

The determination of ClNO₂ via thermal dissociation–tunable infrared laser direct absorption spectroscopy

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Abstract. Nitryl chloride (ClNO₂) is a reservoir species of chlorine atoms and nitrogen oxides, both of which play important roles in atmospheric chemistry. To date, all ambient ClNO₂ observations have been obtained by chemical ionization mass spectrometry (CIMS). In this work, thermal dissociation–tunable infrared laser direct absorption spectrometry (TD-TILDAS) is shown to be a viable method for quantifying ClNO₂ in laboratory and field settings. This technique relies on the thermal dissociation of ClNO₂ to create chlorine radicals, which undergo fast reactions with hydrocarbons to produce hydrogen chloride (HCl) that is detectable by the TILDAS instrument. Complete quantitative conversion of ClNO₂ to HCl was achieved at temperatures > 400 °C, achieving 1 Hz measurement precision of 11 ± 1 pptv (3σ limits of detection of 34 ± 2 pptv) during laboratory comparisons with other ClNO₂ detection methods. After blank and line loss corrections, method accuracy is estimated to be within ± 5 %. Performance metrics of TD-TILDAS during ambient sampling were a 1 Hz precision of 19 ± 1 pptv and 3σ limits of detection of 57 ± 3 pptv, which is directly comparable to previously reported ClNO₂ detec-

tion by quadrupole CIMS. Thus, TD-TILDAS can provide an alternative analytical approach for a direct measurement of ClNO₂ that can complement existing datasets and future studies. The quantitative nature of TD-TILDAS also makes it a potentially useful tool for the calibration of CIMS instruments. However, interpretation of ambient data may be complicated by potential interferences from unaccounted-for sources of thermolabile chlorine, such as ClNO, chloramines, and organochlorides.

1 Introduction

Nitryl chloride (ClNO₂) is an important nighttime reservoir of two highly reactive atmospheric species: atomic Cl and NO₂. Atomic Cl radicals play multifaceted roles in oxidation chemistry throughout the boundary layer (Simpson et al., 2015), including hydrocarbon oxidation (Atkinson et al., 2006a, and references therein), ozone production and destruction (Halfacre and Simpson, 2022; Liao et al., 2014; Sarwar et al., 2012, 2014; Simon et al., 2009; Wang et al., 2016),

and mercury depletion (Driscoll et al., 2013). However, the quantitative magnitude to which they affect these processes remains an open question. On the other hand, NO₂ is one of the principal components of photochemical smog and the major anthropogenic precursor for ozone production. Accounting for all sources of NO₂ is therefore important for accurately informing chemical and air quality models.

The first in situ observation of ambient ClNO₂ was reported by Osthoff et al. (2008) utilizing chemical ionization mass spectrometry (CIMS) in the polluted marine boundary layer. CIMS has since been used in a multitude of studies for additional ClNO₂ observations worldwide, including North America (Jaeglé et al., 2018; Lee et al., 2018a, b; Mielke et al., 2011; Riedel et al., 2012, 2013; Thornton et al., 2010; Wagner et al., 2013; Young et al., 2012), Europe (Bannan et al., 2015; Phillips et al., 2012; Sommariva et al., 2018; Tan et al., 2022), Asia (Le Breton et al., 2018; Liu et al., 2017; Tham et al., 2016, 2018; Wang et al., 2022, 2016, 2017; Xia et al., 2020; Ye et al., 2021; Yu et al., 2020; Zhou et al., 2018), in the presence of snow/ice (Kercher et al., 2009; McNamara et al., 2020), and in indoor air quality studies (Moravek et al., 2022). Limits of detection are often reported at 10⁰ pptv under 25–30 s averaging times (Bannan et al., 2015; Kercher et al., 2009; McNamara et al., 2020; Mielke et al., 2011) and have been recently reported at sub-pptv for 1 s measurements (Decker et al., 2024). Typical observed mixing ratios range from 10¹–10³ pptv, with the highest levels observed in coastal polluted regions, where sources of nitrogen oxides and Cl[−]-rich aerosols are plentiful (Wang et al., 2019, 2021, and references therein).

While CIMS is a highly effective technique, ClNO₂ quantitation involves nontrivial calibration work. A laboratory source of ClNO₂ may be readily generated by flowing a known amount of N₂O₅ across a Cl[−]-containing salt bed (or Cl₂ across an NO₂[−]-containing salt bed), but its quantitation assumes unit conversion out of the salt bed (e.g. Osthoff et al., 2008) or requires additional equipment to observe ClNO₂ thermal dissociation products, such as a thermal dissociation–cavity ring-down spectrometer (TR-CRDS) (Thaler et al., 2011) or a cavity-attenuated phase shift spectrometer (CAPS) (e.g. Tan et al., 2022). Further, I[−]-based CIMS demonstrates variable sensitivities based on the temperature and relative humidity of the ion molecule reactor, thereby requiring substantial laboratory work to develop humidity- and temperature-dependent calibration factors (Lee et al., 2014; Robinson et al., 2022). Thus, there is an opportunity for an innovative method that can detect ClNO₂ directly without the need for supplemental instrumentation.

The advantages of optical methods include analyte specificity and near-absolute detection, utilizing well-defined physical absorption properties, and requiring only infrequent calibrations or method validation procedures. Thaler et al. (2011) previously used a TR-CRDS system to detect ClNO₂ as NO₂ by absorption at 405 nm under laboratory conditions, achieving CIMS-competitive metrics (e.g.

reported a [CE1](#) 20 pptv limit of detection for 1 min averaging). This was achieved by flowing sample air through both an unheated reference pathway and a heated (450 °C) sample pathway, under which ClNO₂ would thermally dissociate into Cl radicals and NO₂ (Reaction R1).



The difference in observed NO₂ signal between the two channels provided a quantitative ClNO₂ measurement. However, its use for conducting field measurements was reported to be limited, as the thermal degradation of alkyl nitrates (i.e. PAN) into NO₂ cannot be distinguished from NO₂ originating from ClNO₂ due to overlapping thermal dissociation profiles.

For this same thermal dissociation setup, product chlorine radicals will react quickly (e.g. Cl radical lifetime of 0.2 s for typical CH₄ mixing ratios of 2 ppmv and $k_{298} = 1 \times 10^{-13} \text{ cm}^3 \text{ molec.}^{-1} \text{ s}^{-1}$; Bryukov et al., 2002) with ambient hydrocarbons (e.g. methane) to form hydrogen chloride (HCl), which is a stable reservoir species for reactive chlorine (Reaction R2).

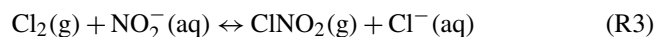


Several optical methods for the high-frequency and precise detection of HCl have recently been reported that overcome historical challenges with its sampling (Furlani et al., 2021; Hagen et al., 2014; Halfacre et al., 2023; Wilkerson et al., 2021), making them attractive candidates for an alternative thermal dissociation approach for the detection of ClNO₂. In this work, we demonstrate the coupling of a thermal dissociation furnace to HCl-TILDAS (TD-TILDAS) for quantitative detection of ClNO₂ as HCl. Compared with CIMS, TD-TILDAS is a lower-time-cost [CE2](#) method for determining ClNO₂ mixing ratios, involving less experimental calibration work and simpler data processing as a direct method.

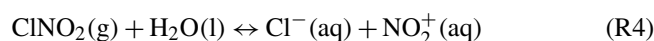
2 Methods

2.1 ClNO₂ generation

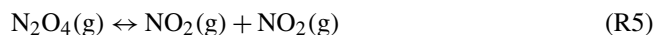
ClNO₂ was synthesized by flowing Cl₂ across a nitrite-rich slurry, as described by Thaler et al. (2011) and shown by Reaction (R3).



However, it is believed the ClNO₂, once produced, may react further by dissolving into the water, hydrolyzing, and producing nitronium and chloride ions (Reaction R4) (Frenzel et al., 1998).



The nitronium ion can then react with NO₂[−] to produce N₂O₄, which exists in equilibrium with NO₂ (Reaction R5).



As detailed by Thaler et al. (2011), this chemistry can be mitigated by minimizing the residence time of ClNO₂ in the reaction vessel and, to a lesser extent, by increasing the Cl[−] content of the slurry to encourage the equilibrium in Reaction (R4) towards ClNO₂. Therefore, we composed our slurry using sodium chloride (> 99.5 % pure, BioXtra, Sigma Aldrich product no. S7653-5KG, USA) and sodium nitrite (99 %, extra pure, Acros Organics Code 196620010, Belgium) at a mole ratio of 100 : 1 Cl[−]:NO₂[−], wetting with 18 MΩ deionized water (Millipore). The slurry was housed in ~ 10 cm of 1.25 cm diameter PFA tubing. Varied flow rates (0.5–5 mL min^{−1}) of 10 ppmv Cl₂ (diluted in nitrogen, BOC product no. 150916-AV-B, United Kingdom) were injected into a dilution flow (ranging from 200–2499.5 mL min^{−1}) of NO_x-scrubbed compressed air (using trap composed of 50 % Sofnofil from Molecular Products Ltd., Essex, United Kingdom, and 50 % activated carbon) that was subsequently passed over the slurry, generating ClNO₂. A portion of the dilution flow was directed into a bubbler containing 18 MΩ deionized water prior to entering the slurry to maintain a humid environment and prevent the slurry from drying out. A schematic diagram of this setup is presented in Fig. 1a.

