Spatial-Temporal Variability and Characterisation of Aerosols in Urban Air Quality of Ahmedabad, India, based on Field and Satellite Data

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Abstract Particulate matter (PM), or aerosol play an important role in Air Quality. Ahmedabad is an urban, industrialized and one of the fastest growing city in Western India. This study presents spatio-temporal variability in mass concentrations of different Particulate Matter as well as their characteristics during different seasons based on field measurements in different micro urban environments of Ahmedabad. The meteorological parameters such as wind speed, temperature, relative humidity and atmospheric pressure played a very important role in the seasonal variation of the PM concentrations. The PM10, PM2.5 and PM1 were found to be highest with 229.4 µg/m3, 82.20 µg/m3 and 53.53 µg/m3, respectively in winter season. These PM concentrations are higher than the National Ambient Air Quality Standard (NAAQS) for PM 10 and PM2.5. The aerosol characteristic based on field measurement of spectral Aerosol Optical Depth indicates significant contribution by coarser mode particles. The Inhalable, Thoracic and Alveolic PM concentrations which are significant for human health impacts of ambient air quality were observed in the increasing order of monsoon<summer<winter seasons. It is interesting to note that rainfall play a scavenging effect only on the coarser particles with mean PM1 values lowest in summer season (20.4 µg/m3) as compared to PM2.5 and PM10 which are lowest during monsoon season. Vertical profile of aerosols from space-borne Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations (CALIPSO) indicated a predominance of polluted dust type aerosols over the study area at height of 3-4 km which is important constituent of inhalable air of resident population. The results of this study may serve as important inputs for policy makers for formulating action plans and improving air quality by adopting suitable air pollution control measures in fast urbanising environment.

Keywords: Particulate matter, seasonal variation, aerosol characterization, aerosol optical depth, space borne lidar data.

1 Introduction

Atmospheric aerosols or ‘Particulate Matter’ (PM) are solid or liquid particles or both suspended in air with diameter ranging from few nanometers to 100 microns in diameter. Aerosols may originate from natural or anthropogenic sources including vehicular emissions, biomass burning, fossil fuels burning etc. Aerosol particle size and size distribution are important physical properties governing aerosol interaction with atmospheric radiation, cloud formation, transport and deposition, and health effects [1]. In spite of significant advances in air pollution control, a serious problem in many cities is control of particulate matter (PM) concentration levels in the urban environment. Many studies have been conducted to monitor the concentrations of particulate matter in the indoor and outdoor environment and also to study their various sources in the developing countries [2-12]. Currently, PM pollution is the most pressing issue in air quality regulation worldwide besides one of the biggest sources of uncertainty in current climate [7]. Furthermore, atmospheric particulates also affect cloud formation and cessation, which regulate heat transfer in the atmosphere, thereby contributing to climate change. The monitoring of PM and its fractions is considered important in order to characterize the physical and chemical properties of particles, identify major particle sources and quantify their contributions, assess the spatial and temporal variability, and investigate the impact of particles on climate and health.
The climatic effect of aerosols depends strongly on their optical properties, namely the aerosol optical depth (AOD), the Angstrom parameter, and the aerosol size distribution functions. All these, emphasize the need for extensive measurements and analyses of aerosol optical properties [13]. The AOD, which is the integral of the atmospheric extinction coefficient from the surface to the top of the atmosphere, is an important parameter for visibility degradation (due to atmospheric pollution), solar radiation extinction, climate effects, and tropospheric corrections in remote sensing [14]. The derivation of α in different wavelength regions is a useful tool for distinguishing and characterizing the different aerosol types [15]. The α parameter is a qualitative indicator of aerosol particle size or fine mode fraction and also provides information on aerosol size distribution. These values have been used to characterize biomass burning aerosols, urban aerosols, maritime aerosol component in islands and desert-dust aerosols [16-25].

