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## Interference from alkenes in chemiluminescent $NO_x$ measurements

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**Abstract.** Nitrogen oxides  $(NO_x=NO+NO_2)$  are critical intermediates in atmospheric chemistry and air pollution.  $NO_x$  levels control the cycling and hence abundance of the primary atmospheric oxidants OH and NO3 and regulate the ozone production which results from the degradation of volatile organic compounds (VOCs) in the presence of sunlight. They are also atmospheric pollutants, and NO2 is commonly included in air quality objectives and regulations.  $NO_x$  levels also affect the production of the nitrate component of secondary aerosol particles and other pollutants, such as the lachrymator peroxyacetyl nitrate (PAN). The accurate measurement of NO and NO2 is therefore crucial for air quality monitoring and understanding atmospheric composition. The most commonly used approach for the measurement of NO is the chemiluminescent detection of electronically excited  $NO_2$  ( $NO_2^*$ ) formed from the  $NO + O_3$  reaction within the  $NO_x$  analyser. Alkenes, ubiquitous in the atmosphere from biogenic and anthropogenic sources, also react with ozone to produce chemiluminescence and thus may contribute to the measured  $NO_x$  signal. Their ozonolysis reaction may also be sufficiently rapid that their abundance in conventional instrument background cycles, which also utilises the reaction with ozone, differs from that in the measurement cycle such that the background subtraction is incomplete, and an interference effect results. This interference has been noted previously, and indeed, the effect has been used to measure both alkenes and ozone in the atmosphere. Here we report the results of a systematic investigation of the response of a selection of commercial NO<sub>x</sub> monitors to a series of alkenes. These  $NO_x$  monitors range from systems used for routine air quality monitoring to atmospheric research instrumentation. The species-investigated range was from shortchain alkenes, such as ethene, to the biogenic monoterpenes. Experiments were performed in the European PHOtoREactor (EUPHORE) to ensure common calibration and samples for the monitors and to unequivocally confirm the alkene levels present (via Fourier transform infrared spectroscopy -FTIR). The instrument interference responses ranged from negligible levels up to 11%, depending upon the alkene present and conditions used (e.g. the presence of co-reactants and differing humidity). Such interferences may be of substantial importance for the interpretation of ambient  $NO_x$ data, particularly for high VOC, low NO<sub>x</sub> environments such as forests or indoor environments where alkene abundance from personal care and cleaning products may be significant.

#### 1 Introduction

Measurement of atmospheric trace constituents is central to atmospheric chemistry research and air pollution monitoring. Key challenges to trace measurements are sensitivity, reactivity and selectivity as many components of interest are only present in parts per billion (ppb;  $10^{-9}$ ) or parts per trillion

(ppt; 10<sup>-12</sup>) mixing ratios; in many cases, their inherent reactivity necessitates in situ detection. Atmospheric trace composition comprises many thousands of different chemical components (Goldstein and Galbally, 2007). Consequently, specific measurement approaches have been developed to measure key atmospheric species within the specific conditions (analyte abundance; presence of other constituents) anticipated (Heard, 2008). This paper reports a systematic study of the interference arising in measurements of nitrogen oxides from the presence of alkenes in sampled air when using their most widespread air quality monitoring technique of chemiluminescence detection.

 $NO_x$  (=  $NO + NO_2$ ) abundance controls the cycling and, hence, the concentration of the primary atmospheric oxidants, hydroxyl (OH) and nitrate (NO<sub>3</sub>) radicals and regulates the ozone production which results from the degradation of volatile organic compounds (VOCs) in sunlight.  $NO_x$ are also atmospheric pollutants in their own right, and NO<sub>2</sub> is commonly included in air quality objectives and regulations (as the more harmful component of  $NO_x$ ; European Environment Agency, 2018; Chaloulakou et al., 2008). In addition to their role in controlling ozone formation,  $NO_x$ levels affect the production of other pollutants, such as the lachrymator peroxyacetyl nitrate (PAN) and the nitrate component of secondary aerosol particles. Consequently, accurate measurement of nitrogen oxides in the atmosphere is of major importance for monitoring pollution levels and assessing consequent health impacts and understanding atmospheric chemical processing. Atmospheric NO and NO2 are formed from natural processes (lightning, soil emissions of NO, biomass burning and even snowpack emissions) and anthropogenic activities (high temperature combustion in air leading to the breakdown of  $N_2$  and  $O_2$ ;  $NO_x$  production via the Zeldovich mechanism), where road traffic is the predominant source in many urban areas (Keuken et al., 2009; Grice et al., 2009; Carslaw and Rhys-Tyler, 2013). Consequently, boundary layer  $NO_x$  abundance varies over many orders of magnitude, from sub-5 ppt levels in the remote marine boundary layer to parts per million (ppm) levels in some urban environments (Crawford et al., 1997).

Techniques used for the measurement of atmospheric  $NO_x$  include laser-induced fluorescence (LIF) spectroscopy for both NO and  $NO_2$ ; absorption spectroscopy (e.g. long path differential optical absorption spectroscopy – LP-DOAS; cavity-enhanced differential optical absorption spectroscopy – CE-DOAS; cavity attenuated phase shift spectroscopy – CAPS; cavity ring-down spectroscopy – CRDS; passive diffusion tubes, primarily for  $NO_2$ ), chemical ionisation mass spectrometry (CIMS) and both on- and offline wet chemical methods, for example, a long path absorption photometer (LOPAP; Heard, 2008; Sandholm et al., 1990; Kasyutich et al., 2003; Kebabian et al., 2005; Cape, 2009; Fuchs et al., 2009; Thalman and Volkamer, 2010; Villena et al., 2011). However, the most commonly employed technique for the measurement of  $NO_x$  species, including for statu-

tory air quality monitoring purposes, is the detection of the chemiluminescence arising from electronically excited  $NO_2$  ( $NO_2^*$ ) formed from the reaction between NO and  $O_3$  (via reaction R1) as follows:

$$NO + O_3 \rightarrow NO_2^* + O_2 \tag{R1}$$

$$NO_2^* \rightarrow NO_2 + h\nu$$
. (R2)

The intensity of the light emitted via reaction (R2) is in the wavelength range  $600-3000 \,\mathrm{nm}$ , peaking at  $\sim 1200 \,\mathrm{nm}$ . Chemiluminescent instruments mix sampled ambient air with a reagent stream containing an excess of ozone to promote the chemiluminescent reaction (see the schematic in Fig. 1); the resulting emission signal is measured using a photomultiplier tube (PMT) and consists of contributions from NO<sub>2</sub>\* formed, as above, but also potentially from other chemiluminescence processes, detector dark counts and other noise contributions. Contributions to the measured emission from other species are minimised by using a red filter on the detector to block emission wavelengths below ca. 600 nm and by employing a background subtraction cycle; chemiluminescent NO<sub>x</sub> monitors commonly acquire a background by increasing the reaction time between NO (from the sampled air) and O<sub>3</sub> (reagent formed within the instrument), using a pre-reactor volume, such that nearly all of the NO present (specifications typically state levels in excess of 99 %) is converted to NO<sub>2</sub>. The difference in PMT signals between the online and background signals is then taken to be proportional to the NO present in the air sample, following the assumption that the abundance of other species, which may contribute to the measured signal, is not affected by the background cycle.

