Abstract. The direct aerosol radiative forcing (DARF) has been estimated for the clear-sky conditions over Delhi from January 2006 to January 2007 using Santa Barbara DISORT Atmospheric Radiative Transfer model (SBDART) in the wavelength range 300–3000 nanometer. The single scattering albedo (SSA) and the asymmetry parameter used in this model were estimated using the Optical Properties of Aerosol and Cloud (OPAC) model. The annual average AOD observed at 500 nm was $\sim 0.86 \pm 0.42$ with an average Angstrom exponent $\sim 0.68 \pm 0.35$. The average monthly AOD throughout the year over Delhi was found to be in the range 0.56 to 1.22 with the Angstrom exponent in the range 0.38 to 0.96. A high monthly average BC concentration in the range 4–15 $\mu$g m$^{-3}$ led to monthly average SSA in the range 0.90±0.4 to 0.74±0.3 during the year. Consequently, the monthly average clear-sky DARF at the surface was found to vary in the range $-46 \pm 8$ W m$^{-2}$ to $-110 \pm 20$ W m$^{-2}$, at TOA in the range $-1.4 \pm 0.4$ to $21 \pm 2$ W m$^{-2}$, whereas in the atmosphere it was in the range $46 \pm 9$ W m$^{-2}$ to $115 \pm 19$ W m$^{-2}$ throughout the year. As the dust concentration in the atmosphere was highest (May–June) the SSA showed an increase with wavelength however when dust concentration was low the SSA decreased with the wavelength.

Keywords. Atmospheric composition and structure (Aerosols and particles)

1 Introduction

The atmospheric aerosols have assumed greater significance recently in climate studies because of their potential to perturb the radiation balance of the earth at regional and global scales (Charlson et al., 1992; Haywood and Shine, 1995; Haywood and Boucher, 2000; Penner et al., 2004). It causes changes in radiation balance of the earth by direct as well as indirect interactions with the solar radiation, giving rise to the aerosol radiation forcings that are generally classified as direct and indirect. The direct aerosol radiative forcing (DARF) involves scattering and absorption of solar radiation by atmospheric aerosols, which, depending upon the nature of aerosols and the surface albedo may produce negative or positive radiation forcing at the surface. The indirect radiative forcing is the mechanism by which aerosols modify the microphysical and hence the radiative properties, amount and lifetime of clouds (IPCC, 2007, and the references therein). The estimation of indirect radiation forcing is far more complex than the estimation of direct radiative forcing. The present study is limited only to the direct aerosol radiative forcing during the clear-sky conditions.

Aerosol radiation forcing at any layer in the atmosphere is defined as the difference in the net fluxes (down minus up) with and without aerosols at that layer. The magnitude and sign of ARF is determined by different aerosol species, their size distribution, and chemical characteristics. Particularly the absorbing aerosols like black carbon and some of the mineral dust aerosols significantly alter the magnitude and sign of the ARF by altering the aerosol single scattering albedo (SSA). The existing scientific literature extensively documents the mineral dust aerosols (Duce, 1995; Miller and Tegen, 1998; Quijano et al., 2000; Prospero et al., 2002; Tegen et al., 2004; Zhu et al., 2007) and the carbonaceous aerosols (Penner, 1995; Jacobson, 2001; Chung and Seinfeld, 2002; Wang, 2004) in the atmosphere and their impacts on aerosol radiation forcing.

The Asian continent accounts for second highest annual mean aerosol optical depth (AOD) in the world, next to Africa (Holben et al., 1998). Several field campaigns have already been undertaken in the Asian region to study the aerosol characteristics and its impacts on aerosol radiative forcing in this continent and the surrounding oceans starting
from the Indian Ocean Experiment (INDOEX), the Bay of Bengal Monsoon Experiment (BOBMEX), the Arabian Sea Monsoon Experiment (ARMEX), the Integrated Campaign for Aerosols, gases and Radiation Budget (ICARB) and the East Asian Studies of Tropospheric Aerosols: An International Regional Experiment (EAST-AIRE). The overview of these campaigns are given by Ramanathan et al. (2001); Bhat et al. (2001); Moorthy et al. (2008), and Li et al. (2007). These experiments have provided a wealth of information on aerosol characterization, size distribution and aerosol radiation forcing in the targeted region during the campaign period.

