Variations in surface ozone and carbon monoxide in the Kathmandu Valley and surrounding broader regions during SusKat-ABC field campaign: Role of local and regional sources

By Piyush Bhardwaj et al., 2017 (ACPD)

We would like to thank the referees for their comments and suggestions and the editor for providing an extended period that allowed us to adequately respond to the comments and improve quality of the manuscript. Here, please find the reviewer’s comments in regular black font and our responses in regular blue font. All the changes made in the revised manuscript are indicated in red color. The line numbers in our response refers to the line numbers in the revised manuscript.

Reviewer-2

The paper presents data collected during the winter and spring of 2012-2013 in the Kathmandu valley region of Nepal of O3 and CO and contemporaneous measurements of these two gases from Nainital and Pantnagar in India. Hydrocarbon data was collected for a few weeks in December and January during the same period in Bode, Nepal. Extensive analysis of this data is presented contrasting O3 and CO for the winter and spring and episode resulting from biomass burning over western parts of India on the air masses observed over the three sites is presented. The dataset is unique and certainly worthy of discussion in a paper.

Thank you very much for appreciating the extensive analysis and uniqueness of data from three sites that brought out contrasting features here. Here, we have addressed all your concerns including the analysis of model results.

One serious short coming of the paper is an absence of model calculations supporting numerous statements in the manuscript. I can’t see how this will make sense without supporting model simulations. I will detail some of them below.

We have now added model (WRF-Chem) simulations performed over the Kathmandu region. The model output is used to support some of the statements, while other general statements without supporting evidence are removed. Aurelia Lupascu, of the IASS, Potsdam, Germany has kindly shared her model simulations with us and so, she has been added as a co-author in this work. A brief description of the model configuration along with discussions of model results has been added in the revised manuscript.

A second un answered question is what connects these three sites besides them being in approximately near and in the Himalayan foothills? Kathmandu is in a valley and that makes it meteorology fairly unique and extensively influenced by drainage flows, flow through mountain passes and other complex flow situations. Nainital site is at the edge of the mountain ranges and high enough that it can be considered a background site. Pantnagar being lower
elevation and in the IGP and is potentially not directly connected to Kathmandu (in terms of transport). I would like to see some discussion why these three sites make a good case for comparing with each other.

We agree with the reviewer that it is very important point to highlight the importance of considering these three sites together. We tried to mention it in the Introduction of the previous version of the manuscript and we apologize that it was not clear to you. The goal of the SusKat field campaign was not only to perform a detailed characterization of air quality in the Kathmandu Valley but also to understand how the magnitude and variability of air pollution in Kathmandu Valley compares with other sites in the region especially with those located in the cleaner Himalayas and the Indo-Gangetic Plain (IGP). To achieve this goal, Indian institutions also participated in the field campaign by deploying some of their instruments in the Kathmandu Valley and by sharing air quality measurements from their own sites in India. Thus, Nainital (a background site representative of the central Himalayan environment) and Pantnagar (representative of the IGP environment) sites were part of the SusKat field campaign. While most of the previous studies published from the SusKat field campaign focused on analysis of measurements performed in Kathmandu Valley, this study takes a step further to present the regional picture during SusKat and attempts to understand the similarities and differences between the air quality of Kathmandu Valley and the Indian sites, to identify the regional emission sources that are common to these sites. This information has been included in the revised manuscript.

Finally, the use of HYSPLIT for boundary layer flow reaching Bode in the central of the Kathmandu valley with a 1 degree x 1 degree GDAS fields is probably not a good idea. It would be reasonable to use this method for trajectories reaching mid troposphere. Due to the complex flow conditions here you need much higher resolution flow fields and may be trajectories that reach the top of PBL at Bode rather than the surface. I recommend they try using higher resolution (0.5 degree?) flow fields or better (generated with WRF simulations for example) to increase the confidence in these trajectories.

Thank you very much for this comment. Earlier, we were unable to obtain higher resolution wind field data. Now, we have managed to get higher resolution (0.5°x0.5°) meteorological fields from the Global Data Assimilation System (GDAS) and same are now used to generate the 5-day back-air trajectories over the Bode region. We have now revised this information in Section 2.4 (Satellite data, model and back-air trajectory). The patterns of the back-air trajectory remain more-or-less similar and do not affect conclusions of the study. We have replaced Figure 4 with the new trajectories and text has also been revised in section 3.2. Back-air trajectories calculated with higher resolution wind fields are also shown below for your ready reference.
Specific Comments:

Page 14, Line 1: photo-dissociation of NO₃ and NO₅ at sunrise Q: Have you measured, NO₃, N₂O₅, HNO₃ or NOy ever during this experiment. Are there model calculations that show how much N₂O₅ can be produced during nighttime? Are there any estimates of PAN produced using models or observations? It is really hard to tell this complicated story using CO and O₃. You will need lot more measurements to constrain your story and these are basic measurements for any air quality study.

Unfortunately, the measurements of NO₃, N₂O₅, HNO₃ or NOy and PAN were not conducted during the campaign, and lack of systematic observations of these species in South Asia remains a long-standing issue. Therefore, as suggested by the reviewer, we use the model output to gain some process-level understanding of this complex system. The model simulated diurnal variations in NO, NO₂, NO₃ and N₂O₅ during February and May of 2013 are shown in Figure 2 (below). NO mixing ratios are close to zero during the nighttime because it rapidly reacts with O₃ to form NO₂, which also explains higher NO₂ levels during nighttime. NO₃ and N₂O₅ also show higher levels during nighttime because of the reactions of NO₂ with O₃, of NO₂ with NO₃, respectively. The sharp morning increase in NO mixing ratios correlates strongly with the sharp decrease in NO₃ and N₂O₅ mixing ratios especially during February indicating that photodissociation of NO₃ (λ < 670 nm) and N₂O₅ (280< λ <380 nm) releases NO back to the atmosphere. There is likely some contribution from the NO₂ photolysis as well. This figure
has been added as Figure 6 in the revised manuscript and discussion has been added in Section 3.3.

Figure 2: Model simulated average diurnal variations in NO, NO$_2$, NO$_3$ and N$_2$O$_5$ during February and May 2013.

Page 14, Line 5 to 10: discussion on ozone mix down Q: Are there any measurements of ozone profiles at Bode? Ozone sondes etc? In its absence running a model may help evaluate these claims. As it stands this is pure speculation and unsupported by any facts and the discussion is very qualitative.

Unfortunately, vertical measurements of ozone are not made at Bode site during this field campaign. Therefore, we have now removed this sentence from the revised manuscript.

Page 15, line 5 to 10: Decrease of CO from morning to evening Q: This is again a fairly qualitative description with no supporting data. Later in the discussion it seems like the photochemistry is active during the early morning hours. What are the OH levels at Bode during early morning and late afternoon time periods? A model will help distinguish between chemical and meteorological phenomena.

We have now added the observations of the boundary layer height and discussion have been revised (section 3.4, page 16) accordingly. OH, observations were not made during the campaign and its direct observations are not existing in the South Asia. But, yes, we agree that CO loss via OH will also contribute and we have modified this accordingly (section 3.4 page 16). We have also added estimates of ventilation coefficient and showed that higher wind speed
leads to lower CO levels

Page 17, Line 1 to 5: Seasonal variability of CO, decrease from winter to Spring Q: Again, I am not sure how much role chemical loss of CO is important here. Having an idea of OH concentration changes between winter and spring in the valley would be useful. This being a valley the CO emitted could stay trapped for much longer times than other places and hence photochemistry plays a bigger role in CO lifetime.

We agree with you and we have revised this part with mention of role of OH chemistry. Model simulated OH levels are found to be higher in May, when compared with February. This would suggest great chemical loss of CO in spring (May).

Page 18, Line 1:10: ozone variation from winter to spring Q: What happens to NO emissions from Winter to Spring? If CO emissions are said to be decreasing will it also not lower NO emissions? Does a decrease in fresh NO at evening hours keep more of the ozone from losses during the night? This would make the ozone issue mostly local.

It is difficult to comment on winter to spring changes in CO and NOx emissions with our dataset because some emission sources (e.g., brick kiln industries and domestic heating) operate only during winter while others (e.g., crop residue burning and forest fires) are more active during the spring. We do not anticipate large variability in other anthropogenic sources such as cooking, traffic and power generation. An accurate characterization of seasonal variability in CO and NOx emissions over this region will require the development of an emission inventory considering the temporal variability in all the emission sources, which is beyond the scope of present study. The increase in biomass burning activity is supported by a detailed analysis of MODIS active fire locations and this discussion has been extensively revised in section 3.7 (Influences of springtime northern Indian biomass burning). Please see our responses to Reviewer#3. We also analyzed OMI and GOME-2 retrieved tropospheric column NO2 (cloud screened 30%) over the Bode region (27-28°N, 85-86°E) to understand variability in the tropospheric column burden of NO2. Both the satellites show similar levels (below figure 3) in winter and spring (except somewhat higher levels in early April which we feel are due to biomass burning in nearby regions).

In addition to changes in emissions, we envisage that variations in the PBL height between winter and summer can significantly affect the surface concentration of different air pollution. The observations of the boundary layer height were made during the campaign using a ceilometer and those clearly show higher boundary layer height during spring (pre-monsoon) i.e. March, April and May (Figure 3 above; please also see Figure 4 from Mues et al., 2017). Similar increase in PBL height is seen in the model results. If we assume that emissions are constant from winter to spring, then increase in the PBL height will lead to lower mixing ratios of species such as CO and NOx by allowing the emissions to mix into a larger volume compared to the winter.

We also agree with the reviewer that lower NO levels during evening hours could reduce the ozone loss. The model simulated NO levels are lower in spring ((Figure 6 of the revised
manuscript) that will lead to reduction in nighttime ozone loss in spring. This discussion has been included in the revised manuscript.

![Graph](image)

**Figure 3:** *OMI tropospheric column NO₂ (cloud screened 30%) and GOME-2 tropospheric column NO₂ over Bode.*

Page 18, Line 11 – 15: negative correlations between CO and ozone Q: what is the explanation for this negative correlation?

Negative correlation during nighttime and early morning time is a manifestation of ozone titration by NOₓ and the lower boundary layer height. We have now revised the text accordingly (section 3.5).

Page 18, Line 20:24: measurements from Nainital and Pantanagar Q: How are these sites connected meteorologically?

Nainital and Pantnagar are meteorologically disconnected during the nighttime and early morning hours because the boundary layer height remains below the altitude of Nainital. However, the boundary layer height is higher the altitude of Nainital during afternoon and thus are connected meteorologically. In the afternoon, Pantnagar acts as a representative of emission sources affecting Nainital. We have now added the discussion in section 2.1 (Observation site) in the revised manuscript. Details of both sites have been provided in previous publications (Kumar et al., 2010, Ojha et al., 2012; Sarangi et al, 2014; Naja et al., 2014; Joshi et al., 2016).

Page 20: Line 2-5: No titration discussion Q: This probably is the explanation for the ratios of CO and ozone

We agree. We have added this in the response of previous comments also.

Page 21, Line 6: Bode are likely Q: do you mean ‘unlikely’?

Sorry for the mistake. We mean, ‘unlikely’ and now the statement is revised accordingly.

Figure 11: ozone, co time series Q: what about the CO and ozone peaks in April? Are they also
from biomass burning?
Yes, they are also due to biomass burning. Now, we have further improved the biomass burning analysis (as suggested by reviewer-3) over the region to identify the role of biomass burning in the CO and O₃ peaks during April. These peaks were identified to be highly correlated to the high biomass burning activity which occurred in the nearby regions surrounding the Kathmandu valley. The HYPLIT calculated back-air trajectories also show transport from the active fire regions to Kathmandu Valley during this period. The air-masses during this period were found to be circulating in the region. A similar episode was also detected during the early May over the region where influences of northern Indian biomass burning was investigated. The entire section (3.7) regarding biomass burning is rewritten to answer this and other similar questions.
Reviewer-3

The manuscript presents observations of ozone, carbon monoxide and some of the hydrocarbons at Bode in the Kathmandu Valley for a period of 6 months. A correlation analysis and comparison with stations in the northern Indian subcontinent are conducted, and effects of biomass burning are studied. However, manuscript in its present form adds limited new insights into the chemistry and dynamics over this region, and the observations of ozone and carbon monoxide shown here for a period of about 6 months are subset of full year data at same station presented in Mahata et al., 2017. The discussions in the present version are qualitative and general, as elaborated in following comments.

