Technical Note: Formal blind intercomparison of HO_2 measurements in the atmosphere simulation chamber SAPHIR during the HOxComp campaign

H. Fuchs¹, T. Brauers¹, H.-P. Dorn¹, H. Harder², R. Häseler¹, A. Hofzumahaus¹, F. Holland¹, Y. Kanaya³, Y. Kajii⁴, D. Kubistin², S. Lou¹⁵, M. Martinez², K. Miyamoto⁴, S. Nishida⁴, M. Rudolf², E. Schlosser¹*, A. Wahner¹, A. Yoshino⁴, and U. Schurath⁶

¹Forschungszentrum Jülich, ICG-2: Troposphäre, 52425 Jülich, Germany
²Max Planck Institute for Chemistry, Atmospheric Chemistry Department, 55020 Mainz, Germany
³Frontier Research Center for Global Change (currently Research Institute for Global Change), Japan Agency for Marine-Earth Science and Technology, Yokohama, 236-0001, Japan
⁴Tokyo Metropolitan University, Department of Applied Chemistry, Tokyo 192-0397, Japan
⁵Shanghai Jiatong University, School of Environmental Science and Technology, Shanghai, China
Karlsruhe Institute of Technology (KIT), IMK-AAF, 76021 Karlsruhe, Germany
*now at: Karlsruhe Institute of Technology (KIT), IMK-AAF, 76021 Karlsruhe, Germany

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Correspondence to: T. Brauers (th.brauers@fz-juelich.de)

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Abstract

Hydroperoxy radical (HO\textsubscript{2}) concentrations were measured during the formal blind intercomparison campaign HOxComp carried out in J"ulich, Germany. Three instruments detected HO\textsubscript{2} via chemical conversion to hydroxyl radicals (OH) and subsequent detection of the sum of OH and HO\textsubscript{2} by laser induced fluorescence (LIF). Instruments sampled ambient air for three days and were attached to the atmosphere simulation chamber SAPHIR during the second part of the campaign. Six experiments of one day each were conducted in SAPHIR, where air masses were homogeneously mixed, in order to investigate the performance of instruments and to determine potential interferences of measurements under well-controlled conditions. Linear correlation coefficients between measurements of the LIF instruments are generally high and range from 0.82 to 0.98. However, the agreement between measurements is variable. The regression analysis of the entire data set of measurements in SAPHIR yields slopes between 0.69 to 1.26 and intercepts are smaller than typical atmospheric daytime concentrations (less than 1 pptv). The quality of fit parameters improves significantly, when data are grouped into data subsets of similar water vapor concentrations. Because measurements of LIF instruments were corrected for a well-characterized water dependence of their sensitivities, this indicates that an unknown factor related to water vapor affected measurements in SAPHIR. Measurements in ambient air are also well correlated, but regression parameters differ from results obtained from SAPHIR experiments. This is most likely caused by sampling different air masses at the slightly distant locations of instruments.

1 Introduction

Hydroperoxy radicals (HO\textsubscript{2}) play an important role in the photochemical degradation of atmospheric trace gases and in the formation of secondary air pollutants. They are produced in the radical recycling of the hydroxyl radical (OH), the major atmospheric
oxidant (e.g., Finlayson-Pitts and Pitts Jr., 2000), via the reaction of OH with CO and organic compounds. Direct sources of HO₂ radicals are the photolysis of organic carbonyl compounds such as acetaldehyde, the reaction of the nitrate radical with organic compounds and the ozonolysis of alkenes (e.g., Geyer et al., 2003; Kanaya et al., 2007). In NOₓ-rich environments, peroxy radicals react predominantly with NO reforming OH and producing NO₂. This reaction and the following photolysis of NO₂ constitute also fundamental steps in the photochemical formation of tropospheric ozone in polluted air. In very clean air at low NO, however, HO₂ has the potential to destroy ozone.

The measurement of HO₂ concentration is challenging for several reasons. (1) Atmospheric concentrations are small (some ten parts per trillion per volume, pptv) (e.g. Monks, 2005; Kanaya et al., 2007; Lelieveld et al., 2008; Hofzumahaus et al., 2009), so that a high detection sensitivity is required to achieve a good signal-to-noise ratio at a reasonable time resolution. (2) Their high reactivity may cause loss of HO₂ in the inlet system of instruments (e.g. Mihele and Hastie, 1998). (3) Calibration of instruments is difficult, because a radical source producing an accurately known radical concentration is required.

Four different techniques have been developed that are applied for atmospheric HO₂ detection (Heard and Pilling, 2003): (1) Peroxy Radical Chemical Amplifier (PERCA), (2) Peroxy Radical Chemical Ionization Mass Spectrometry (ROxMax/PerCIMS), (3) Laser Induced Fluorescence (LIF), and (4) Matrix Isolation Electron Spin Resonance Spectroscopy (MIESR).

PERCA and PerCIMS/ROxMas use chemical amplification by repetitive recycling of HO₂ in a chemical chain reaction to produce an amount of a product species that is well measurable. PERCA instruments produce NO₂, which is detected by a luminol-cheluminescence reaction, whereas PerCIMS/ROxMas techniques detect H₂SO₄ by chemical ionization mass spectrometry. LIF systems convert HO₂ to OH radicals by adding excess NO to the sampled air. OH radicals are then detected via resonant laser-induced fluorescence at 308 nm. MIESR traps radicals in an ice matrix formed by D₂O on a cold finger at a temperature of 77 K. The sample is analyzed by means of
electron spin resonance spectroscopy (ESR) in the laboratory. The MIESR technique is the only one of the four techniques, which does not require calibration.

The experimental difficulties and the large experimental effort of atmospheric HO\textsubscript{2} detection limit the number of instruments used in field experiments. There have been only three attempts to intercompare instruments and all included two instruments only. Thus, the quality and comparability of measurements from different instruments is not well-known. Measurements by LIF and MIESR were compared during the BERLIOZ field campaign (Platt et al., 2002; Mihelcic et al., 2003) and agreed to 3%. The linear correlation coefficient, \( R^2 \), was 0.88. In another, more recent study, HO\textsubscript{2} concentrations were compared between a PerCIMS and an LIF instrument in two phases: (1) by mutual exchange of calibration sources, (2) by ambient air measurements (Ren et al., 2003). In the calibration intercomparison, very good agreement was found within 2% for PerCIMS sampling from the LIF calibration source and within 4% for LIF sampling from the PerCIMS calibration source. Both calibration techniques were based on the production of OH by water photolysis at 185 nm (see below). Good agreement was also shown in a side-by-side intercomparison of ambient HO\textsubscript{2} concentration, which agreed to 4% and exhibited a linear correlation coefficient of \( \hat{R}^2 = 0.85 \). Recently, however, Ren et al. (2008) reported a change of the calibration factor of the LIF instrument. This may possibly require revision of the comparison results. The latest comparison of instruments was conducted between an LIF instrument (ROxLIF), which is also capable of detecting the sum of organic peroxy radicals (RO\textsubscript{2}), and MIESR during two experiments in the atmosphere simulation chamber SAPHIR in Jülich, Germany, after the HOxComp campaign in 2007 (Fuchs et al., 2009). Measurements agreed on average to 2%.