Chemiluminescent instruments typically alternate between two operation modes, namely one that directly measures NO and one that measures  $\Sigma(NO + NO_2)$  by first converting NO<sub>2</sub> to NO. The difference between the two values determines the NO<sub>2</sub> mixing ratio (if only NO and NO<sub>2</sub> are present). This is most commonly achieved by using a molybdenum (Mo) catalyst heated to 300-350 °C. However, the reduction in other NO<sub>7</sub> species to NO has led to the use of these catalysts in chemiluminescent NO<sub>v</sub> monitors to measure total reactive nitrogen rather than  $NO_2$  ( $NO_v = NO_z + NO_x$ ; i.e. NO<sub>7</sub> equals other reactive nitrogen species catalysed by Mo convertors, e.g. HNO<sub>3</sub>, nitrous acid (HONO), N<sub>2</sub>O<sub>5</sub>, HO<sub>2</sub>NO<sub>2</sub>, PAN, NO<sub>3</sub> and organic nitrates but not NH<sub>3</sub>; Navas et al., 1997; Murphy et al., 2007). If atmospheric mixing ratios of NO<sub>7</sub> species are high relative to NO<sub>2</sub>, then NO<sub>2</sub> measurements with monitors equipped with Mo catalysts are increasingly inaccurate. This has led to the adoption of photolytic NO<sub>2</sub> conversion stages in research instruments, where a blue light LED convertor is illuminated in a photolysis cell to convert NO<sub>2</sub> to NO (Lee et al., 2015).

$$NO_2 + h\nu \ (< 420 \,\text{nm}) \to NO + O(^3P).$$
 (R3)

The photolytic conversion technique can have greater specificity than the heated Mo catalyst, as the photolysis wave-

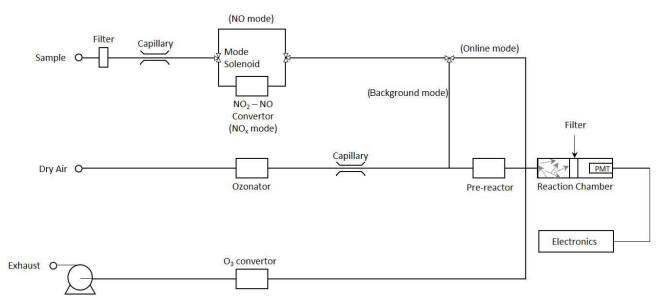


Figure 1. A typical flow schematic of a chemiluminescent NO monitor.

lengths may be selected to match the  $NO_2$  photolysis action spectrum, while potential  $NO_z$  interferents for an  $NO_2$  measurement are thermally unstable and may convert to  $NO_2$  when exposed to heat in the latter approach (Heard, 2008). Despite this, the chemiluminescent analyser with the heated Mo catalyst is the most widely used technique for air quality monitoring of NO and  $NO_2$  worldwide. It is the reference method of measurement specified in the EU directive (BS EN 14211, 2012), providing real-time data with a short time resolution for 212 monitoring sites in the UK, including kerbside, roadside, urban background, industrial and rural locations (AQEG, 2004).

# 2 Origins of interferences in chemiluminescent NO<sub>x</sub> measurements

While  $NO_x$  measurements are sometimes perceived to be straightforward and routine, in practice a number of factors are known to affect the accuracy of the levels obtained using chemiluminescence approaches. A detailed account of the factors affecting atmospheric  $NO_x$  measurement overall is given elsewhere (e.g. Gerboles et al., 2003; Villena et al., 2012; Reed et al., 2016); here we do not focus upon surface sources and losses but rather upon chemical interferences in chemiluminescent  $NO_x$  analysers, which may arise from the following possible general mechanisms:

 Collisional quenching of NO<sub>2</sub>\* by an interferent species with a greater collisional efficiency than the bath gas (e.g. air) used for calibration (this is typically a negative interference, although the magnitude and sign of this depends upon the calibration conditions employed).

- 2. Conversion of other nitrogen-containing species to  $NO_x$  within the  $NO_2$  conversion unit (a positive interference).
- 3. Chemical removal or interconversion of NO and/or NO<sub>2</sub> by an interferent species generated within the instrument (a positive or negative interference).
- 4. Chemiluminescence of other chemical species, which is not fully accounted for during the instrument background cycle (a positive interference).

Collisional quenching of excited species, mechanism (1), results in a reduction in the chemiluminescence intensity, which is to an extent dependent upon the pressure and quenching efficiency, i.e. the efficacy with which the quenching species may accept or remove energy from the excited moiety. In the case of electronically excited NO<sub>2</sub>, effective quenching agents have been shown to include H<sub>2</sub>O, CO<sub>2</sub>, H<sub>2</sub> and hydrocarbons (Matthews et al., 1977; Gerboles et al., 2003; Dillon and Crowley, 2018), of which only quenching by water vapour is considered to be significant under most common (ambient air) conditions, and sensitivity reductions of up to 8% have been reported (Steinbacher et al., 2007). Mechanism (2), the conversion of other nitrogen-containing species to NO, alongside NO<sub>2</sub>, is a recognised issue with heated Mo converters; interferences between 18 % and 100 % have been reported for species such as HONO, HNO<sub>3</sub>, PAN, alkyl nitrates and N<sub>2</sub>O<sub>5</sub> (Dunlea et al., 2007; Lamsal et al., 2008). To address these uncertainties, photolytic converters are now commonly employed in research measurements, although, for most routine air quality monitoring, heated Mo converters are still employed. Recently, it has been shown that a further interference can arise within the photolytic converter stage from the generation of  $HO_x$  radicals through the photolysis of photolabile carbonyl species, such as glyoxal, forming peroxy radicals promoting NO to NO<sub>2</sub> conversion within the instrument (Villena et al., 2012), resulting in a negative NO<sub>2</sub> interference, which may (under some conditions) exceed the positive interference from retrieval of NO<sub>z</sub> species associated with heated Mo converter instruments, i.e. mechanism (3).

