Comparison of temperature dependent calibration methods of an instrument to measure OH and HO$_2$ radicals using laser-induced fluorescence spectroscopy

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Abstract

Laser Induced Fluorescence (LIF) spectroscopy has been widely applied to fieldwork measurements of OH radicals, and of HO$_2$, following conversion to OH, over a wide variety of conditions, on different platforms, and in simulation chambers. Conventional calibration of HO$_x$ (OH + HO$_2$) instruments has mainly relied on a single method, generating known concentrations of HO$_x$ from H$_2$O vapour photolysis in a flow of zero air impinging just outside the sample inlet ($S_{HOx} = C_{HOx}$[HO$_x$], where $S_{HOx}$ is the observed signal and $C_{HOx}$ is the calibration factor). The FAGE (Fluorescence Assay by Gaseous Expansion) apparatus designed for HO$_x$ measurements in the Highly Instrumented Reactor for Atmospheric Chemistry (HIRAC) at the University of Leeds has been used to examine the sensitivity of FAGE to external gas temperatures (266 – 348 K).

The conventional calibration methods give the temperature dependence of $C_{OH}$ (relative to the value at 293 K) of $(0.0059 \pm 0.0015) \text{ K}^{-1}$ and $C_{HO2}$ of $(0.014 \pm 0.013) \text{ K}^{-1}$. Errors are 2σ. $C_{OH}$ was also determined by observing the decay of hydrocarbons (typically cyclohexane) caused by OH reactions giving $C_{OH}$ (again, relative to the value at 293 K) of $(0.0038 \pm 0.0007) \text{ K}^{-1}$. Additionally, $C_{HO2}$ was determined based on the second order kinetics of HO$_2$ recombination with the temperature dependence of $C_{HO2}$, relative to 293 K being $(0.0064 \pm 0.0034) \text{ K}^{-1}$. 
The temperature dependence of $C_{\text{HOx}}$ depends on HOx number density, quenching, relative population of the probed OH rotational level and HOx transmission from inlet to detection axis. The first three terms can be calculated and, in combination with the measured values of $C_{\text{HOx}}$, show that HOx transmission increases with temperature. Comparisons with other instruments and the implications of this work are discussed.

1 Introduction

Hydroxyl radicals (OH) play a key role in our atmosphere, oxidising a broad range of species. OH is the main daytime oxidant in the troposphere and the main sink for methane, a potent greenhouse gas. The OH radical is linked to the HO$_2$ radical through the oxidation of most other non-methane hydrocarbons (NMHCs) and CO in the troposphere and, through reaction with NO$_2$, in the upper troposphere/lower stratosphere. Due to the high reactivity of OH (lifetime ~1 s even in clean air), these radicals undergo minimal transport and local concentrations depend only on the in situ chemistry. Measurements of HOx concentrations, in conjunction with measurements of their sources and sinks are a sensitive test of chemical models. Accurate measurement of [HOx] is therefore paramount, not only for field measurements, (Stone et al., 2012; Heard and Pilling, 2003; Gligorovski et al., 2015), but also for atmospheric simulation chambers where OH/HO$_2$ instruments have been deployed (Karl et al., 2004; Glowacki et al., 2007).

Sensitive detection techniques with high temporal resolution are required for HOx detection and techniques have been reviewed in Stone et al. (2012) and Wang et al. (2021). Fluorescence Assay by Gaseous Expansion (FAGE) (e.g. Hard et al. (1984)) is the most common method used for both field and chamber studies. Here, the sample is expanded to low pressures and OH detected by resonance fluorescence at ~308 nm. The low pressures are required to temporally separate fluorescence from the excitation laser pulse. HO$_2$ is converted to OH by reaction with NO and detected in a separate cell. Both techniques require calibration which is conventionally based on the generation of OH and HO$_2$ from water vapour photolysis at 185 nm at atmospheric temperature and pressure.

Recent studies have demonstrated potential interferences for measurements of both OH and HO$_2$ radicals using the FAGE technique, with the magnitude dependent upon instrument design (Mao et al., 2012; Novelli et al., 2014; Novelli et al., 2017; Fuchs et al., 2011; Whalley et al., 2013; Fuchs et al., 2016; Fittschen et al., 2019). Considerable effort has been made to minimize,
understand and mitigate any interference, with many groups now fitting an external OH scavenger injector to measure OH concentrations using an alternative background signal, \( \text{OH}_{\text{CHEM}} \), alongside the conventional method of measuring OH using a background signal determined by tuning the laser wavelength off-resonant to the transition, \( \text{OH}_{\text{WAVE}} \) (Woodward-Massey et al., 2020; Novelli et al., 2014; Mao et al., 2012). Intercomparison campaigns (e.g. Schlosser et al. (2009), Onel et al. (2017a)) in the controlled environment of an atmospheric chamber are useful to identify systematic errors in different approaches, but if both methods require calibration, the accuracy of the measurements is still compromised by uncertainties in the calibration methods.

In an earlier paper (Winiberg et al., 2015), accurate calibration of a FAGE instrument over a range of external inlet pressures (440 – 1000 mbar) was performed in the Leeds HIRAC (Highly Instrumented Reactor for Atmospheric Chemistry (Glowacki et al., 2007)) chamber. The instrument sensitivity to OH and HO\(_2\) agreed well for the conventional water vapour calibration method (where the external pressure is always 1 bar, and external pressure effects were simulated by altering the pressure in the FAGE detection cell) and alternative methods based on the temporal decay of a hydrocarbon (for OH) or the temporal decay of HO\(_2\) via its second-order self-reaction (for HO\(_2\)) over an external pressure range of 300 – 1000 mbar. For OH, the calibration factor, \( C_{\text{OH}} \), (where \( S_{\text{HOx}} = C_{\text{HOx}}[\text{HOx}] \) and \( S_{\text{HOx}} \) is the FAGE signal) increased by 17% and for HO\(_2\) a slightly greater increase in \( C_{\text{HO2}} \) of 32% was determined as the pressure increased from 350 to 1000 mbar. There was good agreement between the absolute values and their pressure dependence for both calibration methods. Such comparisons are particularly relevant to aircraft operation where external pressures will vary considerably during the flight or for evacuable chambers such as the Leeds HIRAC chamber which can operate from 50 – 1000 mbar. Marno et al. (2020) have also developed the All Pressure Altitude-based Calibrator of HOx Experimentation (APACHE) to allow calibration of their FAGE instrument HORUS (HydrOxyl Radical measurement Unit based on fluorescence Spectroscopy) as a function of pressure, but not temperature.

Little is known on the effect of gas temperature at the inlet upon instrument sensitivity for LIF instruments, despite field instruments being used at extremes of temperature, from day to night, from deserts to the polar regions, and in aircraft, where temperatures change rapidly with altitude. Additionally, ambient conditions influence not only the inlet temperature, but the whole apparatus. For example in the FAGE system associated with HIRAC, based on a design for aircraft use (Commane et al., 2010), the whole inlet tube (~30 cm) is located inside the
HIRAC chamber and so wall loss rates of HOx in the inlet tube will be influenced by the
temperature of the HIRAC chamber. The long inlet is required either to locate the pinhole
outside of the aircraft for the airborne instrument, or to allow sampling across the diameter of
the HIRAC chamber. To date, the only study investigating the effect of inlet temperature on
instrument sensitivity to HOx radicals has been performed by Regelin et al. (2013), who
reported a minor positive dependence of the OH sensitivity (C_{OH}) as a function of decreasing
inlet temperature for the HORUS instrument (possibly due to a cooling effect on the
instrumentation). There was a more marked decrease in the instrument sensitivity to HO2 with
decreasing temperature, most probably due to enhanced wall losses at lower temperatures.

In this paper, instrument sensitivity as a function of external inlet temperature has been
determined for the HIRAC FAGE instrument for both OH and HO2, using the water vapour
photolysis calibration method in an external flowtube (termed ‘conventional method’) and
alternative calibration methods using chemical reactions in the HIRAC chamber (Winiberg et
al., 2015) at varying temperatures. Alternative OH calibrations used the inferred [OH] from the
measured decay of a hydrocarbon (HC), typically cyclohexane, reacting with OH (R1) (termed
‘HC decay method’). The rate of loss of HC is then given by equation (E1).

\[ \text{OH} + \text{HC} \rightarrow \text{products} \quad (R1) \]
\[ \frac{-d[\text{HC}]}{dt} = k_{ba}[\text{OH}][\text{HC}] \quad (E1) \]

In E(1), \( k_{ba} \) is the well-established literature value for the bimolecular rate coefficient between
OH and the monitored hydrocarbon and \( \frac{-d[\text{HC}]}{dt} \) can be measured from the HC time series so
that [OH] is the only unknown parameter and can be calculated and compared with the [OH]
predicted via the conventional calibration method.