2.2 TD-TILDAS

The TILDAS instrument and operation technique have been well-described previously (McManus et al., 2011, 2015). HCl-TILDAS was developed by Aerodyne Research, Inc. and characterized by Halfacre et al. (2023). Briefly, air is sampled at 3.0 L min^{−1} through a heated (50 °C) quartz “inertial inlet”, which is a type of virtual impactor used to remove particles > 300 nm from the sample matrix. Sample air continues its flow through 3 m of heated (50 °C) tubing into the Herriott cell (204 m path length) inside the TILDAS. HCl is then detected via a mid-IR inter-band cascade laser that probes the strong R(1) H³⁵Cl absorption line at 2925.89645 cm^{−1} within the (1-0) rovibrational absorption band (Guelachvili et al., 1981).

Nitryl chloride was converted to HCl for detection by TILDAS via thermal dissociation and the subsequent reaction of Cl radicals with hydrocarbons, namely methane (Reactions R1–R2) (Thaler et al., 2011). While modelling results predicted that ambient mixing ratios of methane (~ 2 ppmv) are sufficient for achieving unit conversion of 1 ppbv ClNO₂ to HCl (Sect. 3.1), the sample flow was additionally spiked with propane (BOC Limited, product no. 34-A) to a mixing ratio of 5 ppmv to both ensure reaction completeness and outcompete Cl wall losses, as the rate constant for the reaction between Cl and propane is 3 orders of magnitude faster than with methane (Atkinson et al., 2006a). Next, the sample was directed to a 90 cm length of quartz tubing (9.5 mm o.d.,

7.5 mm i.d.) housed within a furnace (Carbolite Gero TS1 12/60/450) upstream of the inertial inlet. 60 cm of this tubing is held within the heated region of the furnace, resulting in a residence time of ~ 500 ms under a flow rate of 3 L min^{−1}. The internal temperature of the furnace is monitored using the furnace’s inbuilt temperature sensors and logged using the furnace software. To mitigate HCl surface interactions after ClNO₂ conversion, perfluorobutane-1-sulfonic acid (PFBS; Merck, product no. 562629, United Kingdom) vapour was introduced just after the furnace to actively passivate tubing and inlet surfaces, improving HCl transmission to the TILDAS inlet by (1) displacing HCl sorbed to surfaces and (2) increasing the non-polar character of surfaces by presenting a fluorinated chain to passing analytes (Halfacre et al., 2023; Roscioli et al., 2016). As detailed by Halfacre et al. (2023), a flow (50–75 mL min^{−1}) of oxygen-free nitrogen was flowed into the headspace of a Teflon bubbler containing 5 g of PFBS, thereby flushing the PFBS vapour into the sample line. A schematic diagram of this setup is presented in Fig. 1b.

The major sources of uncertainty with using TD-TILDAS to detect ClNO₂ include the degree of ClNO₂ conversion to HCl, instrument noise, background drifts, and potential line losses of HCl. Unit conversion of ClNO₂ to HCl was confirmed by modelling and laboratory experiments (see Sect. 3.1 and 3.2). Instrument noise and background drifts were assessed regularly from blanks. For laboratory experiments, blanks were performed by sampling the ClNO₂ standard (Sect. 2.1) diluted in NO_x-scrubbed compressed air through the unheated furnace. This dilution air was generated using an air compressor and dehumidifying system (dew point approximately −60 °C, absolute water vapour concentration ~ 0.01 %). To vary sample humidity, carrier gas flow was split such that varied amounts were passed through a bubbler containing deionized water. Concerning line losses of HCl, the only source of HCl will be from ClNO₂ conversion during laboratory experiments, and therefore line losses were assessed between the furnace and the inertial inlet. As detailed in Fig. 1b, 30 mL min^{−1} of flow from a homemade HCl permeation source (Furlani et al., 2021; Halfacre et al., 2023) was injected alternately before the furnace and just before the inertial inlet to determine loss of HCl over this region. So long as unit conversion of ClNO₂ to HCl can be confirmed and blank/line losses are corrected, this method will be as accurate as the TILDAS is for detecting HCl, which was previously found to be within the 5 % tolerance of a commercial HCl cylinder with a certified concentration (Halfacre et al., 2023).

For ambient sampling (Fig. 1c), an additional 5 m of 1.25 cm o.d. PTFE Teflon was added before the tee that splits the CIMS and TILDAS flow paths to sample outside air. A 5 µm PFA Teflon filter was also installed to collect particulates, reducing the potential for HCl displacement through thermodynamic partitioning of particulate Cl[−] that would otherwise enter the heated furnace (Huffman et al., 2009).

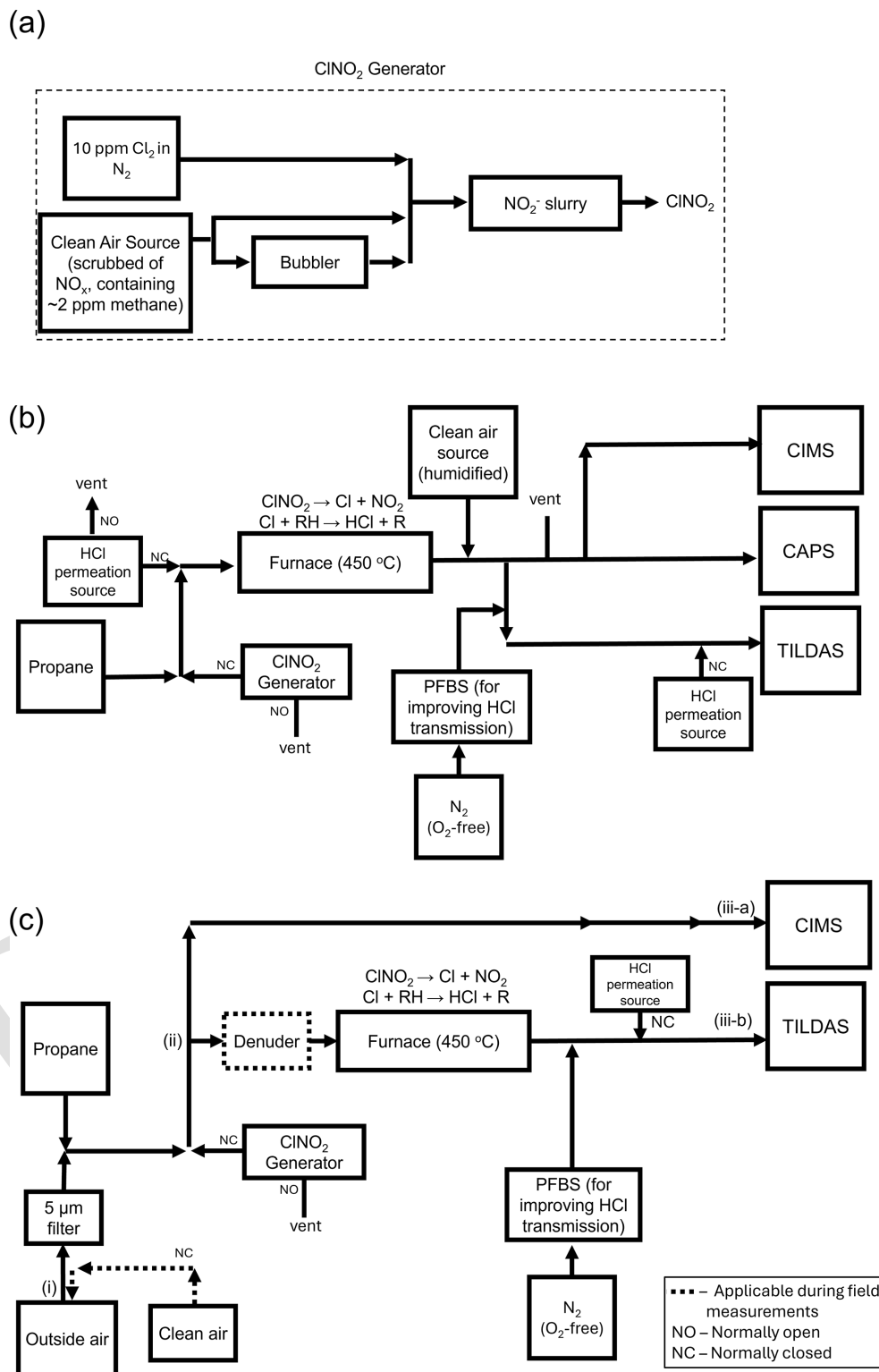


Figure 1. Experimental schematic diagrams for (a) generating ClNO₂, (b) laboratory comparison measurements between CIMS, TILDAS, and CAPS NO₂, and (c) calibration/field sampling between CIMS and TILDAS. Note that “NO” stands for “normally open” and “NC” stands for “normally closed” in reference to solenoid valves that control the flow direction for these items. For panel (c), the approximate distance between point (i) and (ii) is 5 m, from (ii) to (iii-a) is 1.5 m, and from ii to (iii-b) is 2 m.

This Teflon filter was not found to affect the observed mixing ratio of our ClNO₂ standard in measurement comparisons with and without the filter. However, the collection of particulates on the filter could enable heterogeneous chemistry with passing N₂O₅ plumes that may produce corresponding ClNO₂ plumes that are not reflective of ambient chemistry, so frequent replacement of these filters is necessary (e.g. daily). Blank air was generated by pumping ambient air through a 50 % activated carbon/50 % Sofnofil scrubber, which was found to effectively remove ClNO₂ from the sample stream. The pump (KNF model N035.1.2AN.18) was able to overblow the sample inlet at a flow rate of $\sim 25 \text{ L min}^{-1}$. This approach is favoured over the use of synthetic cylinder air as significant changes in sample humidity can result in release of HCl from surfaces (Halfacre et al., 2023). Blanks were performed for 10 min every 30 min to ensure the instrument had enough time to respond and adjust to a stable background value. Additionally, ambient measurements will include HCl, which would act as an interference for ClNO₂ observations. To obviate this, a denuder (coating of 2 % Na₂CO₃ and 2 % glycerol dissolved in 50 % water and 50 % methanol) was installed before the furnace to selectively remove acidic gases (e.g. HCl, HNO₃) that may influence quantitation. Using the denuder for this purpose was found to be effective for at least 1-week periods, after which it was generally replaced to avoid coating exhaustion. The denuder was also found to affect ClNO₂ throughput on shorter-term timescales (e.g. daily), with a freshly coated denuder causing as much as 55 % loss of the ClNO₂ standard mixing ratio. This was determined by calculating the percent difference when sampling the ClNO₂ standard both through and bypassing the denuder. Because ClNO₂ additions during ambient sampling will always be added through the denuder, it was important that the ClNO₂ standard (Sect. 2.1) was sampled in dry air before and after overnight experiments to quantify how this loss evolved over the course of an experiment such that data could be corrected using the percent difference term. Periodic additions of HCl standard were also performed to assess line losses of HCl after conversion in the furnace. In contrast to the laboratory experiment configuration, permeation source HCl in blank air was only injected just downstream of the furnace mid-experiment to reduce exposure of unpassivated sampling surfaces to HCl. Losses were assessed by comparing this observed HCl injection value to pre- and post-experiment injections over dry compressed air. Injections of HCl and ClNO₂ standards was controlled using three-way Teflon solenoid valves (Master-Flex model no. 01540-18, Cole Parmer, United Kingdom).