Similarly, in the north Indian region also there are some studies on aerosol characterization and forcing but they are limited to pre-monsoon season only (Singh et al., 2005; Prasad et al., 2007; Pandithurai et al., 2008; Beegum et al., 2008). Present study, for the first time, is focused on the monthly variation of clear sky direct aerosol radiation forcing (DARF) over a north Indian station, Delhi, for the whole year during January 2006 to January 2007. Being a part of the Indo-Gangetic Basin the aerosol optical properties over Delhi may resemble what it has been reported earlier from the Indo-Gangetic basin (Dey et al., 2008). It is expected that the anthropogenic aerosols rich in its black carbon (BC) content, present in the atmosphere throughout the year, gets mixed with the mineral dust aerosols from deserts during pre-monsoon months, April–June. The DARF in the wavelength range 300–3000 nm has been estimated during clear sky days every month during January 2006 to January 2007 to see its seasonal variation over Delhi. It has been estimated using the Santa Barbara DISORT Atmospheric Radiative Transfer model (SBDART). The input to run this model was obtained using the ground based observations and with the help of aerosol optical properties obtained using the model OPAC (Optical Properties of Aerosol and Clouds). The average surface albedo values were retrieved from the Aura OMI version 3 reflectivity data through Giovanni online data system, developed and maintained by the NASA GES DISC.

2 Experimental site and instruments

Delhi, the capital of India, located between the latitudes of 28°24'17" and 28°53'00" North and longitudes of 76°50'24" and 77°20'37" East, having an area of 1483 sq. km, is one of the most densely populated cities in Asia (10 340 persons km⁻²). It is located in a semi-arid region close to the Thar and Margo desert in the West and experiences extreme weather conditions. Summer temperature peak goes more than 45 °C during daytime and the lowest temperature in winter season reaches 1 °C in the night. The rainy season for Delhi is during July and August when higher humidity levels exist. During the year 2006, the total rainfall was 708 mm out of which a maximum of about 260 mm occurred during the end of July. In the pre-monsoon period (April–June), frequent dust storms from western and north-western desert regions cause large scale loading of dust aerosols over Delhi. This causes considerable reduction in visibility and in the radiation flux reaching the surface (Singh et al., 2005). According to the economic survey report of Delhi 2007–2008 (http://delhiplanning.nic.in/Economic20Survey/ES2007-08/ES2007-08.htm), the average suspended particulate matter (SPM) and respirable suspended particulate matter (RSPM) concentration over Delhi during 2006 was 433 μg m⁻³ and 174 μg m⁻³, much higher than the permissible safe limits. It also states that most of the local pollution in Delhi is attributed to the vehicular emission (~80%). The vehicular density here is more than 10 times the average vehicular density of the country with more than 5.5 Million vehicles. It is further increasing at an average annual rate of 8–10%. The industry and domestic use contribute 12 and 8%, respectively, in the total pollution of Delhi. The estimated vehicular emission load in Delhi for particulate matter alone is about 13 ton per day.

For the present study, the aerosol optical depths (AOD) were measured with recently calibrated MICROTOPS-II sunphotometer (Solar Light Co., USA) at five wavelengths, 380, 500, 675, 870 and 1020 nm (full width at half maximum, FWHM: ±2–10 nm). The MICROTOPS-II observations are susceptible to filter degradation errors and needs periodic filter calibration (Morys et al., 2001). The pointing accuracy of the instrument is better than 0.1° and long-term stability of the filter is better than 0.1 nm year⁻¹. The accuracy of measurements for precision and consistency of the MICROTOPS-II instruments are discussed in detail by Srivastava et al. (2006). The accuracy of the MICROTOPS-II was found within 1.8%. For AOD retrieval from direct sun measurements, as in the case of MICROTOPS-II, theoretical and/or observations error in AOD can vary in the range of 0.002–0.021 in the visible to near infrared wavelength range, maximum being at 340 nm and minimum at 1020 nm (Eck et al., 1999; Schmidl et al., 1999).

