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- 1 Development of an instrument for direct ozone production rate measurements: Measurement
- 2 reliability and current limitations
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11 Abstract

- 12 Ground level ozone (O₃) is an important pollutant that affects both global climate change and regional
- 13 air quality, with the latter linked to detrimental effects on both human health and ecosystems. Ozone
- 14 is not directly emitted in the atmosphere but is formed from chemical reactions involving volatile
- 15 organic compounds (VOCs), nitrogen oxides (NO_x = NO+NO₂) and sunlight. The photochemical
- 16 nature of ozone makes the implementation of reduction strategies challenging and a good
- 17 understanding of its formation chemistry is fundamental in order to develop efficient strategies of
- 18 ozone reduction from mitigation measures of primary VOCs and NOx emissions.
- 19 An instrument for direct measurements of ozone production rates (OPR) was developed and deployed
- 20 in the field as part of the IRRONIC (Indiana Radical, Reactivity and OzoNe production
- 21 InterComparison) field campaign. The OPR instrument is based on the principle of the previously
- 22 published MOPS instrument (Measurement of Ozone Production Sensor) but using a different
- 23 sampling design made of quartz flow tubes and a different O_x (O₃ and NO₂) conversion/detection
- 24 scheme composed of an O₃-to-NO₂ conversion unit and a Cavity Attenuated Phase Shift (CAPS) NO₂
- 25 monitor. Tests performed in the laboratory and in the field, together with model simulations of the

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1 radical chemistry occurring inside the flow tubes, were used to assess (i) the reliability of the

2 measurement principle and (ii) potential biases associated to OPR measurements.

3 This publication reports the first field measurements made using this instrument to illustrate its

4 performance. The results showed that a photo-enhanced loss of ozone inside the sampling flow tubes

5 disturb the measurements. This issue needs to be solved to be able to perform accurate ambient

measurements of ozone production rates with the instrument described in this study. However, an

7 attempt was made to investigate the OPR sensitivity to NO_x by adding NO inside the instrument. This

8 type of investigations does not require measuring ambient OPR but only probing the change in ozone

9 production when NO is added. During IRRONIC, changes in ozone production rates ranging from the

10 limit of detection (3σ) of 6.2 ppbv h⁻¹ up to 20 ppbv h⁻¹ were observed when 6 ppbv of NO was added

11 into the flow tubes.

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12 1 Introduction

Ground-level ozone (O₃) is a primary constituent of photochemical smog that irritates the respiratory

system (WHO, 2013) and damages vegetation (Ashmore, 2005). In addition, ozone is a greenhouse

15 gas and an important precursor of the hydroxyl radical (OH), a key species controlling the

atmospheric oxidative capacity (Monks, 2005;Rohrer et al., 2014;Prinn, 2003). Ozone is a

17 photochemical pollutant formed during daytime and has an average lifetime estimated at 22±2 days

18 (Stevenson et al., 2006), which is long enough to transport it from polluted regions to remote areas

and between continents. The local production of ozone on top of the amount advected from elsewhere

can lead to exceedances of air quality standards in urbanized areas, making ozone pollution an issue

21 of global concern (Akimoto, 2003).

22 In the troposphere, ozone can be rapidly converted to nitrogen dioxide (NO₂) through reaction with

23 nitric oxide (NO), and back to O₃ through NO₂ photolysis. This chemistry does not produce new

24 ozone and is known as the O₃-NO_x PhotoStationary State (PSS), with NO_x being the sum of NO and

NO₂. The production of new ozone is driven by the oxidation of Volatile Organic Compounds

26 (VOCs), which leads to the production of hydroperoxy (HO₂) and organic peroxy (RO₂) radicals. The

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- 1 current understanding of tropospheric ozone chemistry indicates that new ozone is formed via
- 2 reactions of these peroxy radicals with NO, which results in the conversion of NO to NO₂ without
- 3 consumption of ozone (Monks, 2005; Seinfeld and Pandis, 2006).
- 4 When ozone is produced, reactions of peroxy radicals with NO also lead to the formation of OH,
- 5 which can then oxidize other molecules of VOCs to produce more peroxy radicals, and as a
- 6 consequence, more ozone. The propagation chemistry between RO_x (OH, HO₂ and RO₂) radicals,
- 7 which fuels ozone production, is terminated either by NO_x-RO_x reactions or by cross reactions of RO_x
- 8 radicals in NO_x-rich and NO_x-poor environments, respectively. These two types of termination
- 9 reactions lead to different regimes of ozone production referred as NO_x-limited or NO_x-saturated
- 10 when the rate of ozone production increases or decreases with NO_x, respectively. The turnover point
- 11 between the two regimes depends on NO_x concentrations, VOC reactivity, and radical production
- 12 rates (Kleinman, 2005). Since different air quality regulations have to be implemented for the two
- 13 different regimes, i.e either NO_x or VOC emission regulations, the understanding of the complex and
- 14 non-linear radical chemistry is key for the design of efficient emission control strategies.
- 15 The instantaneous ozone production rate, p(O₃), can be calculated from Equation (1) as the rate of
- 16 reactions between peroxy radicals and NO. The instantaneous ozone loss rate, l(O₃), can be calculated
- 17 using Equation (2), based on reaction rates for ozone photolysis, reactions of O₃ with HO_x and
- 18 alkenes, and the reaction of OH with NO₂, since NO₂ is a reservoir molecule for O₃. The net ozone
- 19 production rate, P(O₃), is then computed as the difference between instantaneous production and loss
- 20 rates (Eq. (3)):

21
$$p(O_3) = k_{HO_2 + NO}[HO_2][NO] + \sum_i (k_{RO_2, i + NO}[RO_2, i][NO])$$
 (1)

- $22 l(O_3) =$
- 23 $k_{O(^{1}D)+H_{2}O}[O(^{1}D)][H_{2}O] + k_{OH+O_{2}}[OH][O_{3}] + k_{HO_{2}+O_{3}}[HO_{2}][O_{3}] +$

24
$$\sum_{i} k_{O_1 + Alkene_i}[O_3][Alkene_i] + k_{OH + NO_2}[OH][NO_2]$$
 (2)

$$25 P(O_3) = p(O_3) - l(O_3) (3)$$

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Here k_{X+Y} is the bimolecular reaction rate constant for the two reagents X and Y. Therefore, the 2 calculation of ozone production rates requires peroxy radical concentrations, either from ambient 3 measurements (Green et al., 2006;Liu and Zhang, 2014;Fuchs et al., 2008;Dusanter et al., 4 2009a; Griffith et al., 2016) or box model outputs (Goliff et al., 2013; Stockwell et al., 2011; Saunders 5 et al., 2003). 6 In most urban and suburban environments, where concentrations of NO_x are significant (10-80 ppbv), 7 ozone production rates can reach a few tens of ppbv h-1 (Mao et al., 2010). In highly polluted environments, such as Mexico City or Houston, TX, P(O₃) can even exceed 100 ppbv h⁻¹ (Shirley et 8 al., 2006; Chen et al., 2010). Ozone production rates lower than 10 ppbv h⁻¹ have also been observed in 9 10 urban atmospheres such as Phoenix, AZ (Kleinman et al., 2002), likely due to lower initiation rates of 11 radicals. Ozone production is usually low in more remote areas or forested environments that are not 12 impacted by anthropogenic activities (less than 2-3 ppbv h⁻¹), due to the low NO_x concentrations 13 (Geng et al., 2011). However, if NO_x emission sources are located downwind of a forested area, 14 highly reactive biogenic VOCs (e.g. isoprene) can lead to an enhancement of ozone production (Geng 15 et al., 2011; Thornton et al., 2002). 16 Some studies performed in urban and suburban areas, whose objectives were to test our understanding 17 of the radical chemistry by contrasting measurements and model simulations of HO_x concentrations, 18 showed that models tend to underestimate HO2 for NO mixing ratios higher than a few ppbv (Ren et 19 al., 2013; Chen et al., 2010; Dusanter et al., 2009b; Kanaya et al., 2007; Ren et al., 2003). In contrast, 20 models tend to overestimate HO2 in forested areas and regions characterized by large concentrations 21 of biogenic VOCs (Griffith et al., 2013; Mao et al., 2012; Pugh et al., 2010). Disagreements are also 22 present in the modeling of OH, with the models underestimating the measurements at forested 23 environments (Lelieveld et al., 2008; Tan et al., 2001; Whalley et al., 2011; Hofzumahaus et al., 24 2009; Lu et al., 2013; Pugh et al., 2010), while the agreement may be better when colder temperatures 25 lead to lower concentrations of isoprene and other VOCs (Griffith et al., 2013). The discrepancies 26 between models and measurements question our ability to successfully measure radical species or 27 indicate that there are still unknowns in our understanding of the radical and ozone production

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2 models are widely used for the design of air quality regulations (Rao et al., 2010; Fu et al., 2006) 3 based on emission control strategies. It is therefore essential to ensure that chemical mechanisms used 4 in atmospheric models are accurate enough to simulate the oxidative capacity of the atmosphere and 5 to predict both absolute rates of ozone production and the turnover point between the two ozone 6 production regimes. 7 In order to address these issues, an instrument for direct ozone production measurements (MOPS) was 8 developed by Cazorla and Brune (2010). The principle of MOPS is based on differential ozone 9 measurements between two sampling chambers made of FEP, one exposed to sunlight (referred as 10 sampling chamber) to get an ozone production rate inside the chamber that mimics atmospheric P(O₃) 11 and the other one covered with a UV filter (reference chamber) to suppress the radical chemistry, and 12 as a consequence, ozone production. The difference in ozone between the two chambers divided by 13 the exposure time yields the ozone production rate. However, NO2 can act as a reservoir molecule for 14 O₃ due to the rapid interconversion between these two species and NO₂ has to be converted into O₃ 15 before measuring ozone. The differential O_x ($O_x=O_3+NO_2$) measurements yields $P(O_x)$ values, which 16 represent P(O₃) when NO₂ is efficiently photolyzed during daytime. 17 The first version of the MOPS instrument was tested on the campus of Pennsylvania State University 18 in the late summer of 2008. These tests demonstrated the feasibility of the MOPS technique, as the 19 instrument responded to the presence of solar radiation and ozone precursors and yielded rates of 20 ozone production that were within a range of reasonable values (up to 10 ppbv h⁻¹) for this area. This 21 instrument was then deployed during the Study of Houston Atmospheric Radical Precursors (SHARP, 22 2009) (Cazorla et al., 2012). The measurements were compared to ozone production rates calculated 23 using measurements of HO2 and NO (referred as calculated P(O3)) as well as modeled radical 24 concentrations from a box model (referred as modeled P(O₃)). Measured and calculated P(O₃) had 25 similar peak values but the calculated P(O₃) tended to peak earlier in the morning when NO values were higher. Measured and modeled P(O₃) had a similar diurnal profile, but the modeled P(O₃) was 26 27 only half the measured P(O₃). The MOPS deployment during the SHARP field campaign showed the

chemistry, which in turn could lead to erroneous P(O₃) calculations by atmospheric models. These

