1 Pre-monsoon air quality over Lumbini, a world heritage site

2 along the Himalayan foothills

- 3 Dipesh Rupakheti^{1,2*}, Bhupesh Adhikary³, Puppala S. Praveen³, Maheswar Rupakheti^{4,5},
- 4 Shichang Kang^{6,7*}, Khadak S. Mahata⁴, Manish Naja⁸, Qianggong Zhang^{1,7}, Arnico K. Panday³,
- 5 Mark G. Lawrence⁴
- ⁶ ¹Key Laboratory of Tibetan Environment Changes and Land Surface Processes, Institute of
- 7 Tibetan Plateau Research, Chinese Academy of Sciences, Beijing 100101, China
- 8 ²University of Chinese Academy of Sciences, Beijing 100049, China
- 9 ³International Centre for Integrated Mountain Development (ICIMOD), Kathmandu, Nepal
- ⁴Institute for Advanced Sustainability Studies (IASS), Potsdam 14467, Germany
- ⁵Himalayan Sustainability Institute (HIMSI), Kathmandu, Nepal
- ⁶State Key Laboratory of Cryospheric Science, Cold and Arid Regions Environmental and
- 13 Engineering Research Institute (CAREERI), Lanzhou 730000, China
- ⁷Center for Excellence in Tibetan Plateau Earth Sciences, Chinese Academy of Sciences, Beijing
- 15 100085, China
- ⁸Aryabhatta Research Institute of Observational Sciences (ARIES), Nainital, India
- 17
- 18 **Correspondence to:*
- 19 D. Rupakheti (dipesh.rupakheti@itpcas.ac.cn), S.C. Kang (shichang.kang@lzb.ac.cn)
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21 Abstract

Lumbini, in southern Nepal, is a UNESCO world heritage site of universal value as the 22 birthplace of Buddha. Poor air quality in Lumbini and surrounding regions is a great concern for 23 public health as well as for preservation, protection and promotion of Buddhist heritage and 24 25 culture. We present here results from measurements of ambient concentrations of key air pollutants (PM, BC, CO, O₃) in Lumbini, first of its kind for Lumbini, conducted during an 26 intensive measurement period of three months (April-June 2013) in the pre-monsoon season. The 27 measurements were carried out as a part of the international air pollution measurement 28 29 campaign; SusKat-ABC (Sustainable Atmosphere for the Kathmandu Valley - Atmospheric Brown Clouds). The main objective of this work was to understand and document the level of air 30 pollution, diurnal characteristics and the influence of open biomass burning on air quality in 31 Lumbini. The hourly average concentrations during the entire measurement campaign ranged as 32 follows: BC: 0.3 - 30.0 μg m⁻³, PM₁: 3.6-197.6 μg m⁻³, PM_{2.5}: 6.1 - 272.2 μg m⁻³, PM₁₀: 10.5 -33 $604.0 \ \mu g \ m^{-3}$, O₃: 1.0 - 118.1 ppbv, and CO: 125.0 - 1430.0 ppbv. These levels are comparable to 34 35 other very heavily polluted sites in South Asia. Higher fraction of coarse mode PM was found as compared to other nearby sites in the IGP region. $\Delta BC/\Delta CO$ ratio obtained in Lumbini indicated 36 37 considerable contributions of emissions from both domestic and transportation sectors. The 24-h average PM_{2.5} and PM₁₀ concentrations exceeded the WHO guideline very frequently (94% and 38 85% of the sampled period, respectively), which implies significant health risks for the residents 39 and visitors in the region. These air pollutants exhibited clear diurnal cycles with high values in 40 the morning and evening. During the study period, the worst air pollution episodes were mainly 41 due to agro-residue burning and regional forest fires combined with meteorological conditions 42 43 conducive of pollution transport to Lumbini. Fossil fuel combustion also contributed significantly, accounting for more than half of the ambient BC concentration according to 44 aerosol spectral light absorption coefficients obtained in Lumbini. WRF-STEM, a regional 45 46 chemical transport model, was used to simulate the meteorology and the concentrations of pollutants to understand the pollutant transport pathways. The model was able to reproduce the 47 temporal variation in the pollutant concentrations well; however, estimated values were 1.5 to 5 48 times lower than the observed concentrations for CO and PM₁₀ respectively. Model simulated 49 50 regionally tagged CO tracers showed that the majority of CO came from the upwind region of Ganges Valley. Model needs significant improvement in simulating aerosols in the region.
Given the high pollution level, there is a clear and urgent need for setting up a network of longterm air quality monitoring stations in the greater Lumbini region.

54 1. Introduction

The Indo-Gangetic plain (IGP) stretches over 2000 km encompassing a vast area of land in 55 56 northern South Asia: the eastern parts of Pakistan, most of northern and eastern India, southern 57 part of Nepal, and almost all of Bangladesh. The Himalayan mountains and their foothills stretch 58 along the northern edge of IGP. The IGP region is among the most fertile and most intensely farmed region of the world. It is a heavily populated region with about 900 million residents or 59 60 12% of the world's population. Four megacities - Lahore, Delhi, Kolkata, and Dhaka are located in the IGP region, with dozens more cities with populations exceeding one million. The region 61 has witnessed impressive economic growth in recent decades but unfortunately it has also 62 become one of the most polluted, and an air pollution 'hot spot' of local, regional and global 63 concern (Ramanathan et al., 2007). Main factors contributing to air pollution in the IGP and 64 surrounding regions include emissions from vehicles, thermal power plants, industries, biomass 65 and fossil fuel used in cooking and heating activities, agricultural activities, crop residue burning 66 and forest fires. Air pollution gets transported long distances away from emission sources and 67 across national borders. As a result, the IGP and adjacent regions get shrouded with a dramatic 68 annual buildup of regional scale plumes of air pollutants, known as Atmospheric Brown Clouds 69 70 (ABC), during the long and dry winter and pre-monsoon seasons each year (Ramanathan and Carmichael, 2008). Figure 1 shows the mean aerosol optical depth (AOD) acquired from the 71 MODIS instrument onboard TERRA satellite over South Asia for a period of December 2012-72 June 2013. Very high aerosol optical depth along the entire stretch of IGP reflects severity of air 73 74 pollution over large areas in the region.

Poor air quality continues to pose significant threat to human health in the region. In a new study of global burden of disease released recently, Forouzanfar et al. (2015) estimated that in 2013 around 1.7 million people died prematurely in Pakistan, India, Nepal, and Bangladesh as a result of air pollution exposure, nearly 30% of global total premature deaths due to air pollution. Air pollution also affects precipitation (e.g. South Asian monsoon), agricultural productivity,

ecosystems, tourism, climate, and broadly socio-economic and national development goals of the 80 countries in the region (Burney and Ramanathan, 2014;Shindell, 2011;Ramanathan and 81 82 Carmichael, 2008). It has also been linked to intensification of cold wave and winter fog in the IGP region over recent decades (Lawrence and Lelieveld, 2010 and references therein;Safai et 83 al., 2009; Ganguly et al., 2006). Besides high levels of aerosol loading as shown in Fig. 1, Indo-84 Gangetic plains also have very high levels of ground level ozone or tropospheric ozone (O_3) 85 (e.g., Ramanathan and Carmichael (2008)). It is a toxic pollutant to plant and human health, and 86 a major greenhouse gas (IPCC, 2013;Shindell, 2011;Mohnen et al., 1993). South Asia, in 87 particular IGP region, has been projected to be the most ozone polluted region in world by 2030 88 (Stevenson et al., 2006). Majority of crop loss in different parts of the world results from effects 89 of ozone on crop health and productivity (Shindell, 2011). Burney and Ramanathan (2014) also 90 reported a significant loss in wheat and rice yields in India from 1980 to 2010 due to direct 91 effects of black carbon (BC) and ozone (O_3). BC and O_3 are two key short-lived climate 92 pollutants (SLCP). Because of the IGP's close proximity to the Himalaya-Tibetan plateau region, 93 this once relatively clean region, is now subjected to increasing air pollution transported from 94 95 regions such as the IGP, which can exert additional risks to sensitive ecosystems in the mountain region (e.g., (Lüthi et al., 2015; Marinoni et al., 2013; Duchi et al., 2011). Air pollution transport 96 97 pathways to Himalayas are still not yet fully understood.

Monuments and buildings made with stones are vulnerable to air pollution damage 98 (Brimblecombe, 2003;Gauri and Holdren, 1981). Sulfur dioxide, which forms sulfuric acid upon 99 reaction with water is the most harmful substance for the monuments as it can corrode and 100 101 damage them (Baedecker et al., 1992;Gauri and Holdren, 1981). Indo-Gangetic plains are rich in archeological, cultural and historical sites and monuments and many of them are inscribed as 102 UNESCO World Heritage Site. For example, among many other such sites in IGP are the 103 Archaeological Ruins at Moenjodaro (Pakistan), Taj Mahal in Agra and Mahabodhi Temple 104 Complex in Bodh Gaya (India), Lumbini (Nepal), and ruins of the Buddhist Vihara at Paharpur 105 (Bangladesh) (World Heritage List; UNESCO, website: <u>http://whc.unesco.org/en/list</u>). The Taj 106 Mahal is one of the seven wonders of the modern world and India's greatest landmark. At the 107 end of the last century, the government of India realized the growing problem of air pollution 108 109 damage to Taj Mahal and started a program to save the monument. A recent study has reported that deposition of light absorbing aerosol particles (black carbon, brown carbon) and dust isresponsible for its discoloration (Bergin et al., 2015).

