# Seasonal responses of $\delta 13C$ and $\delta 18O$ of atmospheric CO2 over sub-urban region of India.

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#### Abstract

The seasonal diurnal variability of atmospheric CO2 and driving factors are studied using its continuous isotopic fractionation ( $\delta$ 13C and  $\delta$ 18O) over Shadnagar, a sub-urban location of India with high precision in-situ data from November 2018 to October 2019. The annual averaged atmospheric CO2 concentrations and 13C-CO2 and 18O-CO2 are 415.03±9.77 ppm, -11.18±1.73 CO2 was observed in summer monsoon (17.30 +- 9.29 ppm) and the minimum was noticed in winter (7.19 +- 0.11 ppm) indicating strong seasonality at the study site. To characterize the atmospheric CO2 sources/sinks, an improved model of Miller and Tans was implemented by plotting  $\Delta$ CO2 against  $\Delta$ (CO2 ×  $\delta$ 13C) respectively during day and night. An averaged seasonal  $\delta$ 13C source/sink signature ( $\delta$ s) is -32.84 -26.09related to combustion and dominance of C3 ecosystem respiration respectively. The seasonal relationship between  $\delta$ 18O and  $\delta$ 13C is strongly correlated during pre-monsoon ('r' = 0.93 to 0.95) than post monsoon ('r' = 0.07 to 0.13), which might be due to high vapour pressure deficit. A Lagrangian back-trajectory model confirms the influence of the Indian Summer Monsoon on the variability of atmospheric CO2 concentration during the summer monsoon season.

# Seasonal responses of δ<sup>13</sup>C and δ<sup>18</sup>O of atmospheric CO<sub>2</sub> over sub-urban region of India

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## 24 Key Points:

- Study focused on diurnal variability of seasonal atmospheric CO<sub>2</sub> and its stable isotopes  $(\delta^{13}C \text{ and } \delta^{18}O)$ .
- Determination of  $\delta^{13}$ C signature of atmospheric CO<sub>2</sub> using improved Miller and Tans model.
- Effect of the Indian Summer monsoon circulation on atmospheric CO<sub>2</sub> variation was studied.

#### 31 Abstract

The seasonal diurnal variability of atmospheric CO<sub>2</sub> and driving factors are studied using its 32 continuous isotopic fractionation ( $\delta^{13}$ C and  $\delta^{18}$ O) over Shadnagar, a sub-urban location of India 33 with high precision in-situ data from November 2018 to October 2019. The annual averaged 34 atmospheric CO<sub>2</sub> concentrations and  ${}^{13}$ C-CO<sub>2</sub> and  ${}^{18}$ O-CO<sub>2</sub> are 415.03±9.77 ppm, -11.18±1.73 ‰ 35 and 9.1±13.35 ‰. Seasonal amplitudes of atmospheric CO<sub>2</sub> was observed in summer monsoon 36  $(17.30 \pm 9.29 \text{ ppm})$  and the minimum was noticed in winter  $(7.19 \pm 0.11 \text{ ppm})$  indicating strong 37 38 seasonality at the study site. To characterize the atmospheric  $CO_2$  sources/sinks, an improved model of Miller and Tans was implemented by plotting  $\Delta CO_2$  against  $\Delta (CO_2 \times \delta^{13}C)$  respectively 39 during day and night. An averaged seasonal  $\delta^{13}$ C source/sink signature ( $\delta_s$ ) is -32.84 ‰ in the 40

- 41 day time and -26.09% in night time representing the source of atmospheric CO<sub>2</sub> is related to
- 42 combustion and dominance of  $C_3$  ecosystem respiration respectively. The seasonal relationship
- 43 between  $\delta^{18}$ O and  $\delta^{13}$ C is strongly correlated during pre-monsoon ('r' = 0.93 to 0.95) than post-
- 44 monsoon ('r' = 0.07 to 0.13), which might be due to high vapour pressure deficit. A Lagrangian 45 back-trajectory model confirms the influence of the Indian Summer Monsoon on the variability
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- 46 of atmospheric CO<sub>2</sub> concentration during the summer monsoon season.
- 47 Keywords: atmospheric CO<sub>2</sub>, isotopic fractionation, Miller and Tans, Indian summer monsoon.

## 48 **1. Introduction**

49 The Intergovernmental Panel for Climate Change (IPCC) reported that carbon dioxide (CO<sub>2</sub>) sources from anthropogenic gases in the atmosphere cause more radiative forcing next to water 50 vapor (Smith et al., 1999). CO<sub>2</sub> concentrations are consistently increasing and touched 400 ppm 51 at Mauna Loa, a global reference site during May 2013 (Monastersky, 2013). Globally, CO<sub>2</sub> 52 concentrations are increasing, which could be due to land use land cover changes (LULCC) and 53 54 progress in industrial activities (Ballantyne et al., 2012) especially fossil fuel combustion, cement manufacturing. Due to fossil fuels burnings and LULC emissions, an increase of 40 and 55 150 % in CO<sub>2</sub> and CH<sub>4</sub> concentration respectively is observed since the pre-industrial period 56 (Huang et al., 2016). Emissions of CO<sub>2</sub> by different processes are controlled by varied 57 environmental conditions, in which about half of the CO<sub>2</sub> levels are released into the atmosphere 58 as source and remaining are absorbed by the processes of the terrestrial biosphere and ocean 59 uptake (Andres et al., 1996) as sink. Hence monitoring and maintaining long-term records of 60 atmospheric CO<sub>2</sub> measurements are very important to understand the carbon cycle and to assess 61 the CO<sub>2</sub> mixing ratios in the atmosphere by controlling factors namely photosynthesis, 62 63 respiration, biomass, fossil fuel burning and air-sea exchange processes (Machida et al., 2003).

