Temporal heterogeneity in aerosol characteristics and the resulting radiative impact at a tropical coastal station – Part 1: Microphysical and optical properties

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Abstract. In Part 1 of this two-part paper, we present the results of extensive and collocated measurements of the columnar and near-surface (in the well mixed region) properties of atmospheric aerosol particles at a tropical coastal location, Trivandrum (8.55°N; 76.97°E), located close to the southwest tip of Indian peninsula. These are used to evolve average, climatological pictures of the optical and microphysical properties and to delineate the distinct changes associated with the contrasting monsoon seasons as well as the transition from one season to the other. Our observations show a dramatic change in the columnar aerosol optical depth (AOD) spectra, being steep during winter monsoon season (WMS, months of December through March) and becoming quite flat during summer monsoon season (SMS, June through September). The derived Ångström exponent \( \alpha \) decreases from a mean value of 1.1±0.03 in WMS to 0.32±0.02 in SMS, signifying a change in columnar aerosol size spectrum from an accumulation mode dominance in WMS to a coarse mode dominance in SMS. The composite aerosols near the surface follow suit with the share of the accumulation mode to the total mass concentration decreasing from \( \sim 70\% \) to 34\% from WMS to SMS. The overall mass burden also decreases in tandem. The changes in \( \alpha \) are well correlated to those in the accumulation fraction of the mass concentration. Long-term measurements of the concentration of aerosol black carbon (BC), show prominent annual variations, with its mean value decreasing from as high as 6 \( \mu g m^{-3} \) in WMS to 2 \( \mu g m^{-3} \) in SMS. Correspondingly, its mass mixing ratio to the composite aerosols comes down from 11\% to 4\%. The changes in AOD and \( \alpha \) are significantly positively correlated to those of BC concentration. The columnar properties are, in general well associated with the features near the surface. The implications of these changes to the optical properties and single scattering albedo and the resulting impact on direct radiative forcing are examined in the companion paper (Part 2).

Keywords. Atmospheric composition and structure (Aerosols and particles; Biosphere-atmosphere interactions; Transmission and scattering of radiation)

1 Introduction

Microphysical and optical properties of atmospheric particles (such as aerosol optical depth (AOD), size distribution, single scattering albedo (SSA), phase function and so on) are important in determining their direct radiative forcing or ability to modify the Earth’s radiation budget (e.g., Charlson et al., 1992). Aerosols (used throughout the paper to mean the particulate phase of atmospheric aerosol system) influence the radiative forcing directly by scattering and absorbing the solar radiation (Charlson et al., 1992) and indirectly by altering cloud optical properties, albedo, and lifetime (Twomey, 1977; Lohmann and Feichter, 2005). Aerosol forcing estimates still remain highly uncertain (IPCC, 2001) because of the large variability in their properties (which arises mainly from patchy sources, sinks, and short residence time in the troposphere where they are in abundance), that are not characterized with adequate accuracy (spatially and temporally), despite the concerted efforts in the last few decades (Charlson et al., 1992; Haywood et al., 1997; Satheesh et al., 1999; Russell et al., 1999; Babu et al., 2002; Huebert et al., 2003; Moorthy et al., 2005a). This calls for detailed understanding of how the microphysical properties of these particles undergo systematic changes during the course of a year, over and above the more random variations at considerably shorter time scales. Such systematic changes can be brought about by the changes in the airmass types and prevailing meteorology (e.g., Smirnov et al., 1994; Moorthy et al., 2003a). This calls for long-term and collocated measurements of several,
if not all, aerosol parameters that are important for aerosol radiative forcing estimates. These essentially include spectral AODs, size distribution, SSA, and scattering phase function ($P(\theta)$). Such assessments will provide more realistic inputs to climate modelers.

With this intention, a systematic, all season, investigation using a long database, which will smooth out short-term perturbations, has been undertaken at Trivandrum (8.55° N, 76.9° E, 3 m a.m.s.l. (above mean sea level)), a sparsely industrialized, semi-urban, tropical coastal location at the southwest tip of Indian peninsula. The coastal nature of this station has implications from several standpoints. India has a long coastline extending over 7500 km. More than 50% of its over one-billion population resides in the coastal areas and the resulting anthropogenic activities will contribute significantly to the aerosol loading, which gets modified by the natural processes associated with the monsoons. Sea spray aerosols are produced only at the sea surface, while the primary production of rural, and most of the anthropogenic aerosols take place over the land and these two types have distinct physical and radiative properties. On either side of the coastline, these two types of aerosols are mixed and are supplemented by secondary production (in presence of the high humidity). In addition, the mesoscale processes such as land/sea breezes lead to a thorough vertical and horizontal mixing of these aerosols within the atmospheric boundary layer with consequent changes in their microphysical properties. Above all, the large-scale changes in the atmospheric circulation system and contrasting rainfall pattern occurring at this location associated with the Indian monsoon, as it changes from the northeast winter monsoon season (WMS, months of December through March) to the southwest summer monsoon season (SMS, June through September), will result in contrasting airmass types and significant changes in the aerosol properties. The change over from one to the other is gradual and during this period (inter-monsoon season, (IMS, April and May)) and post monsoon season (PMS, October and November)) the aerosol properties will be more complex.

In Part 1 of this two-part paper, we examine the microphysical properties of aerosols based on the long-term and collocated measurements over a period of ~4 years. These data are used to build-up a climatological model and to delineate the seasonal distinctiveness and the temporal heterogeneity. The impact of these changes in direct radiative forcing will be examined in Part 2 of this paper.

2 Experimental setup and database

Collocated measurements of spectral AOD (at 10 wavelengths from UV (Ultraviolet) through near IR (Infrared)), total mass concentration ($M_T$) and mass size distribution [$m(r)$] of composite aerosols in the well-mixed region near the surface, and mass concentration ($M_B$) of aerosol black carbon (BC) were carried out from Trivandrum (TVM). The actual study site was in a fishing village, Thumba (8.55° N, 76.97° E, 3 m a.m.s.l.) and located within the Thumba Equatorial Rocket Launching Station. The site is characterized by a flat coastal terrain, free from proximity to any major industrial and urban activities. The experimental site was ~500 m due east of the Arabian Sea coast and ~10 km northwest of the urban area of TVM (Fig. 1). Being within the rocket launching station, human activity is highly subdued over an extent of ~±3 km along the coast and about 1 km across it centered about the measurement site. The instruments used for this investigation are listed in Table 1, along with references to earlier publications on the instrument details, data analysis techniques, and error budget, which are not extensively described in this paper. The measurements spanned over 4 years from January 2000 to December 2003. Supplementary data on meteorological parameters such as wind speed, direction, temperature, and relative humidity (RH) were obtained from instruments mounted on a meteorological tower (~17 m above ground level), located near the meteorological facility of the rocket launching station, ~600 m away. Daily total rainfall was also measured at the same meteorological facility.