HO2 was also calibrated by monitoring the HO2 kinetic decay during the recombination
following generation by HCHO photolysis in the presence of O2 (termed ‘HO2 self-reaction
method’).

\[ \text{HCHO} + \text{hv} \rightarrow \text{H} + \text{HCO} \quad (R2) \]
\[ \text{HCO} + \text{O}_2 \rightarrow \text{HO}_2 + \text{CO} \quad (R3) \]
\[ \text{H} + \text{O}_2 + \text{M} \rightarrow \text{HO}_2 + \text{M} \quad (R4) \]
\[ \text{HO}_2 + \text{HO}_2 (+\text{M}) \rightarrow \text{H}_2\text{O}_2 + \text{O}_2 (+\text{M}) \quad (R5) \]
The time dependence of the $[\text{HO}_2]$ in the second-order decay depends on the initial concentration of $\text{HO}_2$ allowing for calibration.

2 Experimental

2.1 The HIRAC chamber

The alternative calibration methods of monitoring hydrocarbon or $\text{HO}_2$ decays were conducted in HIRAC using very similar methods and conditions as described in Winiberg et al. (2015). HIRAC is a stainless steel chamber with a total volume of 2.25 m$^3$ and can operate over a wide range of pressures (50 – 1000 mbar) and temperatures (227 – 343 K). Multiple access ports are available to connect an array of instrumentation and monitoring equipment (pressure gauges, thermocouples etc.). The chamber has been described previously in detail in Glowacki et al. (2007), Malkin et al. (2010) and (Bejan et al., 2018). More recently a temperature control system was installed to further enhance the capabilities of the HIRAC chamber (Section 2.1.1).

Details on the temperature characteristics of HIRAC can be found in Section S1 of the SI.

The photolysis lamps, housed in eight quartz tubes mounted radially inside the reactive volume, were used to initiate photochemistry. The lamps were interchangeable depending on the target molecules; lamps, with primary emissions centred at 254 and 310 nm (GE Optica, GE55T8/HO and Philips, TL40W/12 RS respectively), were used for the alternative OH and $\text{HO}_2$ calibration methods respectively (sections 3.2 and 3.3). The housings were flushed with dry N$_2$ (~3 slm per housing) to help regulate the temperature and remove photolabile species and water, which could condense or freeze around the lamps at lower temperatures. A photolysis lamp induced chamber temperature increase of $\sim$2 – 5 K was seen over the course of a typical experiment (<40 mins), but this variation was reduced if the chamber was temperature controlled. Temperatures were monitored using a series of K-type thermocouples inside the lamp housings (one per lamp) as well as distributed around the inside of the chamber. Thermocouples were placed strategically to allow the temperature to be measured close to the chamber walls, inlets, flanges and in the chamber.

2.1.1 Temperature Control System

During manufacture, square cross section steel tubing (volume $\sim$50 L) was welded directly to the outer skin of HIRAC, allowing a cooling/heating liquid to flow around the chamber,
controlling the temperature inside. The square tubing enabled the temperature control liquid to transfer heat more efficiently to the chamber by offering a larger contact surface area compared to cylindrical tubing. A Huber thermostat unit (model 690W) was used to circulate ~60 L of thermofluid (Huber DW-THERM, 183 - 473 K) around the chamber. Further details are given in the SI (Section S1).

HIRAC was able to sustain a steady temperature (±2 K) across the chamber at any temperature between 227 and 343 K and example temperature profiles are given in the SI. A negligible temperature gradient (<0.5 K, see Figure S2) was observed across the central portion of the chamber, in both the horizontal and vertical axes. Close to the walls of the chamber, however, a change of ~1 K was observed. The flanges around the HIRAC chamber were insulated with ~40 mm of neoprene, however there was no direct temperature control of the flanges or access ports, which was likely responsible for the change in temperature at the large 600 mm access flanges.

2.1.2 HO\textsubscript{x} Instrumentation

The OH and HO\textsubscript{2} radicals were detected using a FAGE instrument based in the HIRAC chamber with a 5 kHz pulse repetition frequency (PRF) laser light source, as described in Winiberg et al. (2015); Winiberg et al. (2016) and Glowacki et al. (2007). Air was sampled at ~6 slm through a 1.0 mm diameter pinhole nozzle and passed down the inlet (length 280 mm, 50 mm diameter) into the OH detection axis maintained at low pressure (typically ~3.85 mbar) using a high-capacity rotary-backed roots blower pumping system (Leybold, Trivac D40B and Ruvac WAU251). The long inlet was used to draw a sample away from the chamber walls where radical losses increase (a maximum of 15% decrease at <10 mm from the chamber wall) and to probe any radical gradients occurring due to spatially inhomogeneous production (Winiberg et al., 2015). The FAGE instrument was coupled to the HIRAC chamber using ISO-K160 flanges, ensuring the pinhole is kept >200 mm from the chamber walls.

Concentrations of HO\textsubscript{2} were measured simultaneously in a second detection axis ~300 mm downstream of the OH detection axis. High purity NO (BOC, N2.5 Nitric Oxide) was added ~20 mm before the HO\textsubscript{2} detection axis into the centre of the FAGE cell in the direction of gas flow through 1/8” stainless steel tubing at a rate of 5 sccm (Brooks 5850S) converting HO\textsubscript{2} to OH. Conversion of some types of RO\textsubscript{2} radicals (in particular β-hydroxyperoxy radicals) to OH upon reaction with NO has been reported in other FAGE instruments (Whalley et al., 2013; Fuchs et al., 2011). However, during the alternative HO\textsubscript{2} calibrations (based on HCHO
photolysis) presented here no β-hydroxyperoxy radicals were generated hence any interference was assumed to be negligible.

A JDSU Nd:YAG pumped Sirah Cobra Stretch system (PRF = 5 kHz) was used to generate the frequency doubled ~308 nm (307.99 nm to excite the Q_1(2) rotational state) light for the fluorescence of OH radicals. Light was directed from the output of the laser and focussed into fibre optic cables (10 m, Oz Optics) which were then attached directly to the FAGE cell arms via collimators (Oz Optics). Fluctuations in laser power were accounted for using a linear response UV sensitive photodiode (UDT-555UV, Laser Components UK) at the exit arm of the OH and HO_2 detection axes to normalise the LIF signal. The laser system provided between 5 – 7 and 2 – 3 mW of 308 nm light to the OH and HO_2 detection axes, respectively.

The OH fluorescence was collected orthogonal to the gas flow onto electronically gated Channeltron PhotoMultiplier tubes (CPM, Perkin Elmer, C943P) via a series of imaging lenses and a narrow bandpass filter (Barr Associates, 308.8 ± 5.0 nm). A spherical concave back reflector was positioned underneath the cell, opposite the detection optics, to optimise light collection onto the CPM. To avoid detector saturation, the CPM was gated (i.e. switched off) for the duration of the laser pulse using a modified gating unit based on the original design by Creasey et al. (1997a). Signals from the CPM were analysed using PC-based photon counting cards (Becker and Hickl PMS-400A).

2.1.3 Other instrumentation

As with the previously published work (Winiberg et al., 2015), a chemiluminescence NO_x analyser (TEC 42C, limit of detection = 50 pptv at 60 s averaging) was used to determine that levels of NO_x (NO + NO_2) in the HIRAC chamber were typically below the detection limit of the apparatus.

Most of the OH calibration experiments using the hydrocarbon decay method were performed monitoring HC decays using a chemical ionization time of flight mass spectrometer (Kore custom build) operating with N_2^+ ionization. Gas was sampled from HIRAC via ~7 m of 1/8" Teflon tubing with the inlet being located close (within 70 cm) to the FAGE inlet. A majority of the experiments were carried out with cyclohexane as the HC (monitored at m/z = 84.15), although other compounds were used. The mass spectrometer signal was calibrated by introducing known HC concentrations into HIRAC. An example of the resulting calibration plot can be found in the SI (Section S2, Figure S3).
2.2 General Chamber preparation

Calibration experiments were conducted at 1000 mbar in an Ultra-High Purity (UHP) 1:4 synthetic air mix of O₂ (BOC, zero-grade, >99.999%) and N₂ (BOC, zero-grade, >99.998%) to match the range of pressures from the water vapour calibration method (section 3.1). Thorough mixing of reaction mixtures within HIRAC was achieved in ≤70 s by four circulation fans mounted in pairs at each end of the chamber. The chamber was evacuated to ~0.05 mbar for ~60 - 120 min following each experiment using the rotary pump backed roots blower to ensure removal of all reactants/products. The combined sampling rate of ~9 slm from the chamber required a counter flow of synthetic air to maintain the desired pressure and resulted in a first order dilution term of \((4.5 \pm 0.2) \times 10^{-5} \text{ s}^{-1}\). The dilution flow was regulated using two Brooks mass flow controllers (N₂ and O₂) and the dilution was taken in account in all analyses.

