Evaluating the sensitivity of radical chemistry and ozone formation to ambient VOCs and NO\textsubscript{x} in Beijing

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Evaluating the sensitivity of radical chemistry and ozone formation to ambient VOCs and NO\textsubscript{x} in Beijing

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but only at NO mixing ratios < 1 ppbv. The modelled ROCl atoms, formed from the photolysis of nitryl chloride, enhanced the heterogeneous loss of HO2 and 

Agreement for OH was fortuitous due to a cancellation of other reactions. On other mornings, when the Cl atom concentration was lower, measured RO2 × 10−4 was calculated to exceed 3 ppbv when the Cl atom concentration was determined. The inclusion of Cl atom chemistry did not enhance the modelled ROCl atoms, formed from the photolysis of nitryl chloride, enhanced the heterogeneous loss of HO2 and 

Atmospheric loss of HO2, as assessed from the ratio of 

10 s−1 during the night, with a minimum during the afternoon of ≈ 22 s−1 on average. An experimental budget analysis, in which the rates of production and destruction of the radicals are compared, highlighted that although the sources and sinks of OH were balanced under high NO concentrations, the OH sinks exceeded the known sources (by 15 ppbv h−1) under the very low NO conditions (< 0.5 ppbv) experienced in the afternoons, demonstrating a missing OH source consistent with previous studies under high volatile organic compound (VOC) emissions and low NO loadings. Under the highest NO mixing ratios (104 ppbv), the HO2 production rate exceeded the rate of destruction by ≈ 50 ppbv h−1, whilst the rate of destruction of total RO2 exceeded the production by the same rate, indicating that the net propagation rate of RO2 to HO2 may be substantially slower than assumed. If just 10% of the RO2 radicals propagate to HO2 upon reaction with NO, the HO2 and RO2 budgets could be closed at high NO, but at low NO this lower RO2 to HO2 propagation rate revealed a missing RO2 sink that was similar in magnitude to the missing OH source. A detailed box model that incorporated the latest Master Chemical Mechanism (MCM3.3.1) reproduced the observed OH concentrations well but over-predicted the observed HO2 under low concentrations of NO (< 1 ppbv) and under-predicted RO2 (both the complex RO2 fraction and other RO2 types which we classify as simple RO2) most significantly at the highest NO concentrations. The model also under-predicted the observed k(OH) consistently by ≈ 10 s−1 across all NOx levels, highlighting that the good agreement for OH was fortuitous due to a cancellation of missing OH source and sink terms in its budget. Including heterogeneous loss of HO2 to aerosol surfaces did reduce the modelled HO2 concentrations in line with the observations but only at NO mixing ratios < 0.3 ppbv. The inclusion of Cl atoms, formed from the photolysis of nitryl chloride, enhanced the modelled RO2 concentration on several mornings when the Cl atom concentration was calculated to exceed 1 × 104 atoms cm−3 and could reconcile the modelled and measured RO2 concentrations at these times. However, on other mornings, when the Cl atom concentration was lower, large under-predictions in total RO2 remained. Furthermore, the inclusion of Cl atom chemistry did not enhance the modelled RO2 beyond the first few hours after sunrise and so was unable to resolve the modelled under-prediction in RO2 observed at other times of the day. Model scenarios, in which missing VOC reactivity was included as an additional reaction that converted OH to RO2, highlighted that the modelled OH, HO2 and RO2 concentrations were sensitive to the choice of RO2 product. The level of modelled to measured agreement for HO2 and RO2 (both complex and simple) could be improved if the missing OH reactivity formed a larger RO2 species that was able to undergo reaction with NO, followed by isomerisation reactions reforming other RO2 species, before eventually generating HO2. In this work an α-pinene-derived RO2 species was used as an example. In this simulation, consistent with the experimental budget analysis, the model underestimated the observed OH, indicating a missing OH source. The model uncertainty, with regards to the types of RO2 species present and the radicals they form upon reaction with NO (HO2 directly or another RO2 species), leads to over an order of magnitude less O3 production calculated from the predicted peroxy radicals than calculated from the observed peroxy radicals at the highest NO concentrations. This demonstrates the rate at which the larger RO2 species propagate to HO2, to another RO2 or indeed to OH needs to be understood to accurately simulate the rate of ozone production in environments such as Beijing, where large multifunctional VOCs are likely present.

1 Introduction

Owing to strict emission controls being implemented across China, a reduction in the levels of PM10, PM2.5 and SO2 has been observed in the country since 2013 (Huang et al., 2018). Similar reductions in these primary pollutants are echoed in other countries across the globe. In the United States this reduction in primary emissions is reflected in a ≈ 0.4 ppbv yr−1 reduction in peak O3 (He et al., 2020). In China, however, despite reductions in primary emissions, the concentration of ground-level ozone gradually increased between 2013–2017 (Huang et al., 2018). The highest peak ozone concentrations in China are observed in the Beijing area (T. Wang et al., 2017), where the highest O3 mixing ratio of 286 ppbv was recorded at a rural site 50 km north of the centre (Wang et al., 2006). During the Beijing Olympic Games, despite emission controls, hourly ozone mixing ratios between 160 to 180 ppbv were frequently observed in central Beijing (Wang et al., 2010). Ozone is a secondary pollutant, primarily formed in the troposphere via OH-initiated volatile organic compound (VOC) oxidation in the presence of NOx. O3 concentrations in megacities worldwide frequently exceed regulatory limits during the summer months, with elevated ozone concentrations shown to have negative impacts on human and crop health. The radical species, OH, HO2 and RO2, play a central role in the catalytic photochemical cycle which removes primary emissions and leads to ozone formation. The OH radical initiates the oxidation of VOCs, leading to the formation of peroxy radicals (HO2 and RO2). Peroxy radicals oxidise NO to NO2, which photoly-
ses and generates ozone. Under high NO$_x$ conditions, OH preferentially reacts with NO$_2$, and both peroxy radical production (via VOC oxidation) and, in turn, ozone production decrease. This non-linear relationship between ozone and NO$_x$ complicates efforts to reduce the ambient ozone levels as, in NO$_x$-saturated environments, reductions in NO$_x$ can lead to increases in the rate of ozone production (e.g., Bigi and Harrison, 2010). Furthermore, a number of studies have highlighted that efforts to reduce PM have the potential to exacerbate O$_3$ due to concomitant increases in HO$_2$ caused by a reduction in the heterogeneous loss of HO$_2$ to aerosol surfaces (Li et al., 2019), although there is continued debate on the magnitude of this effect from field studies (Tan et al., 2020). As well as the central role OH plays in photochemical ozone formation, OH promotes the formation of secondary aerosols (sulfate, nitrate and secondary organic aerosols, SOA), which have negative impacts on human health (Chen et al., 2013). Large, complex RO$_2$ radicals are precursors to highly oxidised molecules (HOMs) (Ehn et al., 2014), which have also been shown to condense and contribute to SOA (Mohr et al., 2019). In China, the fraction of PM attributed to secondary aerosols is significant (between 44 %–71 %, Huang et al., 2014), and so understanding the oxidation chemistry which converts primary emissions to secondary aerosols is an ongoing challenge. There has been an increasing growth in photochemical oxidant studies conducted in China, where radical observations have been performed over the past decade, with the PKU and Juelich groups leading these efforts. The first radical observations took place in the summer of 2006, with observations made in the Pearl River Delta (PRD) region (Hofzumahaus et al., 2009; Lu et al., 2012; PRIDE-PRD2006) and also in suburban Beijing (Lu et al., 2013; CARE-Beijing 2006). These campaigns revealed a strong atmospheric oxidation capacity, with elevated levels of OH and HO$_2$ in these regions, with OH concentrations up to 2.6 $\times$ 10$^7$ molecule cm$^{-3}$ and HO$_2$ concentrations up to 2.5 $\times$ 10$^7$ molecule cm$^{-3}$ reported (Lu et al., 2012). Even during the wintertime, under low levels of solar radiation, concentrations of OH can reach 3 $\times$ 10$^6$ molecule cm$^{-3}$ in Beijing (Slater et al., 2020), which is similar to the OH concentrations observed in other urban centres in European cities during the summer months (Whalley et al., 2018). Similar to findings from radical observations and subsequent modelling activities in forested regions (Whalley et al., 2011), which are characterised by high VOC emissions and relatively low NO$_x$ concentrations, the observations and modelling studies in China in summer (Hofzumahaus et al., 2009; Lu et al., 2012; Lu et al., 2013) revealed that the high OH concentrations could only be explained if an additional source of OH, from recycling peroxy radicals to OH, was added to the model. An updated isoprene scheme (Peeters et al., 2009, 2014), which included isomerisation reactions of the isoprene-derived RO$_2$ radicals, was unable to reconcile the OH observations, however. In a subsequent field study conducted in the PRD region (Tan et al., 2019), RO$_2$ observations were made using the RO$_x$ laser-induced fluorescence (LIF) technique alongside OH, HO$_2$ and OH reactivity, allowing an experimental budget analysis for OH, HO$_2$, RO$_2$ and RO$_x$ (OH + HO$_2$ + RO$_2$) to be performed. The analysis demonstrated a missing OH source of 4–6 ppbv h$^{-1}$ and a missing RO$_2$ sink that was similar in magnitude and, hence, supports the hypothesis of a missing mechanism that converts RO$_2$ species to OH under low NO conditions. The authors calculated that the unknown RO$_2$ to OH conversion that does not involve reaction with NO (and, therefore, does not lead to the formation of ozone) reduced ozone production by 30 ppbv d$^{-1}$, demonstrating that knowledge of the branching ratio between the competitive reactions that RO$_2$ radicals undergo (bimolecular reaction with NO or unimolecular isomerisation), as well as the overall VOC oxidation rate, is important when determining in situ ozone production.