Globally, systematic high precision atmospheric CO<sub>2</sub> observations are accelerated to understand 64 the global carbon cycle. Over the Indian subcontinent, spatio-temporal variability in atmospheric 65  $CO_2$  concentrations are characterized by the terrestrial biosphere and seasonal weather patterns 66 which brings long-range air-masses (Valsala et al., 2013; Tiwari et al., 2014). To understand the 67 seasonal, inter and intra annual variations of atmospheric CO<sub>2</sub> over the Indian subcontinent, high 68 precision CO<sub>2</sub> measurements are being generated across the country from different research 69 institutes (Bhattacharya et al., 2009; Sharma et al., 2014; Mahesh et al., 2016; Nalini et al., 70 2019). A literature survey on atmospheric CO<sub>2</sub> variability over the Indian region is mainly 71 focused on local sources and transport (Sreenivas et al., 2016). However, a need was felt to 72 73 understand the reasons and causes for uncertainty in surface fluxes. Hence an advanced studies of stable isotopic measurements of carbon and oxygen are gaining momentum to understand CO<sub>2</sub> 74 levels, source and sinks of CO<sub>2</sub>, both on regional and global levels. 75

Stable carbon and oxygen isotopes of atmospheric CO<sub>2</sub> can be used as tracers in the carbon cycle, 76 which are affected by the anthropogenic and biogenic  $CO_2$  components. The <sup>13</sup>C-CO<sub>2</sub> and <sup>18</sup>O-77  $CO_2$  are stable isotopes of  $CO_2$  molecules are widely used for source apportionments in the 78 atmosphere, hydrosphere and geosphere as well as interaction between them (Guillon et al., 79 2015). The  ${}^{13}$ C-CO<sub>2</sub> concentration in the atmosphere has been decreasing since pre-industrial 80 times, which indicates the more addition of CO<sub>2</sub> to the atmosphere by fossil fuel burning (Yakri, 81 2011). Many research activities have been carried out on atmospheric  $CO_2$  and its stable isotopes 82  $(\delta^{13}C, \delta^{18}O)$  by various groups (Bhattacharya et al., 2009; Pataki et al., 2003; Clark-Thorne and 83

yapp, 2003; Francey and Tans, 1987; Murayama et al., 2010; Newman et al, 2003; Wada et al., 84 2011; Pataki et al., 2006; Zhou et al., 2005; Zhou et al., 2006; Sturm et al., 2006; Djuricin et al., 85 2010; Guha and Ghosh, 2010; Gorka and Lewicka- Szczebak, 2013; Liu et al., 2014a; Pang et 86 al., 2016). Liu et al. (2014a) studied the atmospheric CO<sub>2</sub>,  $\delta^{13}$ C composition and their 87 relationship to understand the sources and sinks at two stations viz., Waliguan and Shangdianzi 88 in China using observational data for the period from 2007 to 2010. The result of this study 89 indicates that CO<sub>2</sub> and  $\delta^{13}$ C composition possesses long-term trends and seasonal cycles that 90 correlate with each other. An improved model by Miller and Tans (2003) is widely used to 91 determine the source or sink that causes CO<sub>2</sub> variability. Pang et al. (2016) used Keeling plot 92 intercept method for isotopic composition of CO<sub>2</sub> and found increased value in vegetative 93 season and depleted value in heating season. On Indian sub-continent a few studies are carried 94 out on atmospheric CO<sub>2</sub> and its stable isotopes which are limited to discrete sample analysis 95 (Bhattacharya et al., 2009; Guha and Ghosh, 2015). Measurements of the <sup>18</sup>O-CO<sub>2</sub> also play an 96 important role in the carbon cycle to distinguish the photosynthesis and respiration process of 97 CO<sub>2</sub> fluxes (Farquhar et al., 1993; Kato et al., 2004). 98

The objective of the present study is to understand seasonal diurnal variability of 99 atmospheric CO<sub>2</sub> and its stable isotopic composition at the sub-urban region of Telangana. Since 100 the study site is surrounded by multiple sources for  $CO_2$  (e.g., biospheric respiration and fossil 101 fuel emissions), thus an improved model by Miller and Tans (2003) was used to characterize the 102 103 CO<sub>2</sub> sources/sinks. Here, we report continuous high precision CO<sub>2</sub> isotopic measurements, first of their kind from sub-urban region of Telangana. This work has been carried out as part of the 104 Atmospheric CO<sub>2</sub> Retrieval and Monitoring (ACRM) of National Carbon Project (NCP) funded 105 by Climate and Atmospheric Processes of ISRO-Geosphere Biosphere Programme (CAP-IGBP). 106

#### 107 2. Materials and Methods

Observations of CO<sub>2</sub> and its isotopic composition are measured during November 2018 to 108 109 October 2019 by laser based Isotope Ratio Infrared Spectrometer (IRIS) analyser at 8 m height from the surface of Atmospheric Science Laboratory (ASL), NRSC, Shadnagar (Latitude: 17.09 110 °N; Longitude: 78.21 °E and Elevation: 648 m above mean sea level) a sub-urban region of 111 Telangana. Surface meteorological data at the study site are collected from an automatic weather 112 station. An hourly Boundary Layer Height (BLH) were obtained from European Centre for 113 Medium-Weather Range Forecasts (ECMWF-ERA, 114 https://cds.climate.copernicus.eu/cdsapp#!/dataset/reanalysis-era5-single-levels?tab=form). In addition to the 115 above datasets, fire counts with confidence > 70% are obtained from Moderate Resolution 116 Spectroradiometer https://firms.modaps.eosdis.nasa.gov/download/). Imaging (MODIS, 117 Normalized Difference vegetation Index (NDVI) is obtained from an open data archival of 118 119 Bhuvan site (https://bhuvan-app3.nrsc.gov.in/data/download/index.php), which is derived from Oceansat-2 Ocean color monitor sensor. Figure 1a shows that the study area is overlaid with 120 NDVI over Telangana state as well as time series of air temperature, relative humidity and wind 121 122 speed recorded at the study site.

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Figure 1. (a) Study location with prevailing weather conditions and (b) NDVI

Figure 1a also shows *in-situ* atmospheric CO<sub>2</sub> networks across the country established by 150 different research institutes of India. A few of the station's information is obtained from Nalini et 151 al. (2019). Black solid circles are the atmospheric CO<sub>2</sub> measurement stations installed under 152 ACRM of NCP project by NRSC, ISRO. At the present study site (solid black star) long-term 153 CO<sub>2</sub>, CH<sub>4</sub> and H<sub>2</sub>O measurements are complemented by CO<sub>2</sub> and N<sub>2</sub>O isotopic observations. 154 During the study period, air temperature is observed to be high in pre-monsoon (March-May) 155 with maximum air temperature of 43°C and low in winter (January-February) with mininum 156 157 temperature of 9°C. Relative humidity is observed to be high in monsoon (June-September) with maximum value reaching 100 % and low in pre-monsoon with minimum of 12.18 %. However, 158 wind speed is observed to be ranging from 0.005 m s<sup>-1</sup> to 6.0 m s<sup>-1</sup> during the study period. 159 Figure 1b shows minimum, maximum and mean NDVI at 15-day interval at the study site. Thus, 160 seasonal NDVI are computed during the study period for post-monsoon (0.39), winter (0.33), 161 pre-monsoon (0.30) and monsoon (0.41). The study site is about 60 km away from the urban 162 region of Hyderabad (fifth largest city in India) and is associated with 75% air pollution from 163 traffic sector alone (significant anthropogenic impacts) due to increase in population and related 164 factors (Malakshmi et al., 2014). Thus the present study site doesn't fall under single source 165 contributor of atmospheric CO<sub>2</sub>. 166