2.3 Supporting instrumentation

To confirm the efficacy of TD-TILDAS as a valid quantitative method for ClNO₂ detection, testing was performed simultaneously with a cavity-attenuated phase shift (CAPS) NO₂ instrument (Sect. 2.3.1) and time-of-flight chemical ionization

mass spectrometer (Sect. 2.3.2), both of which have previously been reported as ClNO₂ detection methods.

2.3.1 Cavity-attenuated phase shift (CAPS) NO₂

ClNO₂ mixing ratios observed by the TILDAS were confirmed via simultaneous detection of the NO₂ product of ClNO₂ thermal dissociation using a commercial cavity-attenuated phase shift NO₂ detector (Teledyne T500U CAPS). Briefly, emission from an LED (emission centred around 425 nm) is reflected across two spherical mirrors and absorbed by NO₂ in the optical cell. This difference in light is detected by a photodiode and quantified based on its absorbance via the Beer–Lambert law. The instrument was calibrated using gas-phase titration of NO by O₃ to produce varied concentrations of NO₂. A 1 ppm NO in nitrogen cylinder (certified 982 ppb, NPL) was used to verify the concentration of NO in a 25 ppm NO in nitrogen working standard (BOC). A multigas blender (EnviroNics S6100) was used to generate a range of O₃ concentrations (range 0–130 ppbv) for titrating some of the NO (NO in excess, 200 ppbv) into NO₂, and the decrease in the NO concentrations was measured using a calibrated NO_x instrument (Teledyne API Chemiluminescence T200). The NO₂ introduced to the CAPS instrument is thus the sum of the drop in NO from the added ozone and the NO₂ already present in the working standard. The T200 NO_x instrument was also used to measure ambient air alongside the CAPS (range 0–25 ppbv), and these data are presented in Fig. A1. Additionally, the Teledyne T500U includes an internal drying assembly and has a manufacturer-recommended humidity range of 0 %–95 %.

2.3.2 Time-of-flight chemical ionization mass spectrometry (CIMS)

ClNO₂ was additionally detected using a VOCUS high-resolution chemical ionization time-of-flight CIMS (ToFwerk, Switzerland) with a VOCUS AIM reactor and using iodide (I[−]) as a reagent ion gas. A complete description of this instrument and its operational principles can be found in Riva et al. (2024). Briefly, sample gas is drawn into the sampling inlet and pulled through a critical orifice (0.475 mm) and PFA Teflon sample flow guide into a conical ion molecule reactor (IMR) at a flow rate of 1.8 L min^{-1} . The IMR was held at a constant pressure of 50 mbar using a vacuum pump (IDP3, Agilent Technologies) and temperature-controlled to 50 °C. The reagent ion source was a permeation tube containing trace amounts of CH₃I dissolved in benzene (ToFwerk). Ultrahigh-purity, oxygen-free N₂ gas (generated by flowing compressed air through gas with a commercial N₂ generator, Infinity NM32L, Peak Scientific Instruments, UK) is continually flowed over the permeation tube to flush the gaseous CH₃I/benzene mixture into a compact vacuum ultraviolet ion source (VUV). Within the VUV, UV light emitted from a Kr lamp (116.486 and 123.582 nm) is absorbed by benzene,

generating low-energy photoelectrons that can react with CH₃I to produce I[−] (Ji et al., 2020). The I[−] reacts with analytes for approximately 30 ms before being drawn through another critical orifice where the sample travels through four differentially pumped chambers before reaching the drift region of the ToF-CIMS. Ions in the ToF chamber are extracted and converted into mass spectra via an MCP detector with a preamplifier over a mass range of 7–510 Th. The extracted packets are averaged over a period of 1 s and the resolution of the instrument is ≈ 5000 . Data were collected at a rate of 1 Hz. Data averaging, mass calibration, peak assignment, peak fitting, and peak integration are all performed using the software package Tofware (version 4.0.0, TOFWERK) used in Igor Pro 9 software (Wavemetrics). Peak fitting focused on I³⁵ClNO₂[−] (m/z 207.8668) and I³⁷ClNO₂[−] (m/z 209.8638), and isotope abundances were manually confirmed to be $\sim 1:0.32$ based on the natural abundance of chlorine isotopes. CIMS signals were normalized against the sum of the total number of reagent ions, which is equivalent to I[−] + I(H₂O)[−]. Additionally, as the CIMS sensitivity to ClNO₂ varies with humidity in the ion molecule reactor region, we define an additional term equal to the ratio of the iodide water cluster (I(H₂O)[−]) to the reagent ion sum (I[−] + I(H₂O)[−]), hereafter referred to as the iodide water ratio (IWR). Instrument backgrounds were assessed using air scrubbed of ClNO₂, as described in Sect. 2.2.

2.4 Data analysis

Data analysis was conducted using the R language for statistical computing (R Core Team, 2021). Linear regressions were calculated using the York method (Cantrell, 2008) when possible so as to incorporate uncertainties in compared variables.

2.5 Chemical modelling

The 0-D box model Kintecus (Ianni, 2003, 2022) was used to explore the gas-phase chemistry occurring in the heated furnace to predict the timescales of the thermal dissociation of ClNO₂ and the subsequent formation of HCl after reaction with hydrocarbons (Reaction R2). The only hydrocarbon included in these model experiments was methane. The model was also used to identify potential interferents that could prevent unit conversion of ClNO₂ to HCl. The results of the model were used to guide the experimental setup. The modelled species, reaction list, tested interferents (including ClNO and alkenes), and initial concentrations are included in Appendix A (Tables A1–A3). Reaction kinetics were sourced from the NIST Chemical Kinetics Database and IUPAC Evaluated Kinetic Data websites (Manion et al., 2015; Wallington et al., 2021), and primary literature references are listed next to each reaction. No chemical species were held constant or were otherwise constrained outside of initial concentrations. The model integration time was set to

1 ms, and the entire simulation was set to last 150 ms. The model initiated with a temperature of 25 °C (held for 10 ms) before increasing to 450 °C over the course of 22 ms. The temperature was held at 450 °C for 40 ms, before gradually decaying back to 25 °C over 70 ms.

3 Results and discussion

3.1 Modelling TD chemistry

Box model simulations predicted the rapid, virtually CE3 unit conversion of ClNO₂ to HCl after increasing temperature to 450 °C (Fig. 2) under the model conditions outlined in Tables A1–A3. 90 % conversion was calculated to occur within 23 ms from a starting ClNO₂ concentration of 2.46×10^{10} molec. cm^{−3} (1 ppbv at 25 °C), and ambient mixing ratios of methane (i.e. 2 ppmv at 25 °C) were found to be sufficient for facilitating this chemistry. While Cl-mediated hydrocarbon oxidation was shown to produce a modest enhancement of hydroxyl radical concentrations (Fig. 2b), it was not enough to compete meaningfully with Cl to mitigate or retard Reaction (R1). Similarly, an initial O₃ concentration of 9.84×10^{11} molec. cm^{−3} (40 ppbv at 25 °C) did not significantly inhibit the desired chemistry by the direct reaction of O₃ with Cl radicals. Concerning potential interferents, Cl can add to double bonds found on alkenes without producing HCl. Reactions with ethene, propene, and isoprene were included in the model at 1.23×10^{12} molec. cm^{−3} (50 ppbv at 25 °C) each and were found to produce approximately 1×10^6 molec. cm^{−3} of non-HCl product, which is 4 orders of magnitude less than the HCl converted from ClNO₂. As these mixing ratios of alkenes are larger than those typically found in real-world environments (e.g. Budisulistiorini et al., 2015; Hellén et al., 2024; Tripathi et al., 2021), it is unlikely that alkenes will cause meaningful interference for ClNO₂ quantification.