In a recent campaign conducted at a rural site in the North China Plain (Tan et al., 2017), during periods in which NO mixing ratios were below 300 pptv, an additional OH recycling mechanism was again needed to reconcile the OH concentrations observed. The modelled RO$_2$ concentrations were in good agreement with those observed under low NO concentrations typically experienced during the afternoon; however, the model under-predicted the RO$_2$ concentrations by a factor of 3–5 at the higher NO mixing ratios ($>$ 1 ppbv) that were observed during the mornings. Additional sources of RO$_2$ from the photolysis of ClNO$_2$ and subsequent reactions of Cl atoms with VOCs, as well as RO$_2$ from the missing reactivity determined, could explain $\approx$ 10 %–20 % of the model under-prediction but could not fully resolve the missing RO$_2$ source of 2 ppbv h$^{-1}$ under the high NO conditions.

As a result, the model was found to under-predict the nitrate in situ chemical ozone production by 20 ppbv d$^{-1}$. In London, during the ClearfLo campaign (Whalley et al., 2018), under higher NO mixing ratios ($>$ 3 ppbv) a box model constrained to the Master Chemical Mechanism (MCM3.2) was found to increasingly under-predict the RO$_2$ concentrations observed with NO$_x$, and, as a consequence, the rate of ozone production calculated from the modelled peroxy radical concentrations was up to an order of magnitude lower than the ozone production rate calculated from the observed peroxy radicals. The model was able to reproduce the observed levels of HO$_2$ under the high NO concentrations but over-predicted HO$_2$ concentrations when NO mixing ratios were below 1 ppbv, and modest under-predictions of OH were observed under low NO conditions, which demonstrated uncertainties in radical cycling at low NO. Conversely, in other urban studies, models were found to increasingly under-predict HO$_2$ as NO$_x$ levels increased beyond $\approx$ 1 ppbv (Martinez et al., 2003; Ren et al., 2013; Brune et al., 2016), although in some of these earlier studies, the HO$_2$ observations may have been influenced by an RO$_2$ interference (Whalley et al., 2013). Understanding the cause of the model failure under different NO regimes in urban centres is critical to be able to accurately

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predict ozone production and to determine ozone abatement strategies that can be implemented to successfully reduce ozone levels. Measurements of OH, HO₂ and RO₂ as well as OH reactivity are necessary to fully explore a model’s skill to capture the entire atmospheric oxidation cycle and to begin to identify mechanisms that can reconcile the concentration of all radical species.

The integrated Study of AIR Pollution PROcesses in Beijing (AIRPRO) project involved two intensive measurement periods that took place in central Beijing during the winter of 2016 and during the following summer of 2017 and was part of the larger Air Pollution and Human Health (APHH) programme. APHH had the overall aim of better understanding the sources, atmospheric transformations and health impacts of air pollutants in Beijing to improve air quality forecasting capabilities (Shi et al., 2019). In this paper the observations of OH, HO₂, RO₂ and OH reactivity from the summer period are compared to a detailed zero dimensional box model run with the latest Master Chemical Mechanism (MCM3.3.1), and an experimental budget analysis is performed on all radical species. The overall objective of this research was to test the model’s ability to reproduce the radical concentrations and, through the budget analysis, investigate the balance between radical production and destruction rates. Following on from the results of earlier radical observation and modelling studies conducted in urban regions, this research will investigate if there are missing radical sources and sinks under different NO regimes and investigate new chemistry that may improve model predictions. We will assess how uncertainties in the model mechanism influence the rate of in situ ozone production in an environment with large and complex VOC emissions and under highly variable NOₓ concentrations.

2 Experimental

2.1 Site description

The observations took place in central Beijing at the Institute of Atmospheric Physics (IAP), which is part of the Chinese Academy of Sciences. The site was located between the third and fourth north ring roads in Beijing and was within 150 m of several busy roads. All instrumentation was located in close proximity within nine shipping containers that were placed on a grassed area surrounding a large (325 m) meteorological tower. Further details of the measurement site and an overview of all the instrumentation that was run during the campaign can be found in Shi et al. (2019).

2.2 FAGE instrumentation

The University of Leeds fluorescence assay by gas expansion (FAGE) instrument was deployed at the IAP site and made measurements of OH, HO₂, RO₂ radicals and OH reactivity ($k$(OH)). The instrumental set-up was analogous to that used during the ClearfLo project (see Whalley et al., 2016, for the $k$(OH) instrument description and Whalley et al., 2018, for the OH, HO₂ and RO₂ instrument details) and also the winter AIRPRO project (Slater et al., 2020) and so is only briefly overviewed here. Two detection cells, the HO₂ cell and the RO₂ LIF cell, were located on the roof of the Leeds FAGE shipping container at a sampling height of 3.5 m. The $k$(OH) instrument, which was housed inside the container, alongside all other FAGE instrument components (including the laser system), drew air from close by the radical detection cells via a 1/2 in. Teflon line. The HO₂ cell made sequential measurements of OH and then the sum of OH + HO₂, by the addition of NO (Messers, 99.95 %), which titrated HO₂ to OH for detection by laser-induced fluorescence (LIF). In the RO₂ LIF reactor, which is an 83 cm long, 6.4 cm internal diameter flow tube, in HO₂ mode, a flow of CO (10 % in N₂) was added just beneath the sampling inlet, and this rapidly converted any ambient OH sampled to HO₂. Within the RO₂ LIF FAGE cell, a continuous flow of NO (99.95 %) titrated ambient HO₂, the converted OH and also a large percentage of complex RO₂ radicals (see below) to OH for detection. In RO₂ mode, a total RO₂ + HO₂ + OH measurement was made by addition of a dilute flow of NO (500 ppmv in N₂) alongside the CO, which promoted the conversion of all HO₂ and RO₂ radicals to OH; the OH formed was rapidly reconverted to HO₂ by reaction with CO. Within the RO₂ LIF FAGE cell, the HO₂ was titrated back to OH, by reaction with NO, for detection. Using the methodology outlined in Whalley et al. (2013), the sensitivity of both the HO₂ and RO₂ LIF FAGE cells towards HO₂ and complex RO₂ species was assessed before the instrument was deployed to Beijing by sampling isoprene-derived RO₂; the sensitivity of the HO₂ cell towards other RO₂ types such as those derived from ethene, methanol and propane has been previously conducted (Whalley et al., 2013) and compared well with model-predicted sensitivities. The sensitivity of the RO₂ LIF instrument has also been assessed previously towards a range of RO₂ types deriving from methane, isoprene, ethene, toluene, butane and cyclohexane and, again, compared well with model-predicted sensitivities (Whalley et al., 2018). “Complex RO₂” refers to any RO₂ species (primarily those derived from alkene and aromatic hydrocarbons) that have the potential to decompose into OH in the presence of NO on the timescale of the FAGE residence time and, therefore, have the potential to act as an HO₂ interference. The NO flow in the HO₂ cell was kept low to minimise the conversion efficiency of complex RO₂ to OH, and the conversion efficiency was found to be < 5 % when isoprene-derived RO₂ radicals were sampled. In the RO₂ LIF FAGE cell, a higher NO flow was employed to promote the conversion of complex RO₂ to OH, enabling 89 % of isoprene-derived RO₂ radicals to be detected. From the relative sensitivities of the two cells to OH, HO₂ and complex RO₂, and by subtraction of complex RO₂ from total RO₂, the concentration of RO₂ species that do not act as an HO₂ interference (“simple RO₂”) has been derived.
For the entirety of the campaign, the HO$_x$ cell was equipped with an inlet pre-injector (IPI; Woodward-Massey et al., 2020), which, by injection of propane into the ambient air stream directly above the HO$_x$ inlet, removes ambient OH and enables a background measurement from laser scatter, solar scatter and detector dark counts (and potentially any cell-generated OH) to be determined whilst the laser is tuned to the OH transition. The subtraction of this background signal from the ambient OH signal provides the OH$_\text{CHEM}$ measurement, which can be compared to the traditional OH$_\text{WAVE}$ measurement in which the background signal (from laser scatter, solar scatter and detector dark counts only) is determined by tuning the laser wavelength away from the OH transition. Differences between OH$_\text{CHEM}$ and OH$_\text{WAVE}$ can highlight the presence of an OH interference. During the summer AIRPRO campaign, once the known OH interference deriving from laser photolysis of ambient ozone and the subsequent reaction of photogenerated O(1D) atoms with ambient H$_2$O (v) were accounted for (Woodward-Massey et al., 2020), the agreement between OH$_\text{CHEM}$ and OH$_\text{WAVE}$ was generally very good (see Fig. 14 in Woodward-Massey et al., 2020). However, on five afternoons when ozone was extremely elevated (> 100 ppbv) and OH concentrations were high (> 1 × 10$^{6}$ cm$^{-3}$), OH$_\text{WAVE}$ was greater than OH$_\text{CHEM}$ (by up to 18%), highlighting a small unknown interference under these very perturbed conditions. In all the model–measurement comparisons presented in Sect. 3, the interference-free OH$_\text{CHEM}$ measurement is used.