The HOxComp campaign in Jülich, Germany, was an effort supported by the EU program ACCENT to bring together a larger number of instruments, which are capable of detecting OH and HO\textsubscript{2} radicals, in order to intercompare measurements in a formal blind manner. Four LIF instruments were successfully deployed during this campaign: (1) the instrument of the Frontier Research Center for Global Change, Yokohama,
Japan, (FRCGC-LIF), (2) two instruments from the Forschungszentrum Jülich, Germany, one for deployment in field campaigns that took part only during the ambient air sampling period (FZJ-LIF-ambient) and one that is permanently attached to the SAPHIR chamber (FZJ-LIF-SAPHIR), (3) the instrument of the Max-Planck Institute for Chemistry in Mainz, Germany (MPI-LIF). Another LIF instrument from the University of Leeds took part in the campaign, but could not perform measurements, because of a failure of the laser system. The MIESR instrument from the Forschungszentrum Jülich took samples during the campaign, but data were rejected (see below for details). Various instruments measured concentrations of other trace gases such as ozone, nitrogen oxides and volatile organic compounds (VOCs) (see Schlosser et al., 2009 for details). Three days of ambient air sampling were followed by six experiments in the SAPHIR chamber.

The intercomparison of data was done in a formal blind way. Experimental details were known to the referee only and data exchange between the participating groups was not allowed until data were finalized 8 weeks after the campaign. Preliminary data were given to the referee on a daily basis. Questionable data points were marked in the data set. They are excluded from the analysis here. Data of the FZJ-LIF were changed after the data submission deadline, because a systematic error in the calculation of the flow in the calibration source was discovered (see Schlosser et al., 2009 for details). The correction increased HO$_2$ concentrations for the FZJ-LIF-ambient and FZJ-LIF-SAPHIR instrument by 26% and 28%, respectively.

2 Instruments

2.1 Matrix Isolation and Electron Spin Resonance Spectroscopy (MIESR)

For the detection of HO$_2$ radicals by the MIESR technique, radicals are collected during the experiment and samples are analyzed later in the laboratory. A detailed description of the instrument and the analysis procedure is found in Mihelcic et al. (1985, 1990).
Concentrations of \( \text{HO}_2 \), \( \text{RO}_2 \), \( \text{CH}_3\text{C(O)}\text{O}_2 \), \( \text{NO}_3 \) and \( \text{NO}_2 \) can be measured. This instrument took samples of ambient air and was also attached to the SAPHIR chamber for two of the experiments. Although \( \text{HO}_2 \) concentration data were submitted to the referee after the campaign, data were withdrawn later, because they were most likely corrupted by an instrumental problem. This was recognized, when \( \text{NO}_2 \) concentrations by MIESR were compared to \( \text{NO}_2 \) concentrations measured by a chemiluminescence detector (CLD). This showed an irregular behavior of the MIESR instrument. Differences between \( \text{NO}_2 \) concentrations were much larger than expected from earlier comparisons between MIESR and CLD. A similar random relationship between data was found when \( \text{HO}_2 \) concentrations measured by MIESR and all LIF instruments were compared. Again, differences were much larger than observed in previous and later campaign (Platt et al., 2002; Fuchs et al., 2009). Both together indicate that measurements by MIESR were corrupted. The reason for the failure of the analysis of the MIESR samples is not clear. However, the data do not allow a reasonable comparison of \( \text{HO}_2 \) measurements and were rejected.

2.2 Laser-Induced Fluorescence (LIF)

The LIF method for \( \text{HO}_2 \) detection takes advantage of the chemical conversion of \( \text{HO}_2 \) to \( \text{OH} \) radicals, which can be detected by LIF (Heard and Pilling, 2003). Therefore, LIF instruments for ambient \( \text{OH} \) detection usually are also capable of detecting \( \text{HO}_2 \). All LIF instruments deployed in this campaign are similar with respect to their general concept of detection, but differ in technical details described in Kanaya et al. (2001); Kanaya and Akimoto (2006); Holland et al. (2003); Martinez et al. (2010) for FRCGC-LIF, FZJ-LIF and MPI-LIF, respectively. In the following, common properties of FZJ-LIF-ambient and FZJ-LIF-SAPHIR are labeled as properties of FZJ-LIF. Properties of the LIF-instruments are summarized in Table 1 and briefly described in the following.

All instruments sample air through an inlet pin hole into a low pressure fluorescence cell. The diameter of the pin hole, which also determines the flow rate into the cell, differs between the instruments FRCGC-LIF and MPI-LIF (diameter: 1.0 mm, flow rate:
7 slm) and FZJ-LIF (diameter: 0.4 mm, flow rate: 1 slm). The distance between the inlet pin hole and detection volume varies between 23 cm (FRCGC-LIF), 10 cm (FZJ-LIF), and 33 cm (MPI-LIF). OH radicals are excited at a single rovibronic line of the OH $A^2\Sigma^+ (v'0) \leftarrow X^2\Pi(v'' = 0)$ transition by narrow-bandwidth radiation at 308 nm. All instruments use a pulsed, frequency doubled Nd:YAG laser (DPSS) to pump a tunable dye laser, whose wavelength is again frequency doubled, in order to produce the 308 nm radiation. The gas volume in the fluorescence cell is exchanged between two consecutive laser shots to avoid self-generation of OH by ozone photolysis (laser repetition rates: FRCGC-LIF: 1 kHz, FZJ-LIF: 8.5 kHz, MPI-LIF: 3 kHz). The laser beam in the FZJ-LIF and FRCGC-LIF instruments passes the fluorescence cell one time whereas a multi-pass White cell system is used in the MPI-LIF instrument.

After laser excitation, the instruments measure the resonant OH fluorescence (307–311 nm) by time-delayed gated photon counting (time delay 70 to 160 ns, duration 300 to 500 ns). The measured signals also contain contributions from laser excited stray light, from a laser independent dark signal of the detector, and background from solar radiation entering the measurement cell through the inlet orifice. In the FZJ-LIF and FRCGC-LIF instruments, the solar background is quantified after each laser pulse in a second time gate after laser excited signals have decayed to zero. The solar background is then subtracted from the signal. In order to keep the correction small, FRCGC-LIF placed a black aluminum disk coated with halocarbon wax above the nozzle to shade the inlet against direct sunlight. In all instruments the laser excitation wavelength is periodically tuned from the OH absorption line to off-resonant wavelengths, in order to measure the laser stray light and dark signal of the detector. In the MPI-LIF instrument, this signal is also used to account for the signal from solar radiation. Scanning the laser wavelength also allows to lock the laser wavelength to the OH absorption line by observing the signal of a reference cell, in which artificial OH is produced either by a hot filament or by water photolysis at 185 nm.

The conversion of HO$_2$ to OH is accomplished by adding excess NO to the sampled gas. In the FRCGC-LIF instrument, the NO flow is periodically switched on and off.
allowing for alternating detection of OH and the sum of OH and HO$_2$ in the same fluorescence cell. 3 sccm pure NO is injected through a loop (diameter 10 mm) of Teflon tubing with small pinholes downstream of the inlet nozzle (NO mixing ratio 0.04% in the sampled air). The distance between the NO injector loop and the OH detection volume is approximately 21 cm. The other instruments have two detection cells for simultaneous detection of OH and HO$_2$. Both LIF-FZJ instruments have two fluorescence cells with separate inlet nozzle and are passed consecutively by the laser beam. 4 sccm of pure NO is added through a glass ring with small pinholes that is placed 5.5 cm upstream of the fluorescence detection. The resulting NO mixing ratio is 0.6% and the distance to the center of the fluorescence cell gives a reaction time of approximately 0.2 ms for the conversion of HO$_2$. During the period between 18:00 10 July (all times in UTC) and 18:00 11 July the HO$_2$ cell of the LIF-FZJ was equipped with an additional conversion reactor upstream of the HO$_2$ detection cell, in order to detect alternately HO$_2$ and the sum of HO$_2$ and RO$_2$ as described in Fuchs et al. (2008). Regarding HO$_2$ detection, the main differences compared to the previous detection scheme are: (1) A 83 cm long conversion reactor with a residence time of 0.6 s at reduced pressure of 25 hPa is placed upstream of the fluorescence cell. (2) Sampled gas is expanded from the conversion reactor into the low pressure fluorescence cell at 3.5 hPa through a 4 mm diameter nozzle. (3) Excess CO (mixing ratio in the sampled air: 0.17%) is added downstream of the inlet of the conversion reactor. A different approach compared to FRCGC-LIF and FZJ-LIF is applied in the MPI-LIF instrument, where a second fluorescence cell is placed 10 cm downstream of the cell in which OH is detected, in order to measure HO$_x$ concentrations. 6 sccm NO is injected through a loop of perforated tubing placed between both cells, giving a mixing ratio of 0.09% in the sampled air. The distance between NO injection and fluorescence detection is approximately 10 cm corresponding to a reaction time of 6 ms.