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#### 169 **2.1. Continuous stable isotopic measurements**

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In this study, we used commercial laser based IRIS CO<sub>2</sub> Carbon Isotope analyzer Enhanced 171 Performance (CO<sub>2</sub>-CCIA-EP), procured from Los Gatos Research, U.S.A in November 2018. 172 Details of the instrument functioning are given in Baer et al. (2002). The CO<sub>2</sub>-CCIA-EP is 173 capable of simultaneous measurements of dry CO<sub>2</sub>, <sup>13</sup>C, <sup>18</sup>O and H<sub>2</sub>O are measured using the 174 absorption line at 2.05 µm with 1 Hz frequency and uses a performance enhancing off-axis 175 cavity ring down spectroscopy. Mole fraction of isotopic composition also depends on internal 176 cavity pressure and temperature hence maintained constant at 119.14 Torr and 45.36 °C 177 respectively (Mahesh et al., 2015). To keep the moisture as constant and as low as possible in the 178 analyzer, sample and reference gases are passed through a Neflon drying unit. The dry mole 179 fractions of CO<sub>2</sub> where the measured mole fraction of H<sub>2</sub>O, which is also in ppm has been 180 removed as shown in Equation 1. 181

$$CO_{2}(ppm)_{dry} = \left(\frac{CO_{2} (ppm)_{wet}}{(1 - \frac{H_{2}O (ppm)}{10^{6}})}\right)$$
(1)

The <sup>13</sup>C-CO<sub>2</sub> and <sup>18</sup>O-CO<sub>2</sub> compositions are reported as  $\delta^{13}$ C and  $\delta^{18}$ O respectively versus VPDB (Vienna Pee Dee Belemnite) and reported in per mil (‰) as shown in Equation (2) and (3)

$$\delta^{13}C((\%_0) = \left(\frac{C_s}{C_r} - 1\right) \times 1000$$
(2)

$$\delta^{18}O((\%_0) = \left(\frac{O_s}{O_r} - 1\right) \times 1000 \tag{3}$$

185 Where  $C_s, C_r, O_s$ , and  $O_r$  defined as follows;  $C_{s (sample)} = ({}^{13}C/{}^{12}C)_s$ ;  $C_{r (VPDB)} = ({}^{13}C/{}^{12}C)_r$ ;  $O_s$ 186  $(sample) = ({}^{18}O/{}^{16}O)_s$  and  $O_r (VPDB) = ({}^{18}O/{}^{16}O)_r$ 

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#### 188 **2.2. Calibration of CO<sub>2</sub> isotope analyzer**

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The CO<sub>2</sub>-CCIA-EP is calibrated using National Oceanic and Atmospheric Administration 190 (NOAA) supplied CO<sub>2</sub> isotope calibration reference gases (ID: CC718409) towards eliminating 191 instrument drifts and generating high quality data. The precision and accuracy of the instrument 192 are computed by performing the internal calibration on 8<sup>th</sup> January 2020 with a 404.53±0.21 ppm 193 of CO<sub>2</sub>, -8.45±0.01 ‰, VPDB of  $\delta^{13}$ C and -1.28±0.03 ‰, VPDB of  $\delta^{18}$ O respectively. An inflow 194 of reference gas has been passed through the Neflon drying unit for 15 minutes and collected 195 data at 1 Hz frequency. Calibration is performed twice during the study period. Figure 2 shows 196 the time series of CO<sub>2</sub>,  $\delta^{13}$ C-CO<sub>2</sub> and  $\delta^{18}$ O-CO<sub>2</sub> during the calibration process. The shaded area 197 in figure 2 represents  $\pm 1 \sigma$  standard deviation. 198

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Figure 2. Time series of isotopic composition during the calibration process.

The 100 sec (1  $\sigma$ ) average precision of CO<sub>2</sub>,  $\delta^{13}$ C-CO<sub>2</sub> and  $\delta^{18}$ O-CO<sub>2</sub> are 0.20 ppm, 0.05 ‰ and 0.06 ‰ respectively. The result of the calibration report is summarized in table 1.

Cylinder ID	$CO_2$ (ppm, ref) $\pm 1\sigma$	$\delta^{13}$ C of CO $_2$ (‰, ref) $\pm 1\sigma$	$\delta^{18}$ O of CO $_2$ (‰, ref) $\pm 1\sigma$
	$CO_2 (ppm, M) \pm 1\sigma$	$\delta^{13}$ C of CO <sub>2</sub> (‰, M) ± 1 $\sigma$	$\delta^{18}$ 0 of CO <sub>2</sub> (‰, M) ± 1 $\sigma$
NOAA,CC718409	404.53 ± 0.21 (ppm)	-8.45 ± 0.01 (‰, VPDB)	−1.28 ± 0.03 (‰, VPDB)
	404.54 ± 0.20 (ppm)*	-8.98 ± 0.05 (‰, VPDB)*	−1.51 ± 0.06 (‰, VPDB)*
Bias (Ref-M)	-0.01 (ppm)	0.53 (‰)	0.23 (‰)
Precision	0.05%	0.55%	3.97%
Accuracy	$0.00\overline{247} = 0.01\%$	6.27%	17.96%
Ref: Reference; M: Measured		*Indicates $1\sigma$ calculation for 100sec average	

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**Table 1** Calibration report of CO<sub>2</sub>,  $\delta^{13}$ C-CO<sub>2</sub> and  $\delta^{18}$ O-CO<sub>2</sub>.

