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Environmental monitoring of surface ozone and other trace gases over different time scales: chemistry, transport and modeling

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Abstract Increasing concentration of tropospheric ozone (O₃) is a serious air pollution problem faced commonly by the urban people. The present study emphasizes on variations of air pollutant concentrations viz., O₃, nitrogen oxides (NO_x), carbon monoxide (CO), sulfur dioxide (SO₂) and black carbon (BC) at a tropical urban site located in the Deccan plateau region with semi-arid climate. The air monitoring site revealed typical diurnal/seasonal trends attributing to the complex chemistry of surface O₃ formation from its precursors. Role of SO₂ in the formation of free radical (HO₂) and its impact on O₃ concentration is distinguished part of the study. The results showed the highest mean O_3 in summer (57.5 \pm 15.2 ppbv) followed by winter and monsoon. Observations of BC aerosols showed the highest mean value during winter (8.2 \pm 2 μ g m⁻³) and the lowest in monsoon (4.2 \pm 1 μg m⁻³). Besides local influences, long-range transport of air masses were also studied by simulating back trajectories at different elevations during the study period. Furthermore, statistical analysis and modeling was performed with both linear (regression) and nonlinear (neural network) methods.

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Introduction

In the present century, urban areas deprive a healthy air quality environment due to the increase in concentration of tropospheric air pollutants such as sulfur oxides (SO_x), NO_x, CO and O₃. This decrease in air quality is a consequence of accumulation, dispersion and transformation of these air pollutants (Mazzeo et al. 2005). Increased demographic rates and rapid industrialization as well contributed for emissions with high concentrations of air pollutants (Rama Krishna et al. 2005). Air pollution caused by photochemical oxidants (O₃ and NO_x) is one of the serious problems faced by urban areas (WHO 2005). Surface O₃ formed shows detrimental effect on vegetation, human health and various materials (Selvaraj et al. 2010). Abdul-Wahab et al. (1996) reported that anthropogenic sources are responsible for more than 95 % of the O₃ in the lower atmosphere.

 O_3 is a secondary air pollutant and is one of the green house gases in troposphere which causes global warming (Sánchez et al. 2008). O_3 is not emitted directly by any natural source, but is indeed formed through a set of photochemical reactions involving primary precursors [NO_x, CO, volatile organic compounds (VOCs) etc.] under high solar radiation flux conditions (Shavrina et al. 2010). These precursor gases are emitted directly from the agricultural fields and indirectly from oxidation of fossil fuels, biomass burning and anthropogenic activities (Swamy et al. 2012a).

On the other side, destruction of O_3 takes place through several pathways, mainly through surface deposition. Scavenging processes dominate the removal of O_3 by NO





titration and SO₂ oxidation (Abdul-Wahab et al. 2005; Iliadis et al. 2007). Menon et al. (2002) and Akimoto (2003) reported O₃ destruction by air-borne particulates. Black carbon is one such primary aerosol species emitted into the atmosphere through a variety of incomplete combustion of fossil fuels. Role of BC in O₃ reduction and a major contributor to global warming was reported by Latha and Badarinath (2004).

Ground-based monitoring is important in many aspects viz., to clarify local/regional specific sources and sinks of trace gases, to derive the dynamic behavior of air pollutants and to check compliance of statistical models. These models are useful tools for air pollution and climatic studies (Arteta and Cautenet 2007). Furthermore, these models help in the development of an environmental policy, in particular to GHGs emission on a local and regional scale (Shavrina et al. 2010). But, forecasting air pollutant concentration is a difficult task due to complexity of physical and chemical processes involved. By applying multiple linear regressions (MLR), several functions can be fitted to the pollution data in terms of selected predictors. This approach is generally suitable for description of complex site-specific relations between concentrations of air pollutants and potential predictors (Abdul-Wahab et al. 2005; Swamy et al. 2013a).

Recently, artificial neural networks (ANN) model is the focus of attention, as they can handle complex and nonlinear problems better than conventional statistical techniques. Neural network is simple input-output mathematical model which extracts maximum information and reflects out the best prediction. These neural networks are noted to outperform linear regression, because it faces serious difficulties like multi-collinearity (Gardner and Dorling 1999). For regression, functional form is assumed first, such as linear or exponential, and then their coefficients minimize some measure of errors, whereas for neural networks, the method itself extracts functional form from data (McAdams et al. 2000).

