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# Interferences in photolytic NO<sub>2</sub> measurements: explanation for an apparent missing oxidant?

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Measurement of NO<sub>2</sub> at low concentrations is non-trivial. A variety of techniques exist, with the conversion of NO2 into NO followed by chemiluminescent detection of NO being prevalent. Historically this conversion has used a catalytic approach (Molybdenum); however this has been plaqued with interferences. More recently, photolytic conversion based on UV-LED irradiation of a reaction cell has been used. Although this appears to be robust there have been a range of observations in low NO<sub>x</sub> environments which have measured higher NO2 concentrations than might be expected from steady state analysis of simultaneously measured NO, O<sub>3</sub>, JNO<sub>2</sub> etc. A range of explanations exist in the literature most of which focus on an unknown and unmeasured "compound X" that is able to convert NO to NO<sub>2</sub> selectively. Here we explore in the laboratory the interference on the photolytic NO<sub>2</sub> measurements from the thermal decomposition of peroxyacetyl nitrate (PAN) within the photolysis cell. We find that approximately 5% of the PAN decomposes within the instrument providing a potentially significant interference. We parameterize the decomposition in terms of the temperature of the light source, the ambient temperature and a mixing timescale (~0.4s for our instrument) and expand the parametric analysis to other atmospheric compounds that decompose readily to NO<sub>2</sub> (HO<sub>2</sub>NO<sub>2</sub>, N<sub>2</sub>O<sub>5</sub>, CH<sub>3</sub>O<sub>2</sub>NO<sub>2</sub>, IONO<sub>2</sub>, BrONO<sub>2</sub>, Higher PANs). We apply these parameters to the output of a global atmospheric model (GEOS-Chem) to investigate the global impact of this interference on (1) the NO<sub>2</sub> measurements and (2) the NO<sub>2</sub>: NO ratio i.e. the Leighton relationship. We find that there are significant interferences in cold regions with low NO<sub>x</sub> concentrations such as Antarctic, the remote Southern Hemisphere and the upper troposphere. Although this interference is likely instrument specific, it appears that the thermal decomposition of NO<sub>2</sub> within the instrument's photolysis cell may give an explanation for the anomalously high NO<sub>2</sub> that has been reported in remote regions, and would reconcile measured and modelled NO2 to NO ratios without having to invoke novel chemistry. Better instrument characterization, coupled to instrumental designs which reduce the heating within the cell seem likely

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#### 1 Introduction

Accurate quantification of atmospheric nitrogen oxide ( $NO_x$  which is predominantly  $NO + NO_2$  but includes small contributions from  $NO_3$ ,  $N_2O_5$ , HONO,  $HO_2NO_2$  etc.) concentrations is crucial for many aspects of tropospheric chemistry.  $NO_x$  plays a central role in the chemistry of the troposphere, mainly through its impact on ozone ( $O_3$ ) and hydroxyl (OH) radical concentrations.  $O_3$  is a climate gas (Wang et al., 1995), adversely impacts human health (Mauzerall et al., 2005; Skalska et al., 2010) and leads to ecosystem damage (Ainsworth et al., 2012; Ashmore, 2005; Hollaway et al., 2012). It is produced through the reaction of peroxy radicals ( $HO_2$  and  $RO_2$ ) with NO (Dalsøren and Isaksen, 2006; Lelieveld et al., 2004). The OH radical is the primary oxidizing agent in the atmosphere (Crutzen, 1979; Levy II, 1972) as it controls the concentration of other key atmospheric constituents such as methane ( $CH_4$ ), carbon monoxide (CO) and volatile organic compounds (VOCs). It is both produced through the reaction of NO with NO0 and is lost by its reaction with NO1. NO2 itself poses a public health risk (Stieb et al., 2002). Thus understanding the sources, sinks and distribution of  $NO_x$  is of central importance to understanding the composition of the troposphere.

During the daytime there is fast cycling between NO and  $NO_2$ , due to the rapid photolysis of  $NO_2$  and the reaction NO and  $O_3$  to form  $NO_2$  (Kley et al., 1981).

$$NO_2 + hv(< 410 \text{ nm}) \rightarrow NO + O(^3P)$$
 (R1)

$$O_2 + O(^3P) + M \rightarrow O_3 + M$$
 (R2)

$$O_3 + NO \rightarrow O_2 + NO_2 \tag{R3}$$

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$$1 = \frac{k_1[NO][O_3]}{j_{NO_2}[NO_2]} = \phi$$
 (1)

The quantities in the relationship are readily measured and deviations from unity have been interpreted to signify missing (i.e. non ozone) oxidants of NO. These perturbations have been used to infer the existence of oxidants such as peroxy radicals, or halogen oxides in the atmosphere (Bauguitte et al., 2012; Cantrell et al., 2003; Frey et al., 2015).

The concentration of  $NO_x$  varies from > 100 ppb (parts per billion) next to roads (Carslaw, 2005; Pandey et al., 2008) to low ppt (parts per trillion) in the remote atmosphere (Lee et al., 2009). Direct transport of  $NO_x$  from polluted to remote regions is not efficient, because  $NO_x$  is removed from the atmosphere on a timescale of around a day by the reaction of  $NO_2$  with OH and the hydrolysis of  $N_2O_5$  on aerosol surfaces (Brown et al., 2004; Dentener and Crutzen, 1993; Riemer et al., 2003). Instead, reservoir species such as peroxy-acetyl nitrate are made in polluted regions (which are high in both  $NO_x$  and peroxy-acetyl precursors such as acetaldehyde) and are subsequently transported to remote regions where they thermally breakdown to release the  $NO_x$ .

$$CH_3CHO + OH^{\bullet} + O_2 \rightarrow CH_3C(O)OO^{\bullet} + H_2O$$
 (R4)

$$CH_3C(O)OO^{\bullet} + NO_2 + M \rightleftharpoons CH_3C(O)O_2NO_2 + M$$
(R5)

The equilibrium between peroxy-acetyl radicals,  $NO_2$  and PAN (Reactions R4 and R5) is highly temperature sensitive. Thus the PAN lifetime changes from 30 min at 25 °C (Bridier et al., 1991) to 5.36 years at -26 °C (Kleindienst, 1994).

Measurements of  $NO_x$  species in the remote atmosphere have been made over the last 40 years. Multiple in-situ techniques are available such as LIF (Matsumoto and Kajii, 2003), Cavity Ring Down (Osthoff et al., 2006), and QCL – Quantum Cascade Laser (Tuzson et al., 2013). However, probably the most extensively used approach has been based on the chemiluminescent reaction between NO and  $O_3$ . This exploits

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$$NO + O_3 \rightarrow NO_2^* \tag{R6}$$

$$_{5} \text{ NO}_{2}^{*} \rightarrow \text{NO}_{2} + hv \quad (>600 \,\text{nm})$$
 (R7)

This forms the basis of the chemiluminescence analysis of NO (Drummond et al., 1985; Fontijn et al., 1970; Kelly et al., 1980; Peterson and Honrath, 1999). The number of photons emitted by the decay of excited  $NO_2^*$  to  $NO_2$  is proportional to the NO present before reaction with  $O_3$  (Drummond et al., 1985). The photons emitted are detected by a cooled photomultiplier tube (PMT) with the sample under low pressure (to maximize the fluorescence lifetime of the  $NO_2^*$ ), to yield a signal which is linearly proportional to the number density of NO in the sample gas (Fontijn et al., 1970).

