For example, smoke from large forest fires in the western US can be lofted high into the troposphere and transported over the SEUS (Peltier et al., 2007). This contribution to AOD cannot be included if the smoke layer were above the maximum
 altitude of the profile. In this case the AOD calculated from the in situ profiles is a lower limit. Third, we neglect the contribution to the AOD from supermicroeter particles, which we estimate to be less than 10% of the sub-micrometer AOD based on coarse parti-

cle size distribution measurements made during both SENEX and SEAC⁴RS. Fourth, because we have restricted calculated aerosol hygroscopic growth to RH values less than 95 %, the AOD calculated here is only a lower limit.

In addition to the AOD for each profile in the altitude-normalized aggregate, we have also calculated the contributions to AOD from the mixed layer, the transition layer, aerosol water, and the enhancement of aerosol extinction in the transition layer. The median calculated AOD was 0.14 and the interquartile range (IQR) spanned 0.10 to

- 0.20 (Fig. 13a). An idealized profile of extinction during shallow cumulus convection is used to show the contributions to AOD from the transition layer enhancement of extinction (Fig. 13b), aerosol water (Fig. 13c), the transition layer (Fig. 13d), and the mixed layer (Fig. 13e). The contribution of the transition layer enhancement of ambient extinction (median: 7 %, IQR: 4–10 %) is split between the enhancement of dry extinction
- and the aerosol water associated with the additional aerosol loading. The contribution of aerosol water to the whole profile (median: 33 %, IQR: 24–38 %) is sensitive to the aerosol hygroscopicity parameter and ambient RH encountered. The transition layer contribution (median: 45 %, IQR: 33–55 %) was slightly smaller than the mixed layer contribution (median: 48 %, IQR: 38–57 %). The mixed layer's slightly greater vertical
- extent and higher average dry extinction favor a larger contribution to AOD; however, the transition layer also provides a substantial contribution to AOD because of the aerosol water associated with the higher mean RH in the transition layer. The contributions to AOD presented in Fig. 13 have substantial overlap (i. e. aerosol water also



contributes to mixed and transition layer AOD contributions); hence, the contributions do not add to unity.

The altitude-normalized aggregate profiles used in this analysis are drawn from 37 vertical profiles; however, they represent only eight afternoons during the sum-⁵ mer of 2013. For comparison, Fig. 13a (dashed) shows a histogram of 500 nm AOD (measured by an AERONET sun photometer (Holben et al., 2001) at the Centreville SEARCH site) constructed from all available data between 1 May and 30 September 2013. The AOD from the sun photometer is higher than the AOD from the profiles in the normalized aggregate, and this difference likely reflects bias in sampling shallow cumulus convection during the in situ profiles compared to more uniform sampling by

- the sun photometer. The in situ profiles presented here are weighted toward relatively clean early summer conditions in the presence of shallow cumulus convection, and may be biased against late summer biomass burning influence from the western US and Canada.
- Several SEUS studies have noted decreases in anthropogenic emissions (sulfur, nitrogen oxides, and volatile organic compounds) in the first decade of the 21st century (Alston et al., 2012; Attwood et al., 2014; Hand et al., 2012, 2013). Concurrently, particulate sulfate, OA, and AOD have also decreased. Alston et al. (2012) have shown that the summertime mean AOD over Georgia reported by the MISR instrument on the Torre encount decreased from 20.2 in the summer of 0000 to loss than 0.0 in the
- $_{\rm 20}$ Terra spacecraft decreased from ~ 0.3 in the summer of 2000 to less than 0.2 in the summer of 2009, which is in the range of AOD calculated in this work for the summer 2013.

5 Conclusions

Several preceding studies have observed vertical transport of trace gases and aerosol
 ²⁵ from the mixed layer into the cloud-influenced transition layer during shallow cumulus convection (Angevine, 2005; Ching and Alkezweeny, 1986; Greenhut, 1986; Langford et al., 2010). Our observations are consistent with this earlier work. In addition to ver-



tical transport and redistribution of aerosol, we observed a modest enhancement of aerosol loading in the transition layer and conclude that secondary aerosol formation in the transition layer is the likely source of the enhancement. Although we cannot distinguish between condensational and aqueous aerosol formation pathways, the pres-