The focus of this work relates to mechanism (4), namely interference in the chemiluminescent measurements of NO and NO<sub>2</sub> (using both catalytic and photolytical converters) arising from the chemiluminescence of alkenes in the presence of ozone. Alkene-ozone reactions have received substantial attention as a dark source of HO<sub>x</sub> radicals and are a route to the formation of semi-volatile compounds which contribute to secondary organic aerosol (SOA), particularly for biogenic alkenes such as isoprene and the mono- and sesquiterpenes (e.g. Johnson and Marston, 2008; Shrivastava et al., 2017). Rate constants for ozonolysis reactions depend on the alkene structure and are typically larger for biogenic alkenes. Chemiluminescence from the ozonolysis of 14 short chain species at total pressures of 2–10 Torr was first reported by Pitts et al. (1972). Excited formaldehyde (HCHO), vibrationally excited OH and electronically excited OH in the wavelengths 350–520, 700–1100 and 306 nm, respectively, were the identified chemiluminescent species (Finlayson et al., 1974) and, indeed, have been used to perform field measurements of both ozone and alkenes (e.g. Velasco et al., 2007; Hills and Zimmerman, 1990). This combination of alkene-ozone reactions giving rise to a chemiluminescent interference signal and alkene-ozone reactions being sufficiently rapid that alkenes can be appreciably consumed in the background (pre-reactor) cycle and, hence, the interference contribution not being fully subtracted during the background correction gives rise to the potential for interference in  $NO_x$  measurements, which is the focus of this study.

## 3 Experimental approach

#### 3.1 Sampling

Experiments were performed using chamber A of the two  $200 \,\mathrm{m}^3$  simulation chambers of the European PHOtoREactor (EUPHORE) facility in Valencia, Spain, to provide a common, homogeneous air volume for multiple  $\mathrm{NO}_x$  analysers to sample from. The EUPHORE chambers are formed from fluorine-ethene-propene (FEP) Teflon foil fitted with housings that exclude ambient light (Wiesen, 2001; Munoz et al., 2011). The chambers are fitted with large horizontal and vertical fans to ensure rapid mixing (timescale of 3 min). Instrumentation used comprises long path Fourier transform infrared spectroscopy (FTIR; for absolute and specific alkene/VOC measurements), monitors for temperature, pressure, humidity (dew point hygrometer), ozone (UV absorption) and CO (infrared absorption).  $\mathrm{NO}_x$  levels were

measured using four independent chemiluminescent monitors plus (in the case of NO<sub>2</sub>) LP-DOAS absorption spectroscopy. All monitor sampling lines were of similar lengths and attached to one inlet sampling from the centre of the chamber.

Monitors 1 and 2 employed heated Mo catalysts, while 3 and 4 used photolytic  $NO_2$  converters (see Table 1). All  $NO_x$  monitors were calibrated (in the range 0–100 ppb) at the start and end of the 2-week measurement period, using a multipoint calibration derived from a primary NO standard (BOC 5 ppm alpha standard; certified to the National Physics Laboratory (NPL) scale) in addition to single-point calibrations performed on a daily basis.  $NO_2$  calibration was achieved via gas-phase titration, using added ozone within the chamber. In some experiments, the calibrations and interference were confirmed with use of the EUPHORE LP-DOAS system to unequivocally identify and quantify  $NO_2$ .

All experiments were performed with the chamber housing closed (i.e. dark conditions;  $j(NO_2) < 2 \times 10^{-6} \text{ s}^{-1}$ ) at near-atmospheric pressure and ambient temperature. For most experiments, humidity was low (dew point ca. -45 °C). The experimental procedure, starting with a clean flushed chamber, was to add SF<sub>6</sub> (as a dilution tracer) followed by successive aliquots of various alkenes and, in certain cases, additional species (H2O and CO), whilst recording the measured NO and NO<sub>2</sub> levels over periods of 1-3 h. For some systems, ozone was added at the end of the experiment; under such dark, high-O<sub>3</sub> conditions, we can be confident that negligible NO could actually be present in the chamber (e.g. from wall sources) and, hence, that any NO signal observed by the monitors was unequivocally an interference response (as any NO remaining would be rapidly consumed by reaction with  $O_3$ ). The potential interferant species investigated were cis-2-butene (C2B), trans-2-butene (T2B), tetra-methyl ethylene (2,3-dimethyl-butene or TME),  $\alpha$ -terpinene, limonene, methyl chavicol (estragole) and terpinolene, with four to five additions of 20-50 ppb in each case, together with single- or dual-point interference measurements for ethene, propene, isobutene, isoprene,  $\alpha$ -pinene,  $\beta$ -pinene and myrcene. Repeat experiments were performed for T2B, terpinolene and  $\alpha$ -terpinene under conditions of increased humidity (up to ca. 30 % relative humidity – RH). Alkene mixing ratios introduced into the chamber are given in Table S1. Propene, C2B and T2B were supplied by Linde plc. (purity > 99%), isobutene (purity > 99%) and terpinolene (purity > 85 %) were from Fluka Analytical and TME (purity > 98%), isoprene (purity > 99%), limonene (purity > 97 %),  $\alpha$ -pinene (purity > 97 %),  $\beta$ -pinene (purity > 97 %),  $\alpha$ -terpinene (purity > 85 %), estragole (purity > 98 %) and myrcene (purity > 99 %) were from Sigma-Aldrich. All reagents were used as supplied.

Limit of detection (LOD)\* Number Manufacturer Model Institution NO NO<sub>2</sub>NO<sub>2</sub> convertor (ppt) (ppt) 1 Thermo Fisher Scientific TE42i-TL Birmingham Heated molybdenum 210 210 2 **EUPHORE** Teledyne API 200AU Heated molybdenum 190 450 3 Eco Physics AG **EUPHORE** CLD 770 AL ppt Xenon lamp 150 430 and PLC 760 4 60 150 Air Quality Design, Inc. (AQD) York Blue light at 395 nm

**Table 1.** Details of the  $NO_x$  monitoring instruments used. Note: ppt – parts per trillion.

## 3.2 Data analysis

The limit of detection (LOD) for each instrument was determined, under the actual experimental conditions, as three times the standard deviation of the NO and NO2 signal recorded each day from the empty chamber prior to the start of experiments (i.e. before addition of any reactants). The mean LODs determined for NO and NO<sub>2</sub> are shown in Table 1. These LOD values are higher than those quoted by the manufacturers for monitors 1–4 (typically 2–100 ppt) but accurately reflect the actual performance of the instruments as used during these experiments. In the analysis which follows, in order to confirm that any change in the measured NO and NO<sub>2</sub> mixing ratio for each alkene addition was not due to noise or drift and therefore came from signal, the readings were compared to the experimentally determined LOD for each instrument. Only in cases where the measured change was greater than the experimentally determined LOD were these readings used for determining an interference. The interference due to the VOC was determined by means of linear regression (least squares fit), with the slopes and their uncertainty and Pearson's correlation coefficients calculated in IGOR (see Tables 2 and 3).