In this study, environmental monitoring of surface level air pollutants inclusive of meteorological parameters was carried out at an urban site in Hyderabad, India, during the year 2010. The study observed typical temporal distribution of surface O_3 and other trace gases over different time scales. An attempt was also made to understand the chemistry of O_3 formation from its precursors and correlation of O_3 with trace gases during different seasons. In order to know the possible transport pathways of trace gases and aerosols from their potential sources of origin, the trajectory of a hypothetical air parcel into the study site was examined. Statistical assessment between O_3 and other variables was made using Pearson's correlation. Furthermore, O_3 prediction was performed using two modeling strategies viz., linear regression and neural networks.

Materials and methods

The air monitoring station is located at Tata Institute of Fundamental Research-National Balloon Facility (TIFR-NBF) (17.47°N and 78.58°E) Hyderabad, Andhra Pradesh, India. The study site is situated on a Deccan plateau at an altitude of 536 m a.m.s.l. Hyderabad is the capital of Andhra Pradesh state and is fifth largest metropolitan city in India, with population of about 8 million (http://www. censusindia.gov.in). Climate of Hyderabad is a semi-arid type with hot summer from March to early June, followed by monsoon from late June to October and a pleasant winter from November to February. An average rainfall of 810 mm is observed every year. Air masses during monsoon months arrive from southwest directions, while in the other seasons wind flows from southeast direction. The maximum air temperature was observed in May (summer) and minimum in December (winter). Geographical view of the observatory (http://maps.google.co.in/maps) illustrates many industries in south, southeast and southwest directions. Vehicular traffic in Hyderabad is a major contributor to air pollution load during dawn and dusk time. The total vehicular pollution load in the city is about 1,500 T/day. Out of the total vehicular emissions, CO, hydrocarbons, NO_x and SO_2 are 61.4, 34, 3.85 and 0.54 %, respectively (http://www.aptransport.org/html/pollution-control.htm).

Continuous monitoring of air samples were carried out for surface O₃, NO_x, CO and SO₂ during January to December in the year 2010 throughout the day with five-minute interval, and the data were retrieved using Envidas[®] software. The gas analyzers used were O₃ (Model 49i, Thermo Scientific, USA), NO_x (Model 42i, Thermo Scientific, USA), CO (Model 48i, Thermo Scientific, USA) and SO₂ (Model 43i, Thermo Scientific, USA). The O₃ analyzer works on the absorption principle, i.e., O₃ molecule absorbs ultraviolet (UV) light at 254 nm wavelength. The degree to which the UV light is absorbed is directly related to the O₃ concentration as described by the Beer-Lambert law. The O₃ lower detection limit is 1 ppbv with the response time of 20 s. The NO_x analyzer works on the principle of chemiluminescence, where nitric oxide reacts with O₃ to produce a characteristic light, whose intensity is linearly proportional to the NO concentration. Lower detectable limit of NO_x analyzer is 0.40 ppbv, with response time of 40 s. The CO analyzer works on the principle of non-dispersive infrared absorption. The lower detection limit of the CO analyzer is 40 ppbv, and the response time is 30 s. The SO₂ analyzer operates on the principle of light absorption, where the SO₂ molecules are excited by absorbing UV light at one wavelength and later decay to a lower energy state by emitting UV light at a different wavelength which is proportional to the SO₂ concentration. The UV wavelengths emitted by the excited SO₂ is detected by the photomultiplier tube (PMT). The lower





detection limit of the SO₂ analyzer is 2 ppby, and the response time is 80 s. Accuracy of the instruments is sustained by calibrating every week. All the analyzers were zero calibrated with dry air. Span calibration of O₃ analyzer was carried out using multi-point internally assembled O₃ generator. Span calibration for NO_x, CO and SO₂ was done using their respective NIST traceable standard gases through multi-point calibrator cum dynamic dilutor (Model 146i, Thermo Scientific, USA). Black carbon measurements were made using Aethalometer (AE-21, Magee Scientific), and its working details can be obtained elsewhere (http://www.mageesci. com). This instrument used is a self-contained, automatic instrument which requires no calibration other than periodic checks of the air flow meter response. Apart from these air pollutant observations, meteorological parameters viz., temperature (T), relative humidity (RH), wind speed (WS) and wind direction (WD) were measured using automatic weather station (AWS) installed at the site (TIFR-NBF). Solar radiation was recorded using Epply model 8-48 pyranometer. The 8-48 model pyranometer produces a millivolt analog signal that is directly proportional to the irradiance being measured. The instrument is provided with a calibration constant which, when divided into the signal, results in the irradiance in watts per square meter (Wm⁻²). Time-dependent studies and statistical analysis such as correlation, modeling and prediction were also carried out. The strength of O₃ with other measured variables was analyzed by Pearson correlation. Modeling and prediction of O₃ was performed using linear and nonlinear regression methods with observed O₃ as dependent variable and all other factors as independent variables.