Very few studies have been reported in literature which have presented spatio-temporal variability in particulate matter concentration and satellite derived AOD in urban environment of Ahmedabad. A study reported PM$_{2.5}$ and PM$_{10}$ varies in the range of 59 to 18µg/m$^3$ and 144 to 56µg/m$^3$ and satellite derived mean monthly AOD varied from 1.11 to 0.23 over the Ahmedabad district for the study year 2013. The study also reported that estimated mean annual AOD is higher over Ahmedabad city as compared to Ahmedabad district due to higher concentration of PM from urban vehicular exhaust and industrial emissions [26]. Another study presented spatio-temporal variations in MODIS derived AOD for the recent period 2012–2014. Higher AOD values in the range of 0.5–0.9 indicated urban Ahmedabad as “Fairly Polluted Area” [27]. PM concentration and their relationship with meteorology over Ahmedabad during 2005-2008 were studied [28]. Source identification of aerosol over urban Ahmedabad for a period of 3 years (2000-2003) were studied, however PM variations were not presented in this study [29].

The present study was aimed at understanding the spatial-temporal variability in PM concentration and aerosol characterisation based on field measurement over different locations representing different micro environments in urban Ahmedabad. The effects of meteorological parameters on PM mass concentrations were also studied in detail. The in situ AOD measurement were collected to understand the aerosol characteristics. The vertical profile and type of aerosols were studied using space borne lidar satellite data.

2 Study Area

The study area of Ahmedabad is located in Gujarat state with central latitude 23.03° N and longitude 72.58° E. Ahmedabad experiences three distinct seasons of hot and dry summers to more humid in monsoon and moderately cold winters. The relative humidity ranges from 24%-55% in summers and 47% to 67% in winters. Field measurements were collected at 30 locations, chosen differ varieties of locations prevailing in city, like Industrial, Rural, Urban and Suburban, Traffic area of Ahmedabad city during winter (9-11 February, 2016), summer (13-14 May, 20016) and monsoon season (25-26 August, 2016). Figure 1 shows the locations (blue mark) which are selected for monitoring.

Figure 1. Ahmedabad Site with locations covered under this project
3 Instruments and Methods

Different size PM were measured using Portable Grimm Spectrometer Model 11-A. This Spectrometer has wide particle size range 0.25 μm-32 μm. This instrument also provides quantitative estimation of inhalable, thoracic and alveolic components of the total PM concentration. The instrument was kept at the height of about 7-8 feet above the ground that represents inhalable height for human. At each of the location the PM measurements were carried out every 10 minutes. The measurement sites were selected based on expanse and different microenvironments of Ahmedabad city.

AOD was measured at 28 different locations of Ahmedabad using a hand held Microtops-II Sunphotometer (Solar Light Co., USA) [30] at the recommended wavelengths of 340 nm, 500 nm, 870 nm, 675 nm and 1020 nm respectively by the World Meteorological Organization respectively. All the measurements employing the sunphotometer and processing the data were carried out in accordance to a standard protocol. The vertical profile and type of aerosols were studied using NASA’s space borne LIDAR CALIOP (Cloud-Aerosol Lidar with Orthogonal Polarization) sensor on-board CALIPSO (Cloud Aerosol Lidar and Infrared Pathfinder Satellite Observation) satellite [31].

4 Results and Discussion

4.1 Spatial and Seasonal Variability in Particulate Matter (PM10, PM2.5 & PM1) Concentration in Ahmedabad

The detailed analysis of field data indicates large seasonal and spatial variability in PM concentration at different locations in Ahmedabad city. The PM10 and PM2.5 concentrations were observed to be highest during winter followed by summer and monsoon seasons. Mean PM10 and PM2.5 varies from 234 μg/m³ in winter to 50 μg/m³ in monsoon and 82 μg/m³ in winter to 35μg/m³ during monsoon period respectively. However, mean PM1 values are observed to be highest in winter (54 μg/m³) and lowest in summer season (20 μg/m³). The seasonal variations in measured PM concentrations at different locations in Ahmedabad are shown in Figure 2. Also the metrological conditions during the study period are given in the table 1.