We thank the reviewer for a thorough review of our manuscript. However, we strongly disagree with the reviewer’s assessment that this study simply presents a subset of yearlong ozone and CO observations presented in Mahata et al. (2017). This study differs from Mahata et al. (2017) particularly in the sense that Mahata et al. (2017) focuses on observations conducted in and around Kathmandu Valley while this study for the first time provides a broader regional picture by complementing the observations from Kathmandu Valley with simultaneous observations from a high-altitude background site located in the central Himalayas (Nainital) and a semi-urban site representative of the chemical environment of the Indo-Gangetic Plain. Since most of the previous studies have already provided key information about air quality in Kathmandu Valley, this study takes a step further to present the regional picture during SusKat and attempts to understand the similarities and differences between the air quality of Kathmandu Valley and the Indian sites, to identify the regional emission sources that are common to these sites. Furthermore, the revised manuscript complements the observations with model simulations following suggestions from both the reviewers and also discusses the role of biomass burning in greater detail. Below, we respond to all the reviewer comments one by one with the comments reproduced in black font and our response in blue font.

Regional sources are suggested as the driver of springtime ozone enhancement over Bode, however, Fig. 10 clearly shows that during both winter and spring the ozone production is faster over Bode (and Paknajol) as compared to that over IGP stations. For the transport to be the driver, one would expect just an enhancement in the levels with lesser local production, not being evident here.

We agree with the reviewer that in situ ozone production resulting from photooxidation of precursor gases is the main source of ozone at Bode as reflected by faster ozone production rates in Figure 10 (now Figure 12). However, we wanted to point out that regional transport adds ozone on top of the already higher in situ photochemical ozone production at Bode especially during springtime. An increase in regional-scale O₃ levels during the spring season has also been reported in several previous studies (e.g., Kumar et al., 2010, Ojha et al., 2012; Sarangi et al., 2014; Putero et al., 2015), and it seems to be playing an important role at Bode as well. We have revised the statements in the manuscript to reflect this discussion.
Springtime enhancement in ozone at Bode appears primarily due to a broader ozone maxima in the noontime which authors attribute to solar radiation (Page 14, lines: 3-5), and higher ozone levels during the nighttime attributed to lower titration with NOx (Page 20, Line 2). Both of these processes are of local origin.

The reviewer’s question is not clear here, but we believe that the reviewer is advocating that local ozone production is the major contributor at Bode. We agree with the reviewer’s opinion here.

Effect of biomass burning: It would be more appropriate to show a time series of fire counts over potential source region in north India, a running mean of fire counts would be even better as done previously (Kumar et al., JGR, 2011) to classify High and low fire activity periods. Presently, the selected period shows only small enhancements in CO, while larger CO enhancements are seen starting from middle-April.

Thanks for this comment. We have now revised this analysis completely as suggested. We have classified the observations in high and low fire activity periods, and quantified fire induced enhancements in O₃ and CO. During the spring season, two distinct peaks in O₃ and CO over Bode have been studied separately using the same methodology. To improve the quality of discussions, emissions from a high-resolution biomass burning emissions inventory and satellite retrievals are also used. We have now revised the section 3.7 and new figures and tables have also been added in the revised manuscript. The figures and table are reproduced below from the revised manuscript for your ready reference.
Figure 4: Top Left: time series of MODIS daily fire counts (red bar), 3-day running mean (black lines), median fire counts (brown line) for the fire period (3 April-31 May 2013). Total biomass burning emissions (orange line), crop residue burning emissions (dark yellow line) and forest fire emissions (dark green) over 1°×1° grid box around Bode (27-28°N, 85-86°E) are also shown. Top right: average biomass burning emissions for two fire activity periods over Bode region. Center: time series of surface ozone mixing ratios at Bode and Nainital (line plot-Left) and average ozone mixing ratios during two fire periods (as bar plot-Right), respectively. Bottom: time series of surface CO mixing ratios at Bode and Nainital (line plot-Left) and average CO mixing ratios during two fire periods (as bar plot-Right), respectively. The two fire events in April and May are also shown highlighted (in violet boxes).
Figure 5: Spatial distribution of MODIS fire counts during April (red dots) and May (orange dots) 2013. Underneath is daily HYSPLIT backward trajectories during April (blue lines) and May (green lines) initiating from Bode (black star). The two black boxes indicate two hotspots of fire counts during this period.
Figure 6: Top (1a-3a): Spatial distribution of MODIS fire counts over Northern Indian subcontinent during three periods (left to right). The black boxes represent two fire hotspots (refer Fig. S1) over the region with fire counts shown as different colors during the three periods. Center (1b-3c): spatial distribution of AIRS CO mixing ratio at 925 hPa (1b-3b) and 850 hPa (1c-3c) during three periods (left to right). Center (1d-3d): Average biomass burning emissions over Bode region (27-8°E, 85-86°E) using GFED v4.0 inventory during the three periods (left to right). Bottom Panels (1e-3f): changes in average surface mixing ratios of O₃ (1e-3e) and CO (1f-3f) at different sites during the three periods (left to right).
Figure 7: Top (1a-2b): Spatial distribution of MODIS fire counts over Northern Indian subcontinent during two periods (left to right). The black boxes represent two fire hotspots (refer Fig. S1) over the region with fire counts shown as different colors during different periods. Center (1b-2c): spatial distribution of AIRS CO mixing ratio at 925 hPa (1b-2b) and 850 hPa (1c-2c) during two periods (left to right). Center (1d-2d): Average biomass burning emissions over Bode region (27°-8°E, 85°-86°E) using GFED v4.0 inventory during the two periods (left to right). Bottom Panels (1e-2f): changes in average surface mixing ratios of O$_3$ (1e-2e) and CO (1f-2f) at different sites during the two periods (left to right).
**Table 4:** The average $O_3$ and CO mixing ratios at Bode, Nainital, Pantnagar with GFED average biomass burning emissions over Kathmandu region during different periods.

<table>
<thead>
<tr>
<th>Fire Periods</th>
<th>Fire Count</th>
<th>Ozone (ppbv)</th>
<th>CO (ppbv)</th>
<th>Avg. Biomass burning emissions (Tg/day)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Bode</td>
<td>NTL</td>
<td>Bode</td>
</tr>
<tr>
<td>LFAP (Mar 1- Mar 31)</td>
<td>5</td>
<td>37.4</td>
<td>45.2</td>
<td>705</td>
</tr>
<tr>
<td>HFAP (Apr- May)</td>
<td>70</td>
<td>43.7</td>
<td>63.9</td>
<td>504</td>
</tr>
<tr>
<td>Mar (20-25)</td>
<td>3</td>
<td>34.8</td>
<td>46.1</td>
<td>693</td>
</tr>
<tr>
<td>Apr (3-6)</td>
<td>18</td>
<td>46.7</td>
<td>49.2</td>
<td>797</td>
</tr>
<tr>
<td>Apr (11-16)</td>
<td>27</td>
<td>54.0</td>
<td>57.9</td>
<td>762</td>
</tr>
<tr>
<td>Apr (26-30)</td>
<td>26</td>
<td>33.9</td>
<td>45.0</td>
<td>505</td>
</tr>
<tr>
<td>May (2-6)</td>
<td>116</td>
<td>58.8</td>
<td>74.8</td>
<td>625</td>
</tr>
</tbody>
</table>

Page 10, lines 4-5: Why a high-resolution regional-scale model is not used to study transport from such nearby regions of north India to Nepal?

Following the reviewer suggestion, we have now added model (WRF-Chem) simulations results and the discussion has been revised extensively. We have also included figures comparing the model results with the observations.

Page 13: 1.17-20: It should be useful to substantiate the statements by calculating the variations in ventilation coefficients using measured boundary layer height and wind speed.

We have now added observations of the boundary layer height. Information on ventilation coefficients have also been added. The figure of ventilation coefficient is shown below for the ready reference.
Section 3.5: Correlation between ozone and CO: This section needs thorough revision. The anti-correlation is being explained by contrasting seasonality of ozone and CO. Then what is learnt from plotting correlations? Correlating between noontime ozone and CO showing positive relationship is useful. Fig. 7 could be removed.

We have now revised this section and added discussion on the correlation. We would like to retain the figure 7 (now fig 9) as it provides very useful information on changes in correlation with months. It very clearly shows decrease in correlation from winter to spring.

Stratosphere-to-troposphere transport (STT) could be important in ozone variations over the Himalayas during winter (Phanikumar et al., 2017) as well as spring (Sarangi et al., 2014). Smaller enhancements in CO but high ozone levels could have some effects of STT. Mountain regions in Nepal also experience such effects. What is the contribution of STT from northern India, or high altitudes in Kathmandu in the ozone enhancements during May event, and in general from Jan to May?

**Figure 8:** The average diurnal variations in ozone, CO, median MLH and Ventilation coefficient during winter (top) and Spring (bottom).
We have added a complete new section (3.8) to address the stratosphere troposphere exchange in the manuscript. In this section the vertical variations in potential vorticity, CO, relative humidity and ozone were discussed to check the role of stratosphere troposphere transport (STT) from January to May 2013. The updated figures are added below.

**Figure 9:** Vertical distribution of ertel potential vorticity (EPV) calculated by MERRA v2 reanalysis (top), AIRS retrieved ozone mixing ratios (centre), CO mixing ratios (center) and...
relative humidity (RH) during Jan-May 2013.

**Figure 10**: Vertical profiles of EPV, ozone, CO and RH during April (Top Panels) and May (Bottom Panels) biomass burning episodes.

Other comments

Section 2.1.: There should be a description of emission sources, and as paper aims to highlight the differences in the emissions from India and Kathmandu, it should have been substantiated by showing emission maps from recent inventories for some of the chemical tracers.

We appreciate this suggestion and we have added emission maps for CO and NOx (in the supplementary material) from EDGAR-HTAP emission inventory. Details of emission inventories are also available at [http://edgarjrc.ec.europa.eu/htap_v2](http://edgarjrc.ec.europa.eu/htap_v2).
Too general statements should be removed. Such as Page 11: l.17-18: “The solar radiation. . .the valley”

These lines were removed from the manuscript.

Page 11: l.11 “The regional contribution in this regard cannot be ruled out”. Can it be ruled out in other seasons / or other stations?

The high levels of air pollutants in Kathmandu Valley have been suggested to be primarily influenced by local emissions from the valley. In this study, we also observed that except for a few cases the air quality at Kathmandu is constrained by in-situ chemistry and emissions. Therefore, we wanted to emphasize that regional contribution can be important occasionally. Nevertheless, we have now removed this sentence.

With availability of data, there should have been some quantitative discussions. For example: Page 20, l.2-4: “indicating somewhat lesser polluted kind of environment in Bode. . .However this does not necessarily mean that NOx emissions are lower in Kathmandu valley.”

We have now added the information on the previous observations where NO levels are reported to be as high as 60 ppbv. More discussions on this have also been added in section 3.3.

Page 15: l.19: “Methane levels are much higher than the global average”. Mention a global average value for the period or from some reference and indicate by how much % it is found to be higher at Bode?

We have now provided the needed information (section 3.4). Nevertheless, it is quite obvious that methane levels more than 2 ppmv could be considered as higher levels.

