





duction exceeds that of destruction ( $P_{\text{OH}}/D_{\text{OH}} > 1$ ) suggesting uncertainties in either the chemistry used in the model or in the measurements of the precursors of OH under elevated levels of  $\text{NO}_x$  which lead to an overestimation of  $P_{\text{OH}}$ .

OH reactivity measurements themselves are often compared to calculated values simply using Eq. (1) with  $[X_i]$  restricted to those species which are measured in the field, and with knowledge of laboratory kinetic data. In some cases comparison is also made with modelled OH reactivity which also includes unmeasured OH sinks which are often oxygenated VOCs formed following the oxidative degradation of directly emitted hydrocarbon species. Measurements of OH reactivity are typically higher than those predicted (Lou et al., 2010), with the difference between the measured OH reactivity and that either calculated or modelled referred to as the missing OH reactivity. Analysis of the magnitude of this disagreement when calculated and modelled OH reactivity are used to compare with field determined  $k_{\text{OH}}$ , enables some distinction to be made between the fraction of missing OH reactivity due to missing primary emissions and that due to unmeasured oxidised intermediates.

Missing reactivity is often largest in forested environments, e.g. Edwards et al. (2013), with both missing primary emissions (Di Carlo et al., 2004) and oxidised intermediates (Edwards et al., 2013) considered as potential contributors. In contrast, analysis made during a number of urban studies has demonstrated good agreement between measured OH reactivity and that calculated using Eq. (1) from individually measured OH sinks (Ren et al., 2003; Di Carlo et al., 2004; Mao et al., 2010). In almost all urban studies to date, the measured OH reactivity has been compared to calculated OH reactivity only rather than modelled reactivity; the latter would include an estimate of the contribution that model-generated intermediates make to the total loss rate. The fact that closure has previously been obtained using calculated reactivity from primary emissions alone suggests that oxidised intermediates do not always contribute significantly to the total OH reactivity in certain cities. This is not the case at all urban sites, for example Chatani et al. (2009) reported a missing reactivity of  $\sim 30\%$  in Tokyo during the summer determined by comparing the measured OH reactivity with that calculated

31251

from measured sinks, and concluded that oxidised intermediates likely contributed to the missing fraction. Similarly, Dolgorouky et al. (2012) reported a missing reactivity of 75 % when continental air was sampled in Paris during the wintertime MEGAPOLI project and only measured sinks were used to calculate  $k_{\text{OH}}$ . The closure of the OH reactivity budget in certain cities such as Houston (Mao et al., 2010) and New York (Ren et al., 2003) may reflect the dominance of  $\text{NO}_x$  acting as main OH sink; a species from which no model-generated intermediates will derive (in New York the reaction of OH with NO and  $\text{NO}_2$  accounted for 50 % of the total OH reactivity, Ren et al., 2003). The VOC classes which dominate in different cities may also be substantially different (and, hence, their propensity to generate intermediates may differ). For example, aromatics account for up to 15 % of the total reactivity in Houston (Mao et al., 2010) where local petrochemical industries can influence the local air-mass composition. In contrast, aromatics accounted for just 5 % of the total reactivity in Tokyo (Yoshino et al., 2012) during the summer of 2007. During the PRIDE-PRD campaign in southern China, OH reactivity was modelled using the Regional Atmospheric Chemistry Model (RACM) and, alongside calculated OH reactivity from measured sinks, was compared to measured OH reactivity. Lou et al. (2010) found that the calculated OH reactivity was a factor of two less than the measured value, whereas the RACM modelled OH reactivity reproduced the observations well, suggesting that unmeasured secondary species (mainly thought to be OVOCs) do make a significant contribution to the total observed OH reactivity. During the daytime the PRIDE-PRD site was heavily influenced by biogenic emissions, in contrast to some of the urban sites discussed above, which may in part explain the differences of the missing OH reactivity reported.

Care is needed when comparing the missing OH reactivity reported in the literature, as the chemical detail and range of measured VOCs and inorganic (e.g.,  $\text{NO}_x$ , CO,  $\text{O}_3$ ) species which have been used to calculate OH reactivity varies significantly between projects. Oxygenated VOCs or biogenic VOCs (in addition to isoprene) are often measured, although this is not always the case as in the urban studies reported by Lou

31252

et al. (2010), Kovacs et al. (2003), Ren et al. (2003), Mao et al. (2010), Dolgorouky et al. (2012).

This paper presents an extensive dataset of OH reactivity measured in London during the summer of 2012 as part of the Clean air for London (ClearLo) project (Bohnenstengel et al., 2015). Alongside OH reactivity, an extremely comprehensive speciated VOC dataset was measured and used to constrain a zero dimensional box model based on the Master Chemical Mechanism v3.2 in order to model OH reactivity. By comparing modelled and measured reactivity the level of missing OH reactivity is quantified. By also constraining the model with a subset of the VOCs comprising the more commonly reported C<sub>2</sub>-C<sub>8</sub> VOCs (measured using a dual-channel gas chromatography system), we assess which classes of VOC are controlling the OH reactivity in London. Using the model we also investigate the impact of various classes of VOC on the concentration of peroxy radical species and consequently on the magnitude of in situ O<sub>3</sub> production.

## 2 Experimental

### 2.1 Site description

The ClearLo campaign ran from 22 July to 18 August and overlapped with the London 2012 summer Olympics. An extensive suite of instrumentation was deployed and operated from the grounds of Sion Manning School in North Kensington (51°31'61" N, 0°12'48" W), which is located adjacent to a long-term air quality monitoring site in North Kensington (Bigi and Harrison, 2010). Further details on the campaign and location may be found in the ClearLo overview paper (Bohnenstengel et al., 2015). The inlets for the OH reactivity instrument and those for measuring OH sinks were within 5 m of one another horizontally, and 1 m vertically.

