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Variations in O₃, CO, and CH₄ over the Bay of Bengal during the summer monsoon season: shipborne measurements and model simulations

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Abstract. We present shipborne measurements of surface ozone (O₃), carbon monoxide (CO), and methane (CH₄) over the Bay of Bengal (BoB), the first time such measurements have been performed during the summer monsoon season, as a part of the Continental Tropical Convergence Zone (CTCZ) experiment during 2009. O₃, CO, and CH₄ mixing ratios exhibited significant spatial and temporal variability in the ranges of $8-54 \text{ nmol mol}^{-1}$, $50-200 \text{ nmol mol}^{-1}$, and 1.57-2.15 μ mol mol⁻¹, with means of 29.7 \pm 6.8 nmol mol⁻¹, $96 \pm 25 \text{ nmol mol}^{-1}$, and $1.83 \pm 0.14 \text{ µmol mol}^{-1}$, respectively. The average mixing ratios of trace gases over BoB in air masses from central/northern India (O₃: $30 \pm 7 \text{ nmol mol}^{-1}$; CO: $95 \pm 25 \text{ nmol mol}^{-1}$; CH₄: $1.86 \pm 0.12 \,\mu\text{mol}\,\text{mol}^{-1}$) were not statistically different from those in air masses from southern India (O3: $27 \pm 5 \text{ nmol mol}^{-1}$; CO: $101 \pm 27 \text{ nmol mol}^{-1}$; CH₄: $1.72 \pm 0.14 \,\mu\text{mol}\,\text{mol}^{-1}$). Spatial variability is observed to be most significant for CH₄ with higher mixing ratios in the air masses from central/northern India, where higher CH₄ levels are seen in the SCIAMACHY (SCanning Imaging Absorption spectroMeter for Atmospheric CartograpHY) data. O₃ mixing ratios over the BoB showed large reductions (by $\sim 20 \text{ nmol mol}^{-1}$) during four rainfall events. Temporal changes in the meteorological parameters, in conjunction with O₃ vertical profile, indicate that these low-O₃ events are associated with downdrafts of free-tropospheric O3-poor air masses. While the observed variations of O₃ and CO

are successfully reproduced using the Weather Research and Forecasting model with Chemistry (WRF-Chem), this model overestimates mean concentrations by about 6 and 16% for O₃ and CO, respectively, generally overestimating O₃ mixing ratios during the rainfall events. An analysis of modelled O₃ along air mass trajectories show mean en route O_3 production rate of about 4.6 nmol mol⁻¹ day⁻¹ in the outflow towards the BoB. Analysis of the various tendencies from model simulations during an event on 10 August 2009, reproduced by the model, shows horizontal advection rapidly transporting O₃-rich air masses from near the coast across the BoB. This study fills a gap in the availability of trace gas measurements over the BoB and, when combined with data from previous campaigns, reveals large seasonal amplitude (~39 and ~207 nmol mol⁻¹ for O_3 and CO, respectively) over the northern BoB.

1 Introduction

Tropospheric ozone (O₃) is the third most important greenhouse gas, contributing to global warming and climate change with a radiative forcing of 0.40 ± 0.20 Wm⁻² (IPCC, 2013). O₃ is also a pivotal trace gas in tropospheric chemistry, as it is a major source of hydroxyl radical (OH), which removes most of the organic compounds and pollutants from

the atmosphere and controls the oxidation capacity of the troposphere (e.g. Brasseur et al., 1999; Finlayson-Pitts and Pitts, 2003; Seinfeld and Pandis, 2006). Further, enhanced concentrations of surface O₃ have detrimental effects on human health and vegetation (Heagle, 1989; Seinfeld and Pandis, 2006). Approximately 80% of tropospheric O₃ is produced by in situ photochemical reactions in the presence of nitrogen oxides ($NO_x = NO + NO_2$) involving the precursor gases of methane, non-methane hydrocarbons (NMHCs), and CO (Fishman et al., 1979; Crutzen et al., 1999; Seinfeld and Pandis, 2006). The remaining 20 % of tropospheric O₃ is attributed to intrusions of stratospheric air during frontal activities or to tropopause folding events (Lelieveld and Dentener, 2000; Sprenger et al., 2007). Depending upon meteorological conditions and the availability of the aforementioned precursors, a net production or destruction of O₃ prevails. The average lifetime of O₃ is about 1 week in the lower troposphere, which leads to large variability in its spatial and temporal distributions, as compared to the long-lived greenhouse gases. The budget of tropospheric O₃ and its implications for human health, crop yields, and climate are, however, not yet well quantified, especially over regions in Asia. This is mainly due to a lack of in situ measurements (e.g. Cooper et al., 2014; Monks et al., 2015).

Carbon monoxide (CO) is an indirect greenhouse gas which also has adverse effects on the health of humans and animals (WHO, 1999). Although it does not have a direct greenhouse effect like methane and carbon dioxide, its role in atmospheric chemistry is estimated to cause an indirect radiative forcing of 0.23 (0.18–0.29) Wm⁻² (IPCC, 2013). The major sources of CO are fossil fuel combustion, biomass burning, and oxidation of hydrocarbons such as CH₄ and isoprene (e.g. Jacob, 1999; Bergamaschi et al., 2000; Seinfeld and Pandis, 2006).

Methane (CH₄) is one of the major greenhouse gases, with a direct radiative forcing of $0.48 \pm 0.05 \text{ Wm}^{-2}$ (IPCC, 2013). This gas plays a major role in the climate and in atmospheric chemistry. CH₄ is emitted from variety of natural and anthropogenic sources (Jacob, 1999) and is removed primarily through its reaction with OH radicals (Fung et al., 1991; Seinfeld and Pandis, 2006).

The marine regions adjoining South Asia have been observed to have elevated levels of surface O_3 due to the outflow of continental pollution (Lawrence and Lelieveld, 2010) and minimal chemical loss by titration (e.g. Lal and Lawrence, 2001; Ojha et al., 2012). Suggested sources for this elevated O_3 and other trace gases observed over the marine regions surrounding India are anthropogenic, biomass burning, and biogenic emissions over continental India (Naja et al., 2004; Lawrence and Lelieveld, 2010; Nair et al., 2011; David et al., 2011). The air masses influenced by continental emissions undergo chemical transformation, including O_3 production, during their transport to the cleaner marine regions. In situ measurements over the marine regions are required to understand the effects of direct outflow, en route chemical transformation, and the chemistry in the transported air masses (Lawrence and Lelieveld, 2010, and references therein).

The experiments that have been conducted to date over the marine environment adjacent to the Indian region have revealed considerable spatial heterogeneity in the distribution of trace gases and aerosols, influences from source regions such as the Indo-Gangetic Plain (IGP), and radiative impacts (Nair et al., 2010, 2011; David et al., 2011; Mallik et al., 2013; Moorthy et al., 2009). Observations made during the Indian Ocean Experiment (INDOEX; Lal and Lawrence, 2001) and model simulations (Ojha et al., 2012) both found the O₃ mixing ratios over these remote marine regions to be even higher than those over the upwind continental regions due to complex O₃ chemistry. Lawrence and Lelieveld (2010) provided a detailed review of the outflow of trace gases and aerosols from South Asia to the surrounding marine regions. Transport of air masses between the Indian subcontinent and the adjacent marine regions has strong seasonal dependence associated with the monsoonal circulation (e.g. Kumar et al., 2015a).

The marine environment of the Bay of Bengal (BoB), the largest bay in the world, is surrounded by landmasses on three sides, making it highly suitable to observe enhanced concentrations of trace species. Further, seasonal changes in synoptic winds make this a unique region to study variations in trace species due to transport and en route photochemistry. Considering the aforementioned special characteristics of the BoB, as well as the considerable heterogeneity of trace gas and aerosol distribution, in situ measurements covering large areas are essential for investigating the distribution of pollutants and the controlling processes. Extensive in situ measurements of various trace gases over the BoB have been conducted in the following field campaigns: IN-DOEX during the winter months of 1998 and 1999 (Lelieveld et al., 2001; Mühle et al., 2002); the Integrated Campaign for Aerosols, gases, and Radiation Budget (ICARB) during the March-May (pre-monsoon season) of 2006 (Nair et al., 2011; Srivastava et al., 2011, 2012); the winter-ICARB (W-ICARB) during December-January 2009 (Girach and Nair, 2010, 2014; David et al., 2011); the Bay of Bengal Experiment (BOBEX)-I during February-March 2001 (Lal et al., 2006); the Bay of Bengal Process Studies (BOBPS) during September-October 2002 (Sahu et al., 2006); BOBEX-II during February 2003 (Lal et al., 2007); and the other campaign conducted during October-November 2010 (Mallik et al., 2013).

