Emissions of non-methane volatile organic compounds from combustion of domestic fuels in Delhi, India


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Abstract. Twenty-nine different fuel types used in residential dwellings in northern India were collected from across Delhi (76 samples in total). Emission factors of a wide range of non-methane volatile organic compounds (NMVCs) (192 compounds in total) were measured during controlled burning experiments using dual-channel gas chromatography with flame ionisation detection (DC-GC-FID), two-dimensional gas chromatography (GC × GC-FID), proton-transfer-reaction time-of-flight mass spectrometry (PTR-ToF-MS) and solid-phase extraction two-dimensional gas chromatography with time-of-flight mass spectrometry (SPE-GC × GC–ToF-MS). On average, 94% speciation of total measured NMVOC emissions was achieved across all fuel types. The largest contributors to emissions from most fuel types were small non-aromatic oxygenated species, phenolics and furanics. The emission factors (in g kg⁻¹) for total gas-phase NMVOCs were fuelwood (18.7, 4.3–96.7), cow dung cake (62.0, 35.3–83.0), crop residue (37.9, 8.9–73.8), charcoal (5.4, 2.4–7.9), sawdust (72.4, 28.6–115.5), municipal solid waste (87.3, 56.6–119.1) and liquefied petroleum gas (5.7, 1.9–9.8).

The emission factors measured in this study allow for better characterisation, evaluation and understanding of the air quality impacts of residential solid-fuel combustion in India.

1 Introduction

Biomass burning is the second largest source of trace gases to the troposphere, releasing around a half of global CO, ~20% of NO and ~8% of CO₂ emissions (Olivier et al., 2005; Wiedinmyer et al., 2011; Andreae, 2019). Biomass burning releases an estimated 62 Tg yr⁻¹ of non-methane volatile organic compounds (NMVCs) (Andreae, 2019) and is the dominant source of both black carbon (BC) and primary organic aerosol (POA), representing 59% and 85% of global emissions, respectively (Bond et al., 2013). Biomass burning includes open vegetation fires in forests, savannahs, agricultural burning and peatlands (Chen et al., 2017) as well as the biofuels used by approximately 3 billion people to meet their daily cooking and heating energy requirements worldwide (World Bank, 2020). A wide range
of trace gases are released from biomass burning, in different amounts depending on the fuel type and the combustion conditions, meaning that detailed studies at the point of emission are required to accurately characterise emissions. The gases released lead to soil-nutrient redistribution (Ponette-Gonzalez et al., 2016; N’Dri et al., 2019); can themselves be toxic (Naehler et al., 2007); and can significantly degrade local, regional, and global air quality through the photochemical formation of secondary pollutants such as ozone ($O_3$) (Pfister et al., 2008; Jaffe and Wigder, 2012) and secondary organic aerosol (SOA) (Alvarado et al., 2015; Kroll and Seinfeld, 2008). They can also lead to indoor air quality issues (Fullerton et al., 2008).

Emissions from biomass burning and their spatial distribution remain uncertain, and estimates by satellite retrieval vary by over a factor of 3 (Andreae, 2019). Bottom-up approaches use information about emission factors and fuel usage. However, information for many developing countries, where solid fuels are a primary energy source, is particularly sparse. Toxic pollution from burning has been linked to chronic bronchitis (Akhtar et al., 2007; Moran-Mendoza et al., 2008), chronic obstructive pulmonary disease (Denis et al., 1996; Orozco-Levi et al., 2006; Rinne et al., 2006; Ramirez-Venegas et al., 2006; Liu et al., 2007; Perez Padilla et al., 1996), lung cancer (Liu et al., 1993; Ko et al., 1997), childhood pneumonia (Smith et al., 2011), acute lower respiratory infections (Bautista et al., 2009; Mishra, 2003) and low birth weight of children (Boy et al., 2002; Yucra et al., 2011). Smoke from inefficient combustion of domestic solid fuels is the leading cause of conjunctivitis in developing countries (West et al., 2013). The harmful emissions from burning also resulted in an estimated 2.8–3.9 million premature deaths due to household air pollution (Kodros et al., 2018; WHO, 2018; Smith et al., 2014), of which 27% originated from pneumonia, 18% from strokes, 27% from ischaemic heart disease, 20% from chronic obstructive pulmonary disease and 8% from lung cancer, with hazardous indoor air pollution responsible for 45% of pneumonia deaths in children less than 5 years old (WHO, 2018). For this reason, hazardous indoor air pollution from the combustion of solid fuels has been calculated to be the most important risk factor for the burden of disease in South Asia from a range of 67 environmental and lifestyle risks (Lim et al., 2012; Smith et al., 2014).

The emissions from biomass burning fires are complex and can contain many hundreds to thousands of chemical species (Crutzen et al., 1979; McDonald et al., 2000; Hays et al., 2002; Hatch et al., 2018; Stewart et al., 2021c). Measurements of emissions by gas chromatography (GC) have been made (EPA, 2000; Wang et al., 2014; Gilman et al., 2015; Stockwell et al., 2016; Fleming et al., 2018), as it has the potential to provide isomeric speciation of emissions. However, it is of limited use in untargeted measurements from burning due to the complexity of emissions, leading to large amounts of NMVOCs released not being observed. Some of the main issues are that GC does not provide high time resolution measurements, and several instruments with different column configurations and detectors are required to provide information on different chemical classes. Samples can also be collected into canisters or sample bags and then analysed offline (Wang et al., 2014; Sirithian et al., 2018; Barabad et al., 2018), which can increase time resolution but can also lead to artefacts (Lerner et al., 2017).

Recent developments have allowed for the application of proton-transfer-reaction mass spectrometry (PTR-MS) to study the emissions from biomass burning (Warneke et al., 2011; Yokelson et al., 2013; Brilli et al., 2014; Stockwell et al., 2015; Bruns et al., 2016; Koss et al., 2018). PTR-MS uses proton transfer from the hydronium ion ($H_3O^+$) to ionise and simultaneously detect most polar and unsaturated NMVOCs including aromatics, oxygenated aromatics, alkenes, furans and nitrogen-containing volatile organic compounds in gas samples. PTR-MS can measure at fast acquisition rates of up to 10 Hz over a mass range of 10–500 Th (thomson) with very low detection limits of tens to hundreds of parts per trillion by volume (pptv) (Yuan et al., 2016). The more recently developed technique of proton-transfer-reaction time-of-flight mass spectrometry (PTR-ToF-MS) has allowed for around 90% of total measured NMVOC emissions in terms of mixing ratio from burning experiments to be speciated (Koss et al., 2018) and has also been used to study the formation of SOA (Bruns et al., 2016). The main disadvantages of the PTR-ToF-MS technique are its inability to speciate isomers, significant fragmentation of parent ions, only being able to detect species with a proton affinity greater than water and the formation of water clusters needing to be considered (Stockwell et al., 2015; Yuan et al., 2017). More recently, measurements have also been made using iodide chemical ionisation time-of-flight mass spectrometry ($I^-\text{-CIMS}$), which is well suited to measuring acids and multifunctional oxygenates (Lee et al., 2014) as well as isocyanates, amides and organo-nitrate species released from biomass burning (Priestley et al., 2018). Multiple measurement techniques used in concert are therefore complementary, with the use of PTR-ToF-MS and simultaneous gas chromatography often alleviating some of the difficulties highlighted above.