With NO chemiluminescence analysers it is also possible to analyse  $NO_2$  if it is first converted to NO, either catalytically (typically heated Molybdenum) as in Reaction (R8) (Villena et al., 2012), or by converting  $NO_2$  into NO photolytically (Ryerson et al., 2000), exploiting Reaction (R1).

$$Mo + 3NO_2 \rightarrow MoO_3 + 3NO$$
 (R8)

To measure NO and  $NO_2$ , the sample flows through the  $NO_2$  to NO converter (of either type) to the reaction chamber where the  $NO + O_3$  reaction occurs and the decay of  $NO_2^*$  to  $NO_2$  allows the concentration of  $NO + NO_2$  in the air to be measured. Then, the sample flow is switched to bypass the  $NO_2$  to NO converter. Now, only NO present in the sample is detected in the chemiluminescence reaction. The NO signal is then subtracted from the  $NO + NO_2$  ( $NO_x$ ) signal giving the  $NO_2$  signal.

Measurements of NO and  $NO_2$  have been successfully made in a range locations using the chemiluminescence technique (Huntrieser et al., 2007; Lee et al., 2009; Peterson and Honrath, 1999; Zhang et al., 2008). However, measurements made in remote (low  $NO_x$ ) locations, such as Antarctica and in the open ocean have at times

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identified an unexplained imbalance in the Leighton relationship (Cantrell et al., 1997; Hosaynali Beygi et al., 2011). Measured  $NO_2$  concentrations are higher than would be expected from the observed  $NO_3$ ,  $JNO_2$  along with reasonable concentrations of other oxidants (peroxy radicals, halogen oxides). Various explanations have been posited in order to overcome the apparent oxidation gap, typically relying on a mystery unmeasured oxidant, or pushing known chemistry into theoretical realms by theorizing high turnover of short-lived species which may only have been measured in trace quantities (Cantrell et al., 2003; Frey et al., 2013, 2015; Hosaynali Beygi et al., 2011). An alternative explanation would be an unknown interference on the  $NO_2$  measurement increasing its apparent concentration.

Here we explore the potential of PAN to interfere with chemiluminescence  $NO_2$  measurements. In Sect. 2 we provide some details of the experimental studies undertaken. In Sect. 3 we describe the results of experiments introducing differing concentrations of PAN into  $NO_2$  converter/chemiluminescence systems. In Sect. 4 we analyse the potential for errors with different  $NO_x$  systems to investigate the interference on the measurement of  $NO_2$  from PAN. In Sect. 5 we evaluate the impact of this interference on  $NO_2$  measurements and on the Leighton relationship through the use of a global model and provide conclusions in Sect. 6.

#### 2 Experimental details

In Sect. 2.1 we describe the two chemiluminescence instruments used for the analysis. The NO<sub>2</sub> converters are described in Sect. 2.2. In Sect. 2.3 and the LIF instrument used to provide a reference analysis is described. We describe our protocol for production of PAN by acetone photolysis in Sect. 2.4. We provide details of the zero air generation in Sect. 2.5. Then in Sect. 2.6 we describe the experimental methodology of PAN interference tests and residence time tests in Sect. 2.7.

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Chemiluminescent measurements were performed using dual channel Air Quality Design Inc. (Golden, Colorado, USA) instruments equipped with UV-LED based photolytic NO<sub>2</sub> converters - commonly referred to as Blue Light Converters (BLCs). Two similar instruments were employed; the "laboratory" NO<sub>x</sub> analyser (Sect. 2.1.1) – on which the majority of the experiments were performed and the "aircraft" NO, analyser (Sect. 2.1.2) on which only temperature controlled BLC experiments were performed.

Both instruments feature independent mass flow controlled sample flows on each channel (NO and NO<sub>x</sub>). The wetted surfaces of the instrument are constructed of 1/4 inch PFA tubing, with the exception of 316 stainless steel unions/MFC internals.

Both instruments are calibrated for NO by internal, automatic standard addition. Calibration for NO<sub>2</sub> converter efficiency is by internal automatic gas phase titration of NO with O<sub>3</sub> to form NO<sub>2</sub> with the NO signal measured with the BLC lamps active and inactive as described by (Lee et al., 2009). Artefacts in both NO and NO2 are measured whilst sampling zero air.

#### 2.1.1 Laboratory NO<sub>x</sub> analyser

NO and NO<sub>2</sub> measurement – the other 60 % of the duty cycle is devoted to NO (40 %)

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The laboratory NO<sub>x</sub> analyser from Air Quality Design, Inc. (AQD) is a custom dual channel instrument designed for fast response and very low limit of detection (LOD). The dual channel design means that there are effectively two separate NO chemiluminescence instruments working in parallel. Both channels have identical flow paths and share identical duplicate equipment; ozonizers, MFCs, PMTs etc. Both channels share the same vacuum pump - an Edwards XDS 35i. One channel is equipped with a BLC immediately in front of the MFC flow control/low pressure side of the system. It is possible to analyse NO with one channel, and NO, with the other to provide a constant, fast measurement (1 Hz) of NO and NO<sub>2</sub>. Alternatively, a single channel can be used with the BLC in a switching mode so that it is active for only 40 % of the duty cycle to provide

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and to measuring zero (20%). In this case the second channel might be used for NO<sub>V</sub> by connection of a catalytic converter to the inlet as is the set-up at the Cape Verde Atmospheric Observatory GAW station (Lee et al., 2009). In these experiments both modes were used in order to replicate different instrument designs; a switching mode <sub>5</sub> with 40 % duty cycle of the NO<sub>2</sub> converter and a total sample flow of 1 standard L min<sup>-1</sup>, and a parallel mode with 100 % duty cycle of the NO<sub>2</sub> converter with a total sample flow of 2 standard L min<sup>-1</sup>.

The nominal sensitivity of the instrument is 3.5 and 4.0 cps ppt<sup>-1</sup> on channel 1 (NO) and channel 2 (NO and NO $_{v}$ ) respectively. The 1 min LOD is  $\sim$  2.5 pptv.

### 2.1.2 Aircraft $NO_x$ analyser

The aircraft NO<sub>x</sub> analyser, also from AQD, operates similarly to the lab NO<sub>x</sub> analyser with some alterations to make it suited to aircraft operation but which do not affect its use on the ground. It can therefore be considered analogous to the lab NO, analyser with the exception of the BLC which is of a non-standard design in that it uses six more powerful UV-diodes which require active Peltier/forced air-cooling controlled by a PID in order to maintain an operating temperature close to ambient. The special requirements for this NO<sub>2</sub> converter are primarily because of the high sample flow rates needed to measure NO<sub>v</sub> fluxes on an airborne platform at reduced pressure. However, in this study the sample flow rate was a constant 1 standard L min<sup>-1</sup> per channel at ambient temperature and pressure.

The nominal sensitivity of the instrument is 8.3 and 11.6 cps ppt<sup>-1</sup> on channel 1 (NO) and channel 2 (NO<sub>x</sub>) respectively. The 1 min LOD is  $\sim$  1.0 pptv.

#### NO<sub>2</sub> converters

Photolytic converters for the two chemiluminescent systems were supplied by Air Quality Design and manufactured according to their proprietary standards (Buhr, 2004, 2007). Systems have also been developed subsequently (Pollack et al., 2011;

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Photolytic converters employ Reaction (R1) to convert NO2 to NO over a narrow 5 wavelength band, thus providing a more selective NO<sub>2</sub> measurement to that provided by molybdenum catalysts (Ridley et al., 1988; Ryerson et al., 2000). The conversion efficiency is determined by Eq. (2) where t is the residence time within the photolysis cell. Here k[Ox] is the concentration and rate constant of any oxidant that reacts with NO to form NO<sub>2</sub> (Ryerson et al., 2000).