Although earlier studies have covered the spatio-temporal distribution of trace gases during most seasons over the BoB, there is still a lack of observations over the BoB during the summer monsoon season (June–August). The Asian summer monsoon circulation provides a pathway for pollution transport into the stratosphere (Randel et al., 2010), and observations taken during monsoon season capture a time of high water-vapour loading over the BoB. Deep convection during

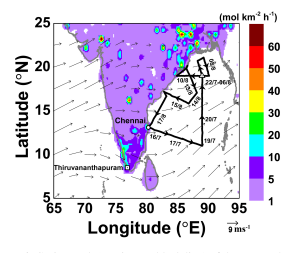


Figure 1. Cruise track (continuous black line) of the Research Vessel *Sagar Kanya* along with synoptic winds at 925 hPa (black thin arrows) and NO_x emissions in background colour map. The dates corresponding to approximate ship positions are marked along the track. The start and end position of the cruise, stationary position of the ship, and Thiruvananthapuram are shown by the circle, triangle and square, respectively.

the summer monsoon can uplift boundary layer pollution to higher altitudes; the pollution is then distributed over a larger region, thereby influencing air quality and climate over much larger regions (Lawrence and Lelieveld, 2010), extending as far as, for example, over the Mediterranean (e.g. Lelieveld et al., 2002; Scheeren et al., 2003). Such in situ measurements are also essential given the fact that satellite remote sensing of boundary layer O_3 has relatively higher uncertainty. The uncertainties in satellite retrievals of trace species are particularly high during the summer monsoon season, as the view of satellite instruments is frequently obscured by thick clouds.

In the present paper, the ship-based measurements of surface O_3 , CO, and CH₄ over the BoB are presented for the summer monsoon season of 2009. These observations were carried out as a part of the Continental Tropical Convergence Zone (CTCZ) experiment (http://www.incois.gov.in/portal/ datainfo/pdctcz.jsp) under the Indian Climate Research Programme (ICRP) of the Government of India. In this study, the spatial and temporal variations of O_3 over the BoB and the effects of transport are analysed. These observations are compared with simulations from a regional model, Weather Research and Forecasting coupled with Chemistry (WRF-Chem). The sharp reductions observed in O_3 during rainfall events are investigated in greater detail.

2 The cruise track and background conditions

Figure 1 shows the cruise track of the Oceanic Research Vessel (ORV) *Sagar Kanya* during the CTCZ campaign (cruise number SK 261). The arrows marked on the track show the direction of the ship, which sailed from Chennai (80.3° E,

13.1° N; marked by a circle) on 16 July 2009. The cruise offered greater coverage in the northern BoB than the southern or central BoB areas. To take time series measurements, the ship was kept stationary for 15 days (22 July to 6 August 2009) at 89° E, 19° N as marked by a triangle in the figure. After several tracks, covering latitude sector 11.0 to 21.1° N and longitude sector 80.3 to 90.1° E, the cruise ended on 17 August 2009 at Chennai, for a total of 32 days of voyage. The average wind pattern at 925 hPa (NCEP/NCAR reanalysis; http://www.esrl.noaa.gov/psd) during the cruise period is shown in Fig. 1. The prevailing westerly and southwesterly winds transport O_3 and its precursors from the Indian landmass to the BoB during the study period. The spatial distribution of emissions of NO_x , an O_3 precursor gas, is also shown as colour map in Fig. 1. NO_x emissions are obtained from the Intercontinental Chemical Transport Experiment Phase B (INTEX-B) inventory (Zhang et al., 2009), which is representative of the year 2006. NO_x emissions are relatively higher over parts of eastern and southern India as compared to central India. The square tagged as Thiruvananthapuram shows the location corresponding to the measurements shown in Fig. 9.

3 Experimental details and data

Surface O_3 measurements were carried out using an online ultraviolet (UV) photometric ozone analyzer (model O3 42), manufactured by Environnement S.A, France. The analyser utilises the absorption of UV radiation by O_3 molecules at 253.7 nm and derives O_3 mixing ratios using the Beer-Lambert law. This UV absorption-based analyser has an uncertainty of about 5% (Tanimoto et al., 2007), corresponding to ~ 1.5 nmol mol⁻¹ for the observed range of O_3 . Zero noise of the instrument is 0.5 nmol mol⁻¹. The instrument has a lower detection limit of 1 nmol mol⁻¹ and a linearity of ± 1 %. An individual measurement is performed at a minimum response time of 10 s. The analyzer was operated on auto-response mode, whereby responses could be 10–90 s depending upon changes in O_3 mixing ratios. However, data were recorded continuously at 5 min intervals.

CO measurements were made using an online CO analyzer (model CO12 Module) manufactured by Environnement S.A, France. This instrument works on the principle of nondispersive infrared (NDIR) absorption by CO molecules at the wavelength of 4.67 µm. The instrument has a lower detection limit of 50 nmol mol⁻¹, a linearity of 1 %, and a response time of 40 s. The overall uncertainty in hourly CO measurements is estimated to be ~ 10 % at a CO value of 150 nmol mol⁻¹ (Sawa et al., 2007; Tanimoto et al., 2007).

Air was drawn from a height of approximately 15 m above the sea surface through a Teflon tube. Before and after the cruise, both analyzers were calibrated, with calibration factors not found to be significantly changed. The calibrations of both analysers were carried out using appropriate calibration

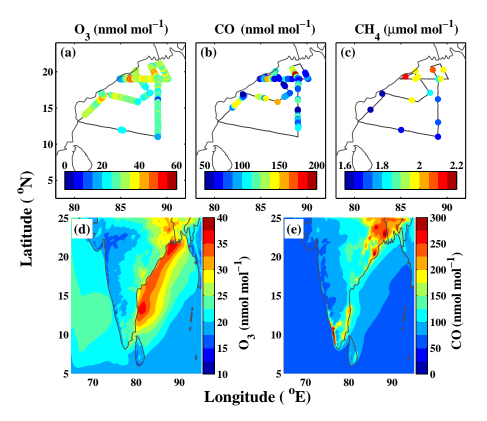


Figure 2. Spatial variation of surface O_3 (a), CO (b), and CH₄ (c) mixing ratios along the cruise track during the CTCZ campaign. WRF-Chem simulated spatial distribution of surface O_3 (d) and CO (e) averaged during 16 July–17 August 2009.

standards traceable to NIST and a multi-channel calibrator, following the procedure mentioned in the manuals of analysers. While the O₃ analyser was calibrated for mixing ratios of 30 nmol mol⁻¹, the CO analyser was calibrated for mixing ratios of 1.1 µmol mol⁻¹. Meteorological parameters such as pressure, temperature, and relative humidity were measured continuously on board the ship. Trace gas measurements affected by the ship exhaust were identified and discarded using on-board wind direction and NO_x measurements.

In addition, a total of 29 air samples were collected in 1L glass flasks during the cruise and were analysed for CH₄ using a gas chromatograph (GC) coupled with a flame ionisation detector (FID), as described in Tiwari and Ravi Kumar (2011). These CH₄ measurements are traceable to the WMO standard scale. CH₄ standards were obtained from the WMO Central Calibration Laboratory (CCL) at the National Oceanic and Atmospheric Administration (NOAA)/Earth System Research Laboratory (ESRL)/Global Monitoring Division (GMD), located in Boulder, Colorado, USA. The precision for CH₄ measurements was approximately $\pm 0.1 \,\mu$ mol mol⁻¹. A detailed description of the analytical procedure for CH₄ measurement and calibration of GC is given in Ravikumar et al. (2014).