A small background signal is generally observed, when NO is added to the sampled gas presumably caused by reactions of impurities in the NO gas that produce artificial OH. The interference is minimized in all instruments by purifying the NO gas.
with Ascarite prior addition in the cell. The value of the background signal is less than 0.3 pptv for the MPI-LIF instrument, between 0.2 and 0.3 pptv for the FZJ-LIF instruments and between 0.16 to 0.27 pptv for the FRCGC-LIF instrument. Only measurements by FZJ-LIF were corrected for this small interference. Measurements by other instruments remained uncorrected, because it was not clear, if this offset was constantly present for all measurements.

It is known that ozone can cause an interference in LIF measurements by its photolysis at 308 nm and subsequent reaction of the excited oxygen atom (O\textsuperscript{1}D) with water vapor to form OH radicals. Since the sampled gas in the detection volume is exchanged between two subsequent laser shots, this effect is minimized. A small background may be still present. Measurements by FZJ-LIF-ambient were corrected for this interference (0.07 pptv per 50 ppbv O\textsubscript{3}), much smaller than HO\textsubscript{2} concentrations encountered during this campaign. All other instruments assumed that this potential interference was negligible.

The calibration of all LIF instruments is accomplished by producing OH and HO\textsubscript{2} radicals by water photolysis at 185 nm in a flow tube, which can be mounted on top of the fluorescence cells (Aschmutat et al., 1994; Schultz et al., 1995). Humidified zero air is irradiated by a low pressure mercury lamp before part of the excess air stream is sampled by the instrument. Flow rate and shape of the flow tube is either chosen to ensure laminar flow condition (FRCGC-LIF, FZJ-LIF) or turbulent mixing (MPI-LIF). Water photolysis in zero air at 185 nm results in the production of equal concentrations of OH and HO\textsubscript{2}. The radical concentration sampled by the instruments depends on the water concentration, the intensity of the 185 nm radiation, duration of irradiation and potential loss of radicals before the air is sampled by the instruments. The water concentration is monitored during calibrations.

Four calibration measurements are taken, in order to calculate the HO\textsubscript{2} sensitivity of the FRCGC-LIF. NO in the fluorescence cell and CO in the calibration source is switched on and off for the different calibration modes (see Kanaya et al., 2001 for details). Light intensity and duration of irradiation is determined by ozone actinometry.
This is done by direct measurement of the ozone concentration in the centerline of the radical source, when the calibration source is not on top of the fluorescence cell for the FRCGC-LIF.

For calibrating the \( \text{HO}_2 \) sensitivity of the FZJ-LIF, OH is quantitatively converted to \( \text{HO}_2 \) by adding excess CO to the calibration gas in the radical sources, so that the sensitivity is given by one measurement. Like for the FRCGC-LIF ozone actinometry gives a measurement of the irradiation parameters of the 185 nm light. However, this is accomplished indirectly by observing the light intensity measured by a phototube, which was calibrated against the ozone production, so that changes in the radical production could be detected during calibration measurements.

The calibration factor for the \( \text{HO}_2 \) sensitivity of the MPI-LIF is calculated from the difference between two calibration measurements, when (1) only OH is detected (NO in the cell turned off) and (2) the sum of OH and \( \text{HO}_2 \) is measured (NO in the cell turned on) instrument. The radical source of the MPI-LIF instrument was characterized by \( \text{N}_2\text{O} \) actinometry before and after the campaign and was found to having been stable for this period.

3 Experiments

The campaign was divided into two parts. During the first three days (9–11 July), the instruments were placed on the paved area between the institute building and the SAPHIR chamber at the Forschungszentrum Jülich, Germany, in order to compare measurements from ambient air sampling. The distance between instruments was approximately 3.2 m (MPI-LIF and FRCGC-LIF) and 4.5 m (FRCGC-LIF and FZJ-LIF-ambient). The sampling height was 3.5 m for all instruments. Trace gas concentrations of NO, NO\(_2\), HONO, O\(_3\), H\(_2\)O, HCHO, VOCs and photolysis frequencies were monitored on site as indicated in Figure 1.

The area is characterized by small buildings, grassland and trees. The Forschungszentrum is surrounded by forest, agricultural areas and main roads.
Meteorological conditions were similar throughout the three days of measurements with northerly winds and moderate summer temperatures with a maximum of 28°C. There was ground fog in the morning and scattered clouds in the afternoon on the first day. The sky was nearly cloud free on the second and third day until a rainstorm evolved in the afternoon of the third day.

Diurnal profiles of trace gases were similar for all three days of ambient air sampling (Fig. 2). Local emissions from traffic within the Forschungszentrum and nearby roads caused high NOx mixing ratios up to 30 ppbv in the morning hours. The CO mixing ratio was almost constant during the campaign (200–300 ppbv) with two short peaks up to 320 ppbv indicating emissions from trucks passing the road next to the instruments. Dominant VOC species were benzene and toluene (mixing ratios 1 ppbv each), indicating anthropogenic emissions. Isoprene mixing ratios were mostly between 0.3 and 0.6 ppbv with peak values up to 2 ppbv in the late afternoon. The ozone mixing ratio exhibited a typical diurnal cycle with peak values of 65 ppbv in the early afternoon and minimum values of 15 to 20 ppbv after midnight.

The instruments were attached to the SAPHIR chamber for six experiments (17–19, 21–23 July), each of which lasted one day. The chamber has been described in more detail elsewhere (e.g. Bohn et al., 2005; Rohrer et al., 2005; Wegener et al., 2007). The chamber was successfully used in a number of intercomparison experiments. These experiments proved that the chamber can serve as a homogeneous source for e.g. OVOCs (Apel et al., 2008), OH (Schlosser et al., 2007, 2009) and NO2 (Fuchs et al., 2010).

The chamber consists of a double wall Teflon film (FEP) of cylindrical shape (length 18 m, diameter 5 m, volume 270 m³). The chamber is maintained at ambient temperature and a small overpressure is applied. A louvre-system can be opened, in order to expose the chamber to natural sunlight, which is well transmitted by the FEP film (transmission of 85% for UV-VIS). The chamber can be filled with ultra pure synthetic air (Linde, quality 6.0), or with polluted ambient air. In both cases, trace gases can be added in known quantities. Turbulent mixing takes place when the chamber is exposed
to sunlight. In addition, a fan can be operated, in order to ensure rapid mixing for example in the dark chamber. OH and HO$_2$ radicals are formed in the chamber by the photolysis of nitrous acid (HONO) and formaldehyde (HCHO), respectively, both of which are presumably produced by photolytic surface reactions on the Teflon film (Rohrer et al., 2005).