Since the start of the century, rapid growth has resulted in India becoming the second largest contributor to NMVOC emissions in Asia (Kurokawa et al., 2013; Kurokawa and Ohara, 2020). However, effective understanding of the relative strength of different sources and subsequent mitigation has been limited by a deficiency of suitably detailed, spatially disaggregated emission inventories (Garaga et al., 2018). A current receptor-model study has shown elevated NMVOC concentrations at an urban site in Delhi to be predominantly due to vehicular emissions, with a smaller contribution from solid-fuel combustion (Stewart et al., 2021a). However, approximately 60% of total NMVOC emissions from India in 2010 were estimated to be due to solid-fuel combustion (Sharma et al., 2015). Other studies have also suggested that

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burning may lead to enhanced concentrations of pollutants such as polycyclic aromatic hydrocarbons in Delhi (Elzein et al., 2020). A need has therefore been identified to measure local source profiles to allow for evaluation with activity data to better understand the impact of unaccounted and unregulated local sources (Pant and Harrison, 2012).

Approximately 25% of worldwide residential solid-fuel use takes place in India (World Bank, 2020), with approximately 25% of ambient particulate matter in South Asia attributed to cooking emissions (Chafe et al., 2014). Despite large government schemes, traditional solid-fuel cookstoves remain popular in India; this is because they are cheaper than ones that use liquefied petroleum gas (LPG), and the meals cooked on them are perceived to be tastier (Mukhopadhyay et al., 2012). The total number of biofuel users has been sustained by an increasing population, despite the percentage use of biofuels decreasing as a proportion of overall fuel use due to increased LPG uptake (Pandey et al., 2014). Cow dung cakes remain prevalent as a fuel, because they are cheap, readily available, sustainable and ease pressure on local fuelwood resources. Few studies have reported emissions data from cow dung cake (Venkataraman et al., 2010; Stockwell et al., 2016; Koss et al., 2018; Fleming et al., 2018), leaving considerable uncertainty over the impact that cow dung cake combustion has on air quality. LPG usage has increased to 500 million users but only reflects around 10% of current rural fuel consumption (Gould and Urpelainen, 2018).

India-specific inventories which include residential burning indicate a considerable emission source of total NMVOCs of around 6000–7000 kt yr⁻¹ (Pandey et al., 2014; Sharma et al., 2015). Burning is likely to have a large impact on air quality in India, but considerable uncertainties exist over the total amount of NMVOCs released, owing to a lack of India-specific emission factors and information related to the spatial distribution of emissions.

Few studies measuring highly speciated NMVOC emission factors from fuels specific to India exist. Recent studies using PTR-ToF-MS to develop emission factors, which are more reflective of the range of species emitted from burning, have focussed largely on grasses, crop residues and peat (Stockwell et al., 2015) as well as fuels characteristic of the western US (Koss et al., 2018). A previous study measured emission factors of NMVOCs from cow dung cake using gas chromatography with flame ionisation detection (GC-FID) of 8–32 g kg⁻¹ (EPA, 2000). Fleming et al. (2018) quantified 76 NMVOCs from fuelwood and cow dung cake combustion using chulha and angithi stoves by collecting samples into Kynar bags; transferring their contents into canisters; and offline analysing samples using GC-FID, GC-ECD (electron capture detector), and GC-MS. The emission factors measured from these 76 NMVOCs were 14 g kg⁻¹ for cow dung cake burnt in chulha stoves, 27 g kg⁻¹ for cow dung cake burnt in angithi stoves and 6 g kg⁻¹ for fuelwood burnt in angithi stoves. An emission factor from one single dung burn measured using PTR-ToF-MS was considerably larger at around 66 g kg⁻¹ (Koss et al., 2018). Emissions from dung in Nepal have also been measured (Stockwell et al., 2016) by sampling into whole air sample canisters followed by offline analysis with GC-FID/ECD/MS and Fourier transform infrared spectroscopy (FTIR). However, very few speciated NMVOC measurements were made, and the emission factors were similar to those measured using just GC (Fleming et al., 2018). Studies have also focussed on making detailed measurements, using a range of techniques, from the burning of municipal solid waste (Christian et al., 2010; Yokelson et al., 2011, 2013; Stockwell et al., 2015, 2016; Sharma et al., 2019) and crop residues (Stockwell et al., 2015; Koss et al., 2018; Kumar et al., 2018).

Detailed chemical characterisation of NMVOC emissions from fuel types widely used in the developing world is much needed to resolve uncertainties in emission inventories used in regional policy models and global chemical transport models. A greater understanding of the key sources is required to characterise and hence understand air quality issues to allow for the development of effective mitigation strategies. In the present study, we measure comprehensive emission factors of NMVOCs from a range of solid fuels characteristic to northern India.

2 Methods

2.1 Fuel collection and burning facility

A total of 76 fuels, reflecting the range of fuel types used in northern India, were collected from across Delhi (see Fig. 1 and Table 1). Cow dung cake usage was prominent in the north and west regions, whereas fuelwood use was more evenly spread across the state. Municipal solid waste was collected from Bhalaswa, Ghazipur and Okhla landfill sites. Collection also included less used local fuel types which were found being burnt, including crop residues, sawdust and charcoal. A low-cost LPG stove, widely promoted across India as a cleaner fuel through government initiatives such as the Pradhan Mantri Ujjwala Yojana and Pratyaksh Hanstantrit Labh schemes, was used for direct emission comparison with other local fuel types.

Fuels were burnt at the CSIR-National Physical Laboratory (NPL), New Delhi, under controlled conditions utilising a combustion chamber based on the design of Venkataraman and Rao (2001). Several previous studies have been based on this chamber design (Venkataraman and Rao, 2001; Venkataraman et al., 2002; Saud et al., 2011, 2012; Singh et al., 2013), which was designed to simulate the combustion-driven conditions of real-world combustion and is displayed in Sect. S1 in the Supplement. The burn cycle used in this study was adapted from the Volunteers in Technical Assistance (VITA) water-boiling test, which was designed to simulate emissions from cooking and included emissions from both low- and high-temperature burning conditions. Fuels
Table 1. Types and numbers of fuel samples burnt, the mean emission factor of total measured NMVOCs (TVOC) in g kg\(^{-1}\) and standard deviation (SD) from all available burns.