Page 16, l.1-3: “. . .hydrocarbons are much higher than at Nainital. . .”. Mention some % or factor.

Yes, now we have provided this information. Nevertheless, values were already given in table 3.

Page 21: l.18-21: This discussion is about aerosol forcing, which is not studied in this paper so it can either be moved to introduction or can be skipped.

These lines are removed from the manuscript.

Page 21: l.22-23: CO enhancement seems small and within the variations, while ozone enhancements are observed in the following days too.

We have now revised the discussion. Some of these ozone enhancements are discussed in the biomass burning analysis section (3.7) and other sections.
Variations in surface ozone and carbon monoxide in the Kathmandu Valley and surrounding broader regions during SusKat-ABC field campaign: Role of local and regional sources

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Key words: Kathmandu, Himalayas, Air Pollution, Ozone, CO, Long Range Transport

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Highlights of the study:

- A comparative study regional picture of the SusKat field campaign based on synergistic analysis of trace gases among sites in the gas observations from Kathmandu Valley and India.

- An important contribution of regional Regional transport also contribute to the springtime ozone enhancement in the Kathmandu Valley.

- The winter-time higher ozone levels in the Kathmandu Valley are largely due to local sources.

- O3, CO and light NMHCs levels are higher in the Kathmandu Valley than the site in IGP.

- Regional pollution resulting from biomass burning in NW IGP led to simultaneous increase in O3 and CO levels in the Kathmandu Valley and two northern Indian sites in India.

- The Kathmandu Valley and the IGP have differences in their emission sources.
Abstract

Air pollutants emitted from rapid urbanization and associated human activities in the Kathmandu Valley of Nepal has been leading to serious public health concerns over the past two decades, are causing serious air quality and health concerns. These concerns led to a multinational field campaign SusKat-ABC (Sustainable atmosphere for the Kathmandu Valley- Atmospheric Brown Clouds) that measured different trace gases, aerosols and meteorological parameters in the Kathmandu Valley and surrounding regions during December 2012 to June 2013 to understand the factors local to regional scale processes influencing air quality of the Kathmandu Valley. This study provides information about the regional distribution of ozone and some precursor gases using simultaneous in situ measurements from a SusKat-ABC supersite at Bode, Nepal and two Indian sites: a high-altitude site Nainital located in the central Himalayan region and a low altitude site Pantnagar located in the Indo-Gangetic Plain (IGP). The diurnal variations at Bode showed a daytime buildup in O\textsubscript{3} while CO shows morning and evening peaks. Similar variations (with lower levels) were also observed at Pantnagar but not at Nainital. Several events of hourly ozone levels exceeding 80 ppbv were also observed at Bode. The CO levels showed a decrease from their peak level of above about 2000 ppbv in January to about 680 ppbv in June at Bode. The hourly mean ozone and CO levels showed a strong negative correlation during winter (r\textsuperscript{2}=0.82 in January and r\textsuperscript{2}=0.71 in February), but this negative correlation gradually becomes weaker, with the lowest value in May (r\textsuperscript{2}=0.12). The background O\textsubscript{3} and CO mixing ratios at Bode were estimated to be about 14 ppbv and 325 ppbv\textsubscript{2} respectively. The rate of change of ozone at Bode showed a more rapid increase (~17 ppbv/hour) during morning than the decrease in the evening (5-6 ppbv/hour), suggesting prevalence of a semi urban kind of environment at Bode. The slower evening time ozone decrease rates and lower CO levels during spring suggests an important contribution of that regional transport than the contribution of local sources to the also contributes appreciably to springtime ozone enhancement in the Kathmandu Valley. The winter time higher ozone levels at Bode are largely due to local sources with relatively less contributions from regional sources on top of the local in situ ozone production. We show that regional pollution resulting from agricultural crop residue burning in north-western IGP led to large (~2 fold) increases simultaneously in O\textsubscript{3} and CO levels at all three sites, i.e., Bode, Pantnagar and Nainital during first week of May 2013. Biomass burning induced increase in ozone and related gases was also confirmed by a global model and balloon borne observations over Nainital. A comparison of surface ozone variations and composition of light non-methane hydrocarbons among different sites indicated
the differences in emission sources of the Kathmandu Valley and the IGP. These results highlight the contribution of that it is important to consider regional sources to air pollution in quality management of the Kathmandu Valley versus the local sources in the Valley.

1. Introduction

The Himalayan region is among the least studied regions in the world despite its known importance in influencing the livelihood of hundreds of millions of about a billion people and agricultural systems. The Himalayan mountain regions are spread over a large region from Afghanistan, Pakistan, India, Nepal, Bangladesh, Bhutan, China, and Myanmar, and provide fresh water to about a billion people living in this region. However, the growing economies, industrialization and increasing population in the region are polluting this pristine environment and perturbing the regional environment, climate and ecosystems. The urban centers in the mountain regions often face severe air pollution problems since the mountains act as a barrier to horizontal ventilation of the pollutants and local mountain valley winds govern the diurnal variations in air pollutants. These processes have been well studied over other parts of the world, such as Mexico City (de Foy et al., 2006; Molina et al., 2007 etc.), Po Valley (Martilli et al., 2002) and Santiago de Chile (Schmitz, 2005; Rappengluck et al., 2005). The Kathmandu Valley, located in the central Himalayas is an ideal natural laboratory to study such processes. However, only a few surface measurements of ozone and related trace species have been reported so far from this region (Pudasainee et al., 2006; Panday and Prinn, 2009; Christofanelli et al., 2010; Putero et al., 2015; Mahata et al., 2017).

The valley has experienced an unprecedented growth as the population increased nearly fourfold from about 0.75 million to about 3 million over the last 25 years. The total vehicle fleet in the
Bagmati Zone, where the Kathmandu Valley is situated, increased by about 22 times from 34,600 in 1989-90 to 755,000 vehicles in 2013-14 (DoTM, 2015; http://www.dotm.gov.np/en). Consequently, the total fossil fuel usage in the valley is about 50% of all of Nepal. The shares of coal, petrol, diesel kerosene and liquefied petroleum gas (LPG) usage in the Kathmandu Valley ranges between 35% and 66% when compared with their respective usage in all of Nepal (Pradhan et al., 2012). These unprecedented growths can have serious implications for the air quality and its impacts in Kathmandu Valley, such as higher occurrences of respiratory problems, skin and eye irritation have already been observed among the people living in the Kathmandu Valley than in other areas (Pradhan et al., 2012). In the past, elevated levels of $O_3$, CO, $NO_x$ and VOCs have been reported over this region during winter and pre-monsoon seasons (Pudaisanee et al., 2006; Pandey and Prinn, 2009).

In order to advance our understanding of atmospheric composition and air quality in the Kathmandu Valley and surrounding broader regions, to understand how the magnitude and variability of air pollution in Kathmandu Valley compares with other sites in the region, the Sustainable Atmosphere for the Kathmandu Valley- Atmospheric Brown Clouds (SusKat-ABC) international air pollution measurement campaign was carried out in Nepal during December 2012-June 2013, with an initial intensive measurement period of two months from December 2012 to February 2013 (Rupakheti et al., 2017). Eighteen international research groups participated and various instruments for the extensive measurements of aerosols, trace gases and meteorological parameters were installed. The campaign covered a total of 23 sites of various measurement capabilities in the region with a supersite at Bode, 5 satellite sites in and on the Kathmandu Valley’s rim, 5 regional
sites (Lumbini, Pokhara, Jomsom, Dhunche and Pyramid) and other collaborating sites in India and China, including Nainital and Pantnagar in India. Measurements of short-lived climate-forcing pollutants (SLCPs, such as ozone and black carbon at Paknajol near the city center of Kathmandu during the SusKat-ABC campaign are reported in Putero et al. (2015). However, that study lacked the collocated measurements of O₃ precursors. Sarkar et al., (2016) presented the measurements of non-methane volatile organic compounds (NMVOCs) at one-second resolution using the Proton Transfer Reaction-Time of Flight-Mass Spectrometry (PTR-TOF-MS) and study on two greenhouse gases are described by Mahata et al., (2017) at Bode during the campaign. These studies provided important information about atmospheric composition in the Kathmandu Valley during the SusKat-ABC period; however, a regional picture of the atmospheric composition variability in ozone and related gases has not been presented so far.

In light of the above conditions, this study aims to provide first information about the regional distribution of ozone and related gases during the SusKat-ABC by analyzing simultaneously in situ measurements of surface ozone and CO at Bode from January to June 2013 with those from two Indian sites, namely Nainital (a high altitude site in the central Himalayas) and Pantnagar (a low altitude site in the Indo-Gangetic Plain (IGP)). Additional observations at the two Indian sites can also help to trace contributions of regional and local pollution are used to understand the similarities and differences between the air quality of Kathmandu Valley and the Indian sites, and identify any regional emission source common to these sites. The previous measurements of ozone in the Kathmandu Valley were only performed near the city centers; however, Bode is on the eastern side of the valley and is generally downwind of the major urban centers of Kathmandu Valley (Kathmandu Metropolitan City and Lalitpur Sub-
metropolitan City). This site also receives regional air masses from west and south especially during afternoons with stronger wind speeds. Therefore, this site can serve as a better representative to suggest the background levels of O\(_3\) and CO in the Kathmandu Valley and the contribution of emissions originating from local to regional scale.

2. Experimental details

2.1. Observation sites

The Kathmandu Valley is an oval-shaped urban basin located in the central Himalayan foothills between the IGP and the Tibetan Plateau (Figure 1). The valley is surrounded by mountain peaks with altitude ranging from 2000 to 2800 m above mean sea level (amsl) and five mountain passes (Nagdhunga, Bhimdhunga, Mudku Bhanjhyang in the West, Sanga and Nagarkot in the East) with altitude ranging from 1500 to 1550 m amsl, and the outlet of the Bagmati River in the southwest corner of the Valley. The flat base area of the Kathmandu Valley is about 340 km\(^2\) with a mean elevation of about 1300 m amsl. There is no river inlet into the Kathmandu Valley and only one narrow river outlet (Bagmati river) in the southwestern side. The spatial extent of the valley is about 25 km in East-West and around 20 km in the North-South direction. The measurement sites located in Kathmandu Valley during the SusKat field campaign are depicted in Figure 1b. In this study observations of O\(_3\), CO and meteorological parameters made at Bode (27.68\(^\circ\)N, 85.39\(^\circ\)E, 1344 m amsl) and at two sites in India viz., ARIES, Nainital; a high altitude
site located on a mountain top (29.36° N, 79.45° E, 1958 m amsl) and Pantnagar; located in the Himalayan foothills in IGP (29.0° N, 79.5° E, 231 m amsl) are discussed.

Two observations sites in India, i.e., Nainital and Pantnagar represent cleaner Himalayan and polluted IGP environments, respectively. Surface ozone levels at Nainital are found to be driven mainly by transport of anthropogenic emissions from the IGP region (Kumar et al., 2010; 2011), while those at Pantnagar are mostly controlled by local emissions (Ojha et al., 2012). In the previous studies, the pollutants level in the Kathmandu valley were reported to be primarily influenced by the local emissions and the unique meteorology in the region (Panday et al., 2009; Putero et al., 2015).

2.2. Ozone and CO instruments

Surface measurements of Surface ozone measurements are made using two types of analyzers from two make viz., i.e., Teledyne M400E (at Bode and Pantnagar) and Thermo Model-49i (at Nainital). The observation principle of both the instruments is based on the commonly used technique of attenuation of UV radiation (~254 nm) by ozone molecules. These instruments are regularly subjected to zero and span tests using an internal ozone generator and ozone observations from both the instruments are also inter-compared by running them side by side and using a common inlet. Further details of such inter-comparisons are reported in Sarangi et al. (2014).