Both detection cells were calibrated every 3 d during the campaign by photolysis of a known concentration of H$_2$O (v) at 185 nm with a Hg lamp in synthetic air (Messer, Air Grade Zero 2) within a turbulent flow tube, which generates an equal concentration of OH and HO$_2$ (Whalley et al., 2018). The product of the photon flux at 185 nm (determined by N$_2$O actinometry before and after the instrument was deployed to Beijing; Commane et al., 2010), [H$_2$O] and irradiance time, was used to calculate [OH] and [HO$_2$]. For calibration of RO$_2$ concentrations, methane (Messer, Grade 5, 99.99%) was added to the humidified air flow in sufficient quantity to completely convert OH to CH$_3$O$_2$. The median limit of detection (LOD) during the campaign was 6.1 × 10$^5$ molecule cm$^{-3}$ for OH, 2.8 × 10$^6$ molecule cm$^{-3}$ for HO$_2$ and 7.2 × 10$^6$ molecule cm$^{-3}$ for CH$_3$O$_2$ at a typical laser power of 11 mW for a 5 min data acquisition cycle (SNR = 2). The field measurements of all species were recorded with 1 s time resolution, and the precision of the measurements was calculated using the standard errors in both the online and offline points. The accuracy of the measurements was ≈ 26% (2σ) and is derived from uncertainties in the calibration, which derive largely from that of the chemical actinometer (Commance et al., 2010).

### 2.3 Experimental budget analysis

An experimental budget analysis has been conducted for OH, HO$_2$, RO$_2$ and total RO$_x$ following the approach outlined in Tan et al. (2019) and which relies only on field-measured quantities (concentrations and photolysis rates) and published chemical kinetic data and not on any model calculated concentrations. The rates of production and destruction of each radical species are calculated using Eqs. (1)–(8) below.

\[
P_{\text{OH}} = j_{\text{HONO}}[\text{HONO}] + (2f \times j_{\text{O}^1D}[O_3]) + \sum_i \left( \phi_i \text{OH}k_i \text{[alkene]}[O_3] \right) + (k_2[\text{NO}] + k_3[O_3])[\text{HO}_2]
\]

\[
D_{\text{OH}} = [\text{OH}]k_{\text{OH}}
\]

\[
P_{\text{HO}_2} = 2j_{\text{HCHO,R}}[\text{HCHO}] + \sum_i \left( \phi_i \text{HO}_2k_i \text{[alkene]}[O_3] \right) + (k_4[\text{HCHO}] + k_5[\text{CO}])[\text{OH}] + \alpha k_6[\text{NO}][\text{RO}_2]
\]

\[
D_{\text{HO}_2} = (k_7[\text{NO}] + k_8[O_3] + k_9[\text{RO}_2] + k_{\text{het}} + 2k_{10}[\text{HO}_2])[\text{HO}_2]
\]

\[
P_{\text{RO}_2} = \sum_i \left( \phi_i \text{RO}_2k_i \text{[alkene]}[O_3] \right) + k_{\text{OH}}[\text{VOC}][\text{OH}]
\]

\[
D_{\text{RO}_2} = \left( (\alpha + \beta) k_6[\text{NO}] + (2k_{11}[\text{RO}_2] + k_9[\text{HO}_2]) \right)[\text{RO}_2]
\]

\[
P_{\text{RO}_x} = j_{\text{HONO}}[\text{HONO}] + 2f \times j_{\text{O}^1D}[O_3] + 2j_{\text{HCHO,R}}[\text{HCHO}] + \sum_i \left( \phi_i \text{OH} + \phi_i \text{HO}_2 + \phi_i \text{RO}_2 \right) k_i \text{[alkene]}[O_3]
\]

\[
D_{\text{RO}_x} = (k_{12}[\text{NO}_2] + k_{13}[\text{NO}])[\text{OH}] + \beta k_{6}[\text{NO}][\text{RO}_2] + 2(k_{11}[\text{RO}_2]^2 + k_9[\text{RO}_2][\text{HO}_2] + k_{10}[\text{HO}_2]^2),
\]

where $j_{\text{HONO}}$ and $j_{\text{O}^1D}$ are the measured photolysis rates of HONO and O$_3$ (forming O$^1$D) respectively; $f$ is the fraction of O$^1$D radicals that react with H$_2$O rather than are collisionally quenched to O(3P) ($f = 0.1$ on average); and $\phi_i$$_{\text{OH}}$, $\phi_i$$_{\text{HO}_2}$, $\phi_i$$_{\text{RO}_2}$, and $k_i$ are the yield of OH, HO$_2$ and RO$_2$ from, and rate coefficients for, individual ozone–alkene reactions taken from the MCM3.1 respectively. $j_{\text{HCHO,R}}$ is the measured HCHO photolysis rate that yields HO$_2$ radicals, and $k_{\text{het}}$ is the first-order loss of HO$_2$ to the measured aerosol surface area, calculated using Eq. (9):

\[
k_{\text{het}} = \frac{\omega A_p}{4}.
\]
where $\omega$ is the mean molecular speed of HO$_2$ (equal to 43 725 cm s$^{-1}$ at 298 K), $\gamma$ is the aerosol uptake coefficient (0.2 is used here as recommended by Jacob, 2000) and $A$ is the measured aerosol surface area in cm$^2$ cm$^{-3}$. $\alpha$ is the fraction of RO$_2$ radicals that upon reaction with NO propagate to HO$_2$ rather than reform another RO$_2$ radical; initially $\alpha = 1$ has been assumed, $\beta$ is the fraction of RO$_2$ radicals that upon reaction with NO form alkyl nitrates and is set to 0.05 as used by Tan et al. (2019) to represent an average alkyl nitrate yield for the various types of RO$_2$ species likely present. All rate coefficients ($k_1$–$k_{13}$) used are listed in Table 1, and the concentrations of species used in the budget analysis are the concentrations that were observed during the campaign.

### 2.4 MCM3.3.1 box model description

A zero-dimensional (box) model incorporating the Master Chemical Mechanism (MCM3.3.1; Jenkin et al., 2015; http://mcm.leeds.ac.uk/MCM/home, last access: 1 February 2021) was used to predict the radical concentrations and OH reactivity for comparison with the observations. The model was constrained by measurements of NO, NO$_2$, NO$_3$, O$_3$, CO, HCHO, HNO$_3$, HONO, water vapour, temperature, pressure and individual VOC species measured by DC-GC-FID (dual-channel gas chromatography with flame ionisation) and PTR-ToF-MS (proton transfer reaction time-of-flight mass spectrometry). Table 2 lists the different VOC species measured. HCHO was measured using a recently developed LIF instrument with 1 s time resolution and LOD of 80 pptv (Cryer, 2016). HONO was measured by a long-path absorption photometer (LOPAP) and broadband cavity-enhanced absorption spectrophotometry (BBCEAS) and the HONO concentration as recommended in Crilley et al. (2019) are used here. Further details on all instrumentation deployed during the campaign are overviewed in Shi et al. (2019).

The model was constrained with the measured photolysis frequencies $j$(O$_1$D), $j$(NO$_2$), and $j$(HONO), which were calculated from the measured wavelength-resolved actinic flux and published absorption cross sections and photodissociation quantum yields. For other species which photolysed at near-UV wavelengths (< 360 nm), such as HCHO and CH$_3$CHO, the photolysis rates were calculated by scaling to the ratio of clear-sky $j$(O$_1$D) to observed $j$(O$_1$D) to account for clouds. For species which photolysed further into the visible, the ratio of clear-sky $j$(NO$_2$) to observed $j$(NO$_2$) was used. The variation of the clear-sky photolysis rates ($j$) with solar zenith angle ($\chi$) was calculated within the model using the following expression:

$$j = l \cos(\chi) m \times e^{-n \sec(\chi)},$$  

with the parameters $l$, $m$ and $n$ optimised for each photolysis frequency (see Table 2 in Saunders et al., 2003). The model inputs were updated every 15 min; the species that were measured more frequently were averaged to 15 min, whilst the measurements with lower time resolution were interpolated.

To estimate how long model-generated intermediate species survive before being physically removed by processes such as deposition or ventilation, the model was left unconstrained to glyoxal, and the rate of physical loss was varied. The model was able to reproduce the observed glyoxal concentrations if a deposition velocity of 0.5 cm s$^{-1}$ was used, combined with a ventilation term that increased with boundary layer depth. As the boundary layer gradually increased in the morning, the lifetime of glyoxal with respect to ventilation was $\approx 1$ h, whilst at night the lifetime gradually increased to $\approx 5$ h; this variable lifetime was applied to all model-generated species. As a further check on the physical loss rate imposed, the model was run unconstrained to HCHO using the same deposition rates and was found to reproduce the observed HCHO concentrations that were observed during the daytime but under-predicted the concentrations at night, potentially indicating that primary emissions of HCHO as well as secondary production contributed to the observed concentrations. In all the model scenarios presented in Sect. 3, the observed HCHO concentration is used. The model was run for the entirety of the campaign in overlapping 7 d 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 and spun up for 2 d before comparison to measurements of OH, HO$_2$, RO$_2$ and k(OH) was made. For comparison of the modelled RO$_2$ to the observed total RO$_2$, complex RO$_2$ and simple RO$_2$, the RO$_x$ LIF instrument sensitivity towards each RO$_2$ species in the model was determined by running a model first under the RO$_x$ LIF reactor and then the RO$_x$ LIF FAGE cell conditions (NO concentrations and residence times) to determine the conversion efficiency of each modelled RO$_2$ species to HO$_2$.