10 CE = 
$$\left[ \frac{jt}{it + k \lceil Ox \rceil t} \right] \left[ 1 - \exp^{(jt - k \lceil Ox \rceil t)} \right]$$
 (2)

The rate constant of photolysis of  $NO_2(j)$ , and so, the rate of production of additional NO beyond that in the original sample is given in Eq. (3).

$$j(T) = \int_{\lambda \min}^{\lambda \max} F(\lambda) \sigma(\lambda, T) \phi(\lambda, T) d\lambda$$
(3)

In Eq. (3) j is the rate constant (s<sup>-1</sup>), F is the spectral photon flux (photons cm<sup>-2</sup> s<sup>-1</sup> nm<sup>-1</sup>),  $\sigma$  is the absorption cross section of NO<sub>2</sub> (cm<sup>2</sup>),  $\phi$  is the quantum yield (dimensionless) of NO<sub>2</sub> photo-dissociation, and T is the temperature (Sander et al., 2011). The j value of the converter is practically determined by the irradiant photolysis power of the UV emitting elements and how efficiently the power is used.

#### Standard BLC 2.2.1

Standard BLCs consist of two ends housing the UV-LEDs (1 W, 395 nm, UV Hex, Norlux Corp) within a heat sink to which is attached a cooling fan. The ends are bolted to a central section with rubber gaskets forming an air-tight seal. Within the centre section 28707

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a propriety Teflon-like material block is housed which serves as a highly UV reflective (> 0.95) cavity thought which the sample gas flows (Buhr, 2007). On two of the opposing sides of the centre section are 1/4 inch Swagelok fittings acting as an inlet and outlet for the sample gas.

The volume of this illuminated sample chamber is 16 mL which, with a standard flow rate of 1 standard L min<sup>-1</sup>, gives a sample residence time of 0.96 s. Additional lamp end units were also supplied by AQD.

The conversion efficiency of the standard BLC with a sample flow of  $1 \operatorname{standard} \operatorname{Lmin}^{-1}$  was 22 to 42% ( $j = 0.2...0.6 \operatorname{s}^{-1}$ ) depending on the combination of lamp units used whilst the external temperature of the converter was typically 34 to 45 °C. All experiments were carried out at with sample gas at ambient temperature and pressure; 20 °C, 1 atm.

#### 2.2.2 High power BLC

The high power BLC of the aircraft instrument is designed to operate at a higher flow rate (1.5 standard L min<sup>-1</sup>), lower pressure (~ 300 Torr) and therefore lower residence time, to that of the standard BLC to allow fast time resolution measurements from an aircraft. For this reason a greater number (six) of more powerful UV-LEDs [2 W, 395 nm, Nichia Corp] are used in order that the conversion efficiency be acceptable under these conditions. The lamps are placed evenly along two sides of a cylindrical cavity of the same highly UV reflective Teflon with inlets at opposing ends. The high power BLC lamps are actively (Peltier) cooled to 47 °C and without Peltier cooling reach 77 °C. It was therefore possible to control the internal temperature of the BLC by varying the power supplied to the Peltier elements via the temperature controller.

The volume of this illuminated sample chamber is  $10 \,\mathrm{mL}$  which, with a standard flow rate of  $1 \,\mathrm{standard} \,\mathrm{L} \,\mathrm{min}^{-1}$ , gives a sample residence time of  $0.60 \,\mathrm{s}$  resulting in a conversion efficiency of  $93 \,\%$  ( $j = 6.5 \,\mathrm{s}^{-1}$ ).

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The Thermal Dissociation Laser Induced Fluorescence (TD-LIF) system is a custom instrument developed for aircraft and ground-based observations of  $NO_2$ ,  $\Sigma PNs$ , SANs, and HNO<sub>3</sub>. A detailed description of the TD-LIF instrument can be found in Di Carlo et al. (2013), with a short description given here. It uses laser induced fluorescence (LIF) to detect NO<sub>2</sub> concentrations directly (Dari-Salisburgo et al., 2009; Matsumoto and Kajii, 2003; Matsumoto et al., 2001; Thornton et al., 2000) and, coupled with a thermal dissociation inlet system, allows measurement of peroxy nitrates  $(\Sigma PNs)$ , alkyl nitrates  $(\Sigma ANs)$ , and HNO<sub>3</sub> after conversion into NO<sub>2</sub> (Day et al., 2002). The TD-LIF comprises four main parts: the laser source, the detection cells system, the inlet system and the pumps. The laser source is a Nd:YAG pulse doubled laser (Spectra-Physics, model Navigator I) that emits light at 532 nm with a power of 3.8 W, a repetition rate of 15 kHz and 20 ns pulse-width. The detection cells system comprises four identical cells, one for each compound class, to allow simultaneous measurements. Each cell is formed by a cube and two arms where the laser beam passes through the sample air flow in the centre of the cell. Perpendicular to both (laser beam and air flow) there is the detector that is a gated photomultiplier with lens and long pass filters to optimize the fluorescence detection, minimizing the non-fluorescence light that reaches the detector (Di Carlo et al., 2013; Dari-Salisburgo et al., 2009). The pump system includes a roots blower coupled to a rotary vane pump to maintain a flow of 6 L min<sup>-1</sup>. The common inlet system is split in four channels: one at ambient temperature to measure NO<sub>2</sub>, and the last three heated at 200, 400 and 550 °C, to thermally dissociate  $\Sigma$ PNs,  $\Sigma$ ANs, and HNO<sub>3</sub> respectively into NO<sub>2</sub> (Di Carlo et al., 2013). To minimize quenching due to atmospheric molecules, and therefore increase the senACPD

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sitivity of the TD-LIF, each cell is kept at low pressure (3-4 Torr). This increases the fluorescence lifetime and facilitates the time-gating of the photomultiplier to further reduce the background (Dari-Salisburgo et al., 2009). The TD-LIF is routinely checked for background by an over-flow of zero air in the detection cells; and calibrated by standard 5 addition of known amount of NO<sub>2</sub> from a cylinder (NIST traceable) diluted in zero air - the flow of both zero air and NO<sub>2</sub> being MFC controlled. The time resolution of the measurements is 10 Hz and the detection limits are: 9.8, 18.4, 28.1, and 49.7 pptv (1 s, S/N = 2) for NO<sub>2</sub>,  $\Sigma$ PNs,  $\Sigma$ ANs, and HNO<sub>3</sub> cells, respectively (Di Carlo et al., 2013).

#### PAN preparation

In order to test the sensitivity of the instrument to peroxy-acetyl nitrate interferences it was prepared by the photolysis in air of acetone and NO as described by (Meyrahn et al., 1987) and later by (Warneck and Zerbach, 1992; Zellweger et al., 2000). Reactions (R9)-(R11) describe the mechanism by which PAN is formed from acetone photolysis.

 $(CH_3)_2CO + O_2 + hv \rightarrow CH_3C(O)O_2^{\bullet} + CH_3^{\bullet}$ (R9)

 $CH_3C(O)O_2^{\bullet} + NO \rightarrow CH_3C(O)O^{\bullet} + NO_2$ (R10)

 $CH_3C(O)O_2^{\bullet} + NO_2 \rightarrow CH_3C(O)O_2NO_2$ (R11)

Here NO<sub>2</sub> reacts stoichiometrically with the acyl radical to form PAN. In practice, an excess of acetone is used to ensure that NO reacts completely. A minor product also found in the photolysis of acetone is methyl nitrate, MeONO2, which is typically approximately 1% of the total yield (Mills et al., 2007). The proposed origin of the methyl nitrate is shown in Reactions (R12)-(R16). Methyl nitrate is also found in the atmosphere through oceanic emission (Moore and Blough, 2002) and as a product of the thermal decomposition of PAN (Fischer and Nwankwoala, 1995; Roumelis and Glavas, 1992; Warneck and Zerbach, 1992). Reactions (R15), (R17) (Fischer and Nwankwoala, 1995) show that the formation of nitro-methane and HONO is also possible from PAN