CO measurements are also conducted using two types of analyzers from two make viz., Horiba APMA-370 (at Bode and Pantnagar) and Thermo 48i (at Nainital). CO instrument, at Bode, was
deployed for the first time in field after factory calibration from the manufacture. Nevertheless, both CO instruments were inter-compared using a common inlet prior to the campaign and correlation coefficient between CO mixing ratios measured by the two instruments is estimated to be ~0.9 with a slope of 1.09. The detection principle of these CO instruments is based on commonly used method of infrared (IR) absorption by CO molecules at 4.6 µm. Regular zero check and span check for CO instruments are performed using a primary calibration mixture from Linde UK (1150 ppbv; Sarangi et al., 2016) and secondary gas from Chemtron Science Laboratories (1790 ppbv). Multipoint calibrations (ultra-pure gases) are also carried out in different observational ranges using a zero air generator (Thermo model 1160) and a dynamic gas calibrator (Thermo model 146i) (Sarangi et al., 2014). The meteorological measurements at Bode are performed using an automatic weather station (Campbell Scientific, UK).

Both the O₃ and CO instruments were installed on the fourth floor of a building in Bode facing eastern side of the Kathmandu Valley (Figure 2; refer Sarkar et al., 2016 for site description). The sampling inlet for these instruments was placed at the top of the building and Teflon (TFE) tubes were used for the air intake. The O₃ and CO instruments at Nainital and Pantnagar were placed in atmospheric science building at Manora Peak [refer Sarangi et al., 2014 for site description] and at the College of Basic Sciences and Humanities (CBSH), G. B. Pant University of Agriculture and Technology (GBPUAT), at Pantnagar (refer Ojha et al., 2012 for site description). Observations at Bode are in Nepalese Standard Time (NST), which is 5:45 hours ahead of GMT, and observations at Nainital and Pantnagar are in Indian Standard Time that is 5:30 hour ahead of GMT.
2.3. Air sampling and analysis for hydrocarbons

Daily, a total of 16 air samples were collected at Bode from 30 December 2012 to 14 January, 2013 at Bode, making total 16 air samples collection with the frequency of one sample per day. These air samples are collected at 1400 hour (two samples at 1200 hour) when the boundary layer is fully evolved and the air is well mixed. Air samples are collected at a pressure of 1.5 bar and analyzed for ethane, ethene, propane, i-butane, n-butane, acetylene and i-pentane using a gas chromatograph (HP 5890 II) equipped with a flame ionization detector and a PLOT column of KCl/Al₂O₃. Helium is employed as carrier gas and H₂ & zero-air are used for flame. Air samples are also analyzed for CH₄ and CO using another GC (Varian Vista, 6000, USA) and employing a molecular sieve 13x, packed column (4 m). CO is measured by converting in to CH₄ using a Ni catalyst heated to about 325°C. Standard mixture from Intergas (International Gases & Chemicals), UK traceable to National Physical Laboratory (NPL), UK, is employed for calibration of NMHCs. Gases from NIST, USA and Linde, UK are used for calibration of CH₄ and CO. More details on sample pre-concentration and calibration can be seen in Lal et al., (2008), Mallik et al., (2014) and Sarangi et al., (2016).

2.4. Satellite data, model emissions, and back-air trajectory

In this study, Ozone Monitoring Instrument (OMI) level-3 daily tropospheric column amount NO₂ data product OMNO2d (cloud screened at 30%) at 0.25° x 0.25° resolution is used to generate spatial maps during biomass burning period. This product is based on the radiance measurements made by OMI instrument in visible (VIS, 405-465nm) channels. A detailed description of measurement principle can be found in Bucsela et al., (2013).
The surface level CO maps at 1.9° x 2.5° spatial resolution are generated using MOZART-4/GEOS-5 simulations. The model is driven by NASA GMAO GEOS-5 meteorological fields, anthropogenic emissions based on David Streets' inventory for ARCTAS and FINN fire emissions (Wiedinmyer et al., 2011) with MOZART-4 chemical mechanism (Emmons et al., 2010). The datasets are provided by the University Corporation for Atmospheric Research (UCAR) which includes its programs, the National Center for Atmospheric Research (NCAR) and its labs. The fire locations during the spring time biomass burning are used from monthly MODIS collection [Giglio, 2010]. For this study fire locations with high detection confidence (>80%) are used and detailed detection principle can be found in Giglio et al. (2003) and Justice et al., (2006). NASA’s Atmospheric Infrared Sounder (AIRS) instrument suite measures atmospheric water vapor and temperature profiles on a global scale. At present the operational instruments in this suite consists of a hyperspectral infrared instrument (AIRS) and a multichannel microwave instrument Advanced Microwave Sounding Unit (AMSU-A). Apart from meteorological datasets these instruments also retrieves the vertical profiles of some of the trace gases such as ozone and CO and detailed description of retrieval algorithms are discussed in Susskind et al., (2003, 2006). In this study the vertical profiles of ozone, CO and relative humidity from AIRS+AMSU joint data product (AIRX3STD v006) at 1 degree spatial resolution are used to understand stratosphere-troposphere transport phenomenon during Jan-May 2013.

The biomass burning emissions of CO were estimated using Global Fire Emissions Database (GFED) version 4 datasets. This data uses satellite information of fire hotspots, vegetation productivity to calculate gridded fire emissions. The data also included fractional contribution of
different types of vegetation to fire emissions. The version 4 dataset has spatial resolution of 0.25 degrees, and detailed description about the dataset and data access could be found from http://www.globalfiredata.org/data.html. The Modern-Era Retrospective analysis for Research and Applications, Version 2 (MERRA-2) is the reanalysis dataset produced by Global Modeling and Assimilation Office (GMAO) of NASA. In this study ertel potential vorticity data is used from this reanalysis and more details can be found in Gelaro et al., (2017).

To understand the synoptic scale atmospheric circulation and role of local and regional scale pollution in the variability of various trace species, four days isentropic back air trajectory analysis is performed using the history of air masses arriving at Bode. Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model (Draxler and Hess, 1999) is used. The model is driven by high resolution (0.5°x0.5°) meteorological fields from Global Data Assimilation System (GDAS) meteorological fields at 1° x 1°-spatial and 3-hour temporal resolution. In order to understand the origins of air masses, 0.5 degree is used to generate a 5-day backward trajectories over Bode. These trajectories are initiated as an ensemble of 9 points separated by 0.25 degrees around Bode at 1 km above ground level for 4 different months.

2.5. WRF-Chem Simulations

WRF-Chem version 3.5.1 with ~0.2° two-nested domains was used for this simulation. The coarse domain that encompasses the area between 16-43° N and 68-107° E, was use with 15-km grid spacing surrounding Bode region, at 2 km altitudes are used, the nested domain that covers the central part of Nepal and the Kathmandu Valley with 3-km grid spacing, and 35 vertically-stretched layers from the ground up to 50 hPa (Mues et al., 2017). The physics options used for
this study include the Lin microphysics scheme (Lin et al., 1983), the Grell cumulus parameterization (Grell and Dévényi, 2002), the Rapid Radiative Transfer Model (Iacono et al., 2008) for longwave and Goddard shortwave scheme (Chou and Suarez, 1994), the Yonsei University boundary-layer parameterization (Hong et al., 2006), and the MM5 scheme for the surface layer (Jimenez et al., 2012). The initial and boundary condition for meteorological fields are using the ERA-Interim data. Anthropogenic emissions were obtained from the HTAP V2 inventory (http://edgar.jrc.ec.europa.eu/htap_v2). Emission maps for CO and NOx are shown in supplementary material (Figure S1 and S2). The RADM2-SORGAM chemical mechanism is used to represent the gas-phase and aerosol chemistry. The photolysis rates were computed using the Fast Tropospheric Ultraviolet and Visible (FTUV) Radiation Model (Tie et al., 2003, Li et al., 2005). The dry deposition was calculated following Wesely (1989) resistance method. Biogenic emissions were computed on-line using the Model of Emissions of Gases and Aerosols from Nature (MEGAN) model (Guenther et al., 2006). The biomass burning emissions are based on Fire INventory from NCAR (FINN) (Wiedinmyer et al., 2011).

3 Results and Discussion

3.1 General Meteorology

The Kathmandu Valley is located in the central Himalayan region due north of the IGP and south of the Tibetan Plateau. The valley is influenced by the South-Asian monsoon and in general receives most of its precipitation during summer (June-September). The remaining seasons are relatively dry with spring or pre-monsoon season (March-May) being the hottest (Panday and Prinn, 2009). Figure 3 shows the diurnal variations in temperature, relative humidity (RH), solar
radiation, wind speed and wind direction at Bode during January and April 2013, which are chosen as representative months for winter (Jan-Feb) and spring (Mar-May), respectively. The average daily RH during January (67%) was higher than in April (62%), with the diurnal maximum during early morning hours. During the month of January, high RH in early morning hours (92% during 5-7 AM) was associated with the foggy conditions during morning hours.

The average temperature and solar radiation were higher in April (20.7°C and ~800 W/m²) than January (11.4°C and ~600 W/m²). The diurnal variations in temperature showed highest values during late afternoons which was few hours after the peak in diurnal solar radiation. The solar radiation together with the topography of Kathmandu Valley were responsible for strongly influence the diurnal mountain flows variations in temperature and out of the valley other meteorological parameters by surface heating causing thermals to rise. The wind speeds were the slowest (<1 m/s) during the night and early morning hours, primarily easterlies, and the highest wind speeds were observed during mid-late afternoon, primarily westerlies (4-6 m/s). These wind patterns were similar in both the months with April having slightly longer duration of daytime westerly flow. These wind flows, together with boundary layer dynamics are responsible for the dispersion/accumulation of pollutants during the course of a day.

The other sites in the Indian region viz., Pantnagar and Nainital also show high values of solar radiation and temperature during spring months and lowest during winter season (Ojha et al., 2012; Sarangi et al., 2014; Naja et al., 2016). The Nainital site which is at a remote mountain top and experiences moderate (2-3 m/s) northwesterly winds (~2-3 m/s) during most of the year with prevalence of southeasterly winds during the summer monsoon period. The Pantnagar site is
situated in the vicinity of Himalayan foothills in the IGP and the similar seasonal changes in temperature, RH and solar radiation are observed (Ojha et al., 2012). In the summer monsoon season, the lowest levels of O\(_3\) and CO are observed due to arrival of cleaner marine air masses.

In winter months, slow winds, lesser ventilation and the lowest boundary layer heights lead to widespread fog particularly are observed, and during the first week of January several cases of widespread fog are also observed.

3.2 Back-Air-trajectories

Four days, nine particles, back-air trajectories at Bode are shown for January, March, May and June (Figure 4). A very strong westerly flow is seen during winter (January) and then air-mass shows a gradual decrease of wind speed in beginning of spring (March). Air masses approach Bode even more slowly during May and pass over parts of the IGP and the Himalayas before arriving at Bode. A dramatic change in air flow is seen from May to June as air masses arrive mostly from the east at Bode. May and June shows very limited influences of long-range transport and air-mass is mostly within the Himalayan mountainous region with some influences from plain regions. The altitude of the air masses suggests for greater contribution from higher heights during winter, while air-masses are shown to be trapped in lower altitude region during spring (Figure 4).

To understand the wind patterns over Bode, the HYSPLIT model is used to generate back-air trajectories. Fig. 4 shows the monthly averaged nine-point trajectories (in different colors) for January, March, May and June around Bode. During winter and early spring months of January and March, a strong westerly flow is observed with air masses passing through IGP prior to entering Nepal region. During late spring (May) these westerly air masses slow down and a slight
change in directions are observed as some of the air masses passes through the central part of India. The air-masses were observed to be mostly arriving from higher altitude during winter than during spring season (not shown in Fig. 4). Similar behavior of air masses is also observed over Nainital (Kumar et al., 2010). However, during June, the monsoon circulations takes over and reversal in wind directions are observed (Fig. 4d). The above features for Bode are similar to those seen over Nainital and Pantnagar. The back air trajectories analysis for Nainital and Pantnagar shows that long-range transport (westerly wind) is a dominant factor during winter. However, air masses mostly circulate over the continental northern Indian region at low altitudes during spring and autumn seasons when local pollution plays an important role (Kumar et al., 2010; Ojha et al., 2012; Naja et al., 2016).