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Bias between reference and measured values of CO<sub>2</sub>,  $\delta^{13}$ C-CO<sub>2</sub> and  $\delta^{18}$ O-CO<sub>2</sub> are -0.01 ppm, 228 0.53 ‰ and 0.23 ‰ respectively. The precision and accuracy of CO<sub>2</sub>,  $\delta^{13}$ C-CO<sub>2</sub> and  $\delta^{18}$ O-CO<sub>2</sub> 229 are deduced with an averaging time of 100 sec. Results of the calibration in precision term are 230 0.05 %, 0.55 % and 3.97% for CO<sub>2</sub>,  $\delta^{13}$ C-CO<sub>2</sub> and  $\delta^{18}$ O-CO<sub>2</sub> respectively. The precision of  $\delta^{18}$ O-231  $CO_2$  is coarse compared  $\delta^{13}C$ - $CO_2$ , which may be improved by performing calibration for longer 232 averaging time. However, one needs to compromise for the precision averaging time (Guillon et 233 al., 2015). The second level quality was applied to the raw data by adjusting the respective 234 biases. 235

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#### 242 **2.3. Isotopic fraction using Improved model by Miller and Tans**

With the calibrated CO<sub>2</sub>, <sup>13</sup>C-CO<sub>2</sub> and <sup>18</sup>O-CO<sub>2</sub> data, implemented smoothing technique proposed by Thoning et al. (1989). This curve fitting function consist of a polynomial and harmonics terms, which accounts for short-term variability in the data. Subsequently, we computed biases ( $\Delta$ ) of CO<sub>2</sub>,  $\delta^{13}$ C-CO<sub>2</sub> and  $\delta^{18}$ O-CO<sub>2</sub> respectively in order to calculate the isotopic signature as explained by Miller and Tans et al. (2003). Following equation is the improved model by Miller and Tans, which was implemented in the present study.

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$$(\delta^{13}C \times CO_2)_{obs} - (\delta^{13}C \times CO_2)_{smooth} = \delta_s (CO_{2_{obs}} - CO_{2_{smooth}})$$
(4)

 $\delta_{s_{s}}CO_{2_{obs}}$  and  $CO_{2_{smooth}}$  in equation (4) are slope representing multiple sources, observational CO<sub>2</sub> and smoothed CO<sub>2</sub> respectively. Further to obtain the slope between  $\Delta CO_2$  versus  $\Delta$ ( $\delta^{13}C \times CO_2$ ) and  $\Delta(\delta^{18}O \times CO_2)$ , ordinary least square (OLS) method was applied on day and night time hours during the study period. The OLS is more commonly known as linear regression which may be simple regression or multiple depending on number of explanatory variables. Generic model of the OLS is defined as shown in Equation (5).

$$y = a_0 + \sum_{i=1} \delta_i x_i + \epsilon$$
(5)

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where y is  $\Delta$  ( $\delta^{13}C \times CO_2$ ), the dependent variable,  $a_0$  is the intercept of the model,  $\delta_i$  and  $x_i$  are the slope and  $\Delta CO_2$  corresponds to the i<sup>th</sup> explanatory variable of the model (i = 1 to n), and  $\epsilon$  is the random error. Similarly, equations 4 & 5 are implemented for computing the slope between  $\Delta CO_2$  versus  $\Delta$ ( $\delta^{18}O \times CO_2$ ). With these methods, further results are discussed in the following sections.

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### 284 **3.Results and Discussion**



285 **3.1. Seasonal Diurnal variation of CO<sub>2</sub>**,  $\delta^{13}$ C and  $\delta^{18}$ O

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**Figure 3.** (a) Time series of (daily averaged) CO<sub>2</sub> (top),  $\delta^{13}$ C (middle) and  $\delta^{18}$ O (bottom). (b) Seasonal diurnal variations of CO<sub>2</sub>,  $\delta^{13}$ C,  $\delta^{18}$ O and boundary layer hight during the study period.

Global to regional carbon cycle can be understood in detail by using the stable isotopic 290 composition of atmospheric  $CO_2$  as a tracer. Hence, the present study is attempted to produce 291 results of atmospheric CO<sub>2</sub> and its controlling factors in the sub-urban region of India using 292 simultaneous stable isotopic measurements. The time series analysis as shown in figure 3a 293 clearly depicts temporal variability in CO<sub>2</sub>, <sup>13</sup>C-CO<sub>2</sub> and <sup>18</sup>O-CO<sub>2</sub> at the study site. Daily means of  $\delta^{13}$ C and  $\delta^{18}$ O show a clear decrease with an increase of atmospheric CO<sub>2</sub>, which indicating 294 295 the associated source-sink mechanisams. The data gaps in CO<sub>2</sub>,  $\delta^{13}$ C and  $\delta^{18}$ O during 23<sup>rd</sup> May 296 to 19<sup>th</sup> June, 2019 was due to instrument technical snag. During the study period, the 297 atmospheric CO<sub>2</sub> concentration ranged from 405 ppm to 450 ppm, with its stable isotopic 298 composition ranging from -7.72 to -14.55 ‰ VPDB for  $\delta^{13}$ C and from -24.63 to 28.91 ‰ VPDB 299 for  $\delta^{18}$ O respectively. Moreover, larger standard devations in CO<sub>2</sub> and its stable isotopes are 300 observed in October, 2019 indicating the large variability. This could be due to varied sources 301 namely change in surface emissions, seasonal biomass burnig and long-range influence along 302 303 with the reversal of monsoonal winds (north-east) at the study site. However, found smaller variability in CO<sub>2</sub> and its stable isotopes during Indian summer monsoon (ISM, June-July-304 August-September) indicating marine fluxes (Tiwari et al., 2014) besides local sources. 305

The study site experience 4 seasons namely post-monsoon (November-December, 306 vegetation-II), winter (January-February), pre-monsoon (March-May, also known as summer 307 months-dry season) and summer monsoon (June-September; vegetation-I). Thus, figure 3b 308 shows diurnal variations of seasonal CO<sub>2</sub>,  $\delta^{13}$ C and  $\delta^{18}$ O against BLH. The height of the 309 Boundary Layer is the vertical extent of air column driven by convection processes associated 310 with the earth's surface heating (Stull, 1988). Due to strong convection and associated surface 311 temperature, the BLH attains maximum height in the afternoon, which modulates the dispersion 312 of air pollutats in the mixed layer. In contrast to this, the BLH quickly collapses and reaches 313 stable form during night hours due to the absence of convection processes. BLH is maximum 314 (minimum) during pre-monsoon (summer monsoon) with 2839 m (1167 m) respectively, which 315 316 indicates strong convection in the dry season. Subsequently, observed low (high)  $CO_2$ concentration during strong (absence) convective hours. The maximal seasonal variation in CO<sub>2</sub> 317 is found during pre-monsoon with a mean value of  $418.88 \pm 4.07$  ppm. The strong afternoon 318 drop of CO<sub>2</sub> during pre-monsoon is associated with the peak BLH of 2839 m. In contrast, 319 minimum seasonal variation in CO<sub>2</sub> is found during summer monsoon due to low BLH (1167 m). 320 Thus, exhibits prominent diurnal variations of CO<sub>2</sub> and  $\delta^{13}$ C against BLH during all the seasons 321 which is also attributed to isotopic fractionation processes during biological activity (Demény 322 and Haszpra 2002). Except for monsoon, diminished diurnal seasonal variability observed with 323  $\delta^{18}$ O (Murayama et al., 2010) compared to CO<sub>2</sub> and  $\delta^{13}$ C at present study site. Besides the 324 influence of BLH, minimum CO<sub>2</sub> and maximum  $\delta^{13}$ C and  $\delta^{18}$ O during afternoon hours are due to 325 the uptake of  $CO_2$  by the plants through photosynthesis process. During night hours, the 326 concentration CO<sub>2</sub> is maximum with low  $\delta^{13}$ C and  $\delta^{18}$ O due to active terrestrial respiration 327 (Sreenivas et al., 2016). Therefore, seasonal diurnal patterns of CO<sub>2</sub> and  $\delta^{13}$ C are anti-correlated 328 during all the seasons with minimum CO<sub>2</sub> in daytime and maximum  $\delta^{13}$ C respectively (figure 329 3b). Similarly, Pang et al. (2016) observed a negative relationship of  $\delta^{13}$ C diurnal cycle with CO<sub>2</sub> 330 mixing ratio at Beijing, in Northern China. 331