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1 potential of this instrument for contributing to the understanding of the ozone-producing chemistry,

2 but was limited by measurement uncertainties due to potential wall effects. The heterogeneous loss of

3 NO₂ under humid conditions (RH> 50%) was reported as a main issue for this technique.

4 Recently, an improved version of the MOPS instrument was deployed during the NASA's

5 DISCOVER-AQ field campaign in 2013, in Houston, Texas (Baier et al., 2015). Wall effects were

reduced by improving the design of the sampling chambers and the airflow characteristics. The

measurements made over one month were consistent with ambient ozone observations and model-

derived P(O₃) values from previous field campaigns in Houston. The authors, however, highlighted a

possible bias due to surface HONO production followed by its photolysis in the sampling chamber, as

10 well as unresolved ozone analyzer issues. HONO concentrations in the sampling chambers were

11 reported as two to five times higher than ambient values, which could cause a bias up to 5-10 ppbv h⁻¹

on the $P(O_3)$ measurements.

13 In this publication, we present the development and the characterization of an Ozone Production Rates

(OPR) instrument. The OPR instrument is based on the principle of the MOPS, using different

15 sampling and detection schemes. This publication describes this new instrument and its

characterization in the laboratory. An emphasis is given to the modeling of the radical chemistry

inside the sampling chambers to assess potential biases on P(O₃) measurements associated to

instrumental characteristics and operating conditions. The publication also reports preliminary field

results from the Indiana Radical, Reactivity and Ozone Production Intercomparison (IRRONIC)

20 campaign, which highlight the current limitations of this instrument.

2 Experimental section

2.1 Description of the OPR instrument

23 The principle of the OPR is based on differential O_x measurements between an "ambient" flow tube,

24 exposed to sunlight to mimic ambient photochemistry, and a "reference" flow tube, covered with an

25 Ultem® film (polyetherimide, 0.25 mm thick, CS Hyde Co, USA) to block wavelengths lower than

26 400 nm, which in turn should suppress ozone production. As mentioned above for the MOPS

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- 1 instrument, the fast partitioning between O₃ and NO₂ requires measuring O_x instead of O₃, assuming
- 2 that $P(O_3)$ is equal to $P(O_3)$ when NO_2 is efficiently photolyzed during daytime. $P(O_3)$ is calculated
- 3 from the difference in O_x between the two flow tubes, ΔO_x , divided by the mean residence time (τ) of
- 4 air inside the tubes:

$$5 P(O_x) = \frac{\Delta O_x}{\tau} = \frac{O_{xamb} - O_{xref}}{\tau} (4)$$

- 6 A detailed schematic of the OPR instrument is shown in Figure 1. The two flow tubes exhibit the
- 7 same geometry and are made of quartz (14 cm-ID and 70 cm long). Each flow tube is connected to the
- 8 inlet and outlet flanges that are made of anodized aluminum and PTFE. Since a major issue previously
- 9 identified for the MOPS instrument was wall effects causing NO₂ losses (Cazorla and Brune, 2010),
- 10 the inner geometry of the flanges was designed based on fluid dynamics simulations using STAR
- 11 CCM+ V.8 (CD-adapco). The geometry was optimized to minimize radial mixing and recirculation
- 12 eddies that could increase wall effects. The design of the flanges can be found in the supplementary
- 13 material (Fig. S1).
- 14 Each flange consists of two parts. For both the inlet and outlet, a conical PTFE piece is screwed inside
- 15 an external aluminum flange. Four holes are drilled symmetrically around the aluminum flanges to
- 16 inject zero air around the PTFE inlet and to extract air around the PTFE outlet. The lengths of the inlet
- 17 and outlet flanges are 25 and 14 cm, respectively. The PTFE inlet has an external diameter of 2.54 cm
- 18 which increases to 7 cm over a length of 20 cm. The PTFE outlet starts from a diameter of 3 cm
- 19 which decreases to 1.27 cm over 10 cm. The aluminum flanges exhibit a curved conical inner surface
- around the PTFE parts.
- 21 Ambient air is sampled through a common inlet (PFA, 1.27 cm-OD) at a flow rate of 4 L min⁻¹ and is
- 22 transferred into both flow tubes through the internal PTFE inlets (2 L min⁻¹), while additional zero air
- 23 (250 mL min⁻¹) is injected at the outer periphery of these inlets inside the flanges. This flow of zero
- 24 air helps keeping the ambient air flow forward, minimizing recirculation eddies, and therefore
- 25 reducing wall effects. The dilution of the sampled air is approximately 10%. At the outlet, air is
- 26 sampled only from the center of the flow tube, through the PTFE outlet (750 mL min⁻¹), while the rest

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1 is extracted by an external pump (1.5 L min⁻¹). Both the injection and extraction of air are regulated

2 by mass flow controllers (MFC in Fig. 1).

3 The Ultern filter is placed on a rectangular aluminum frame outside of the reference flow tube, which

4 enables to flow ambient air between the filter and the flow tube using fans. This setup allows the two

5 flow tubes to be kept at the same temperature by extracting the heat released by the filter. For the

6 same reason, a frame covered by a FEP film (.002" thick, DuPont Teflon® FEP), transparent to the

7 solar radiation, is used for the ambient flow tube to reduce heat dissipation by the wind.

8 The air exiting the two flow tubes is mixed with 10 SCCM of NO (50 ppmv, Indiana Oxygen, USA),

leading to a NO mixing ratio of 650 ppbv in the conversion unit. The mixing of the gases takes place

in two identical pyrex chambers, providing a reaction time of approximately 22 sec at 20°C, which is

long enough to quantitatively titrate O₃ into NO₂. Both the relative humidity and temperature are

monitored in the air flow extracted from the flow tubes and at the O_3 -to- NO_2 conversion unit.

13 Downstream the conversion unit, O_x (O₃ + NO₂) is measured by an Aerodyne Cavity Attenuated

14 Phase Shift Spectroscopy (CAPS) NO₂ monitor (Kebabian et al., 2005; Kebabian et al., 2008). The

detection limit (3σ) for a 1-s integration time is 300 pptv. Since the CAPS is a single-cell monitor, the

measurements from the ambient and reference flow tubes are taken sequentially, using two solenoid

17 valves (SV1 and SV2 in Fig. 1). When air from the ambient (or reference) flow tube is sampled by the

18 CAPS monitor (750 ml min⁻¹), the same flow rate of air is extracted from the other flow tube by a

19 mass flow controller connected to a pump. The valves switch every 1 min, alternating the flows that

are sampled by the CAPS monitor and the pump. ΔO_x is calculated as the difference between an

ambient flow tube measurement and the average of 2 surrounding reference measurements, leading to

a P(O_x) measurement every 2 min. The first 15 seconds of each 1-min measurement are discarded

since they describe a transient regime between ambient and reference flow tube measurements. Ozone

24 production values are calculated from Eq. (4).

25 The measurement sequence is automated and controlled through a National Instruments LabView

26 2013 interface. Three USB data acquisition boards are used (NI-9264, NI-6008, NI-6009) to control

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- 1 the two solenoid valves and the seven mass flow controllers, as well as to record signals from the
- 2 CAPS monitor and sensors setup for humidity and temperature measurements.

3 2.2 Laboratory and field experiments conducted to characterize the OPR

- 4 Experiments conducted to characterize the OPR instrument include measurements of the mean
- 5 residence time, O_x losses, and HONO production rates in the flow tubes and measurements of the O₃-
- 6 to-NO₂ conversion efficiency.
- 7 The mean residence time was quantified in each flow tube by injecting short pulses of toluene (10-s
- 8 in duration) at the inlet of the flow tubes. A PTR-ToFMS (Proton Transfer Reaction-Time of Flight
- 9 Mass Spectrometer, KORE Technology Inc.) was connected at the outlets to measure the time it takes
- 10 for a pulse introduced at the inlets to exit the flow tubes. The pulse experiment was repeated 5 times,
- and the average was calculated as the mean residence time.
- 12 O₃ and NO₂ losses inside both flow tubes were measured in the laboratory and during the field
- deployment described below by sampling mixtures of zero air and O₃ (or NO₂) at known mixing ratios
- and by measuring NO₂ downstream the conversion unit (or directly at the exit of the flow tubes). A
- 15 relative loss was calculated from the difference in concentrations between the inlet and outlet and was
- 16 referenced to the inlet concentration. These tests were performed at relative humidity values ranging
- 17 from 0–65%.
- 18 The release of HONO from the inner surface of the flow tubes was quantified using a Chemical
- 19 Ionization Mass Spectrometer (CIMS, Georgia Tech). Mixtures of NO₂ and humid zero air were
- 20 introduced into the flow tubes, while HONO was measured both at the inlet and outlet. These
- 21 experiments were performed under dark conditions, as well as under various irradiated conditions
- 22 using artificial UV light provided by two types of fluorescent lamps: 4 lamps centered at 312 nm
- 23 (Vilber, T-15.M) and 4 lamps centered at 365 nm (Philips, T12).
- 24 Finally, the O₃-to-NO₂ conversion efficiency was measured by sampling zero air enriched with O₃
- 25 (3-170 ppbv) through the mixing chambers of the conversion unit, varying the flow of NO and
- 26 measuring NO₂ with the CAPS monitor. These tests were performed at various relative humidities

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1 (25-60%). The conversion efficiency at a specific NO level was calculated from the ratio of NO_2

2 measured at this NO level to that measured when 700 ppbv of NO were added, assuming for the latter

3 that 100% of O₃ was converted. This assumption is verified from kinetic considerations

4 $(k_{NO+O3}=1.80\times10^{-14} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1} \text{ and } 23 \text{ s of residence time in the conversion unit)}$ and from the

5 observation of a plateau for NO mixing ratios higher than 500 ppbv.

