1 Nighttime measurements of HO_x during the RONOCO

2 project and analysis of the sources of HO₂

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- 4 H.M. Walker¹, D. Stone¹, T. Ingham^{1,2}, S. Vaughan¹, B. Bandy³, M. Cain^{4,5}, R.L.
- 5 Jones⁴, O.J. Kennedy⁴, M. McLeod⁴, B. Ouyang⁴, J. Pyle^{4,5}, S. Bauguitte^{6,7}, G.
- 6 Forster^{8,9}, M.J. Evans^{10,11}, J.F. Hamilton¹⁰, J.R. Hopkins^{10,11}, J.D. Lee^{10,11}, A.C.
- 7 Lewis^{10,11}, R.T. Lidster¹⁰, S. Punjabi^{10,11}, W.T. Morgan^{12,13}, D.E. Heard^{1,2}

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- 9 [1] {School of Chemistry, University of Leeds, Leeds, UK}
- 10 [2] {National Centre for Atmospheric Science, University of Leeds, Leeds, UK}
- 11 [3] {School of Environmental Sciences, University of East Anglia, Norwich, UK}
- 12 [4] {Department of Chemistry, University of Cambridge, Cambridge, UK}
- 13 [5] {National Centre for Atmospheric Science, University of Cambridge, Cambridge, UK}
- 14 [6] {Facility for Airborne Atmospheric Measurements (FAAM), Cranfield University,
- 15 Cranfield, UK}
- 16 [7] {National Centre for Atmospheric Science, FAAM, Cranfield University, Cranfield, UK}
- 17 [8] {Centre for Ocean and Atmospheric Sciences, School of Environmental Sciences,
- 18 University of East Anglia, Norwich, UK}
- 19 [9] {National Centre for Atmospheric Science, University of East Anglia, Norwich, UK}
- 20 [10] {Wolfson Atmospheric Chemistry Laboratories, Department of Chemistry, University of
- 21 York, York, UK}
- 22 [11] {National Centre for Atmospheric Science, University of York, York, UK}
- 23 [12] {School of Earth, Atmospheric and Environmental Sciences, University of Manchester,
- 24 Manchester, UK}
- 25 [13] {National Centre for Atmospheric Science, University of Manchester, Manchester, UK}

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27 Correspondence to: D. E. Heard (d.e.heard@leeds.ac.uk)

Abstract

Measurements of the radical species OH and HO₂ were made using the Fluorescence Assay by Gas Expansion (FAGE) technique during a series of nighttime and daytime flights over the UK in summer 2010 and winter 2011. OH was not detected above the instrument's 1σ limit of detection during any of the nighttime flights or during the winter daytime flights, placing upper limits on [OH] of 1.8 × 10⁶ molecule cm⁻³ and 6.4 × 10⁵ molecule cm⁻³ for the summer and winter flights, respectively. HO₂ reached a maximum concentration of 3.2 × 10⁸ molecule cm⁻³ (13.6 pptv) during a nighttime flight on 20th July 2010, when the highest concentrations of NO₃ and O₃ were also recorded. Analysis of the rates of reaction of OH, O₃, and the NO₃ radical with measured alkenes indicates that the summer nighttime troposphere can be as important for the processing of VOCs as the winter daytime troposphere. Analysis of the instantaneous rate of production of HO₂ from the reactions of O₃ and NO₃ with alkenes has shown that, on average, reactions of NO₃ dominated nighttime production during winter.

1 Introduction

Trace gases emitted into the atmosphere, including pollutants and greenhouse gases, are removed primarily by oxidation. The hydroxyl radical, OH, is the most important oxidising species in the daytime troposphere, reacting with numerous species including volatile organic compounds (VOCs), CO, SO₂, and long-lived anthropogenic halogenated compounds. During the day, primary production of OH (i.e. initialisation of the radical chain) occurs predominantly via photolysis of ozone at $\lambda \leq 340$ nm followed by reaction of the resulting electronically excited oxygen atom, O(¹D), with water vapour. The OH-initiated oxidation of VOCs leads to the production of the hydroperoxy radical, HO₂, and together the two radicals form the HO_x family. A key reaction in the conversion of OH to HO₂ is the reaction with CO:

$$54 \qquad OH + CO \rightarrow H + CO_{2} \tag{R1}$$

$$55 \qquad H + O_2 + M \rightarrow HO_2 + M \tag{R2}$$

Reaction of OH with VOCs results in the production of organic peroxy radicals, RO₂:

$$57 \qquad OH + RH \rightarrow H_2O + R \tag{R3}$$

$$58 R + O_2 + M \rightarrow RO_2 + M (R4)$$

Reactions of HO₂ and RO₂ with NO propagate the HO_x radical chain, regenerating OH:

$$60 RO2 + NO \rightarrow RO + NO2 (R5)$$

$$61 RO + O_2 \rightarrow R'O + HO_2 (R6)$$

$$62 \qquad HO_2 + NO \rightarrow OH + NO_2 \tag{R7}$$

- The production of OH through photolysis of ozone (and other species at longer wavelengths)
 is limited to daylight hours, and oxidation of trace gases at night proceeds through alternative
- 65 mechanisms. Two mechanisms are known to initiate HO_x radical chemistry and oxidation
- 66 chemistry at night: ozonolysis of alkenes, and reactions of the nitrate radical, NO₃, with
- 67 alkenes.
- 68 Reactions of ozone with alkenes occur via addition of ozone to the double bond to form a 69 five-membered ring called a primary ozonide. The primary ozonide decomposes to form one 70 of two possible pairs of products, each pair consisting of a carbonyl compound and a 71 vibrationally- and rotationally-excited carbonyl oxide termed a Criegee intermediate (CI). 72 The simplest gas-phase CI, CH₂OO, and the alkyl-substituted CH₃CHOO, have been 73 observed directly by photoionisation mass spectrometry (Taatjes et al., 2008; Beames et al., 74 2012; Taatjes et al., 2012; Welz et al., 2012; Beames et al., 2013; Taatjes et al., 2013; Stone 75 et al., 2014a), by infrared absorption spectroscopy (Su et al., 2013), and by microwave 76 spectroscopy (Nakajima and Endo, 2013, 2014). Excited CIs may be stabilised by collision 77 with surrounding molecules (Donahue et al., 2011; Drozd and Donahue, 2011), or may 78 undergo isomerisation or decomposition to yield products including OH, H, and subsequently 79 HO₂ (Paulson and Orlando, 1996; Kroll et al., 2001a; Kroll et al., 2001b; Kroll et al., 2002; 80 Johnson and Marston, 2008). Stabilised CIs (SCIs) are known to react with a variety of 81 compounds, including H₂O, NO₂, SO₂, and a variety of organic compounds (e.g. Mauldin III 82 et al., 2012; Taatjes et al., 2012; Ouyang et al., 2013; Taatjes et al., 2013; Stone et al., 2014a; 83 Taatjes et al., 2014). There is experimental evidence for the formation of OH from thermal 84 decomposition of SCIs, on a much longer timescale than the decomposition or isomerisation of excited CIs (Kroll et al., 2001a; Kroll et al., 2001b). The OH produced through these 85 86 ozonolysis mechanisms will proceed to oxidise other VOC species. Criegee intermediates 87 formed in the ozonolysis of alkenes are known to be an important source of HO_x during the 88 day and at night (Paulson and Orlando, 1996; Donahue et al., 1998; Kanaya et al., 1999;

- 89 Salisbury et al., 2001; Geyer et al., 2003; Ren et al., 2003a; Heard et al., 2004; Harrison et al.,
- 90 2006; Ren et al., 2006; Sommariva et al., 2007). The gas-phase ozonolysis of unsaturated
- VOCs, and in particular the role and subsequent chemistry of the Criegee intermediate, have
- been reviewed in detail by Johnson and Marston (2008), Donahue et al. (2011), Vereecken
- and Francisco (2012), and Taatjes et al. (2014).
- Another key nighttime oxidant, NO₃, is formed primarily by reaction of NO₂ with ozone.
- NO₃ reacts with a range of species in the troposphere, and its reaction with alkenes is known
- to be an important nighttime oxidation mechanism (Salisbury et al., 2001; Geyer et al., 2003;
- 97 Sommariva et al., 2007; Emmerson and Carslaw, 2009; Brown et al., 2011). The reaction
- 98 between NO₃ and an alkene proceeds primarily via addition to a double bond, to form a
- 99 nitrooxyalkyl radical, R-ONO₂. At atmospheric pressure, the main fate of the nitrooxyalkyl
- 100 radical is reaction with O₂ (Berndt and Böge, 1994) to produce a nitrooxyalkyl peroxy
- radical, O₂–R–ONO₂. The nitrooxyalkyl peroxy radical can react with NO₂, HO₂, RO₂, NO
- and NO₃, of which the latter two reactions lead to formation of the nitrooxyalkoxy radical,
- 103 O-R-ONO₂. The nitrooxyalkoxy radical can undergo isomerisation, decomposition, or
- reaction with O_2 . Reaction with O_2 , analogous to the reaction of organic alkoxy radicals,
- 105 yields HO_2 :

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$$O-R-ONO_2 + O_2 \rightarrow O-R'-ONO_2 + HO_2$$
 (R8)

- 107 Thus, nighttime oxidation of hydrocarbons by NO₃ leads to production of HO₂. Reaction of
- 108 HO₂ with NO (Reaction (R7)), O₃ and NO₃ can generate OH:

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$$HO_2 + O_3 \rightarrow OH + 2O_2$$
 (R9)

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$$HO_2 + NO_3 \rightarrow OH + NO_2 + O_2$$
 (R10)

- 111 Atkinson and Arey (2003) published a detailed review of tropospheric degradation of VOCs,
- including reaction with O₃ and NO₃. A comprehensive review of nighttime radical chemistry
- is given by Brown and Stutz (2012).
- The oxidising capacity of the nocturnal troposphere is thought to be controlled by the
- reactions described above, with a limited role for OH and HO₂ due to the absence of their
- photolytic sources. Oxidation of VOCs at night can have significant effects on daytime air
- quality and tropospheric ozone production (Brown et al., 2004; Brown et al., 2006; Wong and
- 118 Stutz, 2010; Brown et al., 2011). Several field measurement campaigns have involved
- nighttime measurements of OH, HO₂, RO₂, and NO₃ (see Table 1), and have highlighted the
- importance of the vertical profile of nighttime radical concentrations and chemistry (Geyer