2.3 Chemical reagents

Known concentrations of precursors (except H₂O₂) and reagents were introduced to the chamber in the vapour phase through a 0.97 L stainless steel delivery vessel. Hydrogen peroxide (50% wt solution, Merck, used as supplied) was directly injected via a syringe. Multiple injections could be made in each run to ensure a wide range of \([\text{OH}]\) was covered.

For the hydrocarbon based OH calibration method, cyclohexane (99%, Fischer Scientific), methylcyclohexane (>99.9%, Sigma Aldrich) and heptane (99%, Fischer Scientific) were purified using freeze-pump-thaw cycles before being introduced into the HIRAC chamber.

For the second-order HO₂ calibration method, formaldehyde (HCHO) was produced in the gas phase by gently heating paraformaldehyde (99.9%, Sigma Aldrich) into the evacuated delivery vessel. This method was sufficient for producing the 2 – 3 ppmv concentrations of HCHO in the HIRAC chamber that were required.

3 Calibration methods

3.1 Flowtube/Water Photolysis Calibration Method

The flowtube calibration method relies on the photolysis of H₂O vapour at 184.9 nm in a fast flow (40 slm) of synthetic air. A mercury penray lamp (LOT-Oriel, Hg-Ar) was used as the photolysis source, placed at the end of a square cross section flow tube (12.7 × 12.7 × 300 mm). Air was humidified by passing a fraction of the bulk air flow through a bubbler containing
deionised water. The $[\text{H}_2\text{O}]$ was measured using a dew-point hygrometer (CR4, Buck Research Instrument) prior to the flow tube and the resulting OH and HO$_2$ concentrations from photolysis can be calculated from equation (E2):

$$[\text{OH}] = [\text{HO}_2] = [\text{H}_2\text{O}] \sigma_{\text{H}_2\text{O}, 184.9 \text{ nm}} \Phi_{\text{OH}} F_{184.9 \text{ nm}} \Delta t$$  \hspace{1cm} (E2)

where $\sigma_{\text{H}_2\text{O}, 184.9 \text{ nm}}$ is the known absorption cross-section of H$_2$O vapour at 184.9 nm $((7.22 \pm 0.22) \times 10^{-20} \text{ cm}^2 \text{ molecule}^{-1}$ (Cantrell et al., 1997; Creasey et al., 2000; Hofzumahaus et al., 1997)), $\Phi_{\text{OH}}$ $(= \Phi_{\text{HO}_2} = 1)$ is the photodissociation quantum yield of OH and HO$_2$ (Fuchs et al., 2011), $F_{184.9 \text{ nm}}$ is the photon flux of 184.9 nm light and $\Delta t$ is the exposure time of the air to the Hg lamp output. The exposure time of the air to the 184.9 nm light, $\Delta t$, was calculated as a function of the known velocity of the air and the cross section of the photolysis region.

The product $F_{184.9 \text{ nm}} \times \Delta t$ was determined for lamp supply currents between 0.2 and 3.0 mA using the N$_2$O actinometry method described in detail in a number of publications (Edwards et al., 2003; Heard and Pilling, 2003; Faloona et al., 2004; Whalley et al., 2007; Glowacki et al., 2007).

The gas output from the flow tube was directed towards the FAGE sampling inlet, where the overfill of the FAGE sample volume from the flow tube stopped the impingement of ambient air. A range of HO$_x$ concentrations ($10^8 - 10^{10} \text{ molecule cm}^{-3}$) were produced by changing the mercury lamp photon flux whilst keeping a constant [H$_2$O] (typically 2000 - 3000 ppmv). The average calculated [HO$_x$] values are compared to their concurrent OH/HO$_2$ signals observed during the same time period, the linear regression of which gives the instrument sensitivity to OH/HO$_2$. A typical calibration plot is shown in Figure 1. Potential systematic errors in the flowtube calibration method have been discussed previously (Winiberg et al. 2015) and are summarized for the current instrument in Table 4 and discussed further in the SI, Section S3, which also contains a schematic of the flowtube calibration apparatus (Figure S4).
Figure 1: Typical room temperature calibration plot from the conventional water photolysis, flow tube method. The total flow rate was 40 slm, with [H\textsubscript{2}O] = 1600 ppmv, the laser power was 9.65 mW and the OH cell was at a pressure of 2.6 Torr. Gradient = (1.266 ± 0.034) × 10\textsuperscript{-8} counts s\textsuperscript{-1} mW\textsuperscript{-1} cm\textsuperscript{3} molecule\textsuperscript{-1}, intercept = 0.28 ± 0.74 counts s\textsuperscript{-1} mW\textsuperscript{-1}. Errors are 2σ.

3.1.1 Calibration for External Inlet Temperature

The FAGE inlet was wrapped with ¼” copper tubing (~ 5 cm between coils) and covered in two layers of aluminium foil to aid thermal contact. A final layer of 10 mm thick neoprene was added to the outside of the foil to aid insulation. The Huber temperature control unit was used to flow DW-THERM thermofluid through the tubing to vary the temperature of the inlet. Temperatures were monitored externally using three K-type thermocouples; two positioned on the inlet and one on the conical pinhole nozzle during the calibration procedure (see Figure 2(a)).

Calibrations were conducted at five external inlet temperatures from 263 – 343 K, representative of the operating temperature range for the HIRAC chamber. During the bulk of the experiments, gases from the flowtube calibration source were maintained at room temperature. However, an additional range of calibration experiments were performed with flowtube gas maintained to within ±5 K of the measured external inlet temperature. This effect was achieved by passing the humidified bulk flow through a 2 m long coil of ¼” copper tubing held at the desired set point using a thermostat controlled water bath (Thermo Fischer Science).
The $[\text{H}_2\text{O}]_{\text{vap}}$ was determined just before the calibration flowtube, with the temperature monitored both before and at the exit of the flowtube. Short gas lines were used between the water bath and the flow tube, which was covered in a thin layer of neoprene to insulate and reduce temperature gradients.

(a)

(b)

(c)

**Figure 2:** (a) Schematic of FAGE Cell showing locations of thermocouples. (b) Internal cell temperatures ($T_{\text{OH}}$ or $T_{\text{HO}_2}$) and inlet temperatures ($T_{\text{inlet}}$) plotted as a function of the external temperature ($T_{\text{ext}}$), when sampling air at 293 K from the calibration flowtube. Slope $T_{\text{inlet}} = 0.558 \pm 0.010$; Slope $T_{\text{OH}} = 0.497 \pm 0.008$; Slope $T_{\text{HO}_2} = 0.236 \pm 0.033$. (c) Internal temperatures as a function of the external temperature when sampling temperature controlled air from the calibration flowtube. Slope $T_{\text{OH}} = 0.890 \pm 0.004$; Slope $T_{\text{HO}_2} = 0.316 \pm 0.007$ (sampling from the HIRAC chamber gave lines with essentially the same gradients).
Prior to the calibration, the internal cell temperatures were measured using three K-type thermocouples positioned in the centre of the gas flow inside the inlet (just after the inlet pinhole), OH and HO₂ fluorescence cells, details of which are discussed in the results section (4.1.1). The thermocouples were inserted into the cell using a ¼” compression fitting port, seal; this allowed the cell to be operated at normal operating pressure during the temperature profile measurements. Thermocouples were held in place temporarily using electrical tape, and OH/HO₂ calibrations were not performed with the thermocouples in place.

3.2 Hydrocarbon decay method

A majority of the hydrocarbon decay OH measurements were made with cyclohexane as the monitored hydrocarbon (HC) (monitored via the m/z = 84.15 peak) and hydrogen peroxide photolysis at 254 nm as the OH source.