2.3 Modeling experiments conducted to characterize the OPR

As previously mentioned, the measurement principle of ozone production rates is based on the

8 assumption that (i) $P(O_x)$ in the ambient flow tube is similar to $P(O_x)$ in the atmosphere and (ii) there

is no significant production of ozone in the reference flow tube. Box model simulations were

10 performed to check whether this assumption is valid. In addition, simulations were also conducted to

11 investigate the impact on OPR measurements of (a) an O₃-to-NO₂ conversion efficiency lower than

12 100%, (b) NO₂ and O₃ losses and (c) HONO production inside the flow tubes, (d) a possible increase

of the temperature in the reference flow tube due to the UV filter, (e) the dilution of ambient air by

injecting zero air inside the flow tubes at the periphery of the inlets, and (f) reactions of OH with NO_z

15 species producing O_x.

16 2.3.1 Selected data and chemical mechanism

17 The simulations were performed using a box model based on the Regional Atmospheric Chemistry

18 Mechanism (RACM) (Stockwell et al., 1997). RACM is a gas-phase chemical mechanism developed

19 for the modeling of regional atmospheric chemistry and includes 17 stable inorganic species, 4

20 inorganic intermediates, 32 stable organic species and 24 organic intermediates for a total of 237

21 chemical reactions. Organic compounds are grouped together to form a manageable set of

22 compounds. Only 8 organic species are treated explicitly (methane, ethane, ethene, isoprene,

23 formaldehyde, glyoxal, methyl hydrogen peroxide and formic acid) and 24 are surrogates that are

24 grouped based on emission rates, chemical structure and reactivity with the OH radical.

25 Measurements from several field campaigns were used for this modeling exercise, including

26 measurements performed in (i) a megacity as part of the 2006 Mexico City Metropolitan Area

27 (MCMA-2006) (Dusanter et al., 2009b) and (ii) an urban area as part of the 2010 California Nexus

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- 1 (CalNex) campaign (Griffith et al., 2016). Two days characterized by elevated and low O_x
- 2 concentrations were selected for each campaign and are presented in the supplementary material
- 3 (Table S1 and Fig. S2). For both campaigns, ozone was higher by approximately a factor 2 on high O₃
- 4 days (≈ 100 ppbv) compared to low O₃ days (≈ 50 ppbv). However, while both high and low ozone
- 5 levels were similar for the selected days of these campaigns, large differences were observed for NO_x
- 6 (6–120 ppbv) and OH reactivity (8–86 s⁻¹). Since OH reactivity and NOx are main drivers of ozone
- 7 production, these modeling results are expected to provide a good assessment of potential biases
- 8 associated to P(O_x) measurement for any urban environments.

9 2.3.2 Modeling of ambient P(O_x) values

- 10 The model was constrained by 10-min (MCMA) or 15-min (CalNex) average measurements of
- 11 temperature, pressure, humidity, organic and inorganic species, and J-values, while the differential
- 12 equation system was integrated by the FACSIMILE solver (MCPA Software Ltd). In total, 24 J-
- 13 values were used to constrain the model, as derived in Dusanter et al. (2009b), together with 7
- 14 inorganic and 17 organic species or surrogates. Tables reporting the constrained species and J-values
- 15 can be found in the supplementary material (Tables S2 and S3). The integration time was set at 30h
- 16 with constrained species reinitialized every two seconds. Ambient ozone production values were then
- 17 calculated from Eq. (1)–(3) and are referred as $P(O_x)_{atm}$ in the following. In total, 18 surrogates of
- 18 RO₂ species were taken into account to calculate p(O₃) from Eq. (1), while 10 unsaturated surrogates
- were used to calculate l(O₃) from Eq. (2) (Table S4).

20 2.3.3 Modeling of $P(O_x)$ values in the ambient and reference flow tubes

- 21 Modeling OPR measurements requires simulating the chemistry inside each flow tube. J-values used
- 22 to model the chemistry in the ambient flow tube were the same as for the ambient modeling since the
- 23 quartz material used to build the flow tubes is transparent to solar irradiation. For the reference flow
- tube, J-values were scaled based on the absorption coefficient of the Ultem film (Philipp et al., 1989)
- as discussed in the supplementary material (section S2.1).
- 26 The model was constrained by the same meteorological parameters and chemical species as for
- 27 $P(O_x)_{atm}$. In addition, modeled concentrations of VOC-oxidation products and peroxy radicals

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- 1 inferred from the modeling of $P(O_x)_{atm}$ were also constrained in these simulations (Table S5),
- 2 assuming that a significant fraction of the latters is not lost in the sampling line. The constrained
- 3 concentrations were initialized once, at the entrance of the flow tubes, and the simulations were run
- 4 for 10 minutes without reinitializing the constraints. The simulations were run separately for each
- 5 flow tube and $P(O_x)$ was calculated every 15 s from Eq. (3). An integrated value of $P(O_x)$ was then
- 6 computed for the flow tube residence time.
- 7 $P(O_x)_{atm}$ is compared to the integrated $P(O_x)$ value from the ambient flow tube (referred as
- 8 $P(O_x)_{amb}$) to check whether ozone production in the ambient flow tube is similar to ambient ozone
- 9 production. The integrated value of $P(O_x)$ in the reference flow tube (referred as $P(O_x)_{ref}$) is also
- scrutinized to check whether ozone production is negligible in this flow tube.

11 2.3.4 Modeling of OPR measurements

- 12 Since the OPR instrument measures O_x after conversion of O₃ into NO₂, NO₂ concentrations at the
- 13 exit of the conversion unit are calculated from the conversion efficiency C as shown in Eq. (5):

14
$$[NO_2]_{conv} = [NO_2]_{\tau} + C[O_3]_{\tau}$$
 : (5)

- 15 Here the concentrations reflect those observed at the exit of the conversion unit (subscript: conv) and
- 16 at the exit of the flow tubes (subscript: τ). The concentrations at the exit of the flow tubes are the
- 17 model outputs at the residence time τ. Based on Eq. (4), the ozone production rate measured by the
- OPR, $P(O_x)_{OPR}$, is then calculated from Eq. (6):

19
$$P(O_x)_{OPR} = \frac{[NO_2]_{conv,amb} - [NO_2]_{conv,ref}}{\tau} = \frac{[NO_2]_{\tau,amb} - [NO_2]_{\tau,ref} + C([O_3]_{\tau,amb} - [O_3]_{\tau,ref})}{\tau}$$
 (6)

- 20 In this equation the subscripts amb and ref indicate the ambient and the reference flow tubes,
- 21 respectively. A Bias in OPR measurements can be quantified by comparing $P(O_x)_{OPR}$ to $P(O_x)_{atm}$
- assuming a conversion efficiency of 100% for the conversion units.

23 2.3.5 Sensitivity tests

- 24 The simulation performed without O_x losses and HONO production in the flow tubes, no dilution, and
- 25 no temperature differences between the tubes will be referred as base simulation in the following. All

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1 simulations performed including sensitivity tests are compared to the results from the base simulation

2 to assess the impact of operating conditions on ozone production measurements.

3 To assess the impact of a conversion efficiency lower than 100%, $P(O_x)_{OPR}$ is calculated from Eq. (6)

4 by varying the conversion efficiency using the model outputs from the base simulation. $P(O_x)$ values

5 inferred when varying the conversion efficiency are compared to values calculated for a conversion

efficiency of 100%. To account for O_x losses, a similar sink of O_3 or NO_2 is introduced in the model

for each flow tube, with a first order loss rate ranging from 1.5×10^{-4} to 1.2×10^{-3} s⁻¹. This range of loss

rates corresponds to a relative loss of 4–28%. The measured $P(O_x)_{OPR}$ is again calculated by Eq. (6)

assuming a conversion efficiency of 100% and compared to the base simulation. Sensitivity tests were

also performed assuming that the loss of NO₂ on the quartz surface led to HONO formation with the

same first order rate as the NO₂ loss, or by including a HONO source in the model, independent of

NO₂, with production rates comparable to experimental observations. Additional sensitivity tests

13 focused on decreasing the constrained species by 5-30% to assess the impact of diluting ambient air in

the flow tubes, as well as increasing the temperature of the reference flow tube by 2% to 20% to

15 simulate a heat release by the UV filter. Finally, sensitivity tests were performed to investigate

16 whether reactions of OH with NO_z species that produce O_x could significantly impact the OPR

17 measurements. NO_z species producing NO₂ or NO₃ (NO₂ reservoir) in the model when reacting with

18 OH are HONO, HO₂NO₂, organic nitrates, HNO₃, PANs and unsaturated PANs. The NO₂ and NO₃

19 products of the reactions mentioned above were removed from the model for the sensitivity test.

2.4 Description of the field measurements

21 The OPR instrument was deployed in the field, as part of the Indiana Radical, Reactivity and Ozone

22 Production Intercomparison (IRRONIC) campaign in Bloomington, Indiana, during July 2015. The

23 measurements were taken at the Indiana University Research and Teaching Preserve (IURTP) field

24 laboratory (39.1908N, 86.502W), 2.5 km northeast of the Indiana University Bloomington campus.

25 The site is a mixed deciduous forest containing northern red oaks and big-tooth aspens, which are

known to be strong emitters of isoprene and monoterpenes (Isebrands et al., 1999;Funk et al., 2005).