A series of model runs have been performed and are summarised in Table 3.

### 3 Results and discussion

#### 3.1 Overview of the chemistry and meteorology during the campaign

As part of the AIRPRO project, gas-phase, aerosol, and meteorological observations were made at the IAP site from 21 May to 26 June 2017. Typically clear skies and elevated temperatures prevailed, with rain on just a few days. Temperatures frequently exceeded 35 °C, whilst $j$(O$_1$D) peaked at just over 3 $\times$ 10$^{-5}$ s$^{-1}$ at noon (Fig. 1). The dominant wind direction reaching the site during the summer was from the south-west, and the measured hourly mean wind speed was 3.6 m s$^{-1}$ (Shi et al., 2019). Despite the close proximity of the measurement site to the heavily trafficked Jingzang highway in Beijing, mixing ratios of NO, which were elevated during the morning hours, often dropped below 500 pptv during the afternoon. The daytime emissions of NO$_x$ that
several VOC (BVOC) as well as chemical conditions more akin to forested environments (low NO, high biogenic VOC (BVOC)). From 9 to 12 June, NO levels were elevated throughout the day, suggesting a local source, whilst from 17 June to the end of the measurement period, NO concentrations dropped, and so, as well as the strong diurnal trend observed in the NO concentration, these periods provide further opportunity to test the model’s ability to predict radical concentrations as a function of NO by removing concomitant variables such as changing boundary layer depth and sunrise which occurred in unison with the morning increase in NO concentration.

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Table 3. Different model scenarios that are discussed in Sect. 3.

| Model name               | Description                                                                 |
|--------------------------|-----------------------------------------------------------------------------|
| Base model               | As described in Sect. 2.4                                                   |
| Base model-SA            | The base model with the inclusion of a first-order loss process of HO₂ to   |
|                          | aerosols calculated using Eq. (9) with an uptake coefficient, \( \gamma = 0.2 \) |
| Base model-Cl            | The base model with the inclusion of Cl atom chemistry, taken from Xue et al.|
|                          | (2015)                                                                      |
| Missing \( k(\text{OH}) \) (OH to \( \text{CH}_3\text{O}_2 \)) | The base model with an additional reaction converting OH to \( \text{CH}_3\text{O}_2 \) at a rate equal to the missing reactivity |
| Missing \( k(\text{OH}) \) (OH to \( \text{HOCH}_2\text{CH}_2\text{O}_2 \)) | The base model with an additional reaction converting OH to \( \text{HOCH}_2\text{CH}_2\text{O}_2 \) at a rate equal to the missing reactivity |
| Missing \( k(\text{OH}) \) (OH to \( \text{CH}_3\text{C(O)O}_2 \)) | The base model with an additional reaction converting OH to \( \text{CH}_3\text{C(O)O}_2 \) at a rate equal to the missing reactivity |
| Missing \( k(\text{OH}) \) (OH to \( \text{C}_9\text{H}_6\text{O}_2 \)) | The base model with an additional reaction converting OH to \( \text{C}_9\text{H}_6\text{O}_2 \) (which is an \( \alpha \)-pinene-derived RO₂ species) at a rate equal to the missing reactivity |

\( \text{C}_9\text{H}_6\text{O}_2 = \) Note that \( \text{C}_9\text{H}_6\text{O}_2 \) is an \( \alpha \)-pinene-derived RO₂ that forms during the ozone-initiated oxidation of \( \alpha \)-pinene. The additional production of \( \text{C}_9\text{H}_6\text{O}_2 \) peroxy radicals in this model scenario was used to investigate the impact of an RO isomerisation mechanism on the modelled radical concentrations.

Figure 1. Time series of ozone, NO, NO₂ isoprene, CO, \( j(O^1D) \) and temperature during the campaign

3.2 Radical concentrations and OH reactivity

The concentrations of RO₂ (OH + HO₂ + RO₂) radicals were high during the campaign (Fig. 2), with OH concentrations frequently exceeding \( 1 \times 10^7 \) molecule cm⁻³ and reaching up to \( 2.8 \times 10^7 \) molecule cm⁻³ on 30 May. These OH levels are amongst the highest measured in an urban environment (Lu et al., 2019) and are comparable to the OH concentrations observed in the Pearl River Delta downwind of the southern Chinese megacity of Guangzhou, where OH concentrations reached \( 2.6 \times 10^7 \) molecule cm⁻³ (Lu et al., 2012). HO₂ concentrations peaked at \( 1 \times 10^8 \) molecule cm⁻³ on 9 June, whilst the highest concentrations of total RO₂ were observed during the latter half of the campaign, peaking at \( 5.5 \times 10^8 \) molecule cm⁻³ on the afternoon of 15 June. RO₂ measurements, alongside OH and HO₂, were, until recently, relatively rare. OH and RO₂ were measured during the MEGAPOLI project in Paris (Michoud et al., 2012), where the average daytime maximum concentrations of RO₂ were \( 1.2 \times 10^8 \) molecule cm⁻³, which is over an order of magnitude lower than the levels observed in Beijing. Since the development of the RO₂ LIF technique (Fuchs et al., 2008), RO₂ observations are now reported by the Leeds, Juelich and PKU FAGE groups. RO₂ concentrations observed in London in the summer reached up to \( 5.5 \times 10^8 \) molecule cm⁻³ in air masses that had previously passed
over central London (Whalley et al., 2018). In Wangdu, a
town situated on the North China Plain, 170 km north-east of
Beijing, summertime RO\textsubscript{2} concentrations reached up to
1.5 \times 10\textsuperscript{9} molecule cm\textsuperscript{-3} (Tan et al., 2017), which, although lower than observed in central Beijing, are much higher than observed in the summertime in European cities, suggesting that there may be significant differences in the urban photochemistry occurring in China and Europe.

As well as the elevated daytime radical concentrations, concentrations of OH, HO\textsubscript{2} and RO\textsubscript{2} remained elevated above the instrumental LOD on most nights. The high nighttime OH concentrations (ranging from the LOD up to 2 \times 10\textsuperscript{9} molecule cm\textsuperscript{-3}) are comparable to the levels of OH observed at night in Yufa (a suburb of Beijing) and downwind of Guangzhou, where night-time OH concentrations ranged from 0.5–3 \times 10\textsuperscript{9} molecule cm\textsuperscript{-3} (Lu et al., 2014). The observations of OH from the earlier Chinese campaigns could be reconciled by a model if an additional RO\textsubscript{2} production process was included which recycled RO\textsubscript{2} to OH via HO\textsubscript{2}. A weak positive correlation is observed between night-time OH and RO\textsubscript{2} during AIRPRO, and the secondary peak in RO\textsubscript{2} occurred when NO\textsubscript{3} was observed to increase rapidly at around 19:30, suggesting that nitrate chemistry was one source of radicals in the evening. Alkyl nitrates, formed from isoprene + NO\textsubscript{3}, were also enhanced at these times at this site (Reeves et al., 2019).

The OH reactivity, typical of urban environments, displayed an inverse relationship with boundary layer height and was highest during the nights when emissions were compressed into a lower boundary layer depth of \approx 150 m. An average maximum of \(k\textsubscript{(OH)} \approx 37 \text{ s}^{-1}\) was observed at 06:00, with OH reactivity reaching 89 \text{ s}^{-1} on 15 June at 03:00. During the daytime, the OH reactivity dropped to a minimum of \approx 22 \text{ s}^{-1} on average at \approx 15:00 when the boundary layer had increased to \approx 1500 m. The magnitude of OH reactivity observed during AIRPRO is comparable to the OH reactivity observed at other urban sites in China in the summer (Lou et al., 2010; Fuchs et al., 2017) and also in Tokyo during the summer (Sadana et al., 2004; Chatani et al., 2009). In London, OH reactivity was approximately \approx 7–10 \text{ s}^{-1} lower than in central Beijing, with \approx 15 \text{ s}^{-1} observed during the day on average and an average maximum of \approx 27 \text{ s}^{-1} at 06:00 (Whalley et al., 2016). Lower OH reactivities are also reported from US urban sites in New York and Texas (Ren et al., 2003; Mao et al., 2010).