121 and Stutz, 2004a, b; Stutz et al., 2004; Volkamer et al., 2010), but prior to the current work there had been no aircraft-based studies of nighttime chemistry involving measurements of 122 123 both NO₃ and HO₂, to enable vertical profiling of the lower atmosphere and full evaluation of 124 the nocturnal radical budget. Table 1 gives details of some previous measurements and 125 modelling of nighttime HO_x concentrations in polluted or semi-polluted environments. 126 Highlights from these studies are discussed here, with particular attention paid to those 127 involving measurements of HO_x, NO₃, and O₃, and in which the contributions made by O₃ and NO₃ to nighttime radical chemistry have been considered. 128 129 Geyer et al. (2003) report radical measurements and modelling from the 1998 Berliner Ozone 130 Experiment (BERLIOZ). Measurements of NO₃, RO₂, HO₂ and OH were made by matrix isolation electron spin resonance (MIESR), chemical amplification (CA), and laser-induced 131 fluorescence (LIF) spectroscopy at a site approximately 50 km from Berlin. HO₂ was 132 detected at night with concentrations frequently as high as 5×10^7 molecule cm⁻³ 133 (approximately 2 pptv), and an average concentration of 1×10^8 molecule cm⁻³ over one hour 134 (02:00 to 03:00) of nocturnal measurements during an intensive period of the study (Holland 135 136 et al., 2003). OH was usually below the limit of detection of the LIF instrument $(3.5 \times 10^5 \text{ molecule cm}^{-3})$. Modelling revealed that nitrate radical reactions with terpenes 137 were responsible for producing 53 % of HO₂ and 36 % of OH radicals in the night, with 138 139 ozonolysis accounting for production of the remaining 47 % of HO₂ and 64 % of OH 140 radicals. A positive linear correlation between RO2 and NO3 was observed and was 141 reproduced by the model. 142 Reactions of O₃ with alkenes were found to be responsible for the majority of formation of OH during the winter PUMA (Pollution of the Urban Midlands Atmosphere) campaign (a 143 low photolysis urban environment) (Heard et al., 2004; Emmerson et al., 2005; Harrison et 144 145 al., 2006). Measurements of OH, HO2 and RO2 were unavailable at night, but model-146 predicted values of these radicals were used to calculate that 90 % of nighttime initiation via 147 HO₂ was from O₃ reactions. Without measurements of NO₃ during the campaign, there was 148 no estimate of its contribution to radical initiation. 149 Modelling results from the MCMA-2003 (Mexico City) field campaign (Volkamer et al., 150 2010) indicate that nighttime radical production at roof-top level (approximately 16 m above 151 the ground) was dominated by ozonolysis of alkenes, and that reactions of NO₃ with alkenes 152 played only a minor role. The measurement site was located in a polluted urban environment, 153 with high levels of NO, NO₂ and O₃. NO₃ was observed at a maximum concentration of 50 154 pptv during the night at a mean height above the ground of 70 m. Roof-top level concentrations of NO₃ were estimated using a linear scaling factor calculated from the 155 156 observed O₃ vertical gradient, and were found to be, on average, 3 times lower than the 157 concentrations measured at 70 m. This predicted vertical gradient accounts for the relative 158 unimportance of NO₃ reactions in radical initiation at roof-top level. Propagation of RO₂ 159 radicals to HO₂ and OH, by reaction with NO₃, was found to be negligible. 160 The 2006 Texas Air Quality Study (TexAQS) involved a series of nighttime flights onboard 161 the NOAA P-3 aircraft over Houston, Texas, and along the Gulf Coast (Brown et al., 2011). Loss rates and budgets of NO₃ and highly reactive VOCs were calculated, but there were no 162 163 measurements of OH, HO₂ and RO₂ during the flights. Budgets for NO₃ show that it was lost 164 primarily through reactions with unsaturated VOCs, but the contribution to NO₃ loss through 165 reaction with peroxy radicals was uncertain because of the lack of direct measurements of 166 RO₂ during the flights. NO₃ dominated VOC oxidation, being 3 to 5 times more important 167 than O_3 . 168 In summary, NO₃ and O₃ have both been found to dominate radical initiation in the nighttime 169 troposphere, and in some situations the two mechanisms were found to be equally important. 170 The relative importance of O₃- and NO₃-initiated oxidation depends on the availability of 171 NO₃, which is determined by the amount of NO_x present in the atmosphere and the ratio of 172 NO to NO₂, and the concentration and species distribution of VOCs (Bey et al., 2001; Geyer et al., 2003). A modelling study by Bey et al. (2001) suggests that nocturnal radical initiation 173 174 is driven by alkene ozonolysis in urban environments or in environments with low NO_x concentrations, while both O₃ and NO₃ contribute to radical initiation in rural environments 175 176 with moderate NO_x levels. It is expected that NO₃ dominates nocturnal radical initiation in air 177 masses containing sufficient NO₂ and O₃ for NO₃ production while being deprived of NO 178 (e.g. air masses downwind of urban areas). Geyer and Stutz (2004b) have found that the 179 effects of suppressed mixing in the nocturnal boundary layer can also control whether NO₃ or 180 O₃ dominates nighttime radical chemistry. 181 In this paper we report airborne measurements of OH and HO₂ made during the RONOCO 182 (ROle of Nighttime chemistry in controlling the Oxidising Capacity of the atmosphere) and 183 SeptEx (September Experiment) projects in 2010 and 2011. The rates of reaction between O₃, 184 NO₃, and OH with the alkenes measured during the flights are investigated. Analysis of 185 radical production from the nighttime reactions of O₃ and NO₃ with alkenes is also given. 186 Comparisons are made between the daytime and nighttime chemistry studied, and between the summer and winter measurement periods. Details and results of a box modelling study, and comparison to the observations, are given by Stone et al. (2014b).

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2 Details of the RONOCO and SeptEx fieldwork

191 RONOCO is a NERC-funded consortium project aimed at improving our understanding of 192 the mechanisms and impact of nocturnal oxidation chemistry over the UK. The RONOCO 193 fieldwork consisted of two measurement campaigns, in July 2010 and January 2011. An 194 additional fieldwork period, SeptEx, was conducted in September 2010. The RONOCO and 195 SeptEx flights were conducted onboard the BAe-146 research aircraft operated by the Facility 196 for Airborne Atmospheric Measurements (FAAM). Both field measurement campaigns were 197 based at East Midlands Airport (52.8° N, 1.3° W) in the UK. During RONOCO the majority 198 of the flying took place at night, with occasional flights beginning or ending in daylight hours 199 to study chemical behaviour at dusk and dawn. Flights during SeptEx were mainly during the 200 day, providing a useful comparison to the nocturnal chemistry. 201 Flights were conducted between altitudes of 50 m and 6400 m, above the UK and the North 202 Sea. Figure 1 shows the flight tracks during the summer, SeptEx, and winter measurements 203 coloured by altitude. Measurements of OH and HO₂ were made using the University of Leeds 204 aircraft-based Fluorescence Assay by Gas Expansion (FAGE) instrument. A suite of 205 supporting measurements, including CO, O₃, NO, H₂O, VOCs, NO₃, and HCHO, were made 206 during the flights and have been used in the current work. Table 2 summarises the techniques 207 used to measure these species. 208 Air mass histories for each flight have been calculated using the UK Met Office Numerical 209 Atmospheric-dispersion Modelling Environment (NAME). NAME is a three-dimensional 210 Lagrangian particle dispersion model (Jones et al., 2007) which is run here using the UK 211 Meteorological Office's Unified Model meteorological fields. Model 'particles', restricted to 212 a 300 m deep layer from the surface, were released along the flight path and were tracked 213 backwards through the modelled atmosphere. Model particle densities were integrated over 214 24 h periods, beginning at 24, 48, 72, and 96 hours before each flight. The resulting 215 'footprint' maps show the regions where the measured air has been in contact with the 216 surface over the 4 days preceding a flight. An example is shown in Fig. 2, which shows 217 model particle densities integrated over the 24 hour period beginning 48 hours prior to flight 218 B535. The majority of the summer flights were characterised by air masses originating from the west and south west of the UK, having Atlantic or continental European influences. The SeptEx flights were predominantly influenced by air masses from the north-east, east, and south-east of the UK, with northern European influences. The winter flights were mainly characterised by air masses arriving from the west of the UK, bringing Atlantic influences.

Table 3 gives mean and maximum mixing ratios of CO, O₃, NO, and NO₂ measured during RONOCO and SeptEx. The mean mixing ratios of NO measured during the summer RONOCO flights are much lower than ground-based nighttime measurements (e.g. 1.0 ppbv during TORCH (Emmerson and Carslaw, 2009), 0–20 ppbv during PMTACS-NY, (Ren et al., 2006)), but are comparable with previous airborne nighttime measurements (e.g. < 30 pptv during TexAQS (Brown et al., 2011)). Mean values of NO up to 14 pptv were reported by Salisbury et al. (2001) for semi-polluted air masses sampled at Mace Head. These comparisons indicate that the RONOCO and SeptEx flights enabled sampling of air masses generally removed from the influence of NO in fresh surface emissions. Table 3 also highlights the unusual chemical conditions encountered during flight B537 on 20th July 2010, discussed further in Sect. 4.1. Nighttime altitude profiles of NO₃, O₃, *trans*-2-butene, and propene (the latter two being illustrative of the alkenes measured) are given in Fig. 3.

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3 Experimental

3.1 The Leeds FAGE aircraft instrument

- 238 The University of Leeds Aircraft FAGE instrument has been described in detail by Commane 239 et al. (2010). A brief description is given here. The instrument, which was designed 240 specifically for use onboard the FAAM BAe-146 research aircraft (Floquet, 2006), is housed 241 in two double-width 19 inch aircraft racks, with the inlet, detection cells, and pump set being 242 separate to the two racks. Ambient air is sampled through a 0.7 mm diameter 'pinhole' into a 243 cylindrical inlet (length = 50 cm, diameter = 5 cm) which extends through a window blank on
- the starboard side of the aircraft.
- Downstream of the inlet are two low pressure fluorescence cells positioned in series, the first for detection of OH and the second for detection of HO₂. During the RONOCO and SeptEx
- 247 flights the pressure inside the cells ranged from 1.9 Torr at ground level to 1.2 Torr at 6 km.
- 248 Laser light at $\lambda \sim 308$ nm is generated by a diode-pumped Nd:YAG-pumped tunable
- 249 Ti:Sapphire laser (Photonics Industries DS-532-10 and TU-UV-308nm) and delivered to the

- fluorescence cells via optical fibres, on an axis perpendicular to the gas flow. A small fraction of the Ti:Sapphire second harmonic ($\lambda = 462$ nm) is directed to the probe of a wavemeter to enable measurement of the laser wavelength to within 0.001 nm. A UV photodiode is positioned opposite the laser input arm on each fluorescence cell to measure laser power.
- The sampled air forms a supersonic gas expansion beam in which the rate of collision between OH radicals and ambient air molecules is reduced. The OH fluorescence lifetime is therefore extended to several hundred nanoseconds, significantly longer than the laser pulse, so that the measured signal can be temporally discriminated from laser scattered light. OH is excited from its ground state, $X^2\Pi_i(v''=0)$, to its first electronically excited state, $A^2\Sigma^+(v'=0)$, at $\lambda \sim 308$ nm. The resulting on-resonance fluorescence is detected by a UV-sensitive channel photomultiplier tube on an axis perpendicular to both the gas flow and the
- laser light. HO₂ is detected by titration with an excess of NO (Reaction (R7)), the resulting
- 262 OH being detected as described.
- The FAGE instrument was calibrated prior to and following each field measurement period,
- using a well-established method (Edwards et al., 2003; Faloona et al., 2004; Commane et al.,
- 265 2010). Light at $\lambda = 184.9$ nm from a mercury pen-ray lamp photolyses water vapour in a flow
- of synthetic air inside an aluminium flow tube, generating OH and HO_2 at known
- 267 concentrations. The aircraft FAGE instrument's limit of detection (LOD) for OH and HO_2 is
- 268 determined by the instrument's sensitivity and the standard deviation of the background
- signal. During the RONOCO and SeptEx fieldwork the 1σ LOD for a 5 minute averaging
- 270 period ranged between 0.64 and 1.8×10^6 molecule cm⁻³ for OH and between 5.9 and
- 271 6.9×10^5 molecule cm⁻³ for HO₂.

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3.2 RO₂-based interference in FAGE measurements of HO₂

- It has recently been shown that reaction of alkene-derived β -hydroxyalkyl peroxy radicals, RO₂, with NO inside the HO₂ detection cell, can lead to interference in FAGE HO₂
- 275 measurements (Fuchs et al., 2011; Whalley et al., 2013). The magnitude of the interference
- depends on the parent alkene, the residence time and mean temperature inside the cell, and
- the amount of NO injected. The interference therefore depends on the chemical environment
- and differs between FAGE instruments. In view of this, the University of Leeds ground-based
- and aircraft FAGE instruments have been tested for RO₂ interference. Thorough descriptions
- of the ground-based experimental method and results, and the results of a modelling study,

are given by Whalley et al. (2013). The strongest interference in the aircraft instrument measurements was observed for ethene-derived RO_2 , amounting to an increase of 39.7 ± 4.8 % in the observed HO_2 signal, with a cell pressure of 1.8 Torr, an estimated detection cell temperature of 255 K (obtained from rotational excitation spectra performed previously), and $[NO]_{cell} = 10^{14}$ molecule cm⁻³.