#### 4 Results

Figures 2–4 give the measured VOC mixing ratios and the retrieved NO and NO<sub>2</sub> measurements of the four monitors during the experiment for selected alkenes, along with the regression analysis for determining the interference levels. Spikes in NO and NO<sub>2</sub> mixing ratios observed after an alkene addition (e.g. Fig. 4) arise from sampling close to the addition point prior to the initial period of mixing in the chamber ( $\sim 3$  min) and were disregarded in the analysis. The slow decay of alkene and NO<sub>x</sub> mixing ratios following each addition arises from dilution effects (with a first-order rate constant of  $\sim 5.7 \times 10^{-5}$  s<sup>-1</sup> derived from the decay of SF<sub>6</sub>).

From Figs. 2 to 4, a clear and systematic response from the monitors to the presence of  $\alpha$ -terpinene, terpinolene and T2B was observed, with the magnitude varying between the mon-

itors. In addition to the alkenes shown in Figs. 2–4, significant interference effects were also observed for C2B, TME and limonene for some of the monitors, as summarised in Tables 2 and 3. No interference was observed, within the detection uncertainty, for ethene, propene, isobutene,  $\alpha$ -pinene,  $\beta$ -pinene, myrcene or methyl chavicol in any of the monitors. For isoprene, no statistically significant interference was observed for monitors 1–3, while monitor 4 observed very small positive interferences of 0.035  $\pm$  0.001 % (NO channel) and 0.076  $\pm$  0.002 % (NO<sub>2</sub> channel).

For the alkenes in which a significant interference was observed, in general a positive interference was observed for NO and a negative interference for NO<sub>2</sub> by monitors 1–4 (Tables 2 and 3), with the exception of TME, in which a negative NO interference was observed by monitor 3 (discussed later). Generally, for monitor 4, a positive NO interference and a mixture of both positive and negative NO<sub>2</sub> interferences was observed. Overall, while the magnitude of interference differed between the monitors, the same trend in the interference was observed, with  $\alpha$ -terpinene having the largest interference effect, followed by terpinolene, TME/T2B, C2B and limonene.

The addition of water (RH ca. 30%) led to the observed NO and NO<sub>2</sub> interference for T2B, terpinolene and  $\alpha$ -terpinene decreasing by 30%–60%, as shown in Tables 2 and 3. The addition of CO resulted in an increase in the NO interference observed for TME from below the LOD to 0.7% for monitors 1 and 2, while monitors 3 and 4 exhibited a larger interference increase (Table 2).

## 5 Discussion

#### 5.1 Interference effects on retrieved NO abundance

Positive NO interferences were observed for those alkenes which reacted most rapidly with ozone and, hence, will be present within the monitor reaction chamber at different levels in the measurement and background modes. This interference is attributed to chemiluminescent emission following the alkene—ozone reaction and may be attributed to a combi-

<sup>\*</sup> Calculated in this study.

**Table 2.** Measured NO interference ( $\% \pm 1$  SD of the slope) for each monitor across a range of different alkenes. Note: LOD – limit of detection; TME – 2,3-dimethyl-butene; T2B – trans-2-butene; C2B – cis-2-butene.

Species	TE42i-TL	Teledyne API 200AU	Eco Physics AG CLD770	Air Quality Design, Inc. (AQD)
C2B	< LOD	< LOD	$0.40 \pm 0.05$	$0.38 \pm 0.01$
TME	< LOD	< LOD	$-0.70 \pm 0.09$	$1.10 \pm 0.01$
T2B	< LOD	< LOD	$1.00 \pm 0.01$	$0.83 \pm 0.01$
Terpinolene	$0.50 \pm 0.05$	< LOD	$1.30 \pm 0.01$	$4.40 \pm 0.15$
$\alpha$ -terpinene	$1.90 \pm 0.05$	$0.50 \pm 0.04$	$2.30 \pm 0.04$	$10.9 \pm 0.06$
Limonene	< LOD	< LOD	< LOD	$-0.10 \pm 0.01$
$TME + H_2O$	< LOD	< LOD	0.60	2.40
T2B and H2O	< LOD	< LOD	$0.48 \pm 0.01$	$0.37 \pm 0.01$
Terpinolene and H <sub>2</sub> O	$0.25 \pm 0.03$	< LOD	$0.88 \pm 0.01$	$1.60 \pm 0.10$
α-terpinene and H <sub>2</sub> O	$1.00 \pm 0.07$	< LOD	$1.30 \pm 0.06$	$6.20 \pm 0.70$
TME and CO	$0.70 \pm 0.01$	$0.66 \pm 0.09$	$1.30 \pm 0.12$	$1.40 \pm 0.02$

**Table 3.** Measured  $NO_2$  interference ( $\% \pm 1$  SD of the slope) for each monitor across a range of different alkenes (LOD – limit of detection).

Species	TE42i-TL	Teledyne API 200AU	Eco Physics AG CLD770	Air Quality Design, Inc. (AQD)
C2B	$-0.60 \pm 0.10$	< LOD	$-1.10 \pm 0.08$	$0.30 \pm 0.02^*$
TME	$-0.63 \pm 0.05$	< LOD	$-0.78 \pm 0.15$	$-0.92 \pm 0.10^{\#}$
T2B	$-0.50 \pm 0.06$	< LOD	$-0.50 \pm 0.03$	$-0.93 \pm 0.02^{\#}$
Terpinolene	$-0.61 \pm 0.02$	< LOD	$-0.18 \pm 0.03$	$-0.94 \pm 0.21^*$
$\alpha$ -terpinene	$-1.90 \pm 0.13$	< LOD	$-1.00 \pm 0.20$	$3.10 \pm 2.10$
Limonene	< LOD	< LOD	< LOD	$0.09 \pm 0.01^{\#}$
TME and H <sub>2</sub> O	-0.60	< LOD	< LOD	-2.00
T2B and H2O	< LOD	< LOD	< LOD	$-0.41 \pm 0.02$
Terpinolene and H <sub>2</sub> O	$-0.29 \pm 0.02$	< LOD	< LOD	-0.25
α-terpinene and H <sub>2</sub> O	$-0.98 \pm 0.06$	< LOD	< LOD	$0.35 \pm 0.10$
TME and CO	$-0.70 \pm 0.01$	< LOD	< LOD	$1.00 \pm 0.30$

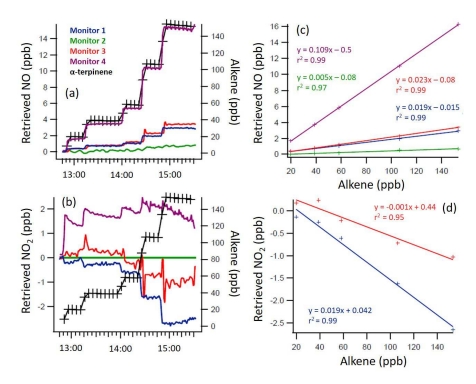
<sup>\*</sup> determined by method (a) – see Sect. 5.3, # determined by method (b) – see Sect. 5.3.

nation of two factors, namely the formation of excited products in the alkene—ozone reaction, which emit chemiluminescence, coupled with the significant removal of some alkenes during the instrument background phase compared with the measurement phase through their reaction with (elevated levels of) ozone within the instrument, i.e. mechanism (4), outlined above.