3.3 Variations in ozone at Bode

The monthly (January to June) average diurnal variations in ozone at Bode are shown in Figure 5. The diurnal variations in ozone show higher levels during daytime. This daytime build-up in ozone is consistently observed throughout the observation period, with relatively lesser buildup during June due to prevailing cloudy/rainy conditions. Additionally, there are only a few days of observations in June, when the campaign ended. The daytime increment in surface ozone is a typical feature of polluted sites and can be associated with daytime photochemical production of ozone from its precursors in the presence of sunlight (e.g., Kleinman et al., 1994) and/or through mixing of. Model simulated normalized average diurnal variations also show a clear daytime buildup in ozone rich air aloft during February and May (Figure S3 in supplementary material).
The very low levels of ozone (during winter months) were observed during night-time which can be attributed to titration of O$_3$ by NO. The boundary layer measurements made using a ceilometer during the campaign suggests lower heights during night time (Figure 6) with very low ventilation coefficients during this season (Mues et al., 2017). The sampling inlet for the gaseous measurements was on the rooftop about 20m from the ground level, the possibility of loss of ozone due to surface deposition in highly stratified nighttime boundary layer should be much less. Just around sunrise, a dip in ozone levels was also observed which is suggested to be due to its reaction with NO and NO$_2$ (which are produced by photo-dissociation of NO$_3$ and N$_2$O$_5$ at sunrise). A similar dip in ozone was also reported from an urban site in India (Lal et al., 2000).

Unfortunately, observations of NO and NO$_2$ are not available during the campaign but some information about the levels of these species can be obtained from the previous studies. Pudasainee et al. (2006) measured the NO, NO$_2$ during winter of 2003-2004. Yu et al (2009) also measured the NO, NO$_2$ and HONO during the similar period and showed two peaks at morning (07:00-08:00) and evening (19:00-20:00) in NO with maximum levels reaching as high as 60 ppb. Although the results are almost a decade old and the difference in O$_3$ magnitude is also observed (Figure S4 in supplementary material) but we expect the increase in NO levels over the valley because satellite retrieved tropospheric column NO$_2$ show an increase over this region (Figure S4).

Just around sunrise, a dip in ozone levels was observed. A similar dip in ozone was observed at an urban site in India (Lal et al., 2000) which is suggested to be due to its reaction with NO and NO$_3$. Since, observations of NO, NO$_2$, NO$_3$ and N$_2$O$_5$ are not made during the campaign, we have employed model results to explain this feature. Figure 7 show model simulated average diurnal variations in NO, NO$_2$, NO$_3$ and N$_2$O$_5$ during February and May 2013. NO mixing ratios are close
to zero during the nighttime because it rapidly reacts with O\textsubscript{3} to form NO\textsubscript{2}, which also explains higher NO\textsubscript{2} levels during nighttime. NO\textsubscript{3} and N\textsubscript{2}O\textsubscript{5} also show higher levels during nighttime because of the reactions of NO\textsubscript{2} with O\textsubscript{3}, of NO\textsubscript{2} with NO\textsubscript{3}, respectively. The sharp morning increase in NO mixing ratios correlates strongly with the sharp decrease in NO\textsubscript{3} and N\textsubscript{2}O\textsubscript{5} mixing ratios especially during February indicating that photodissociation of NO\textsubscript{3} (\textless 670 nm) and N\textsubscript{2}O\textsubscript{5} (280 < \lambda < 380 nm) releases NO back to the atmosphere. It is evident that NO\textsubscript{3} and N\textsubscript{2}O\textsubscript{5} peaks prior to NO and NO\textsubscript{2} and later they are removed.

At Bode, spring time higher levels of ozone with a broader peak was observed when compared with winter months, this can be attributed to the increase in incoming solar radiation which in turn increases the photochemical production of ozone. In addition to photochemical production of ozone, the boundary layer evolution during morning hours also contribute to rapid increase in ozone levels, since ozone rich air aloft gets mixed with near surface ozone depleted air (e.g Rao et al., 2003; Reddy et al., 2012) and thus increases the ozone levels. Photochemical production of ozone. Role of air-masses (local Vs regional contribution) will be discussed later in subsequent sections where differences in winter and spring variations will be clearer. The nighttime ozone mixing ratios during the campaign are the lowest during the winter season. During this period, lower boundary layer height was also observed (Figure 6) (Mues et al., 2017). The ventilation coefficient is a measure of transport and/or mixing of pollutants in the boundary layer. Due to very low wind speeds (<1 m/s) and shallow boundary layer height during night, low ventilation coefficients (<100 m\textsuperscript{2}/s) were observed during this season (Figure S5).

### 3.4 Variations in CO and Hydrocarbons at Bode
The monthly average diurnal variations in CO showed two peaks, one during morning and the other in the evening hours (Figure 5). The monthly mean values of O₃ and CO with standard deviation, maximum and minimum are given in Table 1. Averages of both these gases in four time periods are also given in Table 2. Diurnal variation in CO with two peaks is a typical pattern over a polluted site and such variations have been reported at different South Asian urban sites e.g., Ahmedabad (Lal et al., 2000), Kanpur (Gaur et al., 2014), Pune (Beig et al., 2007), Santiago de Chile (Rappengluck et al., 2005), and Chicago (Pun et al., 2003) etc. The major sources for CO in the valley are vehicular emissions, brick kiln emissions, domestic burning of biofuels for cooking and heating, and garbage burning etc. Out of these, vehicular emissions, cooking and heating occurs largely during two times a day, morning and evening, with some time differences. Additionally, the nighttime and early morning hours are characterized by very slow wind speeds, and a shallower boundary layer along with poor mixing.

It is to be noted that CO levels during morning peaks are greater than those during evening. This could be due to such higher peaks during morning times than those during evening times have also been observed by Panday and Prinn, (2009) and have been explained on the basis of overnight/early morning accumulation of CO emissions, due to poor ventilation and lower height of the boundary layer. After attaining its maximum in the morning, CO starts to decrease. During daytime the CO emissions are countered by the boundary layer evolution and dynamic processes such as flushing of CO and other pollutants by westerly winds (Figures 1 and 3) blowing throughout the afternoon across the valley through the eastern passes. The chemical loss of CO via reaction with OH radical could also contribute slightly in showing CO lower values in the daytime.
to lower daytime CO levels. However, OH measurements are not made during the campaign and it is not possible to confirm this aspect.

The mixing layer starts evolving after sunrise and reaches its peak median values of about 900-1200m during winter and spring seasons (Mues et al., 2017). The ventilation coefficient estimated (with 15 m wind data) during daytime suggested vertical mixing could occur from mid-afternoon till early evening hours when consistently high VC (> 1000 m²/s) was observed (Figure S3). During this period the daily maximum wind speeds (4-5 m/s at 15:00-17:00) were also observed which were mostly westerly and further decreases the CO levels with daily minimum levels (148-218 ppbv) observed during this period. Due to relatively higher wind speeds and associated advection before evening hours, the CO mixing ratios at 19:00-20:00 are about 30-50% lower than morning hours.

In addition to CO, observations of VOCs made at Bode (Sarkar et al., 2016) also showed similar diurnal variations with two peaks and having their levels up to about 15 ppbv. Similar to CO, many VOCs (e.g. Acetonitrile, Benzene, Furan, etc.) showed higher levels during morning, when compared to the evening peaks. We have also collected one air sample almost every day during 30 December 2012 – 14 January 2013 and analyzed them for light non-methane hydrocarbons (C₂-C₅) and for CH₄-CO. Average values along with standard deviation, minimum, maximum and number of samples are given in Table 3. Methane levels are much higher than the global average and Average methane levels for the measurement period at Bode is 2.55±.12 ppmv which are much higher than the global average and about 28% and 27% higher than measured at a Northern Hemisphere background site at Mauna Loa (1.84 ppmv), and at a remote site Mt. Waliguan, China.
(1.87ppmv), respectively (www.esrl.noaa.gov). The average values of other hydrocarbons varied from about 1 ppbv to about 4.4 ppbv. The maximum value is observed to be of propane (15.48 ppbv) and acetylene (14.35 ppbv). These highest mixing ratios are observed on 7 January 2013.

The mixing ratios of methane and all these eight light non-methane hydrocarbons at Bode are much relatively higher (ppbv) than those observed at Nainital (Sarangi et al., 2016 0.8-2.2 ppbv) and somewhat comparable with those at Panthagarr/Haldwani (December data) 0.8-3.7 ppbv and 1.6-4.2 ppbv respectively). Additionally, unlike Bode, none of these sites in India showed higher values of NMHC mixing ratios exceeding 10 ppbv like those at Bode. Methane mixing ratio is also higher at Bode (2.55 ppmv) than those observed at Nainital (1.89 ppmv). Figure 68 shows a comparison of contribution of eight light NMHCs at Bode, Nainital, Panthagarr (including another town Haldwani) (Sarangi et al., 2016) and Kanpur (Lal et al., 2008). Bode data are for December-January months, while data from rest of three sites are in December month. Composition at Bode shows difference with those at Indian sites. Propane (20%) and n-butane (13.5%) show greater contribution, while contribution of i-pentane (4.2%) is significantly lower at Bode when compared with India sites. Greater contribution of propane and n-butane indicates for some leakages of liquefied petroleum gases (LPG) in the Kathmandu Valley.

3.5 Correlation between ozone and CO

It has been discussed in the previous section that O3 and CO show some contrasting diurnal variations. Here we discuss about the correlation between O3 and CO during different months (Figure 79). The highest negative correlation is seen in winter period (r²=0.82 in January and r²=0.71 in February) and this negative correlation reduces gradually with the lowest value in May.
(r²=0.12). The reduction in the tendency of the negative correlation from January to May could be due to changes in emission patterns and the boundary layer mixing. This is discussed further in the subsequent paragraphs. Hourly average CO levels show a systematic decrease from ~2300 ppbv in January to about 680 ppbv in June, whereas ozone shows a tendency of increase. The daytime ozone levels during spring season are slightly higher (~62 ppbv) when compared to winter (~54 ppbv). This spring time increase in ozone levels is also reported by several other studies in northern part of the Indian subcontinent (Kumar et al., 2010; Ojha et al., 2012; Kumar et al., 2013; Gaur et al., 2014). The higher values of CO during winter season can be attributed to an increase in its emissions (domestic and garbage burning to keep warm in winter season) and their inefficient dilution due to poor mixing and shallower boundary layer (Mues et al., 2017). However, during spring season, reduction in some of these emissions and relatively well mixed daytime boundary layer leads to show lower CO levels. As mentioned previously, role of OH chemistry could be an additional contributor in lower CO levels in spring. Model simulated OH levels are found be higher in May, when compared with February. This would suggest great chemical loss of CO in spring (May).

Figure 210 shows daily variations in O₃ and CO during four different times i.e., 0300-0500 hours, 0730-0830 hours, 1300-1500 hours, and 2200-2300 hours. It is considered that 0300-0500 hours and 2200-2300 hours would provide information for the periods when photochemical production of ozone is absent, while 1300-1500 hours can be used to understand the behavior during the periods of high photochemical activity and fully evolved daytime mixed layer. Variations during 0730-0830 hours will provide the information during morning period. The stable nocturnal boundary layer just starts evolving during morning hours and air mass close to surface begins to
mix with air at higher heights. In general, CO levels (blue line) show a decrease from January to June during 0300-0500 hours, 0730-0830 hours, and 2200-2300 hours, while they do not show significant changes during 1300-1500 hours. In contrast, ozone levels (red lines) are increasing from January to May/June during all four time periods. The highest noontime ozone level is observed to be about 80 ppbv during January to March that increases to about 102 ppbv during April-May. The noontime ozone level comes down to about 46 ppbv in June, which is mainly due to beginning of the monsoon season that is characterized by the arrival of cleaner air could be dominating from the oceanic regions of the Bay of Bengal, the Arabian Sea and the Indian Ocean.