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Whalley et al. (2013) show that the chemistry responsible for the observed interferences is well known, and that a model using the Master Chemical Mechanism (MCM, version 3.2: Jenkin al., 1997; Saunders al., 2003; al., 2005, et et **Bloss** et http://mcm.leeds.ac.uk/MCM) can reproduce the interferences once tuned to the conversion efficiency of HO₂ to OH in the FAGE detection cell. Accordingly, Stone et al. (2014b) have applied the results of the ethene-derived RO2 interference testing in a modelling study to assess the effect of the interference on the HO₂ measurements made during the RONOCO and SeptEx campaigns. A box model using a detailed MCM scheme was used to calculate a total potential interference in the RONOCO HO₂ measurements. The model was constrained to the conditions in the detection cell (1.8 Torr, 255 K, [NO] $\sim 10^{14}$ molecule cm⁻³). Equal concentrations of HO_2 and $\sum RO_2$ (sum of all peroxy radicals in the MCM generated from the parent hydrocarbon) were used to initialise the model. The model run time was varied until the model-predicted interference from ethene-derived RO₂ radicals was equal to the experimentally determined interference, thereby tuning the model to the conversion efficiency of HO₂ to OH. An interference factor, f, was calculated for each RO₂ in the MCM as follows:

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$$f = \frac{[OH]_{HO_2 + RO_2} - [OH]_{HO_2}}{[OH]_{HO_2}}$$
 (1)

where $[OH]_{HO_2+RO_2}$ and $[OH]_{HO_2}$ are the modelled concentrations of OH produced from the reactions of RO₂ and HO₂, and the concentration from HO₂ alone, respectively. The greatest interference was calculated to come from isoprene-derived peroxy radicals, followed by aromatic compounds and C₂ to C₅ alkenes. The smallest modelled interference is from the C₁ to C₃ alkanes. The interference factors were applied to model-predicted RO₂ speciation and concentrations for the RONOCO flights. Model-predicted RO₂ species were dominated by CH₃O₂ (33 %; f = 1.1 %) and HO₂ (24 %; f = 0.0 %), with smaller contributions from RO₂ derived from *iso*-butene (12 %; f = 0.5 %), *cis*-2-butene and *trans*-2-butene (10 %; f = 0.05 %), and isoprene (2 %; f = 7.6 %). RO₂ species with high interference factors were a minor

- 312 component of the total RO₂. A modelled value of HO₂ including the total potential
- 313 interference, HO₂*, was calculated using:

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$$[HO_2^*] = [HO_2]_{mod} + f[RO_2]_{mod}$$
 (2)

- 315 Direct comparison between modelled values of [HO₂*] and the FAGE-measured values of
- 316 [HO₂] was therefore made possible. The model-predicted interference during the RONOCO
- campaign is described by $[HO_2]^* = 1.15[HO_2] + 2 \times 10^5$ molecule cm⁻³. The average model-
- 318 predicted interference in the HO₂ measurements is 14 %. The HO₂ measurements made
- during RONOCO and SeptEx were not adjusted since speciated RO₂ measurements were not
- 320 available. The measurements are hereafter referred to as HO_2^* .
- 321 The magnitude of the RO₂ interference can be reduced by reducing the concentration of NO
- in the detection cell. This also reduces the instrument sensitivity to HO₂. Since conversion of
- 323 RO₂ to OH requires at least two NO molecules, while conversion of HO₂ requires only one
- molecule, the ratio of HO₂ signal to RO₂ signal can be made favourable by reducing [NO]
- 325 (Whalley et al., 2013). This effect has been investigated for the ground-based instrument, and
- will be investigated for the aircraft instrument prior to future HO_x measurement campaigns.
- 327 An overview of the laboratory and computational studies of the interference in different
- 328 FAGE instruments is given in a recent review by Stone et al. (2012).

329 3.3 BBCEAS measurements of NO₃ and N₂O₅

- NO₃ and N₂O₅ were measured by the University of Cambridge broadband cavity enhanced
- 331 absorption spectroscopy (BBCEAS) instrument. The instrument was designed and built
- 332 specifically for the RONOCO project and is described in detail in Kennedy et al. (2011). A
- brief description is given here.
- The instrument consists of three 94 cm long high finesse optical cavities formed by pairs of
- 335 highly reflecting mirrors. The cavities are irradiated by incoherent broadband continuous
- wave light sources. Two of the cavities, for the detection of N₂O₅ and NO₃, are irradiated by
- red light emitting diodes (LEDs) centred at 660 nm. The third cavity, for the detection of
- NO₂, is irradiated by a blue LED centred at 460 nm. The light from the LEDs is collimated
- 339 using optical fibres and a focussing lens at the input of each cavity. A spectrometer,
- consisting of a spectrograph and charge couple device (CCD), is positioned at the end of each
- cavity to measure the wavelength-dependent intensity of transmitted light.

Ambient air is sampled through a rear-facing inlet on the aircraft fuselage, positioned approximately 4 m from the aircraft nose and 10 cm from the aircraft body. The air from the inlet is divided into two flows. The flow directed to the N₂O₅ cavity is heated to 120 °C to ensure near complete (> 99.6 %) thermal dissociation of N_2O_5 to NO_2 and NO_3 . The cavity itself is heated to 80 °C and is used to measure the sum of the concentrations of ambient NO₃ plus NO₃ from thermal decomposition of N₂O₅. The second flow is unheated and is directed first through the NO3 cavity and then through the NO2 cavity. Background spectra are recorded at half hour intervals during flights by halting the flow of ambient air and purging the cavities with nitrogen. NO_3 is detected by its strong $B^2E'-X^2A'_2$ electronic transition centred at 662 nm. The concentration of NO₃ is determined by separating the finely structured NO₃ absorption features from the broad features caused by Rayleigh and Mie scattering using a fitting technique analogous to that employed in differential optical absorption spectroscopy (DOAS). A strong water vapour absorption feature that spectrally overlaps with NO₃ absorption around 662 nm is simulated for the pressure and temperature measured in the cavity and is removed from the measured absorption spectrum. The concentration of N₂O₅ is

Contributions to uncertainties in ambient measurements of NO_3 and N_2O_5 , including wall losses of NO_3 and N_2O_5 , temperature- and pressure-dependent absorption cross sections of NO_3 and H_2O , and the length of the cavity occupied by the sample, have been thoroughly investigated in laboratory experiments or addressed in the data analysis routine. In addition, wall losses of NO_3 and N_2O_5 were determined before and after each flight to account for changes in the surface properties of the inlet and detection cell walls, which were found to be negligible. The total uncertainty in the measured concentration of ambient NO_3 was 11 %. The uncertainty in the measured concentration of ambient N_2O_5 is determined for each individual ambient measurement, being dependent on the NO_3/N_2O_5 ratio, and was on the order of 15 %. During RONOCO flights the 1σ limits of detection for NO_3 and the sum of $NO_3 + N_2O_5$ were 1.1 pptv and 2.4 pptv, respectively, for a 1 second integration time.

determined by subtracting the concentration of ambient NO₃ measured in the unheated cavity

from the sum of the concentrations of ambient and dissociated NO₃ measured in the heated

cavity.

4 Overview of OH and HO₂* measurements

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374 FAGE measurements were made on 16 flights during RONOCO and 9 flights during SeptEx. 375 There was insufficient laser power during flights B534 to B536 in the summer campaign to 376 measure both OH and HO₂* by dividing the laser light between the two cells. OH was 377 therefore not measured during these flights. Low laser power throughout the summer fieldwork caused relatively high fluctuations in laser power and therefore higher background 378 variability. This resulted in higher limits of detection for OH $(1.8 \times 10^6 \text{ molecule cm}^{-3})$ and 379 $HO_2* (6.9 \times 10^5 \text{ molecule cm}^{-3}).$ 380 381 Table 4 summarises the OH and HO₂* measurements during RONOCO and SeptEx and gives 382 the instrument's average 1 σ limit of detection for a 5 minute averaging period. OH was not detected above the limit of detection during the summer or winter RONOCO flights, resulting 383 in upper limits of 1.8×10^6 molecule cm⁻³ and 6.4×10^5 molecule cm⁻³ for mean summer and 384 winter concentrations, respectively. These upper limit values are similar to previously 385 386 reported nighttime OH measurements (Geyer et al., 2003; Holland et al., 2003; Ren et al., 2003b; Emmerson and Carslaw, 2009). The mean daytime OH concentration during SeptEx 387 was 1.8×10^6 molecule cm⁻³, which was above the limit of detection. The mean HO₂* 388 mixing ratio was highest during SeptEx (2.9 pptv), and was higher during summer (1.6 pptv) 389 390 than during winter (0.7 pptv). The OH and HO₂* datasets for RONOCO and SeptEx are shown as altitude profiles in Fig. 4 and Fig. 5, respectively. 391 Table 5 gives the mean and maximum HO₂* mixing ratios at different times of day during 392 393 summer, SeptEx and winter. Dawn, day, dusk and night are defined by the solar zenith angle 394 as follows: dawn and dusk are between 90 and 102° and are distinguished by the time of day; day is between 0° and 90° ; night is between 102° and 180° . 395 396 The mean dusk HO₂* mixing ratio in summer was higher than the mean nighttime mixing 397 ratio, suggesting that photochemical production was still active at dusk in summer. The 398 reverse was true for the winter data, with the highest mean HO₂* mixing ratio being at night. 399 This suggests that when photochemical production was suppressed in the winter daytime due to low photolysis rates, production via reactions of NO₃ and O₃ with alkenes was an 400 401 important route to radical initiation. The RONOCO HO2* measurements are similar to 402 nighttime, ground-based, urban measurements. For example, during the TORCH campaign, [HO₂] peaked at 1×10^8 molecule cm⁻³ at night (Emmerson et al., 2007), and during the 403

404 PMTACS-NY 2001 field campaign, nighttime HO_2 concentrations of 8×10^6 molecule cm⁻³

were measured (Ren et al., 2003b).

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4.1 Case study flight B537: high nighttime HO₂* concentrations

- The highest HO_2^* concentration (3.2 × 10^7 molecule cm⁻³; 13.7 pptv) was measured during
- 408 nighttime flight B537 on 20th July 2010. Take-off from East Midlands Airport was at 22:00
- local time (21:00 UTC, sunset at 20:18 UTC). The flight track, coloured by altitude, is shown
- 410 in Fig. 6. The flight involved a profile descent from 3350 m to 460 m down the Norfolk coast
- and a missed approach at Southend Airport (51.6° N, 0.70° E). Plumes from European
- 412 continental outflow (see Fig. 7) were intersected by a series of runs at altitudes between 460
- and the upper boundary of the polluted layer.
- 414 Flight B537 is an unusual flight within the RONOCO dataset, with high concentrations of
- 415 CO, O₃, NO₃, and high temperatures compared to the values measured during other nighttime
- 416 flights (see Table 3). The ambient aerosol surface area was significantly higher during B537
- 417 (nearly 800 µm² cm⁻³) than during other flights (between 100 and 400 µm² cm⁻³), and the
- organic aerosol concentration was significantly enhanced (Morgan et al., 2014). Footprint
- 419 maps for flight B537, indicating regions where the sampled air was in contact with the
- surface prior to the flight, are shown in Fig. 7. The air sampled during the flight originated
- primarily over northern France, Belgium and Germany.
- 422 A region of high surface pressure was positioned over the UK on the 20th July, with a mean
- air pressure of 1012.6 hPa over the 24 hours prior to the flight. The mean air temperature 24
- 424 hours prior to the flight (22:00 19/07/2010 to 22:00 20/07/2010), measured at a number of
- 425 Met Office weather stations in Greater London, was 22.6 °C, and reached a maximum value
- of 28.6 °C. Wind speeds prior to the flight were low, with an average value of 4.7 knots
- 427 (2.4 m s⁻¹). No rainfall was recorded at any of the Greater London weather stations during the
- 428 24 hours prior to the flight. 12.4 hours of sunshine were recorded on the 20th July at
- 429 Heathrow Airport (51.5 °N, 0.45 °W). High temperatures, combined with low wind speed,
- 430 exposure to solar radiation, and little precipitation promote the formation of ozone as a result
- of photochemical processing of VOCs emitted at the surface (e.g. Lee et al., 2006), and offer
- an explanation for the high ozone mixing ratios measured during flight B537. Peak surface
- daytime ozone concentrations measured in Teddington, London, on 20th July were on the
- order of 2.0×10^{12} molecule cm⁻³ (~78 ppbv) (data available at www.airquality.co.uk).
- Similar levels were recorded at a number of locations within Greater London.