In winter the average spatial concentrations of PM10, PM2.5 and PM1 were found to be higher in the month of February. The inhalable, thoracic and alveolic values were also higher (table 1) for this season. Local pollution emissions are responsible for the higher concentrations due to increased human activity and more bonfires to combat cold. Moreover, low wind speed and high humidity 55–78% during this season leads to reduced rate of removal of aerosol particles through scavenging [32].

Table 1. Metrological parameters over ahmedabad and average PM (PM10, PM2.5 and PM1), inhalable, thoracic and alveolic concentration during different seasons

<table>
<thead>
<tr>
<th>Seasons</th>
<th>Parameters</th>
<th>Winter</th>
<th>Summer</th>
<th>Monsoon</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pressure (mb)</td>
<td>1011-1020</td>
<td>998-1013</td>
<td>1003-1009</td>
</tr>
<tr>
<td></td>
<td>Humidity (%)</td>
<td>47 to 67</td>
<td>24 to 55</td>
<td>56 – 78</td>
</tr>
<tr>
<td></td>
<td>Temperature (°C)</td>
<td>25 to 36</td>
<td>32 to 48</td>
<td>25 to 38</td>
</tr>
<tr>
<td></td>
<td>Wind speed (Kts)</td>
<td>5-6</td>
<td>6-8</td>
<td>4-7</td>
</tr>
<tr>
<td></td>
<td>PM10 (µg/m³)</td>
<td>234.8</td>
<td>154.2</td>
<td>50.1</td>
</tr>
<tr>
<td></td>
<td>PM2.5 (µg/m³)</td>
<td>82.20</td>
<td>47.57</td>
<td>35.50</td>
</tr>
<tr>
<td></td>
<td>PM1 (µg/m³)</td>
<td>53.1</td>
<td>19.90</td>
<td>27.27</td>
</tr>
<tr>
<td></td>
<td>Inhalable (µg/m³)</td>
<td>381.15</td>
<td>329.52</td>
<td>97.48</td>
</tr>
<tr>
<td></td>
<td>Thoracic (µg/m³)</td>
<td>234.83</td>
<td>162.89</td>
<td>53.06</td>
</tr>
<tr>
<td></td>
<td>Alveolic (µg/m³)</td>
<td>113.67</td>
<td>70.23</td>
<td>39.40</td>
</tr>
</tbody>
</table>
In summer, during the month of May, the concentrations recorded were lower, as compared to those recorded during winter. This is possibly due to the increase in temperature and moderate wind speed. Particles are dry due to less humidity 35-48%, and so contribute less in increasing their mass concentrations. Moreover during summer the prevailing winds caused by thermal circulations are stronger and the mixing height is deeper. Wind turbulence with greater mixing height results in proper dilution and dispersion of pollutants during summer [33,34]. The doors and windows are also kept open during the summer for proper air exchange rate which also reduces the concentration of PM especially of the fine particulate size indoors [35].

In Monsoon the concentrations are recorded lowest than in the winter and summer. This can be seen from figure 2 and table 1. Lower aerosol concentrations were also observed in monsoon because of the washout effect of particles from the atmosphere. Larger super micron particles are removed faster than submicron particles and have little effect on their concentration [36]. Coarse particles, on absorbing water settles down and therefore re-suspension and re-entrainment of soil is minimal and particulate matter during the monsoon is mainly composed of finer particles [37]. Interestingly the PM₁ concentration was found to be lowest during summer instead of Monsoon. This clearly points to the fact that the scavenging factor does not affect finer particles.
Table 2. PM(PM10, PM2.5 and PM1) and alveolic extremes concentration during different seasons. All the values are in µg/m³