Results and discussion

Temporal study of trace gases

The diurnal variations of trace gases measured during different seasons are shown in Fig. 1. The profile of surface O₃ showed a single-peak with double-trough pattern, which steadily increased from early hours of the day, then attained a peak value in the afternoon and thereafter declined in the evening. Daytime increase in O₃ concentrations is a pronounced feature of urban polluted site, because of the photochemical oxidation of precursors such as CO, and VOCs in presence of sufficient NO_x. Hyderabad being an urban location, NO_x concentrations are commonly observed above the threshold level (~ 10 pptv) which is conducive for O₃ formation (Lal et al. 2000). However, NO_x in the form of NO₂ produces O₃ by photolysis during sunlit hours, and NO in NO_x reduces O₃ during night time. This confirms the role of NO_x in the formation and reduction of O₃. Following are the conventional chemical reactions governing O_3 levels in the lower troposphere.

$$CO + NO + O_2 \rightarrow CO_2 + NO_2 \tag{1}$$

$$NO_2 + h\nu \rightarrow NO + O(^3P)$$
 (2)

$$O(^{3}P) + O_{2} + M(O_{2} \text{ or } N_{2}) \rightarrow O_{3} + M$$
 (3)

$$NO + O_3 \rightarrow NO_2 + O_2 \tag{4}$$

Maximum mean values of O_3 , NO_x , CO and SO_2 in daytime (06:00–18:00) and nighttime (19:00–05:00) during different seasons are given in Table 1. Difference in pollutant mixing ratio during the daytime was found to be significantly different from nighttime. The low O_3 concentrations observed during the night and early morning hours (Fig. 1) are due to the titration of O_3 with NO. Thereafter, O_3 concentration increased and reached to a peak value in the afternoon. Daytime change in O_3 level depends on the intensity of solar flux and boundary layer height (Lal et al. 2000). Increase in boundary layer height in the afternoon due to convective heating and consequent stratification, leads to the mixing of air in the lower altitude with the air in higher altitude rich in ozone concentration.

Besides, the other air pollutants viz., NO_x, CO, SO₂ (Fig. 1) as well showed a typical double-peak profile in reverse relation with O₃ concentration: high values in the morning (08:00-09:00) and at night (21:00-22:00) with low values in the afternoon. These two peaks appear to be associated with rush hour traffic. The morning peak is prominent due to the fumigation effect, a widely discussed phenomenon in the context of dispersion of pollutants (Stull 1988). During evening time, the boundary layer descends due to low temperature and become shallow, thereby restricting the vertical diffusion of anthropogenic air pollutants from lower troposphere causing localization and prolonged evening peak, until the traffic flow subsided at late night, causing the concentration to decrease, accordingly (Tsai et al. 2008). Low wind speed (<2 ms⁻¹) and high relative humidity (>60 %) might have also added up for weak diffusion of gases. However, during afternoon, concentrations of these pollutants reduced due to the increased boundary layer height causing vertical diffusion.

