Implementation of a chemical background method for atmospheric OH measurements by laser-induced fluorescence: characterisation and observations from the UK and China

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Abstract. Hydroxyl (OH) and hydroperoxy (HO2) radicals are central to the understanding of atmospheric chemistry. Owing to their short lifetimes, these species are frequently used to test the accuracy of model predictions and their underlying chemical mechanisms. In forested environments, laser-induced fluorescence–fluorescence assay by gas expansion (LIF–FAGE) measurements of OH have often shown substantial disagreement with model predictions, suggesting the presence of unknown OH sources in such environments. However, it is also possible that the measurements have been affected by instrumental artefacts, due to the presence of interfering species that cannot be discriminated using the traditional method of obtaining background signals via modulation of the laser excitation wavelength (“OHwave”). The interference hypothesis can be tested by using an alternative method to determine the OH background signal, via the addition of a chemical scavenger prior to sampling of ambient air (“OHchem”). In this work, the Leeds FAGE instrument was modified to include such a system to facilitate measurements of OHchem, in which propane was used to selectively remove OH from ambient air using an inlet pre-injector (IPI). The IPI system was characterised in detail, and it was found that the system did not reduce the instrument sensitivity towards OH (<5 % difference to conventional sampling) and was able to efficiently scavenge external OH (>99 %) without the removal of OH formed inside the fluorescence cell (<5 %). Tests of the photolytic interference from ozone in the presence of water vapour revealed a small but potentially significant interference, equivalent to an OH concentration of ∼ 4 × 10^5 molec. cm^{-3} under typical atmospheric conditions of [O_3] = 50 ppbv and [H_2O] = 1 %. Laboratory experiments to investigate potential interferences from products of isoprene ozonolysis did result in interference signals, but these were negligible when extrapolated down to ambient ozone and isoprene levels. The interference from NO3 radicals was also tested but was found to be insignificant in our system. The Leeds IPI module was deployed during three separate field intensives that took place in summer at a coastal site in the UK and both in summer and winter in the megacity of Beijing, China, allowing for investigations of ambient OH interferences under a wide range of chemical and meteorological conditions. Comparisons of ambient OHchem measurements to the traditional OHwave method showed excellent agreement, with OHwave vs OHchem slopes of 1.05–1.16 and identical behaviour on a diel basis, consistent with laboratory interference tests. The difference between OHwave and OHchem (“OHint”) was found to scale non-linearly with OHchem, resulting in an upper limit interference of (5.0 ± 1.4) × 10^6 molec. cm^{-3} at the very highest OHchem concentrations measured (23 × 10^6 molec. cm^{-3}), accounting for ∼ 14 %–21 % of the total OHwave signal.
1 Introduction

The removal of pollutants and greenhouse gases in the troposphere is dominated by reactions with the hydroxyl radical (OH), which is closely coupled to the hydroperoxy radical (HO_2). Comparisons of the levels of OH and HO_2 observed during field campaigns to the results of detailed chemical box models serve as a vital tool to assess our understanding of the underlying chemical mechanisms involved in tropospheric oxidation. Laser-induced fluorescence–fluorescence assay by gas expansion (LIF–FAGE) measurements of OH in forested environments have often been considerably higher than those predicted by models (Carslaw et al., 2001; Lelieveld et al., 2008; Ren et al., 2008; Hofzumahaus et al., 2009; Stone et al., 2011; Whalley et al., 2011; Wolfe et al., 2011). The difficulty in simulating radical concentrations in such environments has prompted a multitude of theoretical (Peeters et al., 2009, 2014; da Silva et al., 2010; Nguyen et al., 2010; Peeters and Muller, 2010), laboratory (Dillon and Crowley, 2008; Hansen et al., 2017), and chamber (Paulot et al., 2009; Crounse et al., 2011, 2012; Wolfe et al., 2012; Fuchs et al., 2013, 2014, 2018; Praske et al., 2015; Teng et al., 2017) studies to help explain the sources of the measurement–model discrepancy through detailed investigations of the mechanism of isoprene oxidation under low NO_x conditions, as well as other biogenic volatile organic compounds (BVOCs) (Novelli et al., 2018). However, another hypothesis is that the LIF measurements have, at least in part, suffered from an instrumental bias in these environments due to interfering species.

Early LIF measurements of OH suffered from significant interferences due to laser-generated OH from ozone photolysis in the presence of water vapour (Hard et al., 1984). While this effect has been reduced by going from 282 to 308 nm laser excitation of OH, it may still be significant, especially at night or with the use of multi-pass laser set-ups (e.g. up to \(\sim 4 \times 10^6\) molec. cm\(^{-3}\) in Griffith et al., 2016). Laboratory experiments conducted by Ren and co-workers using the Pennsylvania State University (PSU) LIF instrument showed negligible interferences in OH detection for a range of candidate species: H_2O_2, HONO, HCHO, HNO_3, acetone, and various RO_2 radicals (Ren et al., 2004). Observations of OH during the PROPHET (Program for Research on Oxidants: Photochemistry, Emissions and Transport) field campaign in summer 1998, located in a mixed deciduous forest in Michigan, USA, revealed unusually high night-time OH concentrations (\(\sim 1 \times 10^6\) molec. cm\(^{-3}\)) but measurement interferences were ruled out (Faloona et al., 2001).

However, the results of more recent studies conducted in forested environments have meant that interferences in the measurement of OH by LIF–FAGE have been revisited. The usual background method of this technique, where the laser wavelength is scanned off-resonance from an OH transition ("OH\(_{\text{wave}}\"), does not discriminate between ambient (i.e. "real") OH and either OH formed inside the FAGE cell (e.g. laser- or surface-generated OH or via unimolecular decomposition in the gas phase to form OH) or fluorescence from other species at \(\lambda \sim 308\) nm (e.g. naphthalene, SO_2), although it is possible to correct for such effects providing the interference has been previously characterised (Martinez et al., 2004; Ren et al., 2004; Griffith et al., 2013; Fuchs et al., 2016). There is an alternative, chemical method ("OH\(_{\text{chem}}\"\)) for obtaining the OH background signal in LIF instruments that allows for interference signals to be determined without their prior characterisation, in order to rule out possible artefacts from unknown species. The OH\(_{\text{chem}}\) method involves the addition of a high concentration of an OH scavenger, such as perfluoropropene (C_3F_6) or propane, just before the FAGE inlet. Ambient OH is quickly titrated away by fast reaction with the scavenger, but any interference should remain in the fluorescence signal, although this must be corrected for reaction of any internally generated OH with the scavenger inside the FAGE cell.

Several LIF–FAGE groups have now made efforts to validate ambient OH measurements through incorporation of the alternative OH\(_{\text{chem}}\) technique, which was first applied for continuous OH measurements by Mao et al. (2012). Since then, field studies of OH measurement interferences have been conducted in forested (Griffith et al., 2013; Novelli et al., 2014a, 2017; Feiner et al., 2016; Lew et al., 2019), rural (Fuchs et al., 2017; Tan et al., 2017), suburban (Tan et al., 2018, 2019), urban (Ren et al., 2013; Brune et al., 2016; Griffith et al., 2016), and coastal (Novelli et al., 2014a; Mallik et al., 2018) locations. Substantial improvement in measurement–model agreement has been possible when OH backgrounds were determined chemically, especially in forested environments, suggesting that understanding of tropospheric oxidation processes in such regions may be better than previously thought (Mao et al., 2012; Hens et al., 2014; Feiner et al., 2016). This is further supported by the positive identification of two new OH interference candidates in laboratory experiments, namely intermediates in alkene ozonolysis reactions, which may (Novelli et al., 2014b, 2017; Rickly and Stevens, 2018) or may not (Fuchs et al., 2016) be related to stabilised Criegee intermediates (SCIs), and the NO_3 radical (Fuchs et al., 2016), although for all cases the observed interferences cannot explain the magnitudes of the OH background signals under ambient conditions. The trioxide species, ROOOH, has also been postulated to explain elevated OH backgrounds in LIF–FAGE measurements made in forested regions (Fittschen et al., 2019).

However, it is not known whether other LIF instruments suffer the same levels of interference, which are likely highly dependent on cell design and operating parameters, especially the residence time of air between sampling and detection (Novelli et al., 2014a; Fuchs et al., 2016; Rickly and Stevens, 2018). Considering the bespoke nature of LIF–FAGE instruments, those of different groups share the same main features but differ in many aspects, such as inlet size and shape, or whether the laser crosses the detection axis...
once (i.e. single-pass) or multiple times (multi-pass). As a result, the magnitude of any interference is likely to vary significantly between different instruments. Because of this, a general recommendation of the 2015 International HOx Workshop (Hofzumahaus and Heard, 2016) was that different groups should incorporate their own chemical scavenger system for use in ambient OH measurements and to test interferences in the laboratory.

Following this recommendation, the Leeds ground-based FAGE instrument was modified to incorporate a chemical scavenger system, through the addition of an inlet pre-injector (IPI). In this work, we describe the design of the IPI system and its thorough characterisation in terms of sensitivity and the degree of external and internal OH removal. Following this, we present the results of interference testing experiments performed using the IPI system, in which we investigated interferences from O₃+H₂O, isoprene ozonolysis, and NO₃ radicals. Finally, we demonstrate the use of the optimised IPI system for measurements of ambient OH made during three separate field campaigns in the UK and China, which encompassed a wide range of chemical and meteorological conditions.

2 Methodology

2.1 Overview of the Leeds ground-based FAGE instrument

The University of Leeds ground-based FAGE instrument, described in detail elsewhere (Creasey et al., 1997a; Whalley et al., 2010, 2013), has participated in 24 intensive field campaigns since its initial deployment in 1996. Measurements of OH, HO₂, and, more recently, RO₂ radicals (Whalley et al., 2013) have been made in a variety of locations, ranging from pristine marine boundary layer (Creasey et al., 2003; Whalley et al., 2010), tropical rainforest (Whalley et al., 2011), and polar (Bloss et al., 2007) environments to coastal (Smith et al., 2006) and semi-polluted regions (Creasey et al., 2001), as well as urban areas (Heard et al., 2004; Emmerson et al., 2007), including a highly polluted megacity (Lee et al., 2016; Whalley et al., 2016).