<table>
<thead>
<tr>
<th>Conc.</th>
<th>Extremes</th>
<th>Winter</th>
<th>Summer</th>
<th>Monsoon</th>
</tr>
</thead>
<tbody>
<tr>
<td>PM10</td>
<td>Max</td>
<td>581.5</td>
<td>286.7</td>
<td>122.5</td>
</tr>
<tr>
<td></td>
<td>Min</td>
<td>99.06</td>
<td>72.50</td>
<td>16.20</td>
</tr>
<tr>
<td>PM2.5</td>
<td>Max</td>
<td>200.76</td>
<td>80.464</td>
<td>93.296</td>
</tr>
<tr>
<td></td>
<td>Min</td>
<td>37.73</td>
<td>24.65</td>
<td>15.134</td>
</tr>
<tr>
<td>PM1</td>
<td>Max</td>
<td>136.14</td>
<td>45.53</td>
<td>70.75</td>
</tr>
<tr>
<td></td>
<td>Min</td>
<td>23.7</td>
<td>9.092</td>
<td>12.64</td>
</tr>
<tr>
<td>Alveolic</td>
<td>Max</td>
<td>272.46</td>
<td>119.15</td>
<td>103.58</td>
</tr>
<tr>
<td></td>
<td>Min</td>
<td>52.09</td>
<td>34.053</td>
<td>15.535</td>
</tr>
</tbody>
</table>

The PM concentration is higher than the NAAQS for PM10 (100 µg/m³) and PM2.5 (60 µg/m³). The PM concentrations are higher in industrial areas (582 µg/m³ in winter to 123 µg/m³ in monsoon) and traffic junction (320 µg/m³ in winter to 72 µg/m³ in monsoon), while open areas like Public Park, colleges, etc., have low PM concentration (99 µg/m³ in winter to 16 µg/m³ in monsoon). During the Monsoon, there is a wet scavenging effect and the concentration steps down, but the scenario remains same for the rest of the year. Ultimately, Alveolic particle concentrations were also seen high in industrial area (272-103 µg/m³), traffic areas and highways (235 µg/m³). This shows that these areas are more harmful for humans as we talk about health impacts, because alveolic (respirable) particles penetrate deep into the respiratory system. According to the survey of Ministry of Health and Family Department (MHFD) and Ahmedabad Municipal Corporation (AMC), 13856, 7168 and 9074 cases of Acute Respiratory Infection (ARI) and Influenza Like Illness (ILI) were recorded during winter, summer and monsoon seasons in Ahmedabad respectively. This is clearly a point of concern because overall average life expectancy is already low (64 years) in India; ranked 150 worldwide in 2012 and future increase in this concentration may worsen the situation [50].

4.2 Ratios of Coarser and Finer PM in Ambient Air

The PM10, PM2.5-10 (PM10 – PM2.5), PM2.5 and PM1 particles are relatively different in their physical and chemical compositions. The different size of atmospheric particles originates from different sources. The ratios of PM2.5/PM10 and PM2.5-10/PM10 have been used for identifying the sources of PM10, PM2.5-10 and PM2.5. Higher PM2.5/PM10 ratios indicate relatively high contribution from secondary particles and from the combustion sources, while lower ratios indicate significant contribution from primary sources such as resuspended soil/road dust and from other mechanical activities [38]. The annual and seasonal statistical summary of the ratios of PM2.5/PM10 and PM2.5-10/PM10 is given in figure 2(d). On average for the 1 year data, the ratio of PM2.5/PM10 for winter and summer was found to be 0.37 and 0.32 respectively, which indicated that ~35% of PM10 was made up of PM2.5 particles. The ratio of PM2.5-10/PM10 during winter and summer was found to be 0.63 and 0.67 during the study period, which is higher than PM2.5/PM10.

The results of the present study denote higher contribution of coarser particles to the PM10 concentration. Clinical and toxicological studies have reported that the coarser particles resulting from road dust (mixed debris from brake wear, tire wear, road abrasion, and biological and geological matter) can be as toxic as fine particles on a mass basis. As the role of particle composition in producing toxicity is uncovered, it becomes increasingly important to investigate the compositional differences between PM2.5 and PM2.5-10 as well as identify their sources. This is of particular importance in urban settings, where traffic-related PM is higher and coarse particles are largely made up of re-suspended road dust [39]. Monsoon is considered to be a special case of wet scavenging.