To further study the observed low- O_3 events over BoB, measurements made at Thumba, Thiruvananthapuram, are

used as a case study. Using the same O_3 analyzer as the one used for surface O_3 measurements over BoB, continuous measurements of surface O_3 were taken at Thumba, Thiruvananthapuram (David and Nair, 2011; Girach et al., 2012), in July 2011. Along with various meteorological parameters, rainfall measurements were also taken at Thumba at 5 min of integration time using an automatic weather station manufactured by Dynalab Weathertech Pvt. Ltd, India. The site, Thumba, is situated just ~ 500 m away from the west coast, with sandy terrain, and is a less populated area in the city of Thiruvananthapuram (8.5° N, 76.9° E) at southern tip of India. For more details about the Thumba site and measurements please see, for example, Nair et al. (2002) and David and Nair (2011).

A vertical profile of O_3 was measured on 28 July 2011 at Thumba using an electrochemical concentration cell ozonesonde (EN-SCI 2ZV7 ECC; Komhyr, 1969; Komhyr et al., 1995). The accuracy of such ozonesondes is reported to be about ± 5 –10 % up to ~ 30 km (Smit et al., 2007). More details of this measurement technique can be found in Ojha et al. (2014).

The accumulated rainfall for every 3 h interval from the Tropical Rainfall Measuring Mission (TRMM; with a horizontal grid size of $0.25^{\circ} \times 0.25^{\circ}$) is also utilised in this study to complement the on-board rainfall measurements.

The 3B42 algorithm is used to calculate precipitation and root-mean-square precipitation-error estimates; these two estimates were then used to compute hourly and daily rainfall estimates (Huffman et al., 1995).

The gridded $(2^{\circ} \times 2^{\circ})$ monthly column-averaged CH₄ (level 3, version 6) retrievals from SCIAMACHY (SCanning Imaging Absorption spectroMeter for Atmospheric CartograpHY) instrument on board the Envisat satellite were used to infer concentrations over Indian land regions. The IMAP-DOAS (iterative maximum a posteriori differential optical absorption spectroscopy) algorithm was used, which retrieves CH₄ utilising the spectra (i.e. 1000–1750 nm) from the near infrared channel no. 6 (Frankenberg et al., 2005).

4 Model simulations

WRF-Chem (Grell et al., 2005) version 3.5.1 was used to simulate meteorological and chemical fields during the campaign period. The model domain (Fig. 2d-e) is defined on the Mercator projection, centred at 80° E, 15.5° N, at a spatial resolution of $15 \text{ km} \times 15 \text{ km}$. The model has 51 vertical levels from surface to 10 hPa. The simulations were conducted for the period of 29 June to 31 August 2009, covering the complete measurement period. The meteorological inputs have been adopted from ERA-Interim reanalyses by the ECMWF. Horizontal winds, temperature, and water vapour are nudged above the planetary boundary layer using a nudging coefficient of 0.0003 s^{-1} (Kumar et al., 2015b), employing the four-dimensional data assimilation (FDDA) technique. Anthropogenic emissions of CO, NO_x , SO₂, and NMVOCs are provided by a regional emission inventory that was developed to support INTEX-B (Zhang et al., 2009; Kumar et al., 2012b; Ojha et al., 2016). This inventory is representative of the year 2006. Aerosol emissions are provided by the Hemispheric Transport of Air Pollution (HTAP v2) inventory (Janssens-Maenhout et al., 2015). Biomass burning emissions from NCAR Fire Inventory (FINN; Wiedinmyer et al., 2011) and biogenic emissions calculated online using MEGAN (Guenther et al., 2006) were used in the simulations.

Gas-phase chemistry in the model is represented by the second-generation Regional Acid Deposition Model (RADM2; Stockwell et al., 1990), and the aerosol module is based on MADE SORGAM (Binkowski and Shankar, 1995; Ackermann et al., 1998; Schell et al., 2001). Initial and boundary conditions for chemical fields are provided by the MOZART-4/GEOS5 data. The options used to parameterise different atmospheric processes are given in Table 1. For more information about meteorological nudging, chemical mechanisms, emissions, boundary conditions, and evaluation of WRF-Chem against in situ measurements and satellite data over the Indian region, please see, for example, Kumar et al. (2012a, b, 2015a, b), Ansari et al. (2016), and Ojha et al. (2016). Model-simulated mean spatial distributions of

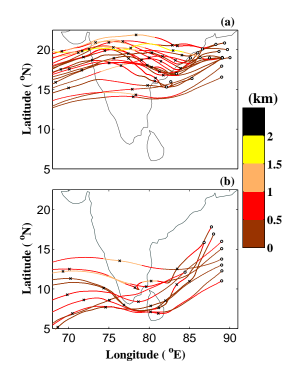


Figure 3. Five-day air mass back trajectories during the study period ending at the measurement locations (small black circles) grouped for corresponding air masses from (a) central/northern India and (b) southern India. The cross symbols along the trajectories represent each back-day. The colour scale shows the height (in km) of the trajectories.

 O_3 and CO over the model domain during the study period are shown in Fig. 2d–e.

5 Results and discussion

5.1 Variations in O₃, CO, and CH₄ over the BoB

Figure 2a-c show the observed variations in O₃, CO, and CH₄ along the ship track during 16 July to 17 August 2009. The mixing ratios of trace gases show large spatio-temporal variations over the BoB. Levels of O₃ and CO varied in the ranges of 8–54 (average of $29.7 \pm 6.8 \text{ nmol mol}^{-1}$) and 50– $200 \text{ nmol mol}^{-1}$ (average of $96 \pm 25 \text{ nmol mol}^{-1}$), respectively. As CO mixing ratios below the detection limit of the instrument are discarded from the analysis, the reported minimum and average values of CO mixing ratios are therefore slightly higher than their actual values. CH₄ mixing ratios are observed to range from 1.57 to $2.15 \,\mu\text{mol}\,\text{mol}^{-1}$, with an average of 1.83 ± 0.14 nmol mol⁻¹. Average CH₄ mixing ratios showed a significant difference of $\sim 0.14 \,\mu mol \, mol^{-1}$ between northern (81-91° E, 16-21.5° N) and central (80-91° E, 11-16° N) BoB during the study period. In addition to sailing across the BoB, the ship was also kept stationary for 15 days, from 22 July to 6 August 2009

Atmospheric process	Scheme used	Features of the scheme			
Cloud microphysics	Lin et al. scheme (Lin et al., 1983)	Sophisticated parameterisation including ice, snow, and graupel processes; suitable for high-resolution simulations.			
Longwave radiation	Rapid Radiative Transfer Model (RRTM; Mlawer et al., 1997)	Accurate scheme utilises look-up tables for efficiency and accounts for multiple bands and microphysical properties.			
Shortwave radiation	Goddard shortwave scheme (Chou and Suarez, 1994)	Two-stream multi-band scheme using O ₃ from climatology and includes cloud effects.			
Surface layer	Monin–Obukhov scheme (Janjic, 1996)	Based on Monin–Obukhov with Zilitinkevich thermal roughness length and standard similarity functions from look-up tables.			
Land surface option	Noah land surface model (Chen and Dudhia, 2001)	Unified NCEP/NCAR/AFWA scheme with soil temperature and moisture in four layers, fractional snow cover and frozen soil physics. This includes the modifications for better representation of processes over ice sheets and snow covered areas.			
Urban surface physics	Urban canopy model	Three-category urban canopy model with surface effects for roofs, walls and streets.			
Planetary boundary layer	Mellor–Yamada–Janjic scheme (Janjic, 2002)	One-dimensional prognostic turbulent kinetic energy scheme; local vertical mixing is included.			
Cumulus parameterisation	Grell 3D ensemble scheme (Grell, 1993; Grell and Devenyi, 2002)	Improved version of the GD scheme suitable for coarse as well as high-resolution simulations.			

Table 1. The WRF-Chem options used for parameterisation of atmospheric processes.

at 89° E, 19° N. During this time period, surface O_3 , CO, and CH₄ mixing ratios are observed to fall into the range of 9–46 nmol mol⁻¹, 58–144 nmol mol⁻¹, and 1.71–1.89 µmol mol⁻¹, respectively, with temporally averaged mixing ratios of 28 ± 7 nmol mol⁻¹, 91 ± 19 nmol mol⁻¹, and 1.81 ± 0.06 µmol mol⁻¹, respectively.

The HYbrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model was used to simulate 5-day backward air mass trajectories arriving at 500 m (a height that falls within the marine atmospheric boundary layer) above the measurement locations (Draxler and Rolph, 2003; Rolph, 2003; http://www.arl.noaa.gov/ready.html), as shown in the Fig. 3. Trajectories are colour-coded to show the altitude variations of the air parcels along their path. The influences of two different air masses are observed over the BoB during the CTCZ experiment. Over the central BoB, the backward air trajectories cross southern India (i.e. $< 13^{\circ}$ N), where a belt of elevated anthropogenic emissions $(5-20 \text{ mol km}^{-2} \text{ h}^{-1} \text{ of } \text{NO}_x$; see Fig. 1) is located. In contrast, most of the air trajectories over northern BoB come across the central Indian region, where anthropogenic emissions are relatively lower. For example, with the exception of a few hotspots, NO_x emissions north of $13^{\circ}N$ are in the range of $1-10 \text{ mol km}^{-2} \text{ h}^{-1}$ (Fig. 1). The O₃ and CO mixing ratios over BoB in air masses from central/northern India (Fig. 3a) are slightly higher or comparable $(O_3: 30 \pm 7; CO: 95 \pm 25 \text{ nmol mol}^{-1})$ to those $(O_3: 27 \pm 5;$ CO: 101 ± 27 nmol mol⁻¹) in air masses from southern India (Fig. 3b).

The observed spatio-temporal variations of trace gases are investigated by calculating the fractional residence time of air masses over land, using HYSPLIT simulated 5-day backward air trajectories. Figure 4a-c show the temporal variations of O₃, CO, and CH₄ during the CTCZ experiment along the cruise track. The percentage of time of air masses over continental India is also shown (Fig. 4d), as estimated by the ratio of residence time over land to the total trajectory time of 5 days. The hours of residence have only been included in the analysis if the altitude along trajectory is less than 1.5 km, as the surface emissions might not been directly influence the air masses aloft. Red vertical bars depict the sharp reductions in O₃ mixing ratios associated with rainfall events (see Sect. 5.3). O₃, CO, and CH₄ show correlated variability with the estimated residence times over the Indian subcontinent with slightly higher correlation in the case of primary species ($R^2 = 0.16$ in case of CO and CH₄), as compared to O_3 ($R^2 = 0.09$). Similar variations in mixing ratios of these trace gases and residence time over continental India indicate the influences of transport from the Indian subcontinent on the observed spatio-temporal variations over the BoB during the summer monsoon season. The occasions on which such a one-to-one correspondence are not observed can be attributed to varying source strengths, vertical mixing or dilution, and en route photochemistry.

Generally, during the summer monsoon season, relatively cleaner marine air masses from the Arabian Sea are transported to the Indian region. These air masses are then exposed to regional emissions and subjected to photochemistry

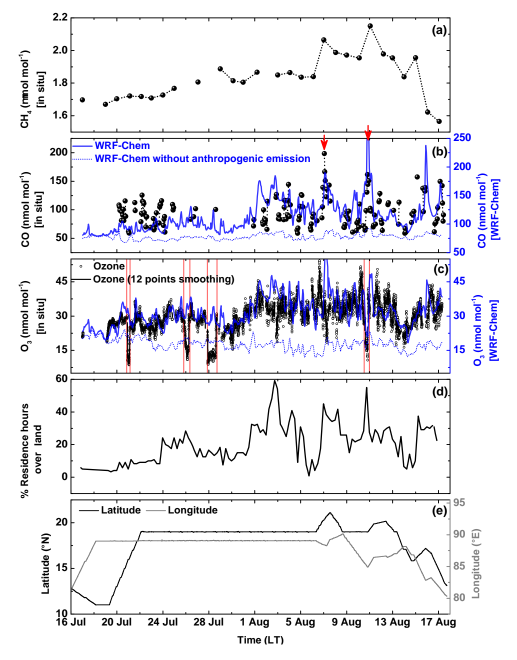


Figure 4. Variations in observed CH₄ (**a**), CO (**b**), O₃ (**c**), and percentage residence time over land (**d**) along with WRF-Chem simulated O₃ and CO (blue line) during the campaign. Events of sharp decrease in O₃ during rainfall are marked by vertical red lines (Fig. 8). (**e**) Variations in measurement locations.

depending upon the availability of solar insolation under the cloudy conditions of monsoon. The air masses in which precursors have accumulated, and to some extent photochemically processed, outflows into the BoB. As a result, the air masses outflowing at the eastern coast of India could have higher O_3 mixing ratios than the background air coming from the Arabian Sea into the western coast of India. The difference between these two values is representative of the O_3 build-up that can be attributed to regional pollution; this difference would also reflect the extent of photochemical processing that would have taken place.

As the observational site Thumba, Thiruvananthapuram, is situated just at the Arabian Sea coast, the monsoon-time observations here could be approximated to represent the background O_3 mixing ratios entering from the Arabian Sea. In August 2009, using only daytime monthly average O_3 , the O_3 at Thumba during the monsoon season was observed to be $23 \pm 7 \text{ nmol mol}^{-1}$. Since the objective of in-

Observation site	Longitude (° E)	Latitude (° N)	Observation period	Surface daytime O ₃ (mean ± standard deviation)	Reference
Arabian Sea					
Arabian Sea	69–76	9–19	July-August 2002	9	Ali et al. (2009)
Western coast of India					
Thiruvananthapuram	76.9	8.5	August 2009	23 ± 7	Present study
Thiruvananthapuram	76.9	8.5	June–August 2008	19 ± 6	David and Nair (2011)
Kannur	75.4	11.9	July 2010–2011	11 ± 4	Nishanth et al. (2014)
MtAbu (1.6 km a.m.s.l.)	72.7	24.6	August 1993–2000	25 ± 9	Naja et al. (2003)
Ahmedabad	72.6	23	July 1991–1995,	22 ± 8 ,	Lal et al. (2000);
			August 1991–1995,	17 ± 4 ,	Srivastava et al. (2012)
			July-August 2003-2007	25*	
Central India					
Anantapur	77.65	14.62	July 2009	30 ± 2	Reddy et al. (2011)
Eastern coast of India					
Bhubaneswar	86.4	20.5	June–August 2011–2012	29 ± 6	Mahapatra et al. (2014
Bay of Bengal					
Bay of Bengal	80.3–90.1	11-21.1	July-August 2009	30 ± 7	Present study

Table 2. A comparison of averaged surface O_3 mixing ratios measured at various sites during summer monsoon period.

* Boundary layer O3 over the Arabian Sea.

vestigation is the additional O₃ over the BoB produced by en route photochemistry, daytime O3 values at Thiruvananthapuram are therefore compared with all the observations over the BoB. The average mixing ratio observed over the BoB during monsoon season for 16 July-17 August 2009 was $30 \pm 7 \text{ nmol mol}^{-1}$, which was $\sim 7 \text{ nmol mol}^{-1}$ higher than the Arabian Sea air mass. This additional amount of \sim 7 nmol mol⁻¹ could be attributed to the effects of regional and en route photochemical O₃ production. Net O₃ production rate in the outflow is estimated to be in the range of $1.5-4 \text{ nmol mol}^{-1} \text{ day}^{-1}$. Note that the O₃ mixing ratio is reported to be $\sim 30 \pm 2$ nmol mol⁻¹ during July 2009 over Anantapur, a rural site in central India, indicating the enhancement due to regional O₃ production. As shown in Table 2, while average O₃ mixing ratios over the west coast of India and the Arabian Sea are in the range of 9- 25 nmol mol^{-1} during the monsoon season, the average O₃ mixing ratio is ~ 30 nmol mol⁻¹ over the central Indian station and the BoB.