Seasonal averages of CO<sub>2</sub> ( $\delta^{13}$ C &  $\delta^{18}$ O) during post-monsoon, winter, pre-monsoon and 332 summer monsoon are 414.70±4.49 ppm (-9.41±0.27 ‰ & 20.03±0.73 ‰), 416.84±3.25 ppm (-333 9.25±0.21 ‰, & 25.95±0.57 ‰), 418.88±4.07 ppm (-11.88±0.13 ‰ & 6.51±0.98 ‰), 334  $412.61\pm7.59$  ppm (-12.76±0.38 ‰ & -3.49±2.87 ‰) respectively. Site specific  $\delta^{13}$ C seasonal 335 values with other study sites in India and across the world are summarized in table 2. Liu et al. 336 (2014a) showed annual means of  $\delta^{13}$ C varying from -8.30 ‰ to -8.35 ‰ at Waliguan station in 337 China. Further, the annual means of  $\delta^{13}$ C are -8.27 ‰ and -8.36 ‰ respectively for 2009 and 338 2010 at Shangdianzi station which is located in a small village about 100 km northeast of Beijing 339 (second populated urban city in China). This site is influenced by strong pollution events from 340 Beijing and surrounding urban areas in the presence of southwesterly winds. Also, the annual 341 mean of  $\delta^{13}$ C were -8.55%, -8.52%, -8.46% and -8.61% during 2007-2010 at Tae-ahn Peninsula 342 which is located on a small Peninsula on the western coast of Korea. Irrespective of the sites 343 background influence, results of worldwide sites mentioned above indicates  $\delta^{13}$ C values are in 344 close proximity with our present observations. 345

An increase and decrease of <sup>18</sup>O/<sup>16</sup>O ( $\delta^{18}$ O) ratio of CO<sub>2</sub> in the atmosphere across the seasons indicates dominance of photosynthesis and ecosystem respiration. During the study period, the atmospheric  $\delta^{18}$ O–CO<sub>2</sub> varies seasonally from -3.49 ‰ to + 25.95 ‰ with large scatter compared to seasonal CO<sub>2</sub> and  $\delta^{13}$ C. The scatter could be due to vegetation cover, leaf water content, retention of soil water and ocean (Zhou et al., 2006). However, interpretation of

seasonal  $\delta^{18}$ O is not that straight as CO<sub>2</sub> and  $\delta^{13}$ C due to the varying fluxes of biospheric CO<sub>2</sub> 351 352 and prevailing meteorology at the study site.

High atmospheric CO<sub>2</sub> in pre-monsoon followed by winter and subsequent increase of  $\delta^{18}$ O-CO<sub>2</sub> 353 in the atmosphere in the absence of relatively low photosynthesis could be attributed to long-354 range air mass transport with enhanced  $\delta^{18}$ O-CO<sub>2</sub> and fossil fuel burning activities (Muravama et 355 al., 2010). Through isotopic exchange,  $\delta^{18}$ O-CO<sub>2</sub> in the atmosphere may affect due to variation in 356 the <sup>18</sup>O from precipitation and soil respiration (Kato et al., 2004). Newman et al. (2008) observed 357 the range of  $\delta^{18}$ O at Los Angeles basin, Southern California during 1972 -1973 (1998-2003) is -358 3.56 ‰ to + 0.21 ‰ (-3.99 ‰ to +0.45 ‰) and average value during 1972-1973 (1998-2003) of 359  $\delta^{18}$ O is -1.28 ‰ (-1.07‰). 360

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262	Study sites	$\delta^{13}$ C values (‰)	References
303		Seasonal	
364	Dallas, USA	-12.0 to -8.1	Clark-Thorne and
504			Yapp, 2003
365	Bern, Switzerland	-14 to -8	Sturm et al.,2006
366	Salt Lake City, USA	-18 to -8	Pataki et al., 2006
300		(December 2004 - January 2005)	
367		-9.3 to -7.5 (Oct)	Djuricin et al., 2010
368	Los Angeles, USA	-12.5 to -8.8 (Dec)	
369		-12.2 to -9.2 (Feb)	
370		-12.5 to -10.2 (April)	
371			
372	Nagoya, Japan	-13.4 to -8.5(May)	Wada et al., 2011
372		-15.0 to -8.5	
373		(December from 2008 - 2009)	71 1 . 1 2004
374	Krakow, Poland	-11 to -9.5	Zimnoch et al., 2004
375	Wroclaw (SW Poland)	-16.4 to -8.2	Gorka et al., 2013
376	Cabo de Rama	-8.4 to -7.8	Bhattacharya et al.,
377	(West Coast of India)		2009
279	NRSC, Shadnagar,	-9.3 to -12.8	Present study
3/8	India	(November 2018 - October 2019)	
379			

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**Table 2**  $\delta^{13}$ C values from different study sites.