Figure 8 shows a time series of altitude, HO_2^* , O_3 , and NO_3 mixing ratios during the flight, demonstrating very similar behaviour between the two radical species. During the missed approach at Southend Airport the mixing ratios of HO_2^* and NO_3 increased with decreasing altitude, to reach values of 4.5 pptv and 35 pptv, respectively, at 50 m above the ground. The maximum HO_2^* and NO_3 mixing ratios were measured over the North Sea east of Ipswich (52.16 ° N, 2.34 ° E) at an altitude of 509 m, in the outflow of the London plume. Figure 9 shows scatter plots of HO_2^* against NO_3 and O_3 during flight B537 and during the other nighttime flights during RONOCO. Strong positive correlation is evident between HO_2^* and NO_3 during B537 (r = 0.97), while during the remaining night flights there is still a significant, though weaker, correlation (r = 0.58). Moderate negative correlation existing for the other nighttime flights (r = 0.19). The data suggest that NO_3 was an important initiator of HO_3 radicals during flight B537, and that O_3 played a limited role overall during the nighttime flights. Further investigation of the roles of NO_3 and O_3 in alkene oxidation and radical initiation at night is described in Sect. 5.

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5 Oxidation of alkenes and production of HO₂: method of analysis

- Following the work of Salisbury et al. (2001), the total rates of reaction, Φ , of O₃ and NO₃
- with the alkenes measured during RONOCO and SeptEx have been calculated:

$$\Phi_{\mathcal{O}_3} = \sum_{i}^{\text{alkene}} k_{\mathcal{O}_3 + \text{alk}_i} \left[\mathcal{O}_3 \right] \left[\text{alkene}_i \right]$$
(3)

- The reactions of O₃ and NO₃ with alkenes yield OH, HO₂, and RO₂ radicals. Consideration of
- 458 the reaction mechanisms of NO₃ and O₃ enables calculation of the rate of instantaneous
- production of HO_2 (P_{HO_3}) from the reactions of NO_3 and O_3 with the alkenes measured
- during RONOCO, using the chemistry scheme, rate constants and branching ratios in the
- 461 MCM (Jenkin et al., 1997; Saunders et al., 2003).
- 462 Figure 10 shows a generalized reaction scheme for the reaction of NO₃ with an alkene. The
- reaction between NO₃ and an alkene proceeds via addition of NO₃ to the double bond to form
- a nitrooxyalkyl radical, followed by rapid reaction with oxygen to yield a nitrooxyalkyl

465 peroxy radical, RO₂ (shown as a single step in Fig. 10). The RO₂ radical can react with a number of species, of which NO, NO₃ and RO₂ lead to production of an alkoxy radical (RO). 466 467 Radical termination occurs via reaction of RO₂ with HO₂ to yield a peroxide (ROOH) or with RO₂ to yield carbonyl (RC(O)CH₃) and alcohol (RCH₂OH) products. Reaction of RO with 468 oxygen proceeds via abstraction of a hydrogen atom to yield HO₂ or an aldehyde (RCHO). 469 This generalised scheme can be applied to the reactions of NO₃ with all the alkenes 470 471 measured. The rate of instantaneous production of HO₂ is found by first calculating the 472 fraction of RO_2 that reacts to produce $RO(F_{RO})$, and the fraction of RO that reacts to produce 473 $HO_2(F_{HO_2})$:

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$$F_{RO} = \frac{k_3[NO] + k_4[NO_3] + 0.6k_5[RO_2]}{k_2[HO_2] + k_3[NO] + k_4[NO_3] + k_5[RO_2]}$$
(5)

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$$F_{\text{HO}_2} = \frac{k_6 [O_2]}{k_7 + k_6 [O_2]}$$
 (6)

where RO_2 represents all peroxy radicals. Average values of F_{RO} for the NO_3 + alkene reactions range between 0.50 for *trans*-2-pentene- and 1-pentene-derived RO_2 species and 0.61 for ethene-derived RO_2 species. F_{HO_2} varies between 0 and 1 for the alkenes studied. Overall, the rate of production of HO_2 (P_{HO_2}) from reactions of NO_3 with alkenes is then given by:

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$$P_{\text{HO}_2} = k_i [\text{NO}_3] [\text{alkene}_i] \times F_{\text{RO}} \times F_{\text{HO}_2}$$
 (7)

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The reaction scheme for the reaction of O_3 with alkenes is more complicated because the number and type of radicals produced in the O_3 + alkene reaction depends on the structure of the alkene. The simplest case is the reaction of ozone with ethene. Ozone adds to the double bond to form a five-membered ring called a primary ozonide. Decomposition of the ozonide yields an excited Criegee intermediate (CH_2OO^*) and a carbonyl compound (in this case formaldehyde, HCHO). The energy-rich Criegee intermediate can be stabilised by collision with a third body or undergo decomposition to yield products including OH, CO, and HO_2 . The primary ozonide produced in the O_3 + propene reaction (see Fig. 11) can decompose via two channels, yielding carbonyls and Criegee intermediates with different structures and different products, including RO_2 . Reaction of RO_2 with RO_3 and RO_4 (all peroxy radicals) yields RO_4 , which in turn yields RO_4 .

493 The rates of production of HO_2 from reactions of O_3 with alkenes (P_{HO_2}) have been

494 calculated as follows:

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$$P_{\text{HO}_3,\text{Direct}} = k_i \left[O_3 \right] \text{ [alkene }_i \times \alpha_{\text{HO}_3}$$
 (8)

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$$P_{\text{HO}_2,\text{RO}_2} = k_i \left[O_3 \right] \left[\text{alkene}_i \right] \times \alpha_{\text{RO}_2} \times F_{\text{RO}} \times F_{\text{HO}_2}$$
 (9)

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$$P_{\text{HO}_2} = P_{\text{HO}_2, \text{Direct}} + P_{\text{HO}_2, \text{RO}_2}$$
 (10)

- 498 where $P_{\text{HO}_2,\text{Direct}}$ is the rate of direct HO₂ production from Criegee intermediate
- 499 decomposition, α_{HO_2} is the branching ratio to HO₂-producing channels from the Criegee
- intermediate, $P_{\text{HO}_2,\text{RO}_2}$ is the rate of HO₂ production from RO₂ radicals produced in the O₃ +
- alkene reaction, α_{RO_2} is the branching ratio to RO₂-producing channels from the Criegee
- intermediate, F_{RO} is the fraction of RO₂ radicals that react to produce RO radicals, and F_{HO} ,
- is the fraction of RO radicals that react to produce HO₂ radicals, which is equal to 1 for all the
- alkenes studied. Average values of F_{RO} for the O_3 + alkene reactions range between 0.54 for
- 505 1-pentene-derived RO₂ species and 0.64 for 1-butene- and trans-2-pentene-derived RO₂
- 506 species.
- The reactions of RO₂ with NO to form RONO₂ have been omitted from the calculations
- because the branching ratio is small (0.001 to 0.02) for the radicals studied (Carter and
- Atkinson, 1989; Lightfoot et al., 1992). The reaction of CH₃O₂ with NO₂ to form CH₃O₂NO₂
- 510 has been omitted from the calculations, since the reverse reaction is much faster than the
- forward direction ($k_f = 6.4 \times 10^{-12} \text{ s}^{-1}$; $k_{rev} = 1.08 \text{ s}^{-1}$ at a mean temperature of 286.5 K during
- 512 RONOCO).
- The primary aims of the analysis presented here are threefold: 1. To calculate the total rate of
- 514 initiation through reactions of NO₃ and O₃ with alkenes; 2. To determine the relative
- importance of NO₃ and O₃ in nighttime HO₂ production; 3. To investigate differences in
- 516 radical production between different seasons and different times of day. The correlation
- between [HO₂*] and [NO₃], especially during flight B537, will be investigated.
- $P_{\mathrm{HO_2}}$ has been calculated for each alkene measured for every 60-second data point where all
- 519 the requisite data were available and where HO₂* was above the limit of detection of the
- 520 FAGE instrument. Concentrations of RO₂ were calculated by scaling the observed HO₂*
- 521 concentrations with the RO₂/HO₂* ratio calculated using a box model constrained to the

- 522 concentrations of long-lived species measured during the flights (Stone et al., 2014b), i.e.
- $RO_{2,obs} = HO_2*_{.obs} \times RO_{2,mod}/HO_2*_{.mod}$. The rates of reaction and rates of production of HO_2 523
- 524 presented hereafter are average values for individual flights, seasons, or times of day.

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6 Results

6.1 Nighttime oxidation of alkenes

- Figure 12 shows histograms of the rate of reaction between O₃ and NO₃ with individual 528 529 alkenes during summer and winter, for the nighttime data only. The reactivity of measured alkenes ($\Phi_{O3} + \Phi_{NO3}$) was greater during summer flights than during winter flights by a factor 530 of 2.2. The reactions of NO₃ are largely responsible for this seasonal difference, since the 531 contribution from O_3 + alkene reactions varies little between summer $(4.1 \times 10^4 \text{ molecule})$ 532 cm⁻³ s⁻¹) and winter $(3.9 \times 10^4 \text{ molecule cm}^{-3} \text{ s}^{-1})$. The factor of 4.1 difference between the 533 rate of NO_3 reactions in summer (9.8 \times 10⁴ molecule cm⁻³ s⁻¹) and winter (2.4 \times 10⁴ 534 molecule cm⁻³ s⁻¹) can be attributed to the higher mean concentration of NO₃ in summer 535 $(5.8 \times 10^8 \text{ molecule cm}^{-3})$ compared to winter $(2.0 \times 10^8 \text{ molecule cm}^{-3})$. This seasonal 536 difference in NO₃ concentrations is attributable to the lower mean nighttime temperature in 537 538 winter (277.7 K) compared to summer (286.7 K) which disfavours NO₃ in the thermal equilibrium $N_2O_5 \rightleftharpoons NO_3 + NO_2$. $K_{eq}[NO_2]$, which determines $[N_2O_5]/[NO_3]$, is calculated to 539 540 be 4.8 in summer and 29.6 in winter. At night in summer, Φ_{NO3} was greater than Φ_{O3} by a 541 factor of 2.4, but in winter Φ_{O3} was a factor of 1.6 greater than Φ_{NO3} . Figure 12 illustrates the 542 importance of the butene isomers (within the VOCs measured) in the reactions of O₃ and NO₃, and therefore radical initiation and propagation. Reactions with iso-butene dominated 543 NO₃ reactivity in summer (42 %) and winter (53 %), with trans-2-butene also contributing 544 significantly (28 % in summer and 32 % in winter). Reactions of O₃ were dominated by 545 546 trans-2-butene (42 % in summer and 34 % in winter) and propene (26 % in summer and 38 % 547 in winter). The importance of these alkenes is attributed to their relatively high abundances 548 compared to the other alkenes measured, during both summer and winter, combined with 549 their fast rates of reaction with O₃ and NO₃. 550 For comparison with the reactions of O₃ and NO₃, the total rate of reaction of measured
- alkenes with OH has been calculated using upper limits on OH concentrations of 1.8×10^6 551
- molecule cm $^{-3}$ and 6.4×10^5 molecule cm $^{-3}$ for the summer and winter flights, respectively, 552
- based on the FAGE instrument's limit of detection. The high upper limits make the total rate 553

of reaction of OH with alkenes, Φ_{OH} , unrealistically high for both summer $(1.6 \times 10^5 \text{ molecule cm}^{-3} \text{ s}^{-1})$ and winter $(7.8 \times 10^4 \text{ molecule cm}^{-3} \text{ s}^{-1})$. However, the OH reactivity will likely be considerably lower than the values calculated using the OH upper limits. A box model constrained to concentrations of long-lived species measured during the flights (Stone et al., 2014b) predicts a mean OH concentration of 2.4×10^4 molecule cm⁻³, significantly lower than the upper limits given by the instrument's limit of detection. Using the mean modelled value for OH gives $\Phi_{OH} = 2.1 \times 10^3$ molecule cm⁻³ s⁻¹ for summer, and $\Phi_{OH} = 2.9 \times 10^3$ molecule cm⁻³ s⁻¹ for winter, indicating a diminished role for OH in alkene oxidation at night in agreement with previous studies (e.g. Geyer et al., 2003; Emmerson et al., 2005).