Possible origins of this signal are the production of excited HCHO, vibrationally excited OH and electronically excited OH (e.g. Finlayson et al., 1974). While the long-pass filters used in chemiluminescence  $NO_x$  monitors should preclude emissions from electronically excited species, vibrationally excited OH produced through the hydroperoxide mechanism is known to emit in the 700–1100 nm wavelength range (Finlayson et al., 1974; Schurath et al., 1976; Hansen et al., 1977; Toby, 1984) and would be detected as  $NO_2$ . Specifications of the long-pass filters used in the chemiluminescence  $NO_x$  monitors in this study are not reported in their respective user manuals but typically block light below ca. 600 nm, while

typical PMT response characteristics are between 400 and 950 nm (Jernigan, 2001). Any chemiluminescence signal in the 600–950 nm wavelength range can therefore cause a potential interference.

The difference in the interference effect among monitors may then reflect differences in the conditions (e.g. ozone abundance, pressure and residence time) within the reaction cell and filter specifications. The relative magnitudes of the positive interference signals observed between the different monitors are consistent with this picture as the reaction chamber pressure is much lower for monitors 3 and 4 (ca. 1–10 Torr) compared with monitors 1 and 2 (ca. 300 Torr), leading to greater collisional quenching. Similarly, the addition of H<sub>2</sub>O, which would be expected to efficiently accept vibrational energy from OH radicals (Gerboles et al., 2003), was found to substantially reduce the apparent interference. In the experiments with higher humidity, a reduced interference (factor of ca. 2; see Table 2) was observed for all NO experiments for all instruments, except for TME for the pho-



**Figure 2.** Time series of the  $\alpha$ -terpinene mixing ratio and indicated/measured NO (**a**, **c**) and NO<sub>2</sub> (**b**, **d**) mixing ratios, as directly retrieved by each monitor (**a**, **b**), with 1 min time resolution, and the regression calculations for the monitors that demonstrated significant interference with the addition of  $\alpha$ -terpinene (**c**, **d**). Note the different y axis scales.

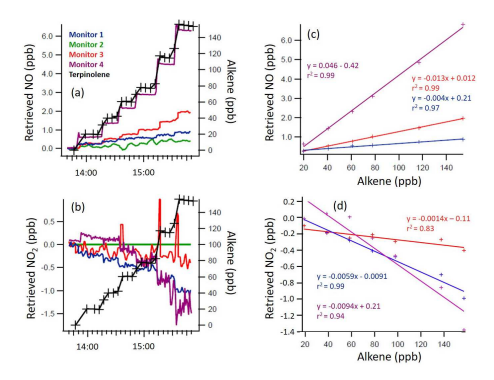
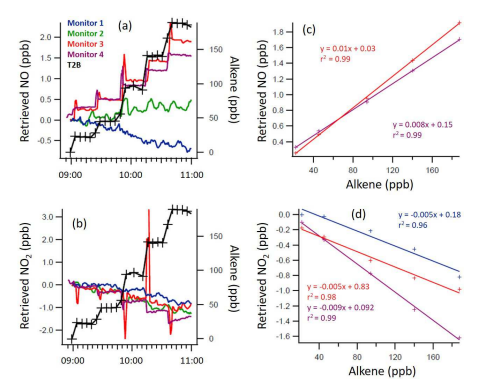


Figure 3. Time series of the terpinolene mixing ratio and measured NO and NO<sub>2</sub> mixing ratios, as retrieved by each monitor  $(\mathbf{a}, \mathbf{b})$ , with 1 min time resolution, and the regression calculations for the monitors that demonstrated significant interference with the addition of terpinolene  $(\mathbf{c}, \mathbf{d})$ . Note the different y axis scales.



**Figure 4.** Time series of the trans-2-butene (T2B) mixing ratio and measured NO (**a**, **c**) and NO<sub>2</sub> (**b**, **d**) mixing ratios, as retrieved by each monitor (**a**, **b**), with 1 min time resolution, and the regression calculations for the monitors that demonstrated significant interference with the addition of T2B (**c**, **d**). Note the different y axis scales.

tolytic converters for which an increase was observed. There is currently no recommended relative humidity under which calibrations should be performed for any of the instruments or within EU and EPA guidelines (AQEG, 2004; USEPA, 2002). However, the installation of permeation driers at the sample inlet should (in principle) reduce the impact of different H<sub>2</sub>O/relative humidity levels upon the quenching of NO<sub>2</sub> or other species and are a common feature of most modern samplers (AQEG, 2004).

## 5.2 Interference magnitude – kinetic and structural effects

The most significant effects are the large positive NO interferences observed for the monoterpenes, namely  $\alpha$ -terpinene and terpinolene, within monitors 1, 3 and 4. The criteria for an alkene to display such a positive interference (i.e. via mechanism 4) are that it reacts with ozone to produce excited products which exhibit a chemiluminescent signal at appropriate wavelengths. In addition, the alkene must have a sufficiently rapid reaction with ozone so that its mixing ratio is substantially reduced during the instrument background phase compared with the measurement phase, thus precluding the correct subtraction of the interference signal. The reaction rate constants for many alkenes with ozone are well known, allowing the calculation of a kinetic interference potential (KIP) ranking for this second factor, and are calculated

$$KIP = 100 \times \left(1 - \exp^{\left(-k't \times \frac{k_{(Alkene+O_3)}}{k_{(NO+O_3)}}\right)}\right),\tag{1}$$

where  $k = k[O_3]t$  and  $\frac{[NO]}{[NO]_0} = 0.01$  (i.e. 1% of NO left after reaction with excess  $O_3$ ; see the Supplement for calculation details). The calculated KIP are shown in Table 4 as the percentage of a given alkene's potential chemiluminescent signal which would not be subtracted in the standard background cycle, under the assumption that the background cycle conditions ( $O_3$  mixing ratio and residence time) would be sufficient to remove 99% of the NO present.

This ranking does not reflect the precise (relative) interference which is observed as it neglects structural features which will affect the yield (and state i.e. electronic or vibrationally excited) of the chemiluminescent products from the ozonolysis reaction, but it is consistent with the trend and relative magnitudes for the substantial positive interferences shown in Tables 2 and 3. For example, a lack of interference is observed for myrcene and limonene, both of which exhibit terminal C=C bonds (see Table 4) and, after reaction with ozone, lead to the production of the CH<sub>2</sub>OO Criegee intermediate (CI), which subsequently decomposes or undergoes rearrangement to form small yields of OH (Alam et al., 2011). The ozonolysis of internal alkenes, such as C2B and

T2B, produce the CH<sub>3</sub>CHOO CI, which predominantly decomposes via the vinyl hydroperoxide mechanism, forming larger yields of OH (Johnson and Marston, 2008; Alam et al., 2013). Such chemically formed OH that produces a detectable signal may also be augmented by contributions from HO<sub>2</sub> and RO<sub>2</sub> converted into OH within the instrument by reaction with NO, especially in the NO<sub>2</sub> channel of photolytic converter instruments.