The principle of the hydrocarbon decay method was outlined in the introduction; the rate of loss of the HC by OH is given by:

$$-\frac{d[HC]}{dt} = k_{bi}[OH][HC]$$  \hspace{1cm} (E1)

The rate coefficient for cyclohexane, c-C₆H₁₂, has received much attention in the literature over the 273 – 343 K temperature range used in this study, and so we use the IUPAC recommended rate expression (Atkinson et al., 2006):

$$k_{OHc-C6H12} = 3.26 \times 10^{-17} T^2 \exp\left((-262\pm66)/T\right) \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$$  \hspace{1cm} (E3)

The calculated [OH] from the hydrocarbon decay can be compared to the corresponding FAGE signal, corrected for the difference in [H₂O] used in the calibration and that present in the HIRAC chamber, to determine the C_OH. In practice, the total HC decay is a combination of reaction with OH and other first order loss processes, primarily dilution (as sampled gas is replenished with air). Therefore

$$-\frac{d[HC]}{dt} = k_{1st}[HC] + k_{bi}[OH][HC]$$  \hspace{1cm} (E4)

where $k_{1st}$ represents the rate coefficient for the sum of all non-OH first order loss processes (e.g. heterogeneous loss and dilution). Gradients were obtained from analysis within the Origin software package. A second order polynomial was fitted to 10 – 40 points (with the separation
of each point being 10 s); the number of points depending on the rate of change of the [HC] and the data points were smoothed via the method of Savitzky and Golay (1964).

$k_{1s}$ was determined from the HC decays in the absence of OH (either with no lamps on, or no OH precursor present). For each injection of HC (typical initial concentration of $3 - 5 \times 10^{13}$ molecule cm$^{-3}$) there were multiple H$_2$O$_2$ injections (~1 ml). FAGE measurements were typically averaged over 30 s (30 data points, with each data point corresponding to accumulated signal over ~1 s) to counteract the noise arising in fluorescence counts. During rapid changes in the observed signal, for example immediately after initial photolysis of hydrogen peroxide in the chamber (see Figure 3(a)), a reduced averaging period was used. The HIRAC FAGE system shows a slight sensitivity to water vapour concentrations due to quenching (Winiberg, 2014). Minor corrections (<5%) were made to account for the different water vapour concentrations in the two calibration methods.

Figure 3(a) shows a typical time series of OH with the black line giving the [OH] derived from the mass spectrometer measurements and the brown line giving [OH] derived from the FAGE signal and converted to [OH] using the conventional flow tube water vapour photolysis calibration at 293 K. Figure 3(b) shows the resulting scatter plot. The slope of the scatter plot gives the correction to be applied to $C_{293K}$ from the conventional calibration to match the [OH] derived from the mass spectrometric measurements.

**Figure 3:** a) Time series of [OH] derived from FAGE measurements and from mass spectrometric measurements of cyclohexane removal recorded following H$_2$O$_2$ photolysis at 293 K and 1000 mbar air. The error bars shown represent absolute uncertainties in the calibration methods, see Table 4. b) Resultant scatter plot where the gradient, $0.998 \pm 0.016$ (2σ) gives $C_{rel}$ for the FAGE apparatus at 293 K for this experiment. The average gradient at 293 K is $1.034 \pm 0.0068$ from five experiments.
3.3 Calibration of HO₂ detection via HO₂ recombination kinetics

The HCHO photolysis/HO₂ recombination kinetics method of HO₂ cell calibration was used as described in Winiberg et al. (2015). Formaldehyde was introduced in a flow of nitrogen into the chamber (containing synthetic air at 1000 mbar) at concentrations of ~2 x 10¹³ molecule cm⁻³. The chamber was irradiated (lamps: Philips TL40W/12 RS) resulting in an almost instantaneous HO₂ signal (reactions R2 – R4). Once a steady state HO₂ concentration was achieved, the photolysis lamps were turned off and the decay of HO₂ was monitored by FAGE for ~120 s (Figure 4). The decay of HO₂ was primarily controlled by the self-reaction (R5), but there was a small first-order contribution from loss to the walls (R6).

The measurement of HO₂ decays was repeated up to six times before the laser wavelength was scanned to the offline position.

\[
\text{HO}_2 + \text{HO}_2 (+\text{M}) \rightarrow \text{H}_2\text{O}_2 + \text{O}_2 (+\text{M}) \quad \text{(R5)}
\]

\[
\text{HO}_2 \rightarrow \text{loss} (k_{\text{loss}}) \quad \text{(R6)}
\]

The chamber mixing fans were used for the first three calibration decays, representative of a typical experimental homogeneous gas mixture. The second series of three calibration decays were conducted without the mixing fans to probe the HO₂ recombination and wall loss kinetics in the absence of effective mixing.

When the fans are on, the loss of HO₂ was characterised by bimolecular self-reactions and a first order wall loss parameter. The solution to this mixed order decay is given by:

\[
(S_{\text{HO}_2})_t = \left( \frac{1}{(S_{\text{HO}_2})_0} + \frac{2 \cdot k_{\text{RO}_2+\text{HO}_2}}{k_{\text{loss}} C_{\text{HO}_2}} \right) \cdot e^{(k_{\text{loss}}) \cdot \left( \frac{2 \cdot k_{\text{RO}_2+\text{HO}_2}}{k_{\text{loss}} C_{\text{HO}_2}} \right) \cdot t} \quad \text{(E5)}
\]

where \((S_{\text{HO}_2})_t\) and \((S_{\text{HO}_2})_0\) are the HO₂ signal at time \(t\) and \(t = 0\) respectively, \((C_{\text{HO}_2})\) is the instrument sensitivity, \(k_{\text{RO}_2+\text{HO}_2}\) is the HO₂ recombination rate coefficient and \(k_{\text{loss}}\) represents the wall loss parameter. Both \(k_{\text{loss}}\) and \(C_{\text{HO}_2}\) were determined by data fitting the \(S_{\text{HO}_2}\) decay using equation (E5) with a Levenburg-Marquardt non-linear least squares algorithm, fixing the initial signal and \(k_{\text{RO}_2+\text{HO}_2}\). The first ~100 s of data were used, ensuring analysis after an almost complete decay of \(S_{\text{HO}_2}\). Figure 4 shows an example of a typical decay and the resulting fit to equation (E5).

For the experimental temperature range (275 – 345 K), \(k_{\text{RO}_2+\text{HO}_2}\) has values between (2.00 – 2.85) x 10⁻¹² cm³ molecule⁻¹ s⁻¹ according to the recommendation given by IUPAC.
The chamber was operated under dry conditions (< 10 ppmv [H2O]vap), and so the enhancement of $k_{\text{HO}_2^+ \text{HO}_2}$ by formation of a pre-reactive complex with H2O was ignored for these analyses. The wall loss rate, $k_{\text{loss}}$, was dependent on daily chamber conditions and was therefore determined as part of the fitting procedure along with $C_{\text{HO}_2}$, typically between 0.032 – 0.073 s\(^{-1}\) with an uncertainty of ±10 % (2σ). Without the fans, the value of $k_{\text{loss}}$ was reduced, but agreement between the HO\(_2\) calibration methods was comparable (within 10%). As HIRAC is generally operated with fans on, we have only reported these data. Wall loss typically contributes 10 – 50% of the initial decay but is well defined in the fitting procedure. As with OH detection, minor corrections have been made for the slightly different sensitivities of the system under the different water concentrations of the two calibration methods (Winiberg, 2014).

**Figure 4:** Typical HO\(_2\) decay recorded at 293 K and 1000 mbar air. The red line is the fit to the data from equation (E5) giving $C_{\text{HO}_2, 293 \text{ K}} = (4.17 \pm 1.66) \times 10^{-8}$ counts cm\(^3\) molecule\(^{-1}\) mW\(^{-1}\) s\(^{-1}\).
4 Results and Discussion

4.1 Conventional Calibration method

4.1.1 Temperature profiles in the FAGE instrument

Temperatures within the FAGE instrument as a function of external temperatures are shown in Figures 2(b) and (c) and tabulated in Table 1. For Figure 2(b) and the first part of Table 1, the temperatures were recorded with FAGE sampling air at 293 K from the calibration flow tube as the FAGE inlet was cooled or heated. Temperatures became closer to ambient (293 K) from the inlet \( T_{\text{inlet}} \) to the OH observation cell \( T_{\text{OH}} \) and finally to the HO\(_2\) observation cell \( T_{\text{HO2}} \).

In Figure 2(c) and the second part of Table 1, the sampled air (either from the calibration flow tube or from HIRAC) matched the external temperature of the inlet tube. For these experiments, there was no thermocouple located inside the inlet to give \( T_{\text{inlet}} \). The temperature in the OH cell was very close to the external temperature of the sampled air. The transmission process through the FAGE inlet following sampling through the pinhole should be similar to when FAGE is in HIRAC, however, even with the temperature controlled air in the wand calibration, it is still difficult to determine the actual temperature and conditions at the pinhole itself.