O₃ mixing ratios were 27 ± 3 and 28 ± 5 nmol mol⁻¹ for 21 July 2009 and 15 August 2009, for which back trajectories (not shown here) crossed Thiruvananthapuram on 20 July and 13 August 2009, with daytime O₃ values of 23 ± 6 and 25 ± 6 nmol mol⁻¹, respectively. The difference of 3– 4 nmol mol⁻¹ between O₃ mixing ratios over the BoB and Thiruvananthapuram represents the en route photochemical production of O_3 in the air masses toward the observation points over the BoB. It is further found that the air masses were typically below 700 m and generally within the marine boundary layer (e.g. mean boundary layer height ~ 897 m during winter over the BoB; Subrahamanyam et al., 2012). The enhancements in O_3 are attributed here to in situ photochemical build-up while moving towards the BoB, which has been noted in previous experiments and model simulations (e.g. Lal and Lawrence, 2001; Ojha et al., 2012).

CO showed a sharp enhancement (denoted with red arrows in Fig. 4b) on 7 and 11 August 2009, coinciding with a longer residence time over the Indian region. Figure 5 shows backward air mass trajectories above the measurement locations, along with the distribution of anthropogenic CO emissions from the INTEX-B inventory, representative of the year 2006. The air masses over the BoB are found to be influenced by emission hotspots (corresponding emission of 250–350 mol km⁻² h⁻¹). The air masses took about half a day to be transported from the emission hotspot to the observation location over the BoB. The CO mixing ratio measured at Bhubaneswar (20.30° N; 85.83° E), a station within the hotspot region, is $\sim 251 \pm 58 \text{ nmol mol}^{-1} \text{ dur-}$ ing the monsoon season (June-August 2011-2012; Mahapatra et al., 2014), with the elevated CO emissions in the Bhubaneswar region being attributed to industrial activities. The higher CO mixing ratio $\sim 200 \,\mathrm{nmol}\,\mathrm{mol}^{-1}$ is in line

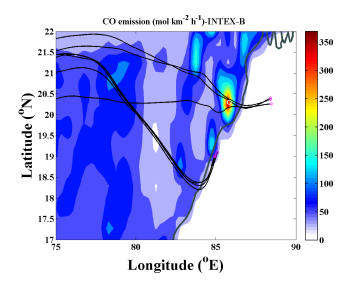


Figure 5. Backward air mass trajectories (black curves) 500 m above the location of higher CO observations as marked by red arrows in Fig. 4b during 7 and 11 August 2009. The background colour map shows the spatial distribution of anthropogenic CO emissions over the Indian region for the year 2006 from INTEX-B inventory. The small circles in magenta represent the points where observations were made, as well as the end point of trajectories. The white circle over the hotspot region denotes an observational site, Bhubaneswar.

with the monsoonal values observed at Bhubaneswar. The CO mixing ratios around $150 \text{ nmol mol}^{-1}$ were sampled on 11 August 2009 near the coastal source regions. Additionally, CO mixing ratios over central BoB (101 nmol mol⁻¹) were only slightly higher than those over northern BoB (95 nmol mol⁻¹). It is suggested that this is partially due to higher emissions over southern India against the shorter residence of air masses over land and the relatively longer lifetime of CO.

The mixing ratios of surface CH₄ were higher in the air masses from central/northern Indian over northern BoB $(1.86 \pm 0.12 \,\mu\text{mol mol}^{-1})$ as compared to those in the air masses from southern India $(1.72 \pm 0.14 \,\mu\text{mol mol}^{-1})$. As CH₄ is a relatively well-mixed trace gas, the average values over the tropospheric column approximates the uniform mixing ratio within the troposphere (Seinfeld and Pandis, 2006). The monthly column-averaged tropospheric CH₄, retrieved from SCIAMACHY, for August 2009 shows (Fig. 6) higher values around $1.85 \,\mu mol \, mol^{-1}$ over central/northern India as compared to that of southern India ($\sim 1.80 \,\mu\text{mol mol}^{-1}$). The higher tropospheric CH₄ over the central/northern Indian landmass during the summer monsoon season has been also reported by Kavitha and Nair (2016). The observed higher CH₄ over the northern BoB are attributed to the influences of emissions from central/northern Indian regions as also suggested by backward trajectories. Owing to the longer lifetime of CH₄, diffusion of CH₄ from a hotspot re-

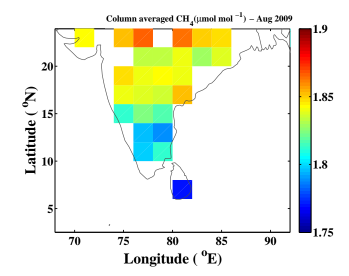


Figure 6. Spatial distribution of column-averaged CH₄ for the month of August 2009 as obtained from SCIAMACHY.

gion over the eastern IGP to northern BoB might be the other source of higher CH₄ levels over northern BoB during summer monsoon season. An emission inventory analysis by sector over the hotspot region (i.e. eastern IGP) indicates that these higher CH₄ emissions are associated with rice cultivation, waste treatment, and livestock. The correlation between the in situ CH₄ measurements and the retrievals from the satellite instrument (AIRS – Atmospheric Infrared Sounder) was found to be statistically insignificant (not shown), which highlights a need of more such in situ measurements in this region to validate the satellite products, especially during the summer monsoon.

5.2 WRF-Chem simulations

WRF-Chem simulations, as described in Sect. 4, are used to evaluate the performance of the model in reproducing the measurements and to investigate the underlying processes that caused the observed variabilities in O₃ and CO. A comparison between model-simulated and measured meteorological parameters shows only small mean biases, such as -1.9 hPa in pressure, -0.6 °C in temperature, and -1.1 % in relative humidity (Table 3). Figure 4b-c compare WRF-Chem simulated O₃ and CO with in situ measurements taken along the cruise track. WRF-Chem is found to reproduce the observed variations of O3 and CO over the BoB during the summer monsoon season with an overestimation of absolute O₃ levels by 1.9 nmol mol⁻¹ (i.e. $\sim 6\%$ of averaged O_3 value, 29.7 nmol mol⁻¹) and absolute CO levels by 18 nmol mol^{-1} (i.e. ~ 16 % of averaged CO value, 96 nmol mol^{-1}). It should be noted that the average CO mixing ratio of 96 nmol mol⁻¹ is slightly higher than its actual value, as data points below the detection limit of the instrument are discarded. Biases in the model simulations can be

Parameter	Observation	Model (WRF-Chem)	Mean bias	R^2
Pressure (hPa)	1001.3 ± 2.1	999.4 ± 2.2	-1.9	0.93
Temperature (°C)	29.3 ± 0.9	28.7 ± 0.6	-0.6	0.13
Relative humidity (%)	87.9 ± 4.2	86.8 ± 2.8	-1.1	0.36
$O_3 (nmol mol^{-1})$	29.7 ± 6.8	31.6 ± 6.6	1.9	0.58
$CO (nmol mol^{-1})$	96 ± 25	114 ± 30	18	0.19

Table 3. A comparison of mean values from observations with model-simulated parameters along with the mean bias. The squared correlation coefficients correspond to the linear regression analysis between daily averaged in situ and simulated parameters.

attributed to the uncertainties in the simulated meteorology and in the emission datasets; however, in the present study, the model fields are used mainly to investigate temporal variations rather than absolute mixing ratios. The squared correlation coefficients between the daily averaged in situ measured and simulated O₃ and CO are 0.58 and 0.19, respectively. The higher value of the squared correlation coefficient for O₃ demonstrates WRF-Chem's ability to reproduce the observed broad features in surface O₃ over the BoB. Note that the sharp reductions that caused very low O3 during rainfall episodes are not captured by WRF-Chem, except during the event of 10-11 August 2009. This will be discussed in detail in Sect. 5.3. WRF-Chem simulated O₃ has also been evaluated against several surface observations in India previously (Kumar et al., 2012b). Modelled O₃ was found to be within 1σ standard deviation of mean from observations at Gadanki in the southern India during summer monsoon (Ojha et al., 2016). At Thumba also, model-simulated O₃ variations correlated reasonably with measurements ($R^2 = 0.6$). More information on evaluation of WRF-Chem simulations of O₃ and CO over India can be found elsewhere (e.g. Kumar et al., 2012b; Ojha et al., 2016).