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$$2CH_{3}C(O)O_{2}^{\bullet} \rightarrow 2CH_{3}CO_{2}^{\bullet} + O_{2}$$

$$CH_{3}CO_{2}^{\bullet} \rightarrow CH_{3}^{\bullet} + CO_{2}$$

$$CH_{3}^{\bullet} + CH_{3}C(O)O_{2}^{\bullet} \rightarrow CH_{3}O^{\bullet} + CH_{3}CO_{2}^{\bullet}$$

$$(R13)$$

$$5 (CH_{3}^{\bullet} + NO_{2} \rightarrow CH_{3}NO_{2})$$

$$CH_{3}O^{\bullet} + NO_{2} \rightarrow CH_{3}ONO_{2}$$

$$(CH_{3}O^{\bullet} + NO_{2} \rightarrow HCHO + HONO)$$

$$(R16)$$

A dedicated "PAN generator", as used by Whalley et al. (2004), was employed to produce a consistent source of PAN. The generator consists of: flow control elements for the NO standard gas, the acetone flow, and the zero air diluent flow; an acetone permeation oven consisting of a reservoir of HPLC grade acetone (ACS grade, Acros) with a silicone permeation tube placed in the headspace through which zero air flows, all thermo-stated at 30 °C; and a Pyrex glass photolysis cell illuminated by UV light centred at 285 nm (Pen-Ray mercury lamp, UVP). The Pyrex functions to filter wavelengths below 290 nm within the photolysis cell thus minimizing PAN photolysis (Mills et al., 2007).

All flow rates within the PAN generator were calibrated using a Gilian Gilibrator-2 Air Flow Calibrator (Sensidyne). The PAN generator is capable of continuously producing 0.1–20.0 ppbv PAN. Linearity and mixing ratio of the PAN output was confirmed by PAN-GC equipped with an ECD detector as described by Whalley et al. (2004), and also by complete reduction back to NO using a heated (325  $^{\circ}$ C) molybdenum catalyst (Thermo Environmental). A Laser Induced Fluorescence instrument, described in Sect. 2.2.3, was used to find if any NO<sub>2</sub> produced directly in the PAN generator with the results presented in Sect. 2.4.1.

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Zero air was generated from dried ( $-40\,^{\circ}$ C dew point) compressed air by subsequent filtering through a cartridge of molecular sieve ( $13\times$ , Sigma Aldrich) to ensure a consistent humidity throughout all experiments and was regenerated by heating to  $250\,^{\circ}$ C for 24 h when necessary. A second filter cartridge placed after the molecular sieve was packed with Sofnofil (Molecular Products) and activated charcoal (Sigma Aldrich) in order to remove ozone,  $NO_x$  and volatile organic compounds (VOC) which may be present in the compressed air. Zero air generated from the compressed air and filter cartridges system and zero air from an Eco Physics AG PAG 003 pure air generator (the industry standard) were both sampled by the NO chemiluminescence analyser. No difference in the counts of the NO analyser was observed between the two sources of zero air. Thus the NO content of both sources of zero air was considered to be comparably low.

The  $NO_2$  content of any zero air used is critical (more so than NO) in this study. In order to determine the  $NO_2$  content of the zero air sources a direct measurement of  $NO_2$  was required in order to avoid biasing the experimental procedure. The LIF instrument described in Sect. 2.2.3 was used to compare the zero air sources (Table 1).

Zero air from both PAG 003 and filter stack was sampled by the NO $_2$  LIF analyser. Additionally, high grade bottled zero air (BTCA 178, BOC Specialty Gasses) was analyzed for NO $_2$ . Table 1 shows the photomultiplier counts per second whilst sampling 1.5 standard L min $^{-1}$  of zero air. The dark counts of the PMT in the absence of laser light are typically less than 3 counts s $^{-1}$ . The counts recorded are therefore the sum of any NO $_2$  fluorescence and scattered laser light. It is clear that the Sofnofil/Carbon filter system has an advantage over both the PAG 003 and BTCA 178 zero air sources in that a lower signal for NO $_2$  fluorescence was observed. Typical sensitivity of the LIF NO $_2$  channel was  $\sim$  180 cps ppb $^{-1}$ , thus a 4–4.5 cps improvement in zero background equates to 22–25 ppts improvement in accuracy. Consequently, all dilution, zeroing, and PAN generation utilized the Sofnofil/Carbon filter system.

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The NO<sub>v</sub> analysers were first calibrated for sensitivity/converter efficiency and sampling zero air by overflowing the inlet from the internal source prior to the experiment for at least 2 h. This is because the PAN from the generator is in zero air which has a very low dew point and the sensitivity of the NO<sub>x</sub> analyser is reduced by high humidity in the sample. This means that switching the NO, analyser from sampling ambient (humid) air to zero air causes the sensitivity to rise slowly and the humidity inside of the reaction cell to decrease. After establishing an NO flow (4.78 ppm NO in N<sub>2</sub>, BOC specialty gases) of  $0.5\,\mathrm{mL\,min}^{-1}$  into the PAN generator, the acetone flow was then adjusted to ~ 10 mLmin<sup>-1</sup> and the diluent flow of zero air adjusted to achieve the desired output mixing ratio. The internal zero air of the NO<sub>x</sub> analyser was then shut off so that the NO<sub>x</sub> analyser was sampling zero air from the PAN generator. Note that the total flow from the PAN generator always exceeded the sample requirements of the NO<sub>x</sub> analyser with excess flow vented to the atmosphere. The system was then allowed to stabilize until a stable NO value was recorded on the NO, analyser. Next, the acetone photolysis lamp of the PAN generator was switched on so that acetone was photolysed in the presence of NO to form PAN. Complete NO conversion to PAN was indicated by the fact that in all cases the NO signal measured by the NO, analyser fell to ~ 0 ppbv after the acetone/NO mixture was illuminated by the photolysis lamp. The diluent flow from the PAN generator was then varied to achieve PAN mixing ratios of between 0.2 to 1.3 ppbv. The corresponding NO<sub>2</sub> signal was recorded once stable. This procedure was repeated for various combinations of BLC lamps/assemblies and analyser operation modes.

In order to investigate any interference from unreacted acetone, the  $NO_x$  analyser was allowed to sample the output of the PAN generator with the photolysis lamp off i.e. a flow of acetone and NO gas. No additional signal relative to zero air during the NO or  $NO_2$  measurement cycle was observed during these experiments at any mixing ratio.

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To investigate any interference from unreacted peroxy radicals left over from photolysis of excess acetone, produced in Reaction (R6), the  $NO_x$  analyser was allowed to sample acetone that had been photolysed within the PAN generator in the absence of NO. NO was added downstream of the photolysis cell also. No additional signal over zero air during the NO or  $NO_2$  measurement cycle was observed in either case, photolysed acetone with or without nitric oxide addition. Thus, it was concluded that acetone does not interfere within the BLC and that peroxy radicals do not exist outside of the broadband photolysis cell of the PAN generator. It should be noted that sampling acetone does cause an increase in the zero count of the  $NO_x$  analyser as acetone does react with ozone; this chemiluminescence interference is known (Dunlea et al., 2007) and is accounted for in the measurements.

To test whether the PAN generator produced any  $NO_2$  directly (i.e. rather than as a consequence of conversion of PAN to  $NO_2$ ), a direct measurement of  $NO_2$  was employed using the TD-LIF described in Sect. 2.2.3. The measured signal relative to pure zero air was measured in the LIF  $NO_2$  channel when sampling various mixing ratios of PAN from the generator as shown in Fig. 1. It is evident that the  $NO_2$  signal observed while sampling PAN from the generator lies within the noise of the zero signal measurement. In this case each point represents a 10 min average as does the zero measurement. With an averaging time of 10 min the theoretical limit of detection is estimated to be less than 0.1 pptv – taking a 10 Hz LOD of 9.8 pptv, and averaging 6000 points (i.e. 10 min) the precision improves by a factor of approximately 1/n where n is the number of points averaged (Lee et al., 2009). It is therefore conservatively estimated that less than 1 ppt  $NO_2$  at 1000 ppt PAN (0.1%) is produced by the PAN generator. This is less than previously estimated (Mills et al., 2007), albeit at higher PAN mixing ratios and lower residence times within the acetone photolysis cell with the same generator.