The increase in ozone from January to May is rather more during nighttime or early morning hours, when photochemical production of ozone is absent. This suggests an enhancement in the background ozone levels. Night time values might have more influences of the daytime air mass when compared to the early morning hours. Hence, probably, average ozone value (13.1 ± 1.2 ppbv) during early morning period of 0300-0500 hours would represent background ozone levels for this region. However, the emissions are directly influencing the CO levels. Hence, the estimated CO mixing ratio (325.4 ± 98.3 ppbv) during noontime (1300-1500 hours) could be considered as background levels for the Bode region. Since the noontime boundary layer at Bode is assumed to be well mixed and the fast westerly flows across the valley reduces the direct sampling of air-masses at Bode from its immediate emission sources scattered in the valley.

Figure 810 also shows correlation between O₃ and CO for different time periods. Weak negative correlation is seen during early morning (0300-0500 and 0730-0830 hours) or night hours (2200-
2300 hours), while a slight positive correlation is seen during noon period (1300-1500 hours). In general, the. During nighttime and early morning hours, lowest boundary layer height (150-200m) was observed. In an urban or semi-urban environment of Kathmandu valley, where NO₃ levels are not lower (Pudasainee et al., 2006) ozone titration takes place throughout nighttime. Figure 10 also indicate the increase in overnight CO levels which peaked during morning hours. These contrasting variations tends to show the negative correlations. However, a slight positive correlation is observed during noon period (1300-1500 hours) which is similar to what is generally, observed at high altitude sites (Kaji et al., 1998; Tsutsumi and Matsueda, 2000; Naja et al., 2003; Sarangi et al., 2014) and cleaner sites (e.g. Island sites, Pochanart et al., 1999) show a positive correlation between O₃ and CO.

3.6 Regional distribution of O₃ and CO during SusKat

Apart from O₃ and CO observations at Bode, simultaneous observations of these two gases were also made at the central Himalayan site in India (Nainital) and a site in Himalayan foothills in the IGP region (Pantnagar) and are discussed in this section (Figure 9). Pantnagar and Nainital, despite being different in altitude, the wind patterns over this region are mostly northerly or northwesterly during winter (Kumar et al., 2010; Ojha et al., 2012; Sarangi et al., 2014). Further, both the sides receive polluted air massed from the IGP during spring season. Whereas the cleaner marine air masses arrive at these sites during the summer-monsoon season. Average diurnal patterns in O₃ and CO mixing ratios are somewhat similar at Bode and Pantnagar having twin peaks in CO and daytime high levels of O₃. However, different variations (Figure 11) are observed at Nainital (green line), which being a remote high altitude site does not show any daytime photochemical buildup or nighttime loss in ozone. Further, the daytime ozone levels at Bode are
higher than those at Pantnagar during winter season, while these are comparable during spring season. Additionally, CO levels are also higher at Bode than those at Pantnagar during winter (Figure 911). A comparison of surface ozone measurements at Kanpur (India) showed a relative better agreement with Pantnagar (India) while measurement at Paknajol (Nepal) showed better agreement with Bode (Nepal) during both seasons (winter and spring) (Figure 911) indicating the differences in emission sources of Kathmandu Valley and the IGP.

The changes in ozone increase/decrease rates (ppbv/hour) are analyzed for all these five sites. Generally, the ozone increase/decrease rates are nearly symmetric during morning and evening at an urban site. However, it is asymmetric with slower changes occurring during afternoon/evening time at a rural or semi-urban sites (Naja and Lal., 2002). Ozone production is strongly dependent on amount of precursor gases and available sunlight. On the contrary, evening time ozone loss depends mainly upon its titration with NO, apart from surface deposition. This rate of change of ozone during morning and evening hours has been used as an indicator of chemical environment (rural or urban) over a site (e.g., Naja and Lal., 2002). Below, we discuss calculated ozone increase/decrease rates.

Figure 1012 shows that the wintertime rate of ozone increase in morning hours is much higher at Bode (about 17 ppbv/hour), when compared to Pantnagar (about 9 ppbv/hour). This suggests a rapid ozone buildup at Bode than at Pantnagar. In contrast, the ozone decrease rate is lower at Bode (5-6 ppbv/hour) when compared to the decrease rate at Pantnagar (about 14 ppbv/hour) during spring. This suggests rather slower ozone loss at Bode via NO titration, indicating somewhat lesser polluted kind of environment in Bode during spring. However, this
does not necessarily mean that NOx emissions are lower in the Kathmandu Valley. NO levels are reported to be as high as 60 ppbv (Yu et al. 2009). Another process driven by diurnal variations in winds could lead to slower evening ozone loss rates. Faster afternoon westerly winds flush the pollutants out of the valley every day, leaving less NOx to titrate ozone during evening hours. In contrast, slower winds at night allow overnight accumulation of precursor gases in nocturnal boundary layer of Kathmandu Valley that in turn can potentially contribute to next morning ozone build-up. Therefore, the slower decrease rate in evening time ozone and lower value of CO during spring, confirms somewhat lesser contribution of local pollutants in spring ozone enhancement in the valley. The regional contribution in this regard cannot be ruled out. Similar to the diurnal variations in average ozone, diurnal pattern in ozone change rates are similar at Pantnagar and Kanpur.

Back-air trajectory assisted analysis of ozone observations at Nainital in the central Himalayas show that the major role of regional/local pollution is in spring when regionally polluted ozone levels are estimated to be 47.1± 16.7 ppbv (Kumar et al., 2010). During spring, net ozone production over the northern Indian Subcontinent is estimated to be 3.2 ppbv/day in regionally polluted air masses in spring but no clear build-up is seen at other times of year. While the role of long-range transport is shown to be dominating in winter with contribution of about 8-11 ppbv of ozone.

Shorter duration of solar heating during winter leads to weaker dynamical processes including convective mixing of pollutants, which in turn confines the pollutants near to the surface. Additionally, the Kathmandu Valley is isolated inside the Himalayas and the only way for
pollutants to reach here is either via upslope flow of polluted air masses through the mountain
valleys or arrival of polluted regional air masses from the air aloft. Thus, we feel that the
wintertime higher levels of O₃ and CO mixing ratios at Bode are least likely influenced by the IGP pollution. Further similar trapping of pollutants during winter season are also reported by previous studies done over this region (Panday and Prinn, 2009). However, intense heating and stronger convective mixing could induce the IGP outflow to influence this valley region during spring season. Spring time ozone enhancement, due to IGP outflow, in the tropical marine region surrounding India has already been observed (Lal et al., 2013).

3.7 Influences of springtime northern Indian biomass burning

Every year northern Indian biomass burning emits large quantities of trace gases and aerosols and significantly affect the regional distribution of several trace species (Kumar et al., 2011; Sinha et al., 2014; Putero et al., 2014; Bhardwaj et al., 2016; Kumar et al., 2016). These studies showed the enhancement in O₃ and CO levels due to crop residue and forest fire burning in the IGP during pre- and post-monsoon seasons under favorable meteorological conditions. It has also been shown that the northern Indian biomass burning induced cooling on the surface (−27 W m⁻²) ozone and top of the atmosphere (−8 W m⁻²) in CO mixing ratios over Bode, a time series of high confidence (confidence > 80%) fire counts over the northern Indian subcontinent (25°-35°N, 70°-95°E) were analyzed. Based on MODIS fire counts, the central Himalayas. This cooling leads to an additional atmospheric warming of 19 W m⁻² (Kumar et al., 2011) from April 3 to May 31, 2013.
During the campaign, high fire activity period (HFAP) is defined when 3-day running mean of fire counts exceeds the median fire counts during the fire activity period (Kumar et al., 2011). The low fire activity period (LFAP) is defined as March 1-31 when very low fire counts are observed over northern Indian subcontinent. The changes in O$_3$-surface ozone and CO levels-mixing ratios during these two periods (HFAP and LFAP) are shown in figure 10 (right panel: center and bottom). The average fire emissions over the Kathmandu region (27°-28°N, 85°-86°E) using GFED v4.0 emission inventory are also calculated and shown in Figure 13 (top right). Since the HFAP is almost 4 weeks long, two different peaks in ozone and CO mixing ratios during April and May are selected (see highlighted peaks in Figure 13). During both the periods a prior-increase in fire counts were observed at Bode, Pantnagar and Nainital (Figure 11) in the first week of May. A global which is followed by higher levels of ozone and CO mixing ratios. Another reason to separately study these cases, is fire locations during these two periods. High CO and O$_3$ case in April are shown to be induced by the fires those are mostly located in Nepal region (dark red), while fire activity was very low in the northern Part of the IGP in April (Figure S6). Whereas, during May event (orange), high increase in fire counts (~ 2.5 fold) is observed over northern Indian subcontinent during the HFAP (Figure S6) and very low fire was seen in Nepal region. Therefore, these two cases could be studied separately to assess impacts of biomass burning over Bode and other sites. The changes during HFAP, LFAP and above two event cases in April and May are shown in Table 4.

During the first half of April daily averaged CO mixing ratios of 861 ppbv were recorded, which were about 200 ppbv higher than the daily averaged CO levels for April. Ozone mixing ratios during this period was about 9 ppbv higher than the daily averaged ozone levels for April. The
spatial maps of MODIS fire counts during the period were also analyzed (Figure 14: top panel) and an increase in fire counts during the period was observed. Based on MODIS fire counts and O₃ and CO time series at Bode, we classified 20-25th March as pre-event period and April 3-6 and April 11-16 as high fire activity periods. During the first high fire period, an increase of 55% and 15% in surface ozone and CO mixing ratios, was observed. The fire maps during this period also indicate that the majority of this fire were occurring near Bode in Nepal region and is shown in orange color in spatial map (Figure 14: Top Panel). The AIRS satellite retrieved CO mixing ratios at 925hPa and 850hPa also indicate an increase in CO levels near Bode (indicated as black star), however no similar increase is observed in CO levels near Pantnagar (or IGP). The majority of air masses at 1km AGL prior to arriving to Bode (or Nainital also) are westerly and passes through IGP/northern Indian subcontinent in general. Since there is almost no fire activity near Punjab region so we argue that these nearby fire and their associated polluted plumes around Kathmandu valley could affect the levels at Bode. Further, GFED biomass burning emissions were used to differentiate different types of biomass burning emissions during this period. During both of the events, the forest fire emissions dominated the total biomass burning emissions over Kathmandu region. The major land use types in black boxes indicated in Figure 14 are agricultural land or forests. The meteorological data observed during this period did not show any noticeable differences and boundary layer height during these events was close to monthly average values. Therefore, we can conclude that forest fires occurring nearby Kathmandu valley were responsible for high levels of pollutants during this period.