An additional simulation was conducted by switching off the anthropogenic emissions over the model domain (Fig. 4b–c; dotted blue curves). Mean O_3 (17.7 nmol mol⁻¹) is found to be lower by 14 with smaller variability of 2.4 nmol mol⁻¹ compared to standard WRF-Chem simulation. Similarly, also the mean CO level is lower by 36 with smaller variability of 4.9 nmol mol⁻¹. This shows that enhanced levels and observed variability in O_3 and CO mixing ratios over BoB are attributable to the regional anthropogenic emissions.

The limited collocated measurements showed en route O_3 production rate of 1.5–4 nmol mol⁻¹ day⁻¹, as discussed in Sect. 5.1. Here, the O_3 production rate is estimated by analysing the model-simulated O_3 mixing ratios along the air mass trajectories ending at few representative locations over the BoB (Fig. 7a). The model-simulated chemical evolution of air parcels clearly shows an increase in the O_3 mixing ratios towards the marine region of BoB. The temporal variations of O_3 mixing ratio averaged corresponding all the trajectories shown in Fig. 7a is shown as a box plot in Fig. 7b. A linear regression analysis (blue curve in Fig. 7b)

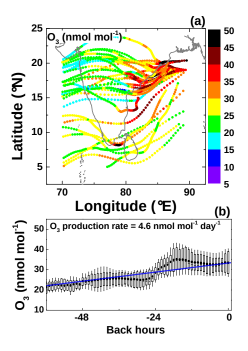


Figure 7. (a) WRF-Chem simulated O_3 along the air mass trajectories ending over a few representative locations over the BoB. (b) Variation of O_3 mixing ratios with time along the trajectories as shown in (a). In box plot, the black dots and lines inside the box represent the mean and median of the data, respectively. While the lower and upper edges of boxes represent the 25th and 75th percentiles, respectively, the whiskers represent standard deviations.

is used to estimate the mean en route O_3 production rate of $\sim 4.6 \text{ nmol mol}^{-1} \text{ day}^{-1}$ in the outflow. The enhancement in average O_3 over the BoB as compared to the continental Indian region is also shown in the Fig. 2d, averaged for the study period.

5.3 Ozone variations during rainfall events

An interesting phenomenon observed during the CTCZ experiment is the abrupt reduction in O_3 mixing ratios that accompanied the onset of heavy rainfall, despite the low solubility of O_3 in water. In this section the possible causes of these low- O_3 events during rainfall are investigated.

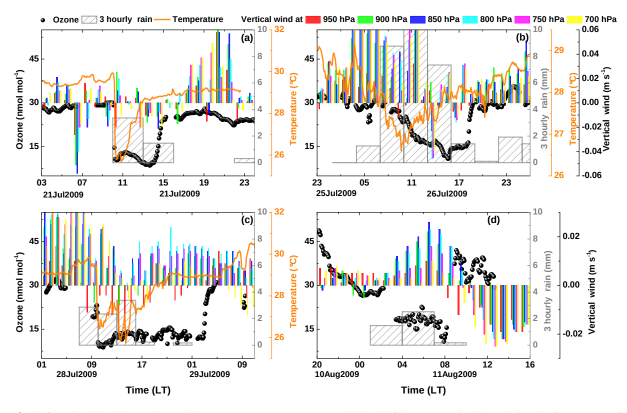


Figure 8. Surface O_3 (black dots) along with temperature (orange curve) and 3 h rainfall (grey vertical bar) during the four events of sharp decline in O_3 (**a**–**d**) as marked in Fig. 4c. Colours indicate the vertical wind as simulated by WRF-Chem.

Table 4. A comparison of average mixing ratios of surface trace gases measured over northern BoB $(81-91^{\circ} \text{ E}, 16-21.5^{\circ} \text{ N})$ and central BoB $(80-91^{\circ} \text{ E}, 11-16^{\circ} \text{ N})$ in different seasons as measured during different experiments. The range of mixing ratios (i.e. minima-maxima) is given in the brackets.

Study period	Name of experiment	Reference	O ₃ (nmol mol ⁻¹) over northern BoB	O ₃ (nmol mol ⁻¹) over central BoB	CO (nmol mol ⁻¹) over northern BoB	CO (nmol mol ⁻¹) over central BoB	CH_4 (µmol mol ⁻¹) over northern BoB	CH ₄ (µmol mol ⁻¹) over central BoB
December 2008– January 2009	W_ICARB	David et al. (2011)	63.0 ± 4.7 (50.8–73.8)	40.9 ± 6.7 (27.7-63.5)	302 ± 68 (140-450)	188 ± 53 (50-320)	No data	No data
February 2003	BOBEX-II	Lal et al. (2007)	\sim 34.1 (15.8–50.4)	~ 26.8 (13.9-35.0)	~ 238 (187–292)	~ 192 (159–224)	~ 1.77 (1.70–1.85)	~ 1.73 (1.68–1.77)
February– March 2001	BOBEX-I	Lal et al. (2006)	~ 23.8 (16.1-38.3)	\sim 38.0 (19.4–62.9)	~ 194 (165–235)	~ 227 (97–339)	~ 1.94 (1.89–2.02)	~ 1.91 (1.74–2.06)
March-April 2006	ICARB	Nair et al. (2011); Srivastava et al. (2012)	27.4 ± 2.9 (21.4-32.6)	13.4 ± 4.2 (3.1–24.6)	~ 189 (157–235)	~ 132 (96–167)	~ 1.84 (1.80–1.88)	~ 1.80 (1.75–1.84)
July-August 2009	CTCZ	Present Study	30.0 ± 6.9 (8.50–54.1)	27.5 ± 5.0 (8.8-40.5)	$95 \pm 25^{*}$ (50-200)*	$101 \pm 27^{*}$ (50–157)*	1.86 ± 0.12 (1.62-2.15)	1.72 ± 0.14 (1.57–1.96)
September– October 2002	BOBPS	Sahu et al. (2006)	~ 27.3 (17.8–33.8)	\sim 30.6 (22.5–35.2)	~ 152 (109–179)	~ 141 (108–211)	~ 1.79 (1.72–1.86)	~ 1.73 (1.68–1.80)
November 2010	-	Mallik et al. (2013)	~46.0 (26.7–59.6)	~ 38.7 (17.8–60.8)	~223 (131–280)	~188 (42–266)	~1.95 (1.85–2.06)	~ 1.79 (1.67–1.93)

* CO mixing ratios below the detection limit (i.e. 50 nmol mol-1) are not considered in the analysis

Figure 8 shows variations in O_3 (black circles) mixing ratios, surface temperature (orange curve), TRMM retrieved rainfall (thick grey vertical bars), and WRF-Chem simulated vertical winds at pressure levels ranging from 950 to 750 hPa (coloured bars) during four such events on 21, 26, and 28–29 July and on 10–11 August 2009. As high time resolution in situ measurements of rainfall were not available aboard ship, Fig. 8 therefore uses 3 h rainfall retrievals from the TRMM, co-located with O₃ measurements. During these events, CO mixing ratios also show a reduction of about $\sim 56 \text{ nmol mol}^{-1}$, with observed values falling below the detection limit of the instrument during the first event of 21 July 2009 (not shown). Although CO measurements are not available for the second and third event, during the fourth event (10–11 August 2009) CO mixing ratios showed an enhancement due to transport from strong source regions (see Sect. 5.1). While the first three low- O_3 events are not captured by WRF-Chem (Fig. 4c), the fourth event is reproduced.

Wet scavenging does not directly reduce O_3 , as its water solubility is low; as a result, some dynamic process could be responsible for the observed reductions in O₃ during rainfall. Air masses could undergo downdrafts during heavy rainfall (Kishore Kumar et al., 2005) through air drag by the falling rain drops and in mesoscale subsidence that compensates convective updrafts. We suggest that, in the presence of O₃-poor air mass aloft, a downdraft would result in reductions in surface O₃ mixing ratios. An opposite scenario leading to O₃ enhancement could take place if downdrafts bring mid-tropospheric air, where typically O₃ is higher than at surface. The model-simulated meteorology shows occurrences of downdrafts at different pressure levels during the first three events on 21, 26, and 28-29 July (Fig. 8a-c), which is further corroborated with measurements of air temperature aboard. Downdrafts of free-tropospheric air could lead to a reduction in near-surface temperature by as much as 10 °C within a few minutes (Ahrens, 2009). Air temperature measured aboard ship showed a sharp decrease of 2-4 °C, coinciding with the first three low-O₃ events (Fig. 8a-c). The reductions in temperature caused by downdrafts are generally short-lived (Ahrens, 2009), as is confirmed in the case of these events (Fig. 8a-c).