The gap between the OH and HO\(_2\) cells means that the sampled air was closer to ambient room temperatures when reaching the HO\(_2\) cell. HO\(_2\) was predominantly exposed to a temperature environment similar to that for OH as it passed through the inlet, which may influence wall loss rates. The variation in \( T_{\text{OH}} \) and \( T_{\text{HO2}} \) relative to room temperature under different calibration regimes means that care has to be taken in comparing \( C_{\text{HOx}} \) values, as a number of processes within FAGE are temperature dependent. Nevertheless, the different calibration methods do yield important insights into the processes in the FAGE apparatus.
Table 1: Temperature Calibration of the FAGE instrument with a) constant temperature (293 K) calibration gas b) with calibration gas at the external temperature.

<table>
<thead>
<tr>
<th>External Temperature/K ($T_{\text{ext}}$)</th>
<th>Inlet Temperature /K, ($T_{\text{inlet}}$)</th>
<th>OH FAGE Cell Temperature /K, ($T_{\text{OH}}$)</th>
<th>HO$<em>2$ FAGE Cell Temperature /K, ($T</em>{\text{HO}_2}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Ambient Calibration Air at 293 K</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>266$^a$</td>
<td>280</td>
<td>284</td>
<td>290.5</td>
</tr>
<tr>
<td>273</td>
<td>282</td>
<td>284</td>
<td>293</td>
</tr>
<tr>
<td>276</td>
<td></td>
<td>285</td>
<td></td>
</tr>
<tr>
<td>293</td>
<td>293</td>
<td>293</td>
<td></td>
</tr>
<tr>
<td>308</td>
<td>301</td>
<td>301</td>
<td>296</td>
</tr>
<tr>
<td>323</td>
<td></td>
<td>308</td>
<td>302</td>
</tr>
<tr>
<td>328</td>
<td>313</td>
<td>312</td>
<td>299</td>
</tr>
<tr>
<td>343</td>
<td></td>
<td>318</td>
<td>308</td>
</tr>
<tr>
<td>353</td>
<td>326</td>
<td>323</td>
<td>313.5</td>
</tr>
<tr>
<td><strong>Calibration Air Matched to FAGE Inlet Tube Temperature</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>273</td>
<td>276</td>
<td>286</td>
<td></td>
</tr>
<tr>
<td>276</td>
<td></td>
<td>278</td>
<td></td>
</tr>
<tr>
<td>278</td>
<td>280</td>
<td>287</td>
<td></td>
</tr>
<tr>
<td>293</td>
<td>293</td>
<td>293</td>
<td></td>
</tr>
<tr>
<td>308</td>
<td>307</td>
<td>297</td>
<td></td>
</tr>
<tr>
<td>323</td>
<td>320</td>
<td>301</td>
<td></td>
</tr>
<tr>
<td>343</td>
<td>338</td>
<td>308</td>
<td></td>
</tr>
</tbody>
</table>

$^a$ – All temperature measurements have uncertainty of ± 0.5 K.

Figures 2(b) and (c) show the linear relationship between the internally measured temperature at the pinhole, OH cell and HO$_2$ cell. For Figure 2(b), the linear regression of the data gives ratios of 0.556 ± 0.002, 0.510 ± 0.002 and 0.195 ± 0.002 for the inlet thermocouple (close to the pinhole), OH cell and HO$_2$ cell. The temperature in the OH cell is controlled by the external temperature. In contrast, in field instruments which have a very different design and where OH is probed very close to the pinhole, there is a significant cooling effect due to the expansion (Creasey et al., 1997b). This is lost in the HIRAC FAGE due to the long inlet prior to probing the OH.

4.1.2 Temperature Dependent Flow Tube Calibration with Air at 293 K

Figure 5 displays the relative $C_{\text{OH}}$ and $C_{\text{HO}_2}$ for the HIRAC FAGE instrument as a function of external temperature between 266 – 343 K, with the data points listed in the top half of Table 2. In these experiments the FAGE inlet was cooled or warmed to give the external temperature ($T_{\text{ext}}$). The air from the calibration flow tube was at a constant 293 K and therefore the temperature in the observation cells (OH or HO$_2$) was varying compared to the inlet air. This
method of investigating the temperature dependence of $C_{\text{HOx}}$ therefore operates under different conditions from the subsequent methods (Sections 4.1.3 and 4.2). Data for $C_{\text{HOx}}$ are presented relative to the calibration factor at room temperature (293 K).

Figure 5: Temperature dependence of the calibration factors ($C_{\text{HOx}}$) as a function of the external temperature with HOx being delivered from the calibration flow tube at a constant temperature. Solid lines are a weighted fit to the data. (a) $C_{\text{OH,obs}}$, slope = (0.0023 ± 0.0007) K$^{-1}$. (b) $C_{\text{HO2,obs}}$, slope = (0.0005 ± 0.0031) K$^{-1}$. Errors are 2σ.

$C_{\text{OH,obs}}$ shows a positive temperature dependence (0.0023 ± 0.0007 K$^{-1}$), for $C_{\text{HO2,obs}}$, the data appear to be more scattered and no systematic trend is observable. The overall temperature dependence of both HOx calibration factors are small compared to the overall uncertainty in the calibration (40%); the relative calibration factor for OH changes by about 20% from 266 – 343 K. However, the error bars in Figure 5 represent the total error in the calibration, much of which will be temperature independent. A full discussion on the temperature dependence of the calibration factors is presented in Section 4.3.
Table 2: Instrument sensitivity to OH, \( C_{\text{OH}} \), and \( \text{HO}_2 \), \( C_{\text{HO}_2} \), determined using the conventional water vapour calibration method.

<table>
<thead>
<tr>
<th>( T_{\text{ext}}/K )</th>
<th>( T_{\text{OH}}/K )</th>
<th>( T_{\text{HO}_2}/K )</th>
<th>( C_{\text{OH,obs}} )</th>
<th>( C_{\text{HO}_2,obs} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ambient Calibration Air at 293 K</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>266</td>
<td>280</td>
<td>284</td>
<td>0.83 ± 0.42</td>
<td>1.11 ± 0.26</td>
</tr>
<tr>
<td>276</td>
<td>285</td>
<td>-</td>
<td>0.92 ± 0.42</td>
<td>-a</td>
</tr>
<tr>
<td>293</td>
<td>293</td>
<td>293</td>
<td>1.00 ± 0.42</td>
<td>1.00 ± 0.50</td>
</tr>
<tr>
<td>308</td>
<td>301</td>
<td>297</td>
<td>0.98 ± 0.41</td>
<td>1.36 ± 0.31</td>
</tr>
<tr>
<td>323</td>
<td>308</td>
<td>302</td>
<td>1.03 ± 0.42</td>
<td>1.40 ± 0.38</td>
</tr>
<tr>
<td>343</td>
<td>318</td>
<td>308</td>
<td>1.03 ± 0.42</td>
<td>1.01 ± 0.32</td>
</tr>
</tbody>
</table>

| Calibration Air Matched to FAGE Inlet Temperature (\( T_{in} \)) |
|--------------------------|------------------|------------------|-------------------|-------------------|
| 276 | 278 | - | 1.06 ± 0.39 | -a |
| 278 | 280 | 287 | 0.91 ± 0.50 | 1.43 ± 0.54 |
| 293 | 293 | 293 | 1.00 ± 0.40 | 1.00 ± 0.45 |
| 323 | 320 | 301 | 1.18 ± 0.39 | 1.91 ± 0.38 |
| 343 | 338 | - | 1.45 ± 0.39 | -a |

The internal temperatures (±0.5 K) for the OH and \( \text{HO}_2 \) fluorescence cells are represented by \( T_{\text{OH}} \) and \( T_{\text{HO}_2} \) respectively. a – determination of \( C_{\text{HO}_2} \) was precluded by a malfunctioning NO mass flow controller.

4.1.3 Temperature Dependent Flow Tube Calibration with Air at Varying Inlet Temperatures

A similar procedure to Section 4.1.2 was carried out, but in this case, the air flowing into the calibration flow tube had been cooled/heated to match the external temperature of the FAGE inlet. This method will give conditions that are more closely matched to those when the FAGE instrument is located in the HIRAC chamber, where the FAGE inlet is at the same temperature as the gas being sampled from HIRAC. The water vapour concentration was measured at a fixed temperature in the dew-point hydrometer and therefore the \([\text{HO}_x]\) emitted from the wand needed to be corrected for the change in [\( \text{H}_2\text{O} \)] and additionally, for the change in \( \Delta t \) in equation (E2).

In this calibration arrangement the temperature of the OH cell (\( T_{\text{OH}} \)) was virtually identical to the external temperature (\( T_{\text{ext}} \)). The \( \text{HO}_2 \) FAGE cell was closer to ambient room temperature.