Lumbini, located near the northern edge of the central Ingo-Gangetic plain, is famous as the birthplace of the Lord Buddha. Lumbini is a UNESCO world heritage site of outstanding universal value to humanity, inscribed in the UNESCO list since 1997. The site, with valuable archaeological remains of the Buddhist *Viharas* (monasteries) and *Stupas* (memorial shrines), as well as modern temples and monasteries, is a center of attraction and visited by hundreds of thousands of pilgrims, scientists, scholars, yogis, and tourists every year. There was no regular air quality monitoring in Lumbini at the time of our measurement campaign.

119 Through this study, we want to understand the level of air pollution, its diurnal characteristic, 120 and the influence of open biomass burning on air quality in Lumbini. We carried out continuous 121 measurements of ambient concentrations of key air pollutants (particulate matter, black carbon, carbon monoxide, ozone) and meteorological parameters during an intensive measurement 122 period of three months (April-June) in the year 2013. These are the first reported pollutant 123 measurements for Lumbini. A regional chemical transport model called Sulfur Transport and 124 dEposition Model (STEM) was used to simulate the variations of meteorological parameters and 125 air pollutants during the observation period to understand pollution source region as well as the 126 contribution of open biomass burning to air quality in Lumbini. Model simulated regionally 127 tagged CO tracers were used to identify emission source regions impacting pollutant 128 129 concentration observed at Lumbini. Satellite data has also been used to understand the high pollution events during the monitoring period. These measurements were carried out as a part of 130 the SusKat-ABC international air pollution measurement campaign (M. Rupakheti, manuscript in 131 preparation for ACPD) jointly led by the International Centre for Integrated Mountain 132 133 Development (ICIMOD), Kathmandu, Nepal and Institute for Advanced Sustainability Studies 134 (IASS), Potsdam, Germany.

- 135 2. Experimental set up
- 136 2.1 Sampling site

137 The Lumbini measurement site (27°29.387' N, 83°16.745' E, elevation: ~100 m above sea level) is located at the premise of the Lumbini International Research Institute (LIRI), a Buddhist 138 library in Lumbini. Lumbini lies in the Nepal's southern lowland plain or *Terai* region, termed as 139 "bread basket of Nepal" due to the availability of very fertile land suitable for crop production, 140 which forms the northern edge of the Indo-Gangetic Plains (IGP). 25 km north of Lumbini the 141 foothills begin, while the main peaks of the Himalayas are 140 km to the north. The remaining 142 three sides are surrounded by flat plain land of Nepal and India. The site is only about 8 km from 143 the Nepal-India border in the south. A three storied 10 m tall water tower was used as the 144 platform for the automatic weather station (AWS) whereas remaining instruments were placed 145 inside a room near the base of the tower. Figure 2 shows the location of Lumbini, the Kenzo 146 Tange Master Plan area of the Lumbini development project, and the sampling tower. An 147 uninterrupted power back up was set up in order to assure the regular power supply even during 148 hours with scheduled power cuts during the monitoring period. The nearby premises of the 149 monitoring site consist of the LIRI main office and staff quarters. Further away is a museum, a 150 local bus park for the visitors to Lumbini, the office of the Lumbini Development Trust, 151 152 monasteries, and thinly forested area with grassland within the master plan area. Outside of the master plan area lie vast area of agricultural fields, village pockets, and several brick kilns and 153 154 cement industries. A local road (black topped), that cuts through the master plan area, lies about 200 m north of the sampling site and experiences intermittent passing of vehicles. According to 155 156 the Ministry of Culture, Tourism and Civil Aviation of Nepal over 130 thousand tourists (excluding Nepalese and Indian citizens) visited Lumbini 2014 157 the area in 158 (http://tourism.gov.np/en).

159 2.2 Monitoring Instruments

160 The summary of instruments deployed in Lumbini is presented in Table 1. All data were 161 collected in Nepal Standard Time (NST) which is GMT +05:45 hour. PM_1 , $PM_{2.5}$ and PM_{10} mass 162 concentrations were monitored continuously with GRIMM EDM164 (GRIMM Aerosol Technik, 163 Germany) which uses the light scattering at 655 nm to derive mass concentrations. Similarly, 164 aerosol light absorptions at 7 wavelengths (370, 470, 520, 590, 660, 880, 950 nm) were 165 measured continuously with an Aethalometer (Model AE-42, Magee Scientific, USA), averaging 166 and reporting data every 5 min. It was operated at a flow rate of 5 l min⁻¹. No cut-off was applied 167 for inlet; hence the reported concentration of BC is total suspended BC particles. As described by the manufacturer, ambient BC concentration is derived from light absorption at 880 nm using a 168 169 specific mass absorption cross section. To obtain BC concentration in Lumbini, we used a specific mass absorption cross-section value of 8 m² g⁻¹ for the 880 nm channel. A similar value 170 has been previously used for BC measurement in the Indo-Gangetic plain (Praveen et al., 2012). 171 172 To remove the filter loading effect, we used correction method suggested by Schmid et al. (2006) 173 which was also used by Praveen et al. (2012) for BC measurements at a rural site in the Indo-Gangetic plain. Surface ozone (O_3) concentration was measured continuously with an ozone 174 analyzer (Model 49*i*, Thermo Scientific, USA) which utilizes UV (254 nm wavelength) 175 photometric technology to measure ozone concentration in ambient air. CO analyzer (Model 48*i*, 176 Thermo Scientific, USA) was used to monitor ambient CO concentration which is based on the 177 principle that CO absorbs infrared radiation at the wavelength of 4.6 microns. The ambient air 178 was drawn through 6-micron pore size SAVILLEX 47 mm filter at the inlet in order to remove 179 the dust particles before sending air into the CO and O₃ analyzers using a Teflon tube. The filters 180 were replaced every 7-10 days depending on particle loading, based on manual inspection. CO 181 182 instrument was set to auto-zero at a regular interval of 6 hours. Local meteorological parameters (temperature, relative humidity, wind speed, wind direction, precipitation, and global solar 183 184 radiation) were monitored with an automatic weather station (AWS) (Campbell Scientific, Loughborough, UK), recording data every minute. 185

186 2.3

Regional chemical transport model

Aerosol and trace gas distributions were simulated using a regional chemical transport model. 187 Sulfur Transport and dEposition Model (STEM), a 3D eulerian model, that has been used 188 extensively in the past to characterize air pollutants in South Asian region was used to interpret 189 observations at Lumbini (Kulkarni et al., 2015;Adhikary et al., 2007). The Weather Research 190 and Forecasting (WRF) model (Skamarock et al., 2008) version 3.5.1 was used to generate the 191 192 required meteorological variables necessary for simulating pollutant transport in STEM. The model domain was centered at 24.94° N latitude and 82.55° E longitude covering a region from 193 3.390° N to 43.308° N latitude and 34.880° E to 130.223° E longitude. The model has 425×200 194 horizontal grid cells with grid resolution of 25×25 km and 41 vertical layers with top of the 195 196 model set at 50 mbar. The WRF model was run from November 1, 2012 to June 30, 2013.

However, for this study, modeled data only from April to June 2013 have been used. The WRF
model was initialized with FNL data available from NCAR/UCAR site
(http://rda.ucar.edu/datasets/ds083.2/).

The tracer version of the STEM model provides mass concentration of sulfate, BC (hydrophilic 200 and hydrophobic), Organic carbon (OC), sea salt (fine and coarse mode), dust (fine PM_{2.5} and 201 PM₁₀), CO (biomass and anthropogenic) and region tagged CO tracers. STEM model domain 202 size, resolution and projection are those of the WRF model. Details about tracer version of the 203 STEM model is outlined elsewhere (Kulkarni et al., 2015;Adhikary et al., 2007). Anthropogenic 204 emission of various pollutants (CH₄, CO, SO₂, NO_x, NMVOC, NH₃, PM₁₀, PM_{2.5}, BC and OC) 205 206 used in this analysis were taken from the EDGAR-HTAP_v2 (http://edgar.jrc.ec.europa.eu/htap_v2/index.php?SECURE=123). 207 Open biomass burning emissions on a daily basis during the simulated period were taken from data obtained from the 208 FINN model (Wiedinmyer et al., 2011). Both these emissions were re-gridded to the STEM 209 210 model domain. As with the WRF model, the STEM model was run from November 2, 2012 to June 30, 2013 however, data presented here are only during the intensive field campaign period. 211

212 3. **Results and discussions**

213 **3.1** Meteorology

Hourly average time series of various meteorological parameters like precipitation in mm hr⁻¹ 214 (Prec), temperature in °C (T), relative humidity in % (RH), wind speed in m s⁻¹ (WS) and 215 direction in degree (WD) during the monitoring period are shown in Figure 3. Meteorological 216 217 parameters were obtained with the sensors at the height of ~ 12 m from the ground. Meteorology results from simulations using WRF model have been used to compare and fill the data gaps. 218 Precipitation data was derived from TRMM satellite (TRMM_3B42_007 at a horizontal 219 resolution of 0.25°) from the Giovanni platform (http://giovanni.gsfc.nasa.gov/giovanni/) as the 220 rain gauge malfuntioned during the sampling period. Precipitation data from TRMM (Figure 3) 221 show that Lumbini was relatively dry in the early portion of the measurement campaign while as 222 the pre-monsoon edged closer to the monsoon onset, the site did experience some rainfall events. 223 This lowered aerosol loading in the later half of the measurement campaign due to washout. 224

Comparison of WRF model outputs with TRMM data shows that the model under-predictsrainfall through out the campaign.

Average observed temperature for the sampling period until the sensor stopped working (on 8th 227 May, 2013, i.e., for 38 days of measurement) was 28.1°C (minimum: 16.5°C, maximum: 40°C). 228 Average temperature from the model, during same period, was 31°C with values ranging 229 between 19 - 40°C. As shown in Figure 3, the model captures the synoptic variability of 230 temperature and is mostly within the range of daily values. However, the model has a high bias 231 and does not capture well daily minimum temperature values. The model does not show any 232 233 large variation in temperature for the period after the sensors stopped working. This insight will 234 be useful to interpret pollution data later on.