The MWR (Multi-Wavelength Radiometer), mounted on the rooftop of the building (~4 m above ground level), was used to estimate spectral AODs, on days when unobstructed solar visibility was available for 3 h or more, following the Langley plot technique (Shaw et al., 1973) and subtracting the contribution due to molecular scattering and absorption from the total atmospheric optical depth. Application of this technique to the MWR data analysis is described in several earlier papers (Satheesh and Moorthy, 1997; Moorthy et al., 1999, 2001; Saha and Moorthy, 2004). In most cases the entire measurements made, with the MWR, over a day is considered as a single set and the mean AOD is deduced...
following the Langley plot method. However, on some days the Langley plots showed distinctly different slopes for the forenoon (FN) and afternoon (AN) parts of the same day and on such days these were considered as two independent data sets and the mean optical depths are deduced for each set (following Langley technique) separately for FN and AN. The temporal stability (statistical) of the Langley intercept was used to ensure the stability of the instrument. The typical uncertainty in the AOD estimates varied from 0.02 to 0.03 at different wavelengths, the higher values occurring at the shorter wavelengths.

For the Quartz Crystal Microbalance (QCM, which is a 10 channel cascade impactor) and the Aethalometer, ambient air was drawn from a height of 4 m above the ground level. The measurements of total mass concentration \( M_T \) and mass size distribution \([m(r)]\) of the composite aerosols (in the size range 0.05 to 25 \( \mu \)m) were made using QCM. Generally, the QCM measurements were made on 4 to 5 days a month and on each day the sampling was done at (nearly) hourly intervals, with the sampling duration varying between 3 to 5 min, depending on the aerosol loading (so that all the channels registered a minimum frequency shift of greater than 2 Hz, Pillai and Moorthy, 2001). Details of the sampling and typical error budget are given in earlier papers (Pillai and Moorthy, 2001; Pillai et al., 2002; Moorthy et al., 2003a). Typical errors in the QCM-estimated total mass concentrations were in the range 15 to 20%; the error tending to the higher side for lower \( M_T \). Due to the affinity of quartz to water, the QCM measurements are not reliable when relative humidity (RH) exceeds 75% and as such, the instrument was operated only during periods when the ambient RH < 75%.

Measurements of the mass concentrations \( M_B \) of atmospheric BC particles were made using an aethalometer (Model AE-21 of Magee Scientific, USA), operated continuously at a flow rate of 31 min\(^{-1}\) (LPM) and a timebase of 5 min; so that \( M_B \) estimates are available at every 5 min, round the clock. The aethalometer estimates \( M_B \) by measuring the change in the transmittance of its quartz fiber filter tape due to deposition of particles aspirated from the ambient. The aethalometer is a simple and rugged instrument, suitable for field experiments of long duration, and provided data almost without break. However, several reports are available in the recent literature on uncertainties in the aethalometer estimated BC (for e.g., Weingartner et al., 2003; Arnott et al., 2005; Sheridan et al., 2005; Corrigan et al., 2005). Most of these studies have shown that there exists no unique method for accurate, field measurement of the mass concentration of BC, and that all the available methods are species-specific and instrument-specific (Hitzenberger et al., 2006). Several experiments were carried out to correct the aethalometer data for its inherent uncertainties and also to compare the estimated values of BC with those from other collocated measurements (Weingartner et al., 2003; Arnott et al., 2005; Hitzenberger et al., 2006). Measurements of BC mass concentration by aethalometry have two known impediments. The first one is the amplification factor arising due to multiple scattering in the quartz fiber matrix of the tape and is normally called the C-factor (Weingartner et al., 2003). The second one arises from the deposition of scattering material along with BC on to the filter tape producing the "shadowing effect" (the R-factor) (Weingartner et al., 2003; Arnott et al., 2005). Other uncertainties due to rapid changes in

Table 1. Instrument details and parameters obtained.

<table>
<thead>
<tr>
<th>Sl. No.</th>
<th>Instrument and reference</th>
<th>Parameter measured/estimated</th>
<th>Conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10 channel multi-wavelength solar radiometer, MWR (Moorthy et al., 1997, 1999, 2001; Saha and Moorthy, 2004)</td>
<td>Spectral AOD (by Langley plot method) at 380, 400, 450, 500, 600, 650, 750, 850, 935 &amp; 1025 nm.</td>
<td>Clear/Cloud free days or partly cloudy days when the part of the sky ( \geq 5^\circ ) with sun at the center was free of any visual clouds.</td>
</tr>
<tr>
<td>2</td>
<td>10-channel Quartz Crystal Microbalance (QCM) impactor: (type PC-2 of California measurements, USA) (Pillai and Moorthy, 2001; Moorthy et al., 2003a)</td>
<td>( M_T ) and ([m(r)]) of ambient aerosols in the size range 0.05 to 25 ( \mu )m, at near real time.</td>
<td>Manually operated and limited to periods when RH &lt; 75%.</td>
</tr>
<tr>
<td>3</td>
<td>Aethalometer (type AE-21 of Magee Scientific, USA) (Babu and Moorthy, 2002; Babu et al., 2004)</td>
<td>Near-real time estimates of ( M_B ) at every 5 min.</td>
<td>Automatic and round the clock</td>
</tr>
</tbody>
</table>
sampling conditions and those due to instrument noise can be considerably reduced by using long averaging periods as in the present case. By applying an empirically derived correction routine based on laboratory and field absorption measurements made in conjunction with a photo acoustic photometer, Arnott et al. (2005) have developed a correction procedure for aethalometer which was used by Corrigan et al. (2006) for aethalometer observations from MCOH (Maldives Climate Observatory at Hanimadhoo) and found that uncertainty in the retrieved absorption coefficients ranges between 5% to 40% depending on the BC mass-load. Intercomparing the aethalometer with other different techniques, Hitzenberger et al. (2006) have shown that even though the instantaneous values are somewhat species-specific (nature of the BC particles) and instrument-specific, the average concentrations inferred by various techniques tend to agree with in the instrumental uncertainties for aged (away from strong sources) aerosols. In our studies, there were no prominent sources of BC anywhere near the sampling site and also we have used very long (monthly and seasonal) averages. Such averaging also helped to reduce the uncertainties due to relative humidity and instrument noise. As detailed in Nair et al. (2007), in our measurements we have used a correction factor of 1.9, which is close to the factor of 2.14±0.21 suggested by Weingartner et al. (2003) for aged, rural aerosols. Considering all the above we attribute an uncertainty of 20% to the long-term averaged (monthly) BC used in this study, but acknowledge that this systematic uncertainty might be higher.