27 A highway (E Matlock Road, State Route 45) is located 1 km southwest, and therefore the site can be

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1 impacted by anthropogenic emissions. The OPR flow tubes were setup on a scaffolding to expose 2 them to the sunlight for the entire day. The conversion units and the CAPS monitor were housed 3 inside the laboratory and were connected to the flow tubes using 4-m long heated 1/4" PFA lines. 4 This campaign included measurements of OH, HO₂* (HO₂+αRO₂), total peroxy radicals (HO₂+RO₂), 5 total OH reactivity, NO_x, O₃, anthropogenic and biogenic VOCs, radiation and meteorological data. 6 For the measurements presented in this publication, VOCs were measured by an online TD-GC/FID, 7 an online TD-GC/FID-MS (Badol et al., 2004; Roukos et al., 2009), and offline samplers for 8 DiNitroPhenylHydrazine (DNPH) cartridges (Waters Sep-Pak) and Sorbent cartridges (Carbopack 9 B/Carbopack C) by IMT Lille Douai. Measurements of NO (chemiluminescence, Thermo model 42i-10 TL), NO₂ (cavity attenuated phase shift spectroscopy, Aerodyne Research), and ozone (2B Tech 11 model 202 sensor) were also conducted by the University of Massachusetts. Measurements of J(NO₂) 12 were performed using a scanning actinic flux spectroradiometer (SAFS, METCON) from the 13 University of Houston, while meteorological data, including temperature, relative humidity, wind 14 speed and wind direction were measured with a meteorological station from Montana State 15 University. 16 The OPR measurements were focused on investigating the sensitivity of P(Ox) to NOx (see section 17 3.3). This was achieved by introducing a certain amount of NO (ppbv range) inside the OPR sampling 18 line for 40 minutes, and then stopping the NO addition for another 40 minutes. This pattern was 19 repeated continuously, keeping the NO level constant for several days. The first 20 minutes of each 20 40-minutes measurements were discarded, since they correspond to a transient regime between the 21 disturbed-undisturbed $P(O_x)$ measurements due to the long air-exchange time in the flow tubes (see 22 section 3.1.1). The addition of NO in the OPR sampling line was performed through a 1/8"-OD 23 stainless steel tube using a NO cylinder (3.75 ppmv in N₂) from Indiana Oxygen and a mass flow 24 controller. After the mixing point, a length of 10 m of 1/2"-OD PFA tube was used as the sampling 25 line to ensure a good mixing of NO with the sampled air, leading to a residence time of approximately 26 10 s in the line at a total flow rate of 4 L min⁻¹.

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Results and discussion

Laboratory characterization

3.1.1 Quantification of the flow tubes residence time

3 4 As described in the experimental section, pulses of toluene were injected in the flow tubes to quantify 5 the mean residence time. One of the 5 experiments that were conducted is shown in Figure 2. The pulse shape is asymmetric and exhibits a long tail, indicating that a large range of residence times is 6 7 observed in the flow tubes. The toluene pulse is treated as a probability distribution of the time 8 variable t, with the average residence time in the flow tubes being the mean of the probability 9 distribution. The latter is calculated as a weighted average of the possible values that the time variable 10 can take. The average residence time from the 5 toluene pulse experiments was $4.52 \pm 0.22 \text{ min } (1\sigma)$. 11 The uncertainty reported for the residence time will lead to a 4.9% error (1σ) on the $P(O_x)$ 12 measurements. Based on the flow tube volume of 10.8 L and a total flow rate of 2.25 L min-1 in each flow tube, a 13 14 laminar plug flow would lead to a residence time of 4.79 min. The measured residence time is 15 approximately 6% lower than the time calculated for laminar flow conditions, which is barely 16 significant considering the 1σ uncertainty of 4.9% determined from the measurements. However, the 17 asymmetry of the peak indicates that the flow rate at the central axis of the tube is larger, with the first 18 molecules of toluene being sampled after approximately 2 minutes (Fig. 2). These observations are 19 similar to that reported by Cazorla and Brune (2010) for sampling chambers exhibiting a different 20 geometry and operated under different flow conditions. In this study, the authors reported a residence 21 time that was 23% lower than plug flow calculations and a similar asymmetric shape for the pulse. Further work is needed to reduce the skewness of the time distribution. 23

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Tests were also performed to quantify the air-exchange time in the flow tubes. These tests were performed by sampling a constant concentration of O_x species with the OPR instrument until a stable O_x signal was measured and by quickly changing the Ox concentration at the inlet. The time needed to reach 95% of a new stable Ox signal was defined as the air-exchange time. The air-exchange time was quantified at approximately 20 minutes, corresponding to a maximum residence time of 1200 s. As

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1 mentioned in section 2.1, a $P(O_x)$ value is recorded every 2 minutes. Since the air-exchange time is 20

2 minutes, the 2-minute P(O_x) values are not independent from each other and therefore the OPR

3 instrument cannot detect rapid changes of P(O_x). In order to get independent measurements of P(Ox),

4 the OPR measurements are therefore averaged over 20 minutes.

3.1.2 Quantification of O_x losses in the flow tubes

6 The principle of the OPR instrument requires that the only difference between the two flow tubes is

the suppression of gas-phase photolytic reactions leading to the formation of free radicals in the

reference tube. All other characteristics, including flow pattern and potential gas-wall interactions

should be the same in the two flow tubes so that they cancel out in the differential O_x measurement.

10 However, if O_x losses were slightly different between the two flow tubes, it could significantly impact

the $P(O_x)$ measurements. For example, a 2% difference in O_x losses between the flow tubes would

lead to a bias of 27 ppbv h⁻¹ on the measurements for an ambient O_x level of 100 ppbv and a residence

13 time of 4.5 min.

14 Figure 3 shows the results of NO₂ and O₃ loss tests for the two flow tubes, performed at different

15 dates during one month of field operation during the IRRONIC campaign and at different relative

humidity values. All NO₂ loss tests were performed under dark conditions, i.e. with both flow tubes

17 covered by an opaque cover. Figure 3-(a, c, e) shows that the NO₂ loss is lower than 5% in both flow

tubes and is close to 3% on average. When the two flow tubes are operated under the same conditions,

19 the relative loss in the reference tube seems to be higher than the loss in the ambient tube by only 1%

at most (Fig. 3-e). For an ambient NO₂ mixing ratio of 30 ppbv, a difference of 1% in NO₂ losses

between the flow tubes would lead to a 4 ppbv h^{-1} bias in the $P(O_x)$ measurements.

22 Cazorla and Brune (2010) reported an uncertainty of $\pm 14\%$ for the MOPS instrument due to potential

differences in relative humidity between the two sampling chambers, which in turn leads to different

NO₂ losses. This was mainly due to a higher temperature in the reference chamber, which is covered

25 by the UV filter. However, the fans used in the OPR instrument to flow ambient air between the UV

filter and the flow tube minimize the temperature differences between the two tubes, leading to

27 relative humidity differences lower than 4%, as observed during the field testing. Figure 3-e also

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1 shows that a decrease in relative humidity from 65% to 0% only leads to a small decrease of the NO₂ 2 loss by 1-2%. A small difference of 4% in relative humidity between the two flow tubes is therefore 3 not expected to lead to additional errors in the P(O_x) measurements. Further analysis of the impact of 4 NO₂ losses on the P(O_x) measurements is discussed in the modeling results section. 5 Ozone loss tests were mainly performed under dark conditions. On 28 July however, O₃ losses were 6 measured with (a) the ambient flow tube exposed to the sunlight and the reference tube covered by the 7 UV filter (orange squares), (b) both flow tubes exposed to the sunlight (orange triangles) and (c) both 8 tubes covered by a dark cover (orange circles). For the first days of the campaign (29 June-8 July), a 9 close inspection of the measurement scatter shown in Figure 3-(b, d) indicates that the relative loss of 10 O₃ is at most close to 5%. However, ozone loss tests performed on 28 July, after one month of 11 operation in the field, reveal an increase of the relative loss up to 13-15%. Additional tests made in 12 the laboratory after the field deployments have shown that this loss can be reduced below 5% if the 13 quartz flow tubes are conditioned with elevated O₃ mixing ratios at high relative humidity for a few 14 days. These results indicate that unsaturated organic species may adsorb on the quartz surface and 15 may react with O₃. 16 Particular attention should be paid to the three different tests performed on 28 July regarding the irradiation conditions. When the losses are quantified under dark conditions (orange circles in Fig. 3-17 18 f), the losses are equal between the two flow tubes and close to 13%. However, when the ambient 19 flow tube is irradiated and the reference is covered by the UV filter (orange squares), it can be seen 20 that the relative loss in the ambient tube is higher than in the reference by approximately 3%. Box 21 modeling has shown that the gas-phase photolysis of O₃ in the ambient flow tube could at most 22 account for 0.05% of this additional ozone loss. Therefore, there seems to be a photo-enhanced ozone 23 loss that takes place when the ambient flow tube is irradiated. For an ambient O₃ level of 50 ppbv, this difference in O₃ losses would lead to a negative P(O_x) bias of approximately 20 ppbv h⁻¹. This photo-24 25 enhanced loss of ozone is further discussed in the field deployment section (3.3).

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production rate to negligible levels.



1 3.1.3 Heterogeneous HONO production in the flow tubes

2 The formation of HONO in the flow tubes was investigated in the laboratory by sampling humid zero 3 air (25-80% RH) enriched with NO₂ at various mixing ratios (0-100 ppbv) and by measuring HONO 4 mixing ratios in the tubes as described above in section 2.2. Both clean and contaminated (used for 5 more than one month during the IRRONIC campaign) flow tubes were tested to assess the magnitude 6 of HONO production rates and to examine whether there is a dependence on NO2 mixing ratios, 7 humidity and irradiation. Mixing ratios of HONO up to 250 and 700 pptv were measured under dark 8 conditions for clean and contaminated flow tubes, respectively. Higher mixing ratios of up to 1.5 ppbv 9 were measured under irradiated conditions in the ambient flow tube (J(NO₂)=1.4×10⁻³ s⁻¹; 10 $J(HONO)=3.1\times10^{-4} \text{ s}^{-1}$). 11 Dividing the measured mixing ratios of HONO by the residence time in the flow tubes (i.e. 4.5 min), 12 an average production rate can be calculated under dark and irradiated conditions. It is important to 13 note, however, that HONO is also photolyzed at the wavelengths emitted by the lamps (312 nm and 14 365 nm) and production rates calculated under irradiated conditions represent lower bounds. It is 15 estimated that for the J(HONO) value mentioned above and a negligible loss of HONO from 16 OH+HONO, the HONO production rate will be underestimated by less than 8%. The dark HONO 17 production is on the order of 9 ppbv h⁻¹ in both flow tubes, while the total HONO production under 18 irradiated conditions (dark + photo-enhanced) can reach up to 20 ppbv h⁻¹ in the ambient flow tube. In 19 the reference flow tube, the UV light did not impact the formation of HONO, since wavelengths 20 below 400 nm are blocked by the UV filter. 21 The HONO production rate was not observed to depend on NO2 or humidity and HONO could be 22 even released when no NO2 was introduced into the contaminated flow tubes. These results strongly

suggest that nitro-containing compounds and organic photosensitizers were adsorbed on the walls of

the flow tubes and that the HONO production rate depends on contamination levels. Indeed, it was

observed that flowing humid zero air in the flow tubes for a few days could reduce the HONO