### 3.3 Experimental radical budget analysis

Owing to the relatively short lifetime of radicals, it can be assumed that their production rates and destruction rates are balanced. A comparison of the rates of production and destruction for each radical species can be used to help identify if all radical sources and sinks are accounted for and if the rates of propagation between radical species are fully understood. In London, the ratio of the OH production rate (Eq. 1) to OH destruction rate (Eq. 2) was generally close to 1 throughout the campaign, demonstrating consistency between the OH, HO\textsubscript{2}, \(k\textsubscript{(OH)}\), HONO and NO\textsubscript{3} observations (Whalley et al., 2018). However, under low NO conditions (< 0.5 ppbv), the rate of OH destruction exceeded the calculated production rate, indicating that Eq. 1 was missing a source term under these regimes (Whalley et al., 2018). A steady-state analysis of HO\textsubscript{2} conducted for the London project, which balanced the HO\textsubscript{2} production terms (Eq. 3) with the first- and second-order loss terms (Eq. 4), highlighted that closure between the production and destruction terms could only be reconciled if the rate of propagation of the observed RO\textsubscript{2} radicals to HO\textsubscript{2} was decreased substantially to just 15%, demonstrating that the mechanism by which RO\textsubscript{2} radicals propagate to other radical species may not be well understood (Whalley et al., 2018). As set out by Tan et al. (2019), analogous budget analyses can be performed for RO\textsubscript{2} species (Eqs. 5 and 6) and for the entire RO\textsubscript{a} budget (Eqs. 7 and 8). Tan et al. (2019) found that the production and destruction terms for RO\textsubscript{2} were balanced in the mornings in the PRD, when the measured OH reactivity was used to calculate the rate of RO\textsubscript{2} production from VOC+OH reactions, but during the afternoon a missing RO\textsubscript{2} sink (2–5 ppbv h\textsuperscript{-1}) was evident. In the PRD study (Tan et al., 2019), the OH destruction rate exceeded the production rate by 4–6 ppbv h\textsuperscript{-1} in the afternoon, but, in contrast to London (Whalley et al., 2018), the HO\textsubscript{2} budget was closed throughout the whole day. The total rates of RO\textsubscript{a} production and destruction were in good agreement in the PRD (Tan et al., 2019).

A comparison of the campaign median production and destruction rates for RO\textsubscript{a}, OH, HO\textsubscript{2} and RO\textsubscript{2} during AIRPRO is presented in Fig. 3. The total rates of RO\textsubscript{a} production and destruction are in good agreement throughout the day from \approx 10:00. A night-time source of radicals of just under 1 ppbv h\textsuperscript{-1} is missing from the budget analysis, likely reflecting missing production from NO\textsubscript{3} + VOC reactions (nighttime radical production is considered further in Sect. 3.5). From 06:00 to 10:00, the RO\textsubscript{a} destruction exceeds the production by up to 4 ppbv h\textsuperscript{-1}, indicating a substantial, \approx 50 %, missing primary RO\textsubscript{a} source at this time. Previous work has suggested that Cl-initiated VOC oxidation may be an important source of RO\textsubscript{2} radicals in urban regions (Riedel et al., 2014; Bannan et al., 2015; Tan et al., 2017) but has not been included in the RO\textsubscript{a} or RO\textsubscript{2} production rate calculations here. Nitryl chloride was measured for part of the AIRPRO campaign, and the impact of this on the modelled RO\textsubscript{2} concentration is investigated in Sect. 3.4. The total RO\textsubscript{a} production and destruction rate is of the order of 6 ppbv h\textsuperscript{-1} at noon, which is slightly faster than in the PRD, where a median peak total radical production rate of \approx 4 ppbv h\textsuperscript{-1} was calculated. The median OH destruction rate is \approx 30 ppbv h\textsuperscript{-1} at noon and is roughly twice as fast the production rate at this time, highlighting a large missing source of OH radicals in the budget (\approx 15 ppbv h\textsuperscript{-1}). Although a missing OH source was also re-
Figure 2. Time series of the measured and modelled OH, HO$_2$, total RO$_2$ and OH reactivity during the campaign.

Figure 3. Campaign median production and destruction rates for OH, HO$_2$, total RO$_2$ and RO$_x$. The shaded areas represent the 1σ standard deviation of the data, representing the variability from day to day.

ported in the PRD (Tan et al., 2019), the missing production rate is $\approx 3$ times faster during AIRPRO. The known OH production rate during AIRPRO is dominated by the reaction of HO$_2$ with NO (contributing $\approx 60\%$ during the day to $P$(OH) in Eq. 1). The median peak HO$_2$ production of $\approx 60$ ppbv h$^{-1}$ is observed in the morning hours and greatly exceeded the known rate of HO$_2$ destruction by $\approx 50$ ppbv h$^{-1}$. HO$_2$ production is driven by the reaction of RO$_2$ with NO, which accounts for 88% of the total. The reaction of OH with CO and HCHO accounts for a further 9%. The total HO$_2$ production...
rate is approximately 4 times faster than that calculated for the PRD (Tan et al., 2019). The total rate of RO₂ destruction mirrors the HO₂ production, in that it is dominated by the reaction of RO₂ radicals with NO. From sunrise to 14:00, the rate of RO₂ destruction is faster than RO₂ production by up to 50 ppbv h⁻¹. After 14:00, the rates of RO₂ production and destruction are in good agreement. This trend contrasts with the budget analysis presented from PRD (Tan et al., 2019), which highlighted a possible missing RO₂ sink during the afternoon hours and budget closure in the morning hours.

Binning the ratio of P(\text{OH}) to D(\text{OH}), P(\text{HO}_2) to D(\text{HO}_2) and P(\text{RO}_2) to D(\text{RO}_2) against the NO mixing ratio (Fig. 4) reveals that the RO₂ budget is in good agreement at the lowest NO mixing ratios, but as NO mixing ratios increase the destruction of RO₂ becomes faster than the production of RO₂ by up to a factor of 10 at the highest NO bin. The trends in the RO₂ and HO₂ ratios are similar in the morning hours, albeit in opposite directions, and suggest that rather than there being a missing primary source of RO₂ and missing sink for HO₂ that happen to balance, instead, as found in London (Whalley et al., 2018), the net propagation rate of RO₂ to HO₂ may be substantially slower than the rate that has currently been used in this analysis. In London (Whalley et al., 2018), the modelled rate of production analysis revealed that only \( \approx 50\% \) of the total RO₂ species propagated to HO₂ following reaction with NO, as a significant fraction of the alkoxy radicals formed (such as those generated during the oxidation of monoterpenes and long-chain alkanes) preferentially isomerised and reformed a more oxidised RO₂ species in the presence of O₂ instead. In the radical flux analysis using the MCM3.2 for London (Whalley et al., 2018), the propagation of alkyl- and acyl-RO₂ species were combined, and so the interconversion of acyl-RO₂ radicals (from the OH-initiated oxidation of aldehydic VOCs, photolysis of ketones and decomposition of PAN species) to alkyl-RO₂ radicals following reaction with NO was not explicitly shown, but this interconversion of one RO₂ species to another would serve to reduce the fraction of RO₂ radicals that propagate to HO₂ further. Thus far for AIRPRO, the experimental budget analysis has assumed that 95 % of the measured RO₂ species, upon reaction with NO, produce HO₂. If, however, a large fraction of the total RO₂ measured derive from long-chain alkanes, monoterpenes or acyl-RO₂ species, the budget analysis will overestimate HO₂ production and also the net RO₂ destruction, as the reaction of these peroxy radicals with NO effectively converts one RO₂ species to another RO₂ species, and so the reaction with NO will be neutral in terms of RO₂ production and destruction. Taking \( \alpha = 0.1 \) leads to a good agreement between the production and destruction rates of HO₂ over the whole day and the observed range of NO. The production and destruction rates of RO₂ agree under high NO conditions, but at NO mixing ratios < 5 ppbv the production of RO₂ exceeds the destruction, highlighting (if this \( \alpha \) value is correct) that there is a missing RO₂ sink at the lower NO concentrations. Tan et al. (2019) also report a missing RO₂ sink under low NO conditions during PRD and suggested that autoxidation of RO₂ species could account for this missing sink and may also possibly act as the missing source of OH identified under the low NO conditions. An additional first-order reaction that converts RO₂ to OH at a rate of 0.1 s⁻¹ brings the P : D(\text{OH}) and P : D(\text{RO}_2) ratios close to 1 at all

Figure 4. The median ratio of the OH, HO₂ and total RO₂ production rates to destruction rates binned over the NO mixing ratio range encountered during the campaign on a logarithmic scale. The box and whiskers represent the 25th and 75th and the 5th and 95th confidence intervals. The number of data points in each of the NO bins is \( \approx 80 \).
NO mixing ratios > 0.3 ppbv, but at low NO mixing ratios (0.1–0.3 ppbv range) an even slower rate of conversion is required, highlighting, as one might expect, that the overall rate of RO₂ isomerisation is variable and likely depends on the specific RO₂ species present at a particular time or location. In the PRD study (Tan et al., 2019), the HO₂ budget was closed when $\alpha = 0.95$ was used, suggesting that acyl peroxy radicals and those derived from long-chain alkanes and monoterpenes only made up a very small fraction of the total RO₂ concentration.

Although revealing, this type of experimental budget analysis coupled with the radical observations is unable to differentiate between different RO₂ types, and so assumptions have to be made on the fraction of the total RO₂ that propagate to HO₂. In the following section, a box model constrained to the latest MCM scheme (MCM3.3.1) is used to predict the radical concentrations. The MCM is a near-explicit model and, as such, treats the production, propagation and destruction of each RO₂ species present discretely and so can provide an insight into the rate at which different RO₂ species convert to HO₂ or to other RO₂ species (or, indeed to OH) and the impact this propagation has on NO to NO₂ conversion and, hence, O₃ production.