Ambient OH is measured using laser-induced fluorescence. Briefly, ambient air is drawn through a 1.0 mm diameter pinhole in a conical turret inlet (4 cm length, 3.4 cm ID; Fig. 1) at ∼7 slm into a stainless steel fluorescence cell, held at ∼2 hPa using a Roots blower (Leybold RUVAC WAU 1001) backed by a rotary pump (Leybold SOGEVAC SV200). An all solid-state laser system is used to excite OH via the $\Delta^2 \Sigma^+ (v'' = 0) \rightarrow X^2 \Pi_{3/2} (v'' = 0)$ electronic transition at $\lambda = 308$ nm (HO₂ is measured by conversion to OH using NO; see details below). The resultant fluorescence at 308 nm is detected by a micro-channel plate photomultiplier (MCP, Photek PMT325/Q/BIG with 10 mm diameter photocathode) equipped with a 50 ns gating unit (Photek GM10-50) and a 2 GHz 20 dB gain amplifier (Photek PA200-10), and the signal is analysed by gated photon counting (Whalley et al., 2010). The background signal is normally obtained by scanning the laser wavelength off-resonance from the OH transition line, yielding the measurement commonly referred to as OH\textsubscript{wave}:

$$[\text{OH}_{\text{wave}}] = COH \times (S_{\text{on-line}} - S_{\text{off-line}}) = COH \times S_{\text{OH}},$$

(1)

where COH is the instrument calibration factor for OH and $S_{\text{on-line}}$ and $S_{\text{off-line}}$ are the OH LIF signals at on- and off-resonance wavelengths, respectively. Similarly, the alternative measurement known as OH\textsubscript{chem} is defined as follows:

$$[OH_{\text{chem}}] = COH \times \left( S_{\text{on-line}} - S_{\text{on-line, scavenger}} \right),$$

$$= COH \times S_{\text{OH, scavenger}},$$

(2)

where $S_{\text{on-line, scavenger}}$ is the OH signal measured at an on-resonance wavelength but in the presence of a scavenger.

HO₂ is detected via its conversion to OH following the addition of NO (BOC, 99.95 %, and Messer, 99.95 %). Although not reported here, RO₂ radicals are measured using the RO₂ LIF method (Fuchs et al., 2008; Whalley et al., 2013), in which their reactions with NO and CO (BOC, 5 % in N₂, and Messer, 10 % in N₂) result in conversion initially to OH (using NO; RO₂ → HO₂ → OH) and subsequently back to HO₂ (using CO; OH → HO₂) that is then detected as described above (via addition of NO inside the FAGE cell; HO₂ → OH). The NO and CO are delivered using mass flow controllers (MFC, MKS Instruments 1179A series), which, unless otherwise stated, were also used to control all other gas flows described in this work. The Leeds FAGE instrument features two fluorescence cells, where laser light (≈10–20 mW at 308 nm, supplied at a pulse repetition frequency of 5 kHz via an optical fibre) enters each cell in
series. For normal operation in the field, the first cell (HO$_x$) measures OH and HO$_2$ (low NO flow, 5 sccm; RO$_2$ interference minimised) sequentially, while the second cell (RO$_x$) measures HO$_2^*$ (high NO flow, 50 sccm; RO$_2$ interference maximised) and then total RO$_2$ (Fuchs et al., 2008; Whalley et al., 2013).

Calibration of the FAGE instrument is achieved by supplying known radical concentrations via a turbulent flow tube (known in Leeds as the “wand”) held at ∼45° to the instrument inlet, where OH and HO$_2$ are formed in a 1 : 1 ratio (Fuchs et al., 2011) by the photolysis of water vapour at 184.9 nm using a Hg(Ar) pen-ray lamp (LOT LSP035) in an excess flow (40 slm) of humidified ultra-high-purity air (BOC, BTCA 178, and Messer, 20.5 % O$_2$ in N$_2$). Chemical actinometry is performed via the photolysis of N$_2$O (BOC, medical-grade 99.5 %) to measure the product of lamp flux and photolysis exposure time to enable calculation of radical concentrations (Edwards et al., 2003; Faloona et al., 2004). The calibration of OH using the water vapour photolysis method has been validated by comparison with other methods, for example the kinetic decay of hydrocarbons (Winberg et al., 2015).

### 2.2 Inlet pre-injector (IPI) design

The Leeds inlet pre-injector (Fig. 1) is similar in concept to the design of Mao et al. (2012) and consists of a 4.0 cm length, 1.9 cm ID perfluoralkoxy (PFA) cylinder embedded inside an aluminium housing, which seals to the FAGE cell via an O-ring base. The scavenger is injected into the centre of the PFA flow tube via four 0.25 mm ID needles, 4.0 cm above the FAGE inlet. The low bore capillary tubing increases the pressure inside the needles, which facilitates mixing of the scavenger into the ambient air stream. In this work propane (BOC, research-grade 99.95 %, and Messer, 99.995 %) was used as the main OH chemical scavenger, with similar results (see Sect. 3.1.2) obtained for C$_3$F$_6$ (Sigma-Aldrich, 99 %). Based on previous investigations of OH interferences (Stevens et al., 1994; Dubey et al., 1996; Faloona et al., 2001; Ren et al., 2004; Mao et al., 2012; Griffith et al., 2013; Rickly and Stevens, 2018), C$_3$F$_6$ was used initially as it reacts quickly and selectively with OH ($k_{298} = 2.2 \times 10^{-12} \text{cm}^3\text{molec.}^{-1}\text{s}^{-1}$ Sander et al., 2011) and does not contain any hydrogen atoms, which could serve as a source of laser-generated OH via abstraction by O(1D) atoms (Stevens et al., 1994; Dubey et al., 1996). However, C$_3$F$_6$ must be diluted in an inert gas before it can be flowed through MFCs, and its availability in the UK became limited in 2015. Following Novelli et al. (2014a), we therefore used propane for most laboratory experiments and all ambient measurements, despite the fact that it reacts more slowly with OH ($k_{298} = 1.1 \times 10^{-12} \text{cm}^3\text{molec.}^{-1}\text{s}^{-1}$ Sander et al., 2011).

As shown in Fig. 2, the scavenger (0–50 sccm) is diluted in a flow of N$_2$ (500 sccm, BOC, 99.998 %) prior to injection, which is controlled using a solenoid valve (Metron Semiconductors). Any dead volume after the valve is purged continuously by the N$_2$ dilution flow, using a narrow-bore injector inserted through the tee after the valve, with the injector tip placed as close to the valve orifice as possible. This enables fast flushing of the system to optimise the response time before and after scavenger injection. Incorporation of the purge system resulted in pre- and post-injection stabilisation times on the order of seconds (data not shown), minimising data loss. The valve state and scavenger flow over the course of the data acquisition cycle are controlled using a custom program nested within the FAGE software.

To reduce radical wall losses, excess ambient air is drawn through the IPI to generate a sheath flow, via four ports spaced evenly around the flow tube housing as shown in Fig. 1. This minimises the FAGE sampling of air from near the walls of the cylinder, housing, and turret. The total flow rate through the IPI is 32 slm, of which 7 slm is sampled by the FAGE cell and the remainder of the flow is maintained by a vacuum pump (Agilent Technologies IDP-3 Dry Scroll Pump) and measured volumetrically using a rotameter (Brooks 2520, 4–50 L min$^{-1}$).

During interference testing experiments using the IPI system (Sect. 3.2), ozone and water vapour concentrations were measured using a commercial UV absorption instrument (Thermo Environmental Instruments Inc. 49C) and a chilled mirror dew point hygrometer (General Eastern 1311DR sensor and 4 × 4 Optica), respectively.
Table 1. Overall meteorological and chemical conditions encountered during each field campaign, including example VOCs. Values are given as the median of all points that coincide with IPI sampling periods, except for $J(O^{1}D)$ and $O_3$, which are reported as diurnally averaged maxima.

|                      | ICOZAa | AIRPRO winterb | AIRPRO summerv |
|----------------------|--------|----------------|----------------|
| **Dates (IPI sampling)** | 3–8 and 12–16 Jul 2015 | 2–7 Dec 2016 | 23 May–25 Jun 2017 |
| **Location**         | 52°57’02” N, 1°07’19” E | 39°58’28” N, 116°22’16” E |
| **Meteorological**   |        |                |                |
| Temperature (°C)     | 16     | 6.1            | 26             |
| $H_2O$ (%)           | 1.5    | 0.4            | 1.6            |
| Wind speed (ms$^{-1}$) | 5.8   | 0.9            | 0.4            |
| $J(O^{1}D)$ ($10^{-6}$ s$^{-1}$) | 16.   | 3.5            | 19             |
| **Chemical**         |        |                |                |
| $O_3$ (ppbv)         | 42     | 15             | 90             |
| NO (ppbv)            | 0.19   | 22             | 0.81           |
| NO$_2$ (ppbv)        | 2.2    | 33             | 17             |
| CO (ppbv)            | 100    | 1120           | 460            |
| Propane (ppbv)       | 0.26   | 6.2            | 3.8            |
| Isoprene (ppbv)      | 0.02   | 0.07           | 0.38           |
| Benzene (ppbv)       | 0.03   | 1.4            | 0.46           |
| $k’_{OH}$ (s$^{-1}$) | 4.4    | 38             | 25             |

*a Integrated Chemistry of OZone in the Atmosphere. b An integrated study of AIR pollution PROcesses in Beijing (Shi et al., 2019).