31253

### 2.2 OH reactivity measurements

OH reactivity measurements were made using the laser flash photolysis pump-probe technique (Sadanaga et al., 2004), and the instrument is described in detail in Stone et al. (2015). Ambient air was drawn into a reaction cell (85 cm in length; 5 cm internal diameter), at a flow rate of 12 slm, by an extraction fan situated at the far end of the reaction cell to the inlet. A flow of 0.5 slm of humidified ultra-high purity air (BTCA 178, BOC Special Gases) was passed across a low pressure Hg lamp, generating ~ 50 ppb O<sub>3</sub>, and mixed with the ambient air flow. Laser photolysis of O<sub>3</sub> at 266 nm gave uniform production of OH along the length of the reaction cell following the reaction of O(<sup>1</sup>D) with water vapour.

Changes in OH radical concentrations owing to pseudo-first-order loss with species present in ambient air were monitored by sampling air from the reaction cell through a 0.8 mm pin-hole into a FAGE (fluorescence assay by gas expansion) detection cell maintained at a pressure of ~ 1.5 Torr by a Roots blower backed by a Rotary pump. The 308 nm probe laser, operating at a pulse-repetition-frequency of 5 kHz, was passed across the gas flow in the FAGE cell to excite OH radicals, with the subsequent laser-induced fluorescence (LIF) signal at ~ 308 nm detected by a gated channel photomultiplier tube (CPM). The OH decay profile owing to reactions with species in ambient air is thus detected in real time.

Decay profiles were averaged for 5 min periods, and fitted to a first-order rate equation to find the rate coefficient describing the observed loss of OH ( $k_{\text{loss}}$ ). The OH reactivity,  $k_{\text{OH}}$ , was determined by subtracting the rate coefficient describing physical losses of OH ( $k_{\text{phys}}$ ) from the observed loss of OH in ambient air, with a small correction to account for the dilution of ambient air with the small flow of O<sub>3</sub>-containing humidified ultra-high purity air. The rate coefficient,  $k_{\text{phys}}$ , was determined by monitoring the loss of OH when sampling ultra-high purity air (BTCA 178, BOC Special Gases, passed through scrubbers to remove trace amounts of NO<sub>x</sub>, H<sub>2</sub>, CO and CO<sub>2</sub>) con-

31254

taining the same small flow of O<sub>3</sub>-containing humidified ultra-high purity air as for the measurements sampling ambient air, and was found to be  $1.1 \pm 1.0 \text{ s}^{-1}$ .

The accuracy of the OH reactivity measurements were verified in the laboratory by measuring the bimolecular rate coefficients for reactions of OH with CO and CH<sub>4</sub> under pseudo-first-order conditions at 293 K. For reaction of OH with CO, a bimolecular rate coefficient of  $(2.4 \pm 0.2) \times 10^{-13} \text{ cm}^3 \text{ s}^{-1}$  was determined, in agreement with the literature value of  $2.3 \times 10^{-13} \text{ cm}^3 \text{ s}^{-1}$  (Atkinson et al., 2006), and for OH + CH<sub>4</sub>, a rate coefficient of  $(5.5 \pm 0.6) \times 10^{-15} \text{ cm}^3 \text{ s}^{-1}$  was determined, in agreement with the literature value of  $5.8 \times 10^{-15} \text{ cm}^3 \text{ s}^{-1}$  (Atkinson et al., 2006). The  $1\sigma$  uncertainty of the measurements was, on average,  $\pm 3.2 \text{ s}^{-1}$  reflecting the combined uncertainties from the fits of the observed OH decays, determinations of  $k_{\text{phys}}$  and uncertainties in the dilution factor.

### 2.3 VOC measurements

Two gas chromatography (GC) instruments were deployed as part of the summer ClearLo campaign for in situ VOC measurements: a dual channel GC, (DC)-GC-FID, measuring the very volatile VOCs (C<sub>2</sub>-C<sub>8</sub> hydrocarbons and a small selection of OVOCs), and a comprehensive two-dimensional GC (GCxGC-FID) which measured the less volatile fraction, (C<sub>6</sub>-C<sub>13</sub> and a large group of OVOCs). The details of these two instruments and the time-series of the VOC measurements during ClearLo are presented in Dunmore et al. (2015). A total of 64 VOCs were individually identified and quantified and used to constrain a zero dimensional box model; these are listed in Table 1. Many hundreds more were included in a carbon number and functionality grouping in an attempt to quantify as much of the reactive carbon loading as possible (Dunmore et al., 2015).

### 2.4 Radical measurements

OH, HO<sub>2</sub> and RO<sub>2</sub> radical concentrations were measured using the FAGE technique (Stone et al., 2012). In the case of HO<sub>2</sub> and RO<sub>2</sub>, the radicals were first titrated with

31255

NO to OH before FAGE detection. The current mode of operation is described further in Whalley et al. (2013), including details of the ROxLIF method used to determine RO<sub>2</sub>. Two measurement axes were operated simultaneously during the ClearLo deployment. In the first, air was sampled through a 1 mm diameter pin-hole into a FAGE cell which was dedicated to sequential measurements of OH and HO<sub>2</sub>. In the second axis, air was sampled, again through a 1 mm diameter pinhole, into an 83 cm long flow reactor coupled to a second FAGE cell via a 4 mm diameter pinhole, which permitted detection of the sum of all RO<sub>2</sub> radicals (Whalley et al., 2013). Both detection FAGE cells were held at 1.1 Torr, whilst the flow reactor, which was differentially pumped, was held at  $\sim 30$  Torr. A fast-flow rate through each fluorescence cell was achieved using a Roots blower backed by a Rotary pump. The two fluorescence cells were joined by a tube containing a quartz window, and so shared a single laser optical axis. The instrument was calibrated twice weekly on average using photolysis of a known concentration of water vapour at 185 nm within a turbulent flow tube to generate OH and HO<sub>2</sub>, with the product of the photon flux at 185 nm and the water vapour photolysis residence time measured using a chemical actinometer (Commane et al., 2010). For calibration of RO<sub>2</sub>, methane (BOC, CP grade, 99.5%) was added to the humidified air flow in sufficient quantity to rapidly convert OH to CH<sub>3</sub>O<sub>2</sub>. The limit of detection (LOD) at a signal-to-noise ratio of one for one data acquisition cycle lasting 7 min was  $\sim 4.5 \times 10^5 \text{ molecule cm}^{-3}$  for OH,  $\sim 2.1 \times 10^6 \text{ molecule cm}^{-3}$  for HO<sub>2</sub> and  $\sim 6.9 \times 10^6 \text{ molecule cm}^{-3}$  for CH<sub>3</sub>O<sub>2</sub> at a typical laser power of 13 mW in each cell. The measurements were recorded with 1 s time-resolution, and the accuracy of the measurements was  $\sim 26\%$  ( $2\sigma$ ). The HO<sub>2</sub> measurements were subject to a small interference from RO<sub>2</sub> (typically  $< 3\%$  for all RO<sub>2</sub>), which was quantified as described in Whalley et al. (2013).