6.2 Daytime oxidation of alkenes

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Figure 13 shows histograms of rates of reaction of O₃ and OH with alkenes during SeptEx, and O₃ and NO₃ with alkenes during winter RONOCO flights, for daytime data only. OH was detected above the limit of detection $(1.2 \times 10^6 \text{ molecule cm}^{-3})$ during the SeptEx flights, so the FAGE OH data were included in the calculations, using a reaction scheme analogous to the one shown in Fig. 10. NO₃ was not detected during the day in SeptEx. NO₃ is not expected to be present at measurable concentrations during daylight hours due to photolysis, but a mean concentration of 8.3×10^7 molecule cm⁻³ (3.3 pptv) was measured during the day in the winter RONOCO flights. These measurements of low mixing ratios of NO₃ may be partly caused by interference from other daytime species as observed by Brown et al. (2005), or by the variability of the instrument baseline, which can be on the order of 1–2 pptv during vertical profiles on the aircraft (Kennedy et al., 2011). This variability is small compared to the range of NO₃ values typically observed during RONOCO flights (0–50 pptv during summer; 0–10 pptv during winter). During SeptEx, Φ_{OH} exceeded Φ_{O3} by a factor of 8. Ethene and propene were the two most abundant alkenes measured during SeptEx and contributed significantly to OH reactivity. O₃ reactivity with alkenes was dominated by propene and trans-2-butene (6th most abundant alkene measured during SeptEx). NO₃ reactivity with alkenes was dominated by trans-2-butene and isobutene (3rd most abundant alkene measured during winter daytime flights). The total rate of reaction of O₃ and OH with alkenes during daytime SeptEx flights $(3.7 \times 10^5 \text{ molecule cm}^{-3} \text{ s}^{-1})$ exceeded the total rate of reaction of O_3 and NO_3 during daytime winter RONOCO flights (6.6 \times 10⁴ molecule cm⁻³ s⁻¹) by a factor of 6, and was more than double the total rate of reaction of O₃ and NO₃ with

- alkenes during nighttime summer flights $(1.4 \times 10^5 \text{ molecule cm}^{-3} \text{ s}^{-1})$. In winter daytime
- flights, Φ_{O3} was greater than Φ_{NO3} by a factor of 2.4.
- 588 Figure 12b and Fig. 13b reveal that reactions of O₃ dominated alkene reactivity during both
- 589 daytime and nighttime winter RONOCO flights. The concentrations of alkenes were
- 590 generally higher at night, with the total alkene concentration (sum of concentrations of
- alkenes measured) being 2.1×10^9 molecule cm⁻³ in the day, and 3.4×10^9 molecule cm⁻³ at
- 592 night. The total measured alkene reactivity ($\Phi_{O3} + \Phi_{NO3}$) was marginally higher during the
- day, by a factor of 1.04. This difference is attributable mainly to the change in Φ_{O3} .
- 594 Comparison of Fig. 12a and Fig. 13b reveals that the total measured alkene reactivity
- 595 $(\Phi_{O3} + \Phi_{NO3})$ was higher during the summer nighttime flights $(1.4 \times 10^5 \text{ molecule cm}^{-3} \text{ s}^{-1})$
- 596 than during the winter daytime flights $(6.6 \times 10^4 \text{ molecule cm}^{-3} \text{ s}^{-1})$, indicating a low
- 597 oxidising environment during winter daytime. The additional contribution to measured alkene
- reactivity from reactions with OH has been calculated using the OH upper limits as described
- in Sect. 6.1. Even with this additional, upper limit OH reactivity $(1.6 \times 10^5 \text{ molecule cm}^{-3} \text{ s}^{-1})$
- and 1.1×10^5 molecule cm⁻³ s⁻¹ for summer nighttime and winter daytime, respectively) the
- 601 total summer nighttime alkene reactivity remains higher than that during winter daytime,
- 602 confirming the importance of the summer nocturnal troposphere for the oxidation of the
- measured alkenes.

6.3 Nighttime production of HO₂ from reactions of O₃ and NO₃ with alkenes

- Table 6 gives total rates (ΣP_{HO2}) of instantaneous production of HO₂ from the reactions of O₃
- and NO₃ with alkenes. NO₃ was not detected during the dawn summer RONOCO flights and
- there were no daytime RONOCO flights during summer. NO₃ dominated HO₂ production
- during dusk and night (68 %), in agreement with Geyer et al. (2003) who found that NO₃ was
- responsible for 53 % of HO₂ production at night in the BERLIOZ campaign. During winter,
- O₃ dominated HO₂ production at all times, with a nighttime contribution of 70 %. This is in
- agreement with the results from the winter PMTACS-NY 2004 field campaign (Ren et al.,
- 612 2006).

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- The total rate of instantaneous production of HO₂ at night was 3.3 times greater in summer
- 614 than in winter, with production from O₃ decreasing by a factor of 1.5, and production from
- NO₃ decreasing by a factor of 7.8, between summer and winter. The mean temperature
- difference between summer and winter of 9 K is thought to be responsible for the lower NO₃
- concentrations in winter $(2.0 \times 10^8 \text{ molecule cm}^{-3}, 8.2 \text{ pptv}, \text{ compared to } 5.8 \times 10^8 \text{ molecule})$

cm⁻³, 24.5 pptv in summer), owing to the increased thermal stability of N₂O₅, and for the 618 619 reduced rate of temperature-dependent reactions between NO₃ and alkenes, and subsequent 620 reactions. There was very little difference between summer and winter mean O₃ mixing concentrations $(9.6 \times 10^{11} \text{ molecule cm}^{-3}, 39.6 \text{ ppbv}, \text{ and } 9.4 \times 10^{11} \text{ molecule cm}^{-3}, 38.6$ 621 622 ppbv, respectively). 623 Production of HO₂ via reactions of NO₃ and O₃ with alkenes is now examined in more detail. 624 The rate of production from individual alkenes was calculated, and plotted in a histogram, as 625 shown in Fig. 14 for the summer and winter nighttime data. During both summer and winter, reactions of O₃ and NO₃ with trans-2-butene were important sources of HO₂, contributing on 626 627 average 62 % to O₃-initiated HO₂ production and 36 % to NO₃-initiated production during the 628 summer and winter flights. Reactions of NO₃ with isoprene were important during summer, 629 contributing 28 % to NO₃-initiated production. The importance of trans-2-butene, despite its 630 relatively low abundance during summer and winter nighttime RONOCO flights (1.8 pptv 631 and 1.7 pptv, respectively, compared to ethene mixing ratios of 55.0 and 104.5 pptv), is 632 attributed to its fast rates of reaction with both O₃ and NO₃ compared to the other alkenes 633 measured. The importance of the isoprene + NO₃ reactions during the summer RONOCO 634 flights is similarly attributed to its fast rate of reaction with NO₃ compared to the other 635 alkenes measured. In addition there is no aldehyde-forming channel from the isoprene-636 derived RO radical (k_7 in Fig. 10), so that the yield of HO₂ from RO is equal to 1. The 637 reaction of isobutene with NO₃ can proceed via one of two channels to produce two different 638 RO₂ radicals but only one channel, with a branching ratio of 0.2, produces HO₂. Isobutene is 639 therefore not a dominant contributor to HO₂ production, despite being the single largest 640 contributor to NO₃ reactivity during daytime and nighttime RONOCO flights (Fig. 12 and 641 Fig. 13). Figure 14 highlights the small change in total production from O₃ between summer 642 and winter, and the dramatic change in total production from NO₃ between summer and 643 winter. 644 Reactions of formaldehyde with NO₃ were included in the analysis where formaldehyde data were available (mean HCHO = 955 pptv). The NO₃ + HCHO reaction contributed a further 645 5.5×10^3 molecule cm⁻³ s⁻¹ (15 %) to HO₂ production from NO₃ reactions, so that production 646 from NO₃ contributed 79 % of the total production. 647

6.4 Production of HO₂ during flight B537

Flight B537, on 20th July 2010, has been identified as an interesting flight, with high 649 concentrations of HO_2 * (3.2 × 10^8 molecule cm⁻³; 13.6 pptv), ozone (peaking at 650 1.8×10^{12} molecule cm⁻³, 89.9 ppbv) and NO₃ (peaking at 4.1×10^9 molecule cm⁻³; 651 176.9 pptv), and a strong positive correlation between HO_2^* and NO_3 (r = 0.97, see Fig. 9). 652 NO, NO₂, and aerosol surface area were also elevated during the flight during flight B537 653 compared to their mean summer values. The highest concentration of ethene $(1.43 \times 10^{10}$ 654 molecule cm⁻³; 0.61 ppbv) during the summer RONOCO flights was measured during B537. 655 $\Sigma P_{\rm HO2}$ from O₃ + alkene reactions (2.6 × 10⁴ molecule cm⁻³ s⁻¹) was higher in flight B537 656 than in all the other summer flights, contributing 42 % of HO₂ production, with NO₃ + alkene 657 reactions contributing 3.6×10^4 molecule cm⁻³ (58 %). The total rate of HO₂ production from 658 O_3 and NO_3 reactions during flight B537 was 6.2×10^4 molecule cm⁻³ s⁻¹. While this is 659 higher than the average value of $\Sigma P_{\rm HO2}$ for the summer flights (5.4 × 10⁴ molecule cm⁻³ s⁻¹) it 660 is not the highest rate of production during the summer flights. During B534 unusually high 661 concentrations of isoprene, cis-2-butene, and 1,3-butadiene contributed to a total rate of HO₂ 662 production of 7.9×10^4 molecule cm⁻³, which is the highest calculated value. 663 Figure 15 shows that the reactions of O₃ and NO₃ with trans-2-butene are once again 664 665 important, contributing 74 % of $\Sigma P_{\text{HO2.O3}}$, and 45 % of $\Sigma P_{\text{HO2.NO3}}$. The correlation between HO₂* and NO₃ is attributed to production of HO₂ by reactions of NO₃ with alkenes, 666 especially trans-2-butene. Figure 16 shows HO₂* versus the total instantaneous rate of 667 production from the reactions of O₃ and NO₃ with alkenes during flight B537, at each 60-668 second data point during the flight for which the requisite data were available. Note that the 669 670 rates plotted in Fig. 16 are higher than those shown in Fig. 15, where the rates of production of HO₂ from each alkene have been averaged across the whole flight. A strong positive 671 correlation exists between HO₂* and both $\Sigma P_{\text{HO2,O3}}$ (r = 0.6) and $\Sigma P_{\text{HO2,NO3}}$ (r = 0.8), 672

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7 Comparison with model results

The observations of OH, HO₂*, NO₃ and N₂O₅ have been interpreted in the context of nighttime oxidation chemistry using a box model constrained to observations of VOCs, NO_x, O₃, CO and other long-lived species measured during the RONOCO flights (Stone et al., 2014b). The Dynamically Simple Model of Atmospheric Chemical Complexity (DSMACC)

indicating the importance of these reactions for production of HO₂ during this flight.