Model-simulated vertical winds and variations in air temperature suggest that downdrafts did occur during the first three rainfall events. As in situ measurements of O3 vertical profiles are not available over the BoB during the summer monsoon season, the observations taken at Thumba, Thiruvananthapuram, in the southern Indian region are used as a case study to investigate this hypothesis. For general details of the typical diurnal and seasonal variations of O₃ at Thumba, please see Nair et al. (2002), David and Nair (2011), and Girach et al. (2012). Figure 9a shows the temporal variation in surface O₃ on 15 July 2011 at Thumba, along with 5 min accumulated rainfall. Here, surface O₃ is observed to decline from 25 to 13 nmol mol^{-1} within 15– 20 min, coinciding with the occurrence of intense rainfall (3.5–0.5 mm rain over a period of 5 min). Measurements of the O₃ vertical profile are not available for this day due to the rainy conditions; a profile measured on 28 July 2011 is therefore shown in Fig. 9b. This profile has lower O₃ mixing ratios aloft (~22 nmol mol⁻¹ at ~1 km) than near the surface ($\sim 42 \text{ nmol mol}^{-1}$). The observed mixed layer height is about 0.15–0.6 km over Thumba, Thiruvananthapuram (Anurose et al., 2016), during July 2011; above this height, O₃ mixing ratios sharply decrease with altitude. The present case study suggests the presence of O3-poor air masses aloft than those near the surface over the south Indian region during summer monsoon. With an O₃ distribution as observed in the present case study at Thumba, the downdraft during intense rainfall could lead to the mixing of free-tropospheric

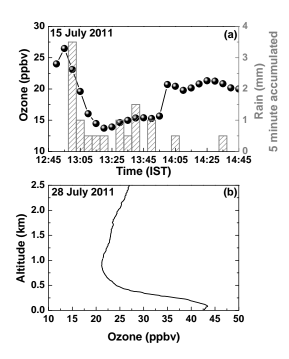


Figure 9. (a) Temporal variation in surface O_3 mixing ratio (black dots) along with 5 min accumulated rainfall (grey vertical bars) over Thumba, Thiruvananthapuram (location of the site shown in Fig. 1), on 15 July 2011. (b) Vertical profile of O_3 mixing ratio over Thumba, Thiruvananthapuram, as measured on 28 July 2011.

air with near-surface air or to the replacement of surface air with free-tropospheric O_3 -poor air.

Although air temperature measurements could not be made during the fourth event (10-11 August 2009) due to a technical problem, model meteorology does not indicate a downdraft during this event (Fig. 8d), indicating the dominance of a different process. As WRF-Chem simulated O₃ variability is in good agreement with observations during this event, various tendency terms (Barth et al., 2012) from WRF-Chem are used to investigate the relative influences of different processes. The variations in instantaneous values for horizontal advection tendency, vertical advection tendency, and net tendency (i.e. the sum of chemical, vertical mixing, convective, vertical advection, and horizontal advection), along with modelled O_3 over the two locations during the event, are shown in Fig. 10. The tendency values shown here are derived by subtracting the accumulated tendencies at (n-1)th hour from the accumulated tendency at *n*th hour. The vertical dotted lines show the time of a low-O₃ event.

Both the horizontal and net tendencies (Fig. 10b–c, e–f) show negative values, indicating that they are contributing towards a reduction in O_3 mixing ratios (Fig. 10a, d). However, as the time of the event approaches, it is the horizontal advection tendency term that is significantly negative (Fig. 10c, f), while other terms are small and close to zero. Horizontal advection is therefore suggested to dominate during the low- O_3 event of 10–11 August 2009. The influence of horizon-

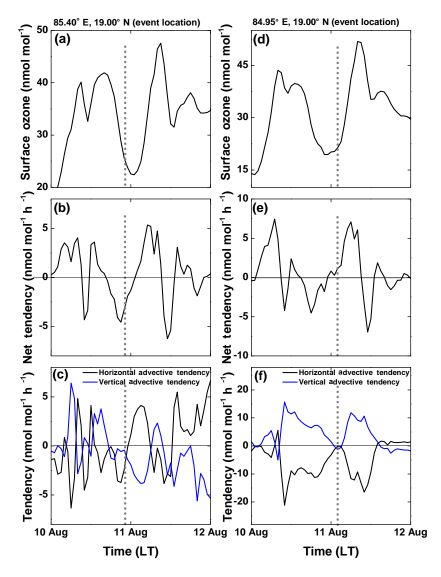


Figure 10. Time series of surface O_3 (**a**) and various tendency terms (**b** and **c**) over the event location during the fourth low- O_3 event, as obtained from WRF-Chem simulations. (**d**)–(**f**) are the same as (**a**)–(**c**), but for another location during the same event. These two event locations are also marked by triangles in Fig. 11. Vertical dotted line shows the time of the event in the in situ observations of surface O_3 over the indicated locations.

tal advection on O_3 during this event is shown more clearly in Fig. 11, which shows the spatial distribution of O_3 and CO from WRF-Chem before the event (16:00 and 19:00 UT) and during the event (22:00 UT) on 10 August 2009. The white triangles show the two locations where the event was observed. During 16:00 and 19:00 UT, a patch of high O_3 mixing ratios (35 nmol mol⁻¹ and higher) is seen to be distributed over a large region surrounding the measurement location. This large patch of elevated O_3 mixing ratios is horizontally advected eastward from 16:00 to 19:00 and then towards 22:00 UT (event time). As a result of this rapid advection, the high- O_3 air masses are transported from the coastal regions to deeper into the BoB; by the time they reached the location of observation, O_3 mixing ratios are observed to be lower (25–35 nmol mol⁻¹) during the event time (22:00 UT). A patch of higher levels of CO (\sim 300 nmol mol⁻¹) was also found to be distributed across the east coast of the Indian region. Transport and dilution of this CO patch is, however, less pronounced than the high-O₃ air masses, possibly due to the relatively longer lifetime of CO. Thus, in a nutshell, the horizontal advection played a key role in transporting O₃-rich air masses deeper into the BoB region, while it diluted O₃ levels near the coastal regions in southern India during the fourth event.

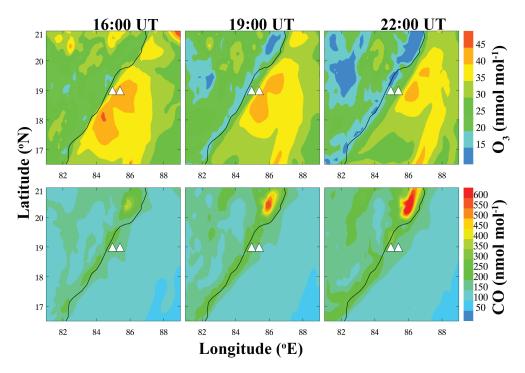


Figure 11. Spatial distribution of surface O_3 (top panel) and CO (bottom panel) at 16:00 and 19:00 UT on 10 August 2009, both prior to and during the fourth event, which took place 22:00 UT on 10 August 2009. White triangles show two locations (85.40° E, 19.00° N; 84.95° E, 19.00° N) corresponding to the event.

5.4 Seasonal variation in trace gases over the BoB

In this section, the monsoon-time measurements of O_3 taken in the present study are combined with data from previous campaigns (see Table 4) to investigate the seasonal variation in O_3 over the BoB (Fig. 12). O_3 , CO, and CH₄ mixing ratios are averaged over northern (81–91° E, 16–21.5° N) and central (80–91° E, 11–16° N) BoB regions.