ClNO₂ was predicted to be the only known inorganic chlorine reservoir to thermally dissociate at 450 °C. This is consistent with the relative bond dissociation energies found for ClNO₂ (142 kJ mol^{−1}) relative to the various other forms of inorganic chlorine simulated (Cl–NO₂ < Cl–Cl < Cl–O < Cl–R < Cl–H) (Darwent, 1970). Production of other inorganic chlorine compounds (e.g. Cl₂, HOCl, ClONO₂, or reformation of ClNO₂) was orders of magnitude less than the resulting HCl and is therefore not believed to influence HCl production. Even so, there are potential inorganic chlorine species that may still act as interferents in this method. Nitrosyl chloride (ClNO) has been previously predicted by modelling to exist at ppbv-level mixing ratios in polluted marine environments and could be an efficient Cl atom source (Raff et al., 2009). Indeed, 1 ppbv (2.46×10^{10} molec. cm^{−3}) of ClNO was found to partially thermally dissociate in our Kintecus model (bond dissociation energy of 159 kJ mol^{−1}) and generate additional HCl, as well as NO that was grad-

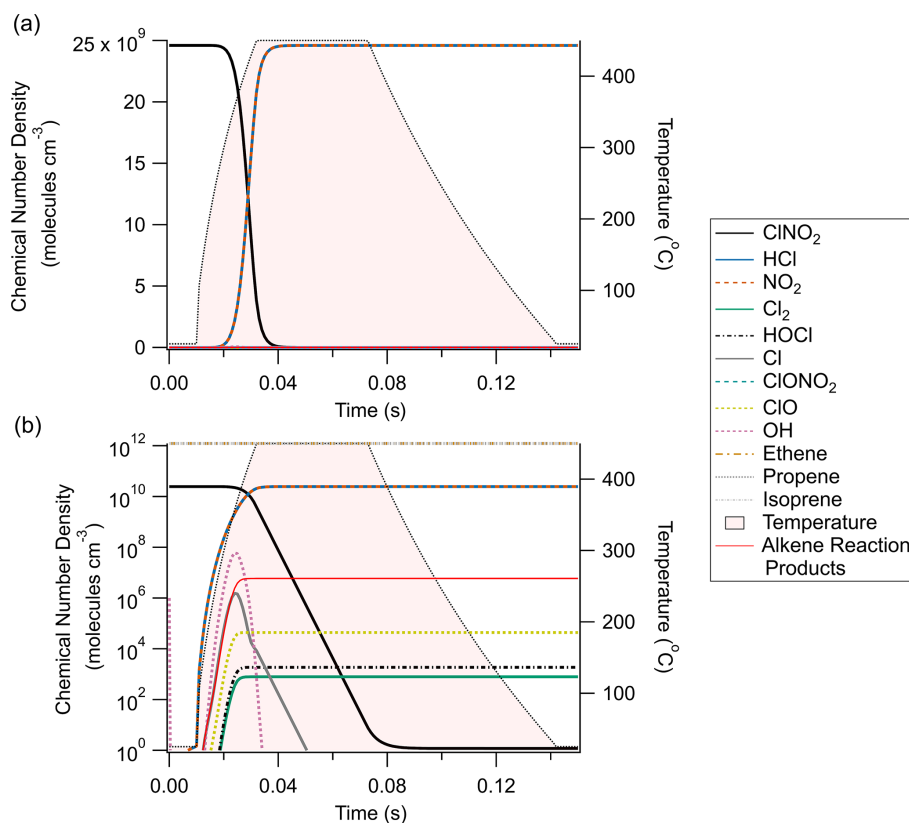


Figure 2. Chemical modelling results of the thermal dissociation of ClNO₂ and its subsequent conversion to HCl. Panel (a) presents results on a linear y axis, while panel (b) features the same data on a logarithmic y axis. Note that ethene, propene, and isoprene are off-scale in panel (a) (1.23×10^{12} molec. cm⁻³ \sim 50 ppbv) to better display the relationships between ClNO₂, HCl, and NO₂ and are shown to remain constant in panel (b).

ually converted to NO₂ (Fig. A2). On the addition of heat, ClNO decreased by 40 %, while HCl increased by an equivalent amount (in addition to the 2.46×10^{10} molec. cm⁻³ generated by ClNO₂ thermal dissociation). While we are unaware of any field measurements that have confirmed the presence of ClNO in the boundary layer to date, it appears likely this method would be sensitive to interference from ClNO if/where its presence is confirmed.

Additionally, one notable class of compounds that could not be simulated were chloramines, which have recently received increased attention as relevant daytime sources of Cl atoms (Angelucci et al., 2023; Wang et al., 2023). Their largest known anthropogenic sources include water disinfection processes, swimming pools, and cleaning products. Trichloramine, dichloramine, and monochloramine have reported bond dissociation energies of 381, 280, and 251 kJ mol⁻¹, respectively (Darwent, 1970) (ClNO₂ bond dissociation energy = 142 kJ mol⁻¹) and would therefore not be expected to produce free Cl radicals in the temperature range simulated herein if its thermochemistry is consistent with the above bond dissociation energy trend. However, to the authors' knowledge no information is available regarding their thermal stability in the gas phase at atmospheric

cally relevant conditions, and this potential source of positive interference for our proposed method cannot be discounted via the model at this time. Similarly, prevalent organochlorides, such as methyl chloride (CH₃Cl), dichloromethane (CH₂Cl₂), chloroform (CHCl₃), and carbon tetrachloride (CCl₄), could cause positive interference if they dissociate and produce Cl atoms in the furnace (World Meteorological Organization, 2022). Global average mixing ratios for CH₃Cl, CH₂Cl₂, CHCl₃, and CCl₄ were \sim 550, \sim 40, 9, and 77 pptv, respectively, during 2020. Appropriate thermal dissociation kinetic parameters could not be sourced for the conditions used herein (i.e. temperatures \leq 450 °C), so these compounds could not be properly simulated by the Kintecus model. Similarly to the chloramines, the bond dissociation energies are much higher than other compounds simulated (339, 310, 346, and 293 kJ mol⁻¹ for CH₃Cl, CH₂Cl₂, CHCl₃, and CCl₄, respectively; Darwent, 1970; Weissman and Benson, 1983).

3.2 Laboratory characterization of TD-TILDAS

For laboratory characterization, a stable source of ClNO₂ was generated (Sect. 2.1) for assessing TD-TILDAS perfor-

mance in comparison with other established ClNO₂ sampling techniques, including CAPS NO₂ and CIMS (Sect. 2.3). One key change between model simulations and this experimental setup is the inclusion of propane in the sample stream (estimated mixing ratio of 5 ppmv within the heated section of sample configuration). While the model predicted the pertinent chemistry will occur in ~ 23 ms using only ambient methane as the hydrocarbon (Sect. 3.1) and the residence time in the heated furnace is ~ 500 ms, adding propane ensures complete conversion of ClNO₂ to HCl and ensures wall losses are negligible, as Cl radicals react with propane approximately 3 orders of magnitude faster than with methane (Atkinson et al., 2006a). The fact that no additional HCl signal was observed on addition of propane at varying levels (not shown) supports our calculations that unit conversion is achieved and competitive loss of Cl radicals to walls is negligible.

A schematic diagram of these experiments is shown in Fig. 1a. Figure 3 represents a typical comparison experiment in which ClNO₂ was sampled by all three instruments simultaneously. First, ClNO₂ was introduced into the flow stream with the furnace unheated, yielding a positive CIMS signal for ICINO₂[−] (~ 1100 ncps for the example in Fig. 3), while TILDAS HCl and CAPS NO₂ mixing ratios remained at background levels. As the furnace temperature approached 450 °C, Reactions (R1)–(R2) began to occur. HCl and NO₂ mixing ratios rose, plateauing at similar values (~ 2.2 ppbv in Fig. 3), while ICINO₂[−] decreased to the instrument baseline, implying that both Reactions (R1)–(R2) proceeded to completion. Signals returned to their original positions once the furnace was allowed to cool back to room temperature (e.g. from 16:45 UTC in Fig. 3). Note that the HCl signal spike during the furnace's temperature ramp was seen consistently across experiments and was most likely caused by a shift in HCl molecule partitioning between the surface of the quartz tubing toward the gas phase (Halfacre et al., 2023). Allan–Werle deviation calculations demonstrate favourable performance metrics for TILDAS while sampling ClNO₂, with 1 Hz precision of 11.8 pptv and as good as 1.2 pptv with an integration time of 96 s (Fig. 4).

A summary of comparison experiments across varied humidities is presented in Fig. 5. The changes in HCl as observed by TILDAS correlated strongly with the changes in NO₂ observed by the CAPS instrument (Pearson correlation coefficients of 0.999, 0.997, and 0.987 for relative humidities of 11 %, 44 %, and 66 %, respectively). However, the slopes were consistently less than unity (0.95 ± 0.01 , 0.93 ± 0.02 , and 0.91 ± 0.02 at 11 %, 44 %, and 66 %, respectively), indicating observed HCl mixing ratios were less than corresponding NO₂ mixing ratios. One potential explanation for this could be loss of Cl radicals in the furnace, but we do not believe this to be the case (as detailed above). While physical losses of HCl to sampling lines would not be unexpected as HCl has a high affinity for sorbing to physical surfaces, experiments were designed to minimize these

interactions, and line loss experiments were performed to quantify any losses observed at tested humidities. Experimentally, a small flow ($50\text{--}75\text{ mL min}^{-1}$) of PFBS vapour was injected into the TILDAS sampling line downstream of the furnace to reduce HCl affinity for surfaces (Sect. 2.2) (note that PFBS was not introduced to the entirety of the flow path to avoid sampling of PFBS by other instruments; additionally, there is evidence that PFBS degrades at temperatures above 400 °C as in Xiao et al., 2020, so its ultimate efficacy and reproducibility within the furnace system would be uncertain). Further, the high operating temperature of the furnace would also be expected to minimize HCl–wall interactions within the quartz tubing. Indeed, no line losses were found at 11 % relative humidity between when the HCl permeation source standard was injected into the sampling line before the heated furnace (2.95 ± 0.02 ppbv) and when HCl was injected just before the inertial inlet (accounting for dilution factors) (2.95 ± 0.02 ppbv), consistent with Halfacre et al. (2023). Similar results were found at 44 % relative humidity (pre-furnace value of 2.68 ± 0.03 ppbv vs. 2.66 ± 0.03 ppbv when HCl was introduced at inlet), and real HCl loss was quantified at 66 % relative humidity (pre-furnace value of 1.87 ± 0.03 ppbv vs. 1.97 ± 0.03 ppbv when HCl introduced at inlet). Having accounted for these line losses, ANOVA calculations found no significant differences between these three slopes as presented in Fig. 3 ($F(2, 19) = 0.10$, $p = 0.902$), indicating consistent performance between TILDAS and CAPS for detecting ClNO₂. However, it does not appear to explain the deviation from unity, which will be discussed below.