680 (Emmerson and Evans, 2009; Stone et al., 2010; Stone et al., 2014b) was initiated with concentrations of measured species, using a chemistry scheme based on the Master Chemical 681 682 Mechanism (MCM, version 3.2: Jenkin et al., 1997; Jenkin et al., 2003; Saunders et al., 2003; 683 Bloss et al., 2005, via http://mcm.leeds.ac.uk/MCM) and was allowed to run to diurnal steady 684 state. The model output includes concentrations of OH, HO₂, NO₃, RO₂, and other species. 685 Data from daytime flights, or during dawn or dusk periods, were not included in the model 686 analysis. Data from flight B537 were also excluded, owing to the atypical observations of HO₂*, NO₃, O₃ and other chemical species made during this flight. The modelling study and 687 688 results are described in more detail by Stone et al. (2014b). The model predicts a mean OH concentration of 2.4×10^4 molecule cm $^{-3}$ for the summer 689 flights, which is consistent with the measured OH concentrations for which the instrument's 690 691 limit of detection is an upper limit only. The base model underpredicts HO₂* by around 692 200 %, and overpredicts NO₃ and N₂O₅ by 80 and 50 %, respectively. These discrepancies 693 were investigated by determining the processes controlling radical production and loss in the 694 model, and using those results to improve model performance. Model production of HO₂ is dominated by reactions of RO + O₂ (42 %), with a significant contribution from OH + CO 695 696 (31 %) despite low OH concentrations at night. RO_x (= RO + RO₂ + OH + HO₂) radical initiation in the model is dominated by reactions of NO₃ with unsaturated VOCs (80 %), with 697 698 a much smaller contribution (18 %) from alkene ozonolysis. Modelled HO₂ loss is dominated by its reactions with NO₃ (45 %) and O₃ (27 %), both of which are radical propagating routes, 699 700 and which are the dominant routes to OH production in the model. In fact NO₃ was found to 701 control both radical initiation and propagation in the model. 702 These results are in general agreement with the results of the analysis presented in Sect. 6.1, 703 though the model predicts a more important role for NO₃ (80 % of RO_x radical production, 704 which is 7.2 times the contribution from O_3 + alkenes) than is predicted by the analysis based 705 on the observations alone (69 % of HO₂ radical production during summer, which is 2.1 706 times the contribution from O_3 + alkenes). The model predicts a relatively small role for O_3 in 707 both summer and winter. The model is constrained to measured values of O₃, but overpredicts 708 NO₃. The mean measured NO₃ nighttime mixing ratio was 24.5 pptv in the summer and 709 8.2 pptv in the winter. The mean modelled summer and winter values are 37.4 pptv and 710 20.7 pptv, respectively. This discrepancy between modelled and measured NO₃ helps to 711 explain the model overprediction of the role of NO₃ in HO_x radical initiation during the 712 RONOCO flights. Modelled NO₃ reactivity was dominated by iso-butene (36 %) and trans2-butene (27 %), and modelled O₃ reactivity was dominated by *trans*-2-butene (51 %), in agreement with the nighttime alkene reactivities presented in Sect. 6.1.

Improvement to the model predictions of NO_3 , N_2O_5 and HO_2* was made by increasing the concentration of unsaturated VOCs in the model. Increasing the total observed alkene concentration by 4 times resulted in a modelled to observed ratio of 1.0 for HO_2* and of ~1.2 for NO_3 and N_2O_5 . Two-dimensional gas chromatography (GC × GC) analysis of the whole air samples taken during RONOCO has revealed a large number of VOCs extra to those routinely measured (Lidster et al., 2014). Calibration standards for the majority of these species are not yet available, and so quantification of their concentrations is not possible, but their detection confirms that the model overprediction of NO_3 and underprediction of HO_2* are attributable to reactions of NO_3 with unquantified unsaturated hydrocarbons.

The presence of unquantified unsaturated VOCs during the RONOCO campaign, suggested by the model and confirmed by the two-dimensional GC analysis, has implications for the conclusions drawn from the analysis based on the observations. The relative contributions of NO₃ and O₃ to nighttime radical initiation will change with the composition of unsaturated VOCs in the sampled air, due to the different rates of reaction of NO₃ and O₃ with different VOC species, and the rates of production of HO₂ following these reactions. The model results indicate that reaction of NO₃ with the unquantified VOCs leads to increased production of HO₂. The role of NO₃ in nighttime radical production would therefore be enhanced by the inclusion of the unquantified VOCs in the observational analysis.

8 Conclusions and future work

Nighttime radical chemistry has been studied as part of the RONOCO and SeptEx campaigns onboard the BAe-146 research aircraft during summer 2010 and winter 2011. NO₃, N₂O₅, OH and HO₂* were measured simultaneously for the first time from an aircraft, with OH and HO₂* being measured by the University of Leeds aircraft FAGE instrument. OH was detected above the limit of detection during the daytime SeptEx flights only, with a mean concentration of 1.8×10^6 molecule cm⁻³. Upper limits of 1.8×10^6 molecule cm⁻³ and 6.4×10^5 molecule cm⁻³ are placed on mean OH concentrations for the summer and winter RONOCO (night, dawn, and dusk) measurement campaigns, respectively. HO₂* was detected above the limit of detection during the summer and winter RONOCO flights and during SeptEx, with a maximum mixing ratio of 13.6 pptv measured during nighttime flight B537 on

745 20th July 2010. Mean nighttime HO₂* mixing ratios were significantly higher in summer than 746 in winter. Significant concentrations (up to 176.9 pptv) of NO₃ were measured during 747 nighttime flights, since the air masses sampled were sufficiently removed from the surface 748 that the loss of NO₃ by reaction with NO was minimised. The RONOCO flights were 749 therefore an excellent opportunity to study the role of NO₃ in nocturnal oxidation and radical 750 initiation.

The rates of reaction of O₃ and NO₃ with the alkenes measured have been calculated. At night during summer, NO₃ dominated alkene reactivity. Several previous nighttime studies have also found NO₃ to be the dominant nocturnal oxidant (e.g. Geyer et al., 2003; Brown et al., 2011). During nighttime winter RONOCO flights the total rate of reaction of NO₃ with alkenes was much reduced, but the rate of reaction of O₃ with alkenes was similar to that in summer. During day and night in winter, O₃ + alkene reactions were faster than NO₃ + alkene reactions. Overall, during RONOCO, the combined rate of alkene oxidation by O₃ and NO₃ was highest at night during summer.

Calculation of rates of instantaneous production of HO₂ from reactions of O₃ and NO₃ with alkenes, using measurements made during the flights, has revealed that nighttime production was dominated by NO₃ in summer and by O₃ in winter. The rate of instantaneous production of HO₂ from reactions of NO₃ with alkenes decreased significantly from summer to winter (87 %), whereas production from O₃ + alkene reactions was similar in summer and winter, decreasing by just 31 %. Strong positive correlation between HO₂* and NO₃, especially during flight B537, is attributed to the production of HO₂ from reactions of NO₃ with alkenes, particularly *trans*-2-butene and other isomers of butene.

Significant concentrations of HO₂* were detected at night, with the highest HO₂* concentration (13.6 pptv) being measured during a summer nighttime flight, indicating that HO_x radical chemistry remains active at night under the right conditions. The role of HO_x is diminished in the low photolysis winter daytime atmosphere, with alkene ozonolysis being primarily responsible for oxidation and radical initiation, in agreement with previous studies (e.g. Heard et al., 2004; Emmerson et al., 2005). Both the analysis presented here and the results of the box modelling study by Stone et al. (2014b) indicate that in air masses removed from sources of NO, NO₃ plays an important role in the oxidation of alkenes and radical initiation at night, in agreement with previous studies (e.g. Brown et al., 2011). Alkene ozonolysis also plays a significant role in nocturnal oxidation in agreement with Salisbury et al. (2001), Geyer et al. (2003), Ren et al. (2003a), Emmerson et al. (2005), Ren et al. (2006),

- and Volkamer et al. (2010). The balance between the roles of NO₃ and O₃ was controlled in
- part by [NO₃], with colder winter temperatures forcing the NO₃–N₂O₅ equilibrium towards
- 780 N_2O_5 .
- 781 The total rate of reaction of O_3 and NO_3 with alkenes during nighttime summer flights (1.4 \times
- 782 10^5 molecule cm⁻³ s⁻¹) was higher than during daytime winter flights $(6.6 \times 10^4 \text{ molecule})$
- $cm^{-3} s^{-1}$) by a factor of 2.1. Whilst it should be remembered that measurements at different
- times of day and in different seasons reflect composition changes in air masses (such as the
- abundance of reactive alkenes) this result supports the hypothesis that oxidation of certain
- VOCs, in particular the reactive alkenes, in the nocturnal summer atmosphere can be as rapid
- as in the winter daytime atmosphere.
- A box model of nighttime chemistry constrained to measurements of long lived species has
- been used to investigate the nighttime chemistry sampled during RONOCO (Stone et al.,
- 790 2014b). The base model underpredicts HO₂* and overpredicts NO₃. These discrepancies were
- 791 minimised by increasing the concentration of alkenes in the model, thereby increasing
- 792 reaction of NO₃ with alkenes, and production of HO₂. The presence of unquantified
- vinsaturated VOCs has been confirmed by 2D-GC analysis, though the exact nature and
- 794 concentrations of the 'missing' species is unclear. The inclusion of these species in the
- analysis presented in this paper would likely increase the role of NO₃ for oxidation of alkenes
- and production of HO₂ at night.

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- 1 Table 1. Examples of modelling studies and observations of HO_x radicals and VOC oxidation at night. PERCA = Peroxy Radical Chemical
- 2 Amplification; LIF = Laser Induced Fluorescence; DOAS = Differential Optical Absorption Spectroscopy; MCM = Master Chemical Mechanism;
- 3 MIESR = Matrix Isolation Electron Spin Resonance; RACM = Regional Atmospheric Chemistry Mechanism; CRDS = Cavity Ring Down
- 4 Spectroscopy; CIMS = Chemical Ionisation Mass Spectrometry; GC = Gas Chromatography; PTRMS = Proton Transfer Reaction Mass Spectrometry;
- 5 FTIR = Fourier Transform Infrared Spectroscopy; DUALER (DUAl channel peroxy radical chemical amplifier); OA-CRD = Off Axis Cavity Ring
- 6 Down Spectroscopy; CRM-PTR-MS = Comparative Reactivity Method Proton Transfer Mass Spectrometry.

Location, Campaign, Date	Methods	Results	Reference
Mace Head, Ireland, EASE97, 1997	Measurements: [HO ₂ +RO ₂] measured by PERCA; HO _x measured by LIF; NO ₃ measured by DOAS). Modelling: Campaign-tailored box model constrained to measurements, based on MCM.	2 nights of HO_x measurements: $HO_2 = 1-2$ and 0.5–0.7 pptv; OH not detected above limit of detection ($\sim 2.5 \times 10^5$ cm ⁻³). NO_3 dominated radical production in westerly (clean) air masses; O_3 dominated in NE, SE, and SW air masses and dominated radical production overall during the campaign.	Salisbury et al. (2001); Creasey et al. (2002)
Pabstthum, Germany, BERLIOZ 1998,	Measurements: HO _x measured by LIF; NO ₃ measured by DOAS and MIESR. Modelling: Zero-dimensional model using lumped VOC reactivity, constrained to measured species.	Nighttime OH = 1.85×10^5 cm ⁻³ , compared to modelled value of 4.1×10^5 cm ⁻³ . Nighttime HO ₂ = 3×10^7 cm ⁻³ , model results in agreement. NO ₃ chemistry responsible for 53 % of HO ₂ and 36 % of OH during the night. O ₃ + alkene responsible for 47 % of HO ₂ and 64 % of OH during the night.	Geyer et al. (2003); Holland et al. (2003)
Birmingham, PUMA, 1999 and 2000	Measurements: HO _x measured by LIF. Modelling: Photochemical box model constrained to measurements, based on MCM.	Daytime OH initiation dominated by O_3 + alkenes, HONO photolysis, and $O(^1D)$ + H_2O during summer. O_3 + alkenes dominated in winter. O_3 + alkenes main radical source at night.	Emmerson et al. (2005); Harrison et al. (2006)
New York, PMTACS- NY, 2001	Measurements: HO _x measured by LIF.	Nighttime OH ~ 7×10^5 cm ⁻³ and nighttime HO ₂ ~ 8×10^6 cm ⁻³ . Increase in HO _x after midnight attributed to increase in O ₃ due to transport. O ₃ + alkenes main source of nighttime HO _x .	Ren et al. (2003a); Ren et al. (2003b)