- ⁵ ence of clouds and elevated relative humidity in the transition layer suggests a potential role for aqueous reactions. Using measurements of particulate oxalate as a tracer for aqueous processing, Wonaschuetz et al. (2012) and Sorooshian et al. (2007) have also observed evidence for secondary aerosol formation in the transition layer during cumulus convection over Texas and near the coast of California. Wonaschuetz et al. (2012)
- show no trends in the OA and particulate sulfate mass fractions with altitude in the mixed and transition layers, which could occur if the production was sufficiently small or if the additional aerosol mass in the transition layer were produced with the same ratio of OA and particulate sulfate that was originally present in the mixed layer. In contrast, our measurements show a distinct difference in composition between the mixed and transition layers and imply a similar magnitude of secondary sulfate and OA production
- in the transition layer, although the production of OA was not statistically significant. Goldstein et al. (2009) and Ford and Heald (2013) hypothesized a layer of aerosol that is primarily composed of SOA, that would be sufficient to explain a significant
- fraction of the observed summertime enhancement of AOD (2–3 times greater than winter), and that does not contribute to aerosol mass at the surface. Although we have observed an enhancement of aerosol in the transition layer, it is not consistent with the hypothesized layer in magnitude or composition. We observed enhancements that were less than 10% of AOD, and sulfate and OA were enhanced by similar magnitude although the OA enhancement was not statistically significant. The seasonality of the
- ²⁵ enhancement of surface aerosol mass (less than 1.6 times greater in summer than winter) compared to the AOD enhancement (2–3 times) was the primary evidence for the hypothesized layer. Given the absence of such a layer, our observations suggest that other factors such as meteorology and transport may influence the seasonality of the relationship of AOD to surface aerosol mass and warrant further investigation.



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Discussion

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Table 1. Measurements aboard the NOAA WP-3-D used in this analysis.

Measurement	Technique	Sample Interval	Lower Limit of De- tection	Accuracy	Reference
Sub-micrometer Aerosol Extinc- tion (dry)	Cavity Ringdown Spectrometer	1 s	0.1 Mm ⁻¹	5%	Langridge et al. (2011)
Sub-mircometer Aerosol Compo- sition	Compact – Time of Flight – Aerosol Mass Spectrometer	10s	OA < 0.8 μg m ⁻³ SO ₄ < 0.1 μg m ⁻³	35 %	Drewnick et al. (2005); Cana- garatna et al. (2007)
Sub-micrometer Aerosol Volume	Optical particle counter	1 s	0.03 µm ³ cm ⁻³	+26%, -12%	Cai et al. (2008)
CO	Vacuum UV fluorescence	1 s	0.5 ppbv	5%	Holloway et al. (2000)
Isoprene	PTR-MS	14 s	< 32 pptv	20 %	de Gouw and Warneke (2007)
RH (Dew Point)	Chilled Mirror Hygrometer	1 s	-	0.2°C	-
Black Carbon Mass	SP2	1s	12 ng m ⁻³	30 %	Schwarz et al. (2008)
SO ₂	Pulsed UV Fluorescence	1 s	250 pptv	20 %	Ryerson et al. (1998)
CH4	Cavity Ringdown Spectrometer	1 s		1.2 ppbv	Peischl et al. (2012)
CO ₂	Cavity Ringdown Spectrometer	1 s	-	0.15 ppmv	Peischl et al. (2012)

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Table 2. Measurements aboard the NASA DC-8 used in this analysis.

Measurement	Technique	Sample Interval	Lower Limit of De- tection	Accuracy	Reference
Sub-micrometer Aerosol Extinc- tion (dry)	Cavity Ringdown Spectrometer	1s	0.1 Mm ⁻¹	5%	Langridge et al. (2011)
Sub-mircometer Aerosol Compo- sition	High Resolution – Time of Flight – Aerosol Mass Spectrometer	1s	OA 0.6 μ g m ⁻³ SO ₄ 0.06 μ g m ⁻³ NO ₃ 0.06 μ g m ⁻³ NH ₄ 0.01 μ g m ⁻³	35 %	Canagaratna et al. (2007)
Sub-micrometer Aerosol Volume	Optical particle counter	1 s	0.03 µm ³ cm ⁻³	+26%, -12%	Cai et al. (2008)
CO	Infrared Absorption	1 s	0.5 ppbv	5%	Sachse et al. (1987)
Isoprene	PTR-MS	14 s	25 pptv	10 %	de Gouw and Warneke (2007)
RH (dew point)	Chilled Mirror Hygrometer	1 s	-	0.2°C	-
Black Carbon Mass	SP2	1 s	12 ng m ⁻³	30 %	Schwarz et al. (2008)
SO ₂	CIMS	1s	9 pptv	15%	S Kim et al. (2007)
CO2	Infrared Absorption	1 s	-	0.2 ppm	Vay et al. (2011)









Figure 2. (a) The number of profiles that contribute to the altitude-binned aggregate, (b) histograms of the altitude of the tops of the transition and mixed layers, and (c) the number of profiles that contribute to the altitude-normalized aggregate.