Chemical conditions during the experiments are summarized in Table 2. The chamber was prepared for daily experiments by flushing out all trace gases. First, Milli-Q water was evaporated and added to the purge flow to adjust the humidity in the chamber on all days with the exception of the first experiment, when measurements started with sampling dry air. The roof of the chamber was periodically opened and closed for approximately one hour each during the first three experiments (Fig. 3). Concentrations of water vapor (17 July), NO$_x$ (18 July) and O$_3$ (19 July), respectively, were increased stepwise, in order to investigate the influence of these compounds on the performance of the instruments. The fan was operated during the injection of trace gases, which took place during the dark periods of experiments. In addition, CO (mixing ratio up to 0.8 ppmv) was added to the chamber air on 18 July. Ambient air was filled into the chamber before the chamber roof was opened for one experiment (21 July, Fig. 4). On this day CO was injected into the chamber at 11:00 (CO mixing ratio 500 ppmv), in order to convert OH to HO$_2$ quantitatively. The ozonolysis of alkenes (1-pentene, trans-butene) were investigated in the dark chamber on 22 July. This experiment allowed to produce approximately constant HO$_2$ concentrations, which were adjusted by a stepwise increase of alkene concentrations during the experiment. The oxidation of a mix of VOCs (5 ppbv benzene, 3 ppbv 1-hexene, 2.5 ppbv m-xylene, 3 ppbv n-octane, 3 ppbv n-pentane, and 1 ppbv isoprene) was investigated in the sunlit chamber during the last experiment (23 July).
4 Results

4.1 Diurnal profiles

During the ambient air sampling period, HO₂ mixing ratios exhibited a typical diurnal profile (Fig. 2). In the morning, HO₂ was suppressed, when NO mixing ratios were high. HO₂ followed the diurnal profile of the radiation during the rest of the day with maximum HO₂ mixing ratios of approximately 25 to 35 pptv. NO mixing ratios were largest in the morning of the first day (12 ppbv), which was a working day and thus local rush-hour traffic was present. Smaller peak values of 2 to 6 ppbv were observed in the mornings of the following weekend. During both nights of the first part of the campaign instruments showed decreasing HO₂ mixing ratios over the course of the night. Mixing ratios (up to 4 to 10 pptv) were well above the detection limits of LIF instruments. This indicates that non-photolytic sources of HO₂ such as ozonolysis could have played a role in the night.

During nearly all chamber experiments, HOₓ was primarily produced by photolysis reactions. Precursors were HONO and HCHO, which were formed in the illuminated chamber, and ozone, which was added to the chamber air and which was photochemically produced over the course of the experiments (Figs. 3, 4). Because of the production of HOₓ by photolytical reactions, HO₂ increased rapidly, when the chamber roof was opened. Radical loss reactions led to a fast decrease of HO₂ once the chamber roof was closed. During periods, when the chamber was exposed to sunlight, HO₂ was well correlated with the pattern of the photolysis rates. In general, the HO₂ concentration was enhanced by trace gases that produce HO₂ such as ozone and water vapor (e.g. 17 and 19 July) and reduced by those which remove HO₂ such as NO (e.g. 18 July).

Only during the ozonolysis experiment on 22 July HO₂ was not formed primarily by photolytic reactions, but by the ozonolysis of 1-pentene (07:30–12:00) and trans-butene (after 12:10). Here, the HO₂ concentration was determined by the balance between production in ozonolysis reactions and radical loss reactions. During the first
part of the experiment, the consumption of ozone and 1-pentene was small on a time scale of an hour, so that nearly constant levels of HO$_2$ were achieved (Fig. 4). HO$_2$ concentrations were varied by a stepwise increase of the alkene concentration. At 12:10 four additional injections of trans-2-butene led to the complete consumption of ozone. HO$_2$ mixing ratios up to 240 pptv were reached, which decreased after each injection due to the decreasing ozone concentration.

4.2 HO$_x$ measurements

FRCGC-LIF and MPI-LIF instruments measured during both parts of the campaign. 505 and 18 805 data points were included in the analysis of the ambient air sampling period from the FRCGC-LIF and MPI-LIF, respectively, and 749 and 16 322 data points for SAPHIR experiments. The FZJ-LIF-ambient instrument measured during the ambient air sampling period (760 data points). The FZJ-LIF-SAPHIR instrument, which was operated during SAPHIR experiments, provided 2582 data points. The different number of total data points were mainly due to differences in the temporal resolution of measurements. FRCGC measured alternately OH and HO$_2$ for 73 s each during the ambient air sampling and the first two SAPHIR chamber experiments. The integration time was shortened to 51 s on 19 July for the rest of the campaign. Short periods between the two measurements were used to measure the background signal and to scan and to lock the wavelength, so that the time resolution was further reduced. The integration time of the FZJ-LIF instrument was varied to improve the detection limit during periods when HO$_2$ and OH concentrations were small. The time resolution ranged between 47 and 355 s (mean 100 s) for ambient air sampling and 24 and 74 s (mean 36 s) for SAPHIR experiments. The time resolution of data from the MPI-LIF instrument was 10 s during the whole campaign.

All LIF instruments ran continuously with two exceptions: (1) FCGCG-LIF was not operated during the first night of ambient air sampling and (2) data from the FZJ-LIF-ambient instrument were invalid between 09:10 and 14:25 on 10 July because of an instrumental failure. Furthermore, data from SAPHIR experiments were also excluded.
from the analysis for all instruments, when a high flow of zero air was used as carrier
gas to add water vapor to the chamber air. During these periods inhomogeneity of
trace gas concentrations in the chamber could have occurred.

The data are averaged to a 1 min time grid for correlation and regression analysis,
when the instrument provides a higher time resolution. For each averaged data point
the standard deviation and the error from the error propagation of the single values are
compared and the larger of both is assigned as an error bar. Figures 6 and 7 show
all 1 min HO\(_2\) concentrations that are included in the following analysis for ambient
sampling and SAPHIR experiments, respectively.

Part of the data from the SAPHIR experiment on 22 July are treated separately
in the analysis, because the HO\(_2\) mixing ratio was increased to maximum values to
very high values (approximately 200 pptv) exceeding all concentrations encountered
during the other experiments. The correlation and regression analysis is sensitive to
the dynamic range of the data set and therefore this small data subset would dominate
the results. In order to avoid that the analysis of the complete data set is biased by
potential artifacts in the data during this short period, these data are treated separately.
As seen in Fig. 6 deviations between ambient HO\(_2\) concentrations are significantly
larger during the second night than observed during the day. Therefore, nighttime data
are discussed separately. Correlation and regression analysis of ambient data are
restricted to daytime data.

4.3 Precision of measurements

The statistical measurement errors specified by each group are checked against the
observed measurement precision when the HO\(_2\) concentration was nearly constant.
This was achieved during the ozonolysis experiment on 22 July (between 1 and
25 pptv), when HO\(_2\) shows only slow changes during several 30 min time intervals.
A linear fit is applied to the data at their original time resolution. No drift or systematic
deviation from a linear function is observed in the fit residual. Accordingly, the standard
deviation of the fit residual gives an estimate of the variability of measurements and is compared to the mean of error bars.

Figure 5 shows the frequency distribution of the residual, which is well described by a gaussian error distribution. The width of the gaussian function agrees with the mean of measurement errors (indicated by the red lines in Fig. 5) for each instrument. This means that error bars of all instruments give a realistic estimate of the measurements precision on a time scale of 30 min.

### 4.4 Statistical analysis

The linear correlation between measurements of the different LIF instruments is generally high (Table 3 and Figs. 8, 9). During ambient air sampling all LIF instruments show similar diurnal and nocturnal profiles (Fig. 6) demonstrated by linear correlation coefficients, $R^2$, within the range of 0.98 for FZJ-LIF and FRCGC-LIF and 0.92 for MPI-LIF and FZJ-LIF. Similar values are reached for the combined data set of all SAPHIR experiments with 0.82 and 0.93 for MPI-LIF and FRCGC-LIF and FZJ-LIF and FRCGC-LIF, respectively. However, values of linear correlation coefficients are more variable from day-to-day, ranging between 0.99 for all instruments on 22 July and 0.62 for MPI-LIF and FRCGC-LIF on 17 July. The $R^2$ values are higher (0.96–0.99) during the last three experiment days (21–23 July) when significant amounts of VOC were present in the chamber. Not only is the similarity of the single diurnal profiles better for these experiments, but also the absolute agreement of the three LIF instruments (Fig. 9).