### 2.5 Model description

A zero dimensional box model, utilising a subset of the chemistry described within the Master Chemical Mechanism MCMv3.2 (Jenkin et al., 2012), has been used to predict OH reactivity for comparison with the measured reactivity. In its entirety, the MCM

31256

treats the degradation of 135 VOCs following oxidation by OH, O<sub>3</sub> and NO<sub>3</sub>, and for alkanes only, oxidation by Cl atoms, and contains ~ 6700 species and ~ 17 000 reactions. Complete details of the kinetic and photochemical data used in the mechanism are available at the MCM website (MCM, <http://mcm.leeds.ac.uk/MCM/home>).

5 The model was constrained to measurements of NO, NO<sub>2</sub>, O<sub>3</sub>, HONO, CO, CH<sub>4</sub>, PAN, HCHO, VOCs (details given in Table 1), water vapour, temperature, pressure, photolysis frequencies (using actinic fluxes measured by a spectral radiometer) and aerosol surface area. A constant H<sub>2</sub> concentration of 500 ppbv was assumed.

10 The model inputs were updated every 15 min. For species measured more frequently, data were averaged to 15 min intervals, whilst those measured at a lower time resolution, for example the VOCs (Table 1), were interpolated. By this method, a model time-series was produced which could be directly compared with the OH reactivity and radical observations and from which diurnal averages were also generated.

15 The loss of all non-constrained, model generated species by deposition was represented by a deposition velocity equivalent to 1 cm s<sup>-1</sup>. The mixing height was constrained to measurements made by a LIDAR instrument (Barlow et al., 2011) and was observed to rapidly increase after sunrise up to a height of ~ 1800 m from a minimum of ~ 330 m at ~ 4 a.m.

20 The model was run for the entirety of the campaign in overlapping 7 day segments. To allow all the unmeasured, model generated intermediate species time to reach steady state concentrations the model was initialised with inputs from the first measurement day (22 July 2012) for 5 days before comparison to measurements were made. Comparison of these 5 spin-up days demonstrated that the concentration of model generated species rapidly converged and there was less than a 1 % difference in (for example) modelled OH concentration by the second spin up day. As a result of this, the model segments were run so as to overlap for 2 days only to reduce the computing time.

31257

### 3 Results

Near continuous OH reactivity measurements were made in London from 22 July to 17 August 2012 and a comprehensive suite of individually measured inorganic and organic OH sink species were made alongside, as detailed in Table 1. The OH reactivity time-series is presented in Fig. 1. Typically, South Westerly winds ranging from less than 1 m s<sup>-1</sup> during the night to between 4–6 m s<sup>-1</sup> in the afternoon were encountered. However, close to the start of the campaign (24–27 July) and also later in the campaign (8–10 August), the wind direction switched to an Easterly flow, bringing air that had passed over Central London to the site, and wind speeds dropped. Fine weather prevailed during these Easterly flows, with enhancements in air temperature and solar radiation observed (Bohnenstengel et al., 2015). During these periods, radical concentrations were elevated, as were the concentration of a number of other species such as NO<sub>x</sub>, formaldehyde, ozone and many of the individually detected VOCs. OH reactivity was also observed to increase during the Easterly flows with a peak reactivity of 116 s<sup>-1</sup> recorded on the morning (7 a.m., GMT, which is local time (BST) – 1 h) of the 24 July.

On all days a distinct diurnal trend was apparent in OH reactivity, which peaked between 6 and 7 a.m. (GMT) and then dropped to a minimum during the afternoon as shown in Fig. 2. A broader, less distinct secondary peak in reactivity was observed after 6 p.m. continuing throughout the night. This trend is typical of the OH reactivity profile observed at urban sites (e.g. Dolgorouky et al., 2012; Sadanaga et al., 2005; Chatani et al., 2009; Lou et al., 2010) and closely follows the observed NO<sub>x</sub> profile. The mean reactivity for the campaign was 18.1 s<sup>-1</sup> and on average the reaction of OH with NO and NO<sub>2</sub> accounted for 4.5 s<sup>-1</sup> (25 %) of the reactivity observed. The reaction with CO accounted for a further 1.3 s<sup>-1</sup> (7 %) and so it is evident, by considering the reactivity that remains unaccounted for, that the combined contribution made by organic compounds must be significant. The model was initially run constrained to the standard set of VOCs measured by the (DC)-GC-FID. The rate of loss of OH by reaction with

31258

each individual VOC compound is extremely small, so by grouping individual VOCs detected by the (DC)-GC-FID by class (alcohol, alkanes, alkenes and alkynes, aromatics, carbonyls and dialkenes) the influence of different VOC types on the removal of OH and hence their influence on the oxidising capacity can be evaluated. Using this approach, for the carbonyl class of VOCs the reactivity towards OH is dominated by formaldehyde (40%) and acetaldehyde (54%), and when considered as a whole, makes a similar contribution as NO<sub>2</sub> to the total OH reactivity. The other organic classes contribute less to total reactivity (between 0.2–0.6 s<sup>-1</sup> on average), with reactivity from most of the classes exhibiting a weak diurnal profile similar to that of NO<sub>x</sub>. In contrast, the contribution to OH reactivity made by the dialkene class, which itself is dominated by isoprene, peaks at noon suggesting a dominance of a temperature/sunlight driven biogenic signature as opposed to a traffic signature for this class.