3.4 MCM modelled radical predictions and comparison with observations

The time series of the model-predicted radical concentrations and a breakdown of the modelled OH reactivity from the base MCM model are overlaid with the observations in Fig. 2. The average diurnal profiles of the measured and modelled radical and $k$(OH) profiles are also provided in Fig. 5. In contrast to the experimental budget analysis, the model-predicted OH is in excellent agreement with the observed OH throughout the campaign. This same model overestimates HO₂, however, particularly during the daytime but also during the evening when a small secondary peak in HO₂ is predicted but not observed. An exception to this trend occurs between 9–12 June when elevated levels of NO were measured at the site during the day, and on these days, the agreement between the observed HO₂ and the model is better. The over-prediction of HO₂ primarily occurs under the lower NO conditions that were typically observed during the afternoon hours; the skill of the model to predict the radical concentrations as a function of NO is discussed further below. The model underestimates total RO₂ throughout the measurement period, although the level of disagreement (in terms of absolute concentration) is most severe (15–22 June) when NO concentrations were at their lowest. During this period, the average NO mixing ratio was $\approx 0.4$ ppbv during the afternoon hours, whilst the average NO mixing ratio for the entirety of the campaign was $\approx 0.75$ ppbv during the afternoons (Fig. S1 in the Supplement). The average peak NO mixing ratio observed in the morning (1–22 June) was just over 6 ppbv, whilst the average peak NO mixing ratio for the entirety of the campaign was close to 16 ppbv. During this period, the observed RO₂ concentrations were most elevated relative to other times during the campaign; however, the model does not predict a similar increase in RO₂ concentrations during this period relative to other times in the campaign. OH reactivity is underestimated by the model, on average by $\approx 10$ s⁻¹. However, between 15–22 June the average missing OH reactivity increases to $\approx 13$ s⁻¹. The model underestimation of OH reactivity may, in part, contribute to the model underestimation of RO₂ as the model is evidently underestimating the rate of OH + VOC reactions which form RO₂. Including an additional reaction between OH and VOC to account for the missing reactivity in the model and the impact this has on the modelled radical concentrations is investigated in Sect. 3.6. Although the model is able to capture the observed OH concentrations reasonably well, the model’s failure to reproduce the observed HO₂ and RO₂ (and in the base model, the OH reactivity) indicates the model is either missing or misrepresenting some key reactions. Furthermore, the discrepancy between the model-predicted OH and OH budget analysis which highlighted a missing OH source, suggests that the over-prediction of HO₂ is masking a missing OH source in the MCM model.

Qualitatively, the model overestimation of HO₂ and underestimation of RO₂ is consistent with the budget analysis, which identified a missing RO₂ production term and missing HO₂ destruction term which could be reconciled, in part, by slowing the rate at which RO₂ propagates to HO₂. However, when the HO₂ measured to modelled ratio is binned against NO, differences between the model and budget analyses become apparent (Fig. 6). The model over-predicts the observed HO₂ concentrations at the lowest NO mixing ratios experienced (0.1–1 ppbv); this over-prediction can be reconciled (under the very lowest NO conditions, < 0.3 ppbv) when a loss of HO₂ to aerosols (calculated using Eq. 9, with an uptake coefficient of 0.2) is included in the model. This demonstrates that a reduction in aerosol surface area has the potential to enhance HO₂ concentrations and thereby increase photochemical ozone formation but only under very low NO conditions. As there was little to no change in the modelled HO₂ concentration upon inclusion of an heterogeneous loss term under the higher NO conditions, efforts to reduce anthropogenic PM when NO is present (which is highly likely to be the case) would not be expected to lead to an increase in HO₂ and, in turn, O₃, as was suggested from earlier modelling studies (Li et al., 2019). Between 1–5 ppbv NO, the model is able to reproduce the observed HO₂ well between 9–12 June, the daytime NO concentrations fell within this intermediate NO range, hence the good agreement between the model and observations on these days. In contrast with the budget analysis, the model under-predicts HO₂ beyond 5 ppbv NO by up to a factor of 10 at the highest NO experienced (see the 52 ppbv NO bin, Fig. 6, which includes NO mixing ratios up to 104 ppbv). The model under-predicts the observed RO₂ over the whole NO range, and, consis-
Figure 5. Average profiles for the observed OH, HO$_2$, total RO$_2$, partially speciated RO$_2$ (in molecule cm$^{-3}$) and OH reactivity at 15 min intervals over 24 h. The error bars represent the 1σ standard deviation of the measurements, representing the variability in the measurements from day to day. The average diurnal profiles for OH, HO$_2$, total RO$_2$, partially speciated RO$_2$ and OH reactivity from the base model are overlaid.

Cl atoms, formed from the photolysis of nitryl chloride (ClNO$_2$), have been shown to act as a source of RO$_2$ (Riedel et al., 2014; Bannan et al., 2015; Tan et al., 2017) and have also been investigated here to see if Cl chemistry can resolve the modelled RO$_2$ under-prediction under the elevated NO concentrations which were typically observed during the mornings. ClNO$_2$ was measured for part of the campaign (Zhou et al., 2018) and reached up to 1.44 ppbv during the night on 12–13 June. The Cl atom concentration exceeded 4 x 10$^4$ atoms cm$^{-3}$ during the morning of 13 June and exceeded 1 x 10$^4$ atoms cm$^{-3}$ on several other mornings (Fig. 7). The Cl atom concentration was calculated from the concentration of ClNO$_2$, its photolysis rate to yield Cl (determined from the observed actinic flux and published absorption cross section of ClNO$_2$) and the VOC loading. During these times, the modelled RO$_2$ concentrations increased, relative to the concentration in the base model, by up to 2.5 x 10$^7$ molecule cm$^{-3}$, which represents close to a 100% increase in the modelled RO$_2$ at these times. On several mornings (4, 5, 7 and 13 June) this increase in RO$_2$ brought the model and measured RO$_2$ into close agreement.
The production rate of RO$_2$ from Cl-initiated VOC oxidation on these mornings would serve to enhance P(RO$_2$) by up to 2.1 ppbv h$^{-1}$. However, on several nights, only low concentrations of ClNO$_2$ were measured, and only very low concentrations of Cl atoms were calculated to be present upon sunrise, and so, on these days, only modest enhancements (1–2×10$^7$ molecules cm$^{-3}$) in RO$_2$ concentrations were predicted by the model, and the large under-prediction in the RO$_2$ concentration on these mornings remained, which may indicate that there are other, overlooked, primary RO$_x$ sources in the experimental budget calculation besides missing Cl + VOC reactions. The Cl atom concentration dropped off rapidly during the mornings with just ≈100 atoms cm$^{-3}$ present by noon on most days and so was unable to reconcile the magnitude of the RO$_2$ underestimation observed throughout the day.

3.5 Rate of production and rate of destruction analysis

A rate of production and rate of destruction analysis on model OH, HO$_2$ and RO$_2$ species (Fig. 8) highlights the main radical sources and sinks in the base model. Consistent with earlier studies of radicals in urban locations, the photolysis of HONO is the dominant primary source of radicals during the daytime, accounting for ≈64% of the primary radical production on average during the day (05:00–19:30) throughout the campaign. The photolysis of O$_3$ and subsequent reaction of O$(^1$D) with H$_2$O vapour accounts for ≈9% of primary production during the day, whilst the photolysis of HCHO and other photo-labile VOCs accounts for ≈11% of the radical production. Ozonolysis and nitrate radical (NO$_3$) reactions account for 9% and 7% of the total radical production during the day, respectively. At night, both ozonolysis (≈18%) and nitrate radical reactions (≈82%) are the source of radicals. The primary source of radicals from VOC + NO$_3$ reactions is ≈1 ppbv h$^{-1}$ during the night, which is sufficient to close the RO$_x$ experimental budget (Fig. 3).

Figure 9 highlights the rates of propagation in the model which transform OH to HO$_2$ and RO$_2$, RO$_2$ to HO$_2$ and HO$_2$ back to OH. The rate of propagation is rapid, and the secondary source of OH from HO$_2$ + NO is more than twice as large as the primary production of OH from HONO photolysis. Approximately one-third of the OH reacts with CO, O$_3$ or HCHO to form HO$_2$, just over one-third reacts with VOCs to form RO$_2$ and just under one-third is lost by reaction with NO$_2$ forming nitric acid. In contrast to London (Whalley et al., 2018), the majority of RO$_2$ formed during AIRPRO propagates to HO$_2$, and subsequently the majority of HO$_2$ propagates back to OH. From the model radical flux analysis, which takes into consideration the different types of RO$_2$ species present, a value of $\alpha = 0.87$ is derived (where $\alpha = 1$ minus the rate at which RO forms RO$_2$ or RC(O)O$_2$ divided by the rate of RO conversion to HO$_2$). Note that this fraction does not consider RO$_2$ and RC(O)O$_2$ termination reac-

Figure 6. The median ratio (−) of the measured to modelled OH, HO$_2$ and total RO$_2$ binned over the NO mixing ratio range encountered during the campaign on a logarithmic scale. The box and whiskers represent the 25th and 75th and the 5th and 95th confidence intervals. The red circles in the middle panel display the measured to modelled HO$_2$ ratio when the model includes a heterogeneous loss of HO$_2$ to aerosols calculated using Eq. (9). The number of data points in each of the NO bins is ≈80.
Figure 7. Time series of the measured and modelled OH, HO\textsubscript{2} and total RO\textsubscript{2} during the campaign when ClNO\textsubscript{2} was also measured. The Cl atom concentration calculated to be present is shown in the top panel. The measured OH concentrations are represented by the blue line, HO\textsubscript{2} by the red line and total RO\textsubscript{2} by the green line. The base model scenario is shown in grey, whilst the base model with Cl atom chemistry included (Xue et al., 2015) is shown in orange (only evident in the RO\textsubscript{2} panel).