The relationship between the KIP (Table 4) and measured NO interference (Tables 2 and 3) is illustrated in Fig. 5 and can be used for predicting the potential interference of a given alkene to the NO signal from a kinetic perspective. For example,  $\alpha$ -humulene has a KIP of 94.54%, which could give rise to a 1.7 %, 2.4 % or 10.2 % NO interference for monitors 1, 3 and 4, respectively. This estimate is, however, based on the rate constant of  $\alpha$ -humulene alone and does not include any structural features such as the presence of terminal and non-terminal C=C bonds.

## Explanation of the interference observed for NO<sub>2</sub>

The above discussion considers only the interference effect arising from alkene chemiluminescent emission; further measurement impacts are also evident in the (negative) interferences apparent for other species/monitors in Tables 2 and 3. An inspection of Tables 2 and 3 shows smaller positive interferences, and some negative interferences, from alkenes in the NO<sub>2</sub> measurements.

NO<sub>2</sub> measurements using chemiluminescence approaches are usually obtained by measuring  $NO_x$  (i.e.  $\Sigma(NO + NO_2)$ , after passing the sampled air through an NO<sub>2</sub> converter) and subtracting the (independently determined) NO contribution. If the actual interference signal (additional chemiluminescence) during the  $NO_x$  measurement mode arises solely from mechanism (4), ozonolysis chemiluminescence, then this would be expected to match that in the NO mode (subject to the alkene abundance not being altered in the NO<sub>2</sub> conversion stage and the detection conditions for the NO and  $NO_x$  phases being identical) and, consequently, would not affect the retrieved NO<sub>2</sub> mixing ratio. Monitors 1, 2 and 3 used a single detection cell, alternating between NO and  $NO_2$  ( $NO_x$ ) modes, and measured the  $NO_2^*$  chemiluminescence signal under identical conditions (optical arrangement, filtering and pressure). The observed negative interference for NO<sub>2</sub>, therefore, may have arisen due to the removal of alkene by the Mo catalyst within the monitors.

For monitor 1 (TE42i-TL), the negative interference observed for NO2 was the same magnitude as that observed for the positive interference for NO, including the experiments with H<sub>2</sub>O and CO (see Fig. 6 and Tables 2–3). This response is thought to arise as a consequence of the calculation methodology, combined with removal of alkenes during the NO<sub>2</sub> conversion by the Mo catalyst.

There are three modes of operation in monitor 1 (TE42i-TL), namely NO measurement, NO<sub>2</sub> / NO<sub>x</sub> measurement and background (pre-reactor) measurement, given by Eqs. (2)–(4), respectively, as follows:

$$sNO = sNO_{real} + X_i,$$
 (2)

$$sNO_x = sNO_{x \text{ real}} + yX_i, \tag{3}$$

$$sP = fX_i, (4)$$

where sNO and sNO<sub>x</sub> are the NO and NO<sub>x</sub> signals produced by the chemiluminescence monitor, respectively, sNO<sub>real</sub> and  $sNO_{xreal}$  are the real NO and  $NO_x$  signals,  $X_i$  denotes the interference alkene i, y is the fraction of the interferant (alkene)  $X_i$  remaining after the Mo convertor, sP denotes signal at the pre-reactor, and f is the fraction of  $X_i$  remaining after the pre-reactor. The mixing ratios of NO, NO<sub>2</sub> and NO<sub>x</sub> are given by the following:

$$[NO] = \frac{sNO - sP}{cNO},\tag{5}$$

$$[NO] = \frac{(sNO_{real} + X_i) - fX_i}{cNO},$$
(6)

$$[NO] = \frac{(sNO_{real} + (1 - f)X_i)}{cNO},$$

$$[NO_x] = \frac{sNO_x - sP}{cNO_x},$$
(8)

$$[NO_x] = \frac{sNO_x - sP}{cNO_x},$$
(8)

$$[NO_x] = \frac{(sNO_{xreal} + yX_i) - fX_i}{cNO_x},$$

$$[NO_x] = \frac{(sNO_{xreal} + (y - f)X_i)}{cNO_x},$$
(10)

$$[NO_x] = \frac{(sNO_{xreal} + (y - f)X_i)}{cNO_x},$$
(10)

$$[NO_2] = \frac{[NO_x] - [NO]}{CE},$$
(11)

$$[NO_2] = \frac{(sNO_{x \text{real}} + (y - f)X_i)}{cNO_x \times CE},$$

$$-\frac{(sNO_{real} + (1 - f)X_i)}{cNO \times CE},$$
(12)

where c is the span factor and CE represents the conversion efficiency. If we assume  $cNO_x \approx cNO \approx c$ , then the following applies:

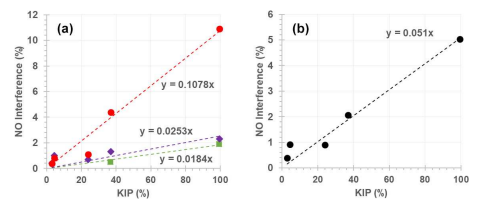
$$[NO_2] =$$

$$\frac{(sNO_{x \text{ real}} + (y - f)X_i) - (sNO_{\text{real}} + (1 - f)X_i)}{c \times CE}.$$
 (13)

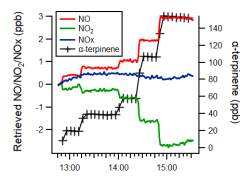
From Eq. (13), it may be seen that if y = 1 (i.e. if the interferant alkene abundance is not affected by passage through the Mo converter), then there would be no interference observed in the retrieved NO<sub>2</sub>, while if the interferant species is subject to removal during its passage through the converter, then y < 1 and a negative interference would be observed. Molybdenum oxide catalysts have been reported to efficiently isomerise alkenes at temperatures between 300 and 400 °C (Wehrer et al., 2003) and are also effective catalysts for the epoxidation of alkenes (Shen et al., 2019). The observed small negative interference effects (for monitors 1 and 2; the Mo converter units), in the absence of significant

**Table 4.** Kinetic ranking of the interference potential, namely the percentage of the potential chemiluminescent signal from the ozonolysis of a given alkene which would not be removed by a standard instrument background cycle under conditions (e.g. ozone mixing ratio and residence time) which would remove 99 % of the NO sampled. Rate constants are taken from Calvert et al. (2000);  $k_{(NO+O_3)} = 1.90 \times 10^{-14}$  cm<sup>3</sup> molecule<sup>-1</sup> s<sup>-1</sup> (298 K). Note: this ranking does not include variations in the yield of chemiluminescent products with an alkene structure which will modulate the values given. Species marked with an \* are investigated in this study.