As discussed in Sect. 2.1, chemistry may occur within the slurry to produce N₂O₄, which can easily degrade at room temperature to produce two NO₂ molecules. If the N₂O₄ output from the NO₂[−] / Cl[−] slurry is constant over the timescale of an experiment (< 1 h), it would be expected that this additional NO₂ is readily accounted for during blank subtraction calculations. While we believe this is largely true for the experiments presented above, discrepancies in ClNO₂ signals were observed as the slurry aged ($> \sim 3$ weeks), with CAPS-observed NO₂ mixing ratios growing in significant excess of TILDAS-observed HCl mixing ratios (Fig. A3). Separate applications of TILDAS- and CAPS-based calibration factors (using data from Fig. 5) to concurrent CIMS ClNO₂ observations show closer resemblance to the TILDAS-observed mixing ratios (Fig. A3), suggesting that additional chemistry may be occurring within the salt bed that produces stable reservoirs of NO₂ that thermally dissociate in the furnace to produce undesired NO₂. This NO₂ artefact serves as a likely explanation for the sub-unity slopes presented in Fig. 5, as it would positively bias the CAPS measurements but not the TILDAS, which is only sensitive to HCl. Thaler et al. (2011) present in great detail strategies for minimizing N₂O₄ production in their study by minimizing the residence time in their ClNO₂ generator (0.3 s herein) and adjusting the molar ratio of Cl[−] : NO₂[−] of their salt bed (100 : 1 herein), but they

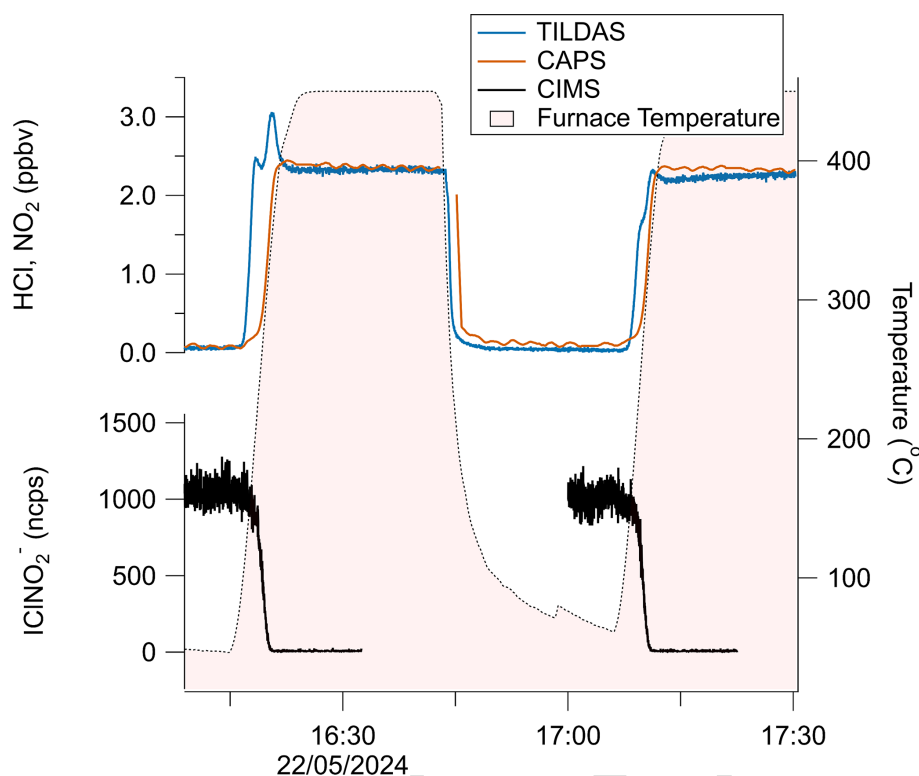


Figure 3. Time series demonstrating the reversible thermal conversion of ClNO₂ to NO₂ (red trace, CAPS) and HCl (blue trace, TILDAS), as evidenced by changes in CIMS-observed ICINO₂⁻ (black). Gaps in CIMS data are from internal CIMS tests not pertinent to this work.

were ultimately unable to completely eliminate it; while we found these strategies helpful for reducing the overall NO₂ background as measured by CAPS, we found they were unsuccessful in eliminating the artefact when sample gas was passed through the heated furnace. We are not aware of such chemistry being addressed in the literature for this ClNO₂ generation method and do not propose potential reactions as it is outside the scope of this paper.

Both HCl and NO₂ mixing ratios independently correlated strongly with the CIMS measurement of ICINO₂⁻ (Fig. 5b, c), and the I⁻ CIMS sensitivity for ICINO₂⁻ was found to vary strongly with humidity, as previously reported (Kercher et al., 2009; Mielke et al., 2011). The weakest Pearson correlation coefficient was for NO₂ and ICINO₂⁻ at 66 % relative humidity ($r = 0.988$), virtually matching that of NO₂⁻ and HCl at the same humidity. Due to the uncertainty/unreliability of the NO₂ as it relates to ClNO₂ quantitation, we do not further consider the relationship between CAPS and CIMS.

The linear equations from Fig. 5a present significant intercepts that suggest a source of positive error for the TILDAS, and the similarity of these intercepts suggests a relatively constant/consistent source (values are statistically the same: $F(4, 19) = 0.624$, $p = 0.546$ per ANOVA). For these experiments, TILDAS blanks were obtained by sampling slurry air flowed through an unheated furnace; in this scenario, Reactions (R1)–(R2) are unable to occur, and therefore any sig-

nal observed by TILDAS could be considered background. It is possible that a small amount of HCl forms in the slurry system from the aqueous disproportion reaction between Cl₂ and H₂O. When the furnace is unheated, some amount of HCl interaction with the quartz tubing is expected given that there is no PFBS flow through this portion of the plumbing, biasing this blank measurement low. Then, once the furnace is heated to 450 °C, this HCl will be liberated from the quartz tubing, possibly then biasing the heated measurement high. This is supported by the presence of a peak in observed HCl as the furnace reheats (e.g. as in the second temperature ramp in Fig. 3), as some of the HCl sorbed to the furnace tube walls under room temperature is forced into the gas phase. The statistical similarity in intercepts implies that this effect is constant across these experiments, leading to a consistent offset. While an ideal blank would sample the gas downstream of the slurry while selectively scrubbing ClNO₂, this was not practical to achieve without simultaneously scrubbing HCl. Therefore, we propose that the y intercept in these cases is a good estimate of the systematic error present in these comparison experiments.

3.3 Applicability as a field instrument

The applicability of TD-TILDAS as a field method for ClNO₂ detection was tested by sampling ambient air from

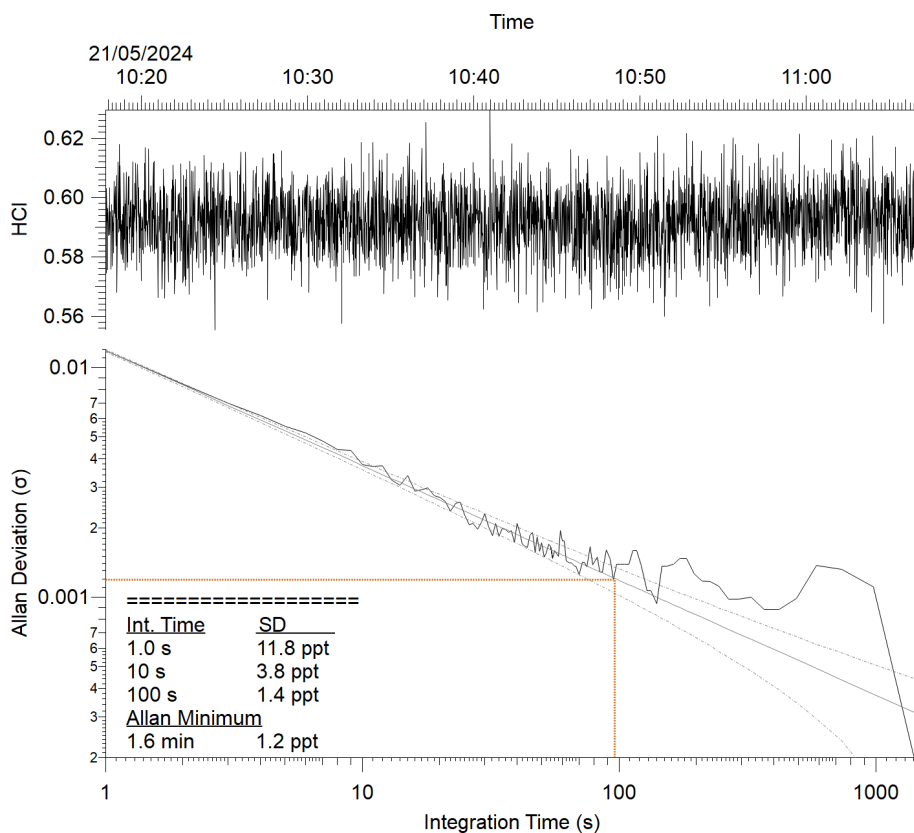


Figure 4. Allan–Werle plot for TD-TILDAS during addition of ClNO₂ standard into the sample line. The Allan minimum is indicated by the dotted red lines.