<table>
<thead>
<tr>
<th>Fuelwood types</th>
<th>(n)</th>
<th>TVOC</th>
<th>SD</th>
<th>Other</th>
<th>(n)</th>
<th>TVOC</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Azadirachta indica</td>
<td>3</td>
<td>18.6</td>
<td>7.9</td>
<td>Cow dung cake</td>
<td>8</td>
<td>61.9</td>
<td>18.4</td>
</tr>
<tr>
<td>Morus spp.</td>
<td>4</td>
<td>27.4</td>
<td>21.1</td>
<td><em>Cocos nucifera</em></td>
<td>2</td>
<td>57.4</td>
<td>23.3</td>
</tr>
<tr>
<td>Melia azedarach</td>
<td>2</td>
<td>23.7</td>
<td>13.1</td>
<td>Charcoal</td>
<td>2</td>
<td>5.1</td>
<td>3.9</td>
</tr>
<tr>
<td>Shorea spp.</td>
<td>2</td>
<td>9.8</td>
<td>2.2</td>
<td>Sawdust</td>
<td>2</td>
<td>71.3</td>
<td>60.8</td>
</tr>
<tr>
<td>Ficus religiosa</td>
<td>2</td>
<td>51.9</td>
<td>63.4</td>
<td>Waste</td>
<td>3</td>
<td>87.3</td>
<td>31.4</td>
</tr>
<tr>
<td>Syzygium spp.</td>
<td>2</td>
<td>8.9</td>
<td>4.9</td>
<td>LPG</td>
<td>3</td>
<td>5.8</td>
<td>5.6</td>
</tr>
<tr>
<td>Ficus spp.</td>
<td>2</td>
<td>7.1</td>
<td>1.2</td>
<td>Cow dung cake mix</td>
<td>1</td>
<td>34.7</td>
<td>–</td>
</tr>
<tr>
<td>Vachellia spp.</td>
<td>2</td>
<td>13.5</td>
<td>9.7</td>
<td><em>Solanum melongena</em></td>
<td>2</td>
<td>13.6</td>
<td>6.5</td>
</tr>
<tr>
<td>Dalbergia sissoo</td>
<td>2</td>
<td>17.9</td>
<td>8.8</td>
<td>Brassica spp.</td>
<td>2</td>
<td>41.0</td>
<td>45.5</td>
</tr>
<tr>
<td>Ricinus spp.</td>
<td>2</td>
<td>8.5</td>
<td>2.5</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Holoptelea spp.</td>
<td>2</td>
<td>6.0</td>
<td>0.8</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed woods</td>
<td>1</td>
<td>6.1</td>
<td>–</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Saraca indica</td>
<td>2</td>
<td>12.9</td>
<td>5.2</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Populus spp.</td>
<td>1</td>
<td>8.5</td>
<td>–</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pithecellobium spp.</td>
<td>2</td>
<td>19.5</td>
<td>5.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Eucalyptus spp.</td>
<td>2</td>
<td>6.9</td>
<td>1.9</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Prosopis spp.</td>
<td>6</td>
<td>14.5</td>
<td>10.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mangifera indica</td>
<td>2</td>
<td>12.4</td>
<td>3.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plywood</td>
<td>8</td>
<td>26.6</td>
<td>24.3</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Processed wood</td>
<td>2</td>
<td>33.7</td>
<td>17.2</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 1. Locations across Delhi used for the local surveys of fuel use and collection of representative biomass fuels. Map tiles by Stamen Design. Data by © OpenStreetMap contributors 2020. Distributed under a Creative Commons BY-SA License.

were collected and stored in the same manner as local customs using expert local judgement. This method was designed to ensure that the moisture content of fuelwood samples was like those being burnt locally and that the combustion replicated real-world burning conditions encountered in local cooking practices, which should consequently give a more reflective NMVOC emission factor.

Fuel (200 g) was placed 45 cm from the top of the hood and rapidly heated to spontaneous ignition, with emissions convectively driven into a hood and up a flue to allow enough dilution, cooling and residence time to achieve the quenching typically observed in indoor environments. These conditions have been previously optimised to ensure that emissions entrainment into the hood did not exert a draught which altered combustion conditions. The mid-point velocity of gases driven up the flue by convection was measured by a platinum hot-wire sensor, calibrated for total flow rate using a standard orifice calibrator. Samples were drawn down a sample line at 4.4 L min\(^{-1}\) (Swagelok, 1/4 in. PFA, < 2.2 s residence time) from the top of the flue, passed through a pre-conditioned quartz filter (\(ø = 47\) mm, conditioned at 550°C for 6h and changed between samples) and held in a filter holder (Cole-Parmer, PFA) which was subsampled for analysis by PTR-ToF-MS, GC × GC-FID and DC-GC-FID instruments at a distance no greater than 5 m from the top of the flue.

Measurements of \(n\)-alkanes from \(n\)-tridecane (C\(_{13}\)) to eicosane (C\(_{20}\)) were also made from a subset of 29 burns using solid-phase extraction discs (SPE, Resprep, C\(_{18}\)). Samples were passed through a cooling and dilution chamber designed to replicate the immediate condensational processes that occur in smoke particles approximately 5–20 min after emission yet prior to photochemistry which may change composition (Akagi et al., 2011). Further details of the SPE sample collection are given in Stewart et al. (2021c).
2.2 PTR-ToF-MS

The PTR-ToF-MS (PTR 8000; Ionicon Analytik, Innsbruck) instrument from the Physical Research Laboratory (PRL), Ahmedabad, was used to quantify 107 masses and sub-sampled the common inlet line using 1/4 in. PFA (perfluoroalkoxy alkane). Additional details of the PTR-ToF-MS system used in this study are given in previous papers (Sahu and Saxena, 2015; Sahu et al., 2016). The sample air was diluted into zero air, generated by passing ambient air (1 L min$^{-1}$) through a heated platinum filament at 550 °C before entering the instrument with an inlet flow of 250 mL min$^{-1}$. Samples were diluted by either 5 or 6.25 times (50 mL min$^{-1}$ in 200 mL min$^{-1}$ zero air or 40 mL min$^{-1}$ in 210 mL min$^{-1}$ zero air). The instrument was operated with an electric field strength (E/N, where N is the buffer gas density and E is the electric field strength) of 120 Td (townsend, 1 Td = 10−17 V cm²). The drift tube temperature was 60 °C with a pressure of 2.3 mbar and 560 V applied across it.

Calibrations were performed twice a week using a gas calibration unit (Ionicon Analytik, Innsbruck). The calibration gas (Apel-Riemer Environmental Inc., Miami) contained 18 compounds: methanol, acetonitrile, acetaldehyde, acetone, dimethyl sulfide, isoprene, methacrolein, methyl vinyl ketone, 2-butanone, benzene, toluene, 2-hexanone, m-xylene, heptanal, α-pinene, 3-octanone and 3-octanol at 1000 ppbv (±5 %), and β-caryophyllene at 500 ppbv (±5 %). This standard was dynamically diluted into zero air to provide a six-point calibration. The normalized counts per second (ncps) per ppbv of the NMVOC (ncps/ppbv) was then determined for each mass using a transmission curve (Tai pale et al., 2008). The maximum error in this calibration approach was shown to be 21 % (Tai pale et al., 2008). Peak assignment was assisted with results reported by previous burning studies and references therein (Brilli et al., 2014; Stockwell et al., 2015; Koss et al., 2018). The results may also contain other indistinguishable structural isomers not mentioned here.

Mass calibration and peak fitting of PTR-ToF-MS data were performed using PTRwid software (Holzinger, 2015). Count rates per second (cps) of each mass spectral peak were normalised to the primary ion (H$_3$O$^+$) and water cluster (H$_2$O·H$_2$O$^+$) peaks, and mixing ratios were then determined for each mass using the normalised sensitivity. Where compounds known to fragment in the PTR-ToF-MS were identified, the mixing ratio of these species was calculated by summing parent ion and fragment ion mixing ratios. Before each burn, ambient air was sampled to provide a background for the measurement.

Petrol and diesel fuel samples were collected from an Indian Oil fuel station in Pusa, New Delhi, and the headspace was analysed to allow for comparison of benzene-to-toluene ratios. This method was designed to analyse the ratios in evaporative emissions, as these have been shown to be an important source of atmospheric NMVOCs (Srivastava et al., 2005; Rubin et al., 2006; Yamada et al., 2015), which for example represented ∼15 % of anthropogenic UK NMVOC emissions in 2018 (Lewis et al., 2020). Fuel samples were placed in a small metal container (1/4 in. Swagelok cap) which was connected to a two-way tap (1/4 in. Swagelok) with a flow of zero air (250 mL min$^{-1}$) passed through it and could be sampled by the PTR-ToF-MS. The tap was then opened to analyse the headspace of fuels.