Species	$k_{(\text{Alkene+O}_3)}$ (298 K)/ cm <sup>3</sup> molecule <sup>-1</sup> s <sup>-1</sup>	Kinetic interference potential (%)	No. of C=C bonds	No. of terminal C=C bonds
Ethene	$1.58 \times 10^{-18}$	0.04*	1	1
1-butene	$9.64 \times 10^{-18}$	0.23	1	1
2,3-dimethyl-1-butene	$1.00 \times 10^{-17}$	0.24	1	1
Propene	$1.01 \times 10^{-17}$	0.24*	1	1
1-pentene	$1.06 \times 10^{-17}$	0.26	1	1
Isobutene	$1.13 \times 10^{-17}$	0.27*	1	1
Isoprene	$1.28 \times 10^{-17}$	0.31*	1	1
2-methyl-1-butene	$1.30 \times 10^{-17}$	0.31	1	1
$\beta$ -pinene	$1.50 \times 10^{-17}$	0.36*	1	1
$\alpha$ -cedrene	$2.80 \times 10^{-17}$	0.68	1	0
3-carene	$3.70 \times 10^{-17}$	0.89	1	0
α-pinene	$8.66 \times 10^{-17}$	2.08*	1	0
C2B	$1.25 \times 10^{-16}$	2.98*	1	0
Cis-3-hexane	$1.44 \times 10^{-16}$	3.43	1	0
Trans-3-hexane	$1.57 \times 10^{-16}$	3.73	1	0
α-coapene	$1.58 \times 10^{-16}$	3.76	1	0
T2B	$1.90 \times 10^{-16}$	4.50*	1	0
Limonene	$2.00 \times 10^{-16}$	4.73*	2	1
2-carene	$2.30 \times 10^{-16}$	5.42	1	0
2-methyl-2-butene	$4.03 \times 10^{-16}$	9.31	1	0
Myrcene	$4.70 \times 10^{-16}$	10.77*	3	2
2,3-dimethyl-2-butene	$1.13 \times 10^{-15}$	23.96*	1	0
Terpinolene	$1.90 \times 10^{-15}$	36.90*	2	0
$\alpha$ -humulene	$1.20 \times 10^{-14}$	94.54	3	0
$\beta$ -carophyllene	$1.20 \times 10^{-14}$	94.54	2	1
$\alpha$ -terpinene	$2.10 \times 10^{-14}$	99.38*	2	0



**Figure 5.** Relationship between measured NO interference (%) and kinetic interference potential (KIP; %) for monitors 1 (green), 3 (purple), 4 (red) and the average of the observed NO interference across all instruments (black).



**Figure 6.** Time series of the  $\alpha$ -terpinene mixing ratio (black) and measured NO (red), NO<sub>2</sub> (green) and NO<sub> $\chi$ </sub> (blue) mixing ratios, as retrieved by monitor 1 (TE42i-TL), with a 1 min time resolution.

sampled  $NO_x$ , may reflect the partial removal of the alkene in the converter.

The negative NO<sub>2</sub> interference apparent for monitors 3 and 4 (photolytic converter instruments) is more difficult to rationalise (as no Mo catalyst is present). Under ambient conditions, where  $NO_x$  is present, mechanism (3) may occur, as outlined below. In reality, the conversion efficiency for photolytic converters is substantially lower than 100 % (Reed et al., 2016) as a consequence of both the finite photolysis intensity achievable and the occurrence of the  $NO + O_3$  back reaction. If the instrument calibration factor for NO<sub>x</sub> is not equal to that for NO (see Eq. 12), or if alkene was removed in the convertor stage, then this will lead to different interferences for NO and NO2, as CE is also (significantly) less than 1. This trend is apparent in the values shown in Table 3, in particular for the instruments fitted with photolytic convertors. However, in the absence of sampled  $NO_x$ , the observed less positive or even negative NO2 interference suggests that less alkene is present in the  $NO_x$  mode. Direct photolysis of alkenes is unlikely to cause such a change, considering the photolytic converter wavelength envelope, but the photolytic production of  $HO_x$  radicals (which then react with the alkene) may be responsible.

Monitor 4 (Air Quality Design, Inc. – AQD) used independent NO<sub>2</sub>\* detection channels; tests were conducted using both channels for C2B and terpinolene systems and revealed significant differences between the two detectors (ca. 40 % lower interference response for NO in the NO<sub>2</sub> detection channel). With two independent detection channels,  $NO_2$  may be determined from the  $NO_x$  measurement by either subtracting the NO level obtained from the NO channel (method a) or via the difference in signal observed in the  $NO_2 / NO_x$  channel when turning the photolysis lamp on and off (method b). Under method (a), as employed for C2B and terpinolene, a lower positive interference from alkene chemiluminescence results as a consequence of the difference in the detection cell conditions (results marked \* in Table 3), while under method (b), as employed for the other alkenes studied here with the AQD system, the interference (from mechanism 4 alone) should cancel out (results marked with a # in Table 3).

## 5.4 Effect of quenching by the alkenes

The data presented in Figs. 2-4 and Tables 2 and 3 show both negative and positive interferences, while mechanism 4 alone would be expected to result in positive interference signals for NO for all alkenes. We therefore conclude that additional mechanisms are occurring. Under the conditions of these chamber experiments, the retrieval of additional NO<sub>v</sub> species can be precluded. The chamber wall source of HONO has been characterised and shown to produce ppt levels of HONO under the dark, dry conditions of these experiments (Zádor et al., 2005) and would be equally present for all experiments. We attribute the negative (or reduced positive) interference effects to a combination of mechanisms (1) and (3), where the quenching of excited OH (produced by alkene and ozone reaction) by alkenes (electron-rich alkenes have been shown to be effective quenchers; Gersdorf et al., 1987; Chang and Schuster, 1987) and the generation of  $HO_x$  radicals within the instrument follow on from the ozonolysis reaction.

The alkene–ozone reactions are known to produce OH,  $HO_2$  and  $RO_2$  radicals both directly (e.g. Johnson and Marston, 2008), following the photolysis of other alkene–ozone reaction products (e.g. carbonyl compounds), and through OH–alkene reactions. Peroxy radicals promote the conversion of NO to  $NO_2$ , altering the abundance of both species (the formation of  $NO_x$  reservoirs, such as nitric acid and organic nitrates, will also occur but will be negligible on the timescale of the operation of most instruments).