Mace Head, NAMBLEX, 2002	Measurements: HO _x measured by LIF; NO ₃ measured by DOAS. Modelling: Zero-dimensional box models constrained to measured species, based on MCM.	Nighttime $HO_2 = 2-3 \times 10^7 \text{ cm}^{-3}$; OH below detection limit $(6 \times 10^4 \text{ cm}^{-3})$. Model overestimated HO_2 . On average, O_3 + alkene reactions contributed 59 % and NO_3 + alkene reactions contributed 41 % to RO_2 production at night, but NO_3 and RO_2 concentrations were always higher in semi-polluted air masses than in clean marine air masses and NO_3 reactions dominated in these conditions.	Fleming et al. (2006); Smith et al. (2006); Sommariva et al. (2007)
Writtle, London, TORCH, 2003	Measurements: HO _x measured by LIF, RO ₂ measured by PERCA, during a heatwave/pollution episode. Modelling: zero-dimensional box model constrained to measured species.	OH and HO_2 observed above the limit of detection on several nights. OH peaked at 8.5×10^5 cm ⁻³ ; HO_2 peaked at 1×10^8 cm ⁻³ . Model overpredicted nighttime OH and HO_2 on average by 24 % and 7 %; underpredicted [$HO_2+\Sigma RO_2$] by 22 %.	Lee et al. (2006); Emmerson et al. (2007); Emmerson and Carslaw (2009)
Mexico City, MCMA 2003	Measurements: HO _x measured by LIF, NO ₃ measured by DOAS. Modelling: Zero-dimensional model based on MCM v3.1, constrained to measured species.	Polluted city location characterized by high levels of NO, NO ₂ and O ₃ . Maximum nighttime OH ~ 1×10^6 cm ⁻³ ; maximum nighttime HO ₂ ~ 6 pptv. Nighttime production of radicals dominated by O ₃ + alkene reactions (76–92 %); NO ₃ + alkene plays a minor role. Daytime radical production ~ 25 times higher than night.	Shirley et al. (2006); Sheehy et al. (2010); Volkamer et al. (2010)
New York City, PMTACS-NY winter 2004	Measurements: HO _x measured by LIF. Modelling: Zero-dimensional model based on RACM and constrained by measurements.	Mean maximum $OH = 0.05$ pptv; mean maximum $HO_2 = 0.7$ pptv. Model under-prediction of HO_2 was pronounced when NO was high. O_3 + alkene reactions were dominant nighttime source.	Ren et al. (2006)
Gulf of Maine, Northeast United States, NEAQS, 2004	Measurements: NO_3 and N_2O_5 measured by CRDS. Modelling: Zero-dimensional model based on MCM v3.1, constrained to measured species. No measurements of OH, HO_2 , or RO_2 .	Ship-based measurements onboard RV Ronald H. Brown in the Gulf of Maine, influenced by unpolluted marine air masses and polluted air masses from USA and Canada. Maximum modelled nighttime $HO_2 = 7.0 \times 10^8$ cm ⁻³ . Base model overestimated NO_3 and NO_2 observations by 30–50 %. In anthropogenic air masses reaction with VOCs and RO_2 each accounted for 40 % of modelled NO_3 loss.	Sommariva et al. (2009)
Houston, Texas, TexAQS, 2006	Measurements: NO ₃ and N ₂ O ₅ measured by CRDS, VOCs measured by CIMS, GC, and PTRMS. No direct measurements of OH, HO ₂ ,	Loss rates and budgets of NO ₃ and highly reactive VOCs calculated. NO ₃ primarily lost through reaction with VOCs. VOC oxidation dominated by NO ₃ , which was 3–5 times more important than O ₃ .	Brown et al. (2011)

or RO_2 .

Pearl River Delta, China, PRIDE-PRD, 2006	Measurements: HO _x measured by LIF; OH reactivity measured by laser-flash photolysis and LIF; VOCs measured by FTIR and GC. Modelling: Box model based on RACM and the Mainz Isoprene Mechanism, and constrained by measurements.	Rural site 60 km downwind of large urban region (Guangzhou), with low local wind speeds favouring accumulation of air pollutants. Maximum nighttime OH (hourly average) = 5×10^6 cm $^{-3}$; maximum nighttime HO ₂ (hourly average) = 1×10^9 cm $^{-3}$. Unknown recycling mechanism required for the model to reproduce measured nighttime values. OH reactivity peaked at night. Missing nighttime reactivity attributed to unmeasured secondary organic compounds.	Lou et al. (2010); Lu et al. (2012); Lu et al. (2013)
Beijing, CAREBEIJIN G2006, 2006	Measurements: HO _x measured by LIF; OH lifetime measured by laser flash photolysis and LIF; VOCs measured by GC. Modelling: Box model based on RACM and the Mainz Isoprene Mechanism, and constrained by measurements.	Suburban rural site south of Beijing, under the influence of slowly moving, aged polluted air from the south. OH reactivity peaked at night. Model generally underestimated observed nighttime OH concentrations.	Lu et al. (2013); Lu et al. (2014)
Cape Verde, RHaMBLe, 2007	Measurements: HO _x measured by LIF. Modelling: Box model based on MCM with added halogen chemistry scheme, constrained to measurements of long-lived species.	Clean tropical Atlantic measurement site with occasional continental influence. OH was not measured at night. HO_2 was detected on two nights, up to 2.5×10^7 cm ⁻³ . Model underprediction of HO_2 was significantly reduced by constraining the model to 100 pptv of peroxy acetyl nitrate (PAN) at night.	Whalley et al. (2010)
Huelva, Spain, DOMINO, 2008	Measurements: $[HO_2+RO_2]$ measured by DUALER; HO_x measured by LIF; NO_3 and N_2O_5 measured by OA-CRD; OH reactivity measured by CRM-PTR-MS. No measurements of anthropogenic VOCs.	Coastal forested site with strong urban-industrial and weak biogenic influences. Maxima in $[HO_2+RO_2]$ and $[HO_2]$ were observed around noon and midnight. Enhanced nighttime $[HO_2+RO_2]$ (up to 80 pptv) was observed in air masses from the urban-industrial region. Maximum nighttime $HO_2=8$ pptv. Measured NO_3 was generally below LOD; calculated NO_3 up to 20 pptv. Calculated production of RO_2 from NO_3 +alkenes accounts for 47–54 % of observed $[HO_2+RO_2]$. Ozonolysis of unmeasured alkenes could account for remaining $[HO_2+RO_2]$.	Andrés- Hernández et al. (2009)

Table 2. Details of supporting measurements.

Species	Instrument, Technique	Time resolution; Limit of detection (LOD)	References
СО	Aero Laser AL5002 Fast Carbon Monoxide Monitor. Excitation and fast response fluorescence at $\lambda = 150$ nm.	1 s; 3.5 ppbv	Gerbig et al. (1999)
O_3	Thermo Scientific TEi49C Ozone analyser. Absorption spectroscopy at $\lambda = 254$ nm.	1 s; 0.6 ppbv	Hewitt et al. (2010)
NO, NO_2, NO_x $(NO + NO_2)$	Air Quality Design dual channel fast-response NO_x instrument. Chemiluminescence from $NO + O_3$ reaction. Conversion of NO_2 to NO by photolysis.	10 s; 3 pptv for NO, 15 pptv for NO ₂	Stewart et al. (2008)
NO ₂ , ΣANs, ΣPNs	TD-LIF (thermal dissociation laser induced fluorescence). Detection of NO_2 by laser-induced fluorescence. Thermal decomposition of ΣANs (total alkyl nitrate) and ΣPNs (total peroxy nitrate) to NO_2 .	1 s; 9.8 pptv for NO_2 , 28.1 pptv for ΣANs , 18.4 pptv for ΣPNs	Dari-Salisburgo et al. (2009); Di Carlo et al. (2013)
Alkenes	Whole air samples (WAS) analysed by laboratory-based gas chromatography with flame ionization detection (GC-FID).	Typically 30 s; variable limits of detection	Hopkins et al. (2003)
NO ₃ , N ₂ O ₅	BBCEAS (broadband cavity-enhanced absorption spectroscopy) of NO_3 at $\lambda = 642-672$ nm. N_2O_5 measured following thermal dissociation to $NO_3 + NO_2$.	1 s; 1.1 pptv for NO_3 , 2.4 pptv for $NO_3 + N_2O_5$	Kennedy et al. (2011)
НСНО	Hantzsch technique: Liquid- phase reaction of formaldehyde followed by excitation, and fluorescence of resulting adduct at $\lambda = 510$ nm.	60 s; 81 pptv	Still et al. (2006)

Table 3. Mean mixing ratios of selected gas phase species, and air temperature, measured during RONOCO and SeptEx. The flight and season during which the maximum values were measured are given in parentheses. NO₂ data are from the TD-LIF instrument. Zero values indicate measurements below the limit of detection.

Species	Summer RONOCO	SeptEx	Winter RONOCO	Maximum
CO / ppbv	102.3	117.1	139.3	256.0 (B537, summer)
O ₃ / ppbv	39.6	40.4	38.6	89.8 (B537, summer)
NO ₃ / pptv	21.1	0	6.2	176.9 (B537, summer)
NO / ppbv	0.05	0	0	18.9 (B539, summer)
NO ₂ / ppbv	1.6	1.7	2.3	18.6 (B568, winter)
Temperature / K	286.5	286.2	276.4	297.5 (B537, summer)

Table 4. Combined daytime and nighttime mean concentrations of OH and mean mixing ratios of HO_2^* with the FAGE instrument's average 1σ limits of detection for a 5 minute averaging period during the RONOCO and SeptEx fieldwork.

	OH / m	OH / molecule cm ⁻³		
	Mean concentration	Limit of detection	Mean mixing ratio	Limit of detection
Summer		1.8×10^6	1.6	0.03
SeptEx	1.8×10^6	1.2×10^6	2.9	0.02
Winter		6.4×10^{5}	0.7	0.02

Table 5. Mean and, in parentheses, maximum HO_2^* mixing ratios measured during RONOCO and SeptEx.

	Mean (maximu	Mean (maximum) HO ₂ * mixing ratio / pptv			
	Summer	SeptEx	Winter		
Dawn	0.74 (1.19)		0.54 (1.81)		
Day		3.78 (11.79)	0.49 (1.68)		
Dusk	2.73 (9.97)		0.32 (0.97)		
Night	1.86 (13.58)		0.98 (2.02)		

Table 6. Average rates of instantaneous production of HO₂ from reactions of O₃ and NO₃ with alkenes.

Measurements	HO_2 production rate $(\Sigma P_{HO2}) / 10^4$ molecule cm ⁻³ s ⁻¹			
	Dawn	Day	Dusk	Night
Summer				
NO_3	0		2.8	3.8
O_3	0.5		2.2	1.7
Total	0.5		5.0	5.5
Winter				
NO_3	0.4	0.4	0.4	0.5
O_3	1.4	1.5	1.2	1.2
Total	1.8	1.9	1.6	1.7

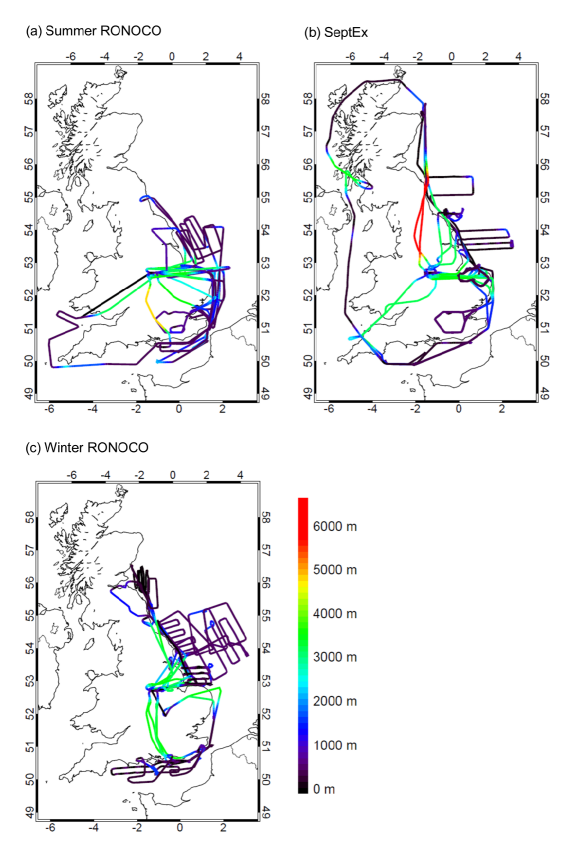


Figure 1. Flight paths for: (a) Summer RONOCO, (b) SeptEx, and (c) winter RONOCO measurement campaigns, coloured by altitude.