The aerosol black carbon (BC) measurements were done with a Magee Scientific Model AE21 Aethalometer, which measures BC using light beams from a high-intensity LED lamp at 880 nm wavelengths by measuring the attenuation of light transmitted through particles that accumulates on a quartz fiber filter strip. A vacuum pump draws air so that particles continuously accumulate on the filter paper. The measurement of the attenuation of light beam is linearly proportional to the amount of BC deposited on filter strip. The BC content of the aerosols deposited was derived at 5 min integrating time interval. The flow was maintained at 21 per minute. The instrument has been factory calibrated and the manufacturer reports an error in BC measurement as ±2%. More details about the instrument can be found elsewhere (Hansen et al., 1984; Hansen, 1996). However, the uncertainty in BC estimates may be much higher due to the other absorbing component (Hematite, Fe₂O₃) in the aerosols (Hansen et al., 1993); due to the reduction in the
optical path in the Aethalometer filter with an increasing filter load, also referred to as the “shadowing effect” (Weingartner et al., 2003); and due to multiple scattering of light in the fiber matrix of the filter (Arnott et al., 2005). In addition, the uncertainty due to change in sampling conditions and instrument noise may also be present but this will be considerably reduced in the present case as we are taking long averaging periods. Weingartner et al. (2003) have suggested a correction factor of 2.14 for multiple scattering corrections. In the present case we have used a value of 1.9 which may lead to an over estimation of about 12% in BC concentration. Considering all other factors the overall uncertainty in the BC measurements may be around 20%.

In addition, the regular measurements on aerosol number size distribution have also been made using a 15-Channel portable aerosol optical particle counter (OPC) from GRIMM (Model: 1.108). The instrument has been widely used for aerosol size distribution measurements (Jayaraman et al., 2006; Pant et al., 2006). This instrument uses a light-scattering technology for single-particle counts, whereby a semiconductor laser serves as the light source. The scattered signal from the particle passing through the laser beam is collected at approximately 90° by a mirror and transferred to a recipient-diode. The signal of the diode passes, after a corresponding reinforcement to a multi-channel size classifier. A pulse height analyzer then classifies the signal transmitted in each channel. The instrument sucks in the ambient air and segregates the aerosols in accordance with the aerodynamic diameter into one of its 15 size modes. The OPC provides cumulative aerosol number concentration in 15 different size modes ranging from >0.3 μm to >20.0 μm with a detection range in 1–10⁶ counts per liter with a sensitivity of 1 particle per liter. The uncertainty in number of counts may depend upon the relative humidity, due to clogging of particles. In order to avoid this, the instrument was kept inside an all weather housing to maintain the humidity inside the instrument at 50% RH. Further precaution was taken in the present case to operate the instrument only till RH was 75%. More details about the instruments can be found at http://www.grimm-aerosol.com.

3 AOD, Angstrom exponent and aerosol size distribution

The aerosol optical depths were measured at five wavelengths 380, 500, 675, 870 and 1020 nm during clear-sky conditions using MICROTOPS-II sunphotometer. The clear sky conditions here imply the cloud free days or the intervals of the day that were cloud free. The microtops observations were taken only when the sun was completely visible. The data was taken nearly every half an hour during the day time 09:00 to 16:00 h on cloud free days. On partially cloudy days data was taken during the intervals that were cloud free. Care was taken so that the field of view remains cloud free during the observations. A minimum of three continuous observations (within a short span <20 s) were made every time the instrument was pointed towards the sun. Out of these three observations the one with minimum AOD at 500 nm was used for further analysis. This was done to ensure better pointing of microtops towards the sun as the minimum AOD will correspond to maximum pointing accuracy at the given time. The daily average was calculated only for those days which had at least six clear sky observations during the day.