3 Prevailing meteorology

Being a tropical, coastal location, situated close to the geographic equator, seasonal changes are not as perceptible at Trivandrum as at the temperate latitudes. The most important synoptic meteorological process relevant to this location is the extensive and widespread rainfall during the Indian summer monsoon and the associated changes in the general circulation pattern (which almost reverses from a weak north-easterly in winter to a strong southwesterly in summer) and the large north south excursion of the Inter Tropical Convergence Zone (ITCZ) (Das, 1986; Asnani, 1993). This is modulated by the mesoscale land/sea breezes arising out of the coastal proximity (Narayanan, 1967), which is perceptible during November to March. While the general circulation is a synoptic scale activity extending from June to October, the land/sea breeze is a mesoscale process occurring almost daily but confined within the boundary layer (Moorthy et al., 1993). Compared to these mesoscale land/sea breezes, the synoptic changes associated with the monsoon activity would produce more significant changes in the aerosol characteristics that would prevail for longer duration. During December to March, the prevailing atmospheric circulation in the lower troposphere is primarily from the inland continental regions directed towards the ocean, constituting a continental airmass type. This flow is generally dry because these winds arrive from the dry continental interiors and will be richer in continental, accumulation mode aerosols, which would have a significant anthropogenic component too. In sharp contrast to this, during the period from June to September, the prevailing winds are directed from the ocean (Arabian Sea). These winds are quite strong (>10 m s\(^{-1}\)) and have a long track over the sea and thus bring in a lot of moisture. This marine airmass will also be richer in marine (sea spray) aerosols that have a different size spectrum and chemical characteristics (with a dominant coarse mode associated with sea salt particles) and are of natural origin. During this season, the skies are generally cloudy and rainfall is strong and widespread resulting in a significant wet removal of aerosols (Moorthy et al., 1991). The annual variation of monthly total rainfall at TVM (averaged over the study period) is shown in the top panel of Fig. 2 followed by monthly mean values of RH, as well as maximum and minimum temperatures respectively in the panels below.

Based on the above, as well as the climatological pattern of synoptic winds (for e.g., Asnani, 1993; Pillai and Moorthy, 2001) the following four distinct seasons are considered for aerosol characterization, namely the winter monsoon
Figure 3. Polar graphs showing wind speed and wind direction for (a) Winter Monsoon Season (WMS; months of December through March) (b) Inter Monsoon Season (IMS; April and May), (c) Summer Monsoon Season (SMS; June through September) and (d) Post Monsoon Season (PMS; October and November). Wind direction refers to the direction of wind arrival. The percentage indicates the percentage of points located in a given quadrant. The clear transformation from WMS to SMS is easily discernible.

Fig. 3. Polar graphs showing wind speed and wind direction for (a) Winter Monsoon Season (WMS; months of December through March) (b) Inter Monsoon Season (IMS; April and May), (c) Summer Monsoon Season (SMS; June through September) and (d) Post Monsoon Season (PMS; October and November). Wind direction refers to the direction of wind arrival. The percentage indicates the percentage of points located in a given quadrant. The clear transformation from WMS to SMS is easily discernible.

3.1 Winter Monsoon Season (WMS)

This is climatologically, the driest season of the year for TVM (as well as the northern hemispheric regions in Indian longitudes), with very little or insignificant rainfall (~5% of the annual). The RH generally remains between 40% and 60% signifying the prevalence of a dry environment despite the coastal proximity. The interiors are still drier with RH dropping to as low as 15% in the central peninsula (Moorthy et al., 2005b). The minimum temperatures are, in general, the lowest observed during the year (in the range 18 to 22°C) and the diurnal variation in temperature ($\Delta T$ = the difference between maximum and minimum temperature) exceeds 15°C on an average, and go as high as 17°C, at times. The prevailing winds (Fig. 3a) are generally weak ($<\sim 6$ m s$^{-1}$).
northerly or northeasterlies (for >70% of the time) signifying an airmass directed from the interior continental India, which is drier. During this season, ITCZ is located climatologically at ∼5° S or beyond (Waliser and Gautier, 1993). Examination of the winds and OLR (Outgoing Longwave Radiation) from the NCEP (National Centers for Environmental Prediction) data also revealed that the ITCZ was between 5° S to 10° S during these months in all the years considered. This season, thus, is considered as the WMS, even though it is much warmer and humid at TVM than the conventional winter experienced at the northern parts of India. The anticyclonic circulation over the continent is conducive for a vertical confinement of aerosols. The sky is mostly clear and cloud free and rainfall is scanty.

3.2 Inter-Monsoon Season (IMS)

The WMS gradually gives way to warmer and more humid IMS extending from April to May during which the winds undergo a transition. The anticyclonic subsidence still prevails and the skies are clear to partly cloudy; but the winds shift more to northerly and northwesterly direction (for >50% of the time) with moderately high speeds ranging from 5 to 15 m s⁻¹ (Fig. 3b). As such, the airmass type is a mixture, transforming from the continental type to the marine type. The ITCZ migrates gradually to the north (Asnani, 1993) and is located between equator and 5° N. The mean land temperature increases and the diurnal variation decreases (∆T~12 to 13°C). Ambient RH is higher and has a positive gradient (at ~50 to 70%). The rainfall increases during this period. The total rainfall during these two months is ~16% of the annual.

3.3 Summer Monsoon Season (SMS)

The onset of Indian summer monsoon occurs over TVM climatologically on June 1, accompanied with the change in winds to northwesterly; directed from the sea to land (for >97% of the period, Fig. 3c). This marine airmass brings with it vast amount of moisture, cloudiness, and precipitation. The wind speeds are high, with mean value of ~12.3±4.7 m s⁻¹ and peak speeds reaching as high as ~25 m s⁻¹. The maximum temperature decreases and ∆T lies in the range 9 to 11°C at times dropping down to 4°C. The RH remains high (and exceeds 80% quite frequently). This season, extending from June to September, provides much of the rainfall (~47% of the annual). During this season ITCZ is located beyond 10° N. The rainfall gradually spreads northward from June, covering entire India by beginning of July.

3.4 Post-Monsoon Season (PMS)

During this season (October and November), the winds start shifting again to northerly or northeasterly (47% of time) with moderate speeds ranging from 5 to 15 m s⁻¹ (Fig. 3d). The extent of cloudiness decreases. The temperature starts increasing and ∆T lies in the range 10 to 12°C and RH decreases. However, rainfall still continues to be significant (~32% of the annual). (In fact the mean rainfall was highest in the month of October during the study period.) This post monsoon season is also a transition season; but it has more intense rainfall associated with it compared to IMS. This season is also characterized by weakening of low-pressure center over the sub-continent. At the same time, the low-pressure center is prominent over the Bay of Bengal making it a seat for generation of depression and cyclones, which bring-in the rainfall over these parts. The ITCZ starts migrating to the south.

4 Results and discussion

In the following we examine how the optical and microphysical properties of aerosols respond distinctly to these large-scale natural processes; which are seasonal and repeat the seasonality from year to year.