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3.1.4 Quantification of the conversion efficiency

Based on kinetic considerations for the titration reaction of O₃ by NO, i.e. a rate constant of 1.80×10⁻¹⁴ 2 3 cm³ molecule⁻¹ s⁻¹ at 298K (Atkinson et al., 2004), a reaction time of 23 seconds, and the addition of 4 500 ppbv of NO in the conversion unit, an O₃-to-NO₂ conversion efficiency of 99.5% is expected. 5 These calculations are shown in Figure 4 (black solid line) for different mixing ratios of NO (50-800 6 ppbv) together with laboratory measurements (symbols) made at different O₃ levels. This figure 7 shows that a plateau of almost 100% of conversion is observed at NO mixing ratios higher than 500 8 ppbv. These experimental results are in good agreement with the calculated curve, although the 9 measurements performed at a low O₃ mixing ratio of 3.5 ppbv slightly underpredict the curve for NO 10 mixing ratios lower than 500 ppbv. However, the conversion plateau is reached for all O_x levels and 11 both conversion units (one for each flow tube) for NO mixing ratios higher than 500 ppbv. During the 12 field deployment of the instrument, an NO mixing ratio of 650 ppbv was used to ensure that the 13 difference in conversion efficiency between the two mixing chambers was lower than 0.1% and could 14 be assumed to be 100% for both chambers. 15 In the first version of MOPS (Cazorla and Brune, 2010) the NO₂-to-O₃ conversion was performed by 16 photolyzing NO₂ using a light-emitting diode, achieving a maximum conversion efficiency of 88% at 17 17 ppbv of NO2. In the most recent version of the instrument (Baier et al., 2015), the conversion 18 efficiency was increased to 88-97% for NO₂ mixing ratios lower than 35 ppbv using a highly-efficient 19 UV lamp that provided ten times more photons than the light-emitting diodes. In the MOPS 20 instrument, however, the conversion efficiency depends on NO₂ levels, as well as on the intensity of 21 the lamp that could drift during a long period of use in the field. In the OPR instrument, the 22 conversion efficiency is stable and does not depend on O₃ mixing ratios. On the other hand, an NO 23 cylinder is required to perform the conversion and possible NO₂ impurities in the cylinder have to be 24 monitored. Indeed, NO2 impurities coming either from the NO mixture or from NO oxidation in the 25 lines were observed, but were kept at low levels of approximately 6-10 ppbv. Since this impurity is

present in both the ambient and reference channel, it does not affect the $P(O_x)$ determination.

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3.1.5 Detection limit of the OPR

- 2 The detection limit (DL) of the CAPS monitor was quantified by sampling zero air for several hours
- 3 after several days of conditioning with ambient air. The time resolution was set to 1 s and the zero
- 4 measurements were averaged over 45 s segments, corresponding to the OPR measurement averaging
- 5 time. The detection limit (3σ) for a 45 s integration time was quantified at 34 pptv. This detection
- 6 limit for NO₂ together with a residence time of 4.5 min in the flow tubes should lead to a detection
- 7 limit of 0.6 ppbv h⁻¹ for 2-min P(O_x) measurements (1-min measurement from each flow tube).
- 8 However, nighttime measurements made during the IRRONIC field campaign revealed that the
- 9 measurement scattering for the complete setup (flow tubes + O₃-to-NO₂ conversion unit + CAPS) was
- significantly larger than that expected from the noise of the CAPS monitor. Based on the observed
- nighttime 1σ variability of 2.1 ppbv h^{-1} , a limit of detection (3 σ) of 6.2 ppbv h^{-1} was inferred for the
- 12 OPR instrument. The scatter in $P(O_x)$ measurements does not only depend on the precision of the
- 13 CAPS monitor, but also depends on how fast each flow tube responds to variations of O_x at the inlet.
- 14 Indeed, if the time constant for the response is slightly different between the 2 flow tubes, fluctuations
- 15 of O_x species at the inlet will introduce some scatter in the OPR measurements. In addition, small
- 16 changes in temperature and humidity may evenly affect O_x losses in each flow tube, leading to
- 17 additional scatter in the P(O_x) measurements.

18 3.2 Numerical Modeling

- 19 As mentioned in the experimental section, several days from different field campaigns were selected
- to model ambient $P(O_x)$, $P(O_x)$ in both flow tubes, and the impact of some operating conditions on the
- 21 OPR measurements. The results from 30 May 2010 of the CalNex field campaign were selected to
- 22 illustrate the discussion and results from the other days are shown in the supplementary material
- 23 (Figs. S4, S5, S7-S9). A detailed analysis of the chemistry occurring in each flow tube is discussed
- below to assess the reliability of OPR measurements.

3.2.1 Radical budget in flow tubes

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- 26 An analysis of the radical budget was performed in each flow tube to gain insights into the processes
- 27 driving radical production and loss routes. Figure 5 shows the production and loss rates of OH (upper

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1 panel) and peroxy radicals (lower panel) for each flow tube on 30 May 2010 during CalNex. The 2 production and loss rates were calculated taking into account initiation, propagation and termination 3 processes as described below. 4 OH production rates were calculated from photolytic reactions involving closed shell molecules (O₃, 5 HONO, H₂O₂, HNO₃, HO₂NO₂ and organic peroxides), reactions of O₃ with alkenes, and the 6 propagation of HO₂ by reaction with NO. Loss routes of OH includes propagation reactions to HO₂ 7 and RO₂ by reaction with CO and VOCs and termination reactions of OH with NO₂ and other species 8 (NO, PANs, HNO₃, HONO and HNO₄). For peroxy radicals, production routes include the photolysis 9 of organic species (carbonyls, organic peroxides and organic nitrates), the ozonolysis of alkenes, PAN 10 decomposition, and the propagation of OH. Loss routes were calculated from reactions of peroxy 11 radicals with NO_x, self or cross reactions between peroxy radicals and propagation of HO₂ to OH. 12 Figure 5 clearly shows that the UV filter covering the reference flow tube leads to a decrease of the 13 initiation rates of all radicals by more than a factor of 10 and a decrease of their propagation rates by 14 at least a factor of 30. In the ambient flow tube, photolytic reactions of OVOCs are the most important 15 initiation routes of peroxy radicals, with a contribution of approximately 95%. HONO and O₃ 16 photolysis are the most important initiation routes of OH, contributing by approximately 45% each. In 17 the reference flow tube, the primary route of radical initiation is O3-alkenes reactions since 18 wavelengths below 400nm are suppressed. 19 The propagation reactions are important in both flow tubes for the production and loss of OH and 20 peroxy radicals. However, the partitioning between initiation and propagation processes is different in 21 the two tubes, which in turn leads to different OH chain lengths. The OH chain length is calculated as 22 the rate of propagation of HO₂ to OH divided by the total initiation of RO_x radicals. As can be seen 23 from Figure 5, the OH chain length is fairly constant at a value of 3 in the ambient flow tube, while in 24 the reference flow tube it quickly decreases to unity for most of the day and to values lower than 1 in 25 the late afternoon. Therefore, in addition to lowering initiation rates of radicals, the UV filter allows to 26 reduce ozone production by lowering the cycling efficiency within the pool of RO_x radicals.

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1 A close inspection of the radical termination rates in Figure 5 indicates that the peroxy-NO_x 2 termination reactions are almost suppressed in the reference flow tube. This observation is also 3 supported by Figure S6, which shows time series of the peroxy radicals (HO₂ and RO₂) and NO in 4 each flow tube at a residence time of 4.5 min. Since NO₂ photolysis is almost eliminated in this tube, 5 the O₃-NO_x PSS is shifted towards NO₂ due to the reaction of NO with O₃. As a result, NO mixing 6 ratios in the reference flow tube are at least one order of magnitude lower than in the ambient flow 7 tube. The propagation rate from HO₂+NO is therefore reduced and the OH+NO₂ loss route is 8 enhanced, leading to the shorter OH chain length discussed above. It is also interesting to note that 9 peroxy radical mixing ratios in the reference flow tube are on the same order of magnitude as in the 10 ambient flow tube. This counterintuitive observation is also due to the consumption of NO in the 11 reference flow tube that leads to a longer lifetime for the peroxy radicals, as shown in Figure S6. 12 Calculating $P(O_x)$ from Equations (1-3) results in ozone production rates in the ambient flow tube, 13 $P(O_x)_{amb}$, in good agreement with the modeled $P(O_x)_{atm}$ values, as shown in Figure 6, with a small underestimation of approximately 10% on average. However, significant ozone production rates are 14 also observed in the reference flow tube, which can reach up to 4 ppbv h⁻¹ on this day, while higher 15 values were observed on other days (e.g. 30 ppbv h⁻¹ on 21 March 2006 of the MCMA-2006 16 17 campaign, Figure S10 in the supplementary material). Ozone production rates in the reference flow 18 tube are about 10–15% of that observed in the ambient flow tube for most of the day. It is important to 19 note, however, that this ozone production is in reality O_x (=O₃+NO₂) production, since NO₂ photolysis 20 is almost suppressed in the reference flow tube. These results indicate that the assumptions initially 21 made on the principle for $P(O_x)$ measurements, i.e that $P(O_x)$ in the ambient flow tube mimics $P(O_x)$ 22 in the atmosphere and P(O_x) in the reference flow tube is not significant, are not completely fulfilled. 23 Based on the modeling results discussed above, the accuracy of the measurements could be 24 significantly impacted by O_x production in the reference flow tube. 25 $P(O_x)_{OPR}$ was calculated from Eq. (6), using an O₃-to-NO₂ conversion efficiency of 100%, and is also shown in Figure 6. As discussed above, $P(O_x)_{OPR}$ underestimates the modeled $P(O_x)_{atm}$, 26 27 mainly due to significant O_x production in the reference flow tube. The scatter plot shown as insert in