The temperature dependence of \( C_{\text{HO}_x,obs} \) relative to 293 K is shown in Figure 6. The calibrations were taken at different times from those in Section 4.1.2, but the absolute \( C_{\text{HO}_x} \) factors at 293 K were in good agreement, within 5%. For OH, the slope of Figure 6(a) is again positive. For \( \text{HO}_2 \) (Fig 6(b)) there are only three datum points and they are somewhat scattered.
Figure 6: Temperature dependence of the calibration factors ($C_{\text{H}Ox,\text{obs}}$) as a function of the external temperature with HOx being delivered from the calibration flow tube at the external temperature. Solid lines are a weighted fit to the data. (a) $C_{\text{OH,obs}}$, slope = $(0.0059 \pm 0.0015)$ K$^{-1}$. (b) $C_{\text{HO}_2,\text{obs}}$, slope = $(0.0143 \pm 0.013)$ K$^{-1}$.

4.2 Alternative Calibration Methods

4.2.1 Hydrocarbon Decay Calibration of OH Sensitivity

The ratio of the conventional water vapour flowtube calibration to the HC decay method derived from scatter plots such as Figure 3 at 293 K was $1.034 \pm 0.068$, where the errors are the statistical errors in the gradient of the scatter plots at the 2σ level. The two methods are therefore in excellent agreement as has been observed in our previous study conducted solely at room temperature (Winiberg et al. (2015), 1.19 ± 0.26). The increased number of data points available for the HC analysis using PTR monitoring increases the precision of this work compared to our earlier studies where [HC] was measured at much lower time resolution by FTIR or gas chromatography.

A potential source of error in the HC decay method is quantifying the removal of the HC by non-OH sources. The effects of dilution and wall loss can be accounted for by suitable blank experiments, however, it is harder to account for any other chemically induced removal by photolytically generated radicals other than OH in such blank experiments. The hydrocarbons chosen for this analysis are simple alkanes with well-established chemistry that should minimize such possibilities i.e., very slow reactions with any photolytically generated O$_3$ or NO$_3$. In addition, when both cyclohexane (CH) and heptane (HEP) were used as the HC, the gradient of the resulting relative rate plot ($\ln([\text{HEP}]_0/\text{HEP}_t)$ vs $\ln([\text{CH}]_0/\text{CH}_t)$, slope = 0.923 ± 0.010) was in good agreement the ratio of the literature rate coefficients for OH reactions.
\( \frac{k_{\text{HEP}}}{k_{\text{CH}}} = 0.97 \pm 0.14 \) at 298 K (Atkinson, 2003). This confirms that OH was the dominant route for chemical removal (see SI, Section S4). A key assumption of the hydrocarbon decay calibration method is that the OH is chemically removed by OH.

**Table 3: Temperature Dependence of \( C_{\text{OH,obs}} \) Determined via the Hydrocarbon Decay Method**

<table>
<thead>
<tr>
<th>Temperature/K (±0.5 K)</th>
<th>( C_{\text{OH,obs}} ) relative to the HC decay method at 293 K</th>
</tr>
</thead>
<tbody>
<tr>
<td>273</td>
<td>0.92 ± 0.17(^a)</td>
</tr>
<tr>
<td>293</td>
<td>1.00 ± 0.18</td>
</tr>
<tr>
<td>323</td>
<td>1.10 ± 0.20</td>
</tr>
<tr>
<td>348</td>
<td>1.21 ± 0.22</td>
</tr>
</tbody>
</table>

\( a \) – errors represent the total uncertainty in \( C_{\text{OH}} \), see Table 4.

Displayed in Table 3 is the instrument sensitivity to OH radicals, \( C_{\text{OH,obs}} \), measured between 273 and 348 K at 1000 mbar HIRAC chamber pressure using the hydrocarbon decay method and Figure 7(a) shows these data as a function of the HIRAC temperature. An increase in \( C_{\text{OH}} \) is observed. As with the experiments carried out in Section 4.1.2, the temperature of the OH cell (\( T_{\text{OH}} \)) is very close to that of the gas being sampled at the inlet.

**Figure 7:** Temperature dependence of \( C_{\text{HO}_x,\text{obs}} \) relative to values at 293 K. Solid lines are a weighted fit to the data. (a) Relative \( C_{\text{OH,obs}} \) from the HC decay method. Slope = \((0.0038 \pm 0.0007) \text{ K}^{-1}\) (b) Relative \( C_{\text{HO}_2,\text{obs}} \) from the HCHO photolysis method. Slope = \((0.0064 \pm 0.0034) \text{ K}^{-1}\). Errors are 2\( \sigma \).
Table 4: The systematic uncertainties in the various parameters that determine the accuracy in
the OH and HO$_2$ calibration factors for the conventional and alternative calibration methods.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Uncertainty</th>
<th>Parameter</th>
<th>Uncertainty</th>
<th>Parameter</th>
<th>Uncertainty</th>
</tr>
</thead>
<tbody>
<tr>
<td>$F_{184.9 \text{ nm} \times t}$</td>
<td>20%$^a$</td>
<td>$k_{OH} - c$-C$<em>6$H$</em>{12}$</td>
<td>12%$^b$</td>
<td>$k_{HO_2 + HO_2}$</td>
<td>38%$^f$</td>
</tr>
<tr>
<td>[H$_2$O]</td>
<td>1%</td>
<td>$k_{Dil}$</td>
<td>2%$^c$</td>
<td>$S_{HO_2 \text{ initial}}$</td>
<td>10%$^g$</td>
</tr>
<tr>
<td>$\sigma_{\text{H}_2\text{O}}$</td>
<td>3%</td>
<td>[c-C$<em>6$H$</em>{12}$]</td>
<td>5%</td>
<td>Laser power</td>
<td>6%</td>
</tr>
<tr>
<td>Laser power</td>
<td>6%</td>
<td>Gradient</td>
<td>10%</td>
<td>Online Position</td>
<td>4%$^e$</td>
</tr>
<tr>
<td>Online Position</td>
<td>4%$^c$</td>
<td>Laser power</td>
<td>6%</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Online Position</td>
<td>4%$^d$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Error</td>
<td>22%$^e$</td>
<td>Error</td>
<td>18%$^e$</td>
<td>Error</td>
<td>40%$^e$</td>
</tr>
</tbody>
</table>

$^a$ – Where the error is statistical, it is reported at the 1σ level.
$^b$ – Error estimated from literature review. Five recent determinations (NIST Kinetics) of the 298 K rate coefficient
give ~5% spread, added some additional uncertainty to account for temperature dependence.
$^c$ – Dilution determined from flow controller measurements.
$^d$ – The online position error is the approximate error in the maximum line intensity that is achieved when
positioning the laser wavelength at the centre of the OH transition.
$^e$ – Total accuracy is taken as the sum in quadrature of the individual uncertainties.
$^f$ – Error in rate coefficient from the IUPAC evaluation.
$^g$ – Uncertainties in the fitting parameters.

Table 4 summarizes the errors associated with the alternative calibration methods. For the
hydrocarbon decay method, the major uncertainties are in the rate coefficient of the
hydrocarbon (~12% for OH + cyclohexane), determination of cyclohexane concentration (5%)
and the gradient of the cyclohexane decay (10%). Other uncertainties are drifts in the laser
power (~6%, determined from monitoring a photodiode) and wavelength position (~4%).

4.2.2 Calibration via HO$_2$ recombination kinetics

Displayed in Table 5 is the instrument sensitivity to HO$_2$, $C_{HO_2,\text{obs}}$, determined using the
alternative calibration method between 273 and 343 K at 1000 mbar chamber pressure. Figure
7(b) shows $C_{HO_2}$ as a function of temperature relative to the instrument sensitivity at 293 K.
Each measurement point represents the weighted average of at least five experimental data sets
and the error bars represent the total uncertainty in the instrument sensitivity to ±2σ. As with
the hydrocarbon decay method, the overall uncertainty is calculated as the sum in quadrature
of fit precision to the decay and the systematic uncertainties listed in Table 4. The largest
uncertainty was in the HO$_2$ self-reaction rate coefficient, dependent on the temperature used.
(38\%). The slope of the linear fit to the $C_{\text{HO2}}$ values is $(0.0064 \pm 0.0034) \text{ K}^{-1}$. The absolute agreement between the conventional and HCHO photolysis methods at 293 K is good with $C_{\text{HO2, conventional}} = (3.38 \pm 1.08) \times 10^{-8} \text{ counts cm}^3 \text{ molecule}^{-1} \text{ mW}^{-1} \text{ s}^{-1}$ and $C_{\text{HO2, HCHO photolysis}} = (3.69 \pm 1.48) \times 10^{-8} \text{ counts cm}^3 \text{ molecule}^{-1} \text{ mW}^{-1} \text{ s}^{-1}$.