4.3 Characterization of Aerosols in Urban Environment of Ahmedabad

The AOD wavelength dependence is suitably expressed by the Angstrom-exponent (α) [40,41,15]. The Angstrom turbidity coefficient (β) is related to the amount of aerosols present in the atmosphere whereas the Angstrom exponent (α) yields information on the predominant size of suspended particles.
and it is also an indicator of the relative fine mode strength. The second-order Angstrom exponent (\(\alpha'\)) further quantifies the deviation from linearity of the spectral AOD variation in logarithmic coordinates. The Angstrom wavelength exponent (\(\alpha\)) can be calculated using the AODs (\(\tau\)) at two different wavelengths by applying Equation 1:

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

\[
\alpha = \frac{d \ln \tau}{d \ln \lambda} = -\frac{\ln \left( \frac{\tau_1}{\tau_2} \right)}{\ln \left( \frac{\lambda_1}{\lambda_2} \right)}
\]

Equation (2) shows that \(\alpha\) is the negative of the slope of \(\tau\) with respect to wavelength in logarithmic scale. We make use of the derivative of \(\alpha\) with respect to the logarithm of wavelength to compute the curvature of the \(\ln \tau\) versus \(\ln \lambda\) curves. The second derivative is a measure of the rate of change of the slope with respect to wavelength and, therefore, is a complement to the Angstrom exponent, which is the negative of the slope (first derivative) of \(\ln \tau\) versus \(\ln \lambda\). We can calculate the second derivative of \(\tau\) as follows:

\[
\alpha'(\lambda) = \frac{d \alpha}{d \ln \lambda} = \left( \frac{2}{\ln \lambda_{i+1} - \ln \lambda_{i-1}} \right) \left( \frac{\ln \tau_{i+1} - \ln \tau_i}{\ln \lambda_{i+1} - \ln \lambda_i} - \frac{\ln \tau_i - \ln \tau_{i-1}}{\ln \lambda_i - \ln \lambda_{i-1}} \right)
\]

The \(\alpha'\) values were derived using the observed AODs at 340 nm, 500 nm, 870 nm and 675 nm wavelengths. The aerosol size distribution is rarely unimodal in the atmosphere due to different sources of origin and the mixing processes. Once the aerosol size distribution is multimodal, the wavelength dependence of AOD does not follow Equation (1) accurately. Consequently, a deviation from the linear behavior of \(\ln \tau\) versus \(\ln \lambda\) is expected and has been reported by several studies. The curvature in the AOD spectra could be expressed as a second-order polynomial fit, written as

\[
\ln \lambda = a_1 (\ln \lambda)^1 + a_2 (\ln \lambda)^2 + a_3
\]

If \(\alpha_2 < 0\) and \(\alpha' > 0\), then the curvature will be concave indicating aerosol size distributions dominated by the fine mode and if \(\alpha_2 > 0\) and \(\alpha' < 0\), then the curvature will be convex indicating size distributions with a significant contribution by the coarse mode. Our result and calculation indicates contribution by coarse mode. The polynomial fit to Equation (4) is more precise than the linear fit to logarithms of Equation (1), while for low turbidity conditions large deviations in the above fits usually occur [42].

| Table 3. AOD and angstrom parameters values |
|-------------------------------|-------------------|------------------|
| AOD 500                      | 0.384             | \(a_{340,870}\) 0.924 |
| AOD675                      | 0.291             | \(\beta\)        0.223  |
| AOD870                      | 0.242             | \(a'\)           -0.573  |
| AOD340                      | 0.573             | \(a_0\)          0.25    |
| \(a_{340,500}\)            | 1.041             | \(a_1\)          -4.10   |
| \(a_{675,870}\)            | 0.734             | \(a_2\)          14.78   |

The average value of \(\alpha\) calculated in the range 340–870 nm is 0.92 with turbidity parameter (\(\beta\)) of 0.22 and average \(\alpha'\) of -0.57. A value of \(\beta > 0.2\) usually indicates heavy pollution while \(\beta > 0.4\) indicates extremely heavy pollution. Large positive values of \(\alpha'\) are attributed to fine-mode dominated aerosol size distribution whereas close to zero and negative values of \(\alpha'\) are characteristic of size distributions with a dominant coarse mode [42]. The results show that in winter, the value of \(\alpha_2\) is positive, signifying domination by the coarse-mode aerosols; these aerosols are of dust origin in the majority of the cases.