4.1 Spectral Aerosol Optical Depth

Spectral AOD is a key parameter in estimating the extent to which aerosols directly influence the radiative balance of the Earth-Atmosphere system and is dependent on the columnar abundance, size distribution and refractive index of aerosols. Spectral AODs, estimated from each MWR data set are examined for the temporal and spectral characteristics.

4.1.1 Annual variation of AOD

The spectral AOD showed significant annual variations, with distinct spectral dependence. Despite, the nature of the variation was similar in all the years, indicating a regular pattern. In view of this, the individual AOD estimates were grouped in to the calendar months, without discriminating them in terms of the year and the monthly mean values and their standard deviations are estimated. This would smooth out the inter-annual variations and the short scale fluctuations and will bring out a climatological mean pattern amenable for inputting into radiation models. This smoothed annual variation is shown in Fig. 4, at four representative wavelengths (380, 500, 750 and 1025 nm) spanning over the spectral range covered by the MWR. Here each point corresponds to the monthly mean AOD, irrespective of year, and the vertical line through each point represents the standard error. When the mean values are to be compared, it is customary to use the standard error, which represents the range over which the mean value would vary, if it were estimated from a number of different ensembles drawn from a population; smaller the error, the more closer is the ensemble mean to the population mean. On the other hand the standard deviation is used to comment on the spread of values with in an ensemble. The shorter lengths of the error bars during WMS indicate smaller
Table 2. Mean AOD at 380, 500, 750 and 1025 nm for different seasons.

<table>
<thead>
<tr>
<th>Season</th>
<th>AOD_{380}</th>
<th>AOD_{500}</th>
<th>AOD_{750}</th>
<th>AOD_{1025}</th>
<th>Number of data points</th>
</tr>
</thead>
<tbody>
<tr>
<td>WMS (Dec, Jan, Feb and March)</td>
<td>0.58±0.02</td>
<td>0.43±0.01</td>
<td>0.27±0.01</td>
<td>0.21±0.01</td>
<td>282</td>
</tr>
<tr>
<td>IMS (April and May)</td>
<td>0.53±0.04</td>
<td>0.40±0.03</td>
<td>0.28±0.02</td>
<td>0.25±0.01</td>
<td>61</td>
</tr>
<tr>
<td>SMS (June, July, Aug and Sep)</td>
<td>0.35±0.03</td>
<td>0.29±0.03</td>
<td>0.29±0.03</td>
<td>0.27±0.02</td>
<td>41</td>
</tr>
<tr>
<td>PMS (Oct and Nov)</td>
<td>0.51±0.06</td>
<td>0.38±0.04</td>
<td>0.23±0.03</td>
<td>0.15±0.02</td>
<td>19</td>
</tr>
</tbody>
</table>

day-to-day and year-to-year variabilities of AOD during this season compared to SMS. Main features are

1. In the visible (short) wavelengths, the AODs are generally high during January, February months. Subsequently they increase gradually to reach a peak in April and dropdown to the low values in May. Thereafter, the AODs remain nearly steady at the low values until October before recovering.

2. In contrast to this, the annual variations are weaker in the NIR wavelength regime (compared to those at the shorter (visible) wavelength regime (such as 380 to 500 nm)). While the AODs at the shorter wavelength show a sharp decrease in May and remain low until October, at the NIR wavelengths the AODs continue to be moderately high. In other words, the AOD spectrum changes its shape from winter to summer.

The seasonal mean values of AOD (obtaining by grouping them into the four seasons considered here) at these four wavelengths are given in Table 2. The contrasting differences in aerosol characteristics as the season changes are seen from the changes in the magnitude and the wavelength variation of the AOD.

4.1.2 Spectral variations and Ångström parameters

The AOD spectra contain information on the columnar size distribution of aerosols as they are related through

\[ \tau_p(\lambda) = \int_{r_1}^{r_2} \pi r^2 Q_{\text{ext}}(r, \lambda, m) n_c(r) \, dr \] (1)

where \( \tau_p(\lambda) \) is the aerosol optical depth, \( Q_{\text{ext}} \) is the Mie extinction efficiency parameter (which is function of particle size \( r \), refractive index \( m \) and wavelength \( \lambda ) \), \( n_c(r) \) is the columnar size distribution, and \( r_1 \) and \( r_2 \) are the lower and upper radii limits. \( Q_{\text{ext}} \) increases from a low value for small values of \( r/\lambda \) to reach a peak value (~4) for \( r \sim \lambda \) and then approaches 2 for higher values of \( r \) (McCartney, 1976). Thus maximum contribution to \( \tau_p \) comes from particles in a size distribution whose radii are comparable to the wavelength of the radiation. Higher values of AOD at shorter wavelength, thus, imply higher relative abundance of smaller particles in the aerosol size spectrum (i.e. compared to the case of lower values of AOD at the same wavelengths). The simplest way to quantify these spectral changes in AOD is to estimate the Ångström parameters (Ångström, 1964) using the equation

\[ \tau_p(\lambda) = \beta \lambda^{-\alpha} \] (2)

The Ångström wavelength exponent (\( \alpha \)) is an indicator of the size distribution, being higher when the columnar aerosol size spectrum has higher relative dominance of accumulation mode aerosols (size < 1 \( \mu \)m), while \( \beta \) is a measure of the aerosol loading. The simplicity of the expression enables its use in radiative forcing calculation. Following Eq. (2), \( \alpha \) and
Table 3. Ångström parameters over various Indian locations as well as marine locations around India along with that observed during the present study.