It was therefore determined that only PAN could be an interfering species in the BLC from the PAN generator. The small percentage of methyl nitrate which may be produced is discounted due to it being less thermally labile that PAN itself. Additionally, the percentage interference observed is significantly greater than any expected or

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#### Residence time

The residence time of PAN in the 2.7 m PFA inlet linking the PAN generator to the NO, analyser was varied by varying the flow rate. This was achieved by altering the sample flows through the each of the NO<sub>x</sub> analyser channels (which share a common inlet).

#### Impact of PAN on NO<sub>2</sub> measurements

In this section we discuss experiments investigating the potential impact of PAN on the two NO<sub>2</sub> instruments. In Sect. 3.1 and 3.2 we explore the interference in the laboratory instrument with a range of BLC convertors, eliminating any possibility for inlet effects in Sect. 3.3, and in Sect. 3.4 we explore the interference in the aircraft instrument which has an active cooling of the convertor. In Sect. 3.5 we investigate whether photolytic decomposition of PAN could lead to the interferences and in Sect. 3.6 we investigate whether thermal decomposition could be the source.

#### 3.1 Standard BLC and laboratory NO<sub>x</sub> analyser in constant mode

PAN was introduced to the NO<sub>x</sub> analyser, equipped with a BLC as described in Sect. 2.2.1, diluted in zero air through a range of mixing ratios. The resulting mixing ratio recorded by the analyser was recorded and is presented in the following sections. Figure 2 shows that the NO<sub>2</sub> signal is proportional to increasing PAN mixing ratios. The measured mixing ratios of NO<sub>2</sub> were 8–25 % of the initial PAN mixing ratio.

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### Standard BLC and laboratory NO<sub>x</sub> analyser in switching mode

Figure 3 shows the NO<sub>2</sub> signal resulting from the same three BLC units operated in switching mode (40 % duty cycle). The percentage PAN conversion observed is lower in all cases than in the corresponding constant mode. This is likely due to the lower lamp temperature as a result of operating only 40% of the time. The relationship between conversion efficiency and signal are not as clearly evident here as for constant mode operation (Fig. 2). It is possible that the greater variation in the measurement due to the lower amounts of NO<sub>2</sub> produced obscured any trend, however it is clear that there is still a significant proportion of PAN measured as NO<sub>2</sub>; an average of 5.8 %.

#### Inlet residence time effects 3.3

It is not clear from either Fig. 2 or 3 whether the PAN decomposition occurs within the BLC exclusively or within the inlet of the system, as has been claimed previously (Fehsenfeld et al., 1987). Previous studies (Fehsenfeld et al., 1990; Ridley et al., 1988; Sadanaga et al., 2010) have also reported a small PAN interference with photolytic converters, while some found the contribution to the NO<sub>2</sub> signal from PAN to be negligible (Ryerson et al., 2000). Others (Val Martin et al., 2008) acknowledge the possibility for an interference and estimate a small (2 to 4 pptv) positive bias. The photolytic converter designs in the aforementioned studies vary greatly in their implementation and do not have the same ubiquity as BLCs used here, i.e. within the GAW network (Penkett et al., 2011).

Table 2 demonstrates that the residence time of PAN within the inlet does not affect the signal arising from PAN decomposition in our system. This rules out any significant contribution from the inlet to PAN decomposition. The inlet in this case consists of ca.

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2.7 m 1/4 inch PFA tubing shielded from light and held at 20 °C. In other applications, for example if the inlet is heated, contaminated, or has a very long residence time, it is quite possible that significant PAN decomposition occurs.

From these experiments it is evident that a significant  $NO_2$  signal is observed when sampling PAN diluted in zero air. The signal seen corresponds to around 5 to 25% of the PAN supplied which represents a significant interference. The possibility of thermal decomposition within the inlet was ruled out.

#### 3.4 High powered and actively cooled photolytic NO<sub>2</sub> converter

Figure 4 shows the difference in  $NO_2$  signal between the cooled and uncooled high powered BLC described in Sect. 2.2.2. In all of the cooled cases the  $NO_2$  measured was significantly lower than in the uncooled case; this accounts for any increased artefact (the signal recorded when sampling zero air) in the uncooled case. The conversion efficiency was 93 % for  $NO_2 \rightarrow NO$ .

The effect of actively cooling the BLC lamps is significant as apparent in the much lower NO<sub>2</sub> concentrations measured whilst sampling a range of PAN mixing ratios (Fig. 4). It is therefore plainly evident that there is a significant effect of cooling the UV-LEDs which acts to mitigate any signal arising from PAN.

#### 3.5 Possible photolytic interferences of BLCs

Spectral radiograms of the UV-LED elements of standard BLCs were obtained using an Ocean Optics QE65000 spectral radiometer coupled to a  $2\pi$  quartz collector. The spectrometer and collector optics were calibrated using an NIST traceable light source (OL FEL-A, Gooch and Housego) and ultra-linear power supply (OL 83A, Gooch and Housego). The light source is a 1000 W quartz-halogen tungsten coiled-coil filament lamp with spectral irradiance standard F-1128. The lamp was operated at 8 Amps DC (125 V), with the lamp-collector distance fixed at 50 cm. Calibration was carried out in

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Figure 5 shows the spectral emission of six different BLC UV-LED units. These units ranged in age from new to nearing the end of their service life i.e. the conversion efficiency of the whole BLC unit had fallen below acceptable limits. As the LED units age, the relative intensity of their outputs declines, this decrease in intensity can be due to dimming of the overall output, or failure of individual array elements determined by visual inspection during operation. It should be pointed out however, that the light intensity of the UV-LEDs is not directly proportional to the NO<sub>2</sub> conversion efficiency of the complete whole BLC. Rather, the conversion efficiency is strongly dictated by the condition of the reflective Teflon-like cavity. For example, disabling one of the two lamps in a BLC does not reduce the conversion efficiency by half, but by a much smaller percentage. Additionally, replacing the UV-LED elements of a converter whose conversion efficiency has dropped below 30 % with new lamps will not lead to a recovery of the conversion. Scrupulous cleaning of the reflective cavity with solvent and mild abrasion of the surface will however recover the conversion efficiency considerably.

Figure 6 depicts the absorption cross sections of atmospheric nitrogen compounds against the measured spectral output of UG5 UV-passing filter glass (Schott, 1997) used in lamp-type PCL optics e.g. Eco Physics PCL 762, and the averaged measured spectral output of six individual BLC UV-LED arrays of varying running hours. Also shown is the NO<sub>2</sub> quantum yield (Gardner, 1987; Koepke et al., 2010). It can be discerned that the UV-LED output overlaps fully with the NO<sub>2</sub> absorption band and the NO<sub>2</sub> quantum yield and is therefore photon efficient. It is also shown that there in minimal overlap with HONO and no overlap in the spectrum at all with PAN. It has been shown (Carbajo and Orr-Ewing, 2010; Talukdar et al., 1997) that there is no overlap either with methyl, ethyl or isopropyl nitrate – methyl nitrate being a minor impurity in PAN synthesis. There is only very minor overlap in the PCL optics spectrum with PAN, methyl ethyl, and isopropyl nitrate. There is a great deal of overlap with HONO which

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#### Possible thermal interferences in BLCs

The thermal and electronic characteristics of the standard BLC lamps were ascertained in bench tests and are summarized in Table 3. Each lamp was run constantly on the bench whilst recording the surface temperature and power draw of the light emitting element. The surface temperature was recorded once a stable maximum had been reached and maintained for at least 10 min - representative of using a BLC in constant mode. The ambient temperature during the experiments was 20°C.