### Table 5: Instrument sensitivity to HO$_2$, $C_{\text{HO2}}$, determined using the HCHO photolysis method over the 273 – 343 K external inlet temperature range.

<table>
<thead>
<tr>
<th>$T_{\text{HIRAC}} / \text{K}$</th>
<th>$T_{\text{HO2}} / \text{K}$</th>
<th>$C_{\text{HO2}}$ (rel. 293 K)$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>273</td>
<td>286</td>
<td>0.89 ± 0.36$^c$</td>
</tr>
<tr>
<td>293</td>
<td>293</td>
<td>1.00 ± 0.40</td>
</tr>
<tr>
<td>308</td>
<td>297</td>
<td>1.38 ± 0.55</td>
</tr>
<tr>
<td>323</td>
<td>302</td>
<td>1.05 ± 0.42</td>
</tr>
<tr>
<td>343</td>
<td>308</td>
<td>1.40 ± 0.56</td>
</tr>
</tbody>
</table>

a – Error in temperature ± 0.5 K.
b – Values are relative to $C_{\text{HO2, 293 K}}$ of $(3.69 \pm 1.48) \times 10^{-8} \text{ counts cm}^3 \text{ molecule}^{-1} \text{ mW}^{-1} \text{ s}^{-1}$.
c – Each $C_{\text{HO2}}$ represents the weighted average of at least 5 individual determinations. All experiments were conducted in 1000 mbar synthetic air mixture.

### 4.3 Discussion of calibration methods and temperature dependence

#### 4.3.1 Comparison of calibration methods

For room temperature, there is excellent agreement between the wand calibration and that for OH based on hydrocarbon decays ([OH]$_\text{wand}$:[OH]$_\text{HC}$ = 1.00:0.97) and HO$_2$ based on HCHO photolysis and the kinetics of the HO$_2$ recombination reaction ([HO$_2$]$_\text{wand}$:[ HO$_2$]$_\text{kinetics}$ = 1.00:1.09). This is consistent with our earlier study (Winiberg et al. 2015) and has also been confirmed in an intercomparison in the HIRAC chamber of the FAGE and NIR – CRDS (near infrared cavity ring down spectroscopy) for HO$_2$ (Onel et al., 2017a) and CH$_3$O$_2$ (Onel et al., 2020;Onel et al., 2017b).

For the hydrocarbon decay method there are several advantages compared to the conventional wand calibration:

1) The [OH] is much closer to the conditions typically used in a chamber experiment ($10^6 – 10^8$ molecule cm$^{-3}$) whereas the lowest [OH] used in the wand calibration performed here is typically $10^8$ molecule cm$^{-3}$. Ideally one should calibrate over the same range as used in an experiment.
2) This work has shown that there is a temperature dependence to the calibration factors.
   Calibrating via the hydrocarbon decay method provides identical conditions (temperature
   and pressure) to that of a real experiment in the HIRAC chamber. Temperature variation
   can be simulated using the conventional wand device, but this introduces additional
   uncertainty.

3) Conventional calibrations always take place with a significant water concentration, whereas
   the water concentration in the hydrocarbon decay can be set at any value.

4) Calibration can be achieved without removing the FAGE apparatus from the HIRAC
   chamber decreasing the time taken for calibration.

   There are some disadvantages too. The calibration for OH is strongly dependent on the
   accuracy of the HC rate coefficient. It is therefore important to use a hydrocarbon with a well-
   characterised rate coefficient; realistically, even the best-characterised rate coefficient is likely
   to have an uncertainly of 5 – 10%. Several HC can be used to give multiple independent
   determinations of [OH]HC, but this may increase the complexity of the analysis (e.g. coincident
   mass spectral peaks, or overlapping FTIR spectra) and reduce the absolute concentration of
   OH. Determination of [OH]HC also relies on an accurate and precise determination of the
   concentration gradient and the [HC] at that time. PTR measurements provide a near continuous
   output, but if the [HC] is measured using systems with lower sampling rates (e.g. FTIR or GC),
   there can be a significant loss in precision of the gradient measurement.

   Many of the advantages and disadvantages of the hydrocarbon decay method also apply to
   HO2 kinetics method for HO2 calibration. The rate coefficient for HO2 recombination has a
   higher degree of uncertainty than many OH + hydrocarbon rate coefficients and is dependent
   on the amount of water present. In the HIRAC chamber the humidity can be kept very low, but
   that may not be possible in all chambers; in these circumstances the humidity would need to
   be measured and the rate coefficient adjusted.

   All calibration methods are subject to systematic uncertainties, the magnitude of which may
   vary with conditions and therefore it is sensible to use a range of calibration methods.

4.3.2 Temperature dependence of C_{HOx}

Table 6 compares the relative observed C_{HOx,obs} calibration factors for the three different
 calibration methods. In all cases, a positive temperature dependence is observed, but for C_{HO2},
 only the alternative calibration method displays a statistically significant positive slope.
The $C_{\text{HOx}}$ factors can be broken down into temperature independent components (laser power, solid angle of fluorescence collection, detector efficiency etc) and temperature dependent terms. Four temperature dependent terms are relevant for $C_{\text{HOx}}$: the number density of OH in the cell, the quenching efficiency of the fluorescence, the population of the probed quantum state of OH and the transmission efficiency through the pinhole and inlet tube (Creasey et al., 1997b). The first three terms can be calculated and hence accounted for. Any residual temperature dependence of $C_{\text{HOx}}$ should then relate to the transmission coefficient through the apparatus.

**HOx number density** – The calculated [HOx] delivered to the FAGE apparatus depends on the temperature of the HOx source, either the wand (operating at a fixed $T = 293$ K (Method 1) or at $T_{\text{ext}}$ (Method 2) or the HIRAC chamber. If the temperature of the HOx cells are different from this temperature, then there will be a change in the number density of HOx, over and above that caused by the pressure changes between the HOx source (1 bar) and the HOx cell (typically 3.6 mbar). As the temperatures of the HOx cells have been measured it is straightforward to correct for the different number density in the observation cells and the resulting contribution to the temperature dependence of $C_{\text{HOx}}$ as summarized in Tables S2-4.

**Quenching** – As shown in Faloona et al. (2004), the quenching parameter, $Q(T)$, is defined by integrating the OH fluorescence decay over the defined sample time, or gated region. The quenching rate coefficients for N$_2$, O$_2$ and H$_2$O have been shown to be dependent on temperature (Copeland and Crosley (1986) and (Bailey et al., 1997) for N$_2$ and O$_2$, and Bailey et al. (1999) for H$_2$O). The total decay intensity is defined by: $[\text{OH}(A^2\Sigma^+, \nu' = 0)]_0 \exp(-\Gamma t)$, where $\Gamma$, the total OH lifetime, is defined approximately as the sum total of the radiative lifetime for OH, $\gamma$, and the non-radiative lifetime due to quenching by the aforementioned bath gases. Bailey et al. (1997) have calculated the impact of temperature on quenching accounting for both the change in the quenching rate coefficients and the change in the number density of the quenchers. Both the rate coefficient for quenching and the quencher number density decrease with increasing temperature and hence quenching overall decreases with increasing temperature (summarized in Table S5), enhancing the fluorescence quantum yield.

**Rotational population** – The rotational population of the probed state in the Q$_1$(2) transition will vary with temperature. The Q$_1$(2) is the transition giving the largest signal between 280 – 340 K, the limits of $T_{\text{OH}}$ explored in the study. Relative to ambient temperature, the rotational population probed by Q$_1$(2) increases by 3.5% at 280 K and decreases by 9.0% at the highest $T_{\text{OH}}$ of 340 K (Table S6).
It is therefore possible to calculate the expected variation in $C_{\text{HOx}}$ for the different calibration methods dependent on OH number density, quenching and rotational population; these can be compared with the observed variation in $C_{\text{HOx}}$ summarized in Table 6. Full details on the temperature dependences of the above components, which vary slightly with the calibration method used are presented in Section S5 of the SI.

The difference between the observed $C_{\text{HOx}}$ and the calculated $C_{\text{HOx}}$ due to the above parameters is attributed to increased transmission of HOx through the pinhole and inlet tube and is given in Table 6. The HOx transmission, to the fluorescence region will depend on the magnitude of heterogeneous loss of radicals to the walls of the FAGE inlet. The wall loss process is a combination of diffusion and uptake at the wall and the actual temperature dependence will depend on the radical, conditions and wall composition (Howard, 1979).