2.3 Field measurement sites

Ambient measurements of $OH_{wave}$ and $OH_{chem}$ were made using the Leeds IPI–FAGE instrument during three separate intensive field campaigns in different locations and seasons. This allowed for the investigation of potential OH interference under markedly contrasting conditions. For all three field campaigns, measurements of $OH$, $H_2O$, and partially speciated RO$_2$ were made using the Leeds FAGE instrument (4 m above ground level), operated in the sequential detection modes described in Sect. 2.1 (Whalley et al., 2013). Total OH reactivity, $k’_{OH}$, was also measured, using the laser flash photolysis–LIF instrument described in detail by Stone et al. (2016). A range of supporting chemical, aerosol, and meteorological parameters were measured, with instruments situated either in buildings or shipping containers at each of the two sites. Gas-phase chemical observations included water vapour, NO$_x$, NO$_y$, O$_3$, CO, SO$_2$, HONO, HCHO (Cryer, 2016), CINO$_2$, VOCs, and OVOCs. Photolysis rates ($J$) for a variety of species, including $O_3$ (→ $O(1D)$), NO$_2$, HCHO, HONO, and CINO$_2$, were measured using a 2$\pi$ actinic receiver optic, Meteorologie Consult GmbH, coupled to an Ocean Optics QE Pro spectrometer, and $J(O^{1}D)$ was also measured using a 2$\pi$ filter radiometer (Meteorologie Consult GmbH) (Bohn et al., 2008). The meteorological and chemical conditions, including some example VOCs, encountered during each campaign are summarised in Table 1 and discussed in further detail below.

The first deployment of the Leeds IPI was during the ICOZA (Integrated Chemistry of OZone in the Atmosphere) project, which took place in July 2015, at the Weybourne Atmospheric Observatory (WAO), Weybourne, located on the northern Norfolk Coast, UK (52°57’02” N, 1°07’19” E, 15 m a.s.l.). The WAO is a Global Atmospheric Watch (GAW) regional station, and the site is impacted by a range of contrasting air masses, from clean Arctic air to processed emissions from the UK (e.g. London, which is located $\sim$ 180 km SSW of the observatory) and northern Europe. The aim of this field campaign was to improve understanding of ozone chemistry through integrated measurements of $P(O_3)$, i.e. the chemical or in situ ozone production rate (OPR) (Cazorla and Brune, 2010; Cazorla et al., 2012), with comparisons to a range of other observational and model approaches. Two continuous IPI sampling periods were conducted in the middle of the campaign, separated by a few days (3–8 and 12–16 July), with a total of 9 d where $OH_{chem}$ measurements are available around midday (fully continuous measurements were not always possible due to multiple instrument problems, e.g. cell blockages and laser power drops). For other times, only measurements of $OH_{wave}$ are available. During the IPI sampling periods, power cuts on the nights of 3–4 and 6–7 July resulted in extended data loss.

In general, the ICOZA campaign was characterised (Table 1) by moderate temperatures (16 °C median), high humidity (RH $\sim$ 80%), and strong wind speeds ($\sim$ 6 m s$^{-1}$), as might be expected at a temperate, coastal location in the
sumertine. The predominant wind sector, based on wind direction measurements at the site, was westerly (∼30 %), followed by southwesterly (∼20 %) and southerly (∼15 %). Back-trajectory analysis showed that during IPI sampling periods, the site was predominantly under the influence of Atlantic air (Cryer, 2016). These air masses had spent a considerable amount of time (∼1 d) over the UK, often encountering emissions from urban areas, which underwent photochemical ageing during their transport to the WAO site. Overall, the levels of pollution observed at the site were moderate, and the lowest of the three field campaigns discussed in this work (Table 1), and levels of isoprene were low. However, ozone mixing ratios were relatively high, with a diel-average maximum of ∼40 ppbv, driven in part by strong UV and near-UV radiation.

The Leeds IPI was deployed during another two campaigns at the Institute of Atmospheric Physics (IAP, 39°58′36.06″N, 116°22′53.69″E), an urban site in Beijing, China, during winter (November–December) 2016 and summer (May–June) 2017, as part of the AIRPRO (an integrated study of AIR pollution PROcesses in Beijing) project. AIRPRO is part of the wider APHH (Air Pollution and Human Health in a Chinese megacity) project (Shi et al., 2019), a joint UK-China programme. The aims of AIRPRO included the assessment of how pollutants are transformed and removed through transport, chemical, and photolytic processes, with a particular emphasis on the identification of the dominant oxidative degradation pathways (i.e. the relative importance of reactions with OH, NO3, and O3). The AIRPRO project allowed for the assessment of OH measurement interferences under the highly polluted conditions of the megacity Beijing, situated on the heavily industrialised North China Plain. In winter, the site is impacted by urban and regional anthropogenic emissions, in particular those from the combustion of fossil fuels for residential heating. During summer, the site is subject to additional biogenic influences, and strong photochemical activity results in high rates of ozone production. In winter, OHwave and OHchem were measured simultaneously for 6 d of the campaign. In summer, almost 1 month of near-continuous IPI data are available, with 1 d of interruptions due to IPI testing (see Sect. 3.1.2).

For both AIRPRO field intensives, the predominant wind sectors were westerly and southerly-southeasterly, which generally result in higher pollutant concentrations (Chen et al., 2015). Indeed, the two campaigns were subject to high pollutant concentrations, as illustrated by the elevated levels of NO2, CO, propane, benzene, and kOH, many of which were over an order of magnitude higher than ICOZA (Table 1). In addition, the biogenic influence during summer is clear from the relatively high isoprene concentrations observed, ∼0.4 ppbv on average but reaching up to 7.9 ppbv, a level considerably higher than those observed in some forested environments. Despite similar J(O1D) values between ICOZA and AIRPRO summer, the higher VOC loadings during the latter resulted in increased production of ozone (90 ppbv diurnally averaged maximum). In contrast, AIRPRO winter was characterised by small ozone mixing ratios (15 ppbv diurnal maximum), as a consequence of high NO levels (median 22 ppbv) and weak UV radiation. In summer, NO levels were high in the morning (∼14 ppbv at 06:00 China standard time, CST) but surprisingly low in the afternoon, with diel-average median levels of ∼0.5 ppbv (15:00–18:00 CST).

For all ambient observation periods, the IPI data acquisition cycle consisted of 5 min of online wavelength and 30 s of offline wavelength (spectral background) integration, where the online period was split into 2 min of OH measurements and 2 min of propane addition to the IPI flow tube (chemical background), followed by 1 min of HO2 measurements (by the addition of NO to the FAGE cell). In terms of instrumental operation, the only difference between ICOZA and the AIRPRO campaigns was the use of different propane flows in the IPI. The propane concentration in the IPI flow tube was ∼110 ppmv (kOH ∼3 000 s−1, τOH ∼0.3 ms) during ICOZA and AIRPRO winter, but after internal removal experiments revealed that the propane level could be increased further (see Sect. 3.1.3), a 10-fold higher concentration (∼1100 ppmv), resulting in a concomitant reduction in the OH lifetime, was used for the AIRPRO summer campaign. On 1 d with high ozone (up to ∼80 ppbv) and moderate isoprene (∼0.5–1 ppbv) levels, the propane mixing ratio was reduced to ∼110 ppmv, but this had no observable effect on the background signals obtained for the summer data. All ambient OHwave data presented here have been corrected for the known interference from O3 in the presence of H2O vapour (see Sect. 3.2.1).

3 Results
3.1 IPI characterisation
3.1.1 Sensitivity

The presence of additional surfaces in the IPI system may result in radical wall losses and therefore reduce the overall FAGE instrument sensitivity. To test for potential OH losses in the IPI flow tube, OH radicals were generated using a 184.9 nm Hg lamp placed ∼19 cm from the instrument inlet, so that ambient air with elevated radical concentrations (∼2–7 × 107 molec. cm−3) was sampled, alternating between IPI and non-IPI sampling (Fig. 3), where for the latter the entire IPI assembly was removed. The dominant source of OH was the photolysis of ambient water vapour at 184.9 nm. In these experiments, the Hg lamp was placed sufficiently far away from each inlet within a large tent enclosure on the container roof, such that it could be assumed that OH concentrations were uniform in the region the inlet sampled from. Otherwise, the difference in inlet height between IPI and non-IPI
Figure 3. Testing of OH losses in the IPI system. Each panel shows repeat measurements of OH signals ($\pm 2\sigma$) over the course of 1 d, where high OH concentrations were generated using a 184.9 nm Hg lamp placed near the instrument inlet. Blue and red markers denote individual measurements (one measurement “loop”, i.e. one wavelength online–offline cycle) performed with (“IPIon”) and without (“IPIoff”) the IPI system, respectively. Solid lines correspond to the average signals for each day, with $2\sigma$ standard deviations (SD) shown by the dashed lines.

Sampling may have resulted in different OH concentrations being sampled, e.g. due to differences in O$_3$ absorption at 184.9 nm (O$_3$ has a high cross section at this wavelength), which would affect the light flux at the point of sampling and hence the concentration of OH generated. Since ambient variability (e.g. in NO$_x$ levels) also affects the atmospheric radical concentrations, the IPI–non-IPI cycle was repeated several dozen times on 3 different days within the tent enclosure to ensure sufficient averaging of the results. Any differences in wind speed or direction during the different days are not important because of the tent enclosure. Based on the averages for each set of repeat measurements in Fig. 3, these experiments yield a mean $\pm 2\sigma$ IPIoff/IPlon ratio of 1.043 $\pm$ 0.023, i.e. a $< 5\%$ sensitivity reduction due to the presence of the IPI. While HO$_2$ loss was not tested, the relative sensitivity is assumed to be closer to unity since it is less reactive than OH. In either case, the correction is smaller than the total instrumental uncertainty ($\sim 26\%$ at $2\sigma$), and as such no corrections were applied to OH or HO$_2$ calibration factors for the final workup of ambient data collected during IPI sampling periods. In other words, we assume negligible transmission losses within the IPI, and the OH calibration factor we applied to ambient data was the same for (1) OH$_{\text{wave}}$ without IPI sampling, (2) OH$_{\text{wave}}$ during IPI sampling, and (3) OH$_{\text{chem}}$ during IPI sampling. However, it
should be noted that in the field, calibrations are normally carried out without the IPI system present.

The lack of OH loss in the IPI system is further supported by another test conducted in the field during the summer 2017 AIRPRO campaign, where on 1 d of the campaign sequential measurements of OH$_{\text{wave}}$ were taken with and without the IPI assembly present. While this was not a formal intercomparison, the summer 2017 campaign provided ideal conditions to assess IPI losses, considering the very high radical concentrations observed (OH was frequently $>1 \times 10^7$ molec. cm$^{-3}$) in Beijing and thus a good signal-to-noise ratio. The results of this experiment are shown for OH in Fig. 4. It can be seen that if no correction for a reduction in sensitivity is applied, adjacent IPIt and IPOn periods of data are qualitatively in agreement, with no discontinuities in the temporal profile, implying that the IPI sensitivity loss is close to zero under field operating conditions. Similar results were obtained for HO$_2$ (data not shown).