The second event took place near first week of May when simultaneous increase in surface O₃ and CO levels at Bode, O₃ levels at Nainital and CO levels at Pantnagar was observed. During this
event, O3 mixing ratios at Bode and Nainital increased by 73% and 67% respectively, and CO mixing ratios at Bode and Pantnagar increased by about 24% and 58% respectively. The MODIS fire maps during these events show a large increase in MODIS fire counts in Punjab region (blue box in top panel of Figure 15) in northwestern Indian subcontinent. During this period, total increase in fire counts is ~350% than that of April 25-30. The spatial distribution of AIRS retrieved CO mixing ratios at 925hPa and 850hPa, also indicate an overall increase in CO levels around northern part of Indian subcontinent. Similar increase is also observed in model (MOZART4/GEOS5) simulations also showed where about two-fold increase in CO levels at 992 hPa in during this period, is observed (Figure S7). Further, OMI tropospheric column NO2 (30% cloud screened) also showed small enhancement during the same period (Figure 12). To study the possible causes, MODIS fire product and the HYPLIT back-air trajectory data are analyzed. MODIS derived fire location showed about 256% increment in fire counts over the Punjab region in the IGP during 2-6 May, 2013 when compared with 28 Apr – 1 May fire counts. (Figure 12). Further, S7). The fire emissions associated during this period indicate that majority of emissions are from crop residue burning unlike the previous case in April. Late April-early May is the harvest period for crops in northern India, and wide spread crop residue burning is common during this period. This leads to release of massive amounts of pollutants over this region (Bhardwaj et al., 2016). The HYPLIT 4-day back-air trajectory analysis shows that the air masses arriving at Bode were mostly circulating over Nepal region (bottom left panel Figure 12) before the event period. However, during the high fire period the trajectories also indicate that air masses were also coming from the active fire region to Bode during the period were arriving from these source regions to the observing sites. During this period, nothing noticeable in wind speeds, directions, temperature, solar radiation, and rainfall is observed so we could conclude that this increase in surface mixing
ratios of O$_3$ and CO at these sites could be related to biomass burning in the northern Indian subcontinent. During this period, the influences are not only limited to surface level but also at higher altitudes where satellite retrieved vertical profiles of CO show high levels during this period. Rupakheti et al (2017) also reported a similar increase in the ambient concentrations of BC, PM, CO and Ozone at Lumbini (regional site of SusKat campaign) during these two episodes were observed (7-9th April and 3-4th May). This site is located southwards of Kathmandu valley Himalayan foothills near IGP. The notable changes in surface O$_3$ and CO mixing ratios and GFED biomass burning emissions over Kathmandu region are shown in Table-4.

An influence of fire activities has been observed in the vertical profile distribution of ozone over the central Himalayas. Details of balloon-borne observations of ozone (Ojha et al., 2014) and meteorological parameters along with inter-comparison of two kinds of meteorological sensors (i-Met and Vaisala) are given in Naja et al. (2016). The weekly balloon borne ozone profiles made from Nainital (on 9th May) also confirmed an enhancement in ozone (~16 ppb) in the lower troposphere (2-4 km) when compared with the ozone profile on 4th May (Figure 4). The enhancement is about 14 ppbv in 4-6 km region. Such events are generally observed during the spring season, when the influence of regionally polluted air masses from the IGP could travel over long distances.

3.8 Influences of Stratosphere-Troposphere exchange (STE) on surface ozone levels at Bode

The tropopause folding events and influences of STE over the Indian subcontinent and Tibetan plateau are more frequent during winter and early spring seasons (Cristofanelli et al., 2010; Chen et al., 2011; Phanikumar et al., 2017) and a few attempts in the past were made to understand the
role of STE over the Indian subcontinent (Mandal et al., 1998; Ganguly and Tzanis, 2011). Here, to understand the role of STE on surface ozone levels at Kathmandu vertical distribution of ertel potential vorticity (EPV) using MERRA v2.0 reanalysis, AIRS satellite retrieved ozone, relative humidity (RH) and CO were observed during Jan-May 2013 (Figure 17). The EPV distribution is represented in potential vorticity unit threshold (EPV > 1.6 PVU = 1.6 x 10^{-6} K m^2/Kg/sec) defined by Cristofanelli et al., (2006). Since the EPV near extratropical tropopause is at about 2 PVU and EPV in the stratosphere is about 1-2 magnitude higher, therefore, any values of EPV greater than 1.6 are suggested to be associated with the downward transport of ozone rich air masses from above (Cristofanelli et al., 2006). Overall, EPV distribution suggests downward transport associated with STE is limited to upper and middle part of the troposphere (Top Panel: Figure 17). The ozone distribution also exhibit similar behavior where increases in ozone levels (downward transport) reaching surface are negligible. However, a seasonal increase in ozone mixing ratios at lower and middle troposphere can be observed during Jan to May. To investigate the role of STE during April and May vertical profiles of EPV, O$_3$, CO and RH were studied for these periods (Figure 18). During April event (Fig. 18, Top Panel) EPV during April 11-16 was found to be increased as compared to prevent period (March) but it was reduced during April 3-6 period. During both the high fire periods ozone, CO were higher during the event which we propose is due to biomass burning since during STE reductions in CO are also observed. So therefore April event do not show any clear evidences of STE based on our analysis. Further, during May (Fig. 18, Bottom Panel) an increase in EPV at higher altitudes and O$_3$ at lower altitude is observed but again CO didn’t showed any decrease during this period. Therefore, we conclude that during both of these high pollution periods, no signs of downward transport is observed.
4. Summary

This study provides information about the regional distribution of O₃ and CO during the SusKat-ABC field campaign (Jan-Jun 2013) by analyzing simultaneous surface measurements of ozone and CO from Bode in the Kathmandu Valley with and from two Indian sites, Nainital and Pantnagar. Results from few air samples and their analysis for eight (C₂-C₅) light non-methane hydrocarbons are also presented. The diurnal variations show higher levels of ozone during daytime and morning/evening peaks in CO. This daytime build-up in ozone is consistent during all months, with a relatively smaller increment during the month of June due to prevailing cloudy or rainy conditions. Such a daytime increase in surface ozone is mainly due to the mixing of ozone rich air above the stable boundary layer and its photochemical production from precursor gases in the presence of sunlight. Very low nighttime levels of ozone were also observed during the winter season, which can be attributed to the titration of O₃ by NO. The diurnal variations in CO showed two peaks during morning and evening hours, due mainly to rush hour traffic sources and cooking activities, and such a similar distribution is also observed in Pantnagar. The evening peak was relatively less prominent at Bode, due to fast westerly winds blowing across the valley during daytime that flush out CO in contrast to calm nighttime winds and a shallow nocturnal boundary layer, resulting in the highest levels being observed during morning time. After reaching its maximum levels during morning time (up to 2300 ppbv in winter months), the levels decrease as the day progresses. This decrease is attributed to the boundary layer evolution and strong winds blowing across the Valley which dilutes the CO levels.
The correlations between $O_3$ and CO are found to be negative in the winter period ($r^2=0.82$ in January and $r^2=0.71$ in February) and this negative correlation becomes weaker gradually, with the lowest value in May ($r^2=0.12$). Hourly average CO levels also show a systematic decrease from its level of about 2100 ppbv in January to about 600 ppbv in June, whereas ozone shows the opposite tendency. A weaker negative correlation is observed during early morning (0300-0500 and 0730-0830 hours) or nighttime hours (2200-2300 hours) while a slight positive correlation is seen during the noon period (1300-1500 hours). The background $O_3$ and CO levels at Bode are found to be about 14 ppbv and 325 ppbv respectively. It is shown that $O_3$, CO and light non-methane hydrocarbon levels are higher at Bode than those at one of the IGP sites (Pantnagar) analyzed here and in the Himalayan foothills, particularly in winter. The rate of change of ozone during morning and evening hours is different at Bode, with a faster ozone increase rate during the day (about 17 ppbv/hour) but a slower ozone decrease rate (5-6 ppbv/hour) in the evening, suggesting Bode as a semi-urban site. The slower decrease rate of ozone in the evening time ozone and the lower value of CO during spring confirm a somewhat lesser contribution of local pollution in the springtime ozone enhancement in the valley, along with a contribution of regional scale pollution, the prevalence of a semi-urban kind of environment at Bode.

During the first week of May, simultaneous increases in $O_3$ and CO levels were observed at Bode, Pantnagar and Nainital. During the spring season, northern Indian biomass burning is found to affect the measurements sites in both India and Nepal. Two distinct events of biomass burning influence corresponding to the first half of April and May 2013 were studied. During both of these periods, an increase in $O_3$ and CO is observed over Bode. A similar increase is also observed at
Nainital during May, but not during April. During first week of April, a sharp increase of (~200 ppbv) in average CO mixing ratios is observed at Bode and increase is also observed in ozone levels. The analysis of spatial distribution of MODIS retrieved active fire locations indicated that the majority of fires took place in the nearby Nepalese regions that are upwind of Bode but downwind of Nainital. Analysis of biomass burning emission inventories indicated these emissions originated primarily from burning of the forests. Satellite retrievals of lower atmospheric CO mixing ratios also indicated an increase in CO levels during the event period near Bode region but not near Pantnagar and Nainital. During the first week of May, simultaneous increases in O₃ and CO levels were observed at Bode, and Nainital. The MOZART simulations during that period also indicate about a two-fold increase in near-surface CO levels. The MODIS-derived fire location showed a ~256% increment over the Punjab region in the IGP, which could emit large amounts. Analysis of precursor gases. Furthermore, biomass burning emission inventories indicated that fires during the event, the May 2013 originated mainly from the crop residue burning.

Analysis of back-air trajectories showed that majority of the air masses passed over the fires in Punjab before arriving at Nainital, Pantnagar and Bode—were coming from the Punjab region. Similar increases in near-surface distribution of satellite retrieved CO mixing ratios around all the sites are also observed. The balloon borne ozone profiles from Nainital also confirmed the significant enhancement in ozone (~16 ppbv) in the lower troposphere between the balloon flights on May 1st and 9th, 2013. Such events are mainly observed during the spring season when the influence of regionally polluted air masses from the IGP region are observed over measurement sites in the Himalayan region. This study has provided the first time measurements regional picture of O₃ and CO downwind of city centers in the Kathmandu Valley. In future, the air quality during the SusKat field campaign. Future studies should focus on long-term continuous and collocated
NOx and NMHCs measurements will be important to help understand the atmospheric chemistry better in this region. The SusKat dataset must also be used to identify and address the weaknesses of state-of-the-science air quality models and emission inventories, which together with observations will be play an important role in developing effective mitigation strategies for this region and ultimately reducing the vulnerability of public health to frequently occurring air pollution episodes.

Acknowledgments

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for their fruitful comments.
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### Table 1: Monthly variations in O₃ and CO over Bode, Nepal during January-June 2013.

<table>
<thead>
<tr>
<th>Month</th>
<th>Ozone (ppbv)</th>
<th>Max/Min (ppbv)</th>
<th>Daytime average ozone ppbv (1100-1700 hours)</th>
<th>CO (ppbv)</th>
<th>Max/Min (ppbv)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jan</td>
<td>23.5 ± 19.9</td>
<td>87.1/1.4</td>
<td>49.8 ± 10.2</td>
<td>832 ± 422</td>
<td>2323/218</td>
</tr>
<tr>
<td>Feb</td>
<td>25.6 ± 20.4</td>
<td>95/1.2</td>
<td>49.9 ± 13.9</td>
<td>717 ± 397</td>
<td>2182/162</td>
</tr>
<tr>
<td>Mar</td>
<td>37.4 ± 23</td>
<td>105.9/1.2</td>
<td>61.8 ± 12.0</td>
<td>698 ± 364</td>
<td>2011/158</td>
</tr>
<tr>
<td>Apr</td>
<td>43.5 ± 26.6</td>
<td>116.2/1.4</td>
<td>67.0 ± 20.4</td>
<td>667 ± 372</td>
<td>1969/175</td>
</tr>
<tr>
<td>May</td>
<td>38.6 ± 21.4</td>
<td>111.1/1.9</td>
<td>55.1 ± 18.9</td>
<td>401 ± 213</td>
<td>1656/146</td>
</tr>
<tr>
<td>Jun</td>
<td>31.1 ± 16</td>
<td>68.4/1.7</td>
<td>46.5 ± 8.5</td>
<td>303 ± 85</td>
<td>676/166</td>
</tr>
</tbody>
</table>
Table 2: Average (avg), standard deviation (std), maximum (max), minimum (min) and daily counts of O₃ and CO values and $R^2$ values during four time periods for the entire observational period (January – June 2013).