For the OH calibrations, there is an increase in OH transmission with temperature across all three calibration methods, consistent with a decrease in OH loss to the walls which has been observed in previous flow tube studies. OH wall loss rate in the inlet tube is usually approximated to a first order process with a rate coefficient, $k_w$, and decreasing values of $k_w$ with temperature have been reported for flow tube studies of OH reactions (Howard, 1979), for example Brown et al. (1990) report $k_w$ decreasing from $35 \text{ s}^{-1}$ at 227 K to $5 \text{ s}^{-1}$ at room temperature.

For HO$_2$ measurements, there is potentially a further temperature dependent component, the conversion of HO$_2$ into OH via R7:

$$\text{HO}_2 + \text{NO} \rightarrow \text{OH} + \text{NO}_2 \quad \text{(R7)}$$

The rate coefficient for this reaction has a negative temperature dependence and the increased number density of NO would further enhance the rate of reaction at lower temperatures. The experiments reported in this work operated with excess NO such that the small variations in the rate of reaction over the range of $T_{\text{HO2}}$ (284 – 313 K) will not alter the conversion of HO$_2$ to OH. However, if one were working at lower HO$_2$ conversions to mitigate against RO$_2$ to OH conversion (Whalley et al. 2013), then variations in the conversion efficiency could change $C_{\text{HO2}}$ as a function of temperature.

Temperature dependent HO$_2$ calibrations based on the conventional wand method give significant scatter, but a positive increase in HO$_2$ transmission is observed for the alternative calibration method based on HO$_2$ kinetics, the magnitude of which is similar to that for OH, albeit with significant error bars. In general, HO$_2$ and RO$_2$ radicals exhibit lower wall loss rate
Table 6: Summary of the temperature dependence of CHO2 with different calibration methods

<table>
<thead>
<tr>
<th>Method</th>
<th>Observed slope of relative C_{OH,obs} with temperature</th>
<th>Calculated contribution$^1$</th>
<th>Difference (relative OH transmission)</th>
<th>Observed slope of relative C_{HO2,obs} with temperature</th>
<th>Calculated contribution$^a$</th>
<th>Difference (relative HO2 transmission)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heated FAGE inlet, ambient air at 293 K</td>
<td>(0.0023 ± 0.0007) K$^{-1}$</td>
<td>(0.0001 ± 0.0010) K$^{-1}$</td>
<td>(0.0022 ± 0.0012) K$^{-1}$</td>
<td>(0.0005 ± 0.0031) K$^{-1}$</td>
<td>(0.0000 ± 0.00010) K$^{-1}$</td>
<td>(0.0000 ± 0.00032) K$^{-1}$</td>
</tr>
<tr>
<td>Heated FAGE inlet, matched air</td>
<td>(0.0059 ± 0.0015) K$^{-1}$</td>
<td>(0.0029 ± 0.0010) K$^{-1}$</td>
<td>(0.0030 ± 0.0018) K$^{-1}$</td>
<td>(0.014 ± 0.013) K$^{-1}$</td>
<td>(0.0033 ± 0.0010) K$^{-1}$</td>
<td>(0.0029 ± 0.0016) K$^{-1}$</td>
</tr>
<tr>
<td>Alternative kinetics based methods</td>
<td>(0.0038 ± 0.0007) K$^{-1}$</td>
<td>(0.0027 ± 0.0010) K$^{-1}$</td>
<td>(0.0011 ± 0.0012) K$^{-1}$</td>
<td>(0.0064 ± 0.0034) K$^{-1}$</td>
<td>(0.0032 ± 0.0010) K$^{-1}$</td>
<td>(0.0032 ± 0.0035) K$^{-1}$</td>
</tr>
</tbody>
</table>

$^a$ - Contribution from the change in number density, quenching and relative rotation population in the probed state.
coefficients, but in our FAGE system, HO$_2$ molecules have to travel further to reach the
titration region where reaction occurs with NO to convert HO$_2$ to OH. Therefore, there is also
potential for OH loss from the titration point to the second detection cell.

4.3.3 Comparison with other instruments

The temperature dependence of the calibration factors will be strongly dependent on the design
of the FAGE apparatus. Our instrument was designed with a long (~ 1 m) inlet such that we
can probe across the diameter of the HIRAC chamber to check for radial distributions of
radicals (Malkin et al., 2010). Hence, we would expect HOx transmission to play a significant
role in the temperature dependence of the calibration factor which is observed. Any similarly
designed instrument would have a contribution from HOx transmission, the magnitude of
which would depend on inlet length/residence time and construction material. Heating the inlet
should reduce transmission losses. The aircraft based instrument, from the Juelich research
group, uses a PID controlled heater to maintain their FAGE inlet at ~300 K, mitigating any
possible temperature effects. They have an in-field calibration system, also, which has shown
negligible deviation from the expected behaviour at 300 K, based on the sample gas altitude
temperature (Marno et al., 2020).

Regelin et al. (2013) have reported a similar temperature dependence study of $C_{OH}$ and $C_{HO2}$
as the current flowtube study with the aircraft based HORUS instrument. Cooling lines were
wound around the inlet to simulate the measured temperature profile and ambient air was
sampled from a calibration flow tube. In contrast to our slight increase in $C_{OH}$ with temperature
in the flow tube experiment, Regelin et al. observed a slight negative dependence of the OH
signal. Regelin et al. report that their calculations have shown that the sample forms a jet
between the pinhole and the OH cell such that there is insignificant interaction with the walls
and therefore transmission will not be a problem.

In contrast, a significant decrease in HO$_2$ signal, $S_{HO2}$, (50%) was observed as the
temperature was decreased from ~295 to ~262 K (slope = 0.017 K$^{-1}$ normalised to $S_{HO2,293}$ K),
i.e. the same qualitative behaviour as we observed, approximately a factor two greater than
measured in our work, based on HO$_2$ recombination kinetics. Beyond the OH cell in the
HORUS experiment, the jet breaks up and Regelin et al. suggest that temperature dependent
wall losses are responsible for the change in $S_{HO2}$. Quantitative comparisons cannot be made
due to the differences in construction. The observed temperature dependence of $C_{OH}$ and $C_{HO2}$
for the HORUS and HIRAC experiments emphasise the important of performing calibrations
for each instrument under conditions as close as possible to those used in measurements.

5 Conclusions

The effect of temperature of the incoming sample on the sensitivity of the HIRAC FAGE
instrument to OH and HO$_2$ has been investigated between 266 and 348 K using a combination
of conventional water vapour photolysis/flow tube method (Faloona et al.) and alternative
calibration methods based on hydrocarbon decays for OH and the HO$_2$ self-reaction for HO$_2$.
In all cases, a positive increase in sensitivity was observed (Table 6) although with large error
bars in the case of HO$_2$ with conventional calibration.

The temperature dependence of the calibration factor can be broken down to four
components. Variations in three parameters: number density, quenching and rotational
population of the probed level, can be accounted for if the temperature and pressure in the LIF
cells are monitored. The difference between the observed and calculated temperature
dependence for the above parameters, has been attributed to HOx transmission from the pinhole
to the relevant detection chamber.

The temperature dependence of $C_{\text{HOx}}$ will depend on the design and construction materials
of the FAGE apparatus. It is therefore difficult to utilise the results of this study to predict
results in other systems. However, for any systems with significant sampling inlet residence
times, such as the HIRAC FAGE described in this work, increased HOx transmission with
increasing temperature should be expected. Therefore, maintaining the inlet at a relatively high
temperature should improve sensitivity in low temperature applications.

The in situ calibration methods (hydrocarbon decay and HO$_2$ recombination kinetics) offer
important advantages in that the FAGE apparatus is calibrated under the physical conditions
and [HOx] that more closely correspond to real experiments. All calibration methods are
subject to significant uncertainty, however, the origins of these uncertainties are different and
hence good agreement between calibration methods should provide confidence that significant
systematic errors are not present.

Supplementary Information

Supplementary information; HIRAC temperature profiles, calibrations, further discussions on
calibration uncertainties, relative rate plots to confirm OH as the key species in hydrocarbon
removal and further discussion on the temperature dependence of the FAGE signal can be found at ******.

Code/Data availability

Any raw data is available either in the Supplementary Information or via contacting Prof. Paul Seakins.

Author Contributions

FAFW and IGB led the initial work on OH temperature dependence performing all experiments with external calibration, WJW, THS and GB completed the experiments with HC decays in HIRAC, CAB and IGB completed experiments on HO₂ temperature dependence. PWS, DEH and DS planned and supervised the experiments and wrote the manuscript with contributions from all co-authors.

Competing Interests

DEH is a member of the editorial board of AMT, otherwise the authors declare that they have no conflict of interest.

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