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Figure 3. Altitude-binned aggregate profiles of (a) the 532 nm dry aerosol extinction, (b) relative humidity, (c) aerosol hygroscopicity, and (d) the calculated ambient extinction. The shaded regions show the interdecile range (light) and the interquartile range (medium), and the solid lines are the median (dark). The dashed line in panel D shows the median dry extinction for comparison.





Figure 4. Comparison of airborne and surface measurements of the dry aerosol extinction coefficient. The airborne measurements are aboard the NOAA WP-3 aircraft. The surface measurements are from the SEARCH monitoring sites.





Figure 5. Aggregate profiles of **(a)** the aerosol mass and **(b)** the mass fractions of nitrate, ammonium, sulfate, and OA. The shaded regions show the interdecile range (light) and the interquartile range (medium), and the solid lines are the median (dark).





Figure 6. An example profile collected over central Georgia in the afternoon of 16 June showing **(a)** the temperature and virtual potential temperature, **(b)** the relative humidity and the water vapor mixing ratio, **(c)** the mixing ratio of CO that has a long lifetime, and **(d)** the mixing ratio of isoprene that has a short lifetime.





Figure 7. Altitude-normalized aggregate profiles of **(a)** CO, and **(b)** isoprene mixing ratios. The shaded regions show the interdecile range (light) and the interquartile range (medium), and the solid lines are the median (dark).





Figure 8. Altitude-normalized aggregate profile of **(a)** fraction of mixed layer air (Eq. 4), **(b)** CH_4 , **(c)**, CO_2 , **(d)** H_2O , and **(e)** black carbon aerosol mass. The dashed line shows the concentrations expected from mixing alone. The shaded regions show the interdecile range (light) and the interquartile range (medium), and the solid lines are the median (dark). These trace gases and black carbon aerosol mass are not expected to be enhanced or reduced in the transition layer. The agreement between the observations and the concentration expected from vertical mixing alone demonstrates that CO can be used to quantify the fraction air from the mixed layer.





Figure 9. Altitude-normalized aggregate profiles of aerosol mass **(a)**, extinction **(b)**, and volume **(c)**. The shaded regions show the interdecile range (light) and the interquartile range (medium), and the solid lines are the median (dark). The dashed line shows the median value expected from mixing alone. The difference between the observed median value and the median value expected from mixing alone indicates an enhancement of aerosol in the transition layer.





Figure 10. Altitude-normalized profiles of the aerosol composition: (a) OA, (b) SO_4 , (c) NH_4 , and (d) NO_3 . The shaded regions show the interdecile range (light) and the interquartile range (medium), and the solid lines are the median (dark). The dashed line shows the median expected concentration from vertical mixing alone.





Figure 11. Altitude-normalized profiles of (a) particulate sulfate, (b) gas phase SO₂, and (c) total sulfur tS. The shaded regions show the interdecile range (light) and the interquartile range (medium), and the solid lines are the median (dark). The dashed line shows the median value expected from mixing alone. Histograms of the transition layer enhancement (E(h)) and the results of the *T* test for (d) particulate sulfate, (e) SO₂, and (f) total sulfur are shown.





Figure 12. Altitude-normalized profiles of (a) particulate sulfate, (b) gas phase SO₂, and (c) total sulfur tS. The shaded regions show the interdecile range (light) and the interquartile range (medium), and the solid lines are the median (dark). The dashed line shows the median value expected from mixing alone. Histograms of the transition layer enhancement (E(h)) and the results of the T test for (e) particulate sulfate, (f) SO_2 , and (g) total sulfur are shown.



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Figure 13. The AOD contributions of dry aerosol, aerosol water, and enhanced extinction in the transition layer are illustrated in an idealized profile. The subpanels show **(a)** histograms of AOD calculated from individual profiles (solid) and AOD measured with a sun photometer (dashed) at the Centreville SEARCH site, and the contributions to AOD from the **(b)** transition layer enhancement of extinction, **(c)** aerosol water, **(d)** the transition layer, and **(e)** the mixed layer. The calculated AOD assumes no contribution from aerosol above the top of the profile and extrapolates the dry extinction and RH to the surface.