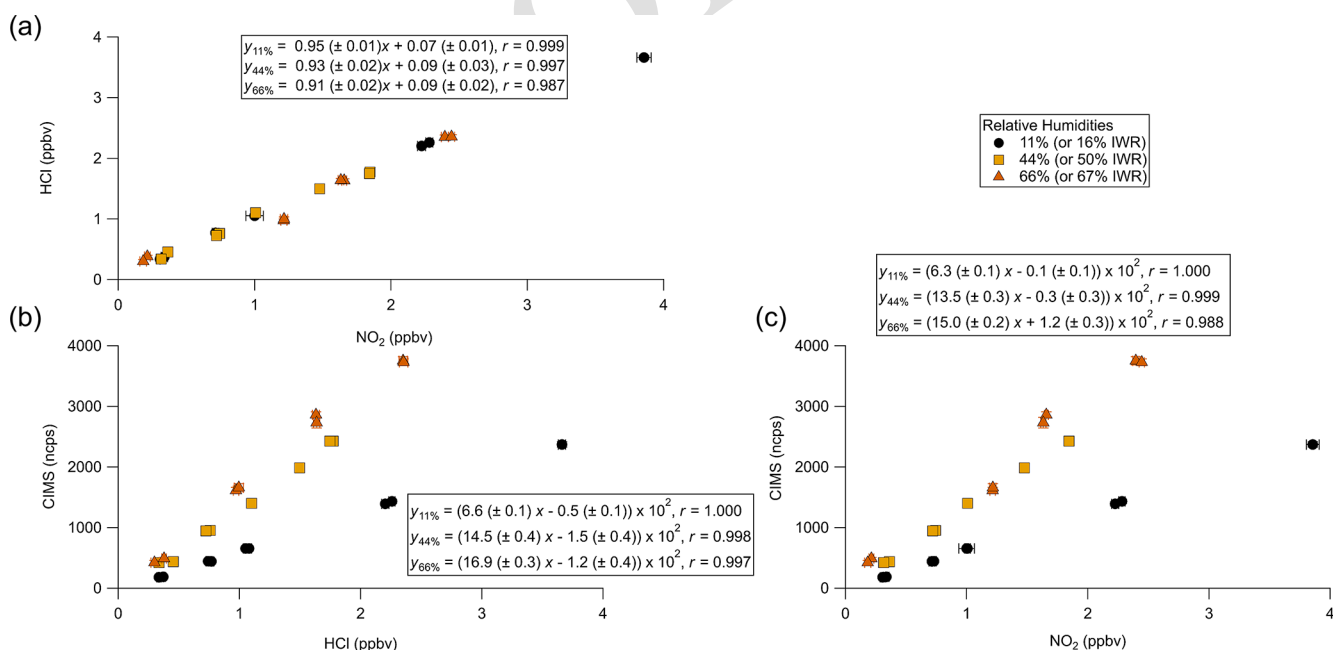


Figure 5. Comparison curves of (a) TILDAS vs. CAPS, (b) CIMS vs. TILDAS, and (c) CIMS vs. CAPS for injections of varied mixing ratios of ClNO₂ across different relative humidities. Regressions involving TILDAS data have been corrected for line losses observed at 66 % relative humidity.

outside the Wolfson Atmospheric Chemistry Laboratory building on the University of York campus (York, United Kingdom) on the morning of 13 January 2025 (Fig. 6). Compared with the laboratory-based configuration described in Sect. 3.2, ambient air will contain varied amounts of HCl that would interfere with accurate quantification of ClNO₂ via the TILDAS method. To address this, a base-coated denuder (Sect. 2.2) was installed in the HCl sampling line. ClNO₂ throughput was found to be hindered when flowed through the denuder but increased over the course of the observation period (pre-experiment estimation of 55 % loss on 10 January vs. 31 % measured directly after the experiment on 13 January). This loss was accounted for by applying a time-varying, linearly interpolated correction factor for the denuder. In addition, line losses affecting HCl between the heated furnace and TILDAS inlet were estimated as 2.7 %, which was added back into the TILDAS measurements. CIMS observations of ICINO₂[−] were calibrated against TD-TILDAS using a mid-experiment ClNO₂ addition, yielding a sensitivity factor of 1982 ncps ppb^{−1} (measured with a corresponding IWR of 42 %). We note that this factor is ∼ 35 % greater than the value of 1450 ncps ppbv^{−1} as presented in Fig. 5b for a comparable IWR (44 %); this is likely due to the replacement of the reagent ion permeation source, repair of reagent ion source heaters, and change in the overall sampling configuration between the experiments from Sect. 3.2 and this section (as illustrated by Fig. 1). Application of this sensitivity factor across this measurement period can be justified as the IWR was found to be stable (38 ± 2 %). Limits of detection, based on instrument blanks, were found to average 10 ± 5 pptv for TD-TILDAS and 1 ± 1 pptv for CIMS (using 60 s data averaging).

As seen in Fig. 6a, TILDAS-observed ClNO₂ and CIMS-observed ClNO₂ demonstrate very good agreement for these ambient observations in both signal magnitude and structure. This is quantitatively supported by regression calculations during this period that yield a slope of 0.97 ± 0.01 (Fig. 6b), which is well within the averaged combined uncertainty for this period of 9 %. While the sub-unity slope could indicate small losses with the TILDAS method, pre- and post-experimental losses were tested and corrected for as detailed above, so this is not believed to be a large source of error in this case. It is otherwise not unexpected that this slope is found to deviate from unity given the uncertainty in the application of a single-point CIMS sensitivity factor. Nevertheless, this agreement gives us confidence that it is appropriate for these measurements and provides a proof of concept for this TILDAS method.

Additional sources of measurement uncertainty include unaccounted-for thermolabile chlorine reservoirs that could cause positive interference in the TILDAS method. As stated above, the TD-TILDAS method functions on the assumption that ClNO₂ is the only major chlorine source that thermally dissociates at 450 °C. As shown by the model, ClNO may be a potential source of interference if present (Fig. A2),

while relevant thermochemistry information was unavailable for organochlorides and chloramines, which therefore cannot be ruled out as possible interferences by modelling. Indeed, while CIMS signals of ClNO and chloramines did not rise above their baselines during the period shown in Fig. 6, a separate measurement period demonstrates multiple occurrences where signal increases in iodide trichloramine and dichloramine adducts (INCl₃[−], INHCl₂[−]) correspond to TILDAS-observed signal increases (Fig. A4). This is most dramatic at ∼ 08:15 UTC, where ∼ 115 ncps of INCl₃[−] and 18 ncps of INHCl₂[−] correspond to an increase of 100 pptv in the TILDAS signal. While these chloramine observations cannot be quantified at this time, trichloramine and dichloramine have previously been detected in downtown Toronto at ≤ 0.104 ppb and ≤ 8 ppbv, respectively (Wang et al., 2023), suggesting that a combined 100 pptv interference contribution from these compounds is realistic. Synthesis and calibration of chloramine standards is a nontrivial task (Wang et al., 2023), so further experiments are required to investigate (1) to what extent the chloramine signals can be quantified by TILDAS and (2) if the chloramine signal can be dissected from the ClNO₂ signal through temperature scans. The results of such experiments may therefore allow this method to be extended for the quantification of both chloramines and ClNO₂.

While organochlorides (e.g. CH₃Cl, CH₂Cl₂, CHCl₃, and CCl₄) were not explicitly measured during the period in Fig. 6, it would be expected that their potential interference in the TILDAS signal (if they dissociate in the furnace) would present as a slowly varying background signal that appears as an offset above a blank, given the ubiquity of these compounds and relatively long tropospheric lifetimes for CH₃Cl, CH₂Cl₂, CHCl₃, and CCl₄ of 1 year, 6 months, 6 months, and 124 years, respectively (World Meteorological Organization, 2022). Such an offset, if present, could be quantified during daytime measurements (i.e. when no ClNO₂ will be present in the boundary layer) and readily subtracted from nighttime measurements if necessary. However, the agreement between TILDAS and CIMS measurements as presented in Fig. 6 suggests that this interference is not present, providing some evidence that these organochlorides are not dissociating in the furnace.

4 Conclusions

This work demonstrates the viability of TD-TILDAS as an independent ClNO₂ detection method for performance metrics comparable to quadrupole CIMS, which are more than adequate for commonly observed mixing ratios in the boundary layer. While modern CIMS instruments can achieve lower limits of detection and higher precision, the major advantage of TD-TILDAS over CIMS is that it does not require external ClNO₂ calibration experiments, as this work demonstrates the unity conversion of ClNO₂ to HCl that is sub-

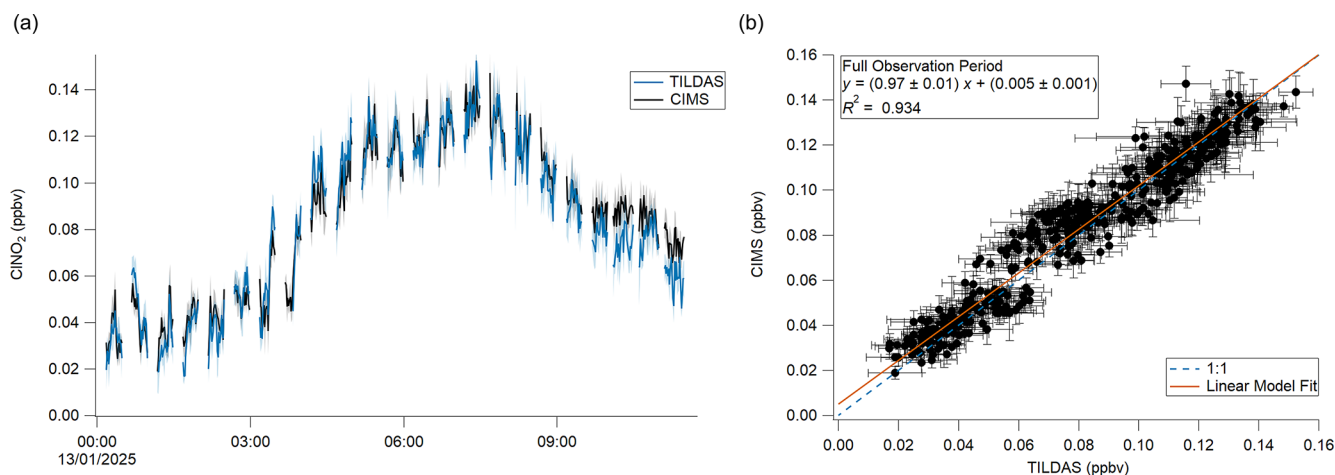


Figure 6. (a) Time series comparison of TILDAS and CIMS observations of CINO₂. (b) Scatter plot of data shown in panel (a). The error shading in panel (a) and bars in panel (b) represent the standard deviation of the 60 s averaged measurements.

sequently detected based on well-understood spectroscopic principles. The TD method described here can thus be used effectively in laboratory settings to measure CINO₂ in related experiments or even to calibrate CIMS for CINO₂ directly without needing to make assumptions regarding Cl₂ conversion on salt slurries. Additionally, use of a denuder allows this method to be readily applied to other HCl optical instruments, such as those based on CRDS.