<table>
<thead>
<tr>
<th>Seasons</th>
<th>Location</th>
<th>( \alpha )</th>
<th>( \beta )</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>WMS</td>
<td>Kanpur (26.43° N, 80.36° E)</td>
<td>1.26±0.25</td>
<td></td>
<td>Singh et al. (2004)</td>
</tr>
<tr>
<td></td>
<td>Dibrugarh (27.3° N, 94.5° E)</td>
<td>~1.3</td>
<td></td>
<td>Bhuyan et al. (2005)</td>
</tr>
<tr>
<td></td>
<td>Vishakapatnam</td>
<td>1.08±0.03</td>
<td>0.19</td>
<td>Ramachandran (2004)</td>
</tr>
<tr>
<td></td>
<td>Trivandrum</td>
<td>1.0±0.08</td>
<td>0.17</td>
<td>Ramachandran, 2004</td>
</tr>
<tr>
<td></td>
<td>Port Blair (11.6° N, 92.7° E)</td>
<td>1.24</td>
<td>0.14</td>
<td>Moorthy et al. (2003b)</td>
</tr>
<tr>
<td></td>
<td>Kaashidhoo (5° N, 73.5° E)</td>
<td>1.23±0.21</td>
<td>0.09</td>
<td>Satheesh et al. (1999)</td>
</tr>
<tr>
<td></td>
<td>Arabian Sea – coastal</td>
<td>0.97±0.23</td>
<td>0.18</td>
<td>Satheesh and Moorthy (1997)</td>
</tr>
<tr>
<td></td>
<td>– far (&gt;800 km)</td>
<td>0.50±0.30</td>
<td>0.19</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Arabian Sea (mean)</td>
<td>0.84±0.38</td>
<td>0.26</td>
<td>Satheesh et al. (2001)</td>
</tr>
<tr>
<td></td>
<td>Coastal India</td>
<td>1.49±0.10</td>
<td>0.14</td>
<td>Ramachandran (2004)</td>
</tr>
<tr>
<td></td>
<td>Arabian Sea</td>
<td>1.50±0.05</td>
<td>0.10</td>
<td>Ramachandran (2004)</td>
</tr>
<tr>
<td></td>
<td>Tropical Indian Ocean</td>
<td>1.52±0.11</td>
<td>0.05</td>
<td>Ramachandran (2004)</td>
</tr>
<tr>
<td></td>
<td>Bay of Bengal</td>
<td>1.23±0.43</td>
<td>0.20</td>
<td>Satheesh et al. (2001)</td>
</tr>
<tr>
<td></td>
<td>Far Indian Ocean</td>
<td>0.61±0.01</td>
<td>0.09</td>
<td>Moorthy et al. (2001)</td>
</tr>
<tr>
<td></td>
<td>Indian Ocean</td>
<td>0.50±0.19</td>
<td>0.24</td>
<td>Satheesh et al. (2001)</td>
</tr>
<tr>
<td></td>
<td>TVM</td>
<td>1.10±0.03</td>
<td>0.20</td>
<td>Present study</td>
</tr>
<tr>
<td>IMS</td>
<td>Bangalore (13° N, 77° E)</td>
<td>1.30±0.01</td>
<td>0.13</td>
<td>Moorthy et al. (2005a)</td>
</tr>
<tr>
<td></td>
<td>Kanpur (26.43° N, 80.36° E)</td>
<td>0.60±0.31</td>
<td></td>
<td>Singh et al. (2004)</td>
</tr>
<tr>
<td></td>
<td>Arabian Sea</td>
<td>1.17±0.03</td>
<td>0.21</td>
<td>Moorthy et al. (2005a)</td>
</tr>
<tr>
<td></td>
<td>TVM</td>
<td>0.85±0.04</td>
<td>0.22</td>
<td>Present study</td>
</tr>
<tr>
<td>SMS</td>
<td>Kanpur (26.43° N, 80.36° E)</td>
<td>0.66±0.45</td>
<td></td>
<td>Singh et al. (2004)</td>
</tr>
<tr>
<td></td>
<td>Dibrugarh (27.3° N, 94.5° E)</td>
<td>~0.6</td>
<td></td>
<td>Bhuyan et al. (2005)</td>
</tr>
<tr>
<td></td>
<td>Arabian Sea</td>
<td>0.35±0.12</td>
<td>0.36</td>
<td>Vinoj and Satheesh (2003)</td>
</tr>
<tr>
<td></td>
<td>TVM</td>
<td>0.32±0.02</td>
<td>0.25</td>
<td>Present study</td>
</tr>
<tr>
<td>PMS</td>
<td>Bangalore (13° N, 77° E)</td>
<td>1.06±0.02</td>
<td>0.12</td>
<td>Babu et al. (2002)</td>
</tr>
<tr>
<td></td>
<td>Kanpur (26.43° N, 80.36° E)</td>
<td>1.12±0.28</td>
<td></td>
<td>Singh et al. (2004)</td>
</tr>
<tr>
<td></td>
<td>TVM</td>
<td>1.20±0.01</td>
<td>0.16</td>
<td>Present study</td>
</tr>
</tbody>
</table>

\( \beta \) are derived from the individual AOD spectra (from January 2000 to December 2003) by least squares fitting the AOD spectra, in a log-log scale. For this we have used AODs at all the 10 wavelengths (in the MWR).

Climatologically monthly mean values of \( \alpha \) and \( \beta \) are then obtained by grouping the individual estimates into monthly ensembles and averaging. The annual variations of mean \( \alpha \) and \( \beta \) are shown in Fig. 5. It very clearly shows the dramatic transformation in the aerosol spectra, from an accumulation-mode domination during WMS, (revealed by high values of \( \alpha \) (>1) and steep AOD spectra) to an increased coarse mode domination in the SMS (characterized by a low value of \( \alpha \) (<0.5) and flat AOD spectra). Interestingly, the highest value of \( \alpha \) (~1.3) occurred in the month of November (PMS). However, \( \beta \) values did not show large changes, even though it tended to be higher during SMS, and lower during PMS. The seasonal transformations of the AOD spectra are shown in Fig. 6, where the individual points are the AODs averaged over the season, irrespective of the year. Performing the regression analysis on the seasonal mean AOD spectrum, seasonal mean Ångström parameters are evaluated and are given in Table 3 along with the values reported, in the recent years, over Indian landmass and the adjoining oceans by several investigators along with relevant references. From Table 3 it is evident that the high values of \( \alpha \) (>1.0) are a consistent feature over most of the Indian landmass during WMS; of course, local variations occur depending on region-specific industrial/urban activities or coastal or arid nature of the sites. In contrast, the low values of \( \alpha \) in SMS is analogous to that seen over far oceanic environments in winter or over the extensively dust infested region of northern India (Kanpur); in both the cases there is an increased influence of natural, coarse mode, aerosols.

Extensive measurements over different environments have shown that the high values of \( \alpha \) are typically associated with continental aerosols (Hess et al., 1998) or over oceans adjacent to continents and influenced by a continental airmass (Hoppel et al., 1990; Satheesh and Moorthy, 1997; Moorthy et al., 2005a), whereas over remote marine environments, far away from continents and under the influence of a marine
airmass, the AOD spectra tend to become flat (Hoppel et al., 1990; Smirnov et al., 2002; Moorthy et al., 2001). Examining our results in the light of the above, it is clear that the columnar aerosol properties over TVM closely resemble that of a typical marine environment during the SMS, while it behaves almost like a polluted and anthropogenically influenced continental site during WMS, with the synoptic transition periods (IMS and PMS) coming in between. We now examine the features of the composite aerosols near the surface, based on the QCM data, in the above perspective.