Combined, the contribution of NO<sub>x</sub>, CO and all measured organic classes cannot reconcile the OH reactivity observed, with an average missing reactivity of 6.7 s<sup>-1</sup> which equates to 37% of the total measured value. Even when model-generated intermediates are considered, which are made up largely by oxygenated compounds such as methylglyoxal, glyoxal and glycoaldehyde which derive from aromatic species, isoprene and methyl vinyl ketone, on average 6.0 s<sup>-1</sup> of OH reactivity (33% of the total measured) remains unaccounted for, as shown in Fig. 2.

An extended range of VOCs (24 additional compounds), consisting of *n*-alkanes, substituted aromatic and carbonyl compounds and the monoterpenes of  $\alpha$  pinene and limonene, were detected using a GCxGC-FID instrument. The missing reactivity determined in the modelling exercise above (which included only VOCs from the (DC)-GC-FID) may derive from these additional VOCs which, historically, are not typically detected by traditional GC methodologies. As shown in Fig. 3, the reaction of OH with the biogenic species of  $\alpha$  pinene and limonene alone (not including any model-derived intermediates) increases the total calculated OH reactivity by 0.25 s<sup>-1</sup>, and similar increases are calculated by inclusion of the additional *n*-alkanes detected (Table 1). Overall, however, calculated OH reactivity increases by just 0.7 s<sup>-1</sup> on average when

31259

these additional contributions are considered without including model-derived intermediates. Interestingly, the contribution that the model-derived intermediates make to OH reactivity when the model is run constrained to the extended VOC suite is much more significant than previously presented in Fig. 2. When combined, the model-generated intermediates contribute 3.2 s<sup>-1</sup> on average to the OH reactivity in the model run constrained to the extended VOC measurements, compared to just 0.7 s<sup>-1</sup> in the model run constrained just with the standard VOC measurements.

In total there are close to 2600 reactions included in the extended VOC model run which destroy OH and the reaction of OH with the different model-generated intermediate species accounts for more than 90% of these reactions. Pinonaldehyde (an oxidation product of  $\alpha$  pinene formed both during ozonolysis and OH-initiated oxidation reactions) is the most destructive model intermediate species with respect to OH and this single species is found to contribute approximately 7.5% to the total OH reactivity of the modelled intermediates at noon. To determine the absolute impact of the biogenic species on OH reactivity, a model was run constrained to all (DC)-GC-FID and GCxGC-FID VOC measurements apart from  $\alpha$  pinene and limonene. The model-generated intermediates from this run contributed 1.04 s<sup>-1</sup> to total OH reactivity on average, demonstrating that although the contribution to total OH reactivity from the measured biogenic class (considering only reaction with the parent VOCs) was small (0.25 s<sup>-1</sup>), the oxidation products from just  $\alpha$  pinene and limonene alone account for 2.2 s<sup>-1</sup> of the OH reactivity. The total concentration of C<sub>9</sub>–C<sub>12</sub> alkanes is ~5 times greater than the combined concentration of the biogenics on average. Despite this, the model intermediates generated during the oxidation of these larger alkanes (C<sub>9</sub>–C<sub>12</sub>) which derive from diesel emissions (Dunmore et al., 2015), and the intermediates generated from the additional aromatic and carbonyl species detected by GCxGC-FID, contribute only 0.34 s<sup>-1</sup> to total reactivity on average. The rate of reaction of OH with  $\alpha$  pinene ( $5 \times 10^{-11}$  cm<sup>3</sup> molecule<sup>-1</sup> s<sup>-1</sup> at 298 K) is approximately 4.5 times faster (and the reaction with limonene ( $1.6 \times 10^{-10}$  cm<sup>3</sup> molecule<sup>-1</sup> s<sup>-1</sup> at 298 K) is an order of magnitude faster) than the analogous reactions with the C<sub>9</sub>–C<sub>12</sub>

31260

alkanes ( $\sim 1.2 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$  at 298 K). Furthermore, the initial reaction of these biogenic species with ozone can also contribute to the formation of the biogenic-derived model intermediates species. The biogenic intermediates themselves remain highly reactive towards OH, with the rate coefficient for the reaction of OH with pinonaldehyde being similar to the rate coefficient for the reaction of OH with  $\alpha$  pinene itself ( $4 \times 10^{-11}$  and  $5 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$  respectively at 298 K). These factors combined lead to the biogenic species strongly influencing OH reactivity.

The contribution that modelled intermediates derived from the larger VOCs (measured by the GCxGC-FID) make to the total OH reactivity is significant, and is able to largely close the gap between modelled and observed OH reactivity, with  $\sim 15\%$  missing reactivity remaining. Good agreement both in terms of the magnitude and variability from day-to-day (Fig. 1) and campaign average diurnal profile (Fig. 3) is achieved by the model when constrained to the extended VOC measurements. This result highlights the impact of biogenic VOCs in London when the oxidised secondary products are considered. The role of biogenic VOCs (excluding isoprene) in increasing the OH reactivity has not previously been demonstrated at urban sites. Often biogenic VOCs other than isoprene are not measured at urban sites when OH reactivity has been reported e.g. (Kovacs et al., 2003; Ren et al., 2003; Mao et al., 2010; Lou et al., 2010; Dolgorouky et al., 2012) and even if they are measured, the contribution they make to total OH reactivity is deemed small if only the reaction of the parent measured biogenic species with OH is considered, e.g. Chatani et al. (2009). The ultimate fate of these oxidised intermediates derived from biogenic VOCs is highly uncertain, but if they remain in the gas-phase, as opposed to partitioning onto particles, their potential to increase the rate of radical propagation and in situ ozone production is significant.

Even when the contribution made to reactivity from the additional VOCs measured by the GCxGC-FID is taken into account, a small missing reactivity ( $\sim 15\%$  on average) remains. This missing reactivity could suggest that the physical loss of model generated intermediates may actually be slower than imposed in the model (see Sect. 3.2) or this under-prediction in OH reactivity could also reflect further missing primary OH sinks,

31261

for example, higher molecular weight carbon ( $\geq \text{C}_{13}$ ) that is not detected by either the (DC)-GC-FID or GCxGC-FID methods employed. It is known that  $\geq \text{C}_{13}$  makes up a large fraction of diesel emissions and diesel-related hydrocarbons have been shown to dominate the gas phase carbon (in terms of mass) in London (Dunmore et al., 2015).