For the same period (until the sensor stopped working), the average (observed) RH was ~ 50% (ranging from 10.5 to 97.5%) whereas the model showed the average RH to be ~ 23% with values ranging between 6 to 78%. RH values are highly underestimated by the model however; the synoptic scale variability is captured by the model.

Average observed wind speed during the study period was 2.4 m s⁻¹, with hourly values ranging 239 between 0.03 - 7.4 m s⁻¹ whereas from the WRF model average wind speed was found to be 3.2 240 m s⁻¹ (range: 0.06 - 11.1 m s⁻¹). Diurnal variation of observed hourly average wind speed 241 suggested that wind speeds were lower during nights and mornings while higher wind speed 242 prevailed during day time, with average winds > 3 m s⁻¹ up to ~ 3.3 m s⁻¹ between 09:00-13:00 243 local time (Supplementary materials, Figure S1, lower panel). High speed strong winds (>4 m s⁻ 244 ¹) were from the NW direction during the month of April which later switched to almost opposite 245 direction, i.e., SE direction from the month of May onwards. Figure 4 shows the monthly wind 246 rose plot (using WRPLOT view from the Lakes Environmental, http://weblakes.com/). 247 Comparing modeled wind direction prediction skills at the surface with one point measurement 248 249 is not sufficient. However, in the absence of other measurements, we also show the comparison 250 of wind direction. Since there are no glaringly large biases in the observed surface wind 251 direction, and the lack of measured upper wind data even from nearby region, we use the model to interpret pollutant transport to Lumbini. Discrepancy on model results might have occurred 252 253 due to various factors inherently uncertain in a weather model. However, we believe that modeled data is vital for understanding pollutant transport in an area where observation data arenon-existent or are incomplete.

The monthly mean synoptic wind for the month of April, May and June is presented in Figure 5. NCEP/NCAR reanalysis monthly data of winds at 1000 mbar were used to study the wind pattern. The red dot in the figure indicates the location of Lumbini. NCEP/NCAR data showed the dominance of calm winds over the measurement site. Similar type of wind directions were observed over Kanpur, India, also in the IGP, during the pre-monsoon season (Srivastava et al., 2011).

262 **3.2** Air Quality

3.2.1 General overview, PM ratios and influence of meteorology on pollution concentrations

Figure 6 shows hourly averaged time series of observed BC, PM₁, PM_{2.5}, PM₁₀, CO and O₃ 265 observed at Lumbini during the study period. Similar temporal behaviour was shown by BC, 266 particulate matter fractions (PM1, PM2.5 and PM10) and CO. BC concentrations during the 267 measurement period ranged between 0.3-29.9 μ g m⁻³ with a mean (±SD) value of 4.9 (±3.8) μ g 268 m⁻³. BC concentrations in Lumbini during pre-monsoon months are lower compared to BC 269 270 concentrations observed in the Kathmandu Valley because of high number of vehicles plying on 271 the street, brick kilns and other industries in Kathmandu valley (Sharma et al., 2012; Putero et al., 2015). The lowest concentration was observed during a rainy day (21-22 April) whereas the 272 highest concentration was observed during a period of forest fire (detailed in Section 3.4). For 273 274 the entire measurement period, we found average (of hourly average values) PM₁: 35.8±25.6 µg m⁻³ (minimum-maximum range: 3.6 - 197.6 μg m⁻³), PM_{2.5}: 53.1±35.1 μg m⁻³ (6.1 - 272.2 μg m⁻³) 275 ³), PM₁₀: 128.9±91.9 μ g m⁻³ (10.5-603.9 μ g m⁻³) and coarse-mode (PM_{10-2.5}): 75.65±61.67 μ g m⁻³ 276 3 (1.98-331.80 µg m⁻³). The coarse-mode fraction was ~ 60% of the PM₁₀. The share of coarse-277 mode aerosol to PM_{10} in Lumbini was higher than that observed in other sites in the IGP; 278 279 Guwahiti, India (42%) (Tiwari et al., 2017) and Dibrugarh, India (9-16%) (Pathak et al., 2013) both in eastern IGP and Delhi (38%) (Tiwari et al., 2015) in western IGP indicating the higher 280 contribution of coarse aerosols in Lumbini, likely lifted from soils from nearby agricultural fields 281 and construction materials by stronger winds during pre-monsoon season. Similar value of 282

coarse-mode fraction, as in Lumbini, has been reported by Misra et al. (2014) at Kanpur for dust
dominated and mixed aerosols events.

The share of BC in PM fractions were found to be ~13% in PM₁, 9% in PM_{2.5} and ~4% in PM₁₀ 285 but the correlation coefficients of BC with three PM fractions were found to be 0.89 (PM₁), 0.88 286 287 $(PM_{2.5})$ and 0.69 (PM_{10}) , indicating the commonality in the sources of these pollutants. The contribution of BC in PM₁ was found to be of ~12% in Kanpur during February-March (Kumar 288 et al., 2016a) similar to Lumbini. Regarding the share of BC in PM_{10} , the share observed in 289 Lumbini (~4%) was similar to that observed over Varanasi (~340 km due south of our site) in 290 central IGP (5%) (Tiwari et al., 2016) and Dibrugarh in eastern IGP (~5%) (Pathak et al., 2013). 291 292 Thus our results indicate that despite our station being located at the northern edge of the IGP along the foothills of the Himalayan range, its aerosol characteristics are similar to those found in 293 294 heavily polluted sites in the central and eastern IGP.

In Lumbini, the average (hourly) share of PM₁ in PM_{2.5}, PM₁ in PM₁₀ and PM_{2.5} in PM₁₀ were 295 found to be $\sim 70\%$, 34% and 47% respectively. The share of average (sampling period) coarse-296 mode aerosols to PM_{10} (60%) was found to be higher as compared to that of average fine mode 297 i.e., PM_{2.5} (40%). Regarding other sites in IGP region, PM_{2.5}/PM₁₀ ratios were reported to be 298 56% in Kanpur (Snider et al., 2016), 60% in Varanasi (Kumar et al., 2015), 57% in Guwahiti 299 (Tiwari et al., 2017), 90% in Dribugarh (Pathak et al., 2013) and 62% in Delhi (Tiwari et al., 300 2015) indicating local differences within IGP as well as suggesting that influence of combustion 301 302 sources at Lumbini is still lower compared to other locations in Indian section of the IGP. A recent study (Putero et al., 2015) reported the PM₁/PM₁₀ during pre-monsoon of 2013 was found 303 to be 0.39 in the Kathmandu Valley of Nepal. Lumbini has significantly lower vehicle emissions 304 and human population than the Kathmandu Valley yet the ratios are similar, indicating the 305 306 importance of regional combustion sources in Lumbini for finer aerosols (PM₁), and soil-based 307 emissions such as road dust in the Kathmandu Valley. Future studies will need to explore the emission sources around Lumbini in much greater detail. Lower PM_{2.5}/PM₁₀ in Lumbini as 308 compared to other regions mentioned earlier could be due to emissions from cement industries 309 located within 15 km distance from the measurement site. Cement factories emit coarse sized 310 311 particles but we are not able to distinguish in our measurement without having an analysis if certain marker species. Trivedi et al. (2014) reported a ratio of 0.39 (during pre-monsoon) over 312

- Delhi, which is lower than the ratio in Lumbini. The lower ratio in Delhi was due to the presenceof coarse sized windblown desert dust and suspended soil materials due to strong winds.
- 315 The observed 24-hour average particulate matter concentrations ($PM_{2.5}$ and PM_{10}) were found
- frequently higher than the WHO prescribed guidelines for $PM_{2.5}$ (25 µg m⁻³) and PM_{10} (50 µg m⁻³)
- 3) with PM2.5: exceeding 94% and PM₁₀: 85% of the measurement period of 53 days.
- Observed CO concentrations ranged between 124.9-1429.7 ppbv with an average value of 318 319 344.1±160.3 ppbv. CO concentration observed in Lumbini is lower than that of Mohali, Western 320 India where the average concentration was 566.7 ppbv during pre-monsoon season due to intense 321 biomass and agro-residue burning over the region (Sinha et al., 2014). Temporal variation of CO 322 concentrations is similar to that of BC as both of these species are emitted during incomplete 323 combustion of fuel. Moreover, a very strong correlation (r = 0.9) was observed between BC and CO. Past studies have shown that the ratio of BC to CO depends upon multiple factors like site 324 location, combustion characteristics (fuel and technology) at the sources, and type of air mass 325 (Girach et al., 2014; Pan et al., 2011; Zhou et al., 2009). Formation of the soot depends on the 326 carbon to oxygen ratio of fuel whereas CO can also be produced naturally due to the oxidation of 327 VOCs (Girach et al., 2014). Figure 7 shows the comparison of the average $\Delta BC/\Delta CO$ ratio 328 (0.021) at Lumbini with that obtained from other sites. Please refer to Figure S2 in the 329 supplementary materials for the time series of $\Delta BC/\Delta CO$ ratio observed in Lumbini. We used the 330 method described by Pan et al. (2011) to calculate the $\Delta BC/\Delta CO$ values. The ratio was 331 calculated using the equation $(BC-BC_0)/(CO-CO_0)$ assuming the background values $(BC_0 \text{ or }$ 332 CO_0) as 1.25 percentile of the data. The $\Delta BC/\Delta CO$ ratio in Lumbini is similar to that obtained at 333 a suburban site, Pantnagar in India (0.017) (Joshi et al., 2016) and in Maldives (0.017) 334 (Dickerson et al., 2002). As compared to Lumbini, the different $\Delta BC/\Delta CO$ ratio obtained over 335 336 megacities such as Beijing and Shanghai are due to the higher number of gasoline and diesel 337 vehicles (Zhou et al., 2009). However, the ratio obtained at Lumbini were within the range of emission ratios from diesel used in transport sector (0.0013-0.055), coal (0.0019-0.0572) and 338 biofuels (0.0087-0.0266) for domestic activities (Verma et al., 2010 and references therein). The 339 hourly averaged observed ozone concentration ranged between 1.0 and 118.1 ppbv with a mean 340 value of 46.6±20.3 ppbv during the sampling period. The 8-hr maximum O₃ concentration 341 exceeded WHO guidelines of 100 $\mu g~m^{\text{-3}}$ (WHO, 2006) during 88% of the measurement period. 342

Our results clearly indicate that the current pollution levels in Lumbini is of great concern to health of the people living in the region as well as over a million visitors who visit Lumbini, as well as ecosystems, particularly agro-ecosystem, especially in warm and sunny pre-monsoon months.