2.3 DC-GC-FID

Gas chromatography was used to analyse entire burns to provide an integrated picture of emissions from fuel types. The DC-GC-FID sampled 51 burns to measure 19 C$_2$–C$_7$ non-methane hydrocarbons (NMHCs) and C$_2$–C$_5$ oxygenated NMVOCs (OVOCs) (Hopkins et al., 2003). A 500 mL sample (1.5 L pre-purge of 100 mL min$^{-1}$ for 15 min, sample at 17 mL min$^{-1}$ for 30 min) was collected (Markes International CIA Advantage), passed through a glass finger at −30 °C to remove water and adsorbed onto a dual-bed sorbent trap (Markes International ozone precursors trap) at −20 °C (Markes International Unity 2). The sample was thermally desorbed (250 °C for 3 min) then split 50:50 and injected into two separate columns for analysis of NMHCs (50 m × 0.53 mm Al$_2$O$_3$ PLOT) and OVOCs (10 m × 0.53 mm LOWOX with 50 µm restrictor to balance flow). The oven was held at 40 °C for 5 min, then heated at 13 °C min$^{-1}$ to 110 °C, and then finally heated at 8 °C min$^{-1}$ to 200 °C with a 30 min hold.

2.4 GC × GC-FID

The GC × GC-FID was used to measure 58 C$_7$–C$_{12}$ hydrocarbons (C$_7$–C$_{12}$ alkanes, monoterpenes and monoaromatics) and collected 3 L samples (100 mL min$^{-1}$ for 30 min) using an adsorption–thermal-desorption system (Markes International Unity 2). NMVOCs were trapped onto a sorbent (Markes International U-T15ATA-2S) at −20 °C with water removed in a glass cold finger at −30 °C, removed and heated to ∼100 °C after each sample to prevent carry-over of unanalysed, polar interfering compounds. The sample was thermally desorbed (250 °C for 5 min) and injected splitless down a transfer line. Analytes were re-focussed for 60 s using liquid CO$_2$ at the head of a non-polar BPX5 held at 50 psi (1 psi = 6.9 kPa) (SGE Analytical, 15 m × 0.15 µm × 0.25 mm), which was connected to a polar BPX50 at 30 psi (SGE Analytical, 2 m × 0.25 µm × 0.25 mm) via a modulator held at 180 °C (5 s modulation, Analytical Flow Products ELDV2-2-MT). The oven was held for 2 min at 35 °C, then ramped at 2.5 °C min$^{-1}$ to 130 °C and held for 1 min with a final ramp of 10 °C min$^{-1}$ to 180 °C and a hold of 8 min. The GC systems were tested for breakthrough to ensure trapping of the most volatile components (see Sect. S2). Calibration was carried out using 4 ppbv gas standards containing alkanes.
alkenes and aromatics purchased from the British National Physical Laboratory and through the relative response of liq-
uid standard injections to toluene for components not in this
gas standard, as detailed elsewhere (Dunmore et al., 2015;
Stewart et al., 2021a). Integration of peak areas was per-
formed in Zoex GC image software (Zoex, USA). Peaks were
individually checked, and where peaks were split in the soft-
ware, they were manually joined. The areas corresponding to
alkane isomers were manually joined within the GC image
software and calibration performed by comparing the areas to
the corresponding n-alkane. For both GC instruments, blanks
of ambient air were made at the beginning, middle and end
of the day, and the mean was subtracted from samples.

2.5 GC × GC–ToF-MS

Measurements of a subset of 29 burns of C_{13}–C_{20} alkanes
were made, as well as other gas-phase species to assist with
qualification of masses measured by PTR-ToF-MS, by ad-
sorbing samples to the surface of SPE discs with analysis
by GC × GC–ToF-MS, as detailed in Stewart et al. (2021c).
Samples of 180 L were adsorbed to the surface of C_{18} coated
SPE discs (Resprep, C_{18}, 47 mm) prewashed with 2 × 5 mL
acetic acid washes and 1 × 5 mL methanol wash. These sam-
ple were collected at 6 L min^{-1} over 30 min using a low-
volume sampler (Vayubodhan Pvt. Ltd), which passed sam-
ple through a cooling and dilution chamber at 46.7 L min^{-1}.
Samples were then wrapped in foil, placed in an airtight bag
and kept frozen until analysis.

SPE extracts were spiked with an internal stan-
ard (EPA 8270 Semivolatile Internal Standard Mix,
2000 µg mL^{-1} in DCM, dichloromethane) and extracted us-
ing accelerated solvent extraction into ethyl acetate. Ex-
tracts were analysed using GC × GC–ToF-MS (Leco Pe-
gasus BT 4D) using a 10:1 split injection (1 µL in-
jection, 4 mm taper focus liner, SHG 560302). The primary
dimension column was a RXI-5SilMS (Restek,
30 m × 0.25 µm × 0.25 mm) connected to a second column of
RXI-17SilMS (Restek, 0.25 µm × 0.25 mm, 0.17 m pri-
mary GC oven, 0.1 m modulator, 1.42 m secondary oven,
0.31 m transfer line) under a He flow of 1.4 mL min^{-1}.
The primary oven was held at 280 °C and then ramped
at 3 °C min^{-1} to 202 °C where it was held for 0.07 min. The
secondary oven was held at 62 °C for 1 min and then ramped
at 3.2 °C min^{-1} to 235 °C. The inlet was held at 280 °C and
the transfer line at 340 °C. A 5 s cryogenic modulation was
used with a 1.5 s hot pulse and 1 s cool time between stages.

Peak assignment was conducted through comparison of
retention times to known standards and comparison to the
National Institute of Standards and Technology (NIST) mass
spectral library. Integration was carried out within the Chro-
maTOF 5.0 software package (Leco, 2019). Eight blank mea-
surements were made at the beginning and end of the day by
passing air from the chamber (6 L min^{-1} for 30 min) through
the filter holder containing a PTFE filter and an SPE disc.

Blank corrections have been applied by subtracting the mean
of blank values closest to measurement of the sample. An
eight-point calibration was performed for n-alkanes using a
commercial standard (C_{7}–C_{40} saturated alkane standard, cer-
tified 1000 µg mL^{-1} in hexane, Sigma-Aldrich 49452-U) di-
luted in the range 0.25–10 µg mL^{-1}.

3 Results

3.1 Comparison of chromatograms from combustion of
different fuel types

Figure 2 shows GC × GC–ToF-MS chromatograms obtained from
collecting the emissions during the combustion of LPG
(Fig. 2a), Saraca indica fuelwood (Fig. 2b), cow dung cake
(Fig. 2c) and municipal solid waste (Fig. 2d). Figure 2d is
labelled to show the position of NMVOCs measured and
displays a homologous series of n-alkanes from n-heptane
(C_{7}) to n-tetradecane (C_{14}) along the bottom, with the 1-
alkenes positioned to the left. Above are more polar species
such as monoterpenes, aromatics from benzene to substitu-
ted monoaromatics with up to five carbon substituents, and at
a higher second dimension retention time even more polar
species, such as styrene.

Many peaks were present in the chromatograms for cow
dung cake and municipal solid waste, and these fuels re-
leased significantly more NMVOCs per unit mass than fu-
elwood and LPG (see Table 1). Cow dung cake and mu-

nicipal solid waste released a range of NMVOCs includ-
ing n-alkanes, alkenes and aromatics. The municipal solid
waste (Fig. 2d) showed a particularly large and tailing peak
numbered 22, owing to large emissions of styrene. Several
unidentified peaks were observed in these complex samples
which were broad in the second dimension. These were as-
sumed to be from polar, oxygenated species formed during
burning such as phenol. These species could not be identified
and were not analysed using the GC × GC–ToF-FID instrumen-
Peaks have been omitted if these species were found to in-
terfere significantly. Analysis has only been carried out using
the DC-GC–ToF-FID from ethane (C_{2}) to n-hexane (C_{6}), owing to
the significant presence of coeluting peaks. The large peak in
the LPG chromatogram (Fig. 2, primary dimension retention
time ~ 6 min, secondary dimension retention time ~ 0.5 s)
was from unresolved propane and butane because of the high
concentrations from this fuel source.