The complexity of London's air can be observed in the GCxGC-FID chromatogram shown in Fig. 4. In addition to the many identified VOCs, there are many others that can be grouped according to chemical class through their position on the chromatogram, but are not identified explicitly. Ten groups are shown in Fig. 4, with areas 1–8 representing branched and cyclic alkanes, area 9 represents unidentified monoterpenes and area 10 is the  $\text{C}_4$ -substituted monoaromatics species. The highly branched alkane species can be incorporated into the model by approximating the reactivity to be the same as for the straight-chain alkane equivalent. For example, for an unidentified branched alkane which contains six carbon atoms it has been estimated that it would have the same reactivity with respect to OH as *n*-hexane (and the same, subsequent, degradation mechanism). For unidentified monoterpenes, a reaction rate coefficient (and the subsequent degradation mechanism) the same as that for the reaction between OH and  $\alpha$  pinene is assumed, whilst for unidentified substituted aromatics the reaction rate coefficient with respect to OH (and degradation mechanism) is taken to be the same as that for propyl-benzene. Inclusion of this additional reactive carbon in the model leads to a  $0.4 \text{ s}^{-1}$  increase in the alkane contribution and only a small increase of  $\sim 0.1 \text{ s}^{-1}$  due to increasing the carbon in the biogenic and aromatic classes combined. Despite the small contribution made by increasing the effective  $\alpha$  pinene present, this increase in biogenic carbon leads to a further  $1.1 \text{ s}^{-1}$  increase in the contribution from model intermediates. Figure 5 compares the total modelled OH reactivity alongside the measured values when these additional unidentified species are included. Overall, this additional carbon (which is known to be present but cannot be fully identified) improves the overall model-to-measured agreement with just 6% of the total reactivity now unaccounted for.

31262

From Fig. 1 it is evident that the largest fraction of missing reactivity occurs during the south-westerly flow conditions encountered between the 28 July–8 August and even considering the  $1\sigma$  uncertainty of the OH reactivity measurements of  $\pm 3.2\text{ s}^{-1}$ , the missing reactivity during this period is significant. The propensity of the model to under-estimate reactivity during the south westerly flow is further highlighted when the modelled averaged diurnal reactivity is taken for south westerly and easterly conditions separately (Fig. 6). The contribution to the total reactivity made by any individual class of compounds does not change significantly between these two regimes, rather, all classes represent bigger OH sinks during the more polluted phase. The diurnal profile is much more distinct during the polluted phase demonstrating a stronger traffic-based source, likely in combination with more significant boundary layer height changes over the course of the day during low wind speed periods. There is a very slight over-prediction during the Easterly flow in OH reactivity during the afternoon and during the morning rush-hour, but overall the agreement between model and measurements is excellent during the Easterlies. In contrast, an under-prediction in the observed OH reactivity throughout the day during south-westerly conditions is apparent suggesting either there are more undetected VOCs during south-westerly flows than during the easterly flows or that the physical loss of the model generated intermediates should be treated differently during these two contrasting air-masses. The sensitivity of model predictions to the physical loss term of the model-generated intermediates is investigated in Sect. 3.2.

### 3.1 Ability of the model to reconcile the observed formaldehyde concentration

Thus far, all model runs have been constrained to the observed formaldehyde (HCHO) concentration. HCHO is formed during the oxidation of nearly every VOC and so may be considered a useful target species against which to assess atmospheric oxidation schemes. A model run constrained to (DC)-GC-FID VOCs only, but now unconstrained to HCHO significantly under-predicts (by 60 % on average) the observed HCHO concentration (Fig. 7). The ability of the model to capture the observed HCHO in terms

31263

of magnitude is substantially improved (to with 32 % of the observations) when the model is constrained to the extended VOC suite (including the contribution from the unidentified GCxGC-FID VOC species). This improved predicative capability of the model in terms of HCHO (as for OH reactivity) is almost entirely due to the inclusion of the biogenics. Even constrained to the extended VOC observations (and the unidentified GCxGC-FID VOC species), the model is unable to capture the HCHO concentration observed with the discrepancy greatest during the morning hours. This under-prediction may reflect inaccuracies in the physical deposition of HCHO in the model (discussed further in Sect. 3.2) but may also reflect a missing primary source of HCHO in the model. Similar findings are reported from field observations in the Po Valley in Italy where the discrepancy between modelled and measured OH reactivity was found to be small, but HCHO concentrations were substantially under-predicted by a 1-D model, demonstrating an unidentified non-photochemical ground-level source of HCHO (Kaiser et al., 2015). The unidentified source of HCHO in the Po Valley was postulated to be a direct emission from agricultural land. In London, the missing HCHO peaks at  $\sim 8$  a.m. and may reflect a missing source directly emitted from vehicular exhausts.

### 3.2 Sensitivity of model-generated intermediates to parameterisation of the rate of physical loss

There is considerable uncertainty associated with the concentration of model species which are generated by reaction in the model rather than constrained to observations. Model species can undergo gas-phase reactions, may be lost via wet or dry deposition (at different rates depending on the particular species), be taken up on aerosol surfaces, or if their lifetime is sufficiently long, there may be a net movement of these species either into or out of the model box.

In the model runs considered above, the deposition rate for all intermediate species (including HCHO in the model runs with HCHO unconstrained) was set equal to  $1\text{ cm s}^{-1}$  which is equivalent to a lifetime of  $\sim 27\text{ h}$  in a 1000 m boundary layer. The

31264

boundary layer varied from a minimum of 330 m at night to 1800 m in the afternoon. By treating the modelled physical loss as a function of boundary layer height, a dry deposition of species to the surface is reasonably well represented, but this method does not represent possible ventilation from the model box very well (as wind-speeds picked up during the afternoon hours for example). Setting the boundary layer height in the model to a constant 300 m throughout effectively enhances the physical loss rate of intermediates during the afternoon when the boundary layer was previously set higher. Figure 8 highlights that this does modestly improve the overall modelled agreement with observed reactivity during the afternoon hours, but the reduction in modelled reactivity is small. This demonstrates that although the treatment of physical loss remains a highly uncertain model parameter, the concentration of model intermediates and their impact on modelled reactivity is not particularly sensitive to this term, with large changes in the lifetime of the intermediates (from  $\sim 50$  to  $\sim 8$  h during the afternoon) only leading to a reduction of  $0.6 \text{ s}^{-1}$  (or 4%) in the modelled reactivity at 2 p.m. It should be noted that this enhanced physical loss of intermediates would lead to further disagreement between the modelled and measured OH reactivity during south westerly flow.