### 3.1.2 External OH removal

The external OH removal efficiency in the IPI system is controlled by the injection height, the choice of scavenger (i.e. the rate coefficient of the reaction of scavenger with OH), the scavenger and N$_2$ dilution gas flows delivered to the injectors, and the sheath flow. A key requirement here is efficient mixing of the scavenger into the ambient air stream, which is difficult considering the fast flow rate and hence short residence time of air in the IPI flow tube. Additionally, it is important to consider that some reaction of the scavenger may occur inside the fluorescence chamber (internal OH removal, Sect. 3.1.3). This would give rise to a positive bias in ambient OH concentrations measured using the OH$_{\text{chem}}$ method, as internal OH removal could result in loss of interfering OH and therefore an apparent reduction in the true background signal.

External OH removal experiments were performed by supplying known concentrations of OH and HO$_2$ to the instrument using the calibration wand described in Sect. 2.1. However, in contrast to normal calibration procedures where the wand is held at 45° to the pinhole (to overfill the pinhole and minimise sampling of pockets of air which may have been in contact with the metal pinhole surface), IPI characterisation experiments were performed with the wand positioned parallel to the direction of flow within the IPI (i.e. 90° relative to the plane of the pinhole), with a distance of $\sim$ 3 cm between the wand exit and the PFA flow tube. The high flow through the calibration wand (40 slm) ensured that an excess of calibration gas was delivered to the IPI system (sample flow $\sim$ 32 slm).

The external OH removal efficiency (RE$_{\text{external}}$) may be calculated from the proportion of OH remaining (R$_{\text{ OH \_ external}}$) after injection of the scavenger, obtained from the ratio of the OH signals in the presence (S$^{\text{OH\_ scavenger}}$) and absence (S$^{\text{OH}}$) of the scavenger:

$$R^{\text{OH\_ external}} = \frac{S^{\text{OH\_ scavenger}}}{S^{\text{OH}}}.$$  

(3)  

$$R^{\text{OH\_ external}} = 100 - R^{\text{OH\_ external}}.$$  

(4)  

Initial tests included variation of the N$_2$ dilution flow; however, the OH removal efficiency was generally low (data not shown), likely due to poor mixing of the scavenger into the sampled air when the flow rate from the injector is small. As a result, the N$_2$ dilution was set to the maximum flow of the MFC used (0.5 slm) for all subsequent experiments. Any further dilution of the ambient air stream would result in a loss of sensitivity towards the detection of radicals; however, at 0.5 slm the dilution flow is virtually negligible compared to the total flow rate in the IPI system (32 slm). In other preliminary experiments, the scavenger was injected closer to the FAGE inlet (1.0 and 2.5 cm), but this also resulted in poor external OH removal owing to the shorter residence time between scavenger injection and FAGE sampling.

The scavenging efficiency was determined for both propane and C$_3$F$_6$, with good agreement between the two scavengers. Figure 5 shows the remaining OH signal as a function of the OH reactivity ($k^{\text{OH}} = k^{\text{OH + scavenger}}$ [scavenger]) calculated in the flow tube, which normalises the scavenger concentrations according to their different reaction rates with OH. The observed removal efficiency is in broad agreement with the theoretical scavenging efficiency, based on the residence time in the flow tube ($\sim$ 20 ms, assuming plug flow) and assuming perfect mixing, suggesting that in the Leeds IPI system the scavenger is well mixed into the gas sampled by the FAGE cell. An optimum removal of virtually 100% (OH remaining $\pm$2σ = (0.030 ± 0.091)%) was observed at $k^{\text{OH}} \sim 3000$ s$^{-1}$, equivalent to $\sim$ 110 ppmv (2.7 $\times$ 10$^{15}$ molec. cm$^{-3}$) propane. This scavenger concentration was used for measurements of OH$_{\text{chem}}$ during the summer 2015 ICOZA project and winter 2016 AIRPRO project. For the summer 2017 AIRPRO project, a 10-fold higher scavenger concentration was used ($\sim$ 1100 ppmv propane), after internal removal experiments revealed no loss of internal OH at this higher concentration, as discussed in detail in the next section.

### 3.1.3 Internal OH removal

Internal removal of OH was quantified by Mao et al. (2012) after forming OH inside the PSU ground-FAGE cell using a Hg lamp and comparing the OH signal with and without the presence of the scavenger (C$_3$F$_6$), which was added externally in the IPI system. It was found that most of the internal removal occurred in the instrument inlet, rather than in the OH detection axis, with a total loss of $\sim$ 20%. Internal removal was not tested in the laboratory by Novelli et al. (2014a) for the Max Planck Institute for Chemistry (MPIC) FAGE instrument, but instead they limited the scavenger (propene and propane) concentration such that the ex-
ternal OH removal efficiency was < 95 %, to minimise possible reaction of the scavenger with OH inside the fluorescence cell. However, during ambient, night-time tests (constant atmospheric OH concentration assumed), no change in the OH background signal was observed after increasing the scavenger concentration by a factor of 7, providing evidence for a lack of internal removal (Novelli et al., 2014a).

In the present study, a novel approach was devised to quantify internal removal of OH in the Leeds IPI–FAGE instrument. First, under otherwise identical experimental conditions to those for external OH removal tests, sufficient CO (75 sccm, 95 ppmv) was added to the calibration wand to verify that the OH formed (alongside HO₂) from the photolysis of water vapour was almost quantitatively converted to HO₂ ((98.0 ± 0.4) %, data not shown). Secondly, in addition to the calibration wand CO flow, a high flow of NO (50 sccm) was injected through 0.125 in. stainless steel tubing inside the FAGE cell, with the injector tip positioned centrally just below the turret pinhole, to reconvert the HO₂ back to OH for LIF detection; these experimental conditions ensured a fairly high HO₂-to-OH conversion efficiency of approximately 30 %. In this manner, OH was only generated inside the FAGE cell, and not in the IPI flow tube, such that any change in the fluorescence signal could be attributed to internal reaction of OH with propane, rather than reaction in the flow tube. The procedure for determination of internal OH removal bears some resemblance to that used for ambient detection of RO₂ using the ROₓ LIF technique (Fuchs et al., 2008; Whalley et al., 2013), i.e. the external conversion of all radical species to HO₂ before internal conversion to OH within the fluorescence cell.

The internal OH removal efficiency (RE_{OH internal}) was quantified in an analogous manner to the external scavenging efficiency, using the total fluorescence signal in the presence (S_{HO² scavenger}) and absence (S_{HO² scavenger}) of the scavenger:

\[ R_{OH internal}(\%) = 100 \times \frac{S_{HO² scavenger}}{S_{HO² scavenger}}, \quad (5) \]

\[ RE_{OH internal}(\%) = 100 - R_{OH internal}. \quad (6) \]

Figure 6 shows a time series of the LIF signal during two example internal removal experiments. Here, the LIF signal represents the sum of signals from OH and HO₂, since they are produced in a 1 : 1 ratio (Fuchs et al., 2011) in the calibration wand. For both of the propane mixing ratios used, which were shown to result in near complete external OH removal in Sect. 3.1.2, there was no obvious decrease in the LIF signal, indicating no significant internal removal of OH. The average ± 2σ internal OH removal observed for repeat experiments was (0.0 ± 4.0) % (Table 2) at a propane mixing ratio of ~110 ppmv (ICOZA and AIRPRO winter conditions). For repeat experiments at the higher propane mixing ratio used during the AIRPRO summer field campaign (~1100 ppmv), the internal removal was still very small and almost insignificant (2.9 ± 6.6) %. Table 2. The observed internal OH removal may be compared to that which might be expected the-
Based on this, an internal OH removal efficiency of between the pinhole and the laser axis is estimated. However, assuming a constant gas velocity, the residence time likely slows down between pinhole sampling and NO detection (Creasey et al., 1997b; Whalley et al., 2013). The gas residence time of 0.9 ms between NO injection and OH detection (i.e. total of 18 cm between the pinhole and 7.5 cm away from the laser axis (i.e. total 25.5 cm) can be used to calculate the scavenging efficiency assuming perfect mixing, using the estimated residence time, $\tau \sim 20$ ms. The propane OH reactivity used for the ICOZA (Integrated Chemistry of OZone in the Atmosphere) and AIRPRO winter campaigns is given but that used for AIRPRO summer is off-scale.

In the ambient pressure flow tube, a propane mixing ratio of 1100 ppmv equates to $k'_{\text{OH}} = 30,000$ s$^{-1}$, but this is a factor of 760/1.5 lower in the detection cell (i.e. the ratio of ambient to cell pressure), 59 s$^{-1}$ (assuming constant gas density and no change in the OH + propane rate coefficient). Under normal operation, NO injection occurs 10.5 cm below the pinhole and 7.5 cm away from the laser axis (i.e. total of 18 cm between the pinhole and detection volume), with a residence time of 0.9 ms between NO injection and OH detection (Creasey et al., 1997b; Whalley et al., 2013). The gas likely slows down between pinhole sampling and NO injection but, assuming a constant gas velocity, the residence time between the pinhole and the laser axis is estimated at $\sim 2$ ms. Based on this, an internal OH removal efficiency of $\sim 12\%$ is calculated, which is higher than observed, likely because the assumption of constant gas velocity is invalid (i.e. the real residence time is closer to $\sim 1$ ms) and mixing between HO$_2$ and NO from the injector is not instantaneous. However, it should be noted that this calculation also neglects the fact that the density is higher in the jet and the perturbation to normal flow caused by moving the NO injector close to the pinhole.