Figure 1 shows the average daytime AOD measured at 500 nm during clear-sky days since 1 January 2006 to 31 January 2007 along with the Angstrom exponent, $\alpha$. The $\alpha$ parameters are retrieved by least square fit on a log-log scale plot of the observed AOD versus wavelength. For a given spectral range the variation of AOD follow Angstrom power law given by $\tau(\lambda) = \beta \lambda^{-\alpha}$ (Angstrom, 1964). In the present case AOD measurements at 340, 500, 675, and 870 nm have been used to estimate $\alpha$. The AOD at 1020 nm may be contaminated due to the absorption of water vapor in the column and therefore it has been excluded in the calculation of $\alpha$. Angstrom exponent is a rough indicator of the size distribution of the aerosol particles in the column. A low $\alpha$ value can be caused by a dominance of bigger-size particles and a high $\alpha$ indicates dominance of smaller-size particles (Dubovik et al., 2002). A close analysis of AOD and $\alpha$ taken together, shows three characteristic variations of aerosols over Delhi during different seasons. During pre-monsoon period high AOD values associated with near zero $\alpha$ values is observed showing dominance of desert aerosols, in the post monsoon to winter period moderate AOD values are associated with the moderate to high $\alpha$ values, indicating dominant anthropogenic aerosols in the atmosphere, and during autumn season moderate to low AOD are associated with the moderate to high $\alpha$ values, indicating mixed aerosol over Delhi. During monsoon (wet) season, as the coarse particles get settled due to downpour, the AOD is generally low and the $\alpha$ values are moderate. A sudden rise in AOD during October end is mainly due to the festival season of Diwali when lots of crackers are burnt and a smoky atmosphere persists for more than a week over Delhi (Attri et al., 2001).
The variation of monthly averaged AOD and $\alpha$ has been plotted in Fig. 2. The monthly averaged data included a minimum of ten days of average (during August) and a maximum of 22 days (in November). The annual average AOD at 500 nm during the period of present observations is $\sim 0.86 \pm 0.42$ with an average Angstrom exponent $\sim 0.68 \pm 0.35$. The average monthly AOD showed maximum values during the months of October ($1.22 \pm 0.20$) and May ($1.12 \pm 0.30$). The high value in May is due to the enhancement in natural desert dust aerosols whereas during October it is mainly due to anthropogenically-enhanced pollution during the festival month. The average monthly Angstrom exponent values during these months were $0.89 \pm 0.27$ and $0.54 \pm 0.32$, respectively, indicating that the fine mode particles dominated more during polluted month of October. On the other hand the minimum monthly average AOD for the whole year occurred during the months of March ($0.56 \pm 0.29$) and August ($0.59 \pm 0.10$) with the average $\alpha$ values being $0.80 \pm 0.19$ and $0.53 \pm 0.21$, respectively. As there are considerable variations in AOD and $\alpha$ throughout the year, it is expected that the average aerosol radiation forcing will also vary during different seasons at Delhi.

The monthly average number-size distribution of aerosols were measured at the surface and it was found that in general for all the months particles with diameter less than 1 $\mu$m dominated in number. Figure 3 shows the four kinds of distinguished number size distribution observed over Delhi. During winters the size distribution is more or less unimodal. For all other seasons the distribution is strongly bimodal. During pre-monsoon period the coarse particle (>1 $\mu$m) concentration increases significantly showing an increase of 1–2 order as compared to other seasons. The autumn season is relatively clean showing lowest concentration in the coarse mode. During monsoon season the concentration of both fine and coarse mode particles decreases. However, the decrease is more pronounced in the coarse mode.

4 Estimation of SSA and asymmetry parameter

The single scattering albedo (SSA) and the asymmetry parameter are very critical to the estimation of aerosol radiation forcing. The SSA is the fraction of energy removed from the incident wave that reappears as scattered radiation whereas the asymmetry parameter is the average or statistically expected value of the cosine of the scattered angle for the scattered radiation. As we do not have the provision of measuring these two parameters directly, it is being estimated in the present case using the OPAC (Optical Properties of Aerosol and Clouds) model. The model, developed by Hess et al. (1998) provides microphysical and optical properties of aerosols and clouds in the solar and terrestrial spectral range of atmospheric particulate matter. The optical properties of aerosol and clouds components are all calculated in the model for 1 particle cm$^{-3}$ and the program allows for a mixture of aerosol components to aerosol types consisting of several components and to calculate a wide range of their optical properties. As the aerosol in the atmosphere is a mixture of different components that can be obtained either through the ground measurements or by the use of typical mixtures, called aerosol types, given in the OPAC model itself. Thus multiplying the stored optical properties data with realistic number densities obtained either with the observed data or the model values available in the OPAC, gives absolute optical properties of aerosol in the atmosphere. The model can give optical properties at up to 61 wavelengths between 0.3 $\mu$m and 40 $\mu$m and for up to 8 relative humidity values.