3.2 PTR-ToF-MS concentration time series analysis

Figure 3 shows an example concentration–time-series plot
measured by the PTR-ToF-MS for a cow dung cake burn.
A sharp rise in NMVOC emissions was seen from the start
of the burn, which decreased as the fuel was combusted.
Emissions of small oxygenated species as well as phenolics
and furanics were dominant throughout most of the burn. At
the beginning, a greater proportion of lower-mass species
were released, as shown in the binned mass spectrum of region A in Fig. 3. At the end of the smouldering phase, emissions were dominated by heavier and lower-volatility species (Fig. 3, region B). A previous study indicated higher-molecular-weight phenolics were from low-temperature pyrolysis (Sekimoto et al., 2018).

Figure 4 shows the cumulative mass of species measured from burns of fuelwood, cow dung cake, municipal solid waste and charcoal as a proportion of the total mass of NMVOCs quantified using PTR-ToF-MS. The results were like those reported by Brilli et al. (2014) and Koss et al. (2018): 65\%–90\% of the mass of NMVOCs at emission originated from around 40 NMVOCs, with around 70\%–90\% identification by mass when quantifying around 100 NMVOCs. The largest contributors to the NMVOC mass from burning of fuelwood and cow dung cake were methanol (m/z 33.034); acetic acid (m/z 61.028); and a peak that reflected the sum of hydroxyacetone, methyl acetate, and ethyl formate (m/z 75.043). For municipal solid waste samples, around 28\% of total mass was from methyl methacrylate (m/z 101.059) and styrene (m/z 105.068), and two of the three municipal solid waste samples released significant quantities of styrene, most likely the result of degradation of polystyrene in the samples.

Figure 5 shows a concentration time series for phenolics and furanics from the burning of an example fuelwood. Most species of similar functionality tracked each other. Stockwell et al. (2015) demonstrated that benzene, phenol and furan could act as tracers for aromatic, phenolic and furanic species released from biomass burning. Figure 5a shows that heavier, more substituted phenolics appeared to be released at cooler temperatures. Guaiacol (dark blue) was released at the start of the flaming phase before the temperature increased and more phenol (red) was released at higher burn temperatures. Later in the burn, a larger proportion of vinyl guaiacol (pink) and syringol (yellow) were emitted. This agreed well with previous results which showed that species emitted from lower temperature depolymerisation had a larger proportion of low-volatility compounds compared to higher temperature processes during flaming (Sekimoto et al., 2018; Koss et al., 2018). Figure 5b shows concentration time series of furanic species, with most species showing similar characteristics throughout the burn. The only species to peak later in the burn was 2-hydroxymethyl-2-furan.

Figure 2. GC × GC-FID chromatograms from burning (a) LPG, (b) Saraca indica (fuelwood), (c) cow dung cake and (d) municipal solid waste samples where 1–7 = n-octane to n-tetradecane, 8–13 = 1-octadecene to 1-tridecene, 14 = benzene, 15 = toluene, 16 = ethylbenzene, 17 = m/p-xylene, 18 = o-xylene, 19 = C3-substituted monoaromatics, 20 = C4-substituted monoaromatics and 21 = C5-substituted monoaromatics and 22 = styrene. Samples (a)–(d) were collected with the same sample collection parameters, and the chromatograms are set at the same contrast level to allow for direct comparison between different fuel types.
3.3 Comparison of emissions data obtained with different instruments

Previous instrument intercomparisons from biomass burning samples were between PTR-MS, GC–MS and open-path FTIR (Gilman et al., 2015) and between PTR-ToF-MS, FTIR, airborne cavity-enhanced spectroscopy (ACES) and I−-CIMS (Koss et al., 2018). Gilman et al. (2015) showed generally good agreement of slopes of measured emission factors between benzene, ethyne, furan, ethene, propene, methanol, toluene, isoprene and acetonitrile using different instruments and/or techniques with slopes of $\sim 1 \pm 0.3\%$ and correlation coefficients $> 0.9$. Koss et al. (2018) showed mean measured values of most NMVOCs from all burns with other instruments compared to the PTR-ToF-MS which agreed within a factor of 2 and had correlation coefficients $> 0.8$ for most species except butadienes, furan, hydroxycetone, furfural, phenol and glyoxal. These previous comparisons indicate the level of consistency expected with instrument comparisons of quantitative NMVOC measurements from burning experiments.

Figure 6 shows a comparison of measurements made using the DC-GC-FID, GC × GC-FID and PTR-ToF-MS techniques. Bar plots show that the mean and lower/upper quartiles of all measurements agreed within a factor of 2 and had correlation coefficients $> 0.8$ for most species except butadienes, furan, hydroxycetone, furfural, phenol and glyoxal. Generally, the mean values measured for the PTR-ToF-MS were slightly larger than using the GC instruments, which was attributed to the presence of other undistinguishable structural isomers measured.
by the PTR-ToF-MS. Comparison between DC-GC-FID and GC × GC-FID measurements were also complicated by high levels of coelution of additional NMVOC species released from combustion with similar retention times ($R_t$) to benzene/toluene ($R_t = 21/25$ min) on the DC-GC-FID instrument. Generally, the smallest values were measured with the GC × GC-FID instrument, consistent with the greatest ability to speciate isomers and limit the impacts of coelution. Significant efforts were made to synchronise the sample periods for the three instruments as best as possible; however, slight uncertainty existed over the exact time each instrument started measuring when calculating mean sample windows (±30 s). These factors combined may help to explain the slight differences observed between different instruments during this study. When multiple instruments have measured the same NMVOC in this study, preference was given to the data from the GC × GC-FID due to the ability of this instrument to resolve coeluting peaks, followed by the DC-GC-FID and then the PTR-ToF-MS.

### 3.4 NMVOC emission factors from biomass fuels

Figure 7 shows a detailed breakdown of the mean NMVOC emission factors by fuel type measured for all 76 burns (see Sect. S3 for values). Emission factors have been determined by calculating the mean NMVOC concentrations up the flue over a 30 min period, in line with the GC sample time, with any small emissions after this sample window not included. This has been related to the total volume of air convectively drawn up the flue and the mass of fuel burnt (see Sect. S4 for details). The data are split by functionality to show trends for different chemical types. This shows that burning released a large amount of different NMVOCs across a wide range of functionalities, molecular weights and volatilities. The large variety of NMVOCs is likely to have different influences on O$_3$ formation, SOA production and the toxicity of emissions.

Figure 7a shows very large emissions of smaller oxygenated species which were driven by methanol; acetic acid; and the unresolved combined peak for hydroxy acetone, methyl acetate, and ethyl formate. For the fuelwood samples, acetic acid/glycolaldehyde (2.6 g kg$^{-1}$), methanol (1.8 g kg$^{-1}$) and acetaldehyde (0.6 g kg$^{-1}$) compared well with mean values reported by Koss et al. (2018) for pines, firs and spruces (2.7/1.3/0.9 g kg$^{-1}$) and the mean values measured by Stockwell et al. (2015) mainly from crop residues, grasses and spruces (1.6/1.3/0.9 g kg$^{-1}$). The emission factor from this study for the unresolved peak of hydroxy acetone, methyl acetate and ethyl formate (1.4 g kg$^{-1}$) was larger than...
Figure 7.
Figure 7. Measured emission factors grouped by functionality.
those previously reported by Koss et al. (2018) and Stockwell et al. (2015) of 0.55 and 0.25 g kg\(^{-1}\), respectively.