Surface SO_2 also influenced O_3 formation, and the chemistry involved is given in reactions (5–9). Role of SO_2 in O_3 formation is dependent on HO_2 produced from SO_2 (Swamy et al. 2013b). CO also acts as OH scavenger and helps in the formation of HO_2 leading to oxidation of NO. Thus, NO_2 formed again participates in O_3 formation during day time.

$$SO_2 + OH' + h\nu \rightarrow HOSO_2'$$
 (5)

$$HOSO_2^{\cdot} + O_2 + h\nu \rightarrow SO_3^{\cdot} + HO_2^{\cdot}$$
 (6)

$$HO_2^{\cdot} + NO \rightarrow NO_2 + OH^{\cdot}$$
 (7)

$$OH^{\cdot} + CO \rightarrow H^{\cdot} + CO_2$$
 (8)

$$H' + O_2 + M \rightarrow HO'_2 + M \tag{9}$$



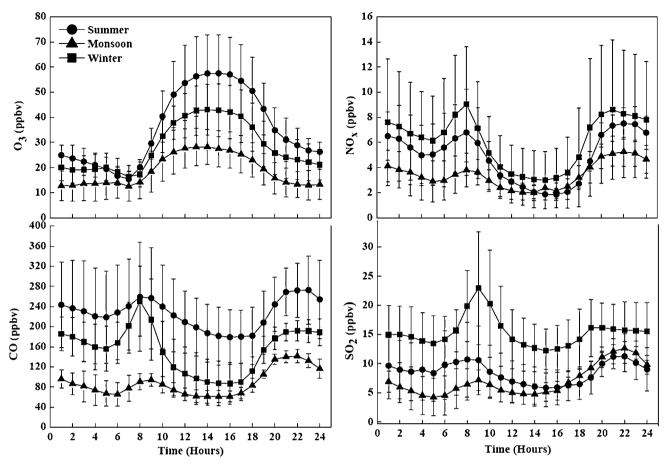


Fig. 1 Diurnal variations of O₃, NO_x, CO and SO₂ during different seasons

Table 1 Maximum mean along with SD values '±' of O₃, NO_x, CO and SO₂

Time (season)	O ₃ (ppbv)		NO _x (ppbv)		CO (ppbv)		SO ₂ (ppbv)	
	Day	Night	Day	Night	Day	Night	Day	Night
Summer	57.5 ± 15.2	50.5 ± 13.4	6.8 ± 3.4	7.5 ± 1.2	260.0 ± 108.0	273.0 ± 68.0	10.7 ± 6.4	11.2 ± 2.6
Monsoon	28.2 ± 7.0	19.4 ± 6.3	3.8 ± 1.3	5.3 ± 1.1	94.0 ± 12.6	141.0 ± 13.5	7.2 ± 1.8	12.6 ± 2.5
Winter	43.0 ± 10.0	29.5 ± 9.0	9.0 ± 4.6	8.6 ± 5.5	251.0 ± 70.0	192.0 ± 20.0	23.0 ± 10.0	16.2 ± 4.3

Day (06:00-18:00); night (19:00-05:00)

Besides diurnal variation, all the measured trace gases showed distinct monthly and seasonal differences (Figs. 1, 2). Seasonal mean concentration of O_3 was highest in summer (57.5 \pm 15.2 ppbv), i.e., in the months of March, April and May, because of the presence of intense solar radiation and high concentrations of precursors. During monsoon seasons, i.e., July, August and September showed low O_3 values (28.2 \pm 6.7 ppbv) due to wet deposition, low sunshine and high wind speed. In winter, the precursor trace gases are entrapped in lower troposphere under humid conditions supported by low wind speed, which resulted in considerably high O_3 concentrations (43 \pm 10 ppbv) than monsoon season.

Seasonal frequency distribution of O_3 during the period showed 90 % of O_3 lie in the range of 15–45 ppbv and the rest in the range of 50–60 ppbv. The amplitude of O_3 is highest in March, moderate in February and December and lowest in July. Sudden increase in amplitude during October is attributed to seasonal change in wind pattern from southwesterly to southeasterly. Wider amplitude observed during winter and summer months is mainly due to thermal inversion (Sánchez et al. 2008). Increase or decrease in amplitude is attributed to seasonal variations and related chemical transformations (Abdul-Wahab and Bouhamra 2004).

Besides O_3 photochemistry, the other factors that influence the temporal/spatial distribution of O_3 and its





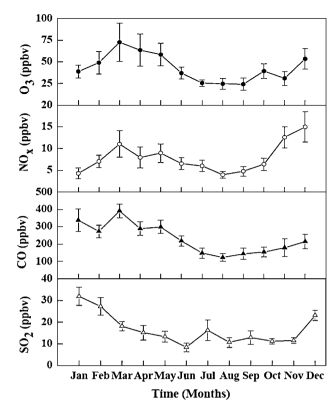


Fig. 2 Monthly variations of O₃, NO_x, CO and SO₂

precursors are concentration and emission rates of precursors ($NO_x/CO/VOCs$) (Kleinman 1991), nature of precursor emission sources (traffic/anthropogenic/biogenic) (Swamy et al. 2012b), boundary layer dynamics, meteorological parameters and heterogeneity of the terrain (i.e., geological site topography) (Jiménez et al. 2006).