347 The relationship of wind speed (WS) with aerosol and gaseous pollutants in Lumbini is shown in Figure S3 (Supplementary information). We were interested in studying the relationship between 348 wind speed and the pollutants since the wind governs the horizontal dilution of the pollutants 349 (Huang et al., 2012) and also likelihood of lifting soil dust. Except ozone, all other pollutants 350 351 exhibited negative correlation with wind speed. BC shows negative correlation (r = -0.42) with the wind speed which is similar with other pollutants as well (as can be seen from the figure). 352 Past studies have also reported a similar negative correlation of BC with wind speed over urban 353 and sub-urban areas (Huang et al., 2012;Cao et al., 2009;Ramachandran and Rajesh, 354 2007;Sharma et al., 2002;Tiwari et al., 2013) indicating that the locally generated BC can 355 356 accumulate in the atmosphere during lower wind speed conditions (Cao et al., 2009). Tiwari et al. (2013) also reported similar negative correlation (r = -0.45) during the pre-monsoon season 357 358 over Delhi. On the other hand, secondary pollutants like ozone exhibited a positive relation with the WS (r=0.38) indicating the WS could be one of the potential factors of high ozone in 359 360 Lumbini. Solar radiation is one of the most important factors for production of ozone in the atmosphere (Naja et al., 2003). The correlation of hourly ozone concentration with solar 361 362 radiation (not shown here) was found to be 0.41 whereas wind speed during the daytime only 363 (06:00-18:00) showed very weak correlation of 0.02 with ozone, indicating the calm condition as 364 conducive to formation and accumulation of ozone in the region.

Interestingly, the highest concentrations of all measured pollutants were obtained when the wind 365 speed was less than 1 m s⁻¹. In a separate analysis (not shown here), we considered only the WS 366 >1 m s⁻¹ and calculated the correlation coefficients to investigate the influence of regional 367 emissions. We found the similar correlation values as previous when all WS values were 368 considered (BC vs WS = -0.41, CO vs WS = -0.42, O₃ vs WS = 0.29, PM₁ vs WS = -0.40, PM_{2.5} 369 vs WS= -0.38, PM₁₀ vs WS= -0.33). The correlation of WS (>1 m/s) with concentration of air 370 pollutants elucidates that air pollution over Lumbini is not only of the local origin, it is rather 371 transported from other nearby regions as well. 372

373 Past studies near this site have been focused on the cities like Kathmandu (Sharma et al., 2012;Ram et al., 2010;Panday and Prinn, 2009;Putero et al., 2015) and Kanpur (Ram et al., 374 375 2010) and agro-residue burning dominated regions of IGP (Rastogi et al., 2016;Sinha et al., 2014;Sarkar et al., 2013) or a remote mountain location in India (Naja et al., 2014). Very high 376 aerosol loading is observed in South Asia during pre-monsoon, mostly over the IGP region 377 (Supplementary materials, Figure S4). As this is the first study over an IGP site located in Nepal, 378 pollution concentrations observed at Lumbini were compared with other sites in the region 379 (Table 2). Different sites located at urban, semi-urban and remote locations were used for 380 comparison to get a clear comparative picture of the situation at Lumbini amongst other locations 381 in the region. Pre-monsoon seasonal average PM_{2.5} concentration in Lumbini has been found to 382 be lower than the megacity like Delhi (Bisht et al., 2015) and north-western IGP (Sinha et al., 383 384 2014), possibly due to higher level of emissions (from traffic and biomass burning, respectively) over those regions. In addition, average BC and CO concentrations in Lumbini were found 385 386 falling in between concentrations observed at rural sites (up to 6 times higher) and cities in the region (see Table 2), indicating that Lumbini can still be considered as semi-urban location. The 387 388 hourly average O₃ concentration in Lumbini were found to be higher than the cities like Kathmandu (Putero et al., 2015) and Kanpur during pre-monsoon season (Gaur et al., 2014). 389 390 However from a mesoscale perspective, the hourly average O_3 concentrations were lower at Lumbini as compared to base camp of Mt. Everest region due to the uplift of polluted air masses 391 392 (Marinoni et al., 2013), stratospheric intrusion (Cristofanelli et al., 2010) and even the regional or long-range transport of the air pollutants (Bonasoni et al., 2010) to the high altitude site. 393

Regarding the monthly average concentration, the concentrations of all measured pollutants decreased as the pre-monsoon months advanced. The monthly average concentrations of the monitored species are shown in the Figure S5 along with the monthly fire hotspots over the region. Reduction in concentration (except PM) during the month of May (as compared to April) could be attributed to the fewer fire events during May as well as previously discussed washout by rainfall. Two peak pollution episodes were observed during the first half of April and May which is discussed in more detail in the next section.

401 **3.2.2** Observation-model inter-comparison

402 Chemical transport models provide insight to observed phenomena; however, interpretation has 403 to take into account model performance before arriving at any conclusion. This section describes 404 pollution concentrations simulated by the WRF-STEM model. A comparison of model calculated pollutant concentration along with the minimum and maximum concentrations of various 405 pollutants (with observation) is shown in Table 3. The model based concentrations used here are 406 instantaneous values for every third hour of the day. BC concentrations ranged between 0.4-3.7 407 μ g m⁻³ with a mean value of 1.8±0.7 μ g m⁻³ for a period of 1st April-15th June 2013. The average 408 model BC concentration was ~2.7 times lower than the observed BC. Regarding PM₁, PM_{2.5} and 409 PM₁₀, the model simulated average concentration was 12.3 ± 5.5 (0.9-41.7) µg m⁻³, 17.3±6.7 (1.9-410 48.3) μ g m⁻³ and 25.4 \pm 12.9 (2.1-68.8) μ g m⁻³ respectively. The model estimated values were 411 lower by the factor of 3 and 5 respectively than the observed concentrations. The data show that 412 model needs much improvement in its ability to adequately predict observed aerosol 413 characteristics. Since pollutant concentration is a function of emissions, transport and 414 transformation and deposition, improvements in any of these areas would improve the model. 415 However, given observation insights by PM ratios, it seems that improvements are much needed 416 417 in the emissions of primary aerosols. Current emissions does not account for trash burning, roadside dust and increasing newer industries, especially emissions from cement factories that 418 419 have propped up in recent years.

420 Average observed CO concentration was 255.7±83.5 ppbv, ranging between 72.2-613.1 ppbv, 421 with average model CO ~1.35 times lower than observed. Comparison of modeled CO versus 422 observation is shown in Figure 6. Apart from two peak episodes the model does a better job in 423 predicting CO concentration over Lumbini. Previous study using the STEM model over Kathmandu valley showed that the model was able to capture annual BC mean value but 424 completely missed the concentrations during pre-monsoon and post monsoon period (Adhikary 425 et al., 2007). Similar behavior is seen this time for CO where the model misses the peak values 426 427 but reasonably captures CO concentration after mid-May. STEM model CO performance can be significantly improved via better constraining emissions of open biomass burning as discussed in 428 Section 3.3. This activity is beyond the scope of this current paper although the improvements 429 430 are underway for all these sectors.

431 **3.2.3** Diurnal variations of air pollutants and boundary layer height

In the emission source region, diurnal variations of primary pollutants provide information about 432 433 the time dependent emission activities (Kumar et al., 2016b). Figure 8 shows the diurnal 434 variation of hourly averaged concentrations of measured pollutants during the sampling period. Primary pollutants like **BC**, **PM** and **CO** showed typical characteristics of an urban environment, 435 i.e., diurnal variation with a morning and an evening peak. However, Lumbini data shows higher 436 concentrations in the evenings compared to morning hours. Elevated concentrations can be 437 linked to morning and evening cooking hours for BC and CO where emission inventory show 438 that residential sector has significant contribution. However, explanation for elevated evening 439 concentration compared to morning needs further investigation. Increase in the depth of 440 boundary layer, reduction in the traffic density on the roads, absence of open biomass burning 441 during mid-day and increasing wind speed often contribute to the dispersion of pollutants 442 resulting in lower concentration during afternoon. Diurnal variation of wind direction 443 (Supplementary information, Figure S1, upper panel) shows the dominance of wind coming from 444 south (mainly during the month of May and till mid-June). Morning and evening period 445 experienced the winds coming from the southeast direction while the winds were predominantly 446 447 from southwest direction during late afternoon. Increase in CO concentrations in the evening hours might be due to transport of CO from source regions upwind of Lumbini which along with 448 449 the local emissions which gets trapped under lower Planetary Boundary Layer (PBL) heights in evening and night time. Ozone concentration was lowest in the morning before the sunrise and 450 451 highest in late afternoon around 15:00 PM after which concentrations started declining, exhibiting a typical characteristic of a polluted urban site. Photo-dissociation of accumulated 452 NO_x reservoirs (like HONO) provides sufficient NO concentration leading to the titration of O_3 453 resulting in minimum O₃ just before sunrise (Kumar et al., 2016b). The PBL height (in meters 454 455 (m)) was obtained from the WRF model as observations were not available. Figure 9 shows the diurnal variation of the model derived PBL height. The study period average PBL height over 456 Lumbini was ~ 910 m (ranging between 24.28 and 3807 m observed at 06:00 and 15:00 457 respectively). As the pre-monsoon month advances, PBL height also increased. The monthly 458 459 average PBL height was 799 m, 956 m and 1014 m respectively during the month of April, May and (1st-15th) June. The fluctuations of modeled PBL height correspond well with the diurnal 460

461 variation of observed pollutants like BC, CO and PM with the period of lower boundary height462 experiencing higher pollution concentration.