<table>
<thead>
<tr>
<th>Time period</th>
<th>0300-0500 hr</th>
<th>0730-0830 hr</th>
<th>1300-1500 hr</th>
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<tr>
<td>Ozone</td>
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<td></td>
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<tr>
<td>Avg (ppbv)</td>
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<td>13.9</td>
<td>58.9</td>
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<tr>
<td>Std (ppbv)</td>
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<tr>
<td>Max (ppbv)</td>
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<td>52.5</td>
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<td>2.0</td>
<td>25.9</td>
<td>1.4</td>
</tr>
<tr>
<td>Counts</td>
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<td>158</td>
<td>158</td>
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<tr>
<td>CO</td>
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<tr>
<td>Avg (ppbv)</td>
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<td>Std (ppbv)</td>
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<td>Max (ppbv)</td>
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<td>Min (ppbv)</td>
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<tr>
<td>Counts</td>
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<tr>
<td>$R^2$</td>
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<td>0.18</td>
<td>0.34</td>
<td>0.11</td>
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</tbody>
</table>
Table 3: Average, standard deviation, minimum and maximum mixing ratios (ppbv, except for CH₄, which is in ppmv) of CH₄, CO and eight (C₂-C₅) light-NMHCs from the analysis of daily air sample collection from 30 December 2012 to 14 January 2013. Percentage contribution of each NMHCs to the total of measured NMHCs is also given.

<table>
<thead>
<tr>
<th>Gases</th>
<th>Average</th>
<th>Standard deviation</th>
<th>Minimum</th>
<th>Maximum</th>
<th>% Contribution</th>
<th>Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methane</td>
<td>2.55</td>
<td>0.12</td>
<td>2.39</td>
<td>2.87</td>
<td>--</td>
<td>15</td>
</tr>
<tr>
<td>CO</td>
<td>392.5</td>
<td>109.3</td>
<td>272</td>
<td>588.8</td>
<td>--</td>
<td>16</td>
</tr>
<tr>
<td>Ethane</td>
<td>3.49</td>
<td>1.24</td>
<td>1.01</td>
<td>6.35</td>
<td>15.8</td>
<td>15</td>
</tr>
<tr>
<td>Ethene</td>
<td>2.84</td>
<td>2.37</td>
<td>0.31</td>
<td>9.69</td>
<td>12.9</td>
<td>15</td>
</tr>
<tr>
<td>Propane</td>
<td>4.41</td>
<td>4.14</td>
<td>0.44</td>
<td>15.48</td>
<td>20.0</td>
<td>13</td>
</tr>
<tr>
<td>Propene</td>
<td>1.06</td>
<td>0.91</td>
<td>0.28</td>
<td>3.86</td>
<td>4.8</td>
<td>13</td>
</tr>
<tr>
<td>i-Butane</td>
<td>2.26</td>
<td>1.93</td>
<td>0.24</td>
<td>7.78</td>
<td>10.3</td>
<td>13</td>
</tr>
<tr>
<td>n-Butane</td>
<td>2.96</td>
<td>1.80</td>
<td>0.18</td>
<td>5.81</td>
<td>18.5</td>
<td>14</td>
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<tr>
<td>i-Pentane</td>
<td>0.92</td>
<td>0.84</td>
<td>0.15</td>
<td>2.63</td>
<td>4.2</td>
<td>14</td>
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</tbody>
</table>
Table 4: The average O$_3$ and CO mixing ratios at Bode, Nainital, Pantnagar with GFED average biomass burning emissions over Kathmandu region during different periods.

<table>
<thead>
<tr>
<th>Fire Periods</th>
<th>Fire Count</th>
<th>Ozone (ppbv)</th>
<th>CO (ppbv)</th>
<th>Avg. Biomass burning emissions (Tg/day)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Bode NTL</td>
<td>Bode PNT</td>
<td>Total</td>
</tr>
<tr>
<td>LFAP (Mar 1- Mar 31)</td>
<td>5</td>
<td>37.4 45.2</td>
<td>705 455</td>
<td>8.96</td>
</tr>
<tr>
<td>HFAP (Apr- May)</td>
<td>70</td>
<td>43.7 63.9</td>
<td>504 374</td>
<td>63.70</td>
</tr>
<tr>
<td>Mar (20-25)</td>
<td>3</td>
<td>34.8 46.1</td>
<td>693 401</td>
<td>7.17</td>
</tr>
<tr>
<td>Apr (3-6)</td>
<td>18</td>
<td>46.7 49.2</td>
<td>797 250</td>
<td>57.74</td>
</tr>
<tr>
<td>Apr (11-16)</td>
<td>27</td>
<td>54.0 57.9</td>
<td>762 265</td>
<td>52.38</td>
</tr>
<tr>
<td>Apr (26-30)</td>
<td>26</td>
<td>33.9 45.0</td>
<td>505 313</td>
<td>22.10</td>
</tr>
<tr>
<td>May (2-6)</td>
<td>116</td>
<td>58.8 74.8</td>
<td>625 495</td>
<td>128.40</td>
</tr>
</tbody>
</table>
Figure 1: (a) Satellite image depicting the location of observation sites viz., Bode (Blue) in Kathmandu Valley, Nepal and Nainital (Green) and Pantnagar (Pink) in India during SusKat field campaign. (b) Satellite image of the Kathmandu Valley (edge on view) with the super site (Bode) and 4 satellite sites (Bhimdhunga, Naikhandi, Nagarkot, and Shivpuri). Figure (b) also indicates the position of five mountain passes surrounding the valley (yellow arrows) and one river outflow location (white). Kathmandu city is also marked by a black square.
### Observation Site Bode

#### Status of Filter

<table>
<thead>
<tr>
<th>Filter #3</th>
<th>Filter #4</th>
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</thead>
<tbody>
<tr>
<td>14-01-13 19:00 NST to 16-01-13 22:20 NST @Bode</td>
<td>16-01-13 22:20 NST to 18-01-13 13:30 NST @Bhimdhunga</td>
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</tbody>
</table>

#### Images

- **Observation Building**
- **O$_3$, and CO Analyzers**
- **Inlets**
**Figure 2:** The observation setup with O$_3$ and CO analyzers (top-left) placed at the fourth floor in a building at Bode, Nepal (bottom-left). The position of sampling inlets are towards eastern side of the valley (bottom-right). Almost black inlet filters are seen in about 2 days of operations at Bode and Bhimdhunga sites (top-right).
Figure 3: Average diurnal variations in temperature, solar radiation, relative humidity, wind speed and direction at Bode during January (upper panel) and April 2013 (bottom panel). These two months are taken as representative for winter and spring season respectively. (right) Wind directions during four time periods (0300-0500, 0900-1100, 1500-1700 and 1900-2100 hours) are shown as pie charts in wind speed plots.
Figure 4: Four Five days nine particles HYSPLIT back-air trajectories, with a monthly average pattern using 9 particles over Bode region during (a) January, (b) March, (c) May and (d) June. The colored trajectories are of monthly averaged for Bode. Altitude variations are also shown in each nine particles during the respective lower panels months.
Figure 5: Monthly variations in average diurnal O₃ and CO mixing ratios with 1-sigma spread at Bode during January-June 2013.
Figure 6: Observed (red) and model (green) simulated boundary layer height during winter and spring at Bode.
Figure 7: Model simulated average diurnal variations in NO, NO$_2$, NO$_3$ and N$_2$O$_5$ during February and May 2013.
Figure 8: Contribution of eight light non-methane hydrocarbons (NMHCs) to the total NMHCs at Bode, Pantnagar/Haldwani, Nainital and Kanpur in December. Bode includes January data too.
Figure 79: Relation between ozone and CO from January 2013 to June 2013 at Bode. Grey dots are hourly average data and black filled dots are 10 ppbv binned averaged with respect to ozone. The spread around the mean value is one sigma value.
Figure 810: Variations in ozone and CO during four-time periods at Bode, Nepal. Correlation between them is also shown (right).
Figure 9: Seasonal averaged diurnal variations in ozone and CO at Bode, Pantnagar, Nainital. Surface ozone observations at Kanpur (data from Gaur et al., 2014) and Paknajol (data from Putero et al., 2015) are also shown for the comparison. CO measurements were not available at Nainital during the spring season.
Figure 10.12: Diurnal variations in the average rate of change of ozone during winter and spring 2013 at Bode (blue), Paknajol (black; Putero et al., 2015), Pantnagar (Pink), Kanpur (brown; Gaur et al., 2014) and Nainital (green).
Figure 11: Time series of ozone and CO observations over the three sites viz., Bode, Nainital and Pantnagar. Red oval indicates the enhancement during the first week of May.
**Figure 12:** (Top panel) MOZART CO levels before the event period (28 Apr—1 May 2013; left) and during the event period (2-6 May 2013; right) at 992 hPa. OMI tropospheric column NO$_2$ during these two periods (Middle panels). HYSPLIT 4-day back air trajectories during April 30$^{th}$ and May 4$^{th}$ 2013. Black symbol in all these figures indicates Bode supersite, Nepal.
**Figure 13:** Top Left: time series of MODIS daily fire counts (red bar), 3-day running mean (black lines), median fire counts (brown line) for the fire period (3 April-31 May 2013). Total biomass burning emissions (orange line), crop residue burning emissions (dark yellow line) and forest fire emissions (dark green) over 1°x1° grid box around Bode (27-28°N, 85-86°E) are also shown. Top right: average biomass burning emissions for two fire activity periods over Bode region. Center: time series of surface ozone mixing ratios at Bode and Nainital (line plot-Left) and average ozone mixing ratios during two fire periods (as bar plot-Right), respectively. Bottom: time series of surface CO mixing ratios at Bode and Nainital (line plot-Left) and average CO mixing ratios during two fire periods (as bar plot-Right), respectively. The two fire events in April and May are also shown highlighted (in violet boxes).
**Figure 14:** Top (1a-3a): Spatial distribution of MODIS fire counts over Northern Indian subcontinent during three periods (left to right). The black boxes represent two fire hotspots (refer Fig. S6) over the region with fire counts shown as different colors during the three periods. Center (1b-3c): spatial distribution of AIRS CO mixing ratio at 925hPa (1b-3b) and 850 hPa (1c-3c) during three periods (left to right). Center (1d-3d): Average biomass burning emissions over Bode region (27-80E, 85-860E) using GFED v4.0 inventory during the three periods (left to right). Bottom Panels (1e-3f): changes in average surface mixing ratios of O3 (1e-3e) and CO (1f-3f) at different sites during the three periods (left to right).
Figure 15: Top (1a-2b): Spatial distribution of MODIS fire counts over Northern Indian subcontinent during two periods (left to right). The black boxes represent two fire hotspots (refer Fig. S1) over the region with fire counts shown as different colors during different periods. Center (1b-2c): spatial distribution of AIRS CO mixing ratio at 925 hPa (1b-2b) and 850 hPa (1c-2c).
during two periods (left to right). Center (1d-2d): Average biomass burning emissions over Bode region (27-8°E, 85-86°E) using GFED v4.0 inventory during the two periods (left to right). Bottom Panels (1e-2f): changes in average surface mixing ratios of O₃ (1e-2e) and CO (1f-2f) at different sites during the two periods (left to right).

Figure 13: The vertical profiles of ozone, temperature, RH, wind speed and direction over Nainital region on 1st (Red) and 9th (Blue) May 2013. The black lines show respective monthly average (May 2013) vertical profiles with bars representing one-sigma variations.
Figure 17: Vertical distribution of ertel potential vorticity (EPV) calculated by MERRA v2 reanalysis (top), AIRS retrieved ozone mixing ratios (centre), CO mixing ratios (center) and relative humidity (RH) during Jan-May 2013.

Figure 18: Vertical profiles of EPV, ozone, CO and RH during April (Top Panels) and May (Bottom Panels) biomass burning episodes.