As a field method, TD-TILDAS demonstrated excellent agreement with a co-located CIMS for CINO₂ detection. The method is reliant on accurate and regular characterization of CINO₂ throughput through the denuder, which was found to increase across 4 d of sampling. Longer-term measurement campaigns would benefit from at least weekly denuder replacements to ensure acidic gases are consistently scrubbed and do not interfere with CINO₂ observations. However, the TD-TILDAS method appears susceptible to positive interference, potentially resulting from chloramines or other unaccounted-for thermolizable chlorine compounds. Care should thus be taken should this method be deployed where large amounts of chloramines are known to be present, such as swimming pools or near water treatment facilities. More work is still required to confirm and quantify the response of this method to chloramine and organochlorides and, if so, identify an appropriate method to mitigate this potential interference. While modelling additionally suggests CINO as an interferent, its presence in the boundary layer has yet to be confirmed through in situ observations. In any case, careful temperature ramps (e.g. Day et al., 2002) performed with the furnace in environments where unknown interferences may be a concern would likely reveal the purity of the CINO₂ signal observed. Experimental adjustments could be further made for the TILDAS to alternate its sampling between a heated channel (as described in this paper) for CINO₂ detection and an unheated pathway that allows for the additional

detection of HCl. Doing so would require careful characterization of physical HCl losses inherent to both sampling pathways, as well as consideration of the likely hysteresis in detected HCl mixing ratios resulting from changes to the sampled air temperature that would affect the partitioning of HCl between surfaces and the gas phase.

Appendix A

Table A1. Bimolecular reactions and parameters used for the modelling described in Sect. 2.5. Reactions follow the rate expression $k(T) = A(T/298)^n e^{-E_a/R^T}$ (Burkholder et al., 2015).

Reaction	A	n	E _a (kJ mol ⁻¹)	Reference
ClNO ₂ + M ==> Cl + NO ₂	9.13 × 10 ⁻¹⁰	0	106	Baulch et al. (1981)
Cl + Cl ==> Cl ₂	6.15 × 10 ⁻³⁴	0	-7.53	Baulch et al. (1981)
M + ClONO ₂ ==> NO ₂ + ClO	2.76 × 10 ⁻⁵ TS1	0	94.78	Anderson and Fahey (1990)
CH ₄ + Cl ==> CH ₃ + HCl	8.24 × 10 ⁻¹³	2.49	5.06	Bryukov et al. (2002)
HCl + OH ==> H ₂ O + Cl	3.74 × 10 ⁻¹²	0	4.27	Baulch et al. (1981)
HCl + M ==> H + Cl	7.31 × 10 ⁻¹¹	0	342	Baulch et al. (1981)
CH ₃ + HCl ==> CH ₄ + Cl	3.89 × 10 ⁻¹³	0	9.64	Baulch et al. (1981)
CH ₃ + NO ₂ ==> CH ₃ O + NO	3.44 × 10 ⁻¹¹	0	0	Srinivasan et al. (2005)
O ₃ + M ==> O + O ₂	7.6 × 10 ⁻¹⁰	0	93.12	Heimerl and Coffee (1979)
CH ₃ + O ==> CH ₂ O + H	2.26 × 10 ⁻¹¹	0	0	Baulch et al. (1992)
HCl + O ==> OH + Cl	7.07 × 10 ⁻¹⁴	2.87	14.72	Mahmud et al. (1990)
OH + CH ₄ ==> CH ₃ + H ₂ O	4.16 × 10 ⁻¹³	2.18	10.24	Srinivasan et al. (2005)
Cl ₂ + M ==> Cl + Cl	3.85 × 10 ⁻¹¹	0	196	Baulch et al. (1981)
Cl + Cl ==> Cl ₂	6.15 × 10 ⁻³⁴	0	-7.53	Baulch et al. (1981)
Cl ₂ + O ==> ClO + Cl	4.17 × 10 ⁻¹²	0	11.39	Baulch et al. (1981)
Cl ₂ + H ==> HCl + Cl	1.43 × 10 ⁻¹⁰	0	4.91	Baulch et al. (1981)
Cl ₂ + OH ==> HOCl + Cl	3.60 × 10 ⁻¹²	0	9.98	Atkinson et al. (2007)
CH ₃ + O ₂ ==> CH ₃ O + O	2.19 × 10 ⁻¹⁰	0	131	Baulch et al. (1992)
ClO + O ==> O ₂ + Cl	2.50 × 10 ⁻¹¹	0	-0.91	Atkinson et al. (2007)
OH + ClO ==> HO ₂ + Cl	6.86 × 10 ⁻¹²	0	-2.49	Atkinson et al. (2007)
OH + ClO ==> HCl + O ₂	4.38 × 10 ⁻¹³	0	-2.49	Atkinson et al. (2007)
CH ₃ O + NO ==> CH ₂ O + HNO	4.00 × 10 ⁻¹²	-0.7	0	Atkinson et al. (1992)
CH ₃ O + O ₂ ==> CH ₂ O + HO ₂	7.20 × 10 ⁻¹⁴	0	8.98	Atkinson et al. (1992)
HOCl + O ==> OH + ClO	1.70 × 10 ⁻¹³	0	0	Atkinson et al. (2007)
ClCO + M ==> CO + Cl	4.10 × 10 ⁻¹⁰	0	24.6	Atkinson et al., (2007)
O ₃ + NO ==> O ₂ + NO ₂	1.40 × 10 ⁻¹²	0	10.9	Atkinson et al. (2004)
CH ₃ O ₂ + NO ==> CH ₃ O + NO ₂	2.30 × 10 ⁻¹²	0	-2.99	Atkinson et al. (2006b)
HO ₂ + NO ==> NO ₂ + OH	3.6 × 10 ⁻¹²	0	-2.24	Atkinson et al. (2004)
CH ₂ O + Cl ==> HCl + HCO	8.20 × 10 ⁻¹¹	0	0.28	Atkinson et al. (1992)
CH ₂ O + OH ==> HCO + H ₂ O	4.73 × 10 ⁻¹²	1.18	-1.87	Baulch et al. (1992)
CH ₃ O ₂ + HO ₂ ==> CH ₃ OOH + O ₂	3.80 × 10 ⁻¹³	0	-6.49	Atkinson et al. (1992)
CH ₃ OOH ==> CH ₃ O + OH	6.00 × 10 ⁻¹⁴	0	177	Baulch et al. (1994)
HCO + O ₂ ==> CO + HO ₂	5.20 × 10 ⁻¹²	0	0	Atkinson et al. (2006b)
CO + OH ==> CO ₂ + H	5.40 × 10 ⁻¹⁴	1.5	-2.08	Baulch et al. (1992)
Cl + HO ₂ ==> HCl + O ₂	1.80 × 10 ⁻¹¹	0	-1.41	Atkinson et al. (1992)
Cl + HO ₂ ==> ClO + OH	6.30 × 10 ⁻¹¹	0	4.74	Atkinson et al. (2007)
Cl + O ₃ ==> ClO + O ₂	2.80 × 10 ⁻¹¹	0	2.08	Atkinson et al. (2007)
CO + Cl ==> ClCO	1.33 × 10 ⁻³³	-3.8	0.00	Atkinson et al. (2007)
OH + HOCl ==> H ₂ O + ClO	5.00 × 10 ⁻¹³	0	0	Atkinson et al. (2007)
ClO + HO ₂ ==> HOCl + O ₂	2.20 × 10 ⁻¹²	0	-2.8	Atkinson et al. (2007)
ClO + ClO ==> Cl ₂ + O ₂	1.00 × 10 ⁻¹²	0	13.22	Atkinson et al. (2007)
ClO + ClO ==> OClO + Cl	3.50 × 10 ⁻¹³	0	11.39	Atkinson et al. (2007)
ClO + ClO ==> ClOO + Cl	3.00 × 10 ⁻¹¹	0	20.37	Atkinson et al. (2007)
ClO + NO ==> Cl + NO ₂	6.20 × 10 ⁻¹²	0	-2.45	Atkinson et al. (2007)
CH ₂ O + O ==> HCO + OH	1.78 × 10 ⁻¹¹	0.57	11.56	Baulch et al. (1992)
OH + NO ₂ ==> HNO ₃	2.70 × 10 ⁻¹¹	0	0	Troe (2012)
CH ₃ Cl + OH ==> CH ₂ Cl + H ₂ O	1.40 × 10 ⁻¹²	1.6	8.65	Cohen and Westberg (1991)
CH ₃ Cl + H ==> CH ₃ + HCl	6.14 × 10 ⁻¹¹	0	38.9	Westenberg and deHaas (1975)
CH ₃ Cl + CH ₃ ==> CH ₄ + CH ₂ Cl	2.09 × 10 ⁻¹²	0	48.6	Macken and Sidebottom (1979)
CH ₃ Cl + Cl ==> CH ₂ Cl + HCl	3.30 × 10 ⁻¹¹	0	10.39	Atkinson et al. (2008)
CHCl ₃ + Cl ==> CCl ₃ + HCl	4.90 × 10 ⁻¹²	0	10.31	Atkinson et al. (2008)
Cl + C ₃ H ₆ ==> Products	2.70 × 10 ⁻¹⁰	0	0	Atkinson et al. (2006b)
Cl + C ₅ H ₈ ==> Products	4.30 × 10 ⁻¹⁰	0	0	Orlando et al. (2003)
ClNO + M ==> Cl + NO	2.16 × 10 ⁻⁹	0	134	Baulch et al. (1981)