Meteorological observations

Most of the atmospheric chemical reactions occur in presence of sunlight; specifically, the shorter wavelength of light with higher energy oxidizes primary pollutants such as NO_x into secondary air pollutants such as O₃. The rate of reaction is dependent on the downward solar flux which is proportional to the latitude-longitude, altitude, time of the day and other radiative factors (Walcek and Yuan 1995). Further, chamber experiments also suggested that the immediate NO₂ photolysis when exposed to irradiation resulted in formation of O₃, and the rate of O₃ formation correlates linearly with radiation flux (Lee et al. 2010). Therefore, measurement of solar radiation at the study site was important, and the observed surface O₃ concentrations showed a positive correlation with solar flux (Table 2). The mean solar radiation received during the entire study (Fig. 3) showed that summer recorded a maximum value of $1{,}116 \pm 94 \text{ Wm}^{-2}$. Low solar flux was observed in monsoon $(842 \pm 68 \text{ Wm}^{-2})$ followed by winter (828 ± 65)

Table 2 Pearson correlation matrix of O₃ with different variables

Variables	Summer	Monsoon	Winter
NO	-0.275**	-0.042	-0.221*
NO_2	-0.353*	-0.260**	-0.461**
CO	-0.241**	-0.237**	-0.438**
SO_2	-0.115^{*}	-0.066	0.164**
SR	0.517**	0.591**	0.644**
T	0.449**	0.547**	0.503**
RH	-0.082	-0.657**	-0.618**
WS	0.069	0.078*	0.508**
BC	-0.240*	-0.254**	-0.445**

^{*} Values with p < 0.005

Wm⁻²). Monthly variation (Fig. 4) showed a maximum mean in April and minimum mean in the month of July.

On diurnal basis (Fig. 3), O_3 showed a positive correlation with temperature, which is a measure of incident solar radiation and consequently the effectiveness of photochemistry (Sillman et al. 1990; Walcek and Yuan 1995; Tu et al., 2007). Annual maximum and minimum temperatures recorded were 42 °C on May 11, 2010 (15:00) and 13 °C on December 22, 2010 (07:00), respectively. Summer recorded a maximum mean temperature of 35 ± 3 °C, while in monsoon and winter, the maximum mean temperatures recorded were 28 ± 1 and 26 ± 4 °C, respectively. Month-wise temperature profile (Fig. 4) showed high values in March, April and May and low values in November and December.

Relative humidity in the atmosphere also played a crucial role in the formation and destruction of O_3 (Chen and Wang 2005), and the possible reactions are given below.

$$O' + H_2O \rightarrow 2OH' \tag{10}$$

$$H' + HO'_2 \rightarrow OH' + OH'$$
 (11)

$$OH' + O_3 \rightarrow HO_2' + O_2$$
 (12)

$$HO_2 + O_3 \rightarrow OH + 2O_2$$
 (13)

Along the diurnal scale, variation of RH depends on temperature (Fig. 3) and is anti-correlated with O_3 and temperature (Table 2). Annual maximum and minimum RH were recorded as 99 % on July 1, 2010 (22:00), and 17 % on April 25, 2010 (16:00), respectively. Highest RH mean of 83 ± 4 % was recorded in monsoon season, while in summer and winter RH recorded were 60 ± 7 and 76 ± 7 %, respectively. Monthly mean RH was highest in August and lowest in January (Fig. 4). David and Nair (2011) reported that monsoonal air hailing from the marine sources is rich in water vapor content and can cause reduction of O_3 Concentration by involving OH radicals. These observations were supported by the air mass back trajectories simulated at the study site, which is discussed in the later part of the text.