Table 3 details the power draw and surface temperature of three BLC UV-LED lamp pairs measured during tests, along with their NO<sub>2</sub> → NO convertor efficiencies when assembled as a complete BLC. The surface temperature of the individual lamps correlate positively with the power drawn by each lamp ( $R^2 = 0.96$ ) and indeed with output intensity (Fig. 2), but with each lamp pair there is only weak correlation ( $R^2 = 0.43$ ) between converter efficiency and temperature. It is worth noting that the power consumption is a combination of the light output, heat dissipation, and power to the cooling fan. It is clear however that the temperature experienced by the sample gas within the NO<sub>2</sub> converter is significantly above ambient. In fact, the entire NO<sub>2</sub> conversion cavity is heated by the lamps leading to external temperatures of the converter of between 34 and 45°C.

It is known that the major product from thermal decomposition of PAN is NO<sub>2</sub>; Reaction (R18) (Roumelis and Glavas, 1992; Tuazon et al., 1991). The NO<sub>2</sub> produced thermally within the converter may then be photolysed to NO and thus be measured as NO<sub>x</sub> and attributed to atmospheric NO<sub>2</sub>.

$$CH_3C(O)OONO_2 \rightleftharpoons CH_3C(O)OO^{\bullet} + NO_2$$
 (R18)

A model of the thermal decomposition of PAN over a range of temperatures within the BLC with a residence time of 0.96 s is shown in Fig. 7. The model run was conducted 28719

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in Facsimile using rate constants from IUPAC Evaluated Kinetic Data (Atkinson et al., 2006).

The model output indicates that measureable PAN decomposition to NO<sub>2</sub> occurs above ca. 50°C. At the maximum LED surface temperature recorded (80°C) the 5 model predicts ca. 30 % decomposition of PAN to NO<sub>2</sub>. However, as only the two UV-LED lamps are at such an elevated temperature we can expect a temperature gradient/heating rate within the BLC so that the average temperature seen by the sample gas over the 0.96 s residence is somewhat lower. Also, it is expected that in a switching mode with only 40 % duty cycle of the lamps, their surface temperature would be lower also. This is borne out when using the external surface temperature of the BLC as a proxy – the temperature was lower in switching mode than in constant mode for the same conversion efficiency. It is shown more clearly in the inset that an average temperature of 60°C would cause a 4.6 % decomposition of PAN and account for the NO<sub>2</sub> measured during experiments with the standard BLCs. Therefore, together with the spectral measurements reported in Sect. 3.5, it seems highly unlikely that the source of the artefact signal is through direct photolysis of PAN, leaving thermal decomposition modelled in Fig. 7 the remaining explanation.

In Sect. 3.1 the percentage conversion of PAN to NO<sub>2</sub> was found to be, on average, highest at the lowest converter efficiency and vice versa. The fact that the convertor temperatures are very similar at different convertor efficiencies (Table 3) suggests that the percentage of PAN thermally dissociated in each case is similar. Explanation for the inverse relationship between percentage conversion of PAN to NO and conversion efficiency (Fig. 2) lies in the way that the NO<sub>2</sub> concentration is derived, which is an inverse function of the assumed conversion efficiency as in Eq. (2) where converter efficiency is expressed fractionally. If in fact the conversion efficiency of PAN to NO in the convertor was not related to the measured NO<sub>2</sub> to NO conversion efficiency but instead a constant value, then the apparent relationship between CE and % conversion would disappear. This explanation is consistent with the fact that when the average conversion percentage in Fig. 2 is normalized to conversion efficiency, the percentage

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for the three BLCs is remarkably similar (Table 4) with the average PAN decomposition of 4.6 % needed to produce the spurious signal observed.

Above a threshold temperature of  $25\,^{\circ}$ C, the  $NO_2$  formed in Reaction (R18) may be excited through the dissipating of internal energy from the parent PAN molecule (Mazely et al., 1995), this  $NO_2^*$  is then more readily photolysed to NO within the BLC than ground state  $NO_2$ . The discussion above suggests that a similar proportion of the  $NO_2$  evolved from thermal dissociation of PAN is converted to NO within the BLC leading to the apparent inverse correlation between conversion efficiency and PAN "artefact". Consequently, the lower  $NO_2 \rightarrow NO$  conversion efficiency of a BLC, the greater the positive error in  $NO_2$  when PAN is present.

### 4 Atmospheric implications

We have shown that a significant proportion of PAN can be decomposed under the normal operating conditions of a BLC equipped NO chemiluminescence instrument leading to a spurious increase in measured NO $_2$  of 5 to 20 % of the PAN supplied. The UV-LED light source employed by was found to reach a temperature of 56 to 80 °C in normal operation with the surface temperature correlating positively with power draw and output intensity.

The positive bias in  $NO_2$  measurements by NO chemiluminescence using BLCs has implications for data in both remote background sites and polluted areas. Figure 8 shows the thermal decomposition profiles of many common  $NO_y$  species. Whereas only a small fraction of PAN is found to convert to  $NO_2$  at the operating temperatures ( $\sim 5\,\%$  at 60 °C) of the instrument, a number of more thermally labile compounds exist which become more important at higher latitudes and altitudes.

The degree of thermal decomposition within the instrument will depend upon the thermal profile of the air (T(t)) as it passes through the instrument. This can be parameterized as a relaxation timescale  $(\tau s)$  for temperature from the ambient temperature

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$$T(t) = T_0 + (T_{\text{BLC}} - T_0) \exp\left(-\frac{t}{\tau}\right) \tag{4}$$

The rate at which thermal equilibrium is reached within the cell,  $\tau$ , is calculated as 0.42 s from the observed PAN decomposition of 4.6% (from which T(t) is derived), at 20°C ambient temperature, a BLC temperature of 75°C and a 1 s residence time. This allows a calculation of the potential interference from other thermally labile NO $_y$  compounds.

Given a 1st order loss of the PAN like compound and a temperature profile within in the instrument as described in 4 we have 5:

$$\frac{d[PAN]}{dt} = -k(T)[PAN] = -k(t)[PAN]$$
 (5)

Given the laboratory observations of the temperature dependence of the rate constant (typically  $k(T) = A \exp(-B/T)$ ), and the parameterized temperature within the instrument (T(t)), the fraction of the compounds that will have decomposed can be found by numerical integration.

Figure 9 uses output from the GEOS-Chem model (version 9.3, http://www.geos-chem.org, Bey et al., 2001) run at  $2^{\circ} \times 2.5^{\circ}$  resolution, plus updates described in Sherwen et al. (2015), to provide an estimate of the interference on NO<sub>2</sub> from the decomposition of NO<sub>y</sub> species within a BLC photolytic converter. The species used for this analysis are: PAN; MPAN; PPN; IONO<sub>2</sub>; BrONO<sub>2</sub>; CIONO<sub>2</sub>; N<sub>2</sub>O<sub>5</sub>; CH<sub>3</sub>O<sub>2</sub>NO<sub>2</sub>, HO<sub>2</sub>NO<sub>2</sub>. Thermal decomposition information are taken from IUPAC evaluated kinetic data (Atkinson et al., 2003, 2006, 2007). Interferences are calculated for each month of a one year simulation and the maximum value shown. The estimate assumes a BLC conversion efficiency of 100 % NO<sub>2</sub>  $\rightarrow$  NO and thus does not include the extra signal from the photolysis of NO<sub>2</sub>\*  $\rightarrow$  NO with converters where conversion is less than unity  $\rightarrow$  in this case a multiplying factor exists.

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Figure 9 shows that in extreme circumstances NO<sub>2</sub> may be over-reported by many hundreds of percent. These are in regions that typically have low NO<sub>x</sub> concentrations and are cold (polar), and in the upper troposphere. Here, compounds such as PAN and HO<sub>2</sub>NO<sub>2</sub> are in high concentrations compared to NO<sub>2</sub> and it is not surprising that thermal decomposition can have an impact. Upper tropospheric over estimates of NO<sub>2</sub> concentrations could be as high as 150 pptv.