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1 this figure indicates that a negative bias of approximately 20% would be observed for P(O_x) 2 measurements performed on this day. A negative bias ranging from 15-20% was observed during the 3 other three days that were modeled (Figure S11). 4 As mentioned in the experimental section, concentrations of peroxy radicals obtained as model 5 outputs from the modeling of $P(O_x)_{atm}$ were constrained for the simulations inside the flow tubes, 6 assuming that most of these species are not lost if a short high-flow rate sampling inlet is used. 7 However, simulations were also performed without constraining the peroxy radicals to assess the 8 impact on the simulation results. These simulations have shown that P(O_x) are lower by 10% and 30% 9 in the ambient and reference flow tubes, respectively, when peroxy radicals are not constrained. 10 Overall, the measured ozone production, which is the difference between P(O_x) in the two flow tubes, 11 would only decrease by 2-4%. Therefore, not constraining peroxy radicals in the simulations does not 12 impact the comparison between $P(O_x)_{atm}$ and $P(O_x)_{OPR}$, with $P(O_x)_{OPR}$ underestimating $P(O_x)_{atm}$ 13 by 15-20 %. 14 However, the reason for this disagreement depends on whether peroxy radicals are constrained. When 15 peroxy radicals are constrained, the disagreement is mainly caused by Ox production in the reference 16 flow tube. On the opposite, when peroxy radicals are not constrained, this disagreement is due to an 17 underestimation of $P(O_x)_{atm}$ by $P(O_x)_{amb}$. This underestimation is the result of a latency in the first 18 part of the ambient flow tube due to the time needed to reproduce the radicals, which is on the order 19 of 1-2 minutes. It is very likely that only a fraction of the peroxy radicals will be transferred to the 20 flow tubes and a combination of the two issues discussed above will lead to the negative bias of 15-21 20%. 22 3.2.2 Sensitivity tests - Assessment of the impact of operating conditions on OPR 23 measurements 24 Figure 7 shows the dependence of P(O_x)_{OPR} on the O₃-to-NO₂ conversion efficiency, O₃ and NO₂ 25 surface-losses, surface-production of HONO, and a dilution of the sampled air. The results are displayed for two different times of the day, characterized by different O₃ and NO₂ mixing ratios, 26 27 which have been identified as upper (orange squares) and lower (blue squares) limits for the impact

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1 on the P(O_x) measurements. In addition, these results are also displayed using daily averaged values 2 (green triangles), which are more representative of the average impact of a particular parameter on 3 P(O_x) measurements. The figures described below are for the CalNex campaign during 30 May 2010. 4 Results from the other days are shown in the supplementary material (Figures S12-S14). 5 Figure 7-a shows that $P(O_x)_{OPR}$ is very sensitive to the O₃-to-NO₂ conversion efficiency. For 6 instance, a conversion efficiency of 85% would lead to an underestimation of the P(O_x) measurements 7 by 20-60% (≈35% on average), depending on the chemical composition of the air mass. It is 8 interesting to see that the change in $P(O_x)_{OPR}$, expressed as the ratio between $P(O_x)_{OPR}$ at a 9 conversion efficiency lower than 100% and $P(O_x)_{OPR}$ at a conversion efficiency of 100% (base 10 simulation), changes linearly with the conversion efficiency. The slope of the straight line can be used 11 as an indicator to gauge the impact of the conversion efficiency on P(Ox) measurements throughout 12 the day. As can be seen from Equation (6), for the limiting case of C=0, the measured P(O_x) is 13 determined by the absolute NO2 difference between the two flow tubes. The O3-NOx PSS is shifted 14 towards NO2 in the reference flow tube, due to the lack of NO2 photolysis, reducing the NO2 15 difference between the two tubes and lowering the measured P(O_x). These results stress out the need 16 to reach a conversion efficiency better than 98% to keep this artifact below 5%. The OPR instrument 17 described in this study exhibits a conversion efficiency higher than 99.9% and is not impacted by this 18 issue. 19 Relative surface-losses of 3% and 5% have been observed for NO₂ and O₃, respectively, during the 20 laboratory and field testing (section 3.1.2). Figure 7-b shows that a relative NO₂ loss of 3% in the flow 21 tubes can lead to an overestimation of up to 8% (≈3% on average). On the other hand, Figure 7-c 22 shows that a 5% relative loss of O_3 can lead to an underestimation of up to 30% (\approx 5% on average). 23 These contrasting effects can be explained as follows; ozone in the reference flow tube is lower than 24 in the ambient flow tube, due to the conjunction of a lower production rate of ozone and a shift of the 25 O₃-NO_x PSS towards NO₂. A similar relative loss of ozone in the two flow tubes will therefore lead to 26 a larger absolute loss of Ox species in the ambient flow tube, which in turn will lead to an 27 underestimation of the P(O_x) measurements (Eq. (6)). In contrast, NO₂ is higher in the reference flow

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1 tube and a loss of NO₂ will lead to a larger absolute loss of O_x species in the reference flow tube, and

2 as a consequence, to an overestimation of the $P(O_x)$ measurements.

3 Figure 7-d shows how an heterogeneous production of HONO can impact the P(O_x) measurements. In

4 these simulations, a HONO source was added in the model, with a production rate of 10 ppbv h⁻¹ in

5 both flow tubes (dark HONO production) and an additional varying production rate in the ambient

flow tube (enhanced HONO production). The x-axis presents the HONO production rate in the

ambient flow tube, where 10 ppbv/h corresponds to the dark production only. Moreover, this figure

8 indicates that HONO production rates of 20 ppbv h⁻¹ in the ambient flow tube, similar to experimental

observations, can lead to an overestimation of the $P(O_x)$ measurements by up to 40% (\approx 27% on

average). This overestimation results from HONO photolysis in the ambient tube, which leads to

additional OH production, which in turn leads to an enhancement of VOC oxidation rates and ozone

12 production. Additional simulations were also performed assuming that NO₂ molecules lost on the

surface were equally converted into HONO in both flow tubes (Fig. 7-f), although it is unlikely that

14 the conversion yield of NO₂ into HONO is 100%. The results indicate that, for a relative NO₂ loss of

3%, P(O_x) could be overestimated by up to 15% (10% on average). Note that the impact of this

16 HONO formation adds up to the previously discussed overestimation due to the NO₂ loss.

17 Figure 7-e displays how the injection of zero air at the periphery on the PTFE inlets impacts P(O_x)

measurement through a dilution of the sampled air. As can be seen from this figure, a 10% dilution

leads to less than 9% underestimation of $P(O_x)$.

20 Additional sensitivity tests (not shown) were performed to test the impact of a temperature increase in

the reference flow tube due to heat release by the UV filter, as well as reactions of OH with NO₂

species that produce NO₂. A temperature increase of 5% in the reference flow tube (1° C increase at

23 20° C) can lead to an underestimation of up to 5%, while the O_x production from reactions of OH with

NO_z species can lead to an overestimation of up to 3%.

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3.2.3 Conclusions on potential biases on $P(O_x)_{OPR}$ measurements

2 From the above discussion, we can conclude that there are two main sources of errors. The first source

3 of errors is due to O_x production in the reference flow tube and the latency for ROx reformation in the

4 ambient flow tube, the extent of each depending on the fraction of ambient peroxy radicals that is

5 transmitted into the flow tubes. The combination of these two issues can lead to an underestimation of

6 ambient $P(O_x)$ by 15-20% on average for the conditions observed during MCMA-2006 and CalNex-

7 2010. The second main source of errors is caused by a surface-production of HONO in the ambient

flow tube. Based on a HONO production rate of 20 ppbv h^{-1} , $P(O_x)$ would be overestimated by

9 approximately 30% on average. Additional sources of errors are due to the 4.9% uncertainty on the

flow tube residence time, 5% O₃ and 3% NO₂ surface-losses, the dilution by 10% of the sampled air, a

possible temperature increase of 5% in the reference flow tube and O_x production from reactions of

12 OH with NO₂ species. Daily averaged values and upper bounds of errors associated with these factors,

as derived from all modeled days, are reported in Table 1.

14 Based on the daily average values reported in Table 1, direct sums of the potential negative and

15 positive biases lead to -44% and +40%, respectively. However, the magnitude of each error will

16 depend on atmospheric composition and positive errors will, to some extent, cancel out with negative

errors. A quadratic sum of all these potential errors leads a range of $\pm 36\%$. The estimation of these

18 errors are based on ambient conditions observed in two different environments, with different air

19 compositions for 4 different days. It is safe to assume that similar error values would be observed in

20 other urban environments.

3.3 Current limitations for field operation

22 As mentioned in section 2.4, OPR measurements were performed during the IRRONIC field

campaign. Figure 8 displays time series for a subset of measurements performed from 10-14 July

24 2015, including two anthropogenic VOCs (toluene and acetylene), a biogenic VOC (isoprene) and

25 inorganic species (O₃, NO and NO₂). It is clear from this figure that the measurement site was mainly

impacted by biogenic emissions, with isoprene reaching at least 5 ppbv most of the days, while

27 anthropogenic VOCs were low (<500 pptv). In addition, NO_x levels were lower than 3 ppbv on these

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1 days, confirming the low impact of anthropogenic emissions. These observations indicate that the 2 photochemistry was mainly driven by the oxidation of biogenic VOCs under low NOx conditions, 3 similar to that observed in other forested areas (Griffith et al., 2013). Isoprene is very reactive with the 4 hydroxyl radical and the strong diurnal variation of this species led to a large range of OH reactivity (a few s⁻¹ up to 30 s⁻¹, not shown). The conjunction of the latter with low levels of NO_x makes this site 5 6 of particular interest to study the sensitivity of ozone formation to NO_x by adding NO_x in the OPR 7 instrument as described in the experimental section (section 2.4). 8 Due to the low levels of ambient NOx, ozone production rates at the site were lower than the OPR detection limit of 6.2 ppbv h⁻¹ (section 3.1.5). Indeed, P(O_x) calculations based on total peroxy radical 9 10 measurements performed using the Peroxy Radical Chemical Amplifier technique indicated peak ozone production rates of approximately 2 ppbv h⁻¹ (not shown). Ambient measurements performed 11 12 by the OPR instrument without addition of NO should therefore be scattered around zero within the 13 measurement precision. Figure 8 also displays ΔO_x values (difference in O_x mixing ratios between the 14 two flow tubes) measured by the instrument without the addition of NO (ΔO_x^{2ero} , blue diamonds). 15 While ΔO_x^{zero} was scattered around zero during nighttime, it consistently exhibited large negative 16 values during daytime (-1 to -5 ppbv), indicating that O_x mixing ratios in the ambient flow tube were 17 lower than in the reference flow tube. 18 It is interesting to note that ΔO_x^{zero} values are anticorrelated with J(NO₂) (Fig. 8). Covering the 19 ambient flow tube with a similar UV filter than the reference flow tube, i.e. operating the two tubes 20 under similar irradiation, showed that ΔO_x increases towards less negative values and ultimately 21 reaches zero. This behavior indicates that the higher loss rate of Ox species in the ambient flow tube is 22 due to the solar irradiation and points towards a photo-enhanced surface loss of O_x species initiated by 23 photons at wavelengths lower than 400 nm. As ambient NO₂ mixing ratios were much lower than the 24 observed loss of O_x, this photo-enhanced loss involves a loss of ozone. For an ambient O₃ level of 40 ppbv, as usually observed during the field measurements, a ΔO_x^{zero} of -3 ppbv corresponds to a 7.5% 25 difference in O₃ losses between the two flow tubes and an ozone loss rate higher by approximately 39 26 27 ppbv h⁻¹ in the ambient flow tube compared to the reference flow tube. This issue was further