^{**} Values with p < 0.001

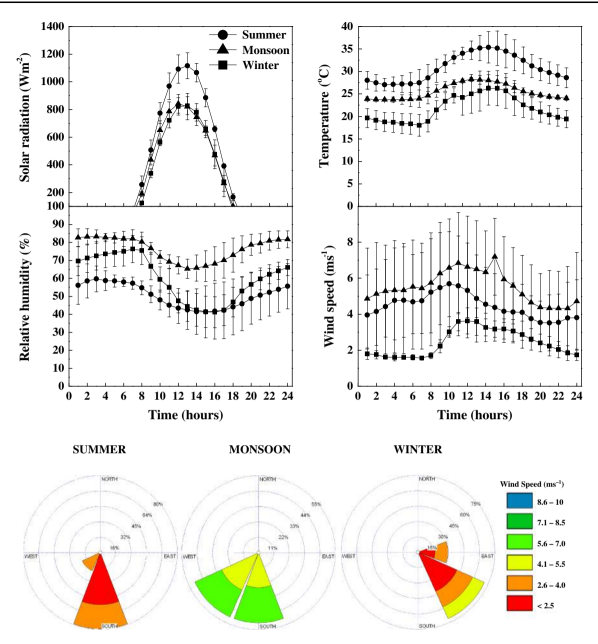


Fig. 3 Diurnal variation of solar radiation, temperature, relative humidity and wind velocity during different seasons

Wind speed and direction also affects trace gas concentrations. It characterizes mechanical turbulence causing dilution/concentration and transport of air pollutants, while wind direction determines the path of the pollutant dispersion (Duenas et al. 2002; Rodriguez and Guerra 2001). Seasonal and monthly variation of wind speed and direction are illustrated in Figs. 3 and 4, respectively. Summer showed calm winds from south direction. Monsoon showed strong winds from southwest direction. In winter, moderate winds prevailed from southeast direction.

Monthly profile of wind speed and direction showed notable changes during February–March, June–July and October–November due to seasonal transitions. A fair

estimate of dispersion of air pollutants in the atmosphere is possible based on the frequency distribution of wind velocity (Manju et al. 2002). Observations made in summer showed 56 % of calm winds (WS < 2.4 ms⁻¹) and 44 % of soft winds (2.4 \leq WS \leq 4.0 ms⁻¹) from the south direction. These calm winds can influence O_3 concentration at the site. Monsoon showed 75 % of strong winds (5.6 \leq WS \leq 8.5 ms⁻¹) and 25 % of moderate winds (4.1 \leq WS \leq 5.5 ms⁻¹) from southwest direction. Low O_3 concentrations was observed in monsoon due to the strong winds that dilute air by carrying O_3 precursors away from the site. In winter, a combination of calm (60 %), soft (30 %) and moderate winds (10 %) arrived from southeast direction.





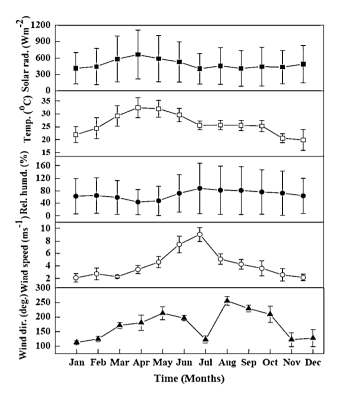


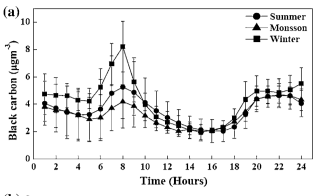
Fig. 4 Monthly variation of meteorological parameters

Therefore, the net O_3 formed is comparatively higher in winter than in monsoon, but lower than summer.

Black carbon

Diurnal pattern of BC mass concentration during different seasons (Fig. 5a) showed two peaks; first sharp peak in the morning (07:00–09:00) and second broad peak at night (19:00–21:00). These peaks are mainly attributed to vehicular traffic emissions. Gradual increase in BC concentrations after evening hours (17:00) was observed from domestic activities (Pathak et al. 2010) and surface-based inversion (Ganguly et al. 2006). The decrease in BC concentration in midnight hours is due reduction in traffic density and sedimentation (02:00–05:00 h). Also, fumigation effect in the boundary layer brings down BC aerosols from the nocturnal residual layer until sunrise and thereafter increases again with traffic (Tripathi et al. 2005).