4.2 Composite aerosol characteristics

The QCM provided near-real-time measurements of size segregated mass concentration as well as total mass concentration \( M_T = \sum_{i=1}^{10} m_{ci} \) where \( m_{ci} \) is the mass concentration in the \( i \)th size-bin. Grouping the individual \( M_T \) estimates into monthly ensembles (irrespective of year) and averaging, the monthly mean (climatological) values of \( M_T \) were estimated. The annual variation of these climatological monthly mean \( M_T \) values (shown in the bottom panel of Fig. 7) closely resembles that of AOD at the shorter wavelengths (shown in the bottom two panels of Fig. 4). High values \( (M_T \geq 60 \mu g m^{-3}) \) occur during the WMS; with \( M_T \) gradually increasing from January to reach as high as \( 70 \mu g m^{-3} \) by March. Subsequently, \( M_T \) decreases rapidly during April–May (IMS) to reach a low value by June. During the entire SMS (June to September) it remains at the low level \( (35–40 \mu g m^{-3}) \) and increases thereafter through the PMS (October–November) to reach high values \( (\sim 60 \mu g m^{-3}) \) again by December.

The dry land condition, low relative humidity, large diurnal variation in the ambient temperature, the rather cloud free skies, and the continental nature of the prevailing airmass are all highly conducive for the high aerosol loading during WMS. The continental nature of the airmass during this period is favorable for the transport of aerosols from drier and densely populated continental regions lying upwind to the site, where the accumulation mode aerosols are known to dominate in winter (Moorthy et al., 2005b). The scanty rainfall \( (\sim 5\% \text{ of annual and even less upwind}) \) and very low cloud cover during this period renders scavenging process to be very weak. Consequently, the aerosols have their longest atmospheric residence time during this season; conducive for long-distance transport (Ramanathan et al., 2001). In contrast, the extensive cloudiness and rainfall during SMS results in an effective wet scavenging of the particles, resulting in a rapid decrease in their concentration. It may be recalled that for aerosols in the size range 0.1 to 1.0 \( \mu m \), wet removal is the chief removal mechanism (Flossmann et al., 1985; Saha and Moorthy, 2004). A part of this loss is replenished by fresh marine aerosol particles, produced by the strong monsoon winds over the ocean, and advected over to the coastal landmass. Moreover, the land remains wet, the sky is generally cloudy and diurnal variation in temperatures is the lowest in the year during this season, all being unfavorable for local production of continental aerosols/dust. Thus the composite aerosols undergo significant seasonal changes in their characteristics.

Further, at the coastal location there would be at least two strong aerosol components; one of continental origin and the other of marine origin; each dominating in distinct size regime and during different parts of the year. The continental component would be richer in accumulation mode aerosols (e.g., Moorthy et al., 2003a) while the marine component will have a strong coarse mode (radius > 1.0 \( \mu m \)). As such, the total aerosol mass concentration \( (M_T) \) is separated into \( M_A \) (mass concentration in the accumulation mode, radius from 0.05 to 1 \( \mu m \)) and \( M_C \) (mass concentration in the coarse mode, > 1 \( \mu m \)) such that \( M_T=M_A+M_C \). The \( M_A \) and \( M_C \), estimated from each QCM samples are averaged to obtain the mean representative values for each month similar to \( M_T \). The month-to-month variation of \( M_A \) and \( M_C \) are shown in the middle panels of Fig. 7. There is a very close similarity in the annual variations of \( M_A \) and \( M_T \), which indicates that \( M_A \) is a significant component of \( M_T \) in all seasons. On the other hand, variation of \( M_C \) is nearly opposite in nature to that of \( M_A \). The above changes are categorized in terms of the 4 seasons considered in this study and the seasonal mean values of \( M_T, M_A \) and the share of \( M_A \) to \( M_T \) are given in Table 4. It very clearly shows that the size spectrum of composite aerosols undergoes a transformation from an accumulation mode dominance (when \( M_A \) accounts for ~70% of \( M_T \)) in WMS to a coarse mode domination in SMS when the share of \( M_A \) to \( M_T \) is only ~34% (Table 4).

From the mass-size distribution, an estimate of the effective radius \( (R_{eff}) \), which is the ratio of the volume to the area of the aerosol distribution is also made (Pillai and Moorthy, 2007).
Fig. 6. The mean spectral aerosol optical depth (AOD) during the four seasons (Winter Monsoon Season – WMS (December through March), Inter Monsoon Season – IMS (April and May), Summer Monsoon Season – SMS (June through September) and Post Monsoon Season – PMS (October and November)) considered in this study. Each point corresponds to the seasonal mean value of AOD at a particular wavelength ($\lambda$) and the vertical line through the point corresponds to the standard deviation of the mean.

Table 4. Seasonal changes in the characteristics of composite aerosols. $M_T$ and $M_A$ refer to total as well as accumulation mass concentrations respectively.

<table>
<thead>
<tr>
<th>Seasons</th>
<th>$M_T$ ($\mu g \ m^{-3}$)</th>
<th>$M_A$ ($\mu g \ m^{-3}$)</th>
<th>$M_A/M_T$</th>
<th>$R_{eff}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>WMS</td>
<td>56.24±1.6</td>
<td>39.29±1.4</td>
<td>0.70</td>
<td>0.17±0.004</td>
</tr>
<tr>
<td>IMS</td>
<td>38.50±1.2</td>
<td>18.71±0.9</td>
<td>0.49</td>
<td>0.22±0.007</td>
</tr>
<tr>
<td>SMS</td>
<td>32.31±0.8</td>
<td>11.00±0.4</td>
<td>0.34</td>
<td>0.24±0.006</td>
</tr>
<tr>
<td>PMS</td>
<td>43.18±2.0</td>
<td>19.86±1.3</td>
<td>0.44</td>
<td>0.22±0.01</td>
</tr>
</tbody>
</table>

2001) and is given in the last column of Table 4. It showed a gradual increase in $R_{eff}$ from 0.17 $\mu m$ in the WMS to 0.24 $\mu m$ in SMS and decrease subsequently, in-line with the changes in the concentration of coarse mode aerosols. As $R_{eff}$ of polydispersion represents the radius of an equivalent monodispersion, which will have the same scattering properties as the polydispersion, the above trend in $R_{eff}$ is in line with that of the AOD spectra. These changes occurring at near the surface are well in line with changes in columnar AOD and $\alpha$ seen in Figs. 4 and 5 thereby indicating a general resemblance in the aerosol columnar properties to those at the near surface.

4.3 Aerosol black carbon

In characterizing the absorbing properties of aerosols at a given region, aerosol black carbon has a very significant role. BC is the most light absorbing aerosol species present in the atmosphere and being mostly in the fine size range and chemically inert, has a fairly long atmospheric life time of about 7 to 10 days (Reddy and Venkataraman, 1999), particularly during dry seasons. Longer residence times would lead to higher concentrations. The aethalometer data was used to characterize BC at TVM.