In the internal removal experiment, OH is not formed instantly at the pinhole but is built up linearly by HO$_2$/NO conversion along the line from the pinhole to the laser axis. Therefore, the experimental internal removal may not be directly compared with the theoretical estimate. In such a sequential reaction system, the OH scavenging is about half as efficient as that for the case where OH is formed as an instant point source at the pinhole. Thus, the experimental value should be doubled to $(5.8 \pm 13)\%$, which is in reasonable agreement with the theoretical value. From this, we cannot rule out a small internal OH removal on the order of 10% at the higher propane level used for the AIRPRO summer campaign. However, no such corrections were applied to the ambient data featured in this work.

### 3.2 Interference testing experiments

#### 3.2.1 O$_3$ + H$_2$O vapour

In LIF–FAGE instruments, there is a known interference due to laser-generated OH from ozone photolysis in the presence of water vapour (Fuchs et al., 2016; Griffith et al., 2016; Tan et al., 2017). This interference was quantified by Whalley et al. (2018) and characterised in further detail in the present work. In these experiments, ozone was generated from the 184.9 nm photolysis of oxygen in a 12–20 slm flow of zero air using a Hg(Ar) pen-ray lamp (LOT LSP035). Another 12–20 slm of zero air was humidified using a water (HPLC-grade) bubbler. The two zero air flows were combined and delivered to the calibration wand, from which the IPI sampled in a manner analogous to the experiments conducted to investigate external and internal OH removal discussed above. Ozone mixing ratios in the range 0–2.5 ppmv were generated by varying the Hg lamp current (0–21 mA), while water vapour volume mixing ratios in the range 0.1 %–1.0 % were produced by varying the flow through the bubbler, or by bypassing it completely, and the total flow (32 slm) was compensated for by changing the dry zero air flow. Laser power (LP) at 308 nm was varied in the range 3–17 mW by varying the ratio of acetone : water in a cuvette placed before the fibre launcher that is used to send laser light to the detection cells.

Figure 7 shows the results of O$_3$ + H$_2$O vapour interference tests. It can be seen that the interference signal ($OH_{\text{int}} = OH_{\text{wave}} - OH_{\text{chem}}$) is linear in both ozone (Fig. 7a) and water vapour (Fig. 7b) mixing ratios. The quadratic dependence of the interference signal on laser power (Fig. 7c) in terms of raw count rates indicates that the interference originates from a two-photon process, as expected. However, since OH...
Table 2. Internal removal of OH (\%, ±2\(\sigma\)) as a function of propane mixing ratio in the IPI flow tube, determined as shown in Fig. 6 (see text for details).

| Propane (ppmv) | Experiment no. | Internal removal (%) |
|----------------|----------------|----------------------|
| 110 (used for ICOZA and AIRPRO winter) | 1 | -0.1 ± 4.8 |
| | 2 | 0.3 ± 7.7 |
| | 3 | -0.9 ± 16 |
| | | Weighted average ±2\(\sigma\) | 0.0 ± 4.0 |
| 550 | 1 | 1.0 ± 9.6 |
| 1100 (used for AIRPRO summer) | 1 | 1.9 ± 12 |
| | 2 | 4.2 ± 11 |
| | 3 | 2.5 ± 11 |
| | | Weighted average ±2\(\sigma\) | 2.9 ± 6.6 |

Figure 6. Time series of the LIF signal during internal OH removal experiments. The raw 1 s data are given by the grey line. NO was continuously added to the FAGE cell during these experiments (to form OH internally), and points where propane was added to the IPI flow tube are indicated by the orange-shaded panels, with the corresponding signal averages (±\(\sigma\)) shown as markers (see text for details). The first experiment (a) corresponds to the propane mixing ratio used for ICOZA, while the second (b) corresponds to that used for AIRPRO. The results of the internal OH removal experiments are summarised in Table 2.

data are normalised to laser power, the equivalent OH concentrations are linear with respect to laser power. Thus, overall, OH\(_{\text{int}}\) is linear in ozone, water vapour, and laser power. Normalisation of the slope in Fig. 7a yields the following relation:

\[
[\text{OH}_{\text{int}}] = (565 ± 42) \text{ molec. cm}^{-3} \text{ ppbv}^{-1} \%^{-1} \text{ mW}^{-1} \times [\text{O}_3] \times [\text{H}_2\text{O}] \times \text{LP},
\]

where [\text{O}_3], [\text{H}_2\text{O}], and LP are in units of ppbv, %, and mW, respectively. Under typical atmospheric conditions of [\text{O}_3] = 50 ppbv and [\text{H}_2\text{O}] = 1 \%, and a typical instrument laser power of 15 mW, the interference signal is equivalent to an OH concentration of 3.9 × 10^5 molec. cm\(^{-3}\). This signal is slightly smaller than the instrumental limit of detection (LOD) of \(\sim 7 \times 10^5\) molec. cm\(^{-3}\) at a signal-to-noise ratio (SNR) of 2, but nonetheless it was used to correct the ambient OH\(_{\text{wave}}\) data presented in Sect. 3.3, using co-located measurements of ozone and water vapour.
Table 3. Summary of interference tests with O$_3$ and isoprene (ISO) in the presence of propane (PROP), based on the data in Fig. 8. n/a – not applicable.

| Test | O$_3$ (ppmv) | H$_2$O (%) | ISO (ppmv) | PROP (ppmv) | Obs. | Correction due to O$_3$ + H$_2$O vapour interference$^b$ | OP3 levels$^b$ | AIRPRO 2017 levels$^c$ |
|------|-------------|------------|------------|-------------|------|--------------------------------------------------|----------------|------------------|
| A    | 1.64        | 0.73       | 0          | 110         | 1.0 \times 10^7 | 0       | n/a               | n/a            |
| B    | 1.86        | 0.07       | 16         | 110         | 1.9 \times 10^7 | 1.8 \times 10^7 | 1               | 430            |
| C    | 1.83        | 0.98       | 16         | 110         | 1.6 \times 10^7 | 8.0 \times 10^5 | 1               | 19             |
| D    | 1.85        | 0.07       | 16         | 1100        | 1.4 \times 10^7 | 1.3 \times 10^7 | 15              | 310            |

$^a$ Corrected using Eq. (7). $^b$ Oxidant and Particle photochemical processes field campaign in Borneo, 2008: average O$_3$ = 10 ppbv; ISO = 3.5 ppbv.

$^c$ Diurnally averaged maximum O$_3$ = 90 ppbv; overall maximum ISO = 7.9 ppbv.

3.2.2 Isoprene ozonolysis

To test for interferences from isoprene (ISO) ozonolysis products, isoprene (∼16 ppmv), and ozone (∼1.8 ppmv) were mixed in the calibration wand and the scavenger (propane, PROP) was injected into the IPI flow tube. The propane concentrations were set to those used for ambient OH$_{chem}$ measurements, such that the tests were representative of normal atmospheric sampling (i.e. to test whether an interference signal would remain in ambient data). However, to generate sufficient OH signal for quantitative analysis, ozone and isoprene were introduced at concentrations that far exceeded their typical ambient levels (Table 3). Unlike previous tests of interferences from alkene ozonolysis (Novelli et al., 2014b), low [O$_3$] : [ISO] ratios were used to suppress the signal contribution from the atmospheric (real) OH generated by ozonolysis (i.e. isoprene acted as an additional OH scavenger). To allow sufficient time for steady-state conditions to develop, the IPI did not sample from the calibration wand directly, but instead a 30 cm flow tube (polycarbonate, ID ∼ 19 mm) was used to extend the IPI (which sampled wand gas at the normal IPI flow rate of ∼ 32 slm, residence time for O$_3$ + isoprene reaction ∼ 0.15 s).

Time series of the interference testing experiments conducted using the IPI are shown in Fig. 8. In Fig. 8a, no isoprene is added, but due to ozone photolysis in the presence of high [H$_2$O] (0.73 %) an interference signal (OH$_{int}$) is observed (i.e. signal in the presence of propane is higher than the offline signal). The magnitude of this signal (OH$_{int}$ ∼ 1.0 \times 10^7 molec. cm$^{-3}$) yields a scale factor of 510 ± 270 molec. cm$^{-3}$ ppbv$^{-1}$ %$^{-1}$ mW$^{-1}$ when linearly extrapolated down from the measured [O$_3$], [H$_2$O], and LP, in agreement with the 565 ± 42 molec. cm$^{-3}$ ppbv$^{-1}$ %$^{-1}$ mW$^{-1}$ in Eq. (7).

In Fig. 8b, ozone and isoprene react under dry conditions, and an interference signal is observed again. The low H$_2$O (0.07 %) suppressed the O$_3$ + H$_2$O interference, such that this cannot explain the magnitude of OH$_{int}$ (∼ 1.9 \times 10^7 molec. cm$^{-3}$, Table 3), suggesting that OH was formed internally from a reaction other than O$_3$ + H$_2$O. Under high-humidity (H$_2$O ∼ 1 %) conditions (Fig. 8c), OH$_{int}$ (∼ 1.6 \times 10^7 molec. cm$^{-3}$) was similar, but in this case the signal can be explained almost entirely by the O$_3$ + H$_2$O interference. Under dry conditions with a 10-fold higher concentration of propane (as used for the AIRPRO summer campaign, Fig. 8c), the interference signal from Fig. 8b was reduced but remained elevated relative to the offline signal (OH$_{int}$ ∼ 1.4 \times 10^7 molec. cm$^{-3}$), where again the contribution from O$_3$ + H$_2$O cannot explain the discrepancy. The decrease in OH$_{int}$ between Fig. 8b and d may be attributed to the suppression of steady-state OH generated from ozonolysis, but the remaining signal implies that OH was also formed internally in both cases. For the dry, low-propane experiment (Fig. 8b), the magnitude of the OH signal is much higher than that calculated from a steady-state model (∼ 1.4 \times 10^6 molec. cm$^{-3}$).