Annual mean BC mass concentrations ranged between 2 and 10 $\mu g \ m^{-3}$ during the entire study period. Highest mean BC concentration was observed in winter (8.2 \pm 2 $\mu g \ m^{-3}$), followed by summer (5.3 \pm 3 $\mu g \ m^{-3}$), and lowest in monsoon (4.2 \pm 1 $\mu g \ m^{-3}$). High mean BC values in winter were due to entrapment of pollutant emissions in the shallow boundary layer. In addition, meteorological parameters such as low temperature and calm winds localize the pollutant concentration (Pathak et al. 2010). In monsoon, low values were recorded due to scavenging effect of rainfall (Kumar



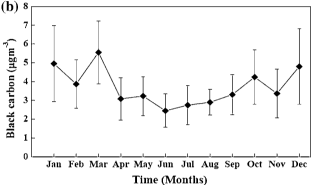


Fig. 5 a Diurnal variation of black carbon during different seasons, b monthly variation of black carbon

et al. 2011) and also due to mechanical turbulence of wind shear (Babu et al. 2002).

Monthly mean BC distribution profile (Fig. 5b) showed high values in January, March, October and December due to biomass burning and air mass transport. Minimum BC values were observed during June, July and August because of wet surface deposition by rain. It was observed that during the morning and evening peak traffic hours (Fig. 5a) high BC mass concentration showed significant O₃ reduction attributing to BC–O₃ heterogeneous chemistry (Latha and Badarinath 2004; Fendel et al. 1995), besides NO titration.

Transport of air pollutants

Air mass back trajectories are calculated at different elevations: 300, 800 and 1,500 m a.g.l such that the trajectory terminated at TIFR-NBF, Hyderabad at 12:30 IST to commensurate with the O₃ observations using the Air Resources Laboratory's Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (v.4.8) (http://www.arl.noaa.gov/ready/hysplit4.html).

Back trajectories were simulated in conjunction with seasonal change to understand the air mass transport and source of origin (Fig. 6). In summer, transport of trace gases occurred at heights <1,000 m a.g.l indicating its origin from local and also from regional surroundings



located in western and southwestern directions. Trajectories in the last week of summer (end of June) indicated onset of monsoon from Arabian Sea. Trajectories above 1,500 m a.g.l indicated BC aerosol transport from intercontinental areas. In monsoon, clean and fresh air originated from Arabian Sea (south-west direction), which implied marine transport. In winter, the air mass approached to the site was mainly of continental type. Venkataraman et al. 2005 reported that air parcel from western, eastern and central India contains trace gases and BC particles in high concentrations. Hence, these attributions made could be considered, which have impacted the change in air pollutant concentrations.

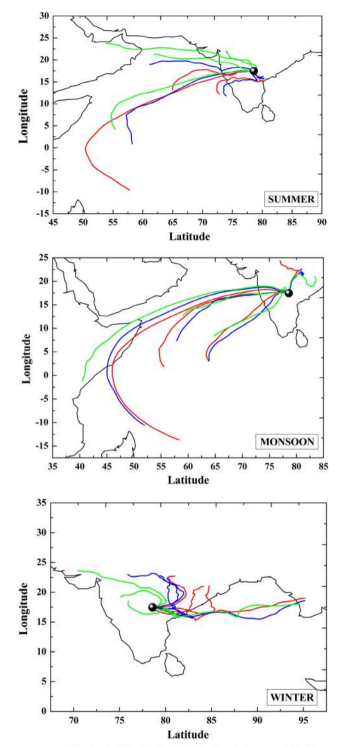
Statistical modeling

Surface O₃ being a secondary pollutant (dependent/response variable), its concentration is influenced by two major factors (independent variables or predictors) namely air pollutant concentrations (NO, NO2, CO, SO2 and BC) and meteorological conditions (SR, T, RH, WS). Assuming the predictors are often linearly correlated with response variable, a stepwise linear regression was employed. In this method, the variables are added iteratively to the model, until no additional variables contributed significantly to explain the variance of response variable. Before regression, the predictors were checked for variable inflation factor (VIF), which estimated multi-colinearity between dependent and independent variables. Only selective variables were considered for regression analysis. Among the total available data records, 1,050, 1,668 and 1,320 were used for summer, monsoon and winter, respectively. The regression process resulted in model equations (14, 15 and 16) and its statistics inferred that variables showed >50 % of overall variance in the observed O_3 for all three seasons (Fig. 7a).