The scatter plots in Figs. 8 (ambient air) and 9 (SAPHIR) emphasize the general high correlation between the data of the LIF instruments, but also show that the data pairs of the instruments cannot be described by a single linear relationship for all days. It is evident that the overall spread of data is not completely represented by the assigned statistical error bars, but systematic effects seem also to play a role. For example, a clear separation of data points (indicated by color code) can be seen in Fig. 9 for those experiments at SAPHIR (e.g., 17 July) which exhibit relatively low $R^2$ values. However, the scatter plot shows also a narrow distribution for single experiments (e.g. 18 July).
The data of the three instruments can be further compared by a linear regression analysis. The resulting fit parameters are given in Table 4 and the fit functions are plotted in Figs. 8 and 9 (gray solid lines). Here, the fit procedure from Press et al. (1992) (FitExy procedure) is used, which accounts for errors in both coordinates. This makes the result invariant of the choice of reference, so that an exchange of the dependent and independent variable gives the inverse result (not shown in Table 4). The large ratios of the sum of squared residuum values and the number of data points, demonstrate that statistical errors of data do not cover the deviations from a linear relationship. For this reason, meaningful errors of the fit parameters cannot be derived by error propagation. However, as shown above, error bars are adequate on a time scale of at least 30 min. Thus, other effects on a longer time scale must have influenced the measurements (see below). Regression parameters can give a hint for systematic differences between calibration factors of instruments.

The linear regression of ambient air data results in slopes of 1.46 for FRCGC-LIF versus MPI-LIF, 0.59 for MPI-LIF versus FZJ-LIF and 1.19 FZJ-LIF versus FRCGC-LIF. Larger deviations between measurements of FZJ-LIF and FRCGC-LIF instruments occur only during the second night, when FZJ-LIF and MPI-LIF instruments measured HO$_2$ mixing ratios up to 10 pptv, whereas FRCGC-LIF measurements show only 3 pptv (Fig. 6). These data behaved systematically different from the daytime data (see color distinction in Fig. 8) and were therefore excluded from the statistical fit. During this night the FZJ-LIF was operated with the additional RO$_x$ converter. However, this was continued during the following day when measurements between FZJ-LIF and FRCGC-LIF were in good agreement like the days before. For data from SAPHIR experiments, slopes of the linear fits are 1.26 (FRCGC-LIF versus MPI-LIF), 1.19 (MPI-LIF versus FZJ-LIF) and 0.69 (FZJ-LIF versus FRCGC-LIF). Overall, the slopes of the regression lines indicate that the calibrations of the instruments were different by factors between 1.2 to 1.7 for the conditions in ambient air and SAPHIR. These factors can only partly be explained by the combined 1 $\sigma$ accuracies of the calibrations (Table 1).
Exceptionally high HO$_2$ concentrations were produced during the second half of the ozonolysis experiment on 22 July by the addition of large concentrations of various alkenes. Although these data are excluded from the analysis, the comparison of them does not give significantly different results (not shown here) compared to the results from the first part of the experiment. This indicates that the sensitivity of the LIF instruments is constant over a wide dynamic range of HO$_2$ concentrations encountered during this experiment.

The offset values of the regression lines are all very small. Their amount is less than 1.3 pptv, which is generally negligible compared to ambient HO$_2$ concentrations. Thus, the regression analysis gives no indication of a general instrumental offset problems that may bias atmospheric HO$_2$ measurements. However, during dark periods of the experiments measurements by MPI-LIF are significantly higher than those by the other two instruments (see discussion below).

5 Discussion

The above results show that the agreement of measurements by the three LIF instruments is variable. The agreement for ambient air is different from the result for SAPHIR experiments (Table 4) and the agreement varies from day to day between different SAPHIR experiments (Fig. 9). Since no absolute HO$_2$ reference is available for comparison, it is not possible to assess absolute concentrations. Only relative deviations can be discussed. Systematic differences between measurements can occur for different reasons:

1. instruments may sample air with different composition (inhomogeneous environment)

2. instrumental calibrations can have systematic errors

3. calibrations may lack of reproducibility
4. intrinsic instrument sensitivities may be variable
5. detection sensitivities may have unknown dependencies on chemical conditions
6. chemical interferences may cause artificial HO\textsubscript{2} signals

The first four reasons are unlikely explanations for the observed HO\textsubscript{2} differences in the SAPHIR chamber, where the same LIF instruments and a DOAS instrument, which provided calibration-free OH concentrations, showed good absolute agreement, within 13\% for OH measurements (Schlosser et al., 2009). One major conclusion of the OH comparison was that the chamber air can be considered to be homogeneously mixed for OH. This is also expected for the less reactive HO\textsubscript{2}. Furthermore, the comparison of OH concentrations from LIF instruments versus the DOAS technique at SAPHIR demonstrated that the LIF calibration sources for OH are accurate and reproducible (Schlosser et al., 2007, 2009). The same radical sources were also used for calibrations of HO\textsubscript{2} sensitivities during HOxComp. The only difference was that CO was added to the calibration gas to convert OH to HO\textsubscript{2} in the radical sources of FZJ-LIF and FRCGC-LIF (see instrument description above). Since the comparison of OH measurements (Schlosser et al., 2009) does not exhibit a day-to-day variability as observed for HO\textsubscript{2}, the conversion of OH to HO\textsubscript{2} in the radical sources of FZJ-LIF and FRCGC-LIF would need to be variable, but complete conversion is ensured by the large excess of CO.

Unstable detection sensitivities due to technical problems of the LIF instruments are also an unlikely explanation for the observations. Similar to the calibration many parts of the instruments that are required for the HO\textsubscript{2} detection are shared with those needed for the OH detection such as the laser used for the excitation of OH. In the FRCGC-LIF instrument even the whole detection cell is the same for OH and HO\textsubscript{2}. Again, the good agreement between measurements of OH concentrations (Schlosser et al., 2009) makes it unlikely that instrumental parameters such as laser performance caused the variability in the agreement of HO\textsubscript{2} mixing ratios. The main difference to the OH detection is the addition of excess NO upstream of the laser excitation, in order
to convert HO\(_2\) to OH. Technical details how the addition is accomplished are different between instruments, but there is no reason to assume that the conversion efficiency exhibited a variability that does not show up in the variability of the daily calibration.

Most likely chemical conditions during experiments affected either the HO\(_2\) detection efficiency or caused interfering signals. As shown in the following, there are two species which correlate with differences in the HO\(_2\) mixing ratios measured by the LIF instruments: (1) ozone and (2) water vapor.

### 5.1 Influence of ozone during SAPHIR experiments

The influence of ozone was studied in the experiment on 19 July by varying ozone stepwise from 0 to 150 ppbv at a relatively constant water vapor mixing ratio of 1.0–1.4%. The measurements by the three LIF instruments agree well when the chamber was illuminated, but an increasing difference between the HO\(_2\) measurements by MPI-LIF and the other two LIF instruments is observed in the dark chamber at increasing ozone concentrations (Fig. 7). The additional HO\(_2\) in the MPI-LIF data reaches about 5 pptv at 150 ppbv ozone relative to FZJ-LIF or FRCGC-LIF during the dark periods, but there is no indication for such an offset in the MPI-LIF data in the illuminated chamber. This is evident in the correlation plot in Fig. 10. A high correlation as well as good absolute agreement and a negligible offset (1 pptv) is observed in the correlation of MPI-LIF versus FZJ-LIF, when the data from the dark periods are excluded, whereas FZJ-LIF and FRCGC-LIF correlate well both for illuminated and dark conditions.