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2 and ozone confirmed that light-induced processes at wavelength lower than 400 nm lead to a loss of 3 ozone at the surface of the ambient flow tube. It was found that this loss depends on ambient ozone 4 levels, J-values and absolute humidity. 5 This version of the OPR instrument is therefore not suitable to perform ambient $P(O_x)$ measurements 6 since the measured ΔO_x is a combination of ambient ozone production and surface-O₃ losses in the 7 ambient flow tube. For this reason, the OPR measurements were focused on investigating the 8 sensitivity of $P(O_x)$ to NO_x , by recording the relative change in $P(O_x)$ when the chemical composition of ambient air was perturbed by an addition of NO. For these measurements, it is assumed that ΔO_x^{zero} 9 is representative of the instrumental zero and ΔO_x^{zero} measurements are referred as "baseline" in the 10 following. ΔO_x measurements performed with an addition of NO are assumed to deviate from ΔO_x^{zero} 11 due to a change in ozone production in the ambient flow tube, while the surface loss of ozone is 12 13 assumed to be unchanged. This measurement step is denoted ΔO_x^{NO} . The difference between ΔO_x^{zero} and ΔO_x^{NO} divided by the residence time in the flow tubes therefore provides a quantification of the 14 15 change in $P(O_x)$, referred as $\Delta P(O_x)$, due to the addition of NO. The validity of the assumption that the 16 O₃ photo-enhanced surface-loss is not disturbed by the addition of NO is discussed below. 17 Investigating the ozone production sensitivity to NO is outside the scope of this paper and we only 18 present measurements performed when 6 ppbv of NO were added in the instrument to illustrate its current performances and limitations. Figure 8 displays time series of ΔO_x^{NO} (orange diamonds) when 19 20 6 ppbv of NO were added in the flow tubes. When NO is added, there is almost no change in ΔO_x 21 during nighttime. In the absence of sunlight, NO only converts O3 into NO2 and the amount of Ox measured by the CAPS monitor does not change. During daytime, ΔO_x^{NO} is higher than ΔO_x^{zero} , 22 suggesting production of ozone in the ambient flow tube. The difference between ΔO_x^{NO} and ΔO_x^{zero} , 23 24 divided by the residence time in the flow tubes, represents the change in ozone production rates and is displayed in the bottom panel of figure 8 as $\Delta P(O_x)$. Changes in ozone production of up to 20 ppbv h 25 ¹, well correlated with J(NO₂), are observed for these days. Ozone production being NO_x-limited in 26

investigated in the laboratory. Tests performed using artificial irradiation and mixtures of humid air

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1 this environment, a positive change in P(O_x) is indeed expected when a small amount of NO_x is added 2 to the flow tubes. 3 However, the assumption that the photo-enhanced surface-loss of ozone does not change when NO is 4 added may breakdown for large NO mixing ratios. Indeed, the addition of NO in the flow tubes leads 5 to the conversion of a significant fraction of O₃ into NO₂, which in turn reduces the absolute loss of O_3 in the ambient flow tube, leading to a shift of the ΔO_x^{zero} baseline to less negative values. $\Delta P(O_x)$ 6 7 values reported in Figure 8 will therefore be the combination of a change in ozone production and a 8 change in the absolute loss of O₃. If the change in the ozone loss rate is significant compared to the 9 change in the ozone production rate, this could lead to an overestimation of the change in ozone 10 production. An assessment of this measurement bias requires modeling the chemistry in both flow 11 tubes to separate the two contributions, i.e the changes in (i) ozone production and in (ii) ozone loss. 12 While this work is outside the scope of this publication, which focuses on the performances and 13 limitations of the OPR instrument, it is interesting to note that preliminary modeling indicates a bias 14 lower than 5 pbbv h⁻¹ when 6 ppbv of NO is added. 15 The field deployment during IRRONIC revealed an additional bias in P(Ox) measurements due to a 16 photo-enhanced loss of ozone at the inner surface of the ambient flow tube and the difficulty to probe 17 changes in P(Ox) when the sampled air mass is perturbed by an addition of NO. Ambient 18 measurements of P(O_x) with the current version of the OPR would necessitates performing frequent 19 zeros of the instrument to track the ozone loss and unfortunately a simple solution to do so was not 20 found. This work shows that the sampling part of the OPR instrument needs to be rethought to remove 21 (or reduce to a negligible level) the photo-enhanced surface-loss of ozone, which is a prerequisite to 22 get an instrument capable of reliable measurements of ozone production rates. 23 Comparison to the MOPS instrument and potential improvements for the OPR 24 instrument 25 Previous studies (Cazorla and Brune, 2010; Baier et al., 2015) have shown that measurements of 26 ambient ozone production rates are feasible. Baier et al. (2015) reported that the zero of their MOPS 27 instrument was achieved by removing the UV filter from the reference chamber for a full day to

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1 record a diurnal profile of ΔO_x , which was then subtracted from the raw ΔO_x measurements on other 2 days. This zeroing procedure was also tested on the OPR instrument, but led to unrealistically high 3 ambient P(Ox) values of approximately 40 ppbv h-1 for the low-NOx forested environment of IRRONIC. This result also suggests that altering the irradiation conditions of the OPR flow tubes 4 5 leads to a wrong zero of the instrument. This zeroing technique seems to provide better results for the 6 MOPS instrument and it is possible that the design used for the MOPS sampling chambers or the 7 material used to build them (FEP) make it less sensitive to light-dependent surface reactions. 8 Since the main artifacts on the OPR instrument are caused by heterogeneous surface-reactions in the 9 flow tubes, i.e. HONO production (section 3.2.2) and ozone losses (section 3.2.2 and 3.3), the flow 10 tubes should be redesigned to reduce the impact of physicochemical processes occurring near the 11 quartz surface on the ozone production chemistry occurring at the center of the tubes. For instance, 12 the diameter of the tubes could be increased to reduce the surface-to-volume ratio, and their lengths 13 could be shortened together with an increase of the total flow rate to reduce the contact-time between 14 trace gases and the walls. A shorter residence time would also lead to a shorter air-exchange-time, 15 which in turn would help minimizing the scatter in ΔO_x measurements and would help improve the 16 time resolution necessary to generate independent P(Ox) measurements. However, a shorter residence 17 time would also lead to a lower detection limit and a tradeoff between these 2 parameters will likely 18 have to be made. 19 Another solution worth investigating would be to minimize surface reactions by coating the inner 20 surface of the flow tubes with Teflon or by applying a chemical treatment on the quartz surface, 21 which should help removing reactive sites. The latter has already been applied for laboratory kinetic 22 experiments to clean reactor surfaces. Interestingly, it was reported that this type of treatment can also 23 reduce HONO production on quartz surfaces (Laufs and Kleffmann, 2016). 24 Regarding the deployment of this OPR instruments in the field, a reliable zeroing method would be 25 suitable for both ambient P(O_x) and P(O_x) sensitivity measurements. An interesting solution would be 26 to introduce a radical scavenger in the flow tubes to supress ozone production, but a suitable 27 compound has yet to be identified.

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4 Conclusions

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2 An instrument dedicated to direct measurements of ozone production rates (OPR) was developed and consists of two quartz flow tubes, an O₃-to-NO₂ conversion unit and an Aerodyne CAPS NO₂ 3 4 monitor. This setup, compared to the NO2-to-O3 conversion approach previously published in the 5 literature, presents the advantage of a conversion efficiency higher than 99.9%, which is independent of ambient O_x levels. Laboratory and field testing performed to characterize the performance of this 6 7 instrument showed that dark losses of O₃ and NO₂ inside the flow tubes are lower than 5% and 3%, 8 respectively. However, it was shown that dark ozone losses can increase after a long exposure of the 9 flow tubes in the field and frequent cleaning steps should be performed during nighttime by flowing 10 humid air and O_3 in the tubes to keep the loss below 5%. 11 A modeling exercise taking advantage of measurements from previous urban field campaigns showed 12 that a latency in ozone production in the ambient flow tube and a net ozone production in the 13 reference flow tube can lead to a 18% measurement underestimation of ambient P(Ox) on a daily 14 average for the conditions of the MCMA-2006 and CalNex-2010 field campaigns. However, the 15 magnitude of this underestimation depends on the chemical composition of ambient air and it is 16 recommended to assess this potential bias for future campaigns. 17 Sensitivity tests performed during the modeling exercise highlighted the importance of a high 18 conversion efficiency, since a conversion of 95%, which is only 5% lower than the maximum, could 19 lead to an underestimation of ambient $P(O_x)$ by approximately 20% on a daily average for the two 20 selected field campaigns. A dark surface loss of ozone in the flow tubes would lead to an 21 underestimation of ambient P(O_x), while a NO₂ loss would lead to an overestimation. On a daily 22 average, an underestimation of 10% and an overestimation of 5% were assessed for an O₃ loss of 5% 23 and an NO2 loss of 2%, respectively. A photo-enhanced production of HONO in the ambient flow tube on the order of 20 ppbv h^{-1} would also lead to an overestimation of ambient $P(O_x)$ by 27% on a 24 25 daily average. Overall, a quadratic sum of these potential biases for the conditions of the two urban 26 field campaigns leads to a range of errors of $\pm 37\%$ on a daily average.