In the present study, as the aerosols over Delhi are generally of anthropogenic origin and may consist of various kinds of sulfates, nitrates, water soluble substances, soot and other water insoluble components, hence the “urban” aerosol type has been chosen in the OPAC model for the estimation of optical properties of aerosols. However, the concentration of these components was varied time-to-time depending upon the observations. The actual measurements of soot particles
using the Aethalometer (described earlier) have been used in the model to estimate the optical properties of the aerosols. Figure 4 shows the monthly average of the daytime measurements of black carbon (soot) concentration along with their standard deviations over Delhi during 2006. Significant variation in the concentration of BC during winter and summer months even during the day time may be noticed, which can cause large variations in the SSA. It is to be mentioned here that as the BC concentration data for the month of November 2006 was not available hence an average value of the same for the month of November 2005 (BC=9.57 µg m⁻³) has been used in the SSA estimation done later on. This was done because the average monthly BC concentrations, particularly during the winter months of year 2005 were not much different from those in the year 2006. The average BC concentration was highest during December 2006 (16.7 µg m⁻³) and lowest during August (3.26 µg m⁻³).

Since aerosol particles in the atmosphere exist as a mixture of several aerosol components, in the present study, the external mixing approach of the OPAC model has been used to estimate the SSA of the aerosol. Generally the “urban” aerosol type of the OPAC model was taken as a base where the different components of aerosols, for example water soluble, insoluble and soot, concentration was varied month to month as per the observation and available data. During the pre monsoon period, due to the abundance of desert dust over Delhi, a combination of two aerosol types “urban” and “desert” of the OPAC model has been used to represent the average aerosol composition that included mineral dust in accumulation and coagulation modes apart from the soot water soluble and insoluble components. The validity of aerosol structure during pre-monsoon period has already been tested as monthly average during January 2006 to January 2007.

As the OPAC model also estimates the AOD along with the SSA and Asymmetry Parameter, it can be used to cross check the assumed aerosol structure. For a given aerosol component and its concentration the OPAC model was used to calculate the AOD at 500 nm and compared with the observed AOD values. The AOD obtained by the model has been constrained by the urban aerosol structure of the model along with the actual BC measurements at surface as described above. In absence of the measurements of vertical profile of aerosols, the default aerosol profile of the model has been used in the calculation. However, the boundary layer height was chosen as 1 km during winters and 2 km during the summers. This assumption is quite reasonable as similar values are also observed by the sodar measurements over Delhi. The AOD values thus obtained from the OPAC model on all clear-sky-days were compared with the observed AOD and the ARF calculations were done for those days where the two AODs at 500nm matched within ±0.05. The Angstrom exponent α was also compared for these days and was found to agree within 5%.

The single scattering albedo thus obtained has been plotted as monthly average during January 2006 to January 2007 along with the Asymmetry Parameter in Fig. 5. The vertical bars denote the ±1 σ standard deviation from the mean values. The average SSA at 500 nm over Delhi during the whole period of observation was found to be 0.79±0.05. It was minimum during January at an average value of 0.74±0.03 and maximum during the month of August with a value of 0.89±0.04. The average asymmetry factor during the period had a value of 0.68±0.03 with a maximum at 0.73±0.04 during June and minimum at 0.65±0.03 during January. In general the SSA over Delhi showed low values, which were mainly due to high concentration of absorbing BC aerosols. Minimum SSA occurred during winter months when high BC concentrations were also observed. The SSA gradually increased from winter month of January to reach a maximum during monsoon month of August when the BC concentration was also found minimum.

The SSA variations in similar ranges have been found at other large cities in India during different campaign observations. For example the SSA at 500 nm at Pune was found to be about 0.81 during 2001–2002 dry season (Pandithurai et
During August, which is the cleanest month in Delhi, SSA increased with the wavelength. Huge differences in SSA values of the order of 0.15 may be noticed for the dust and pollution dominant months at higher wavelengths (1.0 and 1.5 μm). During August, which is the cleanest month in terms of aerosols/BC the SSA at shorter wavelength shows the highest value and also the decrease in SSA with wavelength is steepest (0.13 as λ changes from 0.35 to 1.50 μm). During winters for pollution ridden aerosols the variation is nearly flat with minimum slope. For other months when pollution and dust may be mixed the SSA again decreases with wavelength and lies in between the extreme values found during August and January. Somewhat similar observation of increase in SSA with wavelength during high dust concentration and decrease in SSA during low dust concentration has been reported by Holler et al. (2003) during ACE-Asia (Aerosol Characterization Experiment) campaign at a coastal site near the Sea of Japan. At a nearby location Kanpur, similar observations have also been reported by Prasad and Singh (2007).