Systematic differences between the measurements by MPI-LIF and the other two instruments are also observed in the darkness during other SAPHIR experiments, but the absolute deviations vary for similar ozone mixing ratios on different days (e.g. 21 July during two periods, when the chamber roof was closed (09:15 to 10:00 and 12:00 to 15:00). This indicates that this effect does not only depend on the ozone mixing ratio. There is no offset of HO\(_2\) mixing ratios by MPI-LIF during the ozonolysis of alkenes on 22 July that was also carried out in the dark chamber. The relationship between data is similar to the relationship observed for the other experiments at similar water
concentration. The intercept of the linear fit between data by MPI-LIF and the other two LIF instruments is within the range of intercepts calculated for the other experiments (approximately 1 pptv). The difference between the ozonolysis experiment and all the other experiments is that there are no species in the chamber that are photolytically produced including nitrogen oxides.

The potential interference from self-production of OH from ozone photolysis (see instrument description) is typically small and independent of experimental conditions like darkness, so that this cannot explain the observations. A detailed analysis, which factors affect the sensitivity of the MPI-LIF instrument in the dark beside the observed dependence on ozone, is not possible from measurements of this campaign. As mentioned in Schlosser et al. (2009) OH concentration measurements by MPI-LIF at periods without daylight were rejected, because of an unexplained variable OH background signal. One may speculate that the background signal for both, OH and HO$_2$ detection, has the same source.

5.2 Influence of water vapor during SAPHIR experiments

The largest differences between diurnal profiles measured by the LIF instruments are encountered during the first SAPHIR experiment, when water vapor was stepwise increased to a maximum mixing ratio of 1.8%. Whereas HO$_2$ mixing ratios by FRCGC-LIF are the highest for dry conditions and the lowest for maximum water concentration, HO$_2$ mixing ratios by FZJ-LIF are higher for dry conditions and become similar to those by MPI-LIF for high water vapor concentrations. This behavior is also observed during the other experiments.

In order to investigate the water effect quantitatively, the data from all SAPHIR experiments (17–23 July) are grouped in four classes of water vapor concentrations (according to the water vapor concentrations encountered during the experiment on 17 July, when the water vapor concentration was investigated systematically). For each subgroup, a good linear correlation is found (Fig. 11). Linear fits are then applied to the data pairs of each water vapor class. A data filter is applied, excluding time periods
when the ozone mixing ratio was larger than 30 ppbv in the dark chamber on 18, 19, 21, 23 July. With increasing water vapor concentration slopes of the fits are increasing from 0.35 to 1.26 for measurements by FRCGC-LIF and MPI-LIF, and decreasing from 2.2 to 1.1 and 1.5 to 0.7 for measurements by MPI-LIF and FZJ-LIF, and FZJ-LIF and FRCGC-LIF, respectively.

Apparently, the discrepancies between HO$_2$ measurements are largest (up to a factor 3) at low water vapor mixing ratios (0–0.1% and 0.25–0.6%), while absolute agreement is better (within 30%) at high humidities (0.6–1.2% and 1.2–1.8%). In the latter case, the calibrations of all three instruments agree within their stated 1$\sigma$ accuracies. Furthermore, the ratio of the sum of squared residual and number of data points decreases, if data are restricted to conditions when the water vapor mixing ratio was higher than 0.6% (SAPHIR$^*$ in Table 4), demonstrating that part of the variability indeed depends on the water vapor concentration.

The strong water vapor dependence of the correlation between the LIF measurements is surprising, because the HO$_2$ concentration measurements were corrected by each group for water vapor dependencies prior to data submission. Corrections made for measurements of the FRCGC-LIF and FZJ-LIF instruments take into account the well-known effect of fluorescence quenching of the excited OH radical by water vapor.

A loss of detection sensitivity by 10% can be calculated from published quenching constants for an increase of the water vapor mixing ratio from 0 to 1%. During the campaign, calibrations of the FZJ-LIF-SAPHIR instrument were performed for different water vapor mixing ratios between 0.4 and 1.2%. No water vapor dependence larger than the expected quenching effect (11%) was observed. Since the reproducibility of calibration measurements was within the same range, correction factors were derived theoretically from quenching constants rather than from the calibration measurements. In case of the FRCGC-LIF instrument, calibrations were done at low water vapor mixing below 0.1%. This was necessary, in order to produce radical concentrations in the lower pptv range in the FRCGC-LIF radical source. Measurements at higher water vapor mixing ratios were then corrected for fluorescence quenching calculated
from quenching constants. The validity of this assumption was tested in laboratory experiments. The sensitivity of the MPI-LIF instruments was investigated during the campaign by calibration measurements at different water levels and showed a larger reduction of the sensitivity than expected from fluorescence quenching (i.e. 20% reduction per 1% water vapor concentration). This empirical correction factor was applied to the measurements by MPI-LIF.

The large differences between measurements at low water vapor concentrations cannot be explained by a lack of characterization of the sensitivity with respect to water vapor for several reasons: (1) The validity of the corrections that are applied for fluorescence quenching was demonstrated by the intercomparison of OH measurements (Schlosser et al., 2009). The same corrections are applied for OH and HO₂ measurements. (2) The largest corrections are done for measurements by MPI-LIF, but they are significantly smaller than the differences between measurements observed for conditions with low water vapor concentration. (3) Calibrations of the FRCGC LIF instrument were performed at low water vapor mixing ratios (<0.1%), and MPI-LIF and FZJ-LIF were calibrated over a range of water vapor that includes the mixing ratios encountered during the campaign.

Despite these calibration conditions, measurements of all three instruments show significant disagreement at low humidities. This result suggests that an additional unknown parameter influences the HO₂ detection sensitivity at dry conditions in the chamber experiments, but not in the calibrations. The measurement differences are similar for each pair of instruments. For this reason and due to the lack of an absolute reference, it is not possible to decide which instruments are affected by the unknown parameter. At high humidities (>0.6%), however, all instruments agree within the stated accuracies and exhibit very good linear correlations. It should also be noted that the LIF instruments were mostly deployed for such high water vapor mixing ratios in field campaigns in the lower troposphere in the past, and that one of the instruments (FZJ-LIF) had shown very good agreement with the absolute measurements by MIESR at similar conditions in the field (Platt et al., 2002) and in SAPHIR (Fuchs et al., 2009).
5.3 Comparison of ambient air measurements

During ambient air measurements of the campaign, water vapor concentrations were approximately constant (around 1%) and ozone concentrations were low during the night (between 10 and 30 ppbv). Therefore, the observation of a higher linear correlation between measurements can be expected considering the magnitude of the interferences found in the analysis of the SAPHIR experiments. Whereas a similar slope in the linear fit between data by MPI-LIF and FRCGC-LIF is calculated for ambient air sampling and SAPHIR experiments, which were carried out with a comparable water concentration, HO\(_2\) mixing ratios by FZJ-LIF-ambient are significantly smaller compared to the relationship observed for FZJ-LIF-SAPHIR. Good agreement between measurements by FZJ-LIF-ambient and FRCGC-LIF is observed, but HO\(_2\) mixing ratios by FRCGC-LIF are approximately 30% smaller than those by FZJ-LIF-SAPHIR.

In principle, the different agreement of the instruments under ambient conditions compared to the SAPHIR experiments can have been caused by several reasons. First, the FZJ-LIF data in ambient and SAPHIR air were obtained by two LIF instruments with potentially different systematic measurement errors. This explanation, however, is unlikely, because the two detection systems are constructed in the same way and both shared the same calibration source. Second, all LIF instruments applied different calibration factors during ambient air sampling and SAPHIR measurements. Third, the complex chemical composition of ambient air may have caused additional interferences. However, it is noted that ambient air with a water vapor mixing ratio of about 1% was introduced in the SAPHIR chamber on 21 July. The agreement of the LIF instruments during this particular experiment was not significantly different from the other SAPHIR experiments at similar humidity. Lastly, it is possible that the instruments sampled ambient air of different chemical composition. In fact, the intercomparison of OH measurements during HOxComp has shown worse agreement and less correlation in ambient air compared to the measurements in the SAPHIR chamber. This observation indicates that the ambient air was inhomogeneously mixed (Schlosser et al., 2009),
which may explain at least some of the differences in the measured HO$_2$ concentrations.