The suppression of the interference signal attributable to O$_3$/isoprene only (i.e. O$_3$ + H$_2$O corrected) by the addition of water vapour (H$_2$O ∼ 1 %; see Fig. 8c) suggests that the internal OH may have been formed from SCIs. The simplest C1 and C2 SCIs are known to react quickly with the water vapour dimer (k ∼ 4–7 × 10$^{-12}$ cm$^3$ molec.$^{-1}$ s$^{-1}$) at 298 K for CH$_2$OO: Chao et al., 2015; Lewis et al., 2015) and monomer (k ∼ 1–2 × 10$^{-14}$ cm$^3$ molec.$^{-1}$ s$^{-1}$ for anti-CH$_2$CHOO: Taatjes et al., 2013; Sheps et al., 2014; Lin et al., 2016), respectively. Reaction with the water vapour monomer was also shown to be relatively fast (k ∼ 1.2 × 10$^{-15}$ cm$^3$ molec.$^{-1}$ s$^{-1}$, k$_{loss}$ ∼ 300 s$^{-1}$ at ∼ 1 % H$_2$O) for the ensemble of SCIs, including the C1 SCI, generated from isoprene ozonolysis (Newland et al., 2015).

However, regardless of whether the signal observed at high propane is due to internally formed OH, which may have originated from SCIs, the equivalent OH concentrations are negligible when extrapolated back to ambient chemical conditions (Table 3). Assuming a linear dependence of the interference signal on both ozone and isoprene, the interference (after O$_3$ + H$_2$O correction) is equivalent to <10$^5$ molec. cm$^{-3}$ at the ozone (10 ppbv) and isoprene (3.5 ppbv) levels measured in a low-NOx, biogenic environ-
Figure 7. OH interference (±1σ) from O$_3$ + H$_2$O as a function of (a) O$_3$, (b) H$_2$O, and (c) laser power. The interference signal is linear in O$_3$ and H$_2$O mixing ratios and quadratic in laser power, confirming the two-photon nature of the process. Normalising the slope in panel (a) to O$_3$ = 1 ppbv, H$_2$O = 1 %, and laser power = 1 mW yields an OH interference equivalent to a concentration of 565 ± 42 molec. cm$^{-3}$.

ment during the Oxidants and Particle photochemical processes (OP3) campaign in Borneo, 2008 (Hewitt et al., 2010). Similarly, we have modelled the SCI decomposition in our FAGE cell. We assumed an ambient atmosphere containing 100 ppbv O$_3$ and 10 ppbv trans-2-butene and took the reactions, rate coefficients, and yields from Novelli et al. (2014a) and MCMv3.3.1. The rate of CI to SCI was 1 \times 10^6$ s$^{-1}$ with an SCI yield of 0.18. The unimolecular decomposition of SCI to OH was 3 s$^{-1}$ and the wall loss rate of SCI was 22 s$^{-1}$. From this we calculate an equivalent ambient pressure OH concentration of \sim 4 \times 10^3$ molec. cm$^{-3}$ from the decomposition of SCIs at our FAGE cell residence time of 2 ms. The insignificance of the interference signal for atmospherically relevant O$_3$/alkene concentrations is consistent with the results of previous interference experiments, for which equivalent OH concentrations of \sim 3–4 \times 10^4$ (Novelli et al., 2014b; Fuchs et al., 2016) and \sim 4 \times 10^5$ molec. cm$^{-3}$ (Rickly and Stevens, 2018) can be derived.

3.2.3 NO$_3$ radicals

Fuchs et al. (2016) found that, despite the absence of a hydrogen atom, NO$_3$ radicals were responsible for a small OH interference signal in the Forschungszentrum Jülich (FZJ) LIF–FAGE instrument, equivalent to an OH concentration of 1.1 \times 10^5$ molec. cm$^{-3}$ per 10 pptv NO$_3$. The OH interference scaled linearly with observed NO$_3$ mixing ratios but showed no dependence on inlet length, cell pressure, laser power, or humidity, and the background signal did not change significantly in the presence of CO scavenger, suggesting the OH was indeed being formed internally. It was postulated that the interference was a result of a heterogeneous process involv-
Figure 8. Isoprene (ISO) ozonolysis interference tests: (a) O$_3$/H$_2$O only, (b) O$_3$ and ISO under dry conditions, (c) O$_3$ and isoprene with H$_2$O added, and (d) O$_3$ and isoprene under dry conditions but with a higher concentration of propane (PROP) to remove any steady-state-generated OH. Shaded areas are periods of propane addition, and the light blue lines correspond to the calculated signals from O$_3$ + H$_2$O only (for experiments with isoprene present). The interference signals (“OH no. 2” = “offline”) were used to derive equivalent OH concentrations (OH$_{int}$), which are on the order of $\sim 1$–$2 \times 10^7$ molec. cm$^{-3}$. These experiments are summarised in Table 3. See text for further details.
Figure 9. Equivalent OH concentrations (±1σ) measured during NO$_3$ radical interference tests. NO$_3$ concentrations were calculated using a box model, and OH interference signals were corrected for the interference from O$_3$ + H$_2$O. The OH limit of detection (LOD, 6.3 × 10$^5$ molec. cm$^{-3}$, SNR = 2) is denoted by the dashed red line.

To test for an NO$_3$ interference in OH measurements made by the Leeds FAGE instrument, NO$_3$ was generated from the reaction of ozone and NO$_2$:

$$O_3 + NO_2 \rightarrow NO_3,$$  \hfill (R1)

$$NO_3 + NO_2 \rightarrow N_2O_5,$$  \hfill (R2)

$$N_2O_5 \rightarrow NO_3 + NO_2.$$  \hfill (R3)

In these experiments, ozone was generated by flowing zero air (15 slm) past a Hg lamp (LOT LSP035). A constant 0.5 slm flow of NO$_2$ (BOC, 2 ppmv) was diluted in 25 slm zero air and mixed with the zero air–ozone flow and an additional zero air dilution flow of 10 slm, in order to yield a final mixing ratio of 20 ppbv. Gas was delivered to the IPI system using the calibration wand, with a total residence time of 3.7 s for the O$_3$ + NO$_2$ reaction. Ozone mixing ratios in the range 0–2.8 ppmv (after dilution) were generated by varying the current supplied to the Hg lamp. NO$_3$ radical mixing ratios in the range 0–90 pptv were calculated based on a box model with rate constants taken from the Master Chemical Mechanism (MCM; http://mcm.leeds.ac.uk/MCM, last access: 15 December 2017) version 3.3.1 ($k_{R1} = 3.52 \times 10^{-17}$ cm$^3$ molec.$^{-1}$ s$^{-1}$, $k_{R2} = 1.24 \times 10^{-12}$ cm$^3$ molec.$^{-1}$ s$^{-1}$, and $k_{R3} = 0.045$ s$^{-1}$). These experiments were performed under dry conditions (H$_2$O ∼ 0.07 %), such that only a small correction was applied for the O$_3$/H$_2$O interference.

The results of the NO$_3$ radical interference tests are shown in Fig. 9. It can be seen that the equivalent OH signals were all <8 × 10$^5$ molec. cm$^{-3}$ and almost always below the instrument LOD of 6.3 × 10$^5$ molec. cm$^{-3}$ (SNR = 2). Unlike the dependence found by Fuchs et al. (2016), the interference signal does not increase linearly with NO$_3$. However, based on the point at the highest NO$_3$ mixing ratio of ∼90 pptv, the interference is equivalent to an insignificant ∼6 × 10$^4$ molec. cm$^{-3}$ at 10 pptv NO$_3$, or approximately half of that observed by Fuchs et al. (2016). These experiments suggest that an interference from NO$_3$ radicals is not significant for the detection of OH using the Leeds ground-based FAGE instrument.
3.3 Ambient observations of OH\text{wave} and OH\text{chem}

3.3.1 ICOZA 2015

Figure 10 shows the overall intercomparison of OH\text{wave} (with O$_3$ + H$_2$O interference as given in Eq. 7 subtracted) and OH\text{chem} measurements made during the ICOZA 2015 campaign. It is evident that the raw data (averaged for 4 min periods) are quite noisy, but averaging to 1 h improves the precision and reveals a tight correlation, with the majority of points scattered around the line of 1 : 1 agreement. An orthogonal distance regression (ODR) fit (Boggs et al., 1987), which accounts for errors in both the y and x directions, to the hourly data yields a slope of 1.160 ± 0.058 (2σ) and a negative intercept on the order of the instrumental precision. In a similar manner, an unweighted least-squares linear fit (not shown) gives a slope of 1.060 ± 0.065, an intercept of (0.5 ± 1.5) × 10$^5$ molec. cm$^{-3}$, and a correlation coefficient ($R^2$) of 0.992. Regardless of the fit method, these results show that, on average, the two OH measurements agree within the instrumental uncertainty of ∼ 26 % at 2σ.

Median hourly diurnal profiles of OH\text{wave}, OH\text{chem}, and J(O$^1$D), averaged over both IPI sampling periods, are shown in Fig. 11. The two OH measurements exhibit virtually identical profiles, with peak values of ∼ 3 × 10$^6$ molec. cm$^{-3}$ slightly before solar noon, and relatively high concentrations (∼ 1–2 × 10$^6$ molec. cm$^{-3}$) persisting into the early evening despite the concomitant falloff in J(O$^1$D). Night-time levels were generally below 5 × 10$^5$ molec. cm$^{-3}$. The variability in OH concentrations, shown only for OH\text{chem} for clarity, was high during both daytime and night-time periods.

In Fig. 10 it can be seen that some points lie substantially above the 1 : 1 line, especially for the 4 min averaged raw data. It is possible that, despite the good overall agreement between the median diurnal profiles of OH\text{wave} and OH\text{chem} in Fig. 11, OH\text{int} may have exhibited its own distinct diurnal profile, independent of atmospheric OH concentrations, for example if the interference signal was generated from a particular chemical species. However, the median diurnal profile of individual OH\text{int} measurements (= OH\text{wave} − OH\text{chem}) in Fig. 11 exhibits no obvious structure, with values scattered around zero and a mean ±2σ value of (0.3 ± 3.3) × 10$^5$ molec. cm$^{-3}$, which is well below the LOD for individual OH measurements. Similarly, the average (OH\text{wave} − OH\text{chem}) / OH\text{wave} ratio (i.e. the contribution of interferences to the total OH\text{wave} signal) was zero within error (mean ±2 standard error, SE, = 0.03 ± 0.12).