For the model to better represent the observed HCHO concentrations, a reduction in the physical loss of this species is needed. Reducing the deposition rate from  $1$  to  $0.3 \text{ cm s}^{-1}$  in the model (Fig. 7) enhances the modelled HCHO leading to an improved model representation of this species at night. The model still under-estimates the concentration of HCHO during the day (total underestimate of 16% on average) which may reflect the uncertainty in the treatment of the physical loss of other model-generated intermediates which go on to form HCHO themselves, or may reflect the small under-estimation of OH reactivity (6%) which remains, or may reflect a small direct emission of HCHO not accounted for by the model (as discussed above). Despite the small under-prediction of [HCHO], this analysis demonstrates that the majority of the HCHO observed (84%) can be accounted for by considering the oxidative degradation of the VOCs detected in London's atmosphere, and the HCHO concentration will be signifi-

31265

cantly under-estimated if the VOC model constraints under-represent the actual VOC composition (particularly, if the contribution of the biogenics is neglected).

### 3.3 Impact of the different VOC constraints on modelled OH

As may be expected, with an increase in the modelled OH reactivity moving from the model run with standard VOC constraints to a run with the extended VOC constraints, the OH concentration predicted by the model is concomitantly reduced, as shown in Fig. 9. However, all model runs described above significantly over-predict the OH concentration that was observed during the campaign. A model constrained to the observed  $\text{HO}_2$  concentration, and also constrained with the extended VOC suite including contributions from unidentified GCxGC-FID compounds is much better able to reproduce the observed OH. This improved agreement may indicate the presence of a missing peroxy radical sink that is not considered by the model, and further discussion of this, together with a detailed account of the model-measurement agreement for OH,  $\text{HO}_2$  and  $\text{RO}_2$  radicals, and the radical budgets, will be the subject of future publications. With a lower modelled daytime OH concentration obtained when the model is constrained to measured  $\text{HO}_2$ , the modelled OH reactivity is reduced due to the competition which exists between the rate of OH initiated production of model generated intermediates and physical loss of these species (Edwards et al., 2013). As shown in Fig. 8, the agreement with the observed OH reactivity during the afternoon is improved somewhat, although, despite reductions in modelled OH concentrations by close to a factor of 3, the modelled OH reactivity is only reduced by  $0.24 \text{ s}^{-1}$  (or 2%) at 2 p.m. (GMT). These results demonstrate that the modelled reactivity is not particularly sensitive to the [OH], perhaps as a result of OH influencing both the rate of production and loss of these intermediates (the latter by further reaction with OH).

31266





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31271

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31273

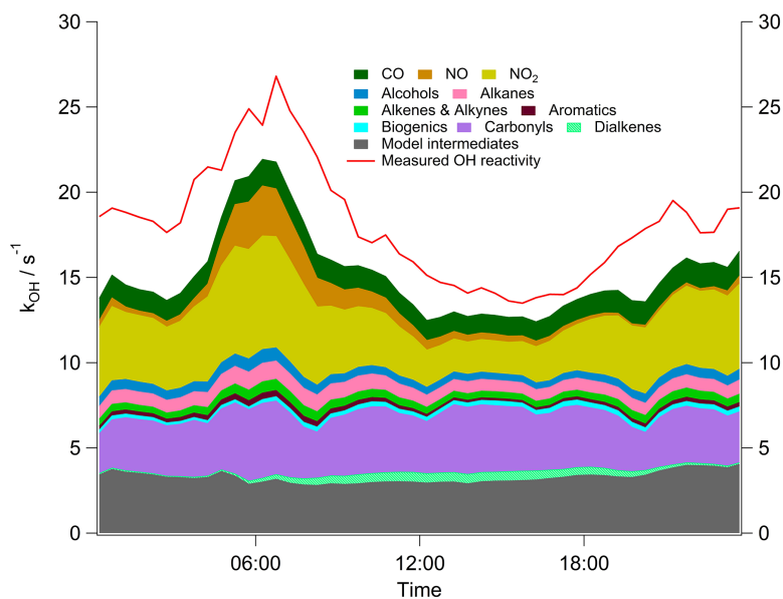
**Table 1.** Listing of the species used as input to the model, the class each species is grouped into, the average contribution (s<sup>-1</sup> and %) of each class to OH reactivity and the detection method for each species. The OH reactivity when the concentration of the unassigned GCxGC-FID peaks is also considered is given in parentheses.

Species (MCM name)	Class	Contribution to OH reactivity (s <sup>-1</sup> )	Contribution to OH reactivity (%)	Instrument
O3	O <sub>3</sub>	0.05 <sup>a</sup>	0.3	Thermo 49 Series
NO	NO	0.76	4.2	Air Quality Design Inc. Chemiluminescence with LED NO <sub>2</sub> converter
NO2	NO <sub>2</sub>	3.82	21.1	
CO	CO	1.31	7.2	Ametek monitor
H2	H <sub>2</sub>	0.08 <sup>a</sup>	0.4	–
HONO	NO <sub>2</sub>	0.04 <sup>a</sup>	0.2	LOPAP
HNO3				CIMS
PAN				GC-ECD
CH3OH	Alcohol	0.54	3.0	(DC)-GC-FID
C2H5OH				(DC)-GC-FID
NPROPOL				(DC)-GC-FID
NBUTOL				(DC)-GC-FID
CH4	Alkane	0.81 (1.22)	4.5 (6.7)	(DC)-GC-FID
C2H6				(DC)-GC-FID
C3H8				(DC)-GC-FID
IC4H10				(DC)-GC-FID
NC4H10				(DC)-GC-FID
IC5H12				(DC)-GC-FID
NC5H12				(DC)-GC-FID
NC6H14				(DC)-GC-FID & GCxGC-FID
NC7H16				(DC)-GC-FID & GCxGC-FID
NC8H18				(DC)-GC-FID & GCxGC-FID
M2PE				(DC)-GC-FID
NC9H20				GCxGC-FID
NC10H22				GCxGC-FID
NC11H24				GCxGC-FID
NC12H26				GCxGC-FID
CH2CL2				GCxGC-FID