### 6 Conclusions

This was the first formal, blind intercomparison of HO$_2$ measurements. Three instruments measured HO$_2$ concentrations via chemical conversion to OH, which was detected by laser-induced fluorescence. The measurements included three days of sampling ambient air and six experiments at the atmosphere simulation chamber SAPHIR in Jülich, Germany.

HO$_2$ concentrations measured by the three LIF instruments are linearly correlated as demonstrated by the range of linear correlation coefficients between 0.82 and 0.98. Although differences between measurements are within the range the combined accuracies of measurements for the entire data set from SAPHIR experiments, there are larger differences during distinct periods of the experiments. The relationship between measurements by the different LIF instruments is variable on a time scale of hours and depends on conditions of the experiments. This cannot be explained by the variability of the instrument sensitivities, because (1) the same calibration factors were applied for several experiment days and the entire period of ambient air sampling, respectively, and (2) the OH sensitivity of instruments was calibrated with the same radical sources, but differences between OH concentrations (Schlosser et al., 2009) do not change in the same way as observed for HO$_2$.

Two chemical species are identified, whose concentrations correlate with changes in the relationship between data sets during SAPHIR experiments. First, the linear correlation between data can be grouped into periods when the water vapor concentration was similar in the chamber. However, the dependence of the instrument sensitivity (mainly caused by fluorescence quenching) on water vapor was well characterized for this campaign. The validity of the correction applied was demonstrated by the intercomparison of OH concentrations measured by the same instruments which does
not exhibit a dependence on water vapor (Schlosser et al., 2009). Furthermore, the observed differences between measurements are larger than the correction factors. Thus, an unknown factor that is related to water vapor in the chamber may have influenced the HO$_2$ instrument sensitivities or may have caused an unknown interference. The results of the linear regression between data sets are significantly improved, if a subset of the data defined by the following conditions is used: (1) the water vapor mixing ratio was > 0.6%, (2) presence of daylight, (3) when there was no daylight the ozone mixing ratio was < 30 ppbv or no photolytically produced species were present. These findings recommend a reevaluation of the dependence of water vapor on the sensitivity of LIF instruments especially for small concentrations and the investigation of potential interferences which may be correlated with water vapor in the SAPHIR chamber. The agreement of measurement is improved for water vapor mixing ratios within the range of concentrations which are typically encountered in the lower troposphere, where many of the HO$_2$ measurements have been done so far.

Second, in one of the LIF instruments (MPI-LIF) a background signal, which is correlated to the ozone concentration in the chamber, is observed during periods without sunlight. The magnitude of this interference does not only depend on the ozone concentration, but is variable for the different experiments. It does not appear during the ozonolysis experiment, when no photolytically produced species were in the chamber. It is known that ozone can cause an interference by its photolysis by the exciting laser beam at 308 nm, but this effect would not depend on the presence of sunlight outside the measurement cell and is expected to be smaller than the signal observed here. The unexplained HO$_2$ background corresponds to the observation of an OH background of this instrument at the same time (Schlosser et al., 2009) and may have the same origin.

The results from SAPHIR experiments are supported by the good correlation between ambient air data. However, the regression analysis results in slopes that are different from slopes observed for SAPHIR experiment and that are partly not within the combined 1σ accuracies of instruments. The data set of ambient air was most likely influenced by inhomogeneities in air masses sampled by the instruments at slightly
different locations leading to deviations between measurements. This emphasizes that it is essential for instrument comparisons that all instruments sample the same air. A simulation chamber like SAPHIR provides an environment which ensures that instruments can sample homogeneously mixed air containing the same trace gas concentrations (Schlosser et al., 2007, 2009; Apel et al., 2008; Fuchs et al., 2010).

The results of this study indicate the need to study the influence of atmospheric components such as water vapor or ozone on the instrument sensitivity under field conditions. Further systematic laboratory and chamber studies may be useful to resolve open questions identified here. Future intercomparison efforts using different instruments could greatly aid to further improve the accuracy and reliability of HO₂ measurements.

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References

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Table 1. Properties of the LIF instruments regarding the HO₂ detection.

<table>
<thead>
<tr>
<th>Property</th>
<th>MPI-LIF</th>
<th>FZJ-LIF</th>
<th>FRCGC-LIF</th>
</tr>
</thead>
<tbody>
<tr>
<td>cell assembly</td>
<td>HO₂-cell downstream of OH-cell</td>
<td>separate cell for HO₂</td>
<td>same cell for OH and HO₂</td>
</tr>
<tr>
<td>distance nozzle - detection</td>
<td>33 cm</td>
<td>10 cm</td>
<td>23 cm</td>
</tr>
<tr>
<td>distance NO addition - detection</td>
<td>10 cm</td>
<td>5.5 cm</td>
<td>21 cm</td>
</tr>
<tr>
<td>cell pressure</td>
<td>3.7 hPa</td>
<td>3.8 hPa</td>
<td>2.9 hPa</td>
</tr>
<tr>
<td>conversion reaction time</td>
<td>6 ms</td>
<td>0.2 ms</td>
<td>no estimate</td>
</tr>
<tr>
<td>conversion efficiency</td>
<td>&gt; 90%</td>
<td>&gt; 90%</td>
<td>&gt; 90%</td>
</tr>
<tr>
<td>NO concentration</td>
<td>0.09%</td>
<td>0.6%</td>
<td>0.04%</td>
</tr>
<tr>
<td>NO purification</td>
<td>Ascarite</td>
<td>Ascarite</td>
<td>Ascarite</td>
</tr>
<tr>
<td>interference from NO addition</td>
<td>&lt; 0.3 pptv, (not corrected)</td>
<td>0.2–0.3 pptv</td>
<td>0.16–27 pptv (not corrected)</td>
</tr>
<tr>
<td>dependence of sensitivity on water</td>
<td>from lab. experiments</td>
<td>fluorescence quenching</td>
<td>fluorescence quenching</td>
</tr>
<tr>
<td>1σ accuracy</td>
<td>± 16%</td>
<td>± 10%</td>
<td>±24%</td>
</tr>
</tbody>
</table>
Table 2. Chemical conditions during the experiments of the HOxComp campaign. SAPHIR experiments on 17–19 and 22–23 July were performed in zero air with added reactants. All mixing ratios are maximum values reached during the experiments.