4.3.1 Annual variations

BC mass concentration near India is known to undergo strong annual variation associated with changes in prevailing meteorology (Babu and Moorthy, 2002). In order to understand the climatological trend, monthly mean values of $M_B$ were estimated by averaging the individual observations on each month irrespective of the year as was done for the composite aerosols. The month-to-month variation of this, shown in the top panel of Fig. 7, reveals very large annual variation, by a factor ~4. High $M_B$ values (6 $\mu g \ m^{-3}$) occur during the months of WMS while the concentration is largely
depleted to 1.5 µg m\(^{-3}\) during the months of SMS. Thereafter \(M_B\) gradually increases during PMS and regain its high value in WMS. During the SMS, when \(M_B\) is lowest, average \(M_B\) level is still at 1.5 µg m\(^{-3}\), which means that the BC concentration is significantly high at this location. (Such high BC has implications to the aerosol forcing, as will be evident in the second part of this paper.) The seasonal mean values of \(M_B\) and its median values are given in Table 5. It shows the distribution of \(M_B\) is skewed; with more frequent occurrence of values lower than the mean. This is mainly due to the exceptionally high concentrations of BC occurring in the morning hours (just after sunrise), associated with fumigation effect in the atmospheric boundary layer (for example, Babu and Moorthy, 2002) and in the evening, during the neutral conditions, before the sea breeze reverses to land breeze.

As our study region is not industrialized, the sources of BC would mainly be automobile exhaust and local domestic activities combined with BC advected from the inland regions. There are no combustion sources that are of seasonal in nature. The advection, of course, has strong seasonal dependence as is seen earlier. Several forest fires of moderate strength occur every year during the WMS period over the forest land lying a few hundred kilometers north-east (upwind) of Trivandrum, mainly due to the dry prevailing conditions and it is likely that a part of these emissions would be advected to this location by the favorable wind direction, during this season.

To examine the role of long-range transport of BC to TVM, from the adjoining regions, 5-day airmass back trajectories were computed using Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT, http://www.arl.noaa.gov/ready/hysplit4.html) model of National Oceanic and Atmospheric Administration (NOAA). Mass plots of these trajectories (arriving at 500 m a.g.l. (above ground level) at TVM) for each season are shown in Fig. 8. In the figure, the panels (a) and (b) show the contrasting seasons WMS and SMS, while (c) and (d) shows the periods of transition. It can be seen that during WMS, when the \(M_B\) values are high, most of the trajectories show advection either from the Indo-Gangetic plain or from the East Asia across the Bay of Bengal. There is some significant advection from the west coast of India and Arabia also. In sharp contrast to this, during SMS all the trajectories are of oceanic nature and reach TVM without any continental overpass. As such, \(M_B\) values are the lowest of the year during this season. During the transition months, the trajectories show different types of advection with consequent impact on \(M_B\). During IMS (transition from WMS to SMS) Fig. 8c shows that majority of the trajectories have a long history over the ocean and only less than 30% of the trajectories have a continental or close to continental history (either West Asia, West Coastal India or over Sri Lanka). Unlike this, in PMS we have significant advection from the Indo-Gangetic plain, central peninsular India, west coast of India and east coast of Africa. Only less than 25% of the trajectories are purely oceanic in nature. This result in seasonal mean BC during PMS being significantly higher than that during IMS, even though both are transition periods. Similar effects of long range transport of aerosols from the IGP to the east coast of India have been reported by Niranjan et al. (2005) and at Ahmedabad in western part of India by Ganguly et al. (2006). Corrigan et al. (2006) have also reported impact of continental advection

\[
\begin{array}{|c|c|c|c|}
\hline
\text{Seasons} & \text{Mean } M_B (\mu g m^{-3}) & \text{Median of } M_B (\mu g m^{-3}) & F_{BC} \\
\hline
\text{WMS} & 5.68\pm0.021 & 4.13 & 10.7 \\
\text{IMS} & 2.62\pm0.014 & 2.15 & 6.5 \\
\text{SMS} & 2.01\pm0.006 & 1.44 & 4.0 \\
\text{PMS} & 3.46\pm0.019 & 2.34 & 6.5 \\
\hline
\end{array}
\]

Fig. 7. Annual variation of monthly mean values of total mass concentration (\(M_T\)), accumulation mode mass concentration (\(M_A\)), coarse mode mass concentration (\(M_C\)) and black carbon mass concentration (\(M_B\)) (respectively in the panels from the bottom to the top) at the near surface level. The vertical line through each point represents the standard deviation of the mean.
Fig. 8. Mass plots of the trajectories (arriving at 500 m a.g.l. (above ground level) at TVM) for each season. In the figure, panels (a) and (b) show the contrasting seasons WMS (December through March) and SMS (June through September), while (c) and (d) shows the periods of transition, IMS (April and May) and PMS (October and November).

4.3.2 BC mass fraction ($F_{BC}$)

The share of BC to the composite aerosol mass is important in assessing the radiative impacts of aerosols. Simultaneous estimates of $M_T$ and $M_B$ are used to obtain the mass fraction (the mass-mixing ratio, $F_{BC}$) of BC ($=M_B/M_T$). Considering the uncertainties in the QCM measurements along with those of $M_B$, the overall uncertainties in the mass fraction would be in the range 14 to 18% (Babu and Moorthy, 2002). The seasonal mean values of $F_{BC}$ are given in the last column of Table 5. It shows that the share of BC to the composite aerosols varies by a factor of $\sim$2.5 over the year. But the more important point is that $F_{BC}$ is high when both $M_B$ and $M_T$ are also high and it occurs during the WMS when the sky is rather clear and cloud free. This implies that BC is a major component of the aerosol system when the aerosol burden is high.

INDOEX measurements have reported 14% of BC in fine aerosol (dry mass at diameters <1 µm) and 10% of BC in coarse aerosol (dry mass at diameters between 1 µm and 10 µm) in the marine boundary layer (Lelieveld et al., 2001; Ramanathan et al., 2001). Based on the campaign mode observation over an urban location Bangalore, during the PMS of 2001, Babu et al. (2002) reported an $F_{BC}$ value of 11%. Similarly, Tripathi et al. (2005) reported a range of values for $F_{BC}$ between 7 and 15% over Kanpur (in the Northern India) during December (WMS) 2004. BC mass fraction in the range of 10 to 15% were reported during ACE-2, INDOEX and TARFOX (Novakov et al., 2000; Ramanathan et al., 2001; Satheesh et al., 2002) while over tropical western Pacific Liley et al. (2002) reported BC to contribute as much as 25% to the composite. From measurements at Bangalore, Babu et al. (2002) reported large amount of BC; both in absolute terms as well as in the fraction to the total aerosol mass ($\sim$11%) and fine aerosol mass ($\sim$23%). In the present study, we notice that, though $F_{BC}$ remain consistently high ($\geq$10%) during WMS the absolute values are lower than those reported for northern Indian region. Moreover, the values in Table 5 indicate that $F_{BC}$ undergoes a rapid decrease (by a factor of $>2.5$) to as low as 4% in SMS. This will have its implications on the aerosol single scattering albedo, which in turn will affect the radiative forcing.