31274

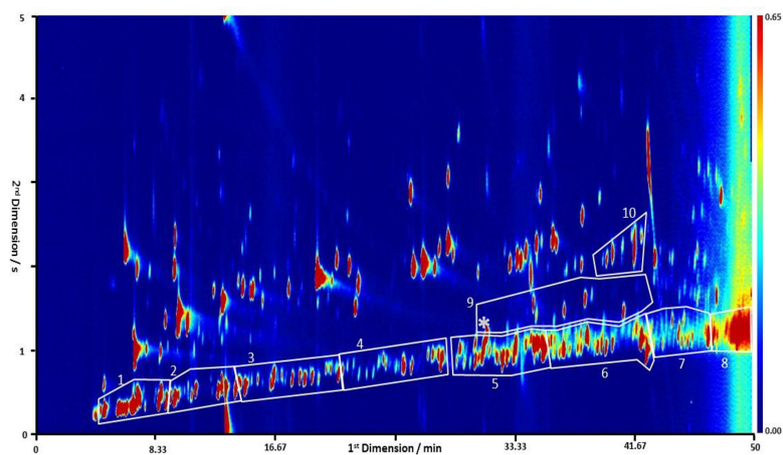






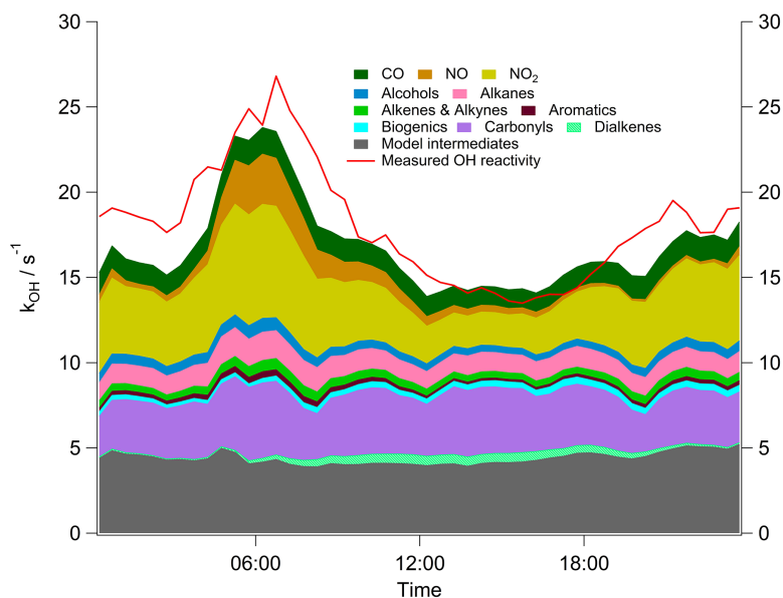
**Figure 3.** Average diurnal profile of measured reactivity and a breakdown of modelled reactivity into inorganic and organic classes when the model is constrained to the standard VOC suite measured by the (DC)-GC-FID and additional VOCs (including  $\alpha$  pinene and limonene, see Table 1 for details) measured with by GCxGC-FID.

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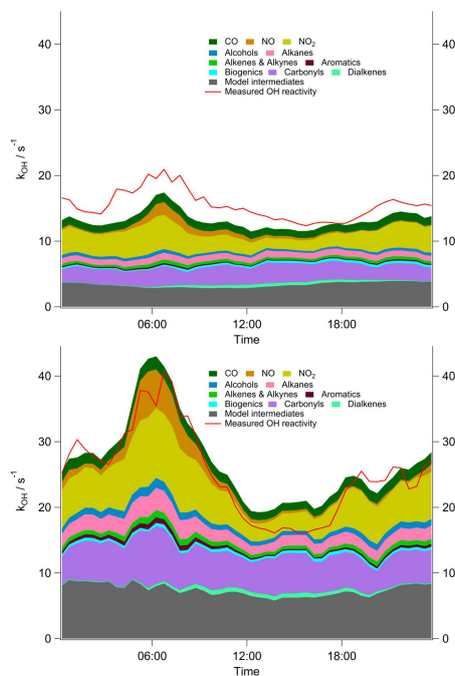
**Figure 4.** A typical GCxGC chromatogram from the summer ClearLo campaign. The retention times from column 1 (separation based on volatility) and column 2 (separation based on polarity) are the  $x$  and  $y$  axis respectively, and compound intensity is the coloured contour. Labelled groups are identified as follows; (1–8) aliphatic groups from  $C_6$  to  $C_{13}$ , (9)  $C_{10}$  monoterpenes with \* corresponding to  $\alpha$  pinene which is the start of that group and (10)  $C_4$  substituted monoaromatics.