Seasonal amplitude (peak to peak  $\pm 1$  SD) of atmospheric CO<sub>2</sub> was 9.82  $\pm 1.39$  ppm, 7.19  $\pm 0.11$ 382 ppm, 9.23  $\pm 2.1$  ppm and 17.30  $\pm 9.29$  ppm in post-monsoon, winter, pre-monsoon and summer 383 monsoon respectively. Weak and strong seasonality was observed during winter and monsoon 384 seasons repectively reflecting enrichment of atmospheric CO<sub>2</sub> in low vegetation months and 385 depleting CO<sub>2</sub> in high vegetation season. The NDVI in figure 1b show maximum during 386 monsoon and minimum during rest of the seasons. The Sinhagad and Cape Rama, two marine 387 stations of western India showed CO<sub>2</sub> seasonal amplitudes between 8-10 ppm during monsoon 388 and >15ppm for remaining seasons (Tiwari et al., 2014), which could be due to the influence of 389 land-sea breeze (Mahesh et al., 2014). The seasonal amplitude of  $\delta^{13}$ C ( $\delta^{18}$ O) are 0.48 ± 0.06 ‰ 390  $(1.91 \pm 0.005 \text{ \%}), 0.44 \pm 0.02 \text{ \%} (1.60 \pm 0.13 \text{ \%}), 0.32 \pm 0.09 \text{ \%} (2.53 \pm 1.003 \text{ \%}) \text{ and } 0.87 \pm 0.07 \text{ \%}$ 391  $0.73 \% (6.58 \pm 4.10 \%)$  in post-monsoon, winter, pre-monsoon and summer monsoon 392 respectively. Thus, modulation in the seasonal amplitude of  $CO_2$  in the atmosphere is attributed 393 to the rate of photosynthesis and respiration besides the impact of local and long-range prevailing 394 meteorology. For better understanding of the relationship between atmospheric  $CO_2$  against its 395

stable isotopic composition, improved model by Miller and Tans was adopted by fitting linear regression using the OLS.

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# 399 **3.2. Seasonal correlation between CO<sub>2</sub>,** $\delta^{13}$ C and $\delta^{18}$ O using the improved model by Miller 400 and Tans

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Since the study site is not a single sourced location and is in proximity of Hyderabad city, a 402 metropolitan city of India, the Miller-Tans (2003) method was applied on the day and night time 403 observations to characterize the effective  $\delta^{13}C({}^{13}C/{}^{12}C)$  source ratios at the study site. In order to 404 estimate the slope ( $\delta_s$ ) for  $\delta^{13}C$  and  $\delta^{18}O$ , the ordinary least square curve fitting was applied to 405  $\Delta CO_2$  against  $\Delta (CO_2 \times \delta^{13}C)$  and  $\Delta (CO_2 \times \delta^{18}O)$  respectively and results shown in Figure 4(a-b). 406 The improved model by Miller and Tans is applied to capture the processes of day and night 407 which are driving the atmospheric CO<sub>2</sub>. During the day (night) time  $\delta^{13}$ C slopes in post-408 monsoon, winter, pre-monsoon and monsoon are  $-10.43 \pm 5.60$  % (-27.08  $\pm 4.21$  %), -36.43  $\pm$ 409 8.11 % (-25.11 ± 5.55 ‰), -32.02±4.57 ‰ (-23.79 ± 2.08 ‰) and -52.50±4.60 ‰ (-28.38 ± 1.29 410  $\infty$ ) respectively with coefficient of determination (R<sup>2</sup>) ranging from 0.06 (0.26) in post-monsoon 411 (winter) to 0.56 (0.83) in monsoon. The strong correlation in summer-monsoon during day and 412 413 night time can be attributed to the long-range air mass sources transported by the monsoonal winds and sinks may be attributed to the terrestrial biospheric activities at the study site. The  $\delta^{13}$ C 414 slope during daytime is largely varied in all the seasons representing the contribution of mixed 415 source emissions at the regional scale and suppressed local sources (Xu et al., 2017). An average 416 value of daytime  $\delta_s$  of  $\delta^{13}$ C during all seasons is -32.84 ‰ indicating the source of atmospheric 417 CO<sub>2</sub> is related to gasoline and natural gas combustion in and around the study site (Clark-Thorne 418 and Yapp, 2003). The estimated slopes are evaluated statistically and found significant (p-value 419 < 0.05) with 95 % confidence interval during all the seasons. The  $\delta^{13}$ C slopes in night time 420 during all the seasons are between -23.79 % to -28.38 %, with an average value of -26.09 %. In 421 general, the average value of  $\delta_s$  of  $\delta^{13}$ C-CO<sub>2</sub> is -26.20 ‰ for C<sub>3</sub> ecosystem respiration (Pataki et 422 al., 2003). Thus, the present study also confirm the dominance of C<sub>3</sub> vegetation at the study site 423 contributes to the emissions of atmospheric CO<sub>2</sub> mixing ratio during night time. The varied  $\delta_s$  of 424 425  $\delta^{13}$ C-CO<sub>2</sub> in different study locations are attributed to the local anthropogenic activities and 426 terrestrial biospheric pathways.

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As shown in figure 4b, improved model by Miller and Tans was implemented on  $\delta^{18}$ O by 428 substituting for  $\delta^{13}$ C in equation 4 to see the relationship between  $\delta^{18}$ O and CO<sub>2</sub> concentration. 429 Figure 4b show strong correlation of  $\delta^{18}$ O with R<sup>2</sup> = 0.75 in night hours of monsoon. During the 430 same period,  $\delta^{13}$ C is also strongly correlated with CO<sub>2</sub> concentration, indicating the active role of 431 biospheric – atmospheric interactions in the exchange of CO<sub>2</sub>. In contrast, varied seasonal slopes 432 of  $\delta^{13}$ C and  $\delta^{18}$ O were observed in daytime representing the study site is also dominated by the 433 regional sources via upwind transport along with the influence of local terrestrial activities 434 namely boundary layer dynamics and photosynthesis. During daytime, the  $\delta_s$  of  $\delta^{18}$ O shows 435 scatter across the seasons for which the driving forces are different (Yakir et al., 2011). With the 436 present study, we understand that interpretation of the seasonal  $\delta^{18}$ O variation is complicated 437 compared to  $\delta^{13}$ C and CO<sub>2</sub> mixing ratio. However, current knowledge on  $\delta^{18}$ O seasonality can be 438 improved with great understanding of varying fluxes of bisopheric CO<sub>2</sub> and role of prevailing 439 meteorology at the study site. 440

The  $\delta_s$  of  $\delta^{13}$ C and  $\delta^{18}$ O analysis at the study site confirms the prominence of C<sub>3</sub> ecosystem activities viz. photosynthesis and respiration in the variability of atmospheric CO<sub>2</sub> during vegetative seasons. During dry seasons (winter and pre-monsoon), the  $\delta_s$  values indicates the important role of local and long-range combustion processes that enrich atmospheric CO<sub>2</sub> at the study site. However, this information is pertained to the total source contributions and individual source apportionment was not carried out in this study.