Extensive analyses based on spectral aerosol measurements indicate that the \(\alpha\) varies with wavelength, and that the spectral curvature of the least-squares fit contains useful information about the aerosol size distribution [43-48]. More specifically, researchers investigated the sensitivity of \(\alpha\) to both monomodal and bimodal aerosol size distributions, and explored the information content in the curvature of alpha.
Different values of the fine-mode fraction strongly modify the curvature changing its sign from negative to positive [49].

![Graph](image.png)

**Figure 3.** ln(λ) vs. ln(τ) graph

Studies show that when $\alpha$ is close to unity, the coarse- and fine-mode aerosol concentrations are nearly equal and it gradually increases as the contribution of fine-mode aerosol concentration increases when one moves toward the more polluted areas. The $\alpha$ calculated using the spectral variation of AOD, has been used in various other previous studies as a tool to estimate particle size distribution and for extrapolating AOD throughout the broad spectral region as well as to distinguish the different aerosol types [42].

### 4.4 Space Borne LIDAR Data Analysis over Study Site

Space borne Lidar data of February, 2016 collected from CALIOP-CALIPSO [31] is given in Figure 4. The image clearly shows the dominance of dust and polluted dust type of aerosols over Ahmedabad and nearby regions at a height of about 3-4 kms which also validates the results of aerosol coefficients indicating that Ahmedabad is dominated by coarse mode aerosols mainly contributed by polluted dust.

![Image](image.png)

**Figure 4.** Vertical profile of aerosols studied using space borne lidar data
5 Conclusions

Particulate Matter Aerosol constitute a significant component of urban air quality. This study presented spatio-temporal variation and characterization of PM aerosol in urban air quality over Ahmedabad, India. Results show that mean PM\textsubscript{10} varies from 234.8 µg/m\textsuperscript{3} in winter to 50.1 µg/m\textsuperscript{3} in monsoon while PM\textsubscript{2.5} varies from 82.2 µg/m\textsuperscript{3} to 35.5 µg/m\textsuperscript{3}. However, mean PM\textsubscript{1} values are observed to be the highest in winter (53.1 µg/m\textsuperscript{3}) and interestingly lowest in summer season (19.9 µg/m\textsuperscript{3}). The PM concentration is higher in industrial areas (582 µg/m\textsuperscript{3} in winter to 123 µg/m\textsuperscript{3} in monsoon) and traffic junction (320 µg/m\textsuperscript{3} in winter to 72 µg/m\textsuperscript{3} in monsoon), while open areas like Public Park, colleges, etc., have low PM concentration (99 µg/m\textsuperscript{3} in winter to 51 µg/m\textsuperscript{3} in monsoon). Alveolar particle concentrations were also seen high in industrial area (272-103 µg/m\textsuperscript{3}), traffic areas and highways (235 µg/m\textsuperscript{3}). According to the data obtained from IDSP and AMC, ARI cases are almost twice during winter than the rest of the year. The PM\textsubscript{2.5}/PM\textsubscript{10} and PM\textsubscript{2.5-10}/PM\textsubscript{10} ratio indicates that the environment of Ahmedabad is dominated by coarser mode which contains mainly dust. The coarse mode domination is also verified through the CALIPO satellite image over Ahmedabad. The AOD calculation further confirms the coarse mode domination and also points towards multimodal behavior of aerosols. Dominance of coarse-mode aerosols, particularly dust and polluted dust type aerosols in urban environment of Ahmedabad is evident from LIDAR and AOD measurements and measurement of PM concentration. This study provides useful insights to current state of sources, contribution and characterization of PM aerosol to urban air quality.

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