Furthermore, OH\text{int} does not exhibit any dependence when binned against various parameters (Fig. S1 in the Supplement), including those previously implicated in LIF–FAGE measurement interferences, such as J(O$^1$D) (Feiner et al., 2016); temperature (Mao et al., 2012; Novelli et al., 2017); OH reactivity (Mao et al., 2012); and O$_3$ (Feiner et al., 2016; Novelli et al., 2017), isoprene (Feiner et al., 2016), and NO (Feiner et al., 2016) mixing ratios.
Figure 11. Hourly median diurnal profiles of OH\textsubscript{wave}, OH\textsubscript{chem}, and $J$($O^1D$) (right axis) from the ICOZA campaign. Also shown (red line and markers) is the hourly median diurnal profile of OH\textsubscript{int} (= OH\textsubscript{wave} − OH\textsubscript{chem}), calculated from individual 4 min data points; the single red marker corresponds to the average $\pm 2\sigma$ of this trace. The variability (interquartile range, IQR) in OH\textsubscript{chem} measurements is denoted by the dashed grey lines and is not shown for others for clarity. OH\textsubscript{wave} data were corrected for the known interference from O\textsubscript{3} + H\textsubscript{2}O.

3.3.2 AIRPRO winter 2016

The overall agreement between the two measurements is presented in the correlation plot in Fig. 12. As with ICOZA (Fig. 10), a tight correlation is revealed after averaging the data to 1 h, and all points are distributed evenly around the line of 1 : 1 agreement. ODR fitting yields an overall slope of 1.051 ± 0.039 and a negative intercept of a similar magnitude to the instrumental precision. An unweighted least-squares linear fit (not shown) gives a slope of 0.997 ± 0.038, an intercept of (−5.12 ± 7.3) × 10\textsuperscript{5} molec. cm\textsuperscript{−3}, and an $R^2$ of 0.97.

The two measurements exhibit the same profile on a diurnal basis (Fig. 13), with a diel maximum of $\sim 3 \times 10^6$ molec. cm\textsuperscript{−3} occurring in the late morning due to the build-up of HONO overnight. At night, OH\textsubscript{chem} concentrations were close to the LOD ($\sim 2 \times 10^5$ molec. cm\textsuperscript{−3}), while OH\textsubscript{wave} measurements were frequently negative, possibly as a result of over-subtraction of the O\textsubscript{3}/H\textsubscript{2}O interference as this is subject to high uncertainty (Fig. 7). The diurnal profile of OH\textsubscript{int} is scattered around zero with a mean $\pm 2\sigma$ difference of (−0.9 ± 2.7) × 10\textsuperscript{5} molec. cm\textsuperscript{−3}, and the mean $\pm 2$ SE contribution of interferences to the total signal was −0.02 ± 0.07.

3.3.3 AIRPRO summer 2017

The intercomparison of OH\textsubscript{wave} and OH\textsubscript{chem} measurements for the AIRPRO summer campaign is shown in Fig. 14. Consistent with ICOZA and the AIRPRO winter results, the 1 h data are scattered around the 1 : 1 line, with an overall ODR slope of 1.103 ± 0.017. However, the intercept is more negative than for the other campaigns, which suggests that the O\textsubscript{3}/H\textsubscript{2}O interference may have been overestimated because it is during this campaign that the highest ozone mixing ratios ($\sim 90$ ppbv diurnally averaged maximum, Table 1) were encountered. Similarly, an unweighted least-squares linear fit to the data (not shown) yields a slope of 1.111 ± 0.029, an intercept of (−3.8 ± 1.7) × 10\textsuperscript{5} molec. cm\textsuperscript{−3}, and an $R^2$ of 0.92 (data not shown).

Again, the two measurements follow the same diurnal profile (Fig. 15), peaking in the afternoon at $\sim 1 \times 10^7$ molec. cm\textsuperscript{−3} with relatively high night-time levels of $\sim 1$−$2 \times 10^6$ molec. cm\textsuperscript{−3}. As with ICOZA and the AIRPRO winter campaign, the OH\textsubscript{int} diurnal profile does not exhibit any obvious structure, with values scattered around zero and a mean $\pm 2\sigma$ difference of (−1.6 ± 4.1) × 10\textsuperscript{5} molec. cm\textsuperscript{−3}. The mean $\pm 2$ SE (OH\textsubscript{wave} − OH\textsubscript{chem}) / OH\textsubscript{wave} ratio was −0.09 ± 0.10. During AIRPRO, measured NO\textsubscript{3} mixing ratios reached up to $\sim 100$ pptv, such that the lack of significant night-time OH interference signals is consistent with the results of NO\textsubscript{3} interference tests (Sect. 3.2.3).
Figure 12. Overall intercomparison of $OH_{\text{wave}}$ and $OH_{\text{chem}}$ observations from the winter 2016 AIRPRO campaign. Grey markers represent raw data (4 min), with 1 h averages ($\pm 2\ SE$) in red. The thick red line is the ODR fit to the hourly data, with its 95 % CI bands given by the thin red lines; fit errors are given at the $2\sigma$ level. For comparison, 1 : 1 agreement is denoted by the dashed blue line. $OH_{\text{wave}}$ data were corrected for the known interference from $O_3 + H_2O$.

Figure 13. Hourly median diurnal profiles of $OH_{\text{wave}}$, $OH_{\text{chem}}$, and $J(O^1D)$ (right axis) from the winter 2016 AIRPRO campaign. Also shown (red line and markers) is the hourly median diurnal profile of $OH_{\text{int}}$, calculated from individual 4 min data points; the single red marker corresponds to the average ($\pm 2\sigma$) of this trace. The variability (IQR) in $OH_{\text{chem}}$ measurements is denoted by the dashed grey lines and is not shown for others for clarity. $OH_{\text{wave}}$ data were corrected for the known interference from $O_3 + H_2O$. 
Figure 14. Overall intercomparison of OH_{wave} and OH_{chem} observations from the summer 2017 AIRPRO campaign. Grey markers represent raw data (4 min), with 1 h averages (±2 SE) in red. The thick red line is the ODR fit to the hourly data, with its 95% CI bands given by the thin red lines; fit errors are given at the 2σ level. For comparison, 1 : 1 agreement is denoted by the dashed blue line. OH_{wave} data were corrected for the known interference from O_{3} + H_{2}O.

Figure 15. Hourly median diurnal profiles of OH_{wave}, OH_{chem}, and J(O^1D) (right axis) from the summer 2017 AIRPRO campaign. Also shown (red line and markers) is the hourly median diurnal profile of OH_{int}, calculated from individual 4 min data points; the single red marker corresponds to the average (±2σ) of this trace. The variability (IQR) in OH_{chem} measurements is denoted by the dashed grey lines and is not shown for others for clarity. OH_{wave} data were corrected for the known interference from O_{3} + H_{2}O.
It can be seen from Fig. 14 that there is a small cluster of points that lie significantly away from both the 1 : 1 and ODR regression lines, which are characterised by high OH\textsubscript{wave} concentrations of $>1.5 \times 10^7$ molec. cm\textsuperscript{-3}. This finding was investigated further, with the results summarised in Fig. 16. Above an OH\textsubscript{chem} threshold of $\sim1.0-1.5 \times 10^7$ molec. cm\textsuperscript{-3}, the OH\textsubscript{int} signal becomes significantly greater than zero and the instrument LOD, reaching $\sim3-5 \times 10^6$ molec. cm\textsuperscript{-3} at OH\textsubscript{chem} levels of around $2 \times 10^7$ molec. cm\textsuperscript{-3}. However, these results should be treated with caution, since only a few points are available for which OH\textsubscript{chem} was present at such high concentrations. The same behaviour was not observed for either the AIRPRO winter or ICOZA campaigns, since OH\textsubscript{chem} levels did not surpass $1 \times 10^7$ molec. cm\textsuperscript{-3}, but the analogous mean values at low OH\textsubscript{chem} concentrations are in agreement with the AIRPRO summer results.

The above results suggest that in the Beijing summertime, the Leeds FAGE instrument is subject to an interference(s) at the highest OH levels, although its contribution of $\sim15\%-20\%$ (Fig. 16) is still below the instrumental accuracy of $26\%$ at $2\sigma$. This finding is consistent with the suggestion of Fittschene et al. (2019) that ROO\textsubscript{H} species, formed from RO\textsubscript{2} + OH reactions, generate an OH interference in LIF–FAGE instruments, since high OH levels would generate high RO\textsubscript{2} concentrations and favour this class of reaction. It is also possible that, for high ambient OH production rates, the scavenger cannot react with the sampled OH sufficiently quickly, leading to elevated but spurious OH\textsubscript{chem} background signals. Although a modelling study of the inlet chemistry would be required to fully assess this hypothesis, it is likely not the case considering that the propane concentration used during AIRPRO results in an OH lifetime of $\sim0.03$ ms in the IPI flow tube, in comparison to a residence time of $\sim20$ ms (i.e. $\sim700$ OH lifetimes).

No clear dependences were found when OH\textsubscript{int} was binned against various parameters previously implicated in OH measurement interferences (Fig. S2), although it can be seen that OH\textsubscript{int} was marginally higher in the highest temperature, $J(O^1\Delta)$, and isoprene bins.

4 Discussion

The results from the three field campaigns that feature in this work demonstrate that, in moderately to highly polluted conditions, the Leeds ground-based FAGE instrument does not suffer from substantial interferences in the measurement of OH using the conventional, wavelength modulation background technique, OH\textsubscript{wave}. This is illustrated best by the slopes of the overall measurement intercomparison plots (Figs. 10, 12, and 14), which ranged from 1.05 to 1.16. However, while the deviations of these slopes from 1 are small, they are still significant, suggesting the presence of unknown OH interferences. Nonetheless, such unknown interferences are well within the instrumental uncertainty of $\sim26\%$ at $2\sigma$.