**Figure 4.** Improved Miller and Tans method applied to the (a) Seasonal  $\Delta CO_2 \text{ vs } \Delta(CO_2 \times \delta^{13}C)$ during day (Top left panel) and nighttime (Top right panel) hours (b) Bottom left panel shows  $\Delta CO_2 \text{ vs } \Delta(CO_2 \times \delta^{18}O)$  during daytime and bottom right panel shows nighttime. The solid line represents the regression fit using ordinary least square method.

	<b>Daytime</b> $\Delta CO_2 \text{ vs } \Delta (CO_2 \times \delta^{13}C)$		Nighttime	
Month			$\Delta CO_2 \text{ vs } \Delta (CO_2 \times \delta^{13}C)$	
	$[\Delta CO_2 \text{ vs } \Delta (CO_2 \times \delta^{18}O)]$		$[\Delta CO_2 \text{ vs } \Delta (CO_2 \times \delta^{18}O)]$	
	ʻr'	Slope	ʻr'	Slope
November, 2018	-0.49 [-0.12]	-18.30 [-23.18]	-0.75[-0.21]	-29.90[-26.40]
December, 2018	-0.52 [-0.57]	-42.30 [-132.37]	-0.56[+0.25]	-24.72[+26.38]
January, 2019	-0.56 [-0.46]	-33.12 [-98.68]	-0.63[-0.80]	-24.18[-176.55]
February, 2019	+0.40 [+0.09]	+28.83 [+22.94]	-0.41[+0.21]	-27.01[+49.61]
March, 2019	-0.49 [-0.14]	-28.51 [-79.64]	-0.66[-0.33]	-18.72[-65.29]
April, 2019	-0.65 [-0.33]	-33.94 [-92.92]	-0.84[-0.44]	-26.10[-77.5]
May, 2019	-0.84 [-0.51]	-37.58 [-200.60]	-0.81[-0.72]	-24.91[-103.5]
June, 2019	-0.15 [-0.87]	-11.31 [-279.45]	-0.95[-0.96]	-24.25[-113.25]
July, 2019	-0.40 [-0.51]	-24.25 [-182.29]	-0.63[-0.71]	-26.29[-126.12]
August, 2019	-0.71 [-0.55]	-52.01 [-239.16]	-0.93[-0.89]	-29.29[-122.94]
September,	-0.94 [-0.79]	-72.46 [-287.80]	-0.93[-0.87]	-28.36[-118.20]
2019				
October, 2019	-0.95 [-0.95]	-51.23 [-227.49]	-0.99[-1.0]	-27.39[-140.52]

#### 488

#### 489 **Table 3** Monthly correlation coefficient (r), and slope $(\delta_s)$ derived from the improved Miller and 490 Tans method

Table 3 shows monthly 'r' and slope values derived between  $\Delta CO_2$  vs  $\Delta$  (CO<sub>2</sub> × $\delta^{13}$ C) and  $\Delta CO_2$ 491 vs  $\Delta$  (CO<sub>2</sub> ×  $\delta^{18}$ O) during day and night hours respectively. The average monthly value of  $\delta^{13}$ C 492 slope during night time (daytime) is -26 ‰ (36.15‰). During nighttime the average value of 493 slope at the study site is close to  $C_3$  ecosystem respiration and during day time the average value 494 of slope is close to sources which are combustion, photosynthesis processes and long range 495 496 transport. The differences in slope value at various study sites is due to photosynthesis pathways and local anthropogenic sources prevailing at the study areas (Pang et al., 2016). In similar 497 studies carried by various authors slope values at their study site is as follows. Liu et al. (2014a) 498 observed annual mean value of slope which is -25.44‰ and -21.70‰ at Waliguan and 499 Shangdianzi stations in China respectively. Zhou et al. (2006) studied the isotopic fractionation 500 at 12 stations of Northern Hemisphere and found slope ( $\delta_s$ ) ranging from -28.85‰ to - 26.50‰ 501 with improved Miller-Tans method. Murayama et al. (2010) observed  $\delta_s$  value of -28.7‰ 502 averaged over the study period at Takayama site in central Japan which is also comparable to the 503 present study site. The  $\delta_s$  of  $\delta^{18}$ O during vegetation - I and II periods is strongly correlated (r > 504 0.7) compared to 'r' value during other months. Thus, vegetation cover is an important 505 component in variability of  $\delta^{18}$ O of atmospheric CO<sub>2</sub>. 506

The correlation coefficient (r) between  $\delta^{18}O$  and  $\delta^{13}C$  during daytime (night time) are 507 0.13 (0.07), 0.64 (0.60), 0.95 (0.93), 0.24 (0.92) for post-monsoon, winter, pre-monsoon and 508 summer monsoon seasons respectively. A very strong positive correlation (0.93 to 0.95) in day 509 and night hours during pre-monsoon between  $\delta^{18}$ O and  $\delta^{13}$ C might be due to high vapour 510 pressure deficit (VPD) than the post monsoon (low VPD). The high VPD and summer conditions 511 are supportive for high transpiration as well as photosynthetic activities as sunshine hours are 512 more during pre-monsoon than post monsoon (approaching towards winter). Low  $\delta^{13}$ C and  $\delta^{18}$ O 513 correlation during post-monsoon indicates high stomatal conductance in vegetation, ensuring 514 enhanced release of water to the atmosphere (Cullen et al., 2008; Liu et al., 2014b). Our study 515

showed clear evidence of an increase or decrease of atmospheric  $CO_2$  is associated with the 516 517 changes of its isotopic composition during photosynthesis and biogenic respiration besides local and long-range anthropogenic influences. Also, Zimnoch et al. (2004) observed good correlation 518 519 between  $\delta^{18}$ O and  $\delta^{13}$ C with 'r' ranging from 0.84 to 0.94 in their observations while studying their diurnal variability from Poland. Further to understand the influence of transport pathways 520 on atmospheric CO<sub>2</sub> via long range air mass is examined in the present study using Hybrid 521 Single Particle Lagrangian Integrated Trajectory (HYSPLIT, 522 https://www.ready.noaa.gov/HYSPLIT.php) Model. 523

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# 525 **3.3. Influence of local and long-range airmass on atmospheric CO<sub>2</sub>**

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527 Study site being a sub-urban region and about 60 km away from the Hyderabad city, the analysis 528 of improved Miller-Tan model depicts atmospheric CO<sub>2</sub> concentration at the study site is mainly

529 controlled by terrestrial biosphere activities during the study period. During the day hours, the

derived  $\delta_s$  of  $\delta^{13}$ C indicating mixed source contribution at the study site, which may be possibly

531 due to combustion activities, biomass burning and the transportation of air mass. Seasonal wind

- vector obtained from the ECMWF at 850 hPa over the Indian region are shown in figure 5a. The
- 533 long-range air mass circulation that is reaching the study site has been analyzed using
- 534 Lagrangian back-trajectory model along with the fire counts during all the seasons (Figure 5b).