463 **3.3** Influence of forest fires on Lumbini air quality

3.3.1 Identification of forest fire influence over large scale using in-situ observations satellite and model data

Forest fires and agricultural biomass burning (mostly agro-residue burning in large scale) are 466 common over the South Asia and the IGP region during pre-monsoon season. North Indo-467 Gangetic region is characterized by fires even during the monsoon and post-monsoon season 468 469 (Kumar et al., 2016b;Putero et al., 2014). These activities influence air quality not only over nearby regions but also get transported towards high elevation pristine environments like Mt. 470 Everest (Putero et al., 2014) and Tibet (Cong et al., 2015a;2015b). So, one of the main objectives 471 of this study was to identify the influence of open burning on Lumbini air quality. Average wind 472 speed during the whole measurement period was 2.4 m s⁻¹. Based on this data, open fire counts 473 within the grid size of 200×200 km centering over Lumbini was used for this analysis assuming 474 475 that the emissions will take a maximum period of one day to reach our monitoring site. Forest 476 fire counts were obtained from MODIS satellite data product called Global Monthly Fire 477 Location Products- MCD14ML. More on this product has already been described by Putero et al. (2014). Figure 10 shows the daily average $\Delta BC/\Delta CO$ ratio, aerosol absorption Ångstrom 478 exponent (AAE) which is derived from Aethalometer data and daily open fire count within the 479 specified grid. The green box in the figure is used to show two peak events (presented earlier in 480 481 Fig. 6) with the elevated BC and CO concentrations observed during the monitoring period. The first peak was observed during 7-9 April and second peak during 3-4 May, 2013. Two pollutants 482 483 having biomass burning as the potential primary source: BC and CO were taken in consideration. AAE values higher during these two events (~ 1.6) are also an indication of presence of BC of 484 485 biomass burning origin. The chemical composition of TSP filter samples collected at Lumbini also showed higher concentration of Levoglucosan, a biomass burning tracer in Lumbini during 486 487 the pre-monsoon season as compared to other seasons of the year (Wan et al., 2016, Manuscript under review for ACPD). Wan et al. (2016) also reported that the higher correlation between K^+ 488

489 with tracers of dust (Ca^{2+} and Mg^{2+}) indicated that dust is the main source of potassium in 490 Lumbini.

491 Contrary to our expectation, we could not observe any significant influence of forest fire within the specified grid of 200x200 km (or the influence of local forest fire on the air quality over 492 Lumbini was not observed). Therefore, a wider area, covering South and Southeast Asian 493 regions, was selected for the forest fire count. Figure 11 (A-B) shows the active fire hotspots 494 from MODIS, over the region, during the peak events which shows the first peak occurred due to 495 the forest fire over the eastern India region whereas the second peak was influenced by the forest 496 497 fire over western IGP region. Moreover, in order to strengthen our hypothesis, we have utilized 498 satellite data products for various gaseous pollutants like CO and NO₂ (Atmospheric Infrared Sounder (AIRS) for CO and Ozone Monitoring Instrument (OMI) for NO₂ both obtained from 499 Giovanni platform). Figure 11 (C-H) shows the daytime total column CO before, during and 500 after occurrence of two events (peaks) as stated earlier. Atmospheric Infrared Sounder (AIRS) 501 satellite with daily temporal resolution and 1°×1° spatial resolution have been utilized to 502 understand the CO concentration over the area. CO concentration over Lumbini during both of 503 504 the peaks confirmed the role of open fires on either sides of the IGP region for elevated concentration of CO over Lumbini. To further strengthen our finding, the aid of wind rose plot of 505 506 local wind speed and direction was taken. Figure 11 (I-J) represent the wind rose plot only for these two events respectively. Wind rose plots also confirm the wind blowing from those two 507 508 forest fire regions affected the air quality in Lumbini region. Figure 11 (K) shows model biomass 509 CO peak coincident with observed CO. Although the magnitudes are significantly different, the 510 timing of the peaks is well captured by the model. This we believe is due to the fact that satellite 511 based open fire detection also has limitation as it does not capture numerous small fires that are prevalent over south Asia which usually burn out before the next satellite overpass. More 512 research is needed to assess the influence of these small fires on regional air quality. 513

In a separate analysis (not shown here), elevated O_3 concentration during these two events were also observed. Average O_3 concentration before, during and after the events were found to be 46.2 ± 20.3 ppbv, 53.5 ± 31.1 ppbv and 50.3 ± 20.9 ppbv respectively (Event-I) whereas it was found to be 54.8 ± 23.8 ppbv, 56.7 ± 35 ppbv and 55.6 ± 13.4 ppbv respectively (Event-II). Increased ozone concentrations during the high peak events have been analyzed using the satellite NO₂ concentration over the region considering the role of NO₂ as precursor for ozone formation. Daily total column NO₂ were obtained from OMI satellite (data available at the Giovanni platform; <u>http://giovanni.gsfc.nasa.gov/giovanni/</u>) at the spatial resolution of $0.25^{\circ}\times0.25^{\circ}$. Figure 12 shows the NO₂ column value before, during and after both events. Even for the NO₂, maximum concentrations were observed during these two special events.

524 **3.3.2** Identifying regional and local contribution

WRF-STEM model has been used to identify the anthropogenic emission source region 525 influencing the air quality over Lumbini. As previously explained, the model is able to capture 526 527 the observed CO concentration when intense open burning events were not present. A recent study (Kulkarni et al., 2015) has explored the source region contribution of various pollutants 528 529 over the Central Asia using similar technique. Figure 13 (A) shows the average contribution from different regions on CO concentration over Lumbini during the whole measurement period. 530 531 Major share of CO was from the Ganges valley (46%) followed by Nepal region (25%) and rest of Indian region (~17.5%). Contribution from other South Asian countries like Bangladesh and 532 Pakistan were ~ 11% whereas China contributed for ~1% of the CO concentration in Lumbini. 533 Regarding the monthly average contribution, the Ganges Valley and Nepal's contribution were 534 535 almost equal during the month of April (~34% and ~37% respectively) but increased for the Ganges Valley region during the month of May (~44%) and got reduced for Nepal region 536 (~25%) (Figure S6). 537

538 Figure 13 (B) is the time series of percentage contribution to total CO concentration during whole measurement period showing different air mass arriving at a 3 hourly intervals. During the 539 whole measurement period, majority of the CO reaching Lumbini were from the Ganges valley 540 (mainly the states of Punjab, Haryana, Uttar Pradesh, Bihar and West Bengal) region with the 541 542 contribution sometimes reaching up to ~80%. Other India (central, south, east and north) regions also contributed significantly. Bangladesh's contribution in CO loading was seen only after mid-543 April lasting for only about a week and after the first week of May. The contribution from 544 Bangladesh was sporadic comparing to other regions. Highest contribution from this Bangladesh 545 region was observed after the first week of June with the arrival of monsoonal air mass. Pakistan 546 also contributed for the CO loading significantly. Others region as mentioned in the figure 547

covered the regions like Afghanistan, Middle east, West Asia, East Asia, Africa and Bhutan.
Contributions from these regions were less than 5%. Contribution from China was not evident
till the first week of June where a specific air mass arrival shows contribution reaching up to
25% of total CO loading.

552 A sensitivity analysis was performed for emission uncertainty in the model grid containing Lumbini. Lumbini and surrounding regions in the recent years has seen significant rise in urban 553 activities and industrial activity and related emissions which may not be accurately reflected in 554 the HTAPv2 emissions inventory. A month long simulation was carried out with emissions from 555 Lumbini and the surrounding four grids off and another simulation with Lumbini and 556 surrounding four grid's emissions increased by 5 times the amount from HTAPv2 emissions 557 inventory. The results are shown in Figure 13 (C) as percentage increase or decrease compared 558 559 to model results using the current HTAPv2 emissions inventory. The black line shows concentration as 100% for the current HTAPv2 emissions inventory. Despite making Lumbini 560 561 and the surrounding grids emissions zero, model calculation shows pollutant concentration on average is still about 78% of the original value indicating dominance of background and regional 562 563 sources compared to local source in the model. Increasing emissions 5 times for the Lumbini and surrounding four grids only increases the concentration on average by 151%. Thus uncertainty in 564 565 emissions are not a local uncertainty for Lumbini rather for the whole region which needs to be better understood for improving model performance against observations at Lumbini. 566

567 **3.4 Does fossil fuel or biomass influence the Lumbini air**?