<table>
<thead>
<tr>
<th>date</th>
<th>NO/ppbv</th>
<th>NO$_2$/ppbv</th>
<th>O$_3$/ppbv</th>
<th>CO/ppmv</th>
<th>H$_2$O/hPa</th>
<th>j(O$_1^D$)/10$^{-5}$ s$^{-1}$</th>
<th>experiment</th>
</tr>
</thead>
<tbody>
<tr>
<td>ambient</td>
<td>9-11</td>
<td>13</td>
<td>20</td>
<td>65</td>
<td>0.3</td>
<td>21</td>
<td>2.5</td>
</tr>
<tr>
<td>SAPHIR</td>
<td>17</td>
<td>0.56</td>
<td>2.8</td>
<td>28</td>
<td>0.02</td>
<td>20</td>
<td>1.7</td>
</tr>
<tr>
<td></td>
<td>18</td>
<td>1.7</td>
<td>8</td>
<td>50</td>
<td>0.8</td>
<td>12</td>
<td>1.5</td>
</tr>
<tr>
<td></td>
<td>19</td>
<td>0.34</td>
<td>4</td>
<td>150</td>
<td>0.02</td>
<td>15</td>
<td>1.5</td>
</tr>
<tr>
<td></td>
<td>21</td>
<td>5</td>
<td>14</td>
<td>47</td>
<td>500$^a$</td>
<td>14</td>
<td>1.5</td>
</tr>
<tr>
<td></td>
<td>22</td>
<td>b</td>
<td>b</td>
<td>100</td>
<td>0.02</td>
<td>30</td>
<td>c</td>
</tr>
<tr>
<td></td>
<td>23</td>
<td>0.25</td>
<td>1</td>
<td>45</td>
<td>0.02</td>
<td>11.5</td>
<td>1.4</td>
</tr>
</tbody>
</table>

$^a$ addition at 11:00, was 150 ppbv before
$^b$ no NO$_x$ addition
$^c$ dark chamber
$^d$ 1-pentene, trans-butene
$^e$ benzene, 1-hexene, m-xylene, n-octane, n-pentane, isoprene
Table 3. Linear correlation coefficient $R^2$ between data measured by different LIF instruments. Results are shown for measurements in ambient air and in chamber experiments (SAPHIR). Numbers in brackets are the number of valid data points (1 min average). Nighttime data from ambient measurements and data after noon on 22 July were excluded (see text for details).

<table>
<thead>
<tr>
<th>Date</th>
<th>MPI-FRCGC</th>
<th>MPI-FZJ</th>
<th>FZJ-FRCGC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ambient daytime 9–11 July</td>
<td>0.97 (334)</td>
<td>0.92 (506)</td>
<td>0.98 (81)</td>
</tr>
<tr>
<td>SAPHIR 17 July</td>
<td>0.62 (85)</td>
<td>0.84 (355)</td>
<td>0.89 (79)</td>
</tr>
<tr>
<td>18 July</td>
<td>0.76 (92)</td>
<td>0.75 (413)</td>
<td>0.91 (82)</td>
</tr>
<tr>
<td>19 July</td>
<td>0.94 (109)</td>
<td>0.92 (406)</td>
<td>0.97 (99)</td>
</tr>
<tr>
<td>21 July</td>
<td>0.96 (123)</td>
<td>0.97 (295)</td>
<td>0.98 (87)</td>
</tr>
<tr>
<td>22 July</td>
<td>0.99 (101)</td>
<td>0.99 (328)</td>
<td>0.99 (101)</td>
</tr>
<tr>
<td>23 July</td>
<td>0.98 (115)</td>
<td>0.99 (404)</td>
<td>0.99 (125)</td>
</tr>
<tr>
<td>SAPHIR 17–23 July</td>
<td>0.82 (625)</td>
<td>0.92 (2201)</td>
<td>0.93 (573)</td>
</tr>
<tr>
<td>SAPHIR* 17–23 July</td>
<td>0.97 (386)</td>
<td>0.98 (1347)</td>
<td>0.98 (362)</td>
</tr>
</tbody>
</table>

SAPHIR*: $\text{H}_2\text{O} > 0.6\%$ (all experiments) and $\text{O}_3 < 30 \text{ ppbv}$ in the dark (18, 19, 21, 23 July).
Table 4. Results of the linear regression analysis between HO2 data of the different instruments. $\frac{\chi^2}{N-2}$ is the sum of squared residuals divided by number of data points. Data are averaged to 1 min time intervals and the standard deviation is taken as error, unless the error propagation of the high resolution data was larger than the standard deviation. Nighttime data from ambient measurements and data after noon on 22 July were excluded (see text for details).

<table>
<thead>
<tr>
<th></th>
<th>x – y</th>
<th>intercept/pptv</th>
<th>slope</th>
<th>$\frac{\chi^2}{N-2}$</th>
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<tbody>
<tr>
<td>ambient daytime</td>
<td>FRCGC-MPI</td>
<td>1.31</td>
<td>1.46</td>
<td>23</td>
</tr>
<tr>
<td></td>
<td>MPI-FZJ</td>
<td>−0.39</td>
<td>0.59</td>
<td>11</td>
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SAPHIR*: same as in Table 3
Fig. 1. Location of instruments during the HOxCOMP campaign. Red dots mark the sampling points of instruments measuring HO$_2$ or OH. The HO$_2$ data in this work were delivered by FRCGC, MPI and FZJ. Numbers indicate positions of instrument inlets measuring: (1) NO$_x$, O$_3$, HCHO, VOC, H$_2$O, CO (red sign: ambient measurements, green sign: SAPHIR measurements); (2) temperature, relative humidity, HONO; (3) ultrasonic anemometer; (4) filter-radiometer; (5) O$_3$. The light path of an OH DOAS instrument is indicated in red within the chamber. A road (closed for traffic) is located to the southeast and the site is bordered in the north and west by bushes and trees (marked by a green line). Liquid nitrogen and oxygen is stored in two tanks northeast of the chamber.
Fig. 2. Time series of HO₂ mixing ratios during ambient air sampling from all instruments at their original time resolution together with the photolysis rate j(Ο¹D) (black line upper panels). In addition, mixing ratios of O₃ (UV absorption photometer), NO₂ and NO (chemiluminescence detector), and water vapor (dewpoint hygrometer) are shown in the lower panels.
Fig. 3. Same as Fig. 2 for the first three SAPHIR chamber experiments.
Fig. 4. Same as Fig. 2 for the last three SAPHIR experiments. In addition, mixing ratios of organic compounds are shown. The scaling of axis showing HO$_2$ mixing ratios changes during the experiment on 22 July, because HO$_2$ mixing ratios were rapidly increased to values, which are ten times higher than encountered during the rest of the campaign. On 21 July the CO mixing ratio was increased to 500 ppmv (not shown here). On 22 July dashed lines in the lower panel show calculated mixing ratios of 1-pentene and trans-butene. The time series is calculated from the times of the addition of alkenes and their dilution and absolute levels are scaled to measurements by GC-FID (dots).
Fig. 5. Frequency distribution of the residuum of a linear fit of measurements on 22 July (1 h periods when HO$_2$ decreased approximately linearly). The frequency distribution was fitted to a normal distribution (black line). The width agrees with the mean of measurement errors (red line) within the precision of the fit for all instruments.
Fig. 6. Time series of HO$_2$ mixing ratios during ambient air sampling. Data were averaged to a 1 min time resolution, if the original data set provided a higher time resolution.
Fig. 7. Same as Fig. 6 for all SAPHIR chamber experiments. Periods during which trace gases were mixed into the chamber with a high flow of zero air were excluded from the analysis (see e.g. 19 July).
Fig. 8. Correlation of \( \text{HO}_2 \) mixing ratios in ambient air on 9–11 July (1 min average). The dashed line is the 1:1 line. The solid lines result of linear fits to the daytime data, only. The scatter of the entire data sets is larger than expected from the precision of single data points.
Fig. 9. Correlation of \(\text{HO}_2\) mixing ratios measured during SAPHIR experiments on 17–23 July. 1 min averaged data are shown. The dashed line is the 1:1 line and the solid lines show the result of linear fits. Like for the ambient data the scatter of the entire data sets is larger than expected from the precision of single data points.
Fig. 10. Correlation of HO₂ mixing ratios on 19 July, when the ozone mixing ratio was varied. The water mixing ratio was 1.0–1.4%. Solid line shows the result of the linear fit. Data taken when the chamber roof was closed (crosses in Figure) were excluded from the fit.
Fig. 11. Correlation of HO$_2$ mixing ratios classified for different water vapor mixing ratios in SAPHIR. All data are included with the exception of data during periods with ozone mixing ratios greater than 30 ppbv in the dark chamber on 18, 19, 20, 23 July. Separate regression analysis were performed for each level of water concentration indicated by the different colors. Solid lines are results from a linear fit.