4.4 Association between near surface and columnar properties

Generally atmospheric aerosols are produced very close to the surface (except the secondary generation process from precursor gases) and are then distributed spatially by winds, turbulence, diffusion and other atmospheric
processes. Hence, the major share of the aerosol loading at a location has a direct dependence on the composite aerosol concentration at the surface, except when transport process becomes significant at higher level. Thus, for estimating the radiative forcing due to aerosols by incorporating the aerosol properties estimated from near surface measurements, it is important to understand how well the near surface aerosol properties are related to columnar aerosol properties. With this view, we examine the association between the composite (surface) aerosol features and AOD, on an average basis.

4.4.1 Composite aerosol characteristics and Columnar AOD

In Fig. 9, the annual variation of the climatological mean of AOD at 500 nm is compared with that of $M_T$ in the bottom panel and the Ångström component $\alpha$ (which is predominantly a measure of the relative abundance of smaller particles) with the accumulation mode fraction $A_f$ (estimated as $M_A/M_T$) in the top panel. A very close association is evident in both the figures all through, except in April, October and November, which are the transition months. This implies that the surface features are broad indicators to the columnar properties for most part of the year. In the transition seasons, particularly during IMS, long-range transport processes have a strong influence on the columnar properties, because advection generally occur at elevated layer at ~2 to 3 km (Muller et al., 2001). Based on shipboard measurements, Moorthy et al. (2005a) have shown that changes in direction of advection significantly affects the AOD over coastal Arabian Sea, adjacent to TVM, but the near surface concentrations are not influenced. Similarly, the extensive rainfall associated with the return monsoon also would probably affect the vertical heterogeneity of aerosol in PMS. Such associations between surface concentration and column AOD have been reported by others also during field campaigns. Based on INDOEX observation, Parameswaran et al. (1999) have reported an association between the aerosol optical depth at 500 nm and near surface total aerosol mass concentration with a correlation coefficient of ~0.74 over the Arabian Sea and northern Indian Ocean during continental airmass period (January to March). From investigation over Barbados, Smirnov et al. (2000) reported that the monthly averages of dust concentration and columnar AOD at 870 nm are strongly correlated with a coefficient of 0.93.

4.4.2 BC concentration and Columnar AOD

It was noticed earlier that BC contributes from ~4% to as much as 10 to 11% to the composite mass during different seasons. Earlier studies have shown that a 6% of BC contribution to the composite aerosols mass can be associated with as much as an 11% contribution to the AOD (Satheesh et al., 1999). Here, we examine the association of AOD and $\alpha$ with $M_B$. Figure 10 shows the scatter plot of daily mean BC concentration ($M_B$) with the simultaneous daily mean AOD at two wavelengths, 380 nm (the shortest wavelength used in MWR) and 1025 nm (the longest wavelength used in MWR).
Fig. 11. Scatter plot showing the variation of Angstrom exponent (\(\alpha\)) with black carbon mass concentration (\(M_B\)). Here \(N\) is the number of days on which simultaneous observation of spectral AOD and \(M_B\) is available and \(\gamma\) is the correlation coefficient.

The plot contains 109 individual observation pairs combining all the seasons. Despite a fair amount of scatter, (which is expected due to several reasons such as, (a) BC is not the only species contributing to AOD; (b) Temporal changes in the composite aerosol concentrations are not one to one correlated with those of BC; (c) BC measurements are made near the surface while AOD is in the column; (d) BC share to the composite aerosols undergoes a large variation with season and (e) the large amount of coarse sea spray aerosols which are added to the composite aerosols in SMS do not have any BC component) some interesting pattern is seen in Fig. 10. At 380 nm, the variation of AOD is positively correlated with \(M_B\) with a correlation coefficient of +0.38 (which is significant) where as at 1025 nm AOD is very weakly negatively correlated with \(M_B\).

Thus BC has a significant role in determining the AOD at short wavelengths, which is influenced by the accumulation mode aerosols (radius <1 \(\mu m\)). At the longer wavelengths, the AOD is more dependent on coarse aerosols (such as sea salt). It is also well established that most of the BC mass resides in the sub-micron size range (particle radius <1 \(\mu m\)). In view of the above observations, in Fig. 11 we examine the association between \(\alpha\) and \(M_B\). It shows a high positive correlation with a coefficient of 0.69. This result is very important because the points in Fig. 11 are representative of all seasons and it implies that BC has a significant role in the enhancement of \(\alpha\) at TVM, irrespective of the season.

5 Conclusions
Collocated measurements of spectral aerosol optical depth; total as well as size segregated mass concentrations of near surface aerosols; and aerosol black carbon mass concentration are made over a period of four years at Trivandrum, a tropical coastal station on the southwestern tip of India. These are used to characterize the optical and microphysical properties of aerosols and to examine the changes in them associated with the changes in prevailing meteorology. The main findings are:

1. Aerosol optical depth showed large annual variation with high values during Winter Monsoon Season (WMS; months of December through March) and low values during Summer Monsoon Season (SMS; June through September) at short (visible) wavelengths and an opposite pattern at longer (near infrared) wavelengths.

2. The columnar AODs transform from a steep spectra during WMS to shallow spectra in SMS with a transition in Inter Monsoon Season (IMS; April and May) and recovering back to WMS through Post Monsoon Season (PMS; October and November). Correspondingly the Angström wavelength exponent (\(\alpha\)) decreases from a high value (~1.1) in WMS to the least value in SMS (0.32). During the transition period \(\alpha\) is moderate in IMS and while it is the highest in PMS. These changes suggest a change in the aerosol environment from a polluted continental type with significant anthropogenic share to a cleaner marine like environment with significant share of natural aerosols.

3. Composite aerosols, in well-mixed region near the surface, also depict distinct changes in their characteristics with changes in the synoptic meteorology. The total mass concentrations decreases drastically by a factor of ~2 in SMS from the high WMS values. Associated with this, the mass size distribution changes from accumulation mode (radius <1 \(\mu m\)) domination in WMS to coarse mode (radius >1 \(\mu m\)) domination in SMS.

4. Although BC mass concentration (\(M_B\)) measurements have considerable systematic uncertainties due to instrument limitations, there is a significant seasonal dependence in \(M_B\). The aerosol BC concentration as well as its mass mixing ratio decrease from very high values in WMS (respectively 5.6 \(\mu g\) m\(^{-3}\) and 11%) to very low values (1.5 \(\mu g\) m\(^{-3}\) and 4%) in SMS.

5. Columnar (AOD and \(\alpha\)) and near surface features (e.g. composite aerosol mass concentration, accumulation mode fraction and BC mass concentration) are well associated during WMS and SMS, but deviate during the transition seasons.

Such distinct and significant temporal changes in the aerosol characteristics definitely have consequent impact on aerosol radiative forcing. This aspect is examined in Part 2 of this paper.
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