1 Reactive chlorine, sulphur and nitrogen containing volatile

2 organic compounds impact atmospheric chemistry in the

3 megacity of Delhi during both clean and extremely polluted

4 seasons

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Abstract. Volatile organic compounds significantly impact the atmospheric chemistry of polluted megacities.
Delhi is a dynamically changing megacity and yet our knowledge of its ambient VOC composition and chemistry

- 15 is limited to few studies conducted mainly in winter before 2020 (all pre-COVID). Here, using a new extended
- 16 volatility range high mass resolution (10000-15000) Proton Transfer Reaction Time of Flight Mass Spectrometer,
- 17 we measured and analyzed ambient VOC-mass spectra acquired continuously over a four-month period covering
- 18 "clean" monsoon (July-September) and "polluted" post-monsoon seasons, for the year 2022. Out of 1126 peaks,
- 19 111 VOC species were identified unambiguously. Averaged total mass concentrations reached $\sim 260 \mu \text{gm}^{-3}$ and
- 20 were >4 times in polluted season relative to cleaner season, driven by enhanced emissions from biomass burning
- 21 and reduced atmospheric ventilation (~2). Among 111, 56 were oxygenated, 10 contained nitrogen, 2 chlorine, 1
- sulphur and 42 were pure hydrocarbons. VOC levels during polluted periods were significantly higher than most
- 23 developed world megacities. Methanethiol, dichlorobenzenes, C6-amides and C9-organic acids/esters, which have
- 24 previously never been reported in India, were detected in both the clean and polluted periods. The sources were
- 25 industrial for methanethiol and dichlorobenzenes, purely photochemical for the C6-amides and multiphase
- 26 oxidation and partitioning for C9-organic acids. Aromatic VOC/CO emission ratio analyses indicated additional
- 27 biomass combustion/industrial sources in post-monsoon season, alongwith year-round traffic sources in both
- 28 seasons. Overall, the unprecedented new information concerning ambient VOC speciation, abundance, variability
- 29 and emission characteristics during contrasting seasons significantly advances current atmospheric composition
- 30 understanding of highly polluted urban atmospheric environments like Delhi.
- 31

32 1 Introduction

33 The national capital territory of Delhi in India is jointly administered by the central and state governments and 34 accommodated more than 32 million people in 2022. For the past several years, its population has grown at the

35 rate of more than 2.7 percent per year, adding about 1 million new inhabitants each year. Thus, the region

- 36 represents a complex dynamically changing emission environment driven by rapid changes in emissions as 37 regulatory authorities make efforts to improve urban infrastructure and public transportation while promoting
- 37 regulatory authorities make efforts to improve urban infrastructure and public transportation while promoting38 cleaner technologies. As a megacity in a developing country with one of the world's highest population densities,
- 39 Delhi exemplifies some of the key challenges faced by many megacities in the global south, where increased
- 40 urbanization and inequitable access to clean energy sources along with unfavourable meteorological conditions
- 41 during cold periods of the year, cause the inhabitants to suffer from extreme air pollution episodes. Lelieveld et
- 42 al. (2015) identified South Asia as one of the global air pollution hotspots in terms of the contribution of outdoor
- 43 air pollution sources to premature mortality due to particulate matter pollution. Reduction of other atmospheric
- 44 pollutants is also deemed necessary to fulfil the UN Sustainable Development Goals (Keywood et al., 2023). Thus,
- 45 the study of Delhi's ambient chemical composition using state of the art technology can offer valuable insights
- 46 and lessons for our understanding of polluted atmospheric environments.
- 47 Previous studies have demonstrated that air pollution in the Delhi-NCR metropolitan area peaks during the post-48 monsoon (October- November) season (e.g. Kulkarni et al., 2020), coinciding with the time of year when large 49 scale paddy stubble burning occurs in the Indo-Gangetic Plain (Kumar et al., 2021). The main air pollutant in 50 exceedance has long been identified to be particulate matter (e.g. PM_{2.5}) and many studies (Gani et al., 2020; Cash 51 et al., 2021; Sharma et al., 2023; Singh et al., 2011) have documented the variability, exceedance and composition 52 of aerosols. Volatile organic compounds (VOCs) are major precursors of secondary organic aerosol, which is a 53 significant component of PM_{2.5} (30-60% in Delhi; Chen et al., 2022; Nault et al., 2021) and surface ozone over 54 Delhi. In fact, in-situ ozone production in Delhi has been reported to be more sensitive to VOCs rather than 55 nitrogen oxides (Nelson et al., 2021). Several VOCs (e.g. benzene, nitromethane, 1,3-butadiene) are also 56 carcinogenic (WHO 2010) at high exposure concentrations and many pose direct health risks (Ho et al., 2006; 57 Espenship et al., 2019; WHO 2019; Weng et al., 2009; Roberts et al., 2011; Durmusoglu et al., 2010). VOCs can 58 also aid source apportionment studies by acting as source fingerprints and valuable molecular markers of specific 59 emission sources (de Gouw et al., 2017; Holzinger et al., 1999; Warneke et al., 2001; Kumar et. al., 2020; Garg et 60 al., 2016; Hakkim et al; 2021; Kumar et al., 2021). In the complex emission environment of cities in the developing 61 world, this can be especially helpful since the energy usage portfolio is such that biomass burning sources are 62 likely to be as significant as fossil-fuel based sources (Bikkina et al., 2019) in influencing the air pollutant burden 63 of VOCs, resulting in ambient air VOC composition that could be quite different from cities like Los Angeles 64 (McDonald et al., 2018).
- Existing knowledge about the abundance and diurnal variability of major ambient VOCs such as methanol,
 acetone, acetaldehyde, acetonitrile, isoprene, benzene, toluene, xylenes and trimethyl benzenes in Delhi, is limited
 to just four previously measured wintertime datasets: Dec-March of 2016 (Chandra et al., 2018; Hakkim et al.,
- 68 2019), Dec-March of 2018 (Wang et al., 2020; Tripathi et al., 2022), few days in October 2018 (Nelson et al.,
- 2017), bee watch of 2010 (wang et al., 2020, inpath et al., 2022), iew days in October 2010 (reison et al.,
- 69 2021; Bryant et al., 2023) and one spanning 145 days of 2019 that reported source apportionment of some VOCs
- 70 for different seasons (Jain et al., 2022). We note that all these were pre-COVID period datasets, and that since
- these observations many new regulations have been put in place e.g. for traffic with the introduction of BS-VI
- 72 (EURO6 equivalent) in 2020 and the Faster Adoption and Manufacturing of hybrid and Electric vehicles (FAME)
- 73 program for promotion of E-vehicles, and for industries with a ban on the use of petcoke in the National Capital
- 74 Region (NCR) and the crackdown on unregistered industries (Guttikunda et al, 2023). After COVID lock-downs
- 75 happened in 2020, a new Commission for Air Quality Management in Delhi National Capital Region and its

- 76 Adjoining Areas (CAQM) was set up in November 2020 (https://caqm.nic.in/index.aspx?langid=1). Under its
- 77 mandate, depending on air quality level, it promulgates immediate graded response action plans (GRAP;
- 78 https://caqm.nic.in/index1.aspx?lsid=4168&lev=2&lid=4171&langid=1) that instruct civic authorities to shut-
- 79 down or restrict particular emission sources. Furthermore, on 7 August 2020, the Delhi government announced a
- 80 new Delhi Electric Vehicle (EV) Policy. In order to address the high-upfront cost of EVs (ICE vehicles), the Delhi
- 81 EV Policy provides demand incentives for purchasing electric vehicles. The incentives help bring cost parity for
 82 EVs and are in addition to those outlined in the Faster Adoption and Manufacturing of Hybrid and Electric
- 83 Vehicles (FAME II). In the budget allocation for 2020, the Government of India allocated \$600 million USD for
- Vehicles (FAME II). In the budget allocation for 2020, the Government of India allocated \$600 million USD for
 clean air measures through the Ministry of Housing and Urban Affairs (MoHUA) to 46 cities across India. These
- 85 have been detailed in a report by Arpan Chatterji (2020). Thus overall, important changes to the transport emission
- 86 sector, construction and urban industrial sector and residential sector were implemented at a policy level after
- 87 2020 to reduce air pollution in the Delhi-NCR region.
- The monsoon season which precedes the post-monsoon season lasts from June to September and is characterized
 by better air quality, aided by favourable meteorological conditions, including higher ventilation co-efficient,
 negligible agricultural waste burning and enhanced wet scavenging (Kumar et al., 2016).
- 91 This study addresses some of the above knowledge gaps pertaining to ambient VOCs during the "clean" monsoon 92 season characterized by baseline pollution levels and the polluted "post-monsoon" season characterized by 93 extreme pollution events and large scale open agricultural biomass waste fires regionally. Employing a new 94 extended volatility range (EVR) high mass resolution (10000-15000) Proton Transfer Reaction Time of Flight 95 Mass Spectrometer 10K (PTR-TOF 10000; Ionicon Analytik GmBH), a technology that has never before been 96 deployed in India, we investigated the ambient VOC speciation, abundance, variability and emission 97 characteristics in the polluted urban environment of Delhi over a 4-month period. This enabled us to discover 98 several low volatility VOCs, many of which are present in fire emissions (Koss et al., 2018), for the time in South 99 Asia, as all previous VOC studies have involved either the older PTR-TOF-MS or PTR-QMS instruments, that 100 have significantly lower mass resolution and lower detection sensitivity and did not possess the extended volatility 101 range components. We first undertook comprehensive and rigorous interpretation of the ambient mass spectra 102 over a four-month period spanning July-Nov of 2022 in Delhi. This was followed by identification and 103 quantification of 111 VOCs, many of which have been discovered and reported for the first time from the South 104 Asian atmospheric environment. Each of these compounds was then classified in terms of oxygenated VOCs, pure 105 hydrocarbons, major nitrogen containing VOCs, chlorine containing VOCs and sulphur containing VOCs, 106 followed by the time series analyses and diurnal profiles of the major VOCs and some new/rarely reported VOCs 107 in both seasons as a function of meteorology and emissions. The atmospheric chemistry implications of some of 108 the newly discovered compounds in this polluted urban environment are discussed. Further, using measured 109 aromatic VOC/CO emission ratios in monsoon and post-monsoon season, a global comparison with reports from 110 megacities in Europe, North America and Asia was undertaken for a nuanced understanding of their levels and
- sources in Delhi relative to megacities across these different continents.