$$\begin{aligned} \text{Summer} : O_3 \\ &= 28.9 - (3.66 * [\text{NO}]) - (0.0251 * [\text{CO}]) \\ &+ (0.0186 * \text{SR}) + (1.14 * \text{T}) \end{aligned} \tag{14} \\ \text{Monsoon} : O_3 &= 122 - (0.013 * [\text{NO}_2]) + (0.0290 * [\text{CO}]) \\ &+ (0.0118 * \text{SR}) - (1.93 * \text{T}) - (0.746 * \text{RH}) \\ &- (1.43 * \text{BC}) \end{aligned} \tag{15}$$

$$\begin{split} \text{Winter}: O_3 \\ &= 41.6 - (0.620 * [\text{NO}_2]) - (0.0165 * [\text{CO}]) \\ &+ (0.403 * [\text{SO}_2]) + (0.0107 * \text{SR}) \\ &+ (0.440 * \text{T}) - (0.276 * \text{RH}) \\ &+ (1.69 * \text{WS}) - (0.462 * \text{BC}) \end{split} \tag{16} \end{split}$$

However, multi-collinearity is always a serious problem with large data points with numerous variables. Although the predictors were checked for VIF value to minimize the multi-collinearity problem, a deeper look in the results still revealed the persistence of the same problem. This could be



300 m (red), 800 m (blue) and 1500 m (green) above ground level

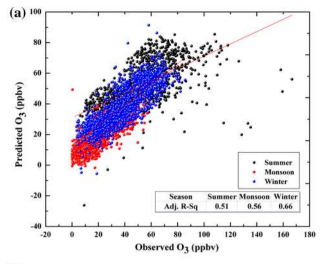
 $\begin{tabular}{ll} Fig. \ 6 \ Air \ mass \ back \ trajectories \ during \ different \ seasons \ and \ different \ heights \end{tabular}$

minimized if all possible conditions and factors influencing the response variable are considered in a nonlinear model.

An optimal nonlinear model using ANN was constructed with automatic architecture by employing







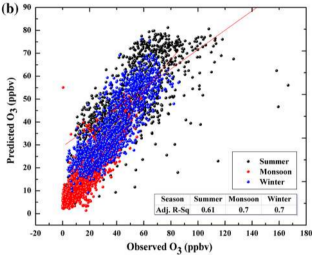


Fig. 7 Scatter plots of O₃—predicted versus observed using a multiple linear regression and **b** multi-layer perceptron

multilayer perceptron (MLP) method, which is commonly cited by many authors and elsewhere (Inal 2010; Kolehmainen et al. 2001). Out of the 4,038 totally available data records, 70 % of data records were used in training process and only 30 % were used for testing process. The model was evaluated with hyperbolic tangent (TanH) transfer function, in batch mode. Sensitivity analysis was carried out to correlate the maximum change in output with changes in the nth predictor. The ANN model used resulted in overall variance of >60 % (Fig. 7b), revealing the influence of NO_x, CO, solar radiation, temperature, relative humidity and black carbon on O₃ concentration.

In both the models, discrepancies between observed and predicted values were due to other factors that are not considered in this study (such as VOCs, dust, particulate matter, etc.). However, ANN model showed higher variance than linear regression. It is clear that the use of ANN model offers greater reliability and high degree of accuracy on the issue of local O_3 modeling using different

predictors, since O_3 formation is a nonlinear process involving complex reactions.

Conclusion

The study reported temporal variations of O₃ and its precursors (viz., NO_x, CO and SO₂) during different months/ seasons for the year 2010. It was observed that O₃ concentration varied with change in concentrations of precursors and local meteorology. Complex atmospheric chemistry of these trace gases has played a significant role in the formation of peroxide free radicals which were involved in O₃ formation. Back trajectories have shown transport of air masses from long distances with different sources of origin during various seasons which have affected air pollutant concentrations at our site. Furthermore, the model equations derived from statistical analysis using MLR and ANN revealed significant variables such as NO_x, CO, SR, T, RH and BC which influenced O₃ concentration. The predicted values obtained showed higher regression coefficients supporting nonlinear formation of O₃. The credible part of this work is to understand the role of O₃ precursor gases on the formation of O₃ with statistical analysis and modeling (linear and nonlinear approaches) at a semi-arid tropical urban site.

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