Figure 7b shows that there were large emissions of furans and furanones from combustion, mainly from methyl furans, furfurals, 2-(3H)-furanone, methyl furfurals and 2-methanol furanone. The World Health Organization considers furan a carcinogenic species of high priority (WHO, 2016) with furan and substituted furans suspected to be toxic and mutagenic (Ravindranath et al., 1984; Peterson, 2006; Monien et al., 2011). Furan emissions originate from the low-temperature depolymerisation of semi-cellulose (Sekimoto et al., 2018) and from large alcohols and enols in high-temperature regions of hydrocarbon flames (Johansson et al., 2016). The OH chemistry of furans has been the subject of several studies (Bierbach et al., 1994; Bierbach et al., 1995; Tapia et al., 2011; Liljegren and Stevens, 2013; Strollo and Ziemann, 2013; Zhao and Wang, 2017; Coggon et al., 2019) and often produces more reactive products such as butenodial, 4-oxo-2-pentenal and 2-methylbutenal (Bierbach et al., 1994; Gómez Alvarez et al., 2009; Aschmann et al., 2011, 2014). Photo-oxidation of furans may also be a potentially important source of small organic acids such as formic acid (Wang et al., 2020). Oxidation can also occur by nitrate (Berndt et al., 1997; Colmaner et al., 2012) or chlorine radicals (Cabañas et al., 2005; Villanueva et al., 2007). As a result, furans have recently been shown to be some of the species with the highest OH reactivity from biomass burning, causing an estimated 10 % of the O\(_3\) produced by the combustion emissions in the first 4 h after emission (Hartikainen et al., 2018; Coggon et al., 2019). Oxidation of furans can lead to SOA production (Gómez Alvarez et al., 2009; Strollo and Ziemann, 2013) with an estimated 8 %-15 % of the SOA caused by furans emitted by burning of black spruce, cut grass, Indonesian peat and ponderosa pine and 28 %-50 % of the SOA from rice straw and wire grass (Hatch et al., 2015), although SOA yields are still uncertain for many species (Hatch et al., 2017).

Phenols are formed from the low-temperature depolymerisation of lignin (Simoneit et al., 1993; Sekimoto et al., 2018), which is a polymer of randomly linked, amorphous high-molecular-weight phenolic compounds (Shafizadeh, 1982). Owing to their high emission factors and SOA formation potentials, phenolic compounds contribute significantly to SOA production from biomass burning emissions (Yee et al., 2013; Lauraguais et al., 2014; Gilman et al., 2015; Finewax et al., 2018). Figure 7c shows that the largest phenolic emissions from fuelwood in this study were methoxyphenols, with significant contributions from phenol, guaiacol, cresols and anisole. Phenolic emissions from sawdust were dominated by guaiacol and cresol. Phenolic emissions from coconut shell were greatest, most likely as a result of the lignin-rich nature of coconut shell (Pandharipande et al., 2018). The larger mean emission of furanics (3.2 g kg\(^{-1}\)) compared to phenolics (1.1 g kg\(^{-1}\)) from fuelwood was consistent with wood being composed of around 75 % cellulose or hemi-cellulose and 25 % lignin (Sjöström, 1993).

Figure 7d shows that the largest alkene emission was styrene from burning municipal solid waste, likely caused by the presence of polystyrene in the fuel. Emissions of alkenes from fuelwood were dominated by ethene and propene, which are species with high photochemical ozone creation potentials (Cheng et al., 2010). Monoterpenes, which are extremely reactive with the OH radical (Atkinson and Arey, 2003), were emitted from combustion of sawdust, cow dung cake and municipal solid waste samples.

Ethane and propane dominated the alkane emissions for fuelwood samples (see Fig. 7e). A wider range of alkanes from C\(_2\)–C\(_{20}\) were observed from combustion of coconut, cow dung cake and municipal solid waste. The largest alkane emission by mass was from LPG due to unburnt propane and butane.

Nitrogen-containing NMVOCs are formed from the volatilisation and decomposition of nitrogen-containing compounds within the fuel, mainly from free amino acids but can also be from pyrroline, pyridine and chlorophyll (Leppalähti and Koljonen, 1995; Burling et al., 2010; Ren and Zhao, 2015). Nitrogen-containing NMVOCs are of interest because nitrogen may be important in the development of new particles (Smith et al., 2008; Kirkby et al., 2011; Yu and Luo, 2014), which act as cloud condensation nuclei (Kerminen et al., 2005; Laaksonen et al., 2005; Sotiropoulou et al., 2006) and alter the hydrological cycle by forming new clouds and precipitation (Novakov and Penner, 1993). They can also contribute to light-absorbing brown carbon (BrC) aerosol formation, effecting climate (Laskin et al., 2015). Additionally, nitrogen-containing NMVOCs can be extremely toxic (Ramírez et al., 2012, 2014; Farren et al., 2015). Cow dung cake was the largest emitter of nitrogen-containing NMVOCs (4.9 g kg\(^{-1}\)), releasing large amounts of acetonitrile and nitriles, likely to have a large impact on the toxicity and chemistry of emissions (see Fig. 7f).

Figure 7g shows that emissions of aromatics from fuelwood, cow dung cake and municipal solid waste were principally benzene, toluene and naphthalenes. Large emissions of benzene were unsurprising as biomass burning is the largest global benzene source (Andreae and Merlet, 2001). Emissions of benzene, toluene, ethylbenzene and xylenes (BTX) from cow dung cake (0.5–1.7 g kg\(^{-1}\)) were in line with previous measurements of 1.3 g kg\(^{-1}\) (Koss et al., 2018) and 1.8 g kg\(^{-1}\) (Fleming et al., 2018), but they were lower than the 4.5 g kg\(^{-1}\) reported from cow dung cake combusted from Nepal (Stockwell et al., 2016). Emissions of BTX from municipal solid waste burning (0.9–2.6 g kg\(^{-1}\)) were comparable to those measured previously (3.5 g kg\(^{-1}\)) (Stockwell et al., 2016).

Figure 7h shows a qualitative comparison of species such as ammonia, HCN and dimethyl sulfide which were measured during experiments, but they could not be accurately quantified as their sensitivity was too different from the
NMVOCs used to build the transmission curve. Cow dung cake emitted significantly more of these species than other fuel types.

Table 2 shows the total measured emission factors of NMVOCs for different fuel types. The total measured emission factor has been calculated as the sum of the PTR-ToF-MS signal, excluding reagent ion peaks (< m/z 31 Th), water cluster peaks (m/z 37 Th) and isotope peaks identified for all masses (SIS, 2016). The emission factors for all alkanes and alkenes measured by the GC instruments were also included, as alkanes up to n-hexane had proton affinities less than water and larger alkanes had proton affinities similar to water (Ellis and Mayhew, 2014; Wróblewski et al., 2006). This low sensitivity meant that no peaks were present in the PTR-ToF-MS spectra for these larger species. Any alkenes measured by the DC-GC-FID were excluded from the PTR-ToF-MS data. Further information on the species included in the calculation of the total measured emission factor is given in Sect. S5.