The aerosol spectral absorption is used to gain insight into nature and potential source of black 568 569 carbon. This method enables to analyze the contributions of fossil fuel combustion and biomass 570 burning contributions to the observed BC concentration (Kirchstetter et al., 2004). Besides BC, 571 other light absorbing (in the UV region) aerosols are also produced in course of combustion, collectively termed as organic aerosols (often also called brown carbon or BrC) (Andreae and 572 Gelencsér, 2006). Figure 14 shows the comparison of normalized light absorption as function of 573 the wavelength for BC observed at Lumbini during cooking and non-cooking hours. Our results 574 are compared with the published data of Kirchstetter et al. (2004) and that observed over a 575 village center site of Project Surya in the IGP (Praveen et al., 2012) (figure not shown). We 576

discuss light absorption data from two distinct times of the day. The main reason behind using 577 data during 07:00-08:00 h and 16:00-17:00 h is these periods represent highest and lowest 578 579 ambient concentration (Fig. 8). Also these period represent cooking and non-cooking or high and low vehicular movement hours (Praveen et al., 2012). To understand the influence of biomass 580 and fossil fuel we plotted normalized aerosol absorption at 700 nm wavelength for complete 581 aethalometer measured wavelengths in Fig. 14. Kirchstetter et al. (2004) reported OC absorption 582 583 efficiency at 700 nm to be zero. Thus we normalized measured absorption spectrum by 700 nm wavelength absorption. Since aethalometer does not provide 700 nm wavelength absorption 584 values, we used methodology followed by Praveen et al. (2012). Our results show that the 585 normalized absorption for biomass burning aerosol is ~3 times higher at 370 nm compared to 586 that at 700 nm whereas fossil fuel absorption is about 2.6 times higher at the same wavelength. 587 588 The normalized curve obtained during both cooking and non-cooking period lies in between the standard curve of Kirchstetter et al. (2004). As shown in Fig. 14, the curve obtained for the prime 589 cooking time is closer towards the published curve on biomass burning whereas that obtained 590 during the non-cooking time is closer towards the published fossil fuel curve. Similar result was 591 592 also observed over the Project Surya village in the IGP region (Praveen et al., 2012;Rehman et al., 2011). This clearly indicates there is contribution of both sources: biomass as well as fossil 593 594 fuel on the observed BC concentration over Lumbini.

In order to identify fractional contribution of biomass burning and fossil fuel combustion to 595 596 observed BC aerosol, we adopted the method described by Sandradewi et al. (2008). Wavelength dependence of aerosol absorption coefficient (b_{abs}) is proportional to $\lambda^{-\alpha}$ where λ is the 597 598 wavelength and α is the absorption Ångstrom exponent. The α values ranges from 0.9-2.2 for fresh wood smoke aerosol (Day et al., 2006) and between 0.8-1.1 for traffic or diesel soot 599 600 (references in Sandradewi et al. (2008)). We have taken α value of 1.86 for biomass burning and 1.1 for fossil fuel burning as suggested by previous literature (Sandradewi et al., 2008). Figure 601 15 shows diurnal variation of the biomass burning BC. Minimum contribution of biomass 602 burning to total BC concentration was observed during 04:00-06:00 local time (only about 30% 603 of the total BC). As the cooking activities start in morning, the contribution of biomass BC starts 604 to increase and reaches about 50%. Similar pattern was repeated during evening cooking hours. 605 Only during these two cooking periods, fossil fuel fraction BC was lower. Otherwise it remained 606

607 significantly higher than biomass burning BC throughout the day. On average, ~40% of BC was from biomass burning whereas remaining 60% was contributed by fossil fuel combustion during 608 609 our measurement period. Interestingly, this is the opposite of the contributions that were concluded by Lawrence and Lelieveld (2010). Lawrence and Lelieveld (2010) concluded that 610 ~60% BC from biomass versus ~40% fossil fuel, based on a review of numerous previous 611 studies to be likely for the outflow from Southern Asia during the winter monsoon. When we 612 compared observed Ångstrom exponent with Praveen et al. (2012), we noticed that Lumbini 613 values were lower than Project Surya Village center site. This implies Surya village center had 614 higher biomass fraction, also it was observed absorption Ångstrom exponent exceeded 1.86 615 during cooking hours which indicates 100% biomass contribution. The difference is attributed to 616 the fact that Lumbini sampling site is not a residential site like Surya village which can capture 617 cooking influence efficiently. Further Lumbini sampling site is surrounded by commercial 618 activities such as a local bus park, hotels, office buildings and industries and brick kilns slightly 619 further away. Although the reason for this difference is not clear, it is an indication of the 620 important role of diesel and coal emissions in the Lumbini and upwind regions. 621

622 4. Conclusions

623 Our measurements, a first for the Lumbini area, have shown very high pollution concentration at Lumbini. Black carbon (BC), carbon monoxide (CO), ozone (O₃) and particulate matter (PM₁₀, 624 PM_{2.5} and PM1) were measured during the pre-monsoon of 2013 as a regional site of the SusKat-625 ABC campaign. Average pollutant concentrations during the monitoring period were found to be: 626 BC: 4.9±3.8 μg m⁻³; CO: 344.1±160.3 ppbv; O₃: 46.6±20.3 ppbv; PM₁₀: 128.8±91.9 μg m⁻³ 627 $PM_{2.5}$: 53.14±35.1 µg m⁻³ and PM_1 : 36.6±25.7 µg m⁻³ which is comparable with other urban sites 628 like Kanpur and Delhi in the IGP region. However, our study finds higher fraction of coarse 629 mode PM in Lumbini as compared to other sites in the IGP region. In addition, $\Delta BC/\Delta CO$ ratio 630 631 obtained in Lumbini was within the range of emission from both domestic and transportation 632 sectors, indicating them as potential key sources of BC and CO, and likely most of PM_1 in Lumbini. The diurnal variation of the pollutants is similar to that of any urban location, with 633 634 peaks during morning and evening. However, our results show higher evening concentration compared to morning concentration values and needs further research to explain this behavior. 635 636 During our measurement period, air quality in Lumbini was influenced by regional forest fires as

637 shown by chemical transport model and satellite data analysis. A regional chemical transport model, WRF-STEM was used to interpret observations. Inter-comparison of WRF-STEM model 638 639 outputs with observations showed that the model underestimated the observed pollutant concentrations by a factor of 1.5 to 5 but was able to capture the temporal variability. Model 640 uncertainties are attributed mostly to uncertainties in meteorology and regional emissions as 641 shown from sensitivity analysis with local emissions. Region-tagged CO as air-mass tracers are 642 employed in WRF-STEM model to understand the anthropogenic emission source region 643 influencing Lumbini. Our analysis shows that the adjacent regions; mostly the Ganges valley, 644 other parts of India and Nepal accounted for the highest contribution to pollutant concentration in 645 the Lumbini. The normalized light absorption curve clearly indicated the contribution to BC in 646 Lumbini from both sources: biomass as well as fossil fuel. On average, ~40% BC was found to 647 be from the biomass burning and ~60% from fossil fuel burning. 648

Various improvements and extensions would be possible in future studies. More reliable 649 650 functioning of the AWS (temperature and RH sensor, rain gauge) would have allowed more indepth analysis of the relationship between meteorological parameters and pollutants 651 652 concentration. Continuous measurements of air pollutants throughout the year would allow for annual and seasonal variation study. Improvements in the model are much needed in its ability to 653 654 simulate observed meteorology. Significant uncertainty lies with regional emissions inventory developed at national and continental scale versus local bottoms up inventory and pollutant 655 656 emissions from small scale open burning not captured by satellites.

There is a clear need for setting up of a continuous air quality monitoring station at Lumbini (UNESCO World Heritage Site) and the surrounding regions for long-term air quality monitoring.

660 Data availability

661 The data used for this manuscript can be obtained by sending an email to the corresponding authors and/or to IASS (Maheswar.Rupakheti@iass.potsdam.de) 662 and/or to ICIMOD (arnico.panday@icimod.org). Modeling data can be obtained Β. Adhikary 663 from (Bhupesh.adhikary@icimod.org). 664

665 Authors' contributions

M.R. and M.L. conceived the Lumbini portion of the SusKat experiment. M.R. and A.K.P.
coordinated the Lumbini field campaign. D.R. and K.S.M conducted the field observations at
Lumbini. B.A. designed and ran the WRF-STEM model. P.S.P., B.A. and D.R. finalized the
manuscript composition. D.R., P.S.P, B.A., M.R. and S.K. conducted the data analysis. D.R. and
B.A. prepared the manuscript with inputs from all coauthors.

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Instrument (Model)	Manufacturer	Parameters	Inlet/sensor height (above ground)	Sampling interval	Sampled period	
Environmental Dust monitor (EDM 164)	GRIMM Aerosol Technik, Germany	PM ₁ , PM _{2.5} , PM ₁₀	5 m	5 min	01/04-15/06	
Aethalometer (AE42)	Magee Scientific, USA	Aerosol light absorption at seven wavelengths, and BC concentration	3 m	5 min	01/04-05/06	
CO analyzer (48 <i>i</i>)	Thermo Scientific, USA	CO concentration	3 m	1 min	01/04-15/06	
O_3 analyzer (49 <i>i</i>)	Thermo Scientific, USA	O ₃ concentration	3 m	1 min	01/04-15/06	
Automatic Weather Station (AWS)	Campbell Scientific, UK	T, RH, WS, WD, Global Radiation, Precipitation	12 m	1 min	01/04-15/06	

Table 1. Summary of instruments deployed during monitoring in Lumbini

Sites	Characteristics	Measurement period	PM _{2.5} (μg m ⁻³)	$\frac{BC}{(\mu g/m^3)}$	CO (ppbv)	O ₃ (ppbv)	References
Lumbini, Nepal	Semi-urban	Pre-monsoon, 2013	53.1±35.1	4.9±3.8	344.1±160.3	46.6±20.3	This study
Kathmandu, Nepal	Urban	Pre-monsoon, 2013	-	14.5±10	-	38.0±25.6	(Putero et al., 2015)
Mt. Everest, Nepal	Remote	Pre-monsoon	-	0.4±0.4	-	61.3±7.7	(Marinoni et al., 2013
Delhi, India	Urban	Pre-monsoon (night-time)	82.3±50.5	7.70±7.25	1800±890	-	(Bisht et al., 2015)
Kanpur, India	Urban	June 2009-May 2013, April-June	-	2.1±0.9	721±403	27.9±17.8	(Gaur et al., 2014) (Ram et al., 2010)
Mohali, India	Semi-urban	May, 2012	104±80.3	-	566.7±239.2	57.8±25.4	(Sinha et al., 2014)
Mt. Abu, India	Remote	Jan 1993-Dec 2000, pre-monsoon	-	0.7±0.14	131±36	39.9±10.8	(Naja et al., 2003) (Da and Jayaraman, 2011

Table 2. Comparison of PM_{2.5}, BC, CO and O₃ concentrations at Lumbini with those at other sites in South Asia

Table 3. Inter-comparison of observed and model simulated hourly average concentrations of air 928 pollutants during the measurement campaign period. Unit: BC and PM in μ g/m³ and CO in ppbv.