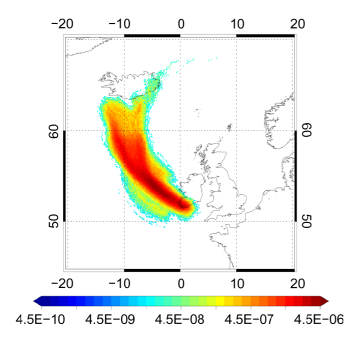


Figure 2. Footprint map for flight B535 on 17th July 2010, showing model particle densities (g s m⁻³) in a 300 m deep layer from the surface, integrated over a 24 hour period beginning 48 hours prior to the flight.

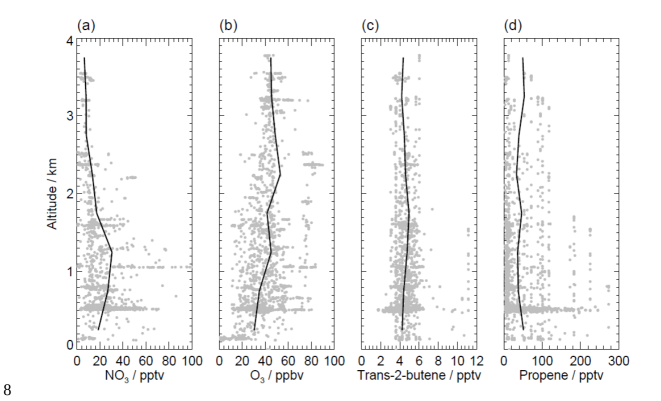


Figure 3. Nighttime altitude profiles of a) NO₃; b) O₃; c) *trans*-2-butene; d) propene, showing 60 second data (grey points) and mean values in 500 m altitude bins (solid black lines).

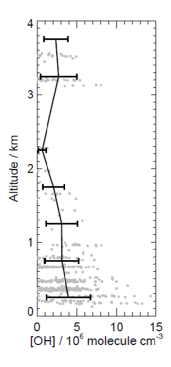


Figure 4. Altitude profile of OH measured during SeptEx showing 60 second data (grey points) and mean values in 500 m altitude bins (solid black lines). Error bars are 1σ.

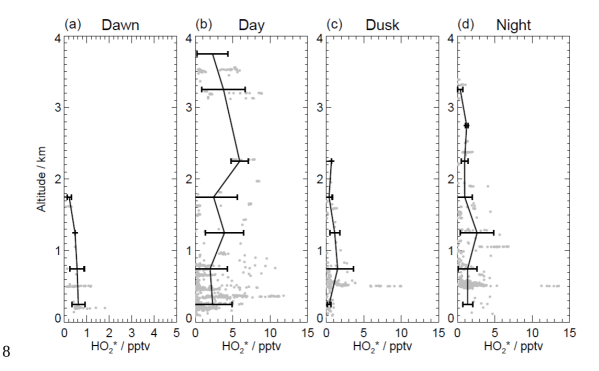
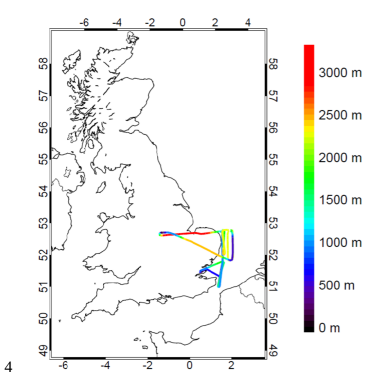


Figure 5. Altitude profiles of HO_2^* measured in RONOCO and SeptEx during: a) dawn; b) day; c) dusk; d) night, showing 60 second data (grey points) and mean values in 500 m altitude bins (solid black lines). Error bars are 1σ .



5 Figure 6. Flight track of flight B537 on 20th July 2010, coloured by altitude.

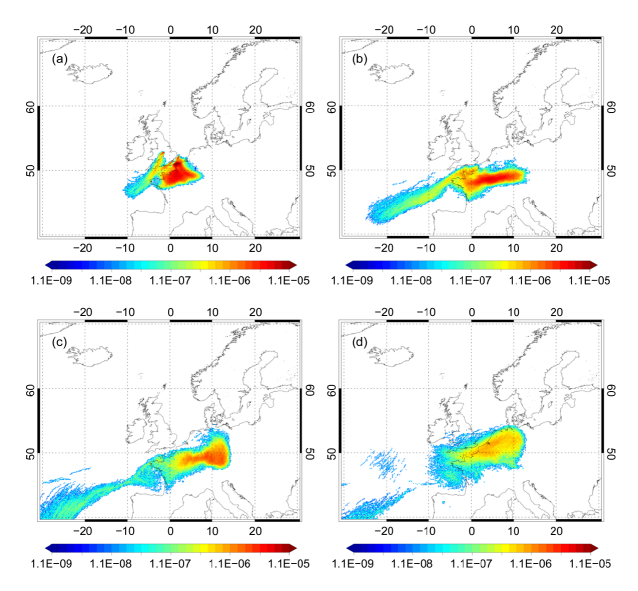


Figure 7. Footprint maps for flight B537 on 20th July 2010, showing model particle densities (g s m⁻³) in a 300 m deep layer from the surface, integrated over 24 hour periods beginning (a) 24 hours, (b) 48 hours, (c) 72 hours, and (d) 96 hours prior to the flight.

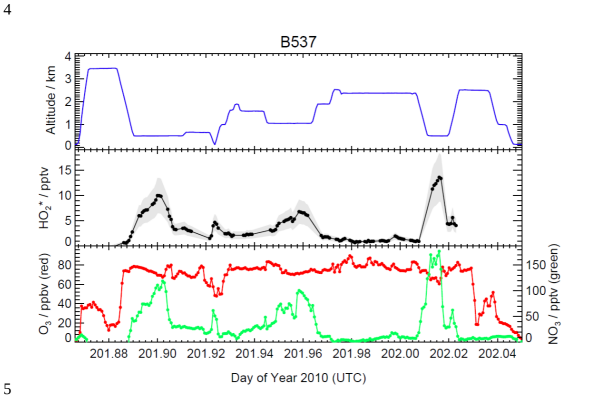


Figure 8. Time series of altitude (top panel, blue), HO_2^* (middle panel, black, with grey shading representing the uncertainty in the measurements), O_3 (bottom panel, red) and NO_3 (bottom panel, green) during nighttime flight B537 on 20^{th} July 2010.

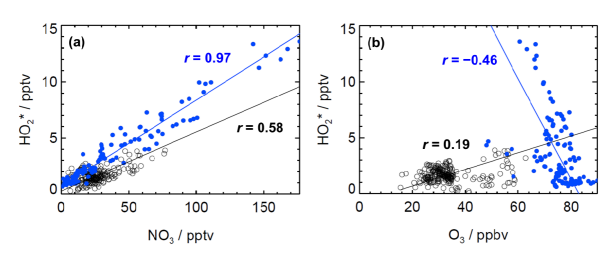
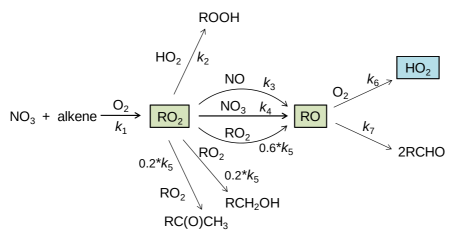
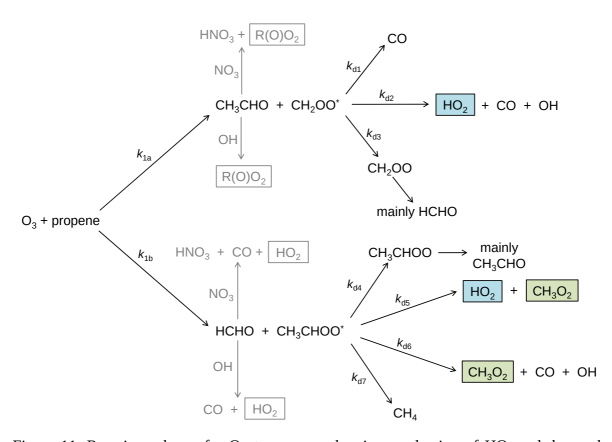


Figure 9. HO₂* versus (a) NO₃ and (b) O₃ during flight B537 (blue, filled circles) and during all other nighttime flights (black, open circles). The solid lines are lines of best fit to the data.



- 2 Figure 10. Generalised reaction scheme for production of RO₂ and HO₂ following reaction of
- NO_3 with an alkene.



- 6 Figure 11. Reaction scheme for O_3 + propene, showing production of HO_2 and the methyl
- 7 peroxy radical, CH₃O₂.

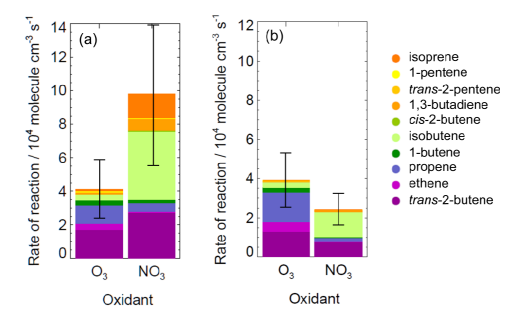


Figure 12. Average nighttime rates of reaction between O₃, and NO₃ with alkenes during: a) summer; and b) winter RONOCO flights. Error bars represent the combined uncertainty in the measurements.

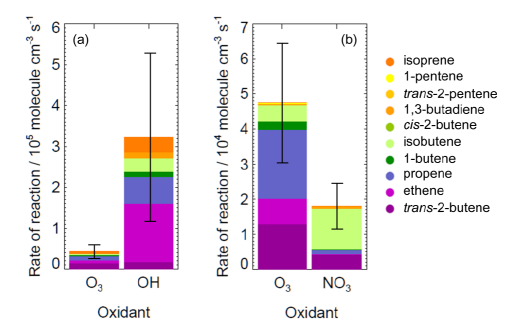


Figure 13. Average daytime rates of reaction of (a) O_3 and OH with alkenes during SeptEx; and (b) O_3 and NO_3 with alkenes during winter. Note the different scales. NO_3 was not detected during daytime SeptEx flights (LOD = 1.1pptv); OH was not detected during daytime winter flights (LOD = 6.4×10^5 molecule cm⁻³). Error bars represent the combined uncertainties in the measurements.

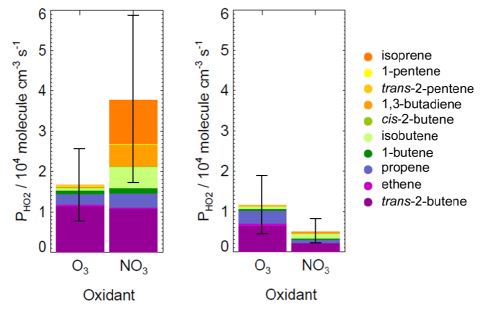


Figure 14. Average rates of instantaneous production of HO_2 from reactions of O_3 and NO_3 with alkenes during: a) summer; and b) winter RONOCO flights. Error bars represent the combined uncertainty in the measurements.

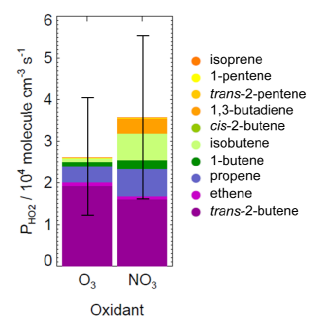


Figure 15. Average rates of instantaneous production of HO_2 from reactions of O_3 and NO_3 with alkenes during flight B537. Error bars represent the combined uncertainty in the measurements.

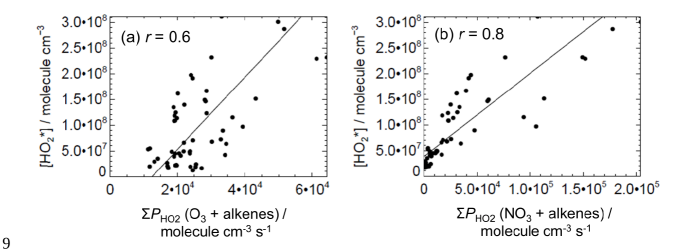


Figure 16. [HO₂*] versus total rate of instantaneous production of HO₂ from reactions of: a) O₃; and b) NO₃ during flight B537. Correlation coefficients (*r*) are given in each plot. Solid lines are lines of best fit to the data.