Overall, higher O_3 mixing ratios are present over both northern and central BoB during the winter, while lower O₃ levels are observed during the spring-summer (with more scatter in the data over central BoB). The O₃ seasonal amplitude (i.e. the range from maxima to minima) is estimated to be ~ 39 over northern BoB and ~ 27 nmol mol⁻¹ over central BoB. The monsoonal surface O₃ mixing ratios $(\sim 30 \pm 7 \text{ nmol mol}^{-1})$ are nearly half those observed during winter $(63 \pm 5 \text{ nmol mol}^{-1})$ over northern BoB. During December 2008-January 2009, February 2003, March 2006, and November 2010, the O₃ mixing ratios were higher (by \sim 3–22 nmol mol⁻¹) over northern BoB than over central BoB. However, over the course of February 2001, O₃ mixing ratios were higher over central BoB (\sim 38 nmol mol⁻¹) than over northern BoB ($\sim 14 \text{ nmol mol}^{-1}$). In contrast, during summer monsoon season, average O₃ mixing ratios are comparable or only slightly higher over northern BoB $(30 \pm 7 \text{ nmol mol}^{-1})$ as compared to over central BoB $(27 \pm 5 \text{ nmol mol}^{-1}).$

As compared with the summer monsoon season, when CO mixing ratios were lower over northern BoB, CO mixing ratios were higher during the winter, while over central BoB CO mixing ratios were higher during the pre-monsoon season. For O₃, spring–summer had the lower mixing ratios in both regions. The seasonal amplitude in CO mixing ratios is estimated to be ~ 205 over northern BoB and ~ 124 nmol mol⁻¹ over central BoB. The monsoonal CO mixing ratio (~95 nmol mol⁻¹) is about one-third that of the winter season (302 nmol mol⁻¹) over northern BoB. During the present study, average CO mixing ratios were comparable over northern (95 ± 25 nmol mol⁻¹) and central BoB (101 ± 27 nmol mol⁻¹).

A clear inference about seasonal patterns is difficult in the case of CH₄, but a tendency of lower levels towards winter can be seen. Higher mixing ratios ~ 1.95 (~ 1.91)µmol mol⁻¹ were observed during November 2010 over northern BoB, and during February–March 2001 over central BoB, as compared to those from other studies. The surface CH₄ observations obtained during the present study show the highest variability (i.e. the difference between maxima and minima) when compared to earlier studies: 0.53 over northern BoB and 0.39 µmol mol⁻¹ over central BoB. This high variability is attributed to the relative source strengths over central and northern India as compared to southern India, highlighting the regional differences in CH₄ variability across India (Kavitha and Nair, 2016).

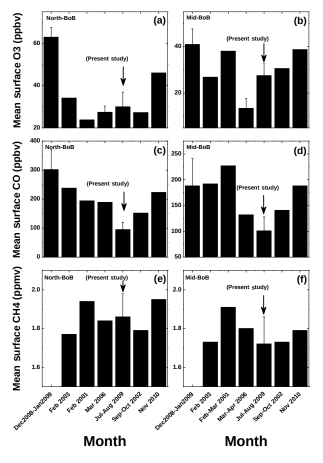


Figure 12. Seasonal variation in average O_3 , CO, and CH₄ mixing ratios over (**a**, **c**, **e**) northern BoB and (**b**, **d**, **f**) central BoB. Except for July–August 2009 (present study period), all average values are obtained from the literature (David et al., 2011; Lal et al., 2007, 2006; Nair et al., 2011; Srivastava et al., 2012; Sahu et al., 2006; and Mallik et al., 2013). Error bars show standard deviations for respective study periods. For any points for which high-resolution measurements are not available, standard deviations are not shown.

Seasonal variations in trace gases over the BoB are attributed to seasonal changes in the meteorological conditions, emissions, and photochemistry over the South Asian region, as well as to synoptic-scale transport patterns. Wintertime stronger westerly winds transport the pollution from South Asia, including that of the Indo-Gangetic basin, to the BoB region. Monsoonal circulation, in contrast, carries cleaner marine air masses to the BoB from the oceanic regions. However, as observed during the CTCZ, polluted continental or coastal air masses can also occasionally be transported deeper over the BoB. Intense monsoonal rainfall generally leads to wet removal of O₃ precursors, while cloudy and rainy meteorological conditions suppress O₃ formation. Along with the importance of monsoonal convection in cloud formation, rainfall, and uplifting the boundary layer pollution, rapid horizontal advection is also an important process during the summer monsoon, especially affecting the nearsurface variability of trace gases over the oceanic regions adjacent to India.

6 Conclusions

In this paper, we presented shipborne in situ measurements of O_3 , CO, and CH₄ that were carried out as a part of the CTCZ experiment over the BoB during the summer monsoon season, July–August 2009. We analysed the spatial and temporal variations in the observations and compared them with results from simulations performed with a regional chemistry transport model (WRF-Chem), as well as with observations from previous campaigns over the BoB. The main conclusions are as follows.

These first monsoonal observations of O₃, CO, and CH₄ show large spatio-temporal variability over the BoB, with mixing ratios varying in the range of 8-54 (mean: 29.7 ± 6.8) nmol mol⁻¹, 50-200 96 ± 25) nmol mol⁻¹, and 1.57–2.15 (mean: (mean: 1.83 ± 0.14) µmol mol⁻¹, respectively. The O₃ and CO mixing ratios in air masses from central/northern India are slightly higher or comparable (O₃: 30 ± 7 , CO: $95 \pm 25 \,\mathrm{nmol \, mol^{-1}}$) to those in air masses from southern India (O₃: 27 ± 5 , CO: 101 ± 27 nmol mol⁻¹). The CH₄ mixing ratios in air masses from central/northern $(1.86 \pm 0.12 \,\mu\text{mol mol}^{-1})$ are higher (~0.14 μ mol mol⁻¹) compared to those in air masses from southern India $(1.72 \pm 0.14 \,\mu\text{mol}\,\text{mol}^{-1})$. This could be due to higher CH₄ levels over central/northern India, also found in SCIAMACHY data.

Back-trajectory analysis shows effects of long-range transport from northern or central India to northern BoB and from southern India to central BoB. The correlated variations of these trace gases and percentage residence time of air parcels over the Indian regions suggest that the enrichment of O_3 and precursors in air parcels over the BoB is associated with both emissions and photochemistry over the Indian region. The trajectory analysis shows that the observed variation in surface O_3 is primarily due to transport and en route photochemistry over the BoB during monsoon season. An analysis of modelled O_3 along air mass trajectories show mean en route O_3 production rate of about 4.6 nmol mol⁻¹ day⁻¹ in the outflow towards the BoB.

The observed spatio-temporal variations of surface O_3 and CO during summer monsoon season are generally reproduced by WRF-Chem, although the absolute mixing ratios of O_3 and CO are typically overestimated by about 6 and 16 %, respectively.

The four low- O_3 events coinciding with intense rainfall were observed over the BoB. After analysing the observed variability in air temperature, model simulations of vertical winds, and an O_3 -profile case study from southern India, we suggest that first three low- O_3 events were due to strong downdrafts of O_3 -poor air masses. Analysis of the fourth event, which is successfully reproduced by the model, shows the pivotal role of horizontal advection in transporting O_3 rich air masses deeper over the BoB.

Finally, the measurements during the monsoon are combined with previous campaigns over the BoB during other seasons to investigate the seasonal variability in trace gases over the BoB. O₃ and CO are shown to have pronounced seasonality, O₃ having amplitudes of about 39 and 27 nmol mol⁻¹ and CO having amplitudes of about 207 and 124 nmol mol⁻¹ over northern and central BoB, respectively.

Our study data fill a gap of observations during the summer monsoon over the BoB, providing information on the extent of seasonal variability. We recommend supplementing these findings with shipborne experiments featuring collocated vertical profile observations from balloon-borne and aircraft-based platforms over the oceanic regions surrounding India to better understand the role of both large-scale dynamics (e.g. Ojha et al., 2016) and of regional influences due to South Asian outflow (see Lawrence and Lelieveld, 2010, and references therein). Such a future study would also improve our understanding of the changes that take place in the atmospheric oxidation capacity during the summer monsoon season.

7 Data availability

The observational data used in the paper can be obtained by contacting the corresponding author (imran.girach@gmail.com). The metadata are available on the website of Space Physics Laboratory (http://spl.gov.in/index.php?option=com_content&view= article&id=394&Itemid=695&lang=en).

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