- 112 2. Methodology
- 113 2.1 Measurement site and meteorological conditions:
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- 115

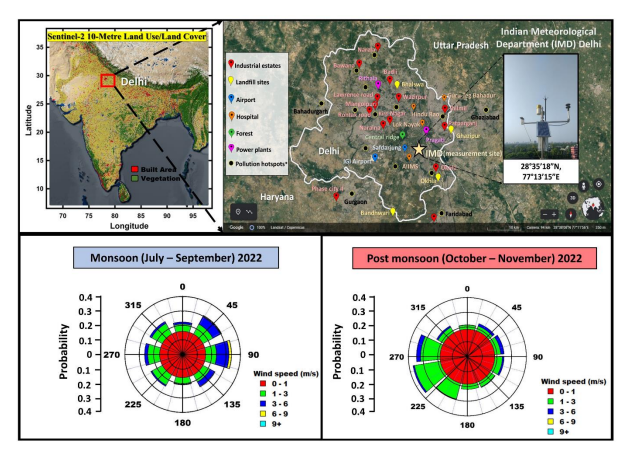


Figure 1: Map of India showing Delhi (1 a) and zoom in of the measurement site (star marked) (1b; Google Earth
 Imagery © Google Earth) with a view from the roof-top of the SatMet Building (28.5896°N-77.2210°E), and wind rose
 plots derived from in-situ one-minute wind speed and wind direction data during monsoon (1c) and post-monsoon
 (1d) 2022 acquired at sampling height of ~35m A.G.L

- 121
- 122 The measurement site was located within the premises of the India Meteorological Department (IMD) which is
- 123 situated in Central Delhi (Fig. 1). Ambient air was sampled at a height of circa 35m above ground level from the
- 124 roof-top of the SatMet building (28.5896°N-77.2210°E), into the instruments which were housed inside a
- 125 laboratory located in the sixth floor of the same building.
- 126 Figure 1 (a) shows the land use/ land cover (Sentinel-2 10m) map of India with a red marked box highlighting
- 127 Delhi. The city is bordered on its northern, western, and southern sides by the state of Haryana and to the east by
- 128 the state of Uttar Pradesh. The star marked in Fig. 1 (b) shows the measurement site (IMD Delhi) and its
- surroundings. The major pollution hotspots include places like Ghaziabad (towards the northeast), Bahadurgarh
- 130 (towards the northwest), Gurgaon (towards the southwest), and Faridabad and Okhla (towards the southeast),.
- 131 Major industrial areas e.g. Okhla industrial area, major landfill sites ,the international airport and some major
- hospitals are also shown in Fig 1 (b).

Meteorological sensors (Campbell Scientific Inc.) were deployed to measure the wind speed, direction, 133 134 temperature, relative humidity and photosynthetic active radiation (model nos.: CS215 for temperature and RH, 135 PAR PQSI sensor, and for rain TE525-L40). Boundary layer height was taken from ERA5 reanalyses dataset 136 (Hersbach et al., 2023) and ventilation coefficient was calculated as the product of the measured wind speed and 137 boundary layer height. Atmospheric ventilation or ventilation coefficient (VC) is a good proxy for the dilution 138 and dispersion of air pollutants near the surface (Hakkim et al., 2019). It is defined as the product of boundary layer height (m) and wind speed (ms⁻¹). The VC represents the rate at which air within the mixed layer is 139 140 transported away from a region of interest and provides information about how concentrations of pollutants are 141 modulated through transport of air over that region. Figures 1 (c) and 1 (d) show the wind rose plot derived from 142 the in-situ one-minute wind speed and wind direction data acquired at the measurement site for monsoon (July 143 2022 - September 2022) and post-monsoon (October 2022 - November 2022) seasons, respectively. The prevalent 144 wind direction changed from easterly flow in monsoon season to westerly flow in the post-monsoon season. 145 During the monsoon season, the major fetch region spanned from the NE to SE-E. These NE, E, and SE winds were associated with high wind speeds ranging from $3 - 6 \text{ ms}^{-1}$, which on occasions reached up to 9 ms⁻¹. During 146 147 the post-monsoon season, the major wind flow was from the NW to the SW-W sector. These wind speeds were 148 lower, ranging from $1 - 3 \text{ ms}^{-1}$ exceeding 6 ms⁻¹ only occasionally. Overall, the site received air from all wind sectors in both seasons. This is also borne by the back trajectory analyses presented in the companion paper 149 150 (Awasthi et al., 2024), which showed that the site is characterized by regional airflow patterns as documented at

- 151 other sites in the Indo-Gangetic Plain (Pawar et al., 2015).
- 152 Fire count data were obtained using the Visible Infrared Imaging Radiometer Suite (VIIRS) 375m thermal
- 153 anomalies/active fire product data from the VIIRS sensor aboard the joint NASA/NOAA Suomi National Polar-
- orbiting Partnership (Suomi NPP) and NOAA-20 satellites for high and normal confidence intervals only.

155 2.2 Measurement of Volatile Organic Compounds using the PTR-TOF-MS 10K

Volatile organic compounds (VOCs) were measured using a new high sensitivity and high mass resolution Proton 156 157 Transfer Reaction Time of Flight Mass Spectrometer (PTR-TOF-MS 10k, model PT10-004 manufactured by 158 Ionicon Analytik GmbH, Austria). While PTR-TOF-MS 8000 series (Tripathi et al., 2022) and PTR-QMS (Sinha 159 et al., 2014) instruments have been previously deployed in India and have mass resolutions of 8000 and 1, 160 respectively, this study marks the first deployment of the PTR-TOF-MS 10K system in India, a system that 161 possesses several unique advantages over the older generation instruments for VOC measurements in polluted 162 and complex emission environments. The first is that this new system is equipped with the extended volatility 163 range technology (Piel et al., 2021), ensuring that even many intermediate volatility range compounds and sticky VOCs can be detected with very fast response times and minimal surface effects. The inlet system of the 164 165 instrument as well as the ionization chamber is fully built into a heated chamber and the inlet capillary is further 166 fed through a heated hose to ensure there are no "cold" spots for condensation. The entire inlet system is made of 167 inert material (e.g. PEEK or siliconert treated steel capillaries to keep surface effects minimal. Additionally, a 7 168 µm siliconert filter just before the drift tube served to minimize clogging/contamination of the system. The second 169 advantage possessed by the PTR-TOF-10K used in this work is the inclusion of an ion booster funnel and hexapole 170 ion guide placed after the drift tube/reaction chamber for improved extraction of ions in a manner that boosts both 171 the mass resolution as well as the sensitivity over its older peers. This helped achieve much higher mass resolution

(> 10000 m/ Δ m), even reaching as high as 15000 m/ Δ m at m/z 330, and detection limits better than 3 ppt for all 172 173 compounds detected in the mass to charge ratio (m/z) 31-330 mass range. These customizations over previously 174 deployed PTR-TOF-MS instruments in Delhi, enabled detection and discovery of several intermediate range-175 volatility compounds (IVOCs) in the gas phase. Other parts of the instrument have already been explained well 176 earlier (Jordan et al., 2009; Graus et al., 2010). During this study, the instrument was operated at a drift tube 177 pressure of 3 mbar, drift tube temperature of 120 °C, and drift tube voltage of 600V, resulting in an operating E/N 178 ratio of ~ 120 Td (1 Td = 10⁻¹⁷ V cm⁻²). These operational instrumental settings are also summarized in Table S1. 179 Ambient air was sampled continuously from the rooftop (~35m A.G.L) through a Teflon inlet line that was 180 protected with a Teflon membrane particle filter (0.2 µm pore size, 47 mm diameter) to ensure that dust and debris 181 did not enter the sampling inlet. The length of the inlet line was 5m and made of Teflon (3m 1/8 inch O.D. and 182 2m 1/4 inch O.D). The total inlet residence time was ~ 2.7 seconds. The part of the inlet that was indoors (3m of 183 1/8 inch O.D.) was well insulated and heated to 80 degree Celsius. We think this short inlet residence time and 184 heated inlet facilitated the detection of IVOCs, relative to previous studies. The instrument background was acquired regularly (typically every 30 min for 5 min), by sampling VOC-free zero air. VOC-free zero air was 185 produced by passing air through an activated charcoal scrubber (Supelpure HC, Supelco, Bellemonte, USA) and 186 a VOC scrubber catalyst (Platinum wool) maintained at 370 °C. Mass spectra covering the m/z 15 to m/z 450 187 range were obtained at 1 Hz frequency. An internal standard comprising 1,3-di-iodobenzene ($C_6H_5I_2^+$) detected at 188 189 m/z 330.848 and its fragment ion [C₆H₃I⁺]) detected at m/z 204.943 were constantly injected to ensure accurate 190 mass axis calibration, so that any drifts in the mass scale were corrected providing for accurate peak detection. 191 Primary data acquisition of mass spectra was accomplished using the ioniTOF software (version 4.2; IONICON 192 Analytik Ges.m.b.H., 6020 Innsbruck, Austria). All the settings related to PTR (Proton Transfer Reaction), TPS 193 (TOF power supply), MPV (Multi-port-valve), and MCP (Multi-channel plate) can be controlled and optimized 194 using this control software. The raw mass spectra and relevant instrumental metadata are stored in HDF5 format. 195 These spectra were further processed using the Ionicon Data Analytik (IDA version 2.2.0.4; Ionicon Analytik 196 GmbH, Innsbruck, Austria) software that has the functionalities for peak search, peak fits and preliminary mass 197 assignments and identification of a broad spectrum of organic compounds. The IDA software employs an 198 automated peak detection routine guided by user-defined sensitivity levels for peak detection, peak fit, and shape. 199 The software then uses chemical composition information based on the exact masses and isotopic patterns and 200 calculates a specific proton transfer rate constant (k-rate) based on the polarizability and dipole moment for the 201 peaks with an assigned chemical formula, instead of using a generic value as was done in previous PTR-TOF-MS 202 measurements in Delhi (Tripathi et al., 2022). We manually compared the values also with the compilation of k 203 rates reported by Pagonis et al., (2019) as an additional check. The user has possibility to define a window for 204 mass accuracy (e.g. 30 ppm). Within this defined range and accuracy window, the software identifies all possible 205 chemical compositions and molecular formulae and calculates the corresponding isotope patterns. These patterns 206 are then compared to find the best-fit chemical composition. The process is carried out iteratively, starting with 207 the lower m/z values, according to the method described in the study by Stark et al., (2015). 208 In this study, a total of 1126 peaks were detected in the raw measured ambient mass spectra. After further