As far as the uncertainty in estimation of SSA using OPAC model is concerned, it may mainly arise from the uncertainty in estimation of composition of aerosol type, state of mixing assumptions and the vertical distribution of aerosols in the atmosphere. Although it is not possible in the present study to ascertain the exact uncertainty due to these factors, however we have tried to do a sensitivity analysis for the effect of BC concentration on SSA as in the present study ‘urban’ aerosol type of the OPAC model has been scaled based on surface BC concentrations measurements. In order to do this the OPAC model was run several times with the BC concentration varying from 2 to 20 μg m⁻³ with an increment of 2 μg m⁻³ keeping all other parameters constant and the optical parameters such as AOD at 500 nm, SSA and Asymmetry parameters.

**Fig. 6.** Variation of single scattering albedo with wavelength as estimated using OPAC model during typical months in different seasons over Delhi.

**Fig. 7.** Effect of change in BC concentration over AOD, SSA and asymmetry factor estimated using OPAC urban aerosol model.
were estimated. Figure 7 shows the result of this exercise where changes in optical properties of the aerosol with varying concentration of BC have been plotted. One can notice that an increase in BC concentration from 2 to 20 $\mu$g m$^{-3}$ leads to a decrease in SSA from 0.90 to 0.60. The change in asymmetry parameter is from 0.68 to 0.63 and an increase in AOD at 500 nm is from 0.57 to 0.91. Thus we can see that on an average in the range of simulation (2–20 $\mu$g m$^{-3}$ BC concentration) an increase of 2 $\mu$g m$^{-3}$ of BC can cause an average change in SSA of about 0.03 (0.05 at 4 $\mu$g m$^{-3}$ and 0.02 at 20 $\mu$g m$^{-3}$). As the maximum BC concentration measured was 16.7 $\mu$g m$^{-3}$ as a result the maximum uncertainty in BC could be about 3.4 $\mu$g m$^{-3}$ which can lead a maximum uncertainty of 0.07 in the SSA estimation. The standard deviation of the monthly averages of SSA also lies in the range 0.03 to 0.06.

5 Direct aerosol radiation forcing

The aerosol radiation forcing at any level in the atmosphere is defined as the difference in the net solar flux (down minus up) with and without aerosol, keeping all other parameters constant. In the present case direct aerosol radiation forcing (DARF) has been estimated in terms of W m$^{-2}$ at the surface and at the top of the atmosphere. The difference of the two therefore gives the DARF in the whole atmosphere. The DARF was estimated using the Santa Barbara DISORT Atmospheric Radiative Transfer model (SBDART). The model was developed at the University of California, Santa Barbara by Ricchiazzi et al. (1998) based upon reliable physical models developed by the atmospheric science community and is being used widely for the radiative transfer calculations. The model uses a large number of parameters to estimate the radiation flux at different layers in the atmosphere. The parameters critical for DARF estimations are the AOD, single scattering albedo (SSA), asymmetry parameter, surface albedo, profiles of atmospheric parameters (temperature, humidity, ozone and other gases), and solar geometry of location. The profiles of atmospheric parameters are generally fixed by once choosing the model atmosphere out of the available options in the model.

On the basis of prevailing weather conditions and the measured meteorological parameters, we have used the mid-latitude summer atmospheric model in the present case. In order to have a better representation of the atmosphere, we have also used the integrated column ozone and the average integrated water vapor values for Delhi. The surface albedo values were obtained for the year 2006 from the Aura OMI version 3 reflectivity data through Giovanni online data system, developed and maintained by the NASA GES DISC. The average monthly values of surface albedo during winters ranged from 0.15 to 0.20 and during the months June–August, it ranged from 0.25 to 0.30. The other two critical parameters SSA and Assymetry parameters were estimated using the OPAC model as described in the above section. Using these parameters the SBDART model was run with eight streams to obtain the TOA and the surface downward and upward fluxes at 1-h interval for a 24-h period with aerosols and without aerosols separately. DARF was then calculated at TOA and surface.