Coconut shell, sawdust, cow dung cake and municipal solid waste released the greatest mass of NMVOC per kilogram of fuel burnt. The mean emission factor for all fuelwood types (18.7 g kg$^{-1}$) was comparable to that for chapparal (16.6 g kg$^{-1}$) measured using PTR-ToF-MS by Stockwell et al. (2015). This may be due to similarities between north Indian fuelwood types with chapparal, which is characterised by hot dry summers, and mild wet winters. The mean fuelwood emission factor was smaller than Stockwell et al. (2015) reported for coniferous canopy (31.0 g kg$^{-1}$). The NMVOC emission measured for cow dung cake (62.0 g kg$^{-1}$) was comparable to that previously reported (66.3 g kg$^{-1}$) in literature using PTR-ToF-MS (Koss et al., 2018) but 2–3 times larger than that measured by GC-FID/ECD/MS likely due to those techniques missing significant amounts of emissions (Fleming et al., 2018). Whilst the total measured emissions reported by Fleming et al. (2018) might therefore be an underestimate, it is noteworthy that the emission factors measured by Fleming et al. (2018) in angithi stoves for cow dung cake were a factor of ∼4 greater than fuelwood under the same conditions. This result was comparable to this study, which showed that cow dung cake emissions were a factor of ∼3 larger than fuelwood, but the techniques used here targeted a greater proportion of total emissions. Moreover, Fleming et al. (2018) reported emission factors from combustion of biomass fuels from a neighbouring state, Haryana, and there may be slight heterogeneity between the different fuels collected in both studies. Venkataraman et al. (2010) and Koss et al. (2018) also showed NMVOC emissions from dung combustion to be greater than from fuelwood. NMVOC emissions from municipal solid waste (87.3 g kg$^{-1}$) were significantly larger than the ∼9 g kg$^{-1}$ (Stockwell et al., 2015) and ∼35 g kg$^{-1}$ (Stockwell et al., 2016) previously reported. This was likely due to differences in composition and moisture content of the fuels collected from Indian landfill sites for the present study, compared with the daily mixed waste and plastic bags collected at a US fire services laboratory (Stockwell et al., 2015) and a variety of mixed waste and plastics collected from around Nepal (Stockwell et al., 2016). It seems noteworthy that combustion experiments of fuels collected from developing countries in Stockwell et al. (2016) had larger emission factors than those collected from and burnt at a laboratory (Stockwell et al., 2015). The mean crop residue combustion emission factor (37.9 g kg$^{-1}$) was comparable to that reported by Stockwell et al. (2015), despite the small number of samples in this study and compositional differences.

Considerable uncertainties exist in consumption estimates for fuels such as cow dung cake and municipal solid waste in India. A previous study estimated that in 1985 in India fuelwood consumption was 220 Tg and cow dung cake consumption 93 Tg (Yevich and Logan, 2003). A different study made an India-wide estimate for 2000 which estimated fuelwood consumption to be 281 (192–409) Tg and cow dung cake consumption to be 62 (35–128) Tg (Habib et al., 2004). A more recent study estimated fuelwood usage at 256 Tg and cow dung cake consumption at 106 Tg for 2007 (Singh et al., 2013). Estimates of the amount of municipal solid waste burnt in India are even fewer than for cow dung cake consumption. Two previous studies have estimated that 81.4 Tg of municipal solid waste was burnt in India in 2010 (Wiedinmyer et al., 2014) and that 68 (45–105) Tg was burnt in 2015 (Sharma et al., 2019). The mean emission factors for cow dung cake and municipal solid waste combustion were considerably larger than for fuelwood, and they highlight that at an India-wide level these may represent significant NMVOC sources.

Intermediate-volatility organic compounds (IVOCs) are defined as having effective saturation concentration, $C^*$, in the range 300–3 × 10$^6$ µg m$^{-3}$ (Donahue et al., 2012). The $C^*$ of several species was estimated using a previously established approach (Lu et al., 2018), with the IVOC boundary defined in this study at benzaldehyde ($m = 106.12$), for which $C^*$ was $\sim 7 \times 10^6$ µg m$^{-3}$. Table 2 also shows an approximation for the mean amount of IVOCs released by fuel type. This approach was approximate as vapour pressures depend on both mass and functionality. The fuels tested in this study showed that mean emissions of IVOC species represented approximately 18%–27% of total measured emissions from all fuel types other than LPG. This demonstrated that domestic solid-fuel combustion is potentially a large global source of IVOCs. In addition, this may represent an underestimate because, the quartz filter placed on the sample line may remove IVOC species which have partitioned to the aerosol phase due to the high aerosol concentrations present during source testing. Further studies are required to better understand the contribution of IVOC emissions from biomass burning to SOA formation. This is evaluated as part of a different study, using the dataset generated as part of this study and provided elsewhere (Stewart et al., 2021b).
Table 2. Mean total measured NMVOC emission factors (g kg\(^{-1}\), including IVOC fraction) where high and low EF represent the largest and smallest emission factors, respectively, measured for a given sample type (g kg\(^{-1}\)); IVOC is the sum of emission factors of species with a mass greater than benzaldehyde (g kg\(^{-1}\)), where \(n\) represents the number of measurements made.

<table>
<thead>
<tr>
<th></th>
<th>Wood</th>
<th>Dung</th>
<th>Waste</th>
<th>LPG</th>
<th>Charcoal</th>
<th>Sawdust</th>
<th>Crop</th>
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<td>5.7</td>
<td>5.4</td>
<td>72.4</td>
<td>37.9</td>
</tr>
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<tr>
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<td>56.3</td>
<td>1.9</td>
<td>2.4</td>
<td>28.3</td>
<td>8.9</td>
</tr>
<tr>
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<td>13.2</td>
<td>0.2</td>
<td>1.4</td>
<td>16.9</td>
<td>8.0</td>
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<tr>
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</table>

Figure 8. Variability in NMVOC emission factor by fuel type. (a) Range of emission factors measured for fuelwood, cow dung cake, crop residue and municipal solid waste samples with boxplots showing the mean, median, interquartile range and range within 1.5IQR and solid circles showing the spread of measured emission factors by fuel type. (b) Zoomed-in green region displaying range of NMVOC emission factors measured for individual species of fuelwood with the grey shaded region indicating repeat samples from the same sample collection location and diamonds indicating the measured NMVOC emission factors.

Figure 8a shows the distribution of total measured NMVOC emission factors for fuelwood, cow dung cake, crop residue and municipal solid waste (MSW). Boxplots show the mean, median, interquartile range and range within 1.5IQR. The solid circles display the spread of measured emission factors by fuel type. The zoomed-in green region given in Fig. 8b specifically focuses on the variability in emission factors of individual species of fuelwood, which has been explored in detail due to the large number of samples. Repeat samples collected from the same location are shaded in grey. For fuelwood, measured NMVOC emission factors varied by over a factor of 20 between 4.3–96.7 g kg\(^{-1}\). The NMVOC emission factors showed a right-skewed distribution with a median of 11.7 g kg\(^{-1}\), mean of 18.7 g kg\(^{-1}\) and an interquartile range of 15.3 g kg\(^{-1}\). For repeat measurements of identical species of fuelwood collected at the same location, except for Ficus religiosa, measured emission factors from repeat experiments varied over a much smaller range, by up to a factor of 2.3. Variation between emissions from these samples were likely due to different moisture contents of actual samples measured and the specific combustion conditions of individual burns. The large variation observed for Ficus religiosa was likely due to the samples being significantly different in terms of composition. Despite the samples for Holoptelea spp. and Eucalyptus spp. coming from different locations, emission factors for these samples were quite reproducible and only varied by a factor of 1.2–1.5. For the remaining identical species of fuelwood collected from different locations, emission factors varied over a much larger range by factors of ~2–9.