additional quality control and assurance steps performed manually as detailed in the Section 3.0, 111 compounds

210 present in ambient air for which the molecular formula could be confirmed unambiguously are reported and for

211 which isotopologues due to molecules of different chemical composition could be ruled out completely, were

212 further analysed in this work. The term "unambiguous" is used in the context of the accurate elemental 213 composition/molecular formula assignment of the ions by leveraging the high mass resolution (8000-13000 over 214 entire dynamic mass range) and detection sensitivity (reaching even 1 ppt or better for many ions; see Table S2) 215 of the instrument. This enabled ensuring peaks due to expected isotopic signals were not construed as new 216 compounds if their height was exactly as expected for a shoulder isotopic peak based on the natural abundance of 217 isotopes of carbon, hydrogen, nitrogen, sulphur, chlorine and oxygen that made up the more abundant molecular 218 ion. Where an ion could occur significantly due to fragmentation of another compound, the same has also been 219 noted in Table S2 during attribution of the compound's name. Figure S1 provides an example of visualization of 220 mass spectra and peak assignment using the IDA software which also illustrate the high mass resolving power of 221 the PTR-ToF-MS 10K, that enables separation of ion signals that differ by less than 0.04 Th, as well the 222 identification of isotopic peaks of parent compounds like methanethiol, dichlorobenzene, C-6 amide and C-9 223 carboxylic acid acid (Fig S2), which are discussed in detail in Section 2.4. Table S2 also provides the limit of 224 detection (LoD) of the compounds as well as the average and interquartile range observed season-wise for each 225 ion. The LoD was calculated by taking the 2σ value of the VOC-free zero air instrument background (Müller et 226 al., 2014). Example of measured data showing the instrumental backgrounds and ambient levels for methanethiol, 227 dichlorobenzene, C-6 amide and C-9 carboxylic acid, over a 3h period are illustrated in Fig S3. A certified VOC 228 calibration gas mixture (Societa Italiana Acetilene E Derviati; S.I.A.D. S.p.A., Italy) containing 11 hydrocarbons 229 at ~100 ppb, namely methanol, acetonitrile, acetone, isoprene, benzene, toluene, xylene, trimethylbenzene, and 230 dichlorobenzene and trichlorobenzene was used during the field deployment for measuring the transmission and sensitivity of compounds covering the mass range (m/z=33 to m/z=181). The instrument was calibrated a total 231 232 of 8 times during the study period: 21.07.2022 after first installation, 26.09.2022, 21.10.2022, 26.10.2022, 233 5.11.2022, 11.11.2022, 16.11.2022 and 30.11.2022. Results were reproducible (~21% or better for all compounds) 234 across all experiments and a transmission curve obtained from one of the calibration experiments is shown in Fig. 235 S4. Measured transmission further allowed for more accurate quantification by accounting for correction of the 236 mass-dependent detection efficiency of the system. Equation S1 (de Gouw et al., 2007) was then used to convert 237 the measured ion signals to mixing ratios. The linearity for compounds available in the VOC standard were also checked independently and was above $r \ge 0.9$ as illustrated in Fig S5 for the tested range of ~ 2 to 8 ppb. The 238 background corrected concentrations of all the detected m/z were exported from IDA in .csv format and further 239 240 analysis of the dataset was carried out using IGOR Pro software (version 6.37; WaveMetrics, Inc.). The overall 241 uncertainty calculated using the root mean square propagation of errors due to the accuracy of gas standard and 242 flow controllers was ~13 % or better for compounds present in the VOC gas standard. For other compounds 243 reported in this work, it is estimated that the combined accuracy of the transmission function and the parameterized 244 k-rates, put the overall uncertainty in the range of $\pm 30\%$ (Reinecke et al., 2024). 245 Carbon monoxide (CO) was measured using IR filter correlation-based spectroscopy air quality analyzer (Thermo 246 Fischer Scientific 48i) while ozone was measured using UV absorption photometry (Model 49i; Thermo Fischer 247 Scientific, Franklin, USA). The overall uncertainty of the measurements was less than 6%. Details concerning

- 248 characterization of the instrument including calibration and data QA/QC protocols have been comprehensively
- described in our previous works (Chandra and Sinha, 2016; Kumar et al., 2016; Sinha et al., 2014).

251 2.3 Mass assignment and compound identification

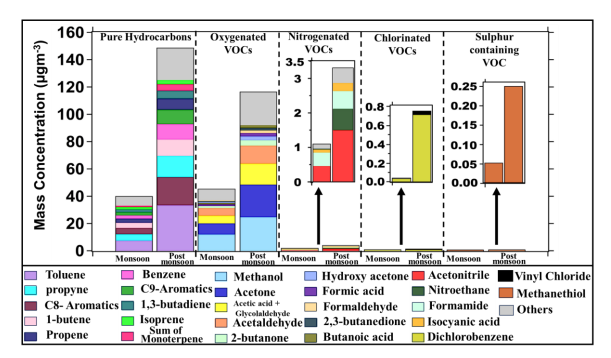
252 A total of 1126 peaks were detected in the raw mass spectra. To identify the ambient compounds of relevance in 253 Delhi from these detected peaks, the following additional manual quality control checks were undertaken. First, 254 peaks attributed to non-ambient compounds such as the impurity ions (e.g. NO⁺), water cluster ion peaks, and 255 peaks associated with internal standards were excluded resulting in 1025 peaks for further consideration. Next, 256 the diel profiles and detection limits of these 1025 ion peaks were perused. Only 319 ions out of the 1025 ions 257 showed some diurnal variability and had values above the detection limit after accounting for the respective 258 instrumental background. Next, we verified the presence and expected theoretical magnitude of the shoulder 259 isotopic peaks based on the natural isotopic distribution abundance of the elemental composition of the ion. Fig 260 S6 provides a visual example. This was feasible for all m/z except the C1 oxygen containing analyte ions, where 261 the shoulder peak was below detection limit. The preceding QA/QC resulted in an unambiguous assignment for 262 111 of the 319 ions. Note that these 111 explained 86% of the total mass concentration (μ gm⁻³) observed due to 263 the 319 detected peaks when accounting for the isotopic peaks as well. Table S2 lists the ion m/z and molecular 264 formula of the corresponding compound, along with the averaged mixing ratios observed in each case during the 265 monsoon and post-monsoon season. Additionally, the characteristic ambient diel profile classification as one of 266 the following: unimodal with daytime peak for biogenic/ evaporative/ photochemical source emitted compounds, 267 bimodal with morning and evening peaks for compounds driven by primary emissions (e.g. toluene) and trimodal 268 which were hybrid of the former two, are also provided for each species. Compound names were attributed to 269 specific ions using assignments reported at that m/z in the compiled peer-reviewed PTR-MS mass libraries 270 published by Yáñez-Serrano et al., (2021) and Pagonis et. al., (2019) as well as previously published pioneering 271 reports by Stockwell et. al. (2015), Sarkar et al. (2016), Yuan et al. (2017) and Hatch et al. (2017).