The NO<sub>2</sub> bias shown in Fig. 9 impacts the modelled Leighton ratio. In Fig. 10 the model is sampled every daylight hour for every surface grid box for the month of March. The calculation shown in red is the Leighton ratio calculated from the modelled concentrations of NO, NO<sub>2</sub>, jNO<sub>2</sub>, O<sub>3</sub>, T, HO<sub>2</sub>, RO<sub>2</sub>, BrO and IO. The model value is in general close to 1. In the blue the same calculation is performed but including the interferences on the NO<sub>2</sub> channel from the instrumental decomposition of a range of BLC lamp temperatures between 60 and 105  $^{\circ}$ C (described by  $\tau$  of 0.42 s and a residency time of 1 s). Here there are significant interferences.

As shown in Fig. 10 the Leighton ratio can be extremely perturbed from what would be predicted by all available measurements given the NO<sub>2</sub> bias we have shown exists with photolytic NO<sub>2</sub> converters which operate above ambient temperature. This is especially true in low NO, environments. This has led to either hypothesizing (a) an unknown, unmeasured, selective oxidant "compound X" or, (b) theoretical mechanisms by which NO is converted to NO2 (Bauguitte et al., 2012; Cantrell et al., 2003; Frey et al., 2013, 2015; Hosaynali Beygi et al., 2011).

As a practical example with an analogous system to that used in this study, the NO<sub>x</sub> observations of Hosaynali Beygi et al. (2011) taken in the South Atlantic during the "OOMPH" project (Ocean Organics Modifying Particles in both Hemispheres) serve as a case study. During this cruise a BLC photolytic NO2 converter (Droplet Measurement Technologies, Colorado, USA), with a residence time of 1 s, as described in this study - coupled to an NO chemiluminescence analyser was deployed aboard the French research vessel Marion Dufresne II. The cruise track covered the South

The results presented by Hosaynali Beygi et al. (2011) show an  $NO_2$ : NO ratio which is a factor c.a. 7 greater than can be explained given the available supporting observations of  $O_3$ , CO, OH,  $HO_2$ ,  $RO_X$  ( $RO + RO_2 + HO_X$ ), i.e. measured  $NO_2$  is much greater than the Leighton relationship would predict. This apparent Leighton ratio is attributed by the author to an "unknown oxidant" which selectively converts NO to  $NO_2$ .

The authors estimate a minimal photolytic interference from atmospheric nitrates using their BLC, consistent with our analysis (Sect. 3.5 and Fig. 6). Only PAN decomposition within the inlet line is considered as a potential interference, but is estimated to be minimal (< 7 ppt) due to the low ambient temperature range of 3 to 11 °C. Our results have shown that thermal, rather than photolytic decomposition of atmospheric NO $_y$  species with the converter itself is the major source of over-reporting in NO $_2$  measurements made with BLCs. Indeed, our model study predicts interference of 50 to 100 % in the region of the cruise, assuming an NO $_2$  converter efficiency of 100 %, whereas in the OOMPH observations the efficiency was 59 % thus, a multiplying factor may apply to any PAN interference of  $\sim$  1.7.

Presented in Fig. 4 of Hosaynali Beygis' paper on page 8503 (Hosaynali Beygi et al., 2011) is the Leighton ratio calculated from their  $NO_2$  data and ancilarry measurments. The ratio in the pristine background appears to be significantly biased towards  $NO_2$ , exceeding what the authors are able to explain given their measurments of  $HO_x$ ,  $RO_x$  and OH. Their values fit well within the modelled Leighton ratios shown in Fig. 10 when an instrumental bias is included however. This is suggestive of an instrumental  $NO_2$  bias cause by a BLC with a lamp temperature of around 90 °C, however does not include any multiplying effect of having an  $NO_2$  conversion efficiency less than unity. As the efficiency is 59 % the actual temperature would not necessarily be so high as 90 °C. Eliminating this  $NO_2$  over-reporting closes the oxidative balance implied by their other data and removes the need to invoke any compound X or mystery oxidant.

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Measurements of NO<sub>2</sub> collected using photolytic converters and chemiluminescence systems may be significantly biased in low NO<sub>x</sub> environments. Thermal decomposition of NO<sub>v</sub> species within the NO<sub>2</sub> converter produces spuriously high readings; this is especially true in pristine environments and at high elevations where the NO<sub>v</sub> to NO<sub>x</sub> ratio may be high. Over-reporting of NO2 has been shown to lead to apparent gaps in oxidation chemistry which cannot be explained with any available measurements. This has led to theorization of an unknown "compound X" which selectively oxidizes NO to NO<sub>2</sub>, however this is likely anomalous and simply due to error in the NO<sub>2</sub> determination. In order to mitigate this overestimation of the NO<sub>2</sub> mixing ratio by the dissociation of PAN and other compounds it is desirable to have the highest possible  $NO_2 \rightarrow NO$ conversion efficiency, i.e. unity, to mitigate the multiplying effect of having lower conversion efficiencies. High conversion efficiency is now achievable thanks to advancements in the power density of UV-LEDs currently available; new generation BLCs have a CE > 95 % for example. Additionally, actively cooling the UV emitting elements or separating them from the gas stream is essential in order that the sample gas should not contact any surface which is above ambient temperature. Failure to achieve this

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may result in spuriously high NO<sub>2</sub> observations.

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**Table 1.** Comparison of the NO<sub>2</sub> signal observed from measurement of three zero air sources. (Eco Physics AG PAG 003, BOC Specialty Gasses BTCA 178, and Sofnofil/Carbon/13× molecular sieve filters) by LIF. The average of ten 1 min averages is shown in raw PMT counts per second as well as the standard deviation of those averages.

	PAG 003	BTCA 178	Sofnofil/Carbon
Signal (cps)	88.10	88.66	83.95
Standard Deviation	1.64	1.56	1.81

**Table 2.** Effect of varying the residence time of PAN (1.0 ppb) within a 2.7 m PFA inlet on measured NO<sub>2</sub> concentrations.

	Inlet residence time (s <sup>-1</sup> )			
	0.84	1.05	1.40	2.10
NO <sub>2</sub> (ppt) NO <sub>2</sub> (%)	60.2 5.4	61.3 5.4	60.6 5.2	61.5 5.3

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**Table 3.** Peak surface temperature and current drawn by BLC lamps in a bench test at 20.0 °C showing converter efficiency, current, and surface temperature.

Lamp No.	Converter efficiency (%) ±1	BLC Lamp Surface Temperature (°C) ±0.05	Current Draw (A) ±0.0005
1 2	41	79.8 75.3	0.969 0.953
3 4	35	77.6 74.0	0.933 0.931
5 6	22	76.2 56.4	0.916 0.567

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Table 4. The average percentage conversion of PAN to NO<sub>2</sub> measured, and normalized to the converter efficiency of each BLC.

	Converter efficiency (%) ±1		
	42	33	22
Measured % Normalized %	10.8 4.4	15.9 5.2	19.6 4.3



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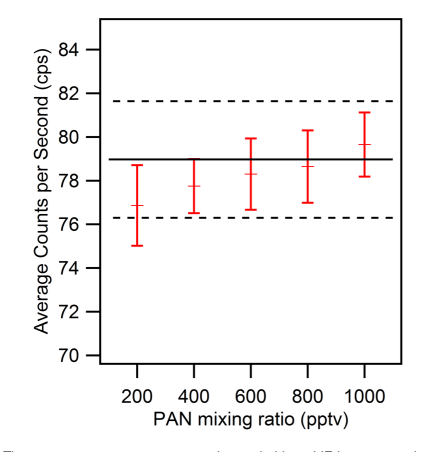


Figure 1. The average raw counts per second recorded by a LIF instrument when sampling various mixing ratios of PAN (in red) and zero air (black). The variance of the zero air signal is also shown (dashed black). The average signal while sampling PAN falls within the noise of the zero signal.