**Figure 5**. (a) Mean winds (m s<sup>-1</sup>) at 850 hPa (b) Seasonal long-range air mass circulation at 2 km altitude using HYSPLIT

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538 During the Indian summer-monsoon season, predominantly the south- westerly (SW) winds are 539 present as shown in figure 5a, which brings maritime air mass (Bhattacharya et al., 2009; Tiwari

- et al., 2014) over the Indian region. The atmospheric  $CO_2$  concentration at the study site is
- observed minimum during the summer-monsoon period indicating the contribution of monsoon

circulation through scavenging effect (Tiwari et al., 2014; Mahesh et al., 2014). The maritime air 542 543 mass during summer-monsoon is relatively pristine due to the absence of anthropogenic sources thus observed low atmospheric CO<sub>2</sub> concentration at the study site. 544

A Lagrangian back trajectory analysis as shown in figure 5b also confirms the observed 546 low atmospheric CO<sub>2</sub> due to influence of maritime air mass (88 %) during the summer-monsoon 547 period. During winter and pre-monsoon, enrichment of atmospheric CO<sub>2</sub> at the study site could 548 be due to the predominant continental sources reaching from the north-east (NE, 76 %) and 549 north-west (NW, 42 %) respectively. Figure 5b shows seasonal long-range air mass circulation 550 over laid with fire count obtained from Moderate Resolution Imaging Spectroradiometer 551 552 (MODIS) during the study period. During pre-monsoon and post-monsoon, agriculture residue burning is commonly observed in Punjab, Haryana and Indo Gangetic Plains (IGBP) areas of 553 India (Liu et al., 2019), which are in NW and NE directions of the study site. Thus, the present 554 study reveals that during these seasons, the biomass burning is one of continental sources 555 contributing to the elevated  $CO_2$  concentration. Therefore the variability of atmospheric  $CO_2$ 556 concentration at the study site is greatly influenced by the maritime air mass during summer-557 558 monsoon and continental sources during other seasons.

#### 4. Conclusion 560

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The present study examined the diurnal and seasonal variations of atmospheric CO<sub>2</sub> and its stable 562 isotopes  $\delta^{13}$ C and  $\delta^{18}$ O at the sub-urban site of Telangana, India during November 2018 to 563 October 2019 using high precision CO<sub>2</sub> isotopic analyzer. 564

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- Following are the salient findings of the present study
- 1. The atmospheric  $CO_2$  concentration ranged from 405 ppm to 450 ppm, with its stable 568 isotopic composition ranging from -7.72 to -14.55 % VPDB for  $\delta^{13}$ C and from -24.63 to 569 28.91 ‰ VPDB for  $\delta^{18}$ O respectively. 570
- 2. The CO<sub>2</sub> and  $\delta^{13}$ C exhibit clear diurnal variation with opposite patterns during all the seasons with minimum CO<sub>2</sub> and maximum  $\delta^{13}$ C in daytime and vice versa. 572
- 3. During strong convective hours (peak BLH), observed low atmospheric CO<sub>2</sub> 573 concentration and high  $CO_2$  concentration in weak convection period indicating strong 574 atmospheric mixing due to the convective boundary layer. 575
- 4. Seasonal amplitude (peak to peak  $\pm 1$  SD) of atmospheric CO<sub>2</sub> at the study site was 9.82  $\pm$ 576 1.39 ppm,  $7.19 \pm 0.11$  ppm,  $9.23 \pm 2.1$  ppm and  $17.30 \pm 9.29$  ppm in post-monsoon, 577 winter, pre-monsoon and summer monsoon respectively. Large seasonality in summer-578 579 monsoon at the study site due to strong influence of monsoonal winds, precipitation and vegetation cover. 580
- 5. To capture the driving processes of atmospheric  $CO_2$  during day and night hours, we thus 581 applied improved Miller-Tans model on seasonal  $\Delta CO_2$ ,  $\Delta$  (CO<sub>2</sub> ×  $\delta^{13}C$ ) and  $\Delta$  (CO<sub>2</sub> × 582  $\delta^{18}$ O) data. During all seasons, mean  $\delta_s$  of  $\delta^{13}$ C value is -32.84 ‰ in daytime indicating 583 the source of atmospheric  $CO_2$  is related to gasoline and natural gas combustion at the 584 study site and nighttime  $\delta_s$  values were between -23.79 ‰ to -28.38 ‰, with an average 585 value of -26.09 % representing the dominance of C<sub>3</sub> ecosystem respiration. 586

- 587 6. Seasonal 'r' value between  $\delta^{18}$ O and  $\delta^{13}$ C during day and night time varied from 0.13 to 588 0.95 and 0.07 to 0.93 respectively. A very strong positive correlation (0.93 to 0.95) in day 589 and night hours during pre-monsoon between  $\delta^{18}$ O and  $\delta^{13}$ C might be due to high VPD 590 than the post monsoon (low VPD).
- 591 7. Upwind transport confirms the influence of biomass burning on enriched atmospheric
   592 CO<sub>2</sub> during pre-monsson and post-monsoon seasons.
- A lagrangian back-trajectories confirms the variability of atmospheric CO<sub>2</sub> concentration
   at the study site is largely influenced by the maritime airmass during summer-monsoon
   and continental sources in other seasons.
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597 Our study showed clear evidence of an increase or decrease of atmospheric  $CO_2$  is 598 associated with the changes of its isotopic composition during photosynthesis and 599 biogenic respiration besides local and long-range anthropogenic influences. The 600 variability in atmospheric  $CO_2$  during monsoon season is strongly associated with the 601 ISM. However, the individual source apportionment of different sources is not discussed 602 in this present study.

603

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605

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615

# 616 **Declaration of competing interest**

- 617618 The authors declare no competing interests.
- 619

620 Data Availabilty Statement

621

The *in- situ* data may be available publicly once archival is completed.

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