272 Fragmentation of certain compounds in specific atmospheric environments can cause significant interferences in 273 the detection of major compounds like isoprene, acetaldehyde and benzene, as reported recently by Coggon et al., 274 2024. We checked these as well as an additional quality control measure. As noted by Coggon et al. 2024, isoprene 275 can suffer significant interferences from higher aldehydes as well as substituted cyclohexanes, which can fragment 276 and add to the signal at m/z 69.067 (at which protonated isoprene $C_5H_9^+$ is also detected). The magnitude depends 277 on the instrument operating conditions (Townsend ratio), instrument design and the mixture of VOCs present in 278 ambient air while co-sampling isoprene. Coggon et al. 2024 very nicely clarified both these aspects and found that 279 when influenced by cooking emissions and oil and natural gas emissions and at higher Townsend ratios, these 280 interferences can be quite significant and even account for upto 50% of the measured signal attributed to isoprene 281 in extreme cases. We operated the PTR-TOF-MS at 120 Td which minimizes fragmentation even if it occurs, 282 compared to when operated at 135-140 Td. Concerning the ambient VOC mixture and emission sources, we note 283 that the type of restaurant cooking emissions present in Las Vegas and over Oil and Natural Gas petrochemical 284 facilities in USA for which Coggon et al. 2024 reported the highest isoprene interferences, were absent/negligible 285 at the study site in Delhi. In the latter, open biomass burning sources such as paddy residue burning in post-286 monsoon season and garbage biomass fires and traffic that occur throughout the year are more significant. Use of 287 more specific though slower analytical techniques based on gas chromatography show that such biomass 288 combustion sources emit significant amounts of isoprene (Andrea et al., 2019; Kumar et al., 2021). The above 289 points and supporting TD-GC-FID measurements of isoprene, benzene and toluene (see Fig S7 and Shabin et al., 290 2024), led us to conclude that such correction is unwarranted for our PTR-TOF-MS dataset. Concerning the

- interference on acetaldehyde detection due to ethanol, we note that even in Coggon et al. 2024 this was reported
- to only be of significance in highly concentrated ethanol plumes such as those encountered on the Las-Vegas strip
- 293 where ~1500 ppb of ethanol was detected. On the contrary, in Delhi as listed in Table S2, ethanol values detected
- at m/z 47.076 were on average only 0.2 ppb (Interquartile range 0.16 ppb) during monsoon and 0.55 ppb
- 295 (Interquartile range 0.5 ppb) in post-monsoon season, respectively, whereas acetaldehyde detected at m/z 45.03
- was significantly higher at 3.34 and 7.75 ppb during monsoon and post-monsoon season, respectively.
- For the same molecular formula, several isomeric compounds with differing chemical structures are possible, with the number of possibilities increasing enormously with an increase in the number of atoms that make up the molecule. In addition, in some instances fragmentation of other compounds can complicate the compound attribution for a given ion. Nonetheless in the interest of stimulating interest and further investigation as many have been previously rarely reported or are being reported for the first time in ambient air, we have made bold to provide one of the many possible chemical structures in the Table S2. We do caution that the chemical structure provided by no means even constitutes a best guess estimate but nonetheless would be appealing to chemists and
- 304 provoke further detailed reporting rather than just the molecular formula.
- 305

306 3. Result and Discussion:

- 307 3.1: Analyses of ambient mass spectra and mass concentration contributions of VOC chemical classes
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Figure 2: Bar graph of 111 compounds class-wise, namely Pure Hydrocarbons, Oxygenated VOCs (OVOCs),
 Nitrogen-containing VOCs (NVOCs), Chlorine-containing VOCs (CIVOCs), and sulphur-containing VOC (SVOC) in

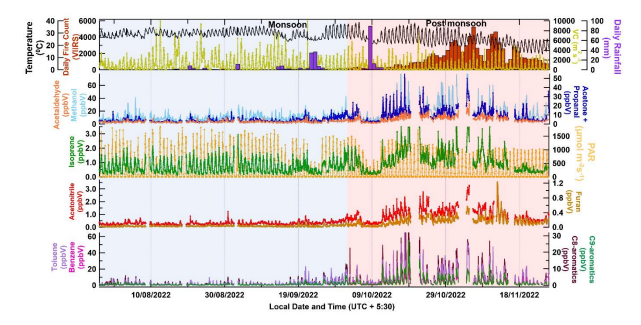
312 both monsoon and post-monsoon periods.

- 313
- 314 A summary of the distribution of the 111 compounds in terms of chemical classes showing their averaged
- measured ambient mass concentration (μ gm⁻³) contributions is shown in Fig. 2 for the monsoon (22^{nd} July 30^{th}

316 September 2022) and post-monsoon seasons (1 October- 26 November 2022). Out of the 111 compounds, 42 were 317 pure hydrocarbons made up only of carbon and hydrogen atoms, 56 were oxygenated volatile organic compounds 318 (OVOCs) made up of only carbon, hydrogen and oxygen, 10 contained nitrogen (NVOCs), 2 contained chlorine 319 (CIVOCs), and 1 contained sulphur (SVOC). The average total mass concentration of the same set of pure 320 hydrocarbons during post-monsoon season was 3.7 times greater than in monsoon season (40 µgm⁻³ vs 148 µgm⁻¹ 321 ³) while the average total mass concentration of OVOCs during post-monsoon was 2.6 times greater than the 322 monsoon season values (44 µgm⁻³). Pure hydrocarbons and OVOCs contributed similarly to the 323 mass concentrations in monsoon season but during the post-monsoon season, the contribution of pure 324 hydrocarbons was significantly higher than that of OVOCs, due to an increase in primary emissions of these 325 compounds. The average mass concentration of NVOCs during post-monsoon was thrice as high relative to the 326 monsoon season (1 µgm⁻³ and 3 µgm⁻³). For the chlorine containing VOCs the post-monsoon, concentrations were 327 20 times higher, though in absolute magnitude, the values were low (1 μ gm⁻³). The average mass concentration of 328 sulphur containing VOCs during post-monsoon was 4 times higher, but again absolute values were low (0.2 µgm⁻ 329 ³). The top 10 pure hydrocarbon compounds by mass concentration ranking were toluene, sum of C8-aromatics 330 (xylene and ethylbenzene isomers), propyne, 1-butene, benzene, sum of C9-aromatics (trimethyl benzene 331 isomers), propene, sum of monoterpenes, isoprene and 1,3 butadiene and contributed to 84% of the total mass 332 concentration due to pure hydrocarbons during both the monsoon and post-monsoon seasons, respectively, while 333 the top 20 contributed to 95% and 96% of the total mass concentration in monsoon and post-monsoon, 334 respectively. The top 10 OVOCs: methanol, acetone, acetic acid+ glycolaldehyde, acetaldehyde, hydroxyl-335 acetone, formaldehyde, 2-butanone, 2,3-butanedione, formic acid, butanoic acid collectively contributed to 84% 336 and 79% of the total mass concentration due to all OVOCs in monsoon and post-monsoon, respectively, while the 337 top 20 contributed to 93% and 90% of the total mass concentration in monsoon and post-monsoon, respectively. 338 The top 4 NVOCs namely acetonitrile, nitroethane, formamide and isocyanic acid contributed to 92% and 91% 339 of the total mass concentration in monsoon and post-monsoon, respectively. Out of 2 identified chlorine containing 340 VOCs, dichlorobenzene (C₆H₄Cl₂) was found to be the major contributor contributing 87% and 95% of the total 341 mass concentration in monsoon and post-monsoon, respectively. The only sulphur containing VOC was 342 methanethiol [CH₄S] detected at its protonated ion m/z 49.007 and confirmed by the shoulder isotopic peak. 343 Overall, there was an increase in the mass concentration of all the classes of VOCs from monsoon to post-344 monsoon. This increase in mass concentration could be attributed to increased emissions from sources that get 345 active in post-monsoon, such as regional post-harvest paddy residue burning, increased open waste burning as 346 well reduced wet scavenging and ventilation coefficient compared to the monsoon season. We examine these in 347 more detail in the next sections.

348 3.2: Time series of VOC tracers during the "clean" monsoon and "polluted post-monsoon" seasons in Delhi

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Figure 3: Time series of hourly data for meteorological parameters like temperature (C) and ventilation coefficient (m²s⁻¹), daily rainfall and daily fire counts (top panel); hourly mixing ratios of methanol, acetaldehyde, and the sum of acetone and propanol (second panel from top); isoprene and PAR (µmolm⁻²s⁻¹) (third panel); acetonitrile and furan (second panel from bottom); and benzene, toluene and the sum of C8 – aromatics (xylene and ethylbenzene isomers) and the sum of C9 – aromatics (isomers of trimethyl benzene and propyl benzene) (bottom panel). The blue and red shaded regions represent the monsoon and post-monsoon periods, respectively.

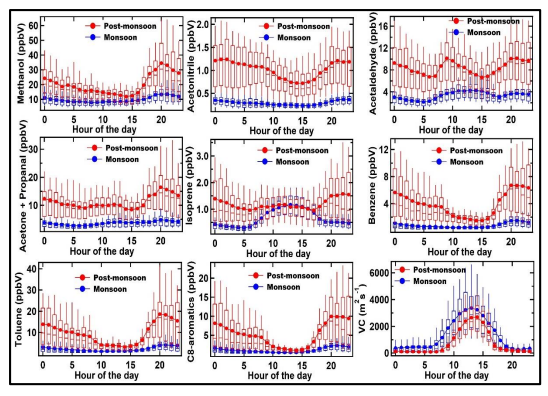
359

360 Figure 3 shows the time series plot of meteorological parameters and the mixing ratios of some key VOC tracer molecules during monsoon (22nd July – 30th September 2022, blue-shaded region) and post-monsoon (1st October 361 -26^{th} November 2022, red-shaded region). The top panel shows the ambient Temperature (0 C), daily VIIRS fire 362 counts on the left side of the top panel and ventilation coefficient (m²s⁻¹), and daily rainfall (mm) on the right side 363 of the top panel during the study period (22^{nd} July $2022 - 26^{th}$ November 2022). A grid (1km × 1km) with latitudes 364 365 between 21°N and 32°N and longitudes between 78°E and 88°E was considered for extracting the fire count data. 366 The second panel from the top represents the time series of mixing ratios of OVOCs which can be formed photochemically as well as be emitted from anthropogenic sources, namely methanol, acetaldehyde, and the sum of 367 368 acetone and propanol; the third panel shows the mixing ratio of isoprene (a daytime biogenic chemical tracer, pure hydrocarbon) and photosynthetic active radiation (PAR) (μ mol photons m⁻² s⁻¹), and the fourth panel shows the 369 370 mixing ratio of acetonitrile (a biomass burning chemical tracer) and furan (a combustion chemical tracer). The 371 bottom panel shows the mixing ratios of benzene, toluene, the sum of C8-aromatics (xylene and ethylbenzene 372 isomers), and the sum of C9-aromatics (trimethylbenzene and propyl benzene isomers). These are some of the 373 most abundant VOCs typically present in any urban megacity environment, due to their strong emission from 374 traffic and industries in addition to biomass burning (Sarkar et al., 2016; Sinha et al., 2014; Chandra et al., 2016; 375 Singh et al., 2023; Dolgorouky et al., 2012; Yoshino et al; 2012; Langford et al., 2010). We note that all the 376 meteorological conditions and fire activity and VOC levels changed significantly between the much "cleaner" 377 monsoon season and "highly polluted" post-monsoon season at the same site. While the average temperature 378 during monsoon season was 29.5±2.8 °C, in the post-monsoon season this changed to 24.8±5.2 °C, while the 379 average ventilation co-efficient was 1.7 times higher during monsoon season relative to the post-monsoon season.