Table A2. Termolecular reactions and parameters used for the modelling described in Sect. 2.5. The effective rate constant is calculated by combining the low- and high-pressure limit expressions into the following formula:

$$k_f(T, [M]) = \left\{ \frac{k_\infty(T)k_0(T)[M]}{k_\infty(T) + k_0(T)[M]} \right\} 0.6 \left\{ 1 + \left[\log_{10} \left(\frac{k_0(T)[M]}{k_\infty(T)} \right) \right]^2 \right\}^{-1}.$$

Reaction	Low-pressure limit		High-pressure limit		Reference
	$k_0 = k_0^{298}(T/298)^{-n}$		$k_\infty = k_\infty^{298}(T/298)^{-m}$		
	k_0^{298}	n	k_∞^{298}	m	
$\text{Cl} + \text{NO}_2 + M \Rightarrow \text{ClNO}_2 + M$	1.8×10^{-31}	2	1.1×10^{-10}	1	Burkholder et al. (2015)
$\text{CH}_3 + \text{O}_2 + M \Rightarrow \text{CH}_3\text{O}_2 + M$	4.1×10^{-31}	3.6	1.2×10^{-12}	-1.1	Burkholder et al. (2015)
$\text{Cl} + \text{C}_2\text{H}_4 + M \Rightarrow \text{Products}$	1.6×10^{-29}	3.3	3.1×10^{-10}	1	Burkholder et al. (2015)

Table A3. Initial concentrations for specified species simulated in model; listed mixing ratios are based on a temperature of 20 °C. Potential interferents were tested in separate model runs according to the groupings on each line below and were otherwise initiated with a concentration of 0 molec. cm⁻³. All other compounds were initialized with a concentration of 0 molec. cm⁻³.

Species	Initial concentration (molec. cm ⁻³)
ClNO ₂	2.46×10^{10} (1 ppbv)
N ₂	1.92×10^{19} (78 %)
O ₂	5.17×10^{19} (21 %)
CH ₄	4.92×10^{13} (2000 ppbv)
OH	1×10^6
O ₃	9.84×10^{11} (40 ppbv)
Potential interferents	
ClNO	2.46×10^{10} (1 ppbv)
Ethene, propene, isoprene	1.23×10^{12} (50 ppbv)

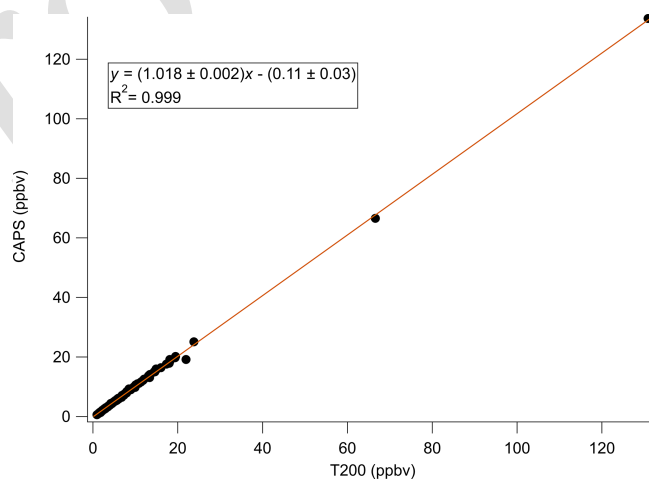


Figure A1. Laboratory calibration curve for CAPS NO₂.

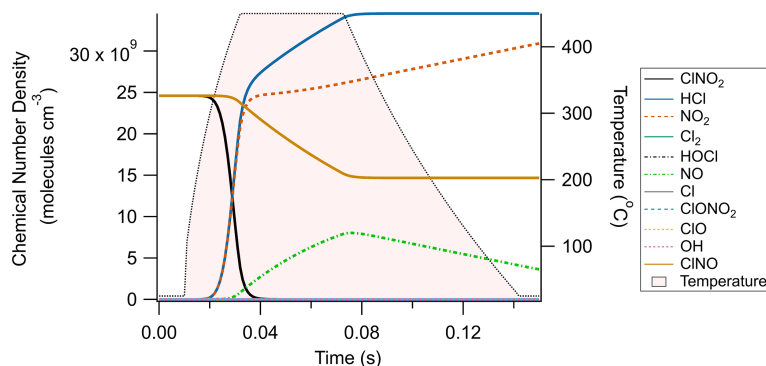


Figure A2. Modelled effects of 2.46×10^{10} (1 ppbv at 20 °C) of ClNO.

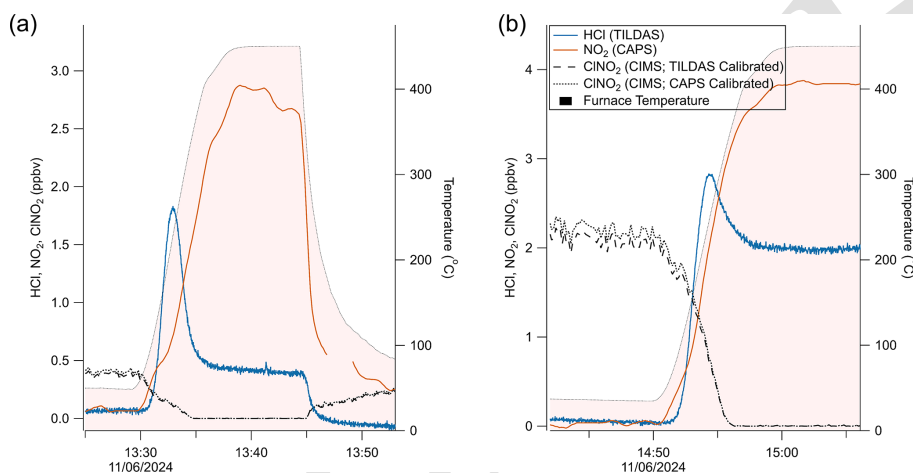


Figure A3. (a) Comparison plot of ClNO₂ observations with an apparent excess of NO₂ formed after ageing/processing of the same slurry used for generating Figs. 3–4. (b) Additional comparison using a freshly made slurry. CIMS signal was calibrated using humidity-dependent calibration factors as presented in Fig. 5.

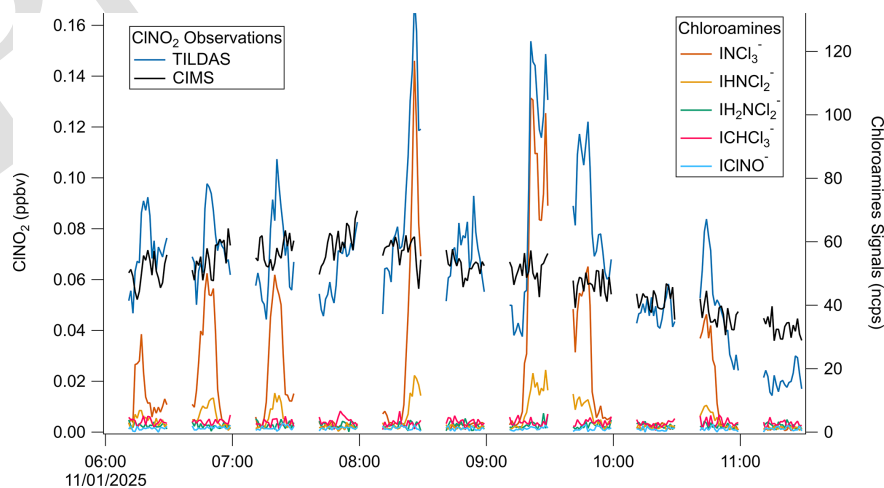


Figure A4. Field data showing apparent coincident signal increases for TILDAS-observed ClNO₂ with CIMS-observed chloramines, ICINO[−], and ICHCl₃[−] (uncalibrated).

Code availability. Code used for this analysis is available from the corresponding author on request.

Data availability. Data are available from the corresponding author on request.

Author contributions. PRV, MAR, and SSB designed and performed proof-of-concept experiments to demonstrate the potential of the method. SCH, HRR, CD, and TIY designed, built, and tested the HCl TILDAS at Aerodyne Research, Inc. SCH, HRR, CD, TIY, and PME designed initial laboratory experiments. JWH and PME designed laboratory and field experiments, and JWH conducted laboratory and field experiments presented in this work. LM, MDS, and LJC provided support for laboratory use of CIMS. EM, TJB, and HC provided support for field CIMS observations. JWH prepared the paper, and all authors reviewed the paper.

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Remarks from the language copy-editor

- CE1** The use of the article “a” is grammatically correct here.
- CE2** Please note the slight edits (something like “a more time-efficient method”, as suggested in the pdf comments, might be clearer).
- CE3** Please double-check and consider revising, as this is grammatically unclear and therefore potentially confusing (because it reads like an adverb modifying a noun). Would something like “near” work?

Remarks from the typesetter

- TS1** Please note changes in values (2.76×10^{-5} to 2.76×10^{-6}) require editor approval. Please provide a short explanation that can be forwarded by us to the editor.