31280



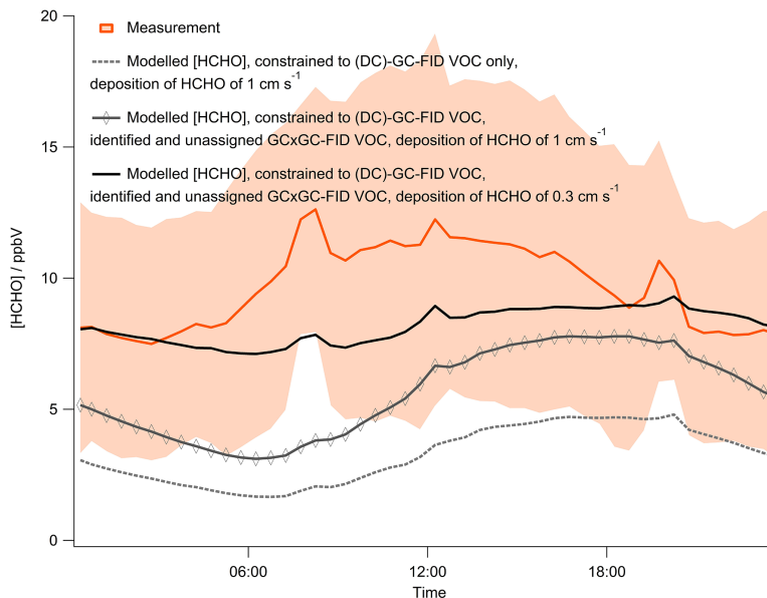
**Figure 5.** Average diurnal profile of measured reactivity and a breakdown of modelled reactivity into inorganic and organic classes when the model is constrained to the standard VOC suite measured by the dual-channel (DC)-GC-FID, the additional identified VOCs (including  $\alpha$  pinene and limonene, see Table 1 for details) measured with by GCxGC-FID and the unassigned GCxGC-FID peaks.

31281



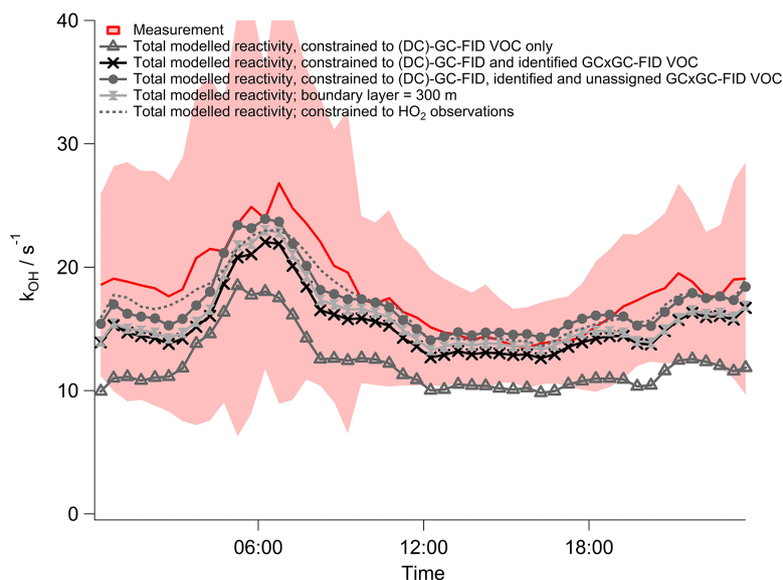
**Figure 6.** Average diurnal profile of measured reactivity and a breakdown of modelled reactivity into inorganic and organic classes when the model is constrained to the standard VOC suite measured by the dual-channel (DC)-GC-FID, additional VOCs (including  $\alpha$  pinene and limonene, see Table 1 for details) measured with by GCxGC-FID and unassigned GCxGC-FID peaks. Top: average diurnal during south westerly flows and Bottom: average diurnal during easterly flows.

31282



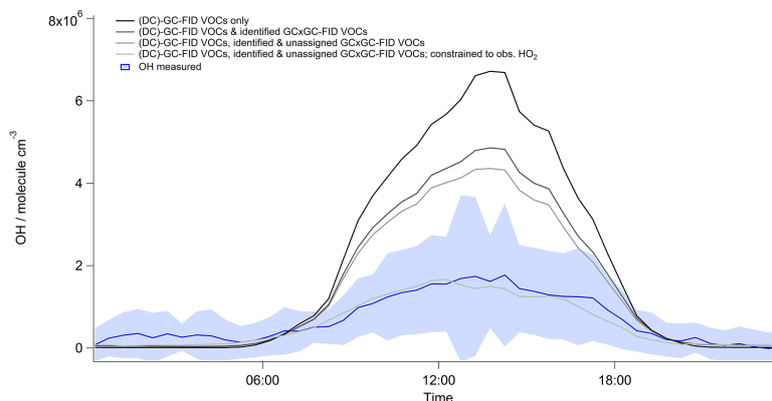
**Figure 7.** Average diurnal profile of the observed HCHO and modelled HCHO for different model scenarios (see text for further details). The shading represents the  $1\sigma$  variability of the measurements; with each data point representing 30 min averaged data.

31283



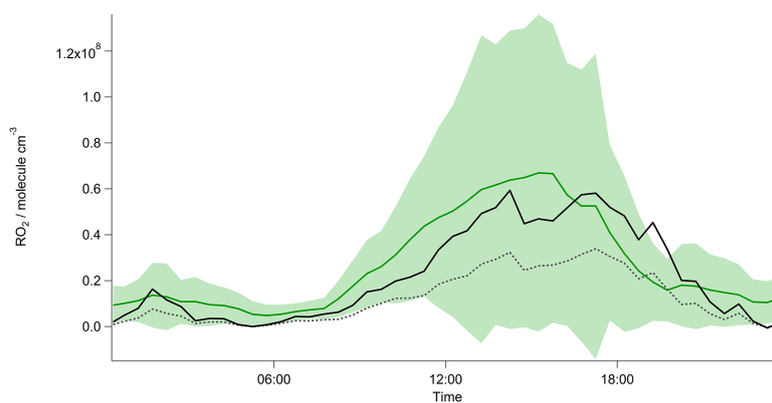
**Figure 8.** Average diurnal profile of the total modelled and measured OH reactivity for different model scenarios (see text for further details). The shading represents the  $1\sigma$  variability of the measurements, with each data point representing 30 min averaged data.

31284



**Figure 9.** Average diurnal profile of modelled and measured [OH]. The shading represents the  $1\sigma$  variability of the measurements; with each data point representing 30 min averaged data. See text for details of the different model scenarios.

31285



**Figure 10.** Average diurnal profile of measured  $RO_2$  (green) and steady state calculated  $RO_2$  using Eq. (3) (black, solid line =  $[RO_2]_{ss}$  derived from the observed total OH reactivity and grey, dashed line =  $[RO_2]_{ss}$  derived from calculated OH reactivity using (DC)-GC-FID VOC only). The shading represents the  $1\sigma$  variability of the measurements; with each data point representing 30 min averaged data.

31286