- **380** Except for the period impacted by heavy rainfall due to western disturbance weather (8th Oct 10th Oct 2022),
- the average mixing ratios for all compounds were considerably higher in the post-monsoon season relative to the
- 382 monsoon season even after accounting for the ventilation coefficient reduction with all the aromatics compounds
- **383** like benzene, toluene, sum of C8 and C9 aromatics, all 4.5 times higher and furan more than 5 times higher and
- acetonitrile, acetone more than 3 times higher and methanol and acetaldehyde 2 times higher. Even isoprene was
- 385 1.7 times higher but its night time mixing ratios were higher than daytime mixing ratios during post-monsoon
- 386 season relative to the monsoon season. The increases clearly exceed what can be accounted for only by the reduced
- 387 ventilation co-efficient (seasonality) and suggests an increase in anthropogenic combustion related sources in
- 388 particular from open biomass burning fire sources, which we investigate in more detail in the subsequent sections.

3.3: Analyses of the diel profiles during the "clean" monsoon and "polluted post-monsoon" seasons in Delhi for discerning major drivers of their ambient values

- 391 Figure 4 shows the diel box and whiskers plot depicting the average, median, and variability (10th, 25th, 75th and 90th percentile) of the same key VOCs like methanol, acetonitrile, acetaldehyde, acetone and propanal, furan, 392 393 isoprene, benzene, toluene and C8 - aromatics for monsoon (derived from ~ 1704 data points, blue markers) and 394 post-monsoon (derived from \sim 1368 data points, red markers) against the hour of the day (the horizontal axis 395 represents the start time of the corresponding hourly bin). This more clearly brings out the season-wise diel 396 variation of the compounds and in turn throws light on the emission characteristics and how they vary for the 397 same compound between seasons. Both in the monsoon and post-monsoon season, methanol mixing ratios seem 398 to be driven by primary emission sources and correlate very well with toluene, a tracer for traffic emissions, with 399 highest increases in the evening hours (17:00 to 20:00 L.T.). Globally the main source of methanol is vegetation 400 but in a megacity like Delhi that possesses more than 150000 compressed natural gas (CNG) vehicles and light 401 duty diesel vehicles, it appears that traffic (see Fig 1 of (Hakkim et al., 2021) emitted methanol controls its ambient 402 abundance. Similarly, based on the correlation with toluene, traffic emissions seem to be a major contributor for 403 acetaldehyde, acetone, sum of C8-aromatics and benzene in the morning and evening hours. All these compounds 404 are among the most abundant VOCs detected in tailpipe exhaust samples (Hakkim et al., 2021). Average ambient 405 mixing ratios of acetonitrile, a compound emitted significantly from biomass burning (Holzinger et al., 1999), 406 were below 0.5 ppb in the monsoon for all hours, with only slight increase at night, but during post-monsoon 407 season, for all hours the values doubled to 1 ppb, with strong increases in the early evening and night time hours. 408 This tendency was mirrored in all the other compounds including isoprene. The diel profile of isoprene and 409 acetaldehyde were the only ones which showed daytime maxima during the monsoon season. 410 This shows that during the monsoon season, the biogenic sources of isoprene majorly drive its ambient mixing
- 411 ratios, whereas acetaldehyde ambient mixing ratios are controlled by photochemical production of the compound
- in the monsoon season. Under the high NOx conditions prevalent in a megacity like Delhi, photo-oxidation of n-
- butane, propene, ethane and propane could be a large photochemical source of acetaldehyde (Millet et al., 2010).
- 414



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Figure 4: Box and whisker plots showing average, median, and variability (10th, 25th, 75th and 90th percentile) for
some major VOCs and the ventilation coefficients (m²s⁻¹) (VC) during monsoon and post-monsoon periods. The blue
and red markers represent the monsoon and post-monsoon periods, respectively.

420 Benzene which is human carcinogen is the only VOC for which there is a national ambient air quality standard (5 421 μ g m⁻³ equivalent to ~1.6 ppb at 298 K) in India. Average mixing ratios in the post-monsoon season (Fig 4) were 422 always above this value no matter what hour of the day, and the seasonal average was twice as high as this value 423 (~4 ppb). The increased biomass burning in post-monsoon season controlled the abundance of benzene, 424 acetaldehyde and acetone and isoprene during this period, due to strong emissions from both biomass burning and 425 traffic. The typical atmospheric lifetimes of all these compounds spans from few hours (e.g. isoprene) to several 426 days (e.g. benzene and methanol) and several months in the case of acetonitrile. The results of the TD-GC-FID 427 measurements along with the average PTR-TOF-MS values presented in Figure 4, are summarized in Figure S7. 428 Even though the TD-GC-FID measurements present only a snapshot as the ambient sampling duration is shorter, 429 the season-wise diel profiles are consistent with those obtained using the PTR-TOF-MS and the average mixing 430 ratios obtained using the PTR-TOF-MS dataset also fall well within the range of mixing ratios observed using the 431 TD-GC-FID. This provides further confidence in the high night-time isoprene observed during the post-monsoon 432 season. The isoprene emissions at night during the post-monsoon season are likely due to combustion sources. 433 Paddy residue burning and dung burning have the highest isoprene emission factors of ~0.2 g/kg (Andreae 2019) 434 and more than 8 Gg of isoprene is released in the space of a few weeks during the post-monsoon season regionally 435 from open paddy residue burning alone (Kumar et al., 2021). Previous studies from the region have also 436 documented isoprene emissions from non-biogenic sources, which are active also at night (Kumar et al., 2020, 437 Hakkim et al., 2021). In 2018 at another site in Delhi, using gas chromatography measurements made in pre-438 monsoon and post-monsoon, Bryant et al. 2023 reported average nocturnal mixing ratios of isoprene that were 5 439 times higher in the post-monsoon compared to the pre-monsoon and showed different diel profiles between the

- 440 seasons. They found that the high night-time isoprene correlated well with carbon monoxide, a combustion tracer 441 and suspected that in addition to the stagnant meteorological conditions, biomass burning sources could be a 442 reason for the significant night time isoprene in Delhi in post-monsoon season and our findings using more 443 comprehensive and high temporal resolution data further substantiate the surprising night-time isoprene.
- 444 As potent precursors of secondary organic aerosol, the aromatic compounds would also enhance secondary
- 445 organic aerosol pollutant formation during the polluted post-monsoon season. When compared with the first PTR-
- 446 MS measurements of these compounds reported from wintertime Delhi (see Fig 2 of Hakkim et al., 2019), the
- 447 average levels of these compounds for the post-monsoon season (Table S2) are lower or comparable, but still
- 448 significantly higher than what has been reported for other major cities of the world like Tokyo, Paris, Kathmandu,
- 449 Beijing, London (Yoshino et al., 2012; Dolgorouky et al., 2012; Sarkar et al., 2016; Li et al., 2019; Langford et
- 450 al., 2010). The monsoon levels on the other hand were comparable to many of the other megacities.
- 451 As the monsoon season is characterized by favourable meteorological conditions for wet scavenging and dispersal
- 452 due to higher ventilation co-efficient, as well as significantly lower open biomass burning due to wet and warm
- 453 conditions, the monsoon levels can be considered as baseline values for the ambient levels of these compounds
- 454 (except isoprene and acetaldehyde) in Delhi, which are driven mainly by year-round traffic and industrial sources
- 455 in Delhi. In monsoon for isoprene, the major driver are biogenic sources whereas for acetaldehyde the major driver
- 456 is photochemistry, a finding that is similar to what has been reported from another site in the Indo-Gangetic Plain
- 457 previously (Mishra and Sinha, 2020).