The ozonolysis of the TME results in the production of OH with a close to unity yield (IUPAC, 2020), and if taking into account the above mechanism (4) only, might be expected to exhibit a large interference in NO mode. Table 2 shows no interference for monitors 1 and 2 (Mo convertor units) and negative and positive interferences for monitors 3 and 4 (photolytic convertor units), respectively, and so it is hard to rationalise (for NO mode). The addition of CO as a scavenger for OH led to an increase in the NO signal for all monitors. A possible origin of this signal is the chemiluminescence production of the excited intermediate HOCO (from the reaction of vibrationally excited OH, which is from the ozonolysis of TME, with CO), which has a temperature and pressure-dependent rate of reaction (Atkinson et al., 2006; Li and Francisco, 2000) and is consistent with the larger NO signal in the photolytic monitors (Table 2).

## 6 Conclusions

The interference in chemiluminescence  $NO_x$  measurements from alkenes has been systematically investigated using four commercially available monitors. Varying degrees of inter-

ference in the NO and NO<sub>2</sub> signals were observed for all monitors investigated, which has been attributed to a combination of mechanisms 1, 3 and 4, particularly for the incomplete subtraction of chemiluminescence from the products of alkene-ozone reactions that manifest due to a significant removal of the alkene during the instrument background cycle. Monoterpenes,  $\alpha$ -terpinene and terpinolene, exhibit the largest interferences, followed by 2,3-dimethyl-2-butene (TME) and trans-2-butene (T2B), which is in line with the calculated kinetic interference potential (KIP; see Table 4). The KIP can be used as a crude indicator of a potential interference of an alkene to an NO signal but has large margins of error as it does not take into account the variation in the yield of chemiluminescent products and other instrumental differences. The alkene interference observed with enhanced RH conditions also indicates the need to accurately calibrate chemiluminescence NO<sub>x</sub> analysers under actual sampling conditions.

The interferences in NO measurements arising from alkenes among the monitors investigated in this study range from 1% to 11%. The varying responses exhibited by the different monitors reflect differences in the conditions within the instrument (ozone abundance, pressure and residence time) within the reaction cell and filter specifications. The magnitude of the NO and NO2 interferences not only vary with different alkenes and commercial monitors but will also be dependent upon sampling environments (and with ambient  $NO_x$  and alkene concentrations). Notably, in these experiments, the alkene abundance is high compared with most ambient air samples; consequently, internally generated OH will react essentially exclusively with the alkene, which may not reflect ambient sampling but which we do not expect to impact the conclusions reached with respect to mechanism 4, i.e. interference in retrieved NO levels. Further research to explore these impacts, and other parameters (e.g. H<sub>2</sub>O abundance), is urgently needed. The chemiluminescence from the monoterpene ozonolysis should also be investigated to identify the emission spectra of possible interfering species; given the varying OH yields and energetics from the ozonolysis of different alkenes, their intensity of emission are likely to vary. A combination of selective long-pass filters and detector characteristics can then be exploited within chemiluminescence NO<sub>x</sub> monitors to eliminate such interferences with similar emission spectra to  $NO_2^*$ .

Mixing ratios of  $NO_x$  vary from > 100 ppb in some urban areas, e.g. Marylebone Road (Carslaw, 2005), < 300 ppt in biogenic environments (Hewitt et al., 2010) and < 35 ppt in remote areas (Lee et al., 2009). For typical urban environments where alkene mixing ratios are relatively low (< 2 ppb; e.g. von Schneidemesser et al., 2010), the interferences identified here are not likely to be significant ( $\sim$  1 % of the NO signal). However, for biogenic environments where monoterpenes and sesquiterpenes, which react rapidly with ozone, are abundant, the interference could be significantly larger. For example, the average mixing ratios for isoprene

( $\sim$  1 ppb), five monoterpenes ( $\sim$  220 ppt), three short chain alkenes ( $\sim$  240 ppt) and NO (0.14 ppb) were measured within a southeast Asian tropical rainforest (Jones et al., 2011). Using the relationship between KIP and NO interference, an overestimation of NO levels of to up to 58 % would result, with very significant implications for the prediction of other atmospheric chemical processes involving NO<sub>x</sub>. Given that NO<sub>x</sub> mixing ratios are relatively small in biogenic and remote environments, these interferences could lead to their substantial overestimation. Such alkene interference may contribute to the relatively high NO and low NO<sub>2</sub> reported in the tropical rainforests at night, which could not otherwise be accounted for (Pugh et al., 2010).

Within indoor environments,  $NO_x$  primarily arises from outdoor sources or indoor combustion sources (Young et al., 2019). Typically, in the absence of a known indoor combustion source, indoor NO levels are low (ca. 13 % of outdoor levels), with NO<sub>2</sub> comprising the majority of the NO<sub>x</sub> (Zhou et al., 2019). There are multiple sources of alkenes indoors, such as fragranced volatile personal care products (Nemafollahi et al., 2019; Yeoman et al., 2020) and cleaning products (Kristenson et al., 2019), resulting in much larger levels of alkenes than NO<sub>x</sub> (McDonald et al., 2018; Kristenson et al., 2019). Consequently, monoterpenes are among the most ubiquitous VOCs reported for indoor air, with the main species including linalool,  $\alpha$ -pinene,  $\beta$ -myrcene and limonene (Krol et al., 2014; Nematollahi et al., 2019). Peak limonene mixing ratios may be a factor of ca. 50 higher indoors than in outdoor environments (Lerner et al., 2012). Although the monoterpenes,  $\alpha$ -pinene, myrcene and limonene show no significant NO interferences in chemiluminescence  $NO_x$  monitors, other fast-reacting monoterpenes (with  $O_3$ ), such as  $\alpha$ -terpinene and terpinolene which are not generally reported in the literature, exhibit quite large interferences and may lead to very substantial overestimations in indoor NO<sub>x</sub> measurements. Monoterpene mixing ratios in indoor environments can be further enhanced by cleaning activities (Singer et al., 2006; Kristenson et al., 2019; Weschler and Carslaw, 2018). Indoor  $\alpha$ -terpinene and  $\alpha$ -pinene mixing ratios have exceeded 10 and 68 ppb, respectively (Singer et al., 2006; Brown et al., 1994). These relatively large monoterpene mixing ratios may lead to substantial interferences in chemiluminescence NO<sub>x</sub> monitors; their incorrect retrieval, as measured  $NO_x$ , will impact assessments of indoor air chemistry, indoor air quality and, hence, health.

Data availability. Experimental data are available at https://doi.org/10.25500/eData.bham.00000566 (Alam et al., 2020) and also from the Eurochamp Data Center at https://data.eurochamp.org/data-access/chamber-experiments/ (last access: 4 August 2020).

*Supplement.* The supplement related to this article is available online at: https://doi.org/10.5194/amt-13-5977-2020-supplement.

Author contributions. MSA, WJB and JDL conceived and planned the experiments. MSA, JDL, MVM, AM and MR performed the experiments. LRC, LJK and MSA performed the data analysis. LRC, LJK, MSA, CP and WJB contributed to the data investigation and curation. MSA wrote the original draft, and all co-authors contributed to the reviewing and editing of the paper.

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

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