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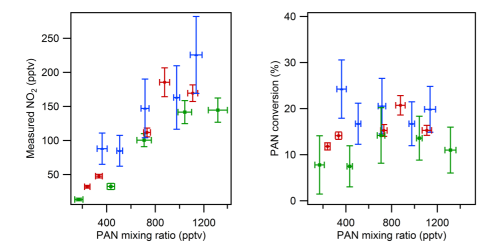


Figure 2. The measured absolute NO<sub>2</sub> signal (left panel) of the supplied PAN mixing ratio, and as a percentage (right panel), for three BLC units operating in constant mode. Green = 41 % CE, Red = 35 % CE, Blue = 22 % CE.



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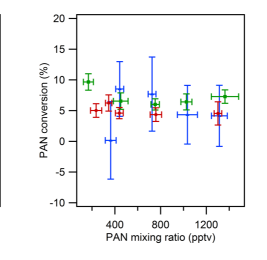


Figure 3. The measured absolute NO<sub>2</sub> signal as a function of the supplied PAN mixing ratio (left panel), and as a percentage (right panel), for three BLC units operating in switching mode. Green = 41 % CE, Red = 35 % CE, Blue = 22 % CE.

100

80

60

20

400

800

PAN mixing ratio (pptv)

1200

Measured NO<sub>2</sub> (pptv)



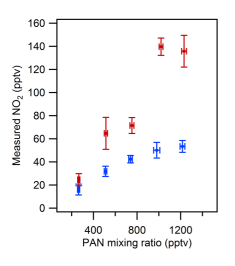
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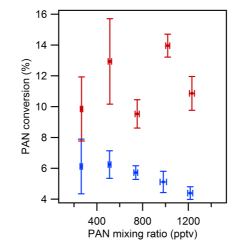
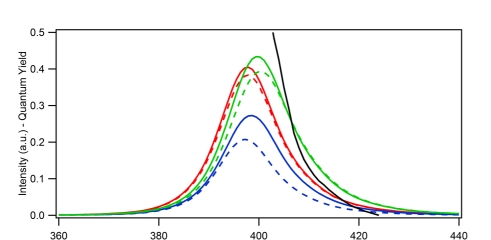


Figure 4. The measured NO<sub>2</sub> signal as a function of the supplied PAN mixing ratio (left panel), and as a percentage (right panel), for the cooled (blue) and uncooled (red) high powered BLC.



**Figure 5.** Shown is the spectral output vs. wavelength of two new, previously unused BLC lamps No. 1 (solid) and 2 (dashed) in green, two used lamps No. 3 (solid) and 4 (dashed) in red; still within acceptable conversion efficiency, and two which fall below acceptable limits No. 5 (solid) and 6 (dashed) in blue. The  $NO_2$  quantum yield is shown in black.

Wavelength (nm)

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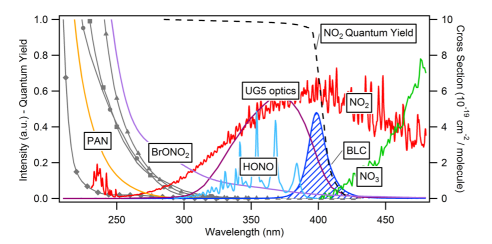
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**Figure 6.** Absorption cross section (red) and quantum yield (dashed black) of  $NO_2$ , presented with the spectral output of UG5 optical filtering (purple) and an average of the six BLC lamps used in this study output (dark blue). Shown are interfering species;  $NO_3$  radicals (green), HONO (light blue), BrONO<sub>2</sub> (lilac) which are overlapped significantly by the UG5 optics – completely in the case of HONO, whilst much less overlap is exhibited by the UV-LEDs of the BLC. Also shown is PAN (gold), which clearly is not overlapped by either UG5 or BLC light sources. Additional non-interfering species;  $CIONO_2$  (triangles),  $N_2O_5$  (squares),  $CIONO_2$  (circles),  $CIONO_3$  (diamonds) shown for reference.

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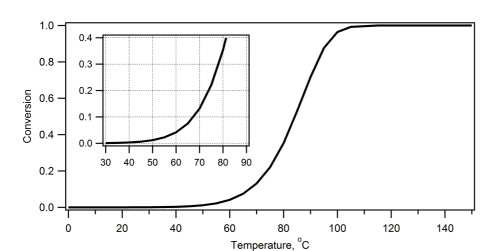
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**Figure 7.** Model of thermal decomposition of PAN to NO<sub>2</sub> with a residence time of 0.96 s at temperatures between 0 to 150 °C. Inset is detail of 30 to 90 °C.

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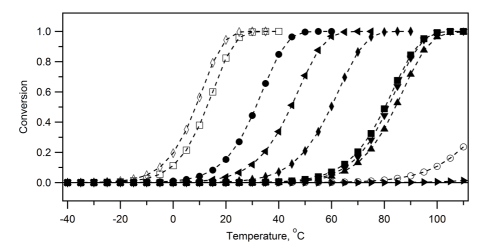
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**Figure 8.** Thermal decomposition profiles of IONO₂ ♦; BrONO₂ ○; CIONO₂ ▶; HO₂NO₂ •;  $N_2O_5$  ◀;  $C_2H_5O_2NO_2$  ⋄;  $CH_3C(O)CH_2O_2NO_2$  □; MPAN ■; PAN ▲; PPN ▼. Calculated from IUPAC recommended kinetic data using FACSIMILE software based on 1 s residence time. Note  $CH_3O_2NO_2$  is not shown but has the same profile as  $HO_2NO_2$ .

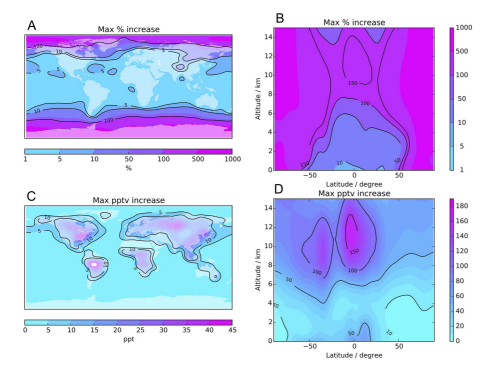
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**Figure 9.** GEOS-Chem model showing the monthly maximum percentage over-reporting of NO<sub>2</sub> (a) zonally and (b) by altitude in any month of a 1 year simulation. Panels (c) and (d) show the same in absolute pptv values. Note; the area over the Amazon in plot (c) is over the maximum range of the colour scale. Surface values are the maximum over-reporting in any month, zonal values are the maximum over reporting in any month and in any of the longitudinal grid boxes.

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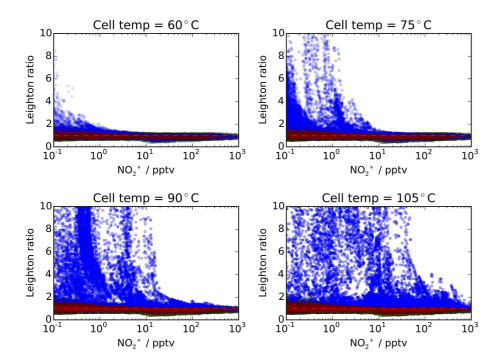


Figure 10. Leighton ratio calculated for each model grid-box for each daylight hour for March by the model as a function of the grid box NO<sub>2</sub> concentration. The instrument interference is characterized by a numerical solution of Eq. (5) with  $\tau = 0.42 \, \mathrm{s}$  and a residence time of 1 s. Red shows the values calculated without the interference on the NO<sub>2</sub> concentration and the blue indicated the values calculated with the interference. The interferences are calculated for different lamp temperatures, 65 to 105 °C.

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