458 3.4: Discovery of methanethiol (CH₃SH), dichlorobenzenes (C₆H₄Cl₂), and C6-amides (C₆H₁₃NO₂) and C9-459 organic acids (C₉H₁₈O₂) in ambient Delhi air

- 460 Figure 5 shows the average diel profile of four compounds present in both monsoon and post-monsoon periods
- that have to our knowledge never been reported from Delhi or any site in South Asia and only rarely been reported
- in the gas phase in any atmospheric environment in the world. Except for methanethiol detected at m/z 49.007
- 463 (also called methyl mercaptan), all the other compounds namely dichlorobenzene ($C_6H_4Cl_2$) detected at at m/z
- 464 146.977, C6-amides like hexanamide ($C_6H_{13}NO_2$) and its isomers detected at m/z 116.108 and C9- carboxylic
- 465 acid/ester such as nonanoic acid ($C_9H_{18}O_2$) and its isomers detected at m/z 159.14, are all intermediate volatility
- 466 range organic compounds. The saturation mass concentration (C₀) of methanethiol, C6-amide, dichlorobenzene,
- 467 and C9 organic acid, were calculated using the method described in Li et al. 2016 using the following equation:

468
$$C_0 = \frac{M \, 10^6 \, p_0}{760 \, R \, T}$$
 (1)

- 469 wherein M is the molar mass [g mol⁻¹], R is the ideal gas constant [8.205 x 10⁻⁵ atm K⁻¹ mol⁻¹ m³], p₀ is the 470 saturation vapor pressure [mm Hg], and T is the temperature (K). Organic compounds with $C_0 > 3 \times 10^6 \,\mu g \,m^{-3}$ 471 are classified as VOCs while compounds with $300 < C_0 < 3 \times 10^6 \,\mu g \,m^{-3}$ as Intermediate VOCs (IVOCs).
- 472 The presence of such reactive organic sulphur, chlorine and nitrogen containing compounds in the gas phase
- 473 provides new insights concerning the chemical composition and secondary chemistry occurring in air, during the
- 474 extremely high pollution events. Below we examine the sources and chemistry of these compounds in further
- 475 detail.
- 476 The diel profiles of both methanethiol and dichlorobenzene in both the monsoon and post-monsoon seasons were
- 477 similar (bimodal with afternoon minima), and controlled by the ventilation coefficient diel variability (see Fig. 4),
- 478 and in fact even the difference in their average magnitudes (50 ppt Vs 130 ppt for CH₃SH and 25 ppt Vs 100 ppt

- for dichlorobenzene between monsoon and post-monsoon seasons), can largely be explained by the reduction in ventilation co-efficient (~2 reduction). Further, the conditional probability wind rose plots for both compounds shows that the high values come from the same wind sector upwind of the site spanning north-east to south during early morning and evening hours, which is actually where a variety of industrial sources are located. Previously, Nunes et al., (2005) and Kim et al., (2006) have reported methanethiol from petrochemical industries and landfills in Brazil and Korea, respectively. Toda et al., (2010) reported high (tens of ppb) methanethiol mixing ratios from a pulp and paper mill industry in Russia.
- 486
- 487

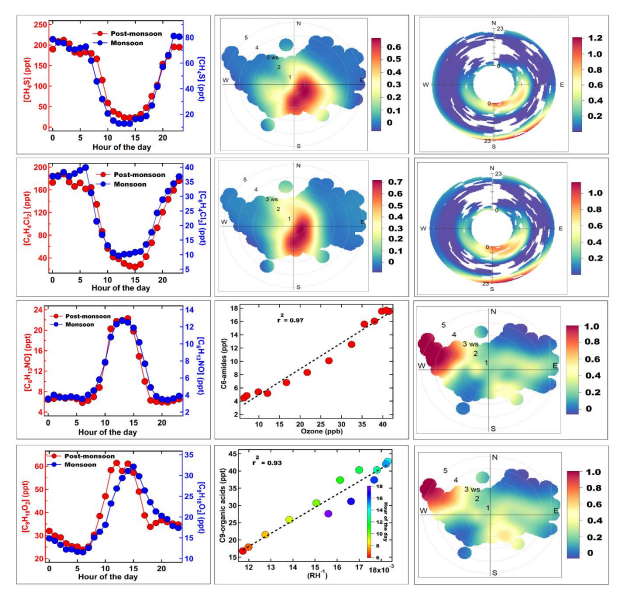




Figure 5: Average diurnal profile of methanethiol, isomers of dichlorobenzene, C6-amides, and C9- organic acid in the left panel for both monsoon (blue marker) and post-monsoon (red marker) periods. The second panel shows the wind rose plot of methanethiol and isomers of dichlorobenzene, plot of C6-amide vs ozone and C9-organic acid vs RH⁻¹ colour coded by the hour of the day. The third panel shows the polar annulus plot of methanethiol, isomers of dichlorobenzene and wind rose plot of C6-amides and C9-organic acid.

- 494
- 495 Both compounds are also used in the deodorant and pesticide products as reagents (Chin et al., 2013) and although 496 large scale pesticide manufacturing facilities were shifted out of Delhi, there are still units that sell and distribute

- 497 these products in those areas, from which fugitive emissions are likely happening. Methanethiol is further used as 498 a precursor in methionine production (Francois, 2023) an essential amino acid used in manufacture of pesticides, 499 and fragrances industry uses methanethiol for its distinct sulphur-like aroma (Bentley et al., 2004), contributing 500 to the creation of savory flavors and unique fragrances. In the Delhi environment, a combination of such industries, 501 in particular paper and pulp industries, are likely candidate sources. Figure 5 confirms that elevated methanethiol 502 values in the windrose had a clear directional dependence from the area spanning the north to south east sector. 503 This is the region where various manufacturing facilities and industrial areas of Delhi like Patpargani (north-east) 504 Okhla, Faridabad (south) are situated and these industrial estates were earlier marked in Figure 1 b. Methanethiol 505 is an extremely reactive molecule reacting primarily with the hydroxyl radical (OH) during daytime with an 506 estimated lifetime of 4.3 h (Wine et al., 1981). Its photo-oxidation in daytime with hydroxyl radicals produces 507 sulphur dioxide, methanesulfonic acid, dimethyl disulphide and sulphuric acid (Kadota and Ishida et al., 1972; 508 Hatakeyama et al., 1983), all of which play key roles in aerosol formation pathways. Dimethyl disulphide has a 509 very short atmospheric lifetime spanning from 0.3 to 3 hours (Hearn et al., 1990), because of its high reactivity $([1.98 \pm 0.18] \times 10^{-10} \text{ cm}^3 \text{ molecule}^{-1}\text{s}^{-1})$ (Wine et al., 1981) with OH radicals. Although dimethydisulphide is the 510 511 major product of the photo-oxidation of methanethiol (yield 50%; Wine et al., 1981), since methanethiol itself 512 was on average only 48 ppt (monsoon) and 128 ppt (post-monsoon), and plumes occur only at night we 513 hypothesize that the ambient concentrations of DMDS were too low to be detected by the mass spectrometer. 514 Further it can also react with nitrate radicals (Berreshiem et al., 1995) and participate in night-time chemistry. 515 More recently, Reed et al. (2020) performed laboratory experiments and observed that even trace amounts of 516 organosulphur compounds, such as thiols and sulfides, can significantly enhance the organic aerosol mass 517 concentration and its particle effective density. Though there has not been any relevant data set attributing the 518 enhancement of organic aerosols to methanethiol in Delhi specifically, previous studies have found enhanced 519 secondary aerosol formation rates during haze and fog episodes (Acharja et al., 2022). These studies collectively 520 suggest an increase in the haze events in Delhi is linked to sulphur chemistry in which methanethiol due to its 521 high reactivity and atmospheric chemistry could also be a contributor along with ammonia and other sulphur 522 containing molecules. Wine et al. (1981) had further predicted that the very rapid rate at which methanethiol reacts 523 with OH would result in low steady-state concentrations in ambient air, even though reasonably large-scale 524 sources may exist.
- 525 Several recent studies have reported high chloride in sub-micron aerosol of Delhi (Gani et al., 2020; Acharja et 526 al., 2023; Pawar et al., 2023). Dichlorobenzene is an intermediate range volatile organic compound (IVOC) which 527 can partition between gas and aerosol phase. However, till date no gaseous IVOC chlorinated organic compound 528 have been reported in ambient air from India. p-dichlorobenzene (PDCB) also called 1. 4-dichlorobenzene, one 529 of the dichlorobenzene isomers is known for its use as a pest repellent and deodorant in indoor environments. 1,4-530 dichlorobenzene in outdoor air in various locations of North America and Europe ranged from 30 ppt to 830 ppt 531 (Chin et al., 2013). It is emitted only from anthropogenic sources as there are no known natural sources. Its 532 emission sources include consumer and commercial products containing PDCB, waste sites, and manufacturing 533 facilities for flavour and as insect repellent products (ATSDR. 2006). Its atmospheric lifetime is estimated to be 534 21-45 days (Mackay et al., 1997). It has been reported as a precursor of secondary organic aerosol in indoor conditions (Komae et al., 2020). Due to its long lifetime, dichlorobenzene can be transported to upper regions of 535 536 the atmosphere where some release of some reactive chlorine through photolysis can occur, but this is not likely

- 537 to be of large consequence. Instead, reaction with hydroxyl radicals would convert it more readily to phenolic
- 538 compounds that would readily partition to aqueous aerosol phase and also undergo nitration to form nitrophenolics

539 (Hu et al., 2021), which are a component of brown carbon (Lin et al. 2015, 2017).