Figure 8. The average diurnal rates of primary production and termination for RO\textsubscript{x} radicals in parts per billion per hour (ppbv h\textsuperscript{-1}) in the base model scenario. CH\textsubscript{3}C(O)O\textsubscript{2} + NO\textsubscript{2} (Net) represents the net rate (forward minus backward) for all RC(O)O\textsubscript{2} + NO\textsubscript{2} ↔ PAN species.

In London, the model-derived \(\alpha\) was \(\approx 0.5\), reflecting the presence of long-chain alkane-derived RO\textsubscript{2} species from diesel emissions and monoterpenes. In Beijing, measurement of such long-chain VOC species could not be attempted, but these could have been present. A lumped monoterpene signal was measured by PTR-ToF-MS and is included in the model, split equally between \(\alpha\)-pinene and limonene. The base model, on which the radical flux analysis was performed, under-predicts OH reactivity and so is likely missing RO\textsubscript{2} species from additional OH+VOC reactions, which, depending on the RO\textsubscript{2} type, may serve to reduce \(\alpha\).

3.6 OH reactivity and missing OH reactivity

NO\textsubscript{2} was the single biggest contributor to the OH reactivity in Beijing, with a campaign average contribution of 18.6 % (Fig. 5). This is similar to the NO\textsubscript{2} contribution to OH reactivity observed in London (Whalley et al., 2016). NO contributed just 1.3 % to the total reactivity in Beijing, compared to a 4.2 % contribution in London (Whalley et al., 2016). In London, measured carbonyl species accounted for close to 20 % of the OH reactivity budget, largely due to the high concentrations of HCHO (Whalley et al., 2016). In contrast, in Beijing, carbonyls accounted for just 3.8 % of the mea-
Figure 9. A model reaction flux analysis, showing the mean rate of reaction for formation, propagation and termination of radicals (pptv h$^{-1}$) (day and night) during the whole campaign.

sured $k$(OH). Alkenes and dialkenes were more prevalent in Beijing than in London, and the dialkene group of VOCs (dominated by isoprene) accounted for 10.5 % of the OH reactivity in Beijing compared to 1.8 % in London (Whalley et al., 2016). Owing to the faster physical loss of secondary species in Beijing by ventilation compared to London (see Sect. 2), the contribution that model-generated intermediate species made to the observed OH reactivity was 2.7 % in Beijing vs. 23.8 % in London (Whalley et al., 2016). In contrast to Beijing, where approximately 30 % of the measured reactivity remains unaccounted for, in London, the OH reactivity budget was largely closed (Whalley et al., 2016).

In Beijing during the measurement period when the missing OH reactivity reached on average 13 s$^{-1}$ (15–22 June), isoprene concentrations were elevated relative to earlier in the campaign (Fig. 1). Overall, much higher concentrations of isoprene were observed in Beijing than in London (Whalley et al., 2016), and so this may indicate that other biogenic species that were not measured, along with their oxidation products, may account for some of the missing OH reactivity in Beijing.

A series of model simulations have been performed, whereby an additional OH to RO$_2$ reaction has been included to account for the missing reactivity at a given time (Fig. 10); the RO$_2$ formed has been varied to investigate the influence of different RO$_2$ types on the modelled radical concentrations. When OH converts to methyl peroxy radicals, the modelled RO$_2$ concentration increases by close to a factor of 2 on average, but just over a factor-of-2 under-prediction of the observed RO$_2$ radicals remains. Unsurprisingly, it is the modelled fraction of RO$_2$ radicals that do not act as an HO$_2$ interference (simple RO$_2$) that increase in this scenario, and the model now only underestimates this class of RO$_2$ species by a factor of 1.45, whilst complex RO$_2$ is still underestimated by a factor of 6.2. When OH converts to HOCH$_2$CH$_2$O$_2$ (an RO$_2$ species that does act as an HO$_2$ interference, formed from the reaction of OH with ethene), the modelled complex RO$_2$ fraction increases, and the model underestimation of complex RO$_2$ is reduced to a factor of 1.8 on average, with the largest under-predictions observed during the evening hours. In both these model simulations, the modelled over-prediction of HO$_2$ increases from the base model scenario as CH$_3$O$_2$ and HOCH$_2$CH$_2$O$_2$ both rapidly propagate to HO$_2$. The modelled OH concentration displays a modest decrease with the additional OH sink; however, this is largely compensated for by the increase in modelled HO$_2$, which enhances the secondary source of OH from HO$_2$ + NO, and so, overall, the modelled OH concentration is largely buffered by the inclusion of missing OH reactivity in the form of additional methane (leading to CH$_3$O$_2$) or ethene (leading to HOCH$_2$CH$_2$O$_2$).

Model simulations (not shown) which include an additional source of CH$_3$C(O)O$_2$, for example, from additional CH$_3$CHO + OH reactions, do predict substantially less HO$_2$
(and can reconcile the observed HO$_2$ to with 25%), but modelled RO$_2$ concentrations do not increase as a large fraction of the acyl-RO$_2$ radicals react with NO$_2$ to form PAN and are, therefore, lost. These missing reactivity model simulations and measurement comparisons suggest that the missing RO$_2$ may be a species, which, upon reaction with NO, converts from one RO$_2$ species to another and, therefore, competes with RO$_2$ to HO$_2$ propagation rather than a RO$_2$ radical, which leads to RO$_2$ termination. This suggests that the overall lifetime of RO$_2$ radicals is longer than currently estimated and that multiple conversions of one RO$_2$ species to another may be occurring to sustain the high concentrations observed. As identified in London, larger, more complex VOC species such as monoterpenes or long-chain alkanes deriving from diesel emissions do undergo multiple RO$_2$ to RO$_2$ conversions in the presence of NO as the alkoxy radical formed preferentially undergoes isomerisation rather than an external H atom abstraction by O$_2$. If an additional reaction which converts OH to an RO$_2$ species formed during the oxidation of $\alpha$-pinene, and which undergoes four reactions with NO before eventually forming HO$_2$, is added to the model at a rate sufficient to reconcile the missing OH reactivity, the model predicts significantly more total RO$_2$ and now only modestly under-predicts the observed RO$_2$ concentrations (by a factor of 1.8). The modelled radical concentrations predicted from the “missing k(OH) (OH to C9602)” scenario are overlaid with the radical observations and modelled radicals from the base model scenario in Fig. S2. The additional VOC reactivity which produces RO$_2$ radicals that isomerise after reaction with NO is able to increase the modelled total RO$_2$ concentration, both under the lower NO conditions experienced between 15–22 June and on the higher NO days (9–12 June), indicating that NO is still at sufficient concentrations to dominate the fate of RO$_2$ between 15–22 June, despite NO concentrations being lower. The median measured to modelled (missing k(OH) (OH to C9602)) ratio vs. NO (Fig. S3) highlights that the inclusion of alkoxy isomerisation following the RO$_2$ + NO reaction increases the modelled RO$_2$ across the entire NO range but, considering the log scale, has the biggest impact on the ratio (from the measured to modelled (base) ratio) at the highest NO concentration. Both the simple and complex RO$_2$ species are enhanced, as the first three generations of RO$_2$ species formed would be detected during the RO$_2$ mode in the RO$_1$ LIF instrument and, hence, contribute to simple RO$_2$. The final RO$_2$ species formed, that does propagate to HO$_2$ via RO upon reaction with NO, would be detected during the HO$_2$ mode in the RO$_2$ LIF instrument and, as such, contributes to the complex RO$_2$ fraction. In this scenario, the HO$_2$ concentration is now only modestly overestimated by a factor of 1.4. The RO$_1$ LIF instrument relies on the conversion of RO$_2$ species to HO$_2$ (and ultimately to OH) for detection, so one might expect the instrument to be insensitive to RO$_2$ species that do not directly propagate to RO then to HO$_2$ upon reaction with NO. However, given the RO$_2$ LIF flow tube conditions (NO concentration of 4 $\times$ 10$^{13}$ molecule cm$^{-3}$ and residence time of just under 1 s), RO$_2$ species that require several reactions with NO before HO$_2$ is produced should still be detected. These types of RO$_2$ species that require more than one reaction with NO before HO$_2$ forms may be generated via the additional VOC + OH reactions identified as missing OH reactivity (as presented here). They may also be present due to a missing primary source of RO$_2$ such as decomposition of a complex PAN species, VOC photolysis, a Cl atom + VOC reaction or an alkene ozonolysis product. The experimental peroxy radical budget analysis highlighted that budget closure could only be achieved if $\alpha$ was reduced to 0.1, which suggests that the model breakdown of peroxy radical species present (e.g. the fraction of acyl-RO$_2$, long- vs. short-chained alkyl-RO$_2$ species) may be incomplete. In the scenario in which OH converts to an $\alpha$-pinene-derived RO$_2$ species, consistent with the experimental budget analysis, the model under-predicts the observed OH by a factor of 1.8, revealing that there is a missing source of OH under the low NO conditions in Beijing that was previously masked by the model over-prediction of HO$_2$.