For the crop residue species studied here, NMVOC emissions were right skewed with a median of 29.5 g kg\(^{-1}\), which was less than the mean of 37.9 g kg\(^{-1}\) and varied from 8.9–73.8 g kg\(^{-1}\) with an interquartile range of 53.9 g kg\(^{-1}\). Cocos nucifera and Solanum melongena were repeat measurements of fuel collected from the same location and varied by factors of 1.8–2. NMVOC emissions from Brassica spp. fuel, which was collected from different locations, varied by a factor of ~8. Cow dung cake and MSW samples were all collected from different locations and varied by up to factors of 2.4 and 2.1, respectively.

Figure 9a shows the mean total emissions measured in this study for different fuel types split by functionality. Large variability in total measured emissions were observed for fuelwood, with emission factors from individual burns varying by a factor of ~20. Figure 9b shows the mean

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emissions by functionality as a proportion of total measured emissions averaged by overall fuel type. Oxygenates were the largest emission (33 %–55 %), followed by furanic compounds (16 %–21 %), phenolics (6 %–12 %) and aromatics (2 %–9 %) for all fuel types except LPG. LPG emissions were mainly alkanes, with a small emission of furanic species. These have previously been reported to be produced in hydrocarbon flames (Johansson et al., 2016).

Figure 9a and b also show the amount of NMVOCs which remained unidentified (black). On average 94 % of all measured NMVOCs emitted across all burns were speciated. Speciation was greater than 90 % for all sample types, except Vachellia spp. due to several large unidentified peaks (see Sect. S6). Mean speciation by fuel type was between 93 %–96 % for all other fuels, except LPG where speciation was > 99 %.

3.5 Emission ratios

The ratio of the mixing ratios of NMVOCs in the emitted gas can be a useful indicator of their source(s) in ambient air. Ratios can be specific to sources and can allow one source to be distinguished from another. The ratio of i-/ n-pentane can be a useful indicator of whether emissions are anthropogenic or from biomass burning, with a ratio 2.2–3.8 indicative of vehicular emissions, 0.8–0.9 for natural gas drilling, 1.8–4.6 for evaporative fuel emissions and < 1 from burning (Stewart et al., 2021a). Benzene/toluene ratios can also be useful and have been reported from traffic exhaust to be around 0.3 (Hedberg et al., 2002).

\[ \frac{i-/n\text{-pentane}}{} \]

\[ \frac{\text{Benzene/toluene}}{} \]

\[ i-/ \text{n-Pentane indicator ratios have been evaluated for fuelwood sources, propane/butane ratios for LPG, and benzene/toluene ratios for fuelwood and cow dung cake (see Fig. 10). The range of values for multiple different burns has been evaluated rather than just reporting mean and median ratios. The median of } i-/ \text{n-pentane ratios from biomass samples measured during this study was } \sim 0.7 \text{ (see Fig. 10). The mean ratio was } \sim 1.0, \text{ with an interquartile range (IQR) of } \sim 0.5–1.5, \text{ which suggests caution is required when assigning burning sources based on emission ratios due to considerable variability. Despite this, the ratio from solid-fuel combustion sources was often less than expected from petrol emissions. The mean ratio of propane/butane from LPG burning was measured to be } \sim 3.1. \text{ The ratios of benzene/toluene varied considerably between different sources and was measured for fuelwood combustion (2.3), cow dung cake combustion (0.94), petrol liquid fuel (0.40) and diesel liquid fuel (0.20). The range of benzene/toluene ratios for fuelwood was large, with an IQR of } \sim 1.5–2.8 \text{ and the range within 1.5IQR shown by the whiskers in Fig. 10 from } \sim 0.9–4.2. \text{ Despite the variability of ratios from specific source types, the considerable range of benzene/toluene ratios could potentially be a useful indicator of the origin of unaged (fresh) ambient emissions in Delhi. However, further study would be required to assess whether these ratios were also true in the exhaust of petrol and diesel vehicles in India or just limited to fugitive emissions. These findings agree well} \]

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with literature which report mean benzene/toluene ratios of 1.4–5.0 from fuelwood and 0.3 from automotive emissions (Hedberg et al., 2002), indicating that on average biomass burning releases a greater molar ratio of benzene than toluene when compared to automotive emissions.

4 Conclusions

This study is based on comprehensive measurements of NMVOC emissions using a range of detailed and complementary techniques across a large range of functionalities and volatilities. It presents detailed burning emission factors for different NMVOCs from a range of fuels used in the Delhi area of India for residential combustion. This work allows for a better understanding of the impact of residential combustion on air quality and shows that fuelwood, cow dung cake and municipal solid waste burning sources release significantly more NMVOCs than LPG.

A range of areas where future studies are required to better improve and understand emissions from burning are highlighted here:

1. Better understanding of stove burn conditions on emissions

   The impact of stove conditions on NMVOC emissions remains poorly understood. Experiments in this study were carried out using expert local judgement to attempt to ensure that laboratory conditions reflected real-world burning conditions. A range of stoves are used in India for combustion of local fuels, such as chulha and angithi stoves, and an evaluation of the impact of these on emissions and their relative use and spatial distribution requires further study.

2. Better understanding of the effect of moisture content on modified combustion efficiency

   Fuels in this study were collected and stored in a manner designed to be reflective of local practices to ensure that laboratory combustion conditions, and in turn emissions, reflected local burning practices. Future studies should conduct detailed compositional analysis of fuel types and moisture content prior to burning. These studies should also measure CO and CO$_2$ to allow for an evaluation of the impact of modified combustion efficiency on emissions from different fuel types.

3. Limited measurements of some fuel types

   Few measurements were made from domestic, commercial and industrial waste, and the emission factors measured in this study were higher than those observed in previous studies. The effect of moisture content on waste burning has been suggested to impact emissions of particulate matter by around an order of magnitude (Jayarathne et al., 2018). Furthermore, only one LPG stove was used to evaluate emissions from this fuel source, with emissions likely to vary by the type of burner used. Future studies should also make more measurements from waste burning to better understand the effect of composition on emissions. Comprehensive measurements should also be made of emissions from combustion of a range of additional crop residues, as these are an important NMVOC source in India (Jain et al., 2014).

4. Evaluation of the impact on O$_3$ and SOA production as well as the toxicity of emissions

   Better understanding of the drivers of photochemical O$_3$ and SOA production from burning emissions is required. A large variety of high-molecular-weight species with likely low volatilities, such as phenolic and furanic compounds, were released from burning. These NMVOCs are expected to have a large influence on subsequent atmospheric chemistry, and a detailed understanding of this chemistry is required to truly assess the impact of biomass burning on air quality. The SOA production potential and OH reactivity of emissions from these fuels are examined as part of a further study (Stewart et al., 2021b).

5. Evaluation of the relative importance of fuel types to air quality in India

   Detailed evaluation of fuel use across India is required to evaluate the relative impact of emissions from fuelwood, municipal solid waste, cow dung cakes and LPG.
The emission factors measured for cow dung cake and municipal solid waste in this study were much higher than for fuelwood and LPG and indicated that these sources are likely to contribute significantly to poor air quality.

The comprehensive characterisation of emissions from fuel types in this study should be used to produce spatially disaggregated local emission inventories to provide better inputs into regional policy and global chemical transport models. This should allow for a better understanding of the key drivers of poor air quality in India and could allow for meaningful mitigation strategies to alleviate the poor air quality observed.

Data availability. The emission factors measured as part of this study are provided in the Supplement.

Video supplement. An illustrative example of the burn conditions used in this study are given in the video supplement at https://doi.org/10.5446/50203 (Stewart, 2020).

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/acp-21-2383-2021-supplement.

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