With respect to previous studies during which OH has been measured by a LIF instrument equipped with a scavenger injector, the significance of interferences during the campaigns that feature in this work are amongst the lowest observed (Table 4). This can likely be attributed to two main factors: environment and instrumental. In terms of the former, none of the field campaigns described in the present study took place in forested environments, where the most significant interferences have been observed (Mao et al., 2012; Novelli et al., 2014a; Feiner et al., 2016). However, as mentioned previously, the AIRPRO summer campaign did share some characteristics, in that high BVOC and low NO mixing ratios were observed in the afternoon. Despite this, OH\textsubscript{wave} and OH\textsubscript{chem} were in good agreement. Although AIRPRO summer took place in a city, its results do provide confidence in previous measurements of OH using the same instrument, and support the hypothesis that there are unknown OH sources in the atmosphere.

The insignificance of daytime interferences during the AIRPRO campaigns is consistent with results of another urban study, CalNex-LA (Research in California at the Nexus of Air Quality and Climate Change) (Griffith et al., 2016). The O$_3$/H$_2$O interference is much higher (up to $\sim4 \times 10^6$ molec. cm\textsuperscript{-3} OH equivalent during CalNex-LA) in the Indiana University (IU) LIF instrument (Dusar et al., 2009), such that the daytime contributions of $\sim33\%$ can be explained entirely by this known interference. However, measurements made at a nearby site during the same study (CalNex-SJV) showed daytime contributions of $\sim20\%$ (Brune et al., 2016), although this may be related to instrumental differences, as discussed below. On average, interferences were not observed in the daytime during ICOZA, but they were observed in other coastal campaigns, namely DOMINO (Diel Oxidants Mechanisms In relation to Nitrogen Oxide), HO$_x$ ($\sim50\%$) (Novelli et al., 2014a), and CYPHEX (CYPHotochemistry ExPeriment, $\sim45\%$) (Malik et al., 2018), as well as in rural regions, such as HOPE (Hohenpeißenberg Photochemistry Experiment, $20\%$–$40\%$) (Novelli et al., 2014a). Studies of the North China Plain have revealed small interferences on the order of $0\%$–$10\%$ (Tan et al., 2017, 2018), with slightly higher but variable contributions of $0\%$–$20\%$ in the Pearl River Delta (Tan et al., 2019).

The second major reason for the differences in contributions between the studies listed in Table 4 is likely instrumental effects. For the campaigns in which the highest OH interferences have been observed (Mao et al., 2012; Novelli et al., 2014b; Feiner et al., 2016), OH measurements were made using the Max Planck Institute (MPI) (Martinez et al., 2010) and PSU (Faloona et al., 2004) LIF instruments. These instruments feature laser multi-pass detection cells, which give rise to larger detection volumes and increased UV fluence, although this may not be relevant considering that the interference signals did not display any laser power dependence for these instruments. The Leeds instrument also differs in
terms of cell geometry, where the HOX cell is composed of a short (5 cm) turret inlet on top of a large fluorescence cell (additional ~ 8 cm to laser axis, ~ 13 cm total length, and a cell diameter of 25 cm). In contrast, the MPI and PSU instruments feature flow-tube-like inlets (14–17 cm from the pinhole to laser axis) mounted on smaller fluorescence cells, facilitating the interaction of sampled gas with the cell walls, which may promote the generation of internal OH. For the measurements listed in Table 4, the Peking University (PKU) instrument (Tan et al., 2017) is most similar to the Leeds FAGE (i.e. single-pass detection, ~ 10 cm total length from sampling inlet to laser axis), for which similar daytime interferences on the order of ~ 0–20% were observed.

For the ICOZA campaign, nothing could be inferred about the origin of the OH interference signal when one was observed, as it did not exhibit any characteristic diurnal profile (Fig. 11), and showed no obvious dependences on a variety of meteorological and chemical parameters. This finding is in contrast to previous studies in which diel profiles (Mao et al., 2012; Feiner et al., 2016) and dependences (Mao et al., 2012; Feiner et al., 2016; Novelli et al., 2017) of the interference have been observed. The occurrence of large (i.e. > 1 × 10⁶ molec. cm⁻³) background OH signals (OH_{int} = OH_{wave} − OH_{chem}) after instrumental problems (e.g. power cuts, data not shown) implies that the differences may have been instrumental rather than as a result of a species present in ambient air, although the data at these times did pass all quality control filters and therefore could not be rejected. Nonetheless, any differences are still a concern, regardless of their cause; the IPI system thus serves as an additional check on measurement accuracy and operational stability, and is perhaps most useful for fieldwork sites where power supplies are unreliable, for example in more remote areas.

It is possible that, even though the background OH had a flat diurnal profile in each field campaign, the species responsible for any interference observed were different between daytime and night-time periods. Thus, analysis of the daytime and night-time data separately, as a function of the same parameters, might reveal more information. Considering the recent identification of NO₃ radicals as an internal

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Figure 16. The relationship between OH_{int} and OH_{chem} using binned data. The error bars for AIRPRO summer (a, blue markers) denote 1 SD and are not shown for AIRPRO winter and ICOZA for clarity. The dashed black line corresponds to a sigmoid fit and is used to guide the eye only. The contribution of interferences to the total OH_{wave} signal (OH_{int} / OH_{wave} × 100%) for the AIRPRO summer campaign is shown in (b), with the number of points in each bin shown in (c). All OH_{int} data used here have been corrected for the known interference from O₃ + H₂O.
OH source in LIF instruments (Fuchs et al., 2016), and that OH concentrations have often been under-predicted at night (Faloona et al., 2001; Mao et al., 2012; Ren et al., 2013; Hens et al., 2014; Lu et al., 2014; Tan et al., 2017), this is perhaps the most interesting period for further study. However, for the data presented in this work, robust quantitative night-time analyses are not possible due to OH measurements being below or close to the instrument LOD.

In this work, there are several key findings that stand out. First, OHwave and OHchem were in good agreement even at the very low NO concentrations of < 100 pptv during ICOZA and the moderate afternoon levels (∼500 pptv on average) but often <100 pptv; Shi et al., 2019) during the AIRPRO summer campaign. While the role of isoprene could not be assessed for ICOZA, due to the limitation range of concentrations observed (<0.2 pptv), it reached high levels during AIRPRO summer (up to 7.9 ppbv, larger than seen in some forested regions) but did not seem to perturb the agreement between the two measurements. In addition, very high levels of aromatic VOCs were observed during both AIRPRO winter and summer, where the agreement between OHwave and OHchem suggests that the intermediates of aromatic oxidation, such as exotic bicyclic species (Birdsall et al., 2010) and highly oxygenated molecules (HOMs) (Wang et al., 2017; Molteni et al., 2018; Tsiligiannis et al., 2019), do not give rise to OH interferences, which is postulated to be the case for intermediates (SCIs) in the ozone-oxidation of alkenes (Novelli et al., 2014a, 2017; Rickly and Stevens, 2018). However, the large alkene and ozone concentrations observed during AIRPRO summer should favour the formation of these SCIs, but significant interferences were not observed, consistent with laboratory investigations of the isoprene interference and casting doubt on the SCI hypothesis. Although, the AIRPRO SCI concentrations also depend on the magnitude of the SCI loss rates, which could be high if elevated levels of SO2 (Welz et al., 2012; Sheps et al., 2014) or organic acids (Welz et al., 2014) were present.

Considering the success of the first three field deployments of the IPI system and given that it does not reduce the instrument sensitivity towards OH, it is suggested that the system is adopted for permanent use in ambient studies, although conventional sampling should still be performed from time-to-time to check for potential artefacts caused by the IPI system itself. Another advantage of the IPI system is that it reduces the amount of solar light entering the pinhole, which reduces the size and variability of daytime background signals and therefore improves signal-to-noise and hence detection limits. It is recommended that the IPI propane concentration is kept the same as the summer AIRPRO campaign, as it is possible that the slightly poorer agreement between OHwave and OHchem during ICOZA was because of the lower propane flow used (i.e. the flow was not sufficient to ensure that OH generated from all steady-state sources was removed), although this cannot be verified.

Future field campaigns using the IPI will allow for the assessment of interferences in the Leeds FAGE instrument for a range of different environments. From these, the contribution of interferences for previous studies in similar environments, where measurements were made prior to the discovery of significant interferences in the LIF measurement of OH reported.
by others, may be inferred. The measurement–model comparisons may then be reassessed in light of any new information regarding the accuracy of OH measurements. Regardless of the reasons for any differences between the two measures of OH (i.e. chemical interferences or instrumental problems such as during recovery periods after power cuts), the IPI system serves as an additional check on OH observations, increasing confidence in the validity of the data obtained.

5 Conclusions

The addition of an IPI system to the Leeds ground-based FAGE instrument allowed for a comprehensive investigation of OH measurement interferences in both the laboratory and the field. Following its optimisation and thorough characterisation in terms of sensitivity and external and internal OH removal efficiency, laboratory experiments were conducted to assess potential interferences from (1) the photolysis of O₃ in the presence of H₂O vapour, (2) the intermediates and products of isoprene ozonolysis, and (3) NO₃ radicals. For O₃ + H₂O, a small but potentially significant interference (at high O₃ levels) was found, but interferences from isoprene ozonolysis products and NO₃ radicals were shown to be insignificant under typical atmospheric conditions.

Field campaigns conducted in the UK and China showed that, on average, the Leeds ground-based FAGE instrument does not suffer from significant interferences in the detection of OH. It was only under the very high OH levels of > 1.5 × 10⁶ molec. cm⁻³ sometimes observed during the AIRPRO summer campaign that interferences were found consistently, although their contributions (~ 15%–20%) were smaller than the instrumental accuracy of 26% at 2σ. Large interference signals (> 1 × 10⁶ molec. cm⁻³) were occasionally observed during the ICOZA campaign but always after instrumental problems such as power cuts, suggesting that the OHchem method serves as an additional tool for verifying instrument stability and validating measurements. The Leeds IPI system will find continued use in future fieldwork.

Data availability. Data presented in this study are available from the authors upon request (l.k.whalley@leeds.ac.uk and d.e.heard@leeds.ac.uk).

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Author contributions. RWM, TI, LKW, and DEH designed the IPI system. RWM, EJS, JA, LMS, CY, and LKW performed laboratory experiments. RWM, EJS, DRC, CY, LKW, and DEH were responsible for field measurements. RWM, EJS, and LKW analysed the data. RWM wrote the manuscript with input from all co-authors.

Competing interests. The authors declare that they have no conflict of interest.

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