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1 As shown from the first deployment of the OPR instrument, there is an additional bias due to a photo-

2 enhanced loss of O₃ taking place in the ambient flow tube. This requires improving the sampling

3 design to be able to perform reliable ambient measurements. The first field deployment of the OPR

4 instrument was performed in a low NO_x environment, allowing focusing the study on the sensitivity

5 of ozone production to NO_x. Significant changes in ozone production rates were observed (up to 20

6 ppbv h⁻¹) when 6 ppbv of NO_x were added in the flow tubes, consistent with a NO_x-limited production

7 regime.

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Table 1. Sources of errors on $P(O_x)$ measurement. Upper limits and campaign averages of errors assessed from modeling the selected days of the MCMA-2006 and CalNex-2010 field campaigns (see text). FT: Flow Tube

Sources of errors	Value	Negative bias on P(O _x) average (upper limit)		Positive bias on P(O _x) average (upper limit)	
Residence time (s)	271 ± 13*	-4.9%*	(-4.9%*)	+4.9%*	(+4.9%*)
O ₃ production in ref. FT & latency in amb. FT		-18%**	(-20%**)		<u>—</u>
O ₃ loss	5%*	-10%**	(-25%**)		_
NO ₂ loss	<3%*		_	5%**	(+11%**)
HONO production	up to 20 ppbv/h*		_	+27%**	(+40%**)
Dilution of sampled air	10%*	-8%**	(-9% ^{**})		_
Temperature increase in ref. FT	5%***	-3%**	(-5%**)		_
O _x formation from OH+NO _z	_		_	+3%**	(+3%**)
Conservative sum of biases		-44%	(-64%)	+40%	(+59%)

^{*}from laboratory testing; **from model simulations; ***from estimation

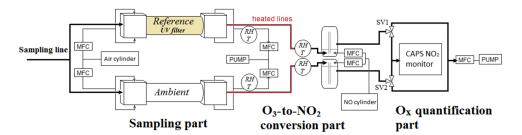


Figure 1. Schematic of the OPR instrument. O_3 converted into NO_2 by reaction with NO. Difference in O_x mixing ratios between the two flow tubes quantified by CAPS. SV: Solenoid Valves. MFC: Mass Flow Controller.

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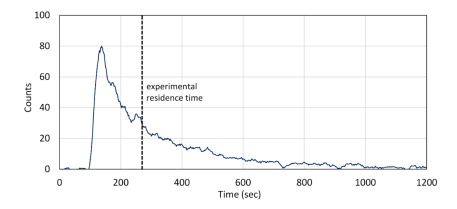
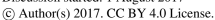


Figure 2. Example of pulse experiments for the quantification of the flow tubes residence time. Pulse of toluene generated at the entrance of the flow tube at t=0s.

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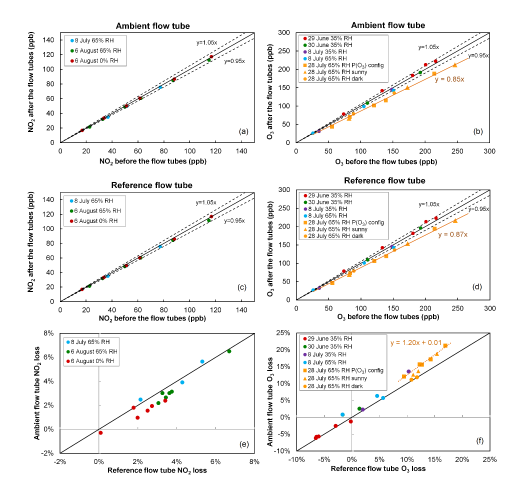


Figure 3. NO_2 and O_3 relative losses measured during the IRRONIC field campaign at different relative humidity values. Losses in the ambient and reference flow tubes are shown in the top and middle panels, respectively. The bottom panel reports the difference in relative losses between the 2 flow tubes. On 28 July O_3 losses were measured under sunny conditions (orange squares: ambient flow irradiated and reference flow tube covered by the UV filter; orange triangles: both flow tubes irradiated), and dark conditions (orange circles: both flow tubes covered by an opaque cover).

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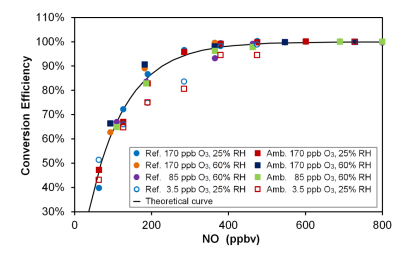
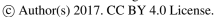


Figure 4. O_3 -to- NO_2 conversion efficiency for various NO mixing ratios, O_x levels and relative humidity values. The black curve was calculated from the reaction rate constant between O_3 and NO and a reaction time of 23 s. Open symbols (3.5 ppbv O_3) are hidden behind the plain symbols for NO>500 ppbv. "Ref." and "Amb." refer to the conversion units coupled to the reference and ambient flow tubes, respectively.

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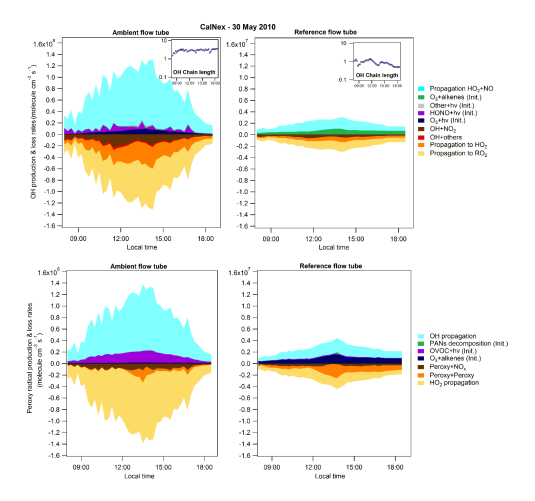
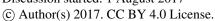


Figure 5. OH (top) and total peroxy (HO_2+RO_2 , bottom) radical budgets for 30 May 2010 of the CalNex-2010 campaign. Radical budgets modeled for the ambient (left) and the reference (right) flow tubes. The OH chain length is also presented in an insert (top) for each flow tube. The note (Init.) in the legend indicates initiation reactions.

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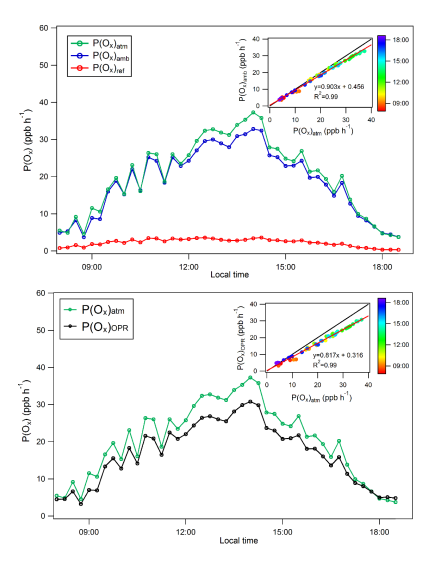


Figure 6. Modeling comparison of P(Ox) values. Top: ozone production rates modeled for the atmosphere, $P(O_x)_{atm}$, the ambient flow tube, $P(O_x)_{amb}$, and the reference flow tube, $P(O_x)_{ref}$ for 30 May 2010 of the CalNex-2010 campaign. Bottom: comparison of modeled ozone production rates for the OPR, P(Ox)OPR, and the atmosphere, $P(O_x)_{atm}$, for 30 May 2010. Inserts: correlations between $P(O_x)_{atm}$ and $P(O_x)_{amb}$ (top), and $P(O_x)_{atm}$ and P(O_x)_{OPR} (bottom), color-coded by the time of day.

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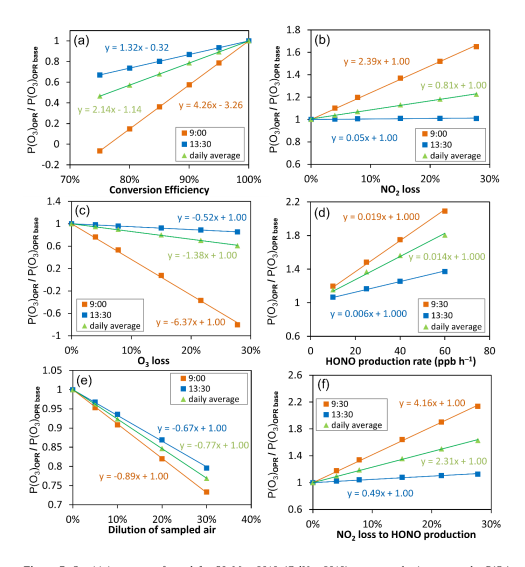
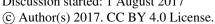


Figure 7. Sensitivity tests performed for 30 May 2010 (CalNex-2010) to assess the impact on the $P(O_x)$ measurements of (a) the O_3 -to- NO_2 conversion efficiency, (b) NO_2 and (c) O_3 dark losses, (d) heterogeneous HONO formation, (e) dilution of ambient air, and (f) NO_2 loss towards HONO production in the flow tubes. The results presented here correspond to the two hours of the day identified as lower (blue squares) and upper (orange squares) limits of the impact on the $P(O_x)$ measurements. The daily average behavior is also shown using green triangles.

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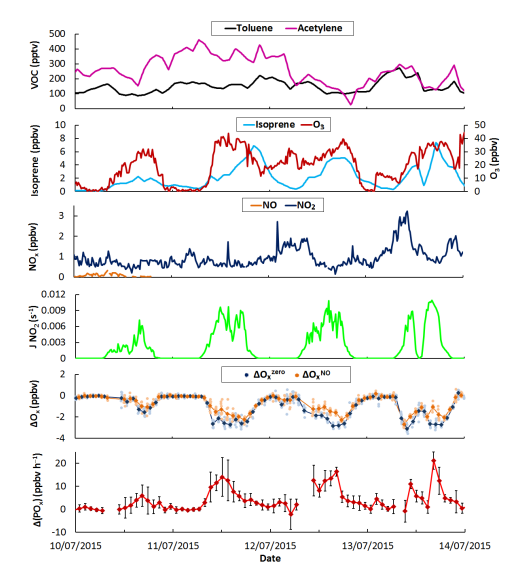


Figure 8. Time series of selected trace gases, $J(NO_2)$, measured ΔO_x and $\Delta P(O_x)$ values during four days of the IRRONIC campaign when 6 ppbv of NO was intermittently added in the flow tubes. The light colors on ΔO_x correspond to 2-min measurements while the darker colors are 20-min averaged values. Error bars on $\Delta P(O_x)$ are 1σ on the averaged 20-min measurements.