Pollutants	Observed (mean and range)	Modeled (mean and range)	Ratio of mean (observed/modeled)	
BC	4.9 (0.3-29.9)	1.8 (0.4-3.7)	2.7	
PM ₁	36.6 (3.6-197.6)	12.3 (0.9-41.7)	3	
PM _{2.5}	53.1 (6.1-272.2)	17.3 (1.9-48.3)	3	
PM ₁₀	128.8 (10.5-604.0)	25.4 (2.1-68.8)	5	
СО	344.1(124.9-1429.7)	255.7 (72.2-613.1)	1.4	

931 Figures

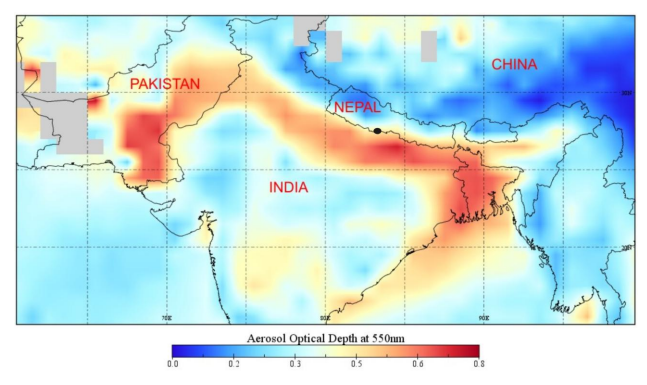


Figure 1. Aerosol optical depth in South Asia acquired with the MODIS instrument aboard
TERRA satellite averaged over the winter and pre-monsoon season (December 2012-June 2013).
High aerosol loading can be seen over the entire Ingo-Gangetic Plains (IGP). An aerosol hotspot
south of Lumbini (small black mark nearby the border of Nepal with India) is clearly visible.
Light grey color used in the figure represents the absence of data.

938

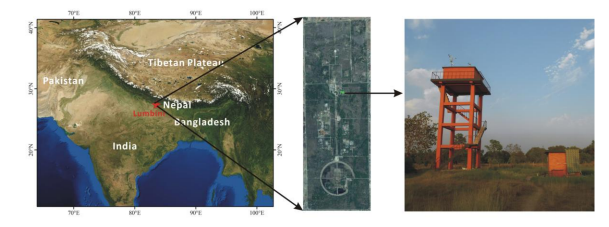


Figure 2. Location of sampling site in Lumbini in southern Nepal (left panel). The middle panel

- shows the Kenzo Tange Master Plan Area of Lumbini while the right panel shows the sampling
- 942 tower in the Lumbini Master Plan Area.

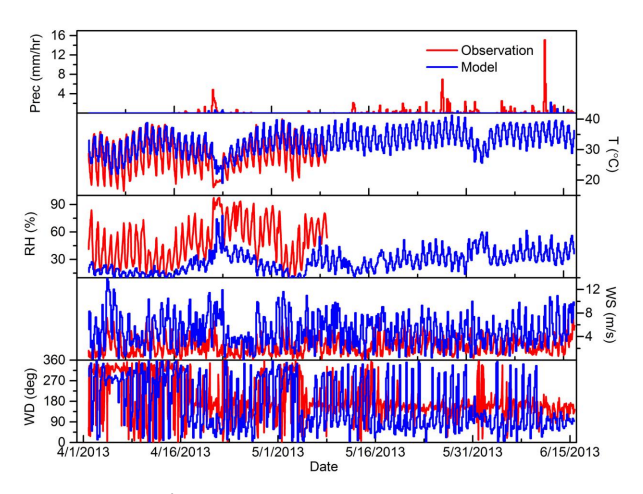


Figure 3. Time series of hourly average observed (red line) and model estimated (blue line)
meteorological parameters at Lumbini, Nepal for the entire sampling period from 1 April to 15
June 2013.

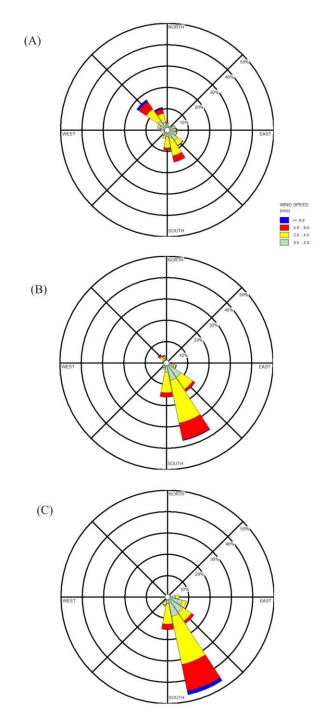


Figure 4. Wind rose of wind speed and wind direction observed at Lumbini during the month of
(A) April, (B) May, and (C) (1st-15th) June 2013.

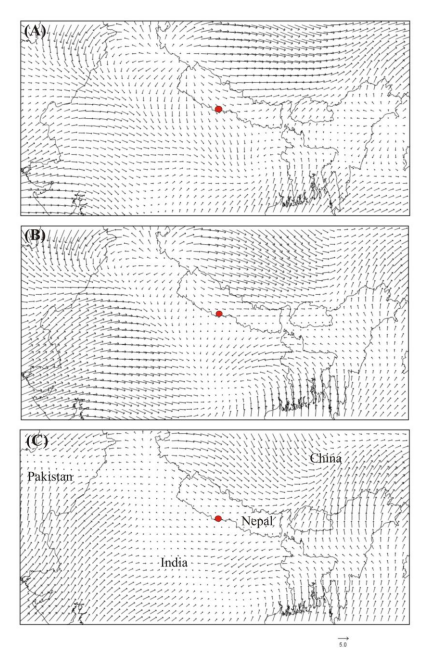


Figure 5. Monthly synoptic surface winds for the month of (A) April, (B) May and (C) June
2013, based on NCEP/NCAR reanalysis data. Orientations of arrows in the figures refer to wind
direction whereas the length of arrows represents the magnitude of wind speed (m/s). Red dot in
the map represents the location of Lumbini.

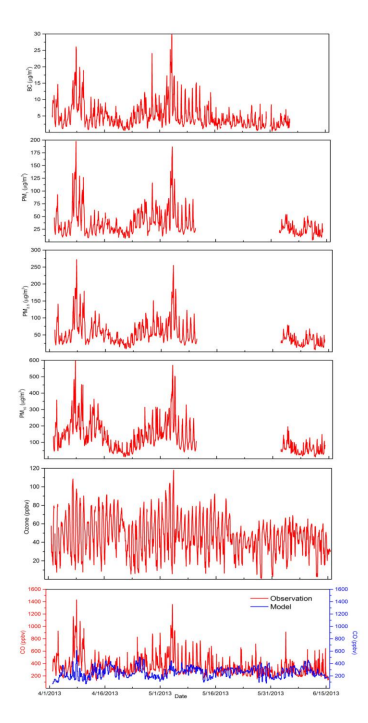
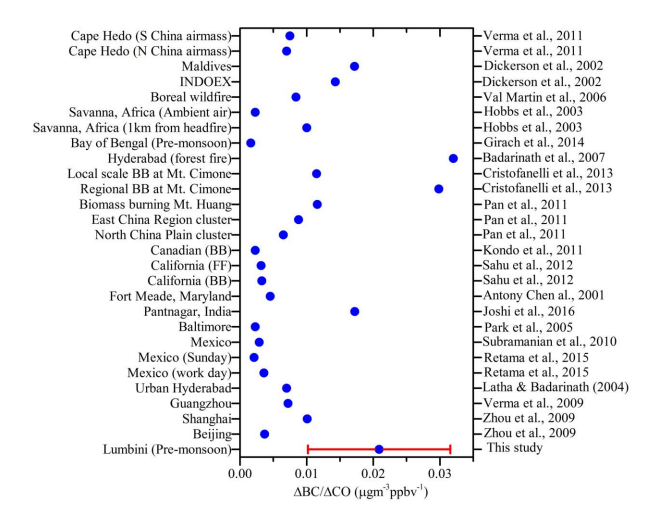


Figure 6. Time series of the observed (red line) and model estimated (blue line) hourly average
concentrations of BC, PM₁, PM_{2.5}, PM₁₀, O₃ and CO at Lumbini, Nepal for the entire sampling
period from 1 April to 15 June 2013.



967 Figure 7: Comparison of BC concentrations to CO concentrations (Δ BC/ Δ CO) ratios obtained 968 for Lumbini with other sites. The red horizontal bar represents standard deviation.