Figure 6 shows the average variation in DARF along with the standard deviations obtained over Delhi at the surface, TOA and in the atmosphere every month. The average forcing for the whole period of observations at the surface was of the order of $-67 \text{W m}^{-2}$, at the top of the atmosphere it was about $4 \text{W m}^{-2}$, giving rise to an atmospheric forcing of about $+71 \text{W m}^{-2}$. The DARF at the surface is found to be maximum (maximum negative) during the pre-monsoon months of May ($-110 \pm 20 \text{W m}^{-2}$) and minimum (least negative) during August ($-46 \pm 8 \text{W m}^{-2}$). The TOA forcing was maximum during June at $21 \pm 2 \text{W m}^{-2}$ and minimum at $-1.4 \pm 0.4 \text{W m}^{-2}$ during November. The DARF in the atmosphere showed maximum value in the month of June ($115 \pm 19 \text{W m}^{-2}$) and minimum during August ($46 \pm 9 \text{W m}^{-2}$). The DARF values were generally high during summers and low during winters. The value increased from March to reach a maximum during May as the contribution of dust aerosols generally increased in the atmosphere. Afterwards, as the monsoon started during July–August the values decreased and remained more or less constant during winters (after September), except for the month of October, which also had high AOD due to the increase in smoke and particulate matter owing to fire cracker burning during this festival months.

The uncertainty in the DARF calculation in the present case may arise from the difference in the used mid-latitude-summer atmospheric condition and the real atmosphere, deviations in the surface albedo values, uncertainty in AOD measurements and the uncertainty in the SSA estimation of aerosols. The combined maximum uncertainty due to the atmospheric condition and the deviation in surface albedo values is $\sim 7\%$. The uncertainty in AOD introduces an additional uncertainty of about 8%. It was estimated by repeatedly running the SBDART model to compute the fluxes
by varying one property at a time while keeping the others constant. It is thus used to find the sensitivity to the range of observed values and expressed as the radiative forcing efficiency. It is found that for 0.1 change in SSA the TOA forcing changes by $-8.7 \text{ W m}^{-2}$ and surface forcing by $14.7 \text{ W m}^{-2}$. Similarly, for AOD the radiative forcing efficiency at TOA is $7.4 \text{ W m}^{-2}$ and at the surface it is $-12.7 \text{ W m}^{-2}$.

6 Conclusions

The monthly mean variations of the aerosol optical properties like, AOD, SSA and the asymmetry factor along with the surface albedo were studied over the mega-city of Delhi during the year 2006. Their subsequent impacts on the DARF at TOA and at the surface were estimated. We have observed a high AOD value and a very high load of BC aerosols throughout the year. The AOD showed peaks during pre-monsoon months of May–June and again in October. The former was due to the heavy load of desert dust aerosols arising due to sandstorms during pre-monsoon and the latter was due to burning of crackers during festivities. Consequently they have a bearing on the DARF estimation over Delhi during the year. From the present study of aerosols optical characteristics over Delhi during January 2006 to January 2007 and the estimation of DARF, the following conclusions can be made:

1. The annual average AOD at 500 nm during the period of present observations is $\sim 0.86 \pm 0.42$ with an average Angstrom exponent $\sim 0.68 \pm 0.35$. The average monthly AOD and Angstrom exponent throughout the year are found to be in the range 0.56 to 1.22 and 0.38 to 0.96, respectively.

2. The monthly variations of aerosols over Delhi have three characteristic features: (a) During pre-monsoon period high AOD values associated with very low $\alpha$ values are observed showing dominance of desert aerosols, (b) in the post monsoon to winter period moderate AOD values are associated with the moderate to high $\alpha$ values, indicating dominant anthropogenic aerosols in the atmosphere, and (c) during autumn season moderate to low AOD are associated with the moderate to high $\alpha$ values, indicating mixed aerosol over Delhi.

3. The monthly average SSA over Delhi is found to be in the range 0.70 to 0.89 which is largely affected by the BC concentration that lies in the range $4$–$15 \text{ mg m}^{-3}$ as monthly average during daytime.

4. When the dust concentration was highest (May–June) the SSA increased with wavelength, however, when dust concentration was low the SSA decreased with the wavelength.

5. The monthly average clear-sky DARF at the surface varied in the range $-45 \text{ W m}^{-2}$ to $-110 \text{ W m}^{-2}$ throughout the year. The value of monthly DARF at TOA was found to be in the range $-1$ to $21 \text{ W m}^{-2}$ and in the atmosphere it was in the range $46 \text{ W m}^{-2}$ to $115 \text{ W m}^{-2}$ throughout the year.

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