Figure 10. Average diel profiles for the observed OH, HO$_2$, total RO$_2$ and partially speciated RO$_2$ (black lines) at 15 min intervals over 24 h. The error bars represent the 1σ standard deviation of the measurements. The average OH, HO$_2$, total RO$_2$ and partially speciated RO$_2$ model profiles when the missing reactivity observed at a given time is accounted for by different OH to RO$_2$ reactions are overlaid (yellow, blue and green lines); the base model predictions are in red. See text for details.
3.7 Impact on ozone production

Previous work, for example, by Tan et al. (2017), suggested that the addition of a primary RO$_2$ source could help reconcile the model under-prediction of RO$_2$. However, as demonstrated in Sect. 3.6, the identity of the primary RO$_2$ is important, and in Beijing a complex RO$_2$ species that has a large enough carbon skeleton such that the RO radical formed upon reaction with NO preferentially isomerises to another RO$_2$ (and undergoes multiple RO$_2$ to RO$_2$ conversions before eventually forming HO$_2$) is needed to reconcile both the observed RO$_2$ and HO$_2$ concentrations. These types of RO$_2$ species may also preferentially isomerise rather than undergo the bimolecular reactions with NO if NO concentrations are low enough. For example, laboratory studies have shown that the monoterpenes, following an initial attack by ozone or OH, form highly oxidised RO$_2$ radicals within a few seconds via repeated H shift from C–H to an R–O–O bond and subsequent O$_2$ additions (Ehn et al., 2014; Jokinen et al., 2014; Berndt et al., 2016). Recently, autoxidation has also been shown to occur during the oxidation of aromatic VOCs too (S. N. Wang et al., 2017). Autoxidation reactions may generate OH directly from RO$_2$ and, therefore, may also resolve the missing OH source reported under low NO conditions (here and in the literature). These types of autoxidation reactions lead to the generation of HOMs, which have also been shown to condense and contribute to SOA (Mohr et al., 2019). Mass spectrometric signals relating to these highly oxidised RO$_2$ species were observed during the AIRPRO campaign (Breen et al., 2019; Mehra et al., 2021) suggesting that autoxidation was occurring at the Beijing site. Unimolecular H atom shifts are represented within the MCM3.3.1 for isoprene oxidation. Autoxidation reactions for other RO$_2$ radicals are currently not included within the MCM3.3.1, although improved representation of RO$_2$ radical chemistry is a focus for the next generation of explicit detailed chemical mechanisms (Jenkin et al., 2019).

In addition to missing unimolecular RO$_2$ reactions, the model may be missing other RO$_2$ reaction pathways, for example, RO$_2$ accretion reactions, as identified by Berndt et al. (2018). Although it is difficult to fully assess how competitive these RO$_2$ + RO$_2$ reactions may be compared to RO$_2$ + NO reactions from the total RO$_2$ observations made (the concentration of each individual RO$_2$ would be needed), the inclusion of accretion reactions in the MCM would serve to reduce the modelled RO$_2$ concentration under low NO$_x$ conditions as the reaction represents an overall RO$_3$ sink. This suggests that the missing RO$_2$ source identified here may be even larger under the lower NO conditions.

The model measurement comparisons above suggest that our understanding of the rate at which the larger RO$_2$ species propagate to HO$_2$ (or to OH directly) and the possible reactions they undergo (which have not undergone substantial laboratory study) is far from complete and highlights that RO$_2$ chemistry warrants further study. One important finding, however, is that the underestimation of the observed RO$_2$ may be caused by missing reactions that compete with the RO$_2$ + NO reactions that form HO$_2$. These competing reactions are effectively slowing the rate at which RO$_2$ species convert to HO$_2$, but if, as suggested here, these reactions are RO$_2$ + NO reactions that reform another RO$_2$ radical, they will still be relevant in terms of ozone production. Under low NO conditions there is emerging evidence that unimolecular isomerisation reactions occur for a range of RO$_2$ radicals (Ehn et al., 2014; Jokinen et al., 2014; Berndt et al., 2016; S. N. Wang et al., 2017) as well as RO$_2$ accretion reactions (Berndt et al., 2018). These reactions will effectively remove RO$_2$ radicals without conversion of NO to NO$_2$ and so also have implications for modelling in situ O$_3$ production, if models rely only on the rate of VOC oxidation when investigating O$_3$ production.

By approximating the rate of ozone production to the rate of NO$_2$ production from the reaction of NO with HO$_2$ and RO$_2$ radicals, urban radical measurements can be used to estimate chemical ozone formation (Kanaya et al., 2007; Ren et al., 2013; Brune et al., 2016; Tan et al., 2017; Whalley et al., 2018).

\[
P(O_3) = \left( k_{\text{HO}_2+\text{NO}[\text{HO}_2][\text{NO}]} + k_{\text{RO}_2+\text{NO}[\text{RO}_2][\text{NO}]} \right) 
\]

Losses of O$_3$ ($L(O_3)$) include chemical losses such as the reaction of NO$_2$ with OH, net PAN formation, the fraction of O($^1D$) (formed by the photolysis of O$_3$) that react with H$_2$O and the reaction of O$_3$ with OH and HO$_2$. Physical loss processes, such as O$_3$ deposition and ventilation out of the model box (see Sect. 2.4), will also contribute to L(O$_3$). Physical processes such as advection of O$_3$ into the model box would also need to be considered in the model to make a direct comparison to the observed O$_3$ concentrations.

Considering the chemical production of O$_3$ (Eq. 11), recent studies, in which OH, HO$_2$ and RO$_2$ observations (via RO$_2$, LIF) were made, demonstrated that models may underpredict ozone production at high NO due to an underestimation of the RO$_2$ radical concentrations at high NO concentrations (Tan et al., 2017; Whalley et al., 2018). Figure 11 displays the mean ozone production calculated from the radical observations (red line) as a function of NO, and, consistent with the earlier ozone production calculations from the Wangdu (Tan et al., 2017) and London (Whalley et al., 2018) studies, the in situ ozone production calculated from the modelled OH and peroxy radicals (black line) is lower than from the observed radicals, most significantly at the higher NO concentrations. To accurately simulate ozone production and to understand how emission reduction policies may impact ozone levels, it is essential that the model accurately reflects the types of RO$_2$ species present and how fast they propagate to another RO$_2$ species or to HO$_2$ or to OH.
Figure 11. Mean $\text{O}_3$ production (ppbv h$^{-1}$) calculated from observed (red line) and modelled (black line) RO$_2$ concentrations using Eq. (11) binned over the NO mixing ratio range encountered during the campaign on a logarithmic scale. The shading represents the 25th and 75th percentile confidence limits. The number of data points in each of the NO bins is $\approx$ 80.

4 Conclusions

Measurement and model comparisons of OH, HO$_2$, complex RO$_2$, simple RO$_2$ and total RO$_2$ in Beijing have displayed varying levels of agreement as a function of NO$_x$. Under low NO conditions, consistent with previous studies in low NO$_x$ but high VOC environments, a missing OH source is evident. Radical budget analysis has demonstrated that this missing OH source could be resolved if unimolecular reactions of RO$_2$ radicals generate OH directly. Under the low NO conditions (< 1 ppbv), the MCM over-predicted HO$_2$, although this over-prediction could be resolved at very low NO mixing ratios (< 0.3 ppbv) by including a heterogeneous loss term to aerosol surfaces. This highlights that a reduction in aerosol surface area has the potential to enhance HO$_2$ concentrations and thereby increase photochemical ozone formation but only under very low NO conditions. The model under-predicted RO$_2$, most severely under high NO conditions (> 1 ppbv). Although Cl atoms could increase the concentration of RO$_2$, this enhancement was limited to times when the Cl atom concentration was elevated and could not resolve the RO$_2$ under-prediction observed at all times. In the presence of NO, the model overestimates the rate at which RO$_2$ propagates to HO$_2$, and we hypothesise that larger RO$_2$ species likely undergo multiple bimolecular reactions with NO, followed by isomerisation of the RO radical to another RO$_2$ species, before a HO$_2$ radical forms. By this process, the lifetime and the concentration of total RO$_2$ radicals are extended. The ozone production efficiency of large, complex VOCs from which these RO$_2$ species are formed may be greater than currently appreciated, and so further efforts to understand the rate at which the larger RO$_2$ species propagate to HO$_2$ (or to OH directly) and all the possible reactions they undergo are necessary to accurately model ozone levels in urban centres such as Beijing and to fully understand how emission controls will impact ozone.

Data availability. Data presented in this study are available from the author upon request (l.k.whalley@leeds.ac.uk).

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Author contributions. LKW, EJS, RWM, CY and DEH carried out the measurements. LKW and EJS developed the model and performed the calculations. JDL, FS, JRH, RED, MS, JFH, ACL, AM, SDW, AB, TJB, HC, BO, RLJ, LRC, LJK, WJB, TV, SK, SG, YS, WX, SY, LR, WJFA, CNH, XW and PF provided logistical support and supporting data to constrain the model. LKW prepared the manuscript, with contributions from all the co-authors.
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