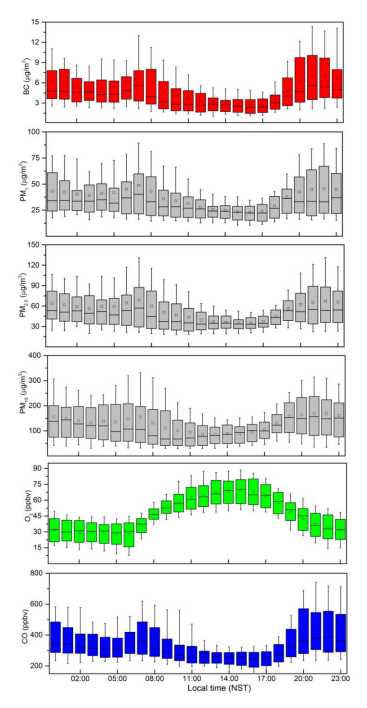


Figure 8. Diurnal variations of hourly average ambient concentrations of BC, PM_1 , $PM_{2.5}$, PM_{10} , O₃ and CO at Lumbini during the monitoring period (1 April -15 June 2013). In each box, lower and upper boundary of the box represents 25th and 75th percentile respectively, top and bottom of the whisker represents 90th and 10th percentile respectively, the mid-line represents median, and the square mark represents the mean for each hour.

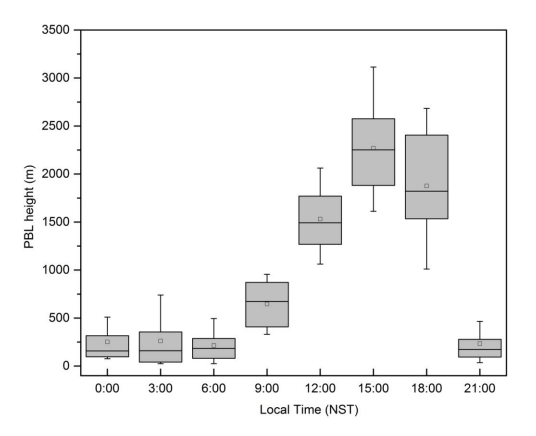
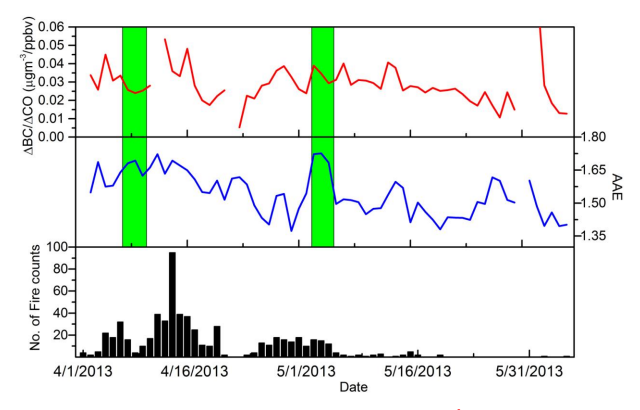


Figure 9. Diurnal variation of the planetary boundary layer (PBL) height at Lumbini obtained
for every three hours of each day from the WRF-STEM model for the sampling period. The
square mark in each box represents the mean PBL height, bottom and top of the box represents
25th and 75th percentile, top and bottom of the whisker represents 90th and 10th percentile
respectively.



982

Figure 10. Time series of daily average $\Delta BC/\Delta CO$ ratio, absorption Ångstrom exponent (AAE), along with fire counts acquired with the MODIS instrument onboard TERRA satellite for a 200×200 km grid centered at Lumbini. Two rectangular green boxes represent time of two episodes with high peaks in CO and BC concentrations as shown in earlier figures.

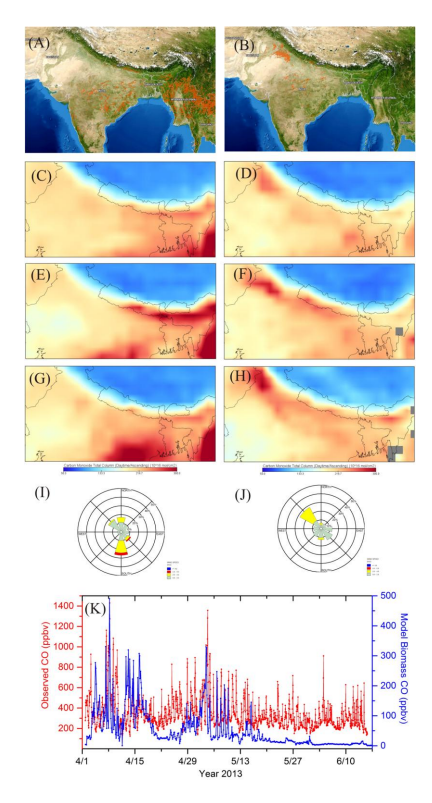


Figure 11. Active fire hotspots in the region acquired with the MODIS instrument on TERRA
satellite during (A) Event-I (7-9 April) and (B) Event-II (3-4 May). CO emissions, acquired with

- AIRS satellite, in the region 2 days before (3-5 April), during (7-9 April) and 2 days after (10-12
- April) the Event-I are shown in panels (C), (E) and (G), respectively while panels (D), (F) and
- (H) show CO emissions 2 days before (1-2 May), during (3-4 May) and 2 days after (5-6 May)
- the Event-II. Panels (I) and (J) represent the average wind rose plot of observed wind direction
- and wind speed during Event I and II, respectively. (K) Observed CO versus Model open
- burning CO illustrating contribution of forest fires during peak CO loading.

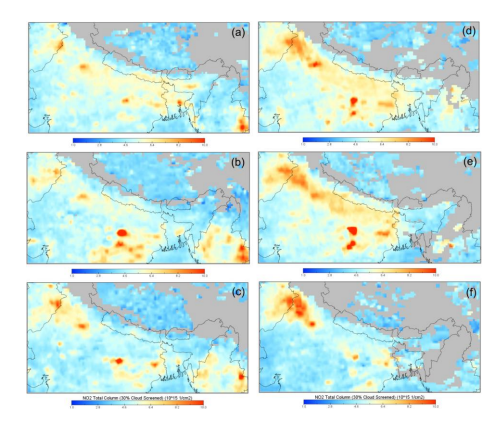
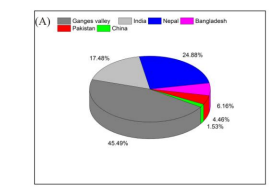


Figure 12. NO₂ total column obtained with OMI satellite over the region (a) before, (b) during,
and (c) after the Event- I. The panels (d), (e), (f) show NO₂ total column before, during and after
the Event- II.





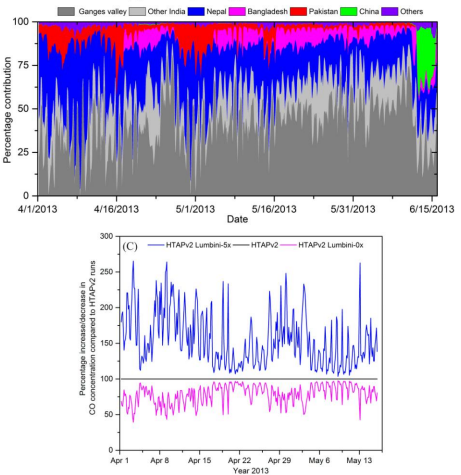


Figure 13. (A) WRF-STEM model estimated contributions of various source regions to average
 CO concentration in Lumbini for the sampling period, (B) time series of region tagged CO tracer
 during the whole measurement period using HTAP emission inventory and (C) Figure showing
 percentage increase/decrease in CO concentration with different emissions scenario.

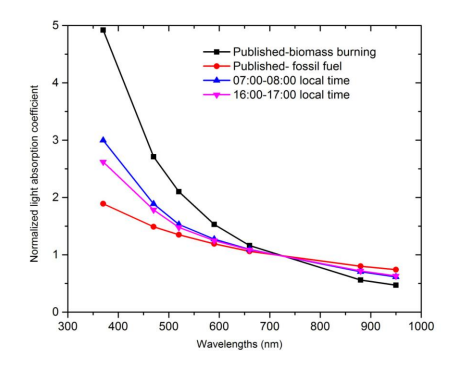


Figure 14. Comparison of normalized spectral light absorption coefficients obtained during the
prime cooking (07:00-08:00 local time) and non cooking time (16:00-17:00 LT) at Lumbini with
published data from Kirchstetter et al. (2004).

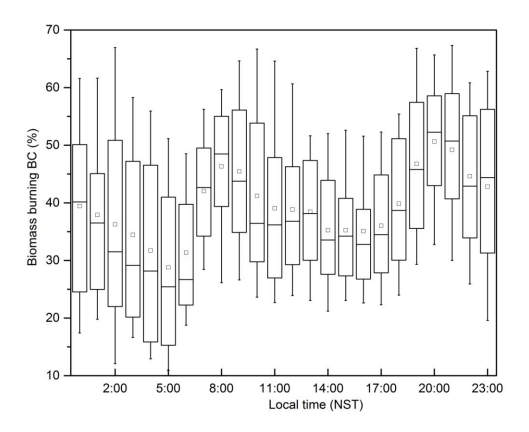


Figure 15. Diurnal variation of the fractional contribution of biomass burning to ambient BC concentration at Lumbini for the measurement period. In each box, lower and upper boundary of the box represent 25th and 75th percentile, respectively, top and bottom of the whisker represents 90th and 10th percentile, respectively. The mid-line in each box represents median while the square mark represents the mean for each hour.