- 540 In contrast, the diel profile of the average mixing ratios of $C_6H_{13}NO$ (Fig. 5), likely hexanamide or isomers of C6-541 amides measured at m/z 116.108, was similar in both monsoon and post-monsoon season and characteristic of a 542 compound with a purely photochemical source with no evening time peaks even during the enhanced biomass 543 burning in post-monsoon season. As observed for several other compounds in this study, the difference in 544 magnitude between both seasons (peak value 22 ppt in post-monsoon season vs 12 ppt in monsoon season) could 545 be accounted for almost completely by the reduced ventilation co-efficient in post-monsoon season (factor of \sim 2). 546 The presence of photochemically formed formamide and acetamide from OH oxidation of alkyl amine precursors 547 has been previously reported (Chandra et al., 2016; Kumar et al., 2018), from another site in the Indo-Gangetic 548 Plain which experiences strong agricultural waste burning. In the literature we could only find only one report for 549 presence of C6 amides in the ambient air in the gas phase (Yao et al., 2016), who reported ~14 ppt in summertime 550 air of Shanghai using an ethanol reagent ion CIMS, the source of which was both industrial and photochemical 551 origin. However, to our knowledge this is the first study world-wide to detect and report only photo-chemically 552 formed C6-amides in the gas phase. C6-amides are IVOCs, which can easily partition to aerosol phase depending 553 on environmental conditions and also act as a new source of reactive organic nitrogen to the atmospheric 554 environment. We found the highest values in air masses arriving in the afternoon from the north-west direction at 555 high wind speeds (see Fig 5) during the post-monsoon season, which indicated that paddy stubble burning emissions of amines (Kumar et al., 2018) were its likely precursors. The mechanism of amide formation through 556 557 photochemical reactions has been elucidated in several previous laboratory studies (Bunkan et al., 2016, Barnes et al., 2010; Nielsen et al., 2012; Borduas et al., 2015). When correlated with daytime ozone hourly mixing ratios, 558 559 the very high correlation (r^{2} > 0.97), confirmed its purely photochemical origin. Being an amide, further gas phase 560 oxidation products are likely to result in organic acids or condensation on existing aerosol particles which could 561 add to the reactive organic nitrogen in aerosol phase and neutralize acidity just like ammonia, as ammonium ion 562 is formed from hydrolysis of amides (Yao et al., 2016). However, the exact role of these amides in nucleation and 563 aerosol chemistry will warrant further investigations.
- Finally, the last row of Fig. 5 shows the average mixing ratios of the compound with molecular formula $C_9H_{18}O_2$ which is likely due to isomers of C9- carboxylic acids (e.g. nonanoic acid), although one cannot rule out contributions from isomers of esters such as methyl octanoate or 2-methylbutyl isobutyrate also detected at m/z 159.14. Hartungen et al. (2004) and more recently the insightful study by Salvador et al. (2022), have highlighted that carboxylic acids (RCOOH) can undergo dissociation reactions within the drift tube in addition to protonation, and form acylium ions as per the following reaction below (Hartungen et al., 2004):

570
$$H_3O^+ + RCOOH \rightarrow RCOOHH^+$$
 (protonated ion) + H_2O (R1)

571 $H_3O^+ + RCOOH \rightarrow RCO^+ (acylium ion) + 2 H_2O$

572 We detected the corresponding acylium ion of C9-carboxylic acid ($C_8H_{17}CO^+$ detected at m/z 141.13) in the

- 573 measured ambient spectra (Figure S9) and found that not only was it present but that it also correlated well in the
- ambient data with the protonated ion (r=0.83). The presence of the fragment ion and its correlation, provides
- additional confirmation concerning the attribution of m/z 159.14 to the C9-organic acid and for quantification the

)

(R2)

ion signals due to the protonated and acylium ions, were summed together for greater accuracy. Although here 576 577 also there is a daytime peak, the timing of the peak is much later in the day (15:00 local time). The peak hourly 578 values reached 60 ppt in post-monsoon season. It showed high correlation ($r^{2} > 0.93$) with the inverse of the 579 ambient daytime relative humidity indicating that it partitions back and forth between the gas phase and aerosol 580 phase depending on the environmental conditions of temperature and RH. n-alkanoic acids in general and 581 nonanoic acid in particular have long been reported as major organic acids present in biomass burning emitted organic aerosol (Oros et al., 2006; Fang et al., 1999). The corresponding wind rose plot (Fig. 5) shows that the 582 583 highest values were in air masses arriving at high wind speeds in the afternoon from the north-west during post-584 monsoon season, which is a major source region of biomass burning emitted organic aerosols. It is also possible 585 that photochemical oxidation through ozonolysis of precursors and hydroxyl radical initiated oxidation can form 586 such carboxylic acids as an advanced oxidation product (Kawamura et al., 2013). In both cases, biomass burning 587 emissions and evaporation from aerosol phase, appear to be the major source of this compound. Carboxylic acids 588 in the aerosol phase would serve to neutralize some of the excess ammonia in the atmospheric environment of the 589 Indo-Gangetic Plain (Acharja et al., 2022) and would be important for night-time aerosol chemistry in Delhi.

590 3.5: Comparison of ambient mixing ratios and VOC/CO emission ratios for aromatic VOCs in Delhi with 591 some megacities of Asia, Europe and North America

592 Aromatic compounds are among the most important class of compounds in urban environments due to their

593 direct health effects (e.g. benzene is a human carcinogen), and reactivity as ozone and secondary organic aerosol

- 594 precursors. Therefore, these compounds have been widely investigated in many cities and information
- 595 concerning their ambient levels and emission ratios to carbon monoxide is often used for assessing similarities

and differences in the sources of these compounds in varied urban environments (Warneke et al., 2007' Borbon

- et. al., 2013). In Figure 6, we show the emission ratios (ER) derived for benzene, toluene and the sum of C8 and
- C9 aromatic compounds (VOC / CO ppb/ppm) using night-time monsoon (left panel) and post monsoon (right
 panel) measurements made in Delhi. The method is based on a linear regression fit to determine the slope of the
- night-time scatterplot data (from 20:00 to 06:00 L.T.) between a VOC (ppb) and CO (ppm) (de Gouw et. al.,
- 601 2017, Borbon et. al., 2013). Using night-time hourly data (18:00 to 06:00 L.T.) provides the advantage of
- 602 minimizing complications due to daytime oxidative losses of the compounds. It can be noted from Fig. 6, that
- during the monsoon season (from 18:00 to 23:00 and 00:00 to 06:00 local time) and post-monsoon season
- 604 (18:00 to 23:00), the observed emission ratios as inferred from the slopes and fits are not statistically different
- from each other (all highlighted by oval circles) with values for benzene/CO, toluene/CO, sum of C8-aromatics,
- sum of C9-aromatics/CO in the range of 1.2-2.43, 3.14-6.76, 1.97-3.84, and 1.05-2.07, respectively. All these
 emission ratios fall within the range of what has been reported for typical petrol 2 and 4 wheeler vehicles in
- 607 emission ratios fall within the range of what has been reported for typical petrol 2 and 4 wheeler vehicles in608 India in tail pipe emissions (Hakkim et al., 2021). For the monsoon season, although two linear fits are observed

from 18:00 to 23:00 and 00:00 to 06:00, the values of the emission ratios as inferred from the respective slopes

- 610 for all compounds overlap or are very close to each other and within the uncertainties for all compounds. We
- 611 hypothesize that the two fits are due to the change in relative numbers of 2 wheelers and 4 wheelers. In the post-
- monsoon season however, for the time period in the second half of the night (00:00-06:00), the emission ratios
- 613 derived from the slopes are statistically different from the ones observed in monsoon season and the first half of
- night in post-monsoon season (18:00-23:00). When we examined the wind rose plots for the same night-time
- data of the aforementioned compounds for each season (Figure S8), we noted that during the post-monsoon

- 616 season more pollution plumes from the south east sector which has industrial facilities and the north west sector
- 617 (a major fetch region for biomass burning plumes from regional paddy residue burning in Punjab and Haryana)
- 618 occurred. During the post-monsoon season due to dip in temperatures at night, the heating demand (Awasthi et
- al., 2024) and associated open biomass burning (Hakkim et al., 2019) also goes up, relative to the monsoon
- 620 period nights. Hence overall we think that these additional sources in the post-monsoon season, do add to the
- burden of these mainly traffic emitted aromatic compounds and could help explain atleast partially the higher
- 622 emission ratios observed during the post-monsoon season (00:00- 06:00), wherein values for benzene/CO,
- toluene/CO, sum of C8-aromatics , sum of C9-aromatics/CO values range from 3.15-3.27, 7.72-8.68, 5.03-5.37,
- 624 2.6-2.76, respectively, and are statistically different from the others (ones marked by oval circles).
- 625

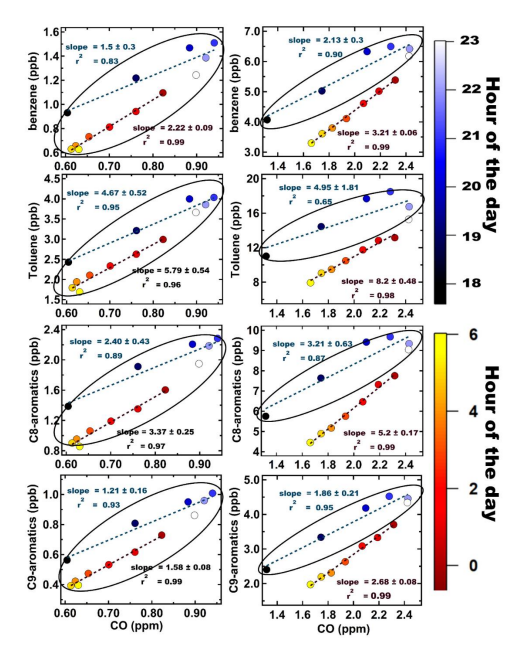


Figure 6: Emission ratios (VOC (ppb)/CO (ppm)) of benzene, toluene, C8 aromatics and C9 aromatics for both
monsoon (left panel) and post monsoon (right panel) periods respectively. The data points for each period are colour
coded with the hour of the day (18:00 L.T to 06:00 L.T).

630 631 Table 1 provides a comparison of the ambient mixing ratios and emission ratios that have been reported in some 632 other major megacities of Asia, Europe and North America for these compounds. Although, the year of 633 measurements and seasons are not the same, nonetheless such comparison helps put the 2022 levels of these 634 compounds in Delhi in a global context. It may be further noted that we took care to calculate the emission ratios 635 using only night-time data when chemical loss of these compounds is negligible as their main oxidation is through 636 OH radicals during daytime, as also noted by de Gouw et al., 2017. Further, the other studies referred to in Table 637 1 for comparison, have also reported emission ratios derived using only nighttime data. 638

639 Table 1: Comparative summary of the average mixing ratio (ppb) and Emission Ratios of VOC/ CO (ppb/ppm) of 640 Delhi (in parentheses) with other megacities of Asia, Europe and North America

641	` `	,	0	,	•					
VOC	Delhi*	Langzhou Valley ¹	Sao Paulo ²	London ³	Los	Paris ^{5(a)}	Mexico City ^{6(b)}	New York ^{4(c)}	Beijing ^{7(d)}	Lahore ⁸
					Angeles ⁴ (a)					
Benzene	2.02	0.54	0.67	0.31	0.48	0.38	0.80	0.74	1.79	28.20
	(2.65)	(1.37)	(1.03)	(1.59)	(1.30)	(1.07)	(1.21)	(1.09)	(1.24)	(5.08)
Toluene	5.15	0.72	2.11	0.60	1.38	1.40	3.10	0.19	1.98	32.40
	(7.03)	(1.41)	(3.1)	(3.09)	(3.18)	(12.30)	(4.20)	(3.79)	(2.41)	(6.67)
Sum of C8	2.74	0.61	1.52	0.63	1.03	1.30	1.10	0.88	2.66	29.40
aromatics	(4.20)	(1.42)	(2.15)	(3.69)	(2.45)	(4.75)	(4.30)	(1.11)	(2.15)	(6.04)
642 *	* This work (2022) ¹ Zhou et al., (2019)		² Brito et al., (2015)		³ Valach et al., (2014)		⁴ Baker et al., (2008)			

643

^(a)Borbon et al., (2013) ⁵Gros et al., (2011) ⁶Garzón et al., (2015) ⁷Yang et al., (2019a) 644 (2016) ^(b)Bon et al., (2011) ^(c)Warneke et al., (2007) ^(d)Wang et al., (2014) +*Apel et al.*, (2010) 645

646 Except for Lahore, where benzene levels were 10 times higher, benzene levels in Delhi were comparable to Beijing 647 and about three times higher than those that have been reported from other megacities like Sao Paulo, London, 648 Los Angeles, Paris, Mexico City and New York. The annual averaged national ambient air quality standard for 649 benzene is 5 μ g m⁻³ in India which is approximately 1.6 ppb at room temperature. Thus, the data suggest that 650 sources in the investigated period (Monsoon and Post-monsoon season) would contribute to violation of the annual 651 averaged values. Similarly, toluene and the sum of C8 aromatic compounds (e.g. xylene and ethyl benzene 652 isomers) were 6 to 10 times higher in Lahore compared to Delhi and more than twice as high relative to the 653 aforementioned megacities, except for Beijing, where the sum of C8 aromatic compounds were comparable to 654 Delhi. Overall, this indicates that Delhi has much higher levels of aromatic VOC pollution than many other 655 megacities. When we peruse the emission ratios (ER) that have been reported for these compounds in these other 656 megacities (shown in parentheses in Table 1), barring few exceptions (e.g. Lahore and Paris), the ERs were 657 generally much higher in Delhi with an average value of 2.65, as compared to cities like Sao Paulo (Brito et al., 658 2015), London (Valach et al., 2014) and Los Angeles and Paris (Borbon et al., 2013), Mexico City (Bon et al., 659 2011) and several US cities (Baker et al., 2008). The ER of toluene was highest in Paris (12.3) followed by Delhi. 660 Overall, the mixing ratios and ERs indicate that the influence of non -traffic sources (e.g. biomass burning and 661 industries) is more significant in Delhi compared to many other megacities of the world. The companion paper on 662 source apportionment based on this dataset (Awasthi et al., 2024) will focus more on the quantitative contributions 663 of the different sources.

⁸Barletta et al.,

664 4. Conclusion

665 This study has provided unprecedented characterization of the VOC chemical composition of ambient air in Delhi666 for the clean monsoon and extremely polluted post-monsoon seasons. The total average mass concentration of the

reactive carbon in the form of the 111 VOC species identified unambiguously was $\sim 260 \ \mu gm^{-3}$ and more than 4

- times higher during the polluted post-monsoon season mainly due to the impact of large scale open fires and
- reduced ventilation relative to the "cleaner" monsoon season. Of the 111, 42 were pure hydrocarbons (CH), 56
 were oxygenated volatile organic compounds (OVOCs; CHO), 10 were nitrogen containing compounds (NVOCs;
- 671 CHON), 2 were chlorinated volatile organic compounds (CIVOCs), and 1 namely methanethiol, contained 672 sulphur. The detection of new compounds that have previously not been observed discovered in Delhi's air, under both the clean and polluted periods such as methanethiol, dichlorobenzenes, C6-amides and C9-organic acids in 673 674 the gas phase was very surprising, considering there have been several PTR-TOF MS studies in Delhi earlier 675 (Wang et al., 2020; Tripathi et al., 2022; Jain et al., 2022). Our data points to both industrial sources of the sulphur 676 and chlorine compounds, photochemical source of the C6-amides and multiphase oxidation and chemical 677 partitioning for the C9-organic acids. To our knowledge this is the first reported study world-wide to detect and 678 observe only photo-chemically formed C6-amides in the gas phase. C6-amides are IVOCs, which can easily 679 partition to aerosol phase depending on environmental conditions and also act as a new source of reactive organic
- 680 nitrogen to the atmospheric environment.
- 681 The monsoon season VOC abundances for major compounds were comparable to several other megacities of the 682 world showing that the baseline VOC levels for the city of Delhi due to year-round active sources, helped by 683 favourable meteorological conditions for removal of VOCs through ventilation and wet scavenging, can lead to 684 comparable air quality as observed in other megacities. The VOC levels during the polluted post-monsoon season 685 when severe air pollution events occur leading to shutdowns and curbs, on the other hand were significantly (2-3 686 times) higher. Overall, for many important aromatic VOCs, the levels measured in Delhi were even higher (> 5 687 times) than many other megacities of the world located in Europe and North America. Generally these aromatic 688 compounds in megacities are primarily due to traffic and industrial emission sources, and this source is of course 689 common to Delhi and megacities in Europe and North America. In Delhi, the highest ambient mixing ratios of 690 these aromatic compounds occurred in the post-monsoon season. This is the period when enhanced open biomass 691 burning occurs due to heating demand increase owing to dip in temperatures (Hakkim et al., 2019; Awasthi et al., 692 2024) and open fire emissions due to the seasonal post-harvest paddy stubble biomass burning in which more than 693 1 billion ton of biomass is burnt regionally (Kumar et al., 2021) within few weeks during mid-October to end of 694 November occur. This adds significantly to the atmospheric burden of these compounds, compared to megacities 695 in developed countries where open biomass burning is better and more strictly regulated. Secondly, the 696 meteorological conditions during post-monsoon season due to shallower boundary layer height and poor 697 ventilation, and lack of wet scavenging due to absence of rain also slow down atmospheric removal of these 698 compounds compared to megacities in Europe, wherein it rains more frequently throughout the year compared to 699 Delhi.
- The presence of such a complex mixture of reactant VOCs adds to the air pollutant burden through secondary pollutant formation of aerosols. The reactive gaseous organics, which reached total averaged mass concentrations of ~85 μ gm⁻³ (monsoon season) and ~265 μ gm⁻³ (post-monsoon season) were found to rival the high mass
- concentrations of the main air pollutant in exceedance at this time, namely PM_{2.5} during the extremely polluted

- periods (post-monsoon season average: $\sim 145 \ \mu gm^{-3}$ which exceeds the 24h national ambient air quality standard
- of 60 μ gm⁻³). The data of the time series of the PM_{2.5} hourly data along with acetonitrile (a biomass burning VOC
- tracer) measured at the same site is provided in Figure S10. While the present study has quantified the molecules
- in the gas phase that are important for the air chemistry driving the high pollution events in Delhi in unprecedented
- detail, the implications on secondary pollutant formation will require building up on this new strategic knowledge
- 709 and further investigations. Moreover, the unique primary observations will yield quantitative source
- apportionment of particulate matter and VOCs in a companion study (Awasthi et al., 2024), that is being been co-
- submitted to this journal to enrich the scientific insights.
- All previous VOC studies in the literature from a dynamically growing and changing megacity like Delhi were
- reported for periods before 2020 (pre-COVID) times, without the new enhanced volatility VOC quantification
- technology deployed for the first time in a complex ambient environment of a developing world megacity like
- 715 Delhi. These have resulted in unprecedented new information concerning the speciation, abundance, ambient
- variability and emission characteristics of several rarely measured/reported VOCs. The significance of the new
- vunderstanding concerning atmospheric composition and chemistry of highly polluted urban atmospheric
- environments gained from this study, will no doubt be of global relevance as they would aid atmospheric chemistry
- 719 investigations in many megacities and polluted urban environments of the global south, that are in similar
- development and growth trajectory as Delhi and experience extreme air pollution and air quality associated
- 721 challenges, but remain understudied.

722 Data availability

The primary VOC, CO and Ozone and meteorological data presented in this manuscript can be downloaded by
 accessing the following Mendeley doi link: <u>https://data.mendeley.com/preview/pb6xs2fzwc?a=7658dfde-2ca0-</u>
 46c8-b89b-54ba8211e1de

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727 Author Contribution

Sachin Mishra: Data curation, Formal analysis, Investigation, Software, Visualization, Writing – original draft
preparation. Vinayak Sinha: Conceptualization, Data curation, Formal analysis, Methodology, Project
administration, Software, Supervision, Validation, Writing – review & editing. Haseeb Hakkim: Data curation,
Formal analysis, Investigation, Writing – review & editing. Arpit Awasthi : Data curation, Formal analysis,
Investigation. Sachin D. Ghude: Writing – review & editing. Vijay Kumar Soni: Writing – review & editing. N.
Nigam: resources. Baerbel Sinha: Conceptualization, Data curation, Supervision, Writing – review & editing. M.
Rajeevan: Writing – review & editing.

735 Competing Interests

736 The authors declare that they have no conflict of interest.

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