Low levels of nitrily chloride at ground level: nocturnal nitrogen oxides in the Lower Fraser Valley of British Columbia

Hans D. Osthoff1, Charles A. Odame-Ankrah1, Youssef M. Taha1, Travis W. Tokarek1, Corinne L. Schiller2, Donna Haga3, Keith Jones2, and Roxanne Vingarzan2

1Department of Chemistry, University of Calgary, Calgary, Alberta T2N 1N4, Canada
2Applied Science Division, Prediction and Services West, Meteorological Service of Canada, Environment and Climate Change Canada, Vancouver, British Columbia V6C 3S5, Canada
3British Columbia Ministry of Environment and Climate Change Strategy, Cranbrook, British Columbia V1C 7G5, Canada

Correspondence: Hans D. Osthoff (hosthoff@ucalgary.ca)

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Abstract. The nocturnal nitrogen oxides, which include the nitrate radical (NO₃⁻), dinitrogen pentoxide (N₂O₅), and its uptake product on chloride containing aerosol, nitrily chloride (ClNO₂), can have profound impacts on the lifetime of NOₓ (NO + NO₂), radical budgets, and next-day photochemical ozone (O₃) production, yet their abundances and chemistry are only sparsely constrained by ambient air measurements. Here, we present a measurement data set collected at a routine monitoring site near the Abbotsford International Airport (YXX) located approximately 30 km from the Pacific Ocean in the Lower Fraser Valley (LFV) on the west coast of British Columbia. Measurements were made from 20 July to 4 August 2012 and included mixing ratios of ClNO₂, N₂O₅, NO, NO₂, total odd nitrogen (NOₓ), O₃, photolysis frequencies, and size distribution and composition of non-refractory submicron aerosol (PM₁).

At night, O₃ was rapidly and often completely removed by dry deposition and by titration with NO of anthropogenic origin and unsaturated biogenic hydrocarbons in a shallow nocturnal inversion surface layer. The low nocturnal O₃ mixing ratios and presence of strong chemical sinks for NO₃⁻ limited the extent of nocturnal nitrogen oxide chemistry at ground level. Consequently, mixing ratios of N₂O₅ and ClNO₂ were low (< 30 and < 100 parts-per-trillion by volume (pptv) and median nocturnal peak values of 7.8 and 7.9 pptv, respectively). Mixing ratios of ClNO₂ frequently peaked 1–2 h after sunrise rationalized by more efficient formation of ClNO₂ in the nocturnal residual layer aloft than at the surface and the breakup of the nocturnal boundary layer structure in the morning. When quantifiable, production of ClNO₂ from N₂O₅ was efficient and likely occurred predominantly on unquantified supermicron-sized or refractory sea-salt-derived aerosol. After sunrise, production of Cl radicals from photolysis of ClNO₂ was negligible compared to production of OH from the reaction of O(^1D) + H₂O except for a short period after sunrise.

1 Introduction

The Lower Fraser Valley (LFV) is prone to episodes of poor air quality, in part because of its geography, which facilitates stagnation periods and accumulation of airborne pollutants through processes such as the Wake-Induced Stagnation Effect (Brook et al., 2004), and also because of continued growth of human population and associated emissions from urban, suburban, agricultural, and marine sources. Of special concern have been repeated exceedances of the Canada-wide standard and, as of 2012, the Canadian Ambient Air Quality Standards (CAAQS) for fine particulate matter (PM₂.₅) and ozone (O₃) at Chilliwack and Hope, located in the eastern part of the LFV downwind of Vancouver (Ainslie et al., 2013). These exceedances have occurred in spite of ongoing declines in emissions of both nitrogen oxides (NOₓ = NO + NO₂) and volatile organic compounds (VOCs) resulting from the introduction of new vehicle standards and (now discontinued) local vehicle emission testing programs (Ainslie et al., 2013). Previous large-scale studies
in the LFV such as Pacific 1993 (Steyn et al., 1997), the Regional Visibility Experimental Assessment in the Lower Fraser Valley (REVEAL) I and II (Pryor et al., 1997; Pryor and Barthelme, 2000), and Pacific 2001 (Vingaran and Li, 2006) have added important information regarding atmospheric processes, leading to O3 and aerosol formation and visibility issues. However, the transformation of primary (e.g., NOx, VOCs, SO2, NH3) to secondary pollutants (i.e., O3 and fine particulate matter) is highly complex, and the scientific understanding of these highly non-linear processes remains incomplete.

A complicating factor in the LFV is the interaction of anthropogenic emissions with marine-derived sea salt aerosol. While sea spray aerosol is a primary source of particulate matter (PM) and hence directly affects particle concentrations and mass loadings (Pryor et al., 2008) and aerosol chloride concentrations (Anlauf et al., 2006) in the LFV, there is now considerable evidence from modelling (Knipping and Dabdub, 2003), laboratory (Raff et al., 2009), and field studies (Tanaka et al., 2003; Osthoff et al., 2008) that “active chlorine” species released from sea salt can negatively affect air quality and promote O3 and secondary aerosol formation in coastal regions.

In an analysis of 20 years of O3 air quality data in the LFV region, Ainslie and Steyn (2007) concluded that precursor buildup, prior to an exceedance day, plays an important role in the spatial O3 pattern on exceedance days. Secondary processes involving active chlorine produced from the interaction of marine aerosol with anthropogenic pollution would fit this profile but are not currently constrained by measurements.

One pathway to activate chlorine from sea salt is the reactive uptake of dinitrogen pentoxide (N2O5) on chloride containing aerosol to yield nitryl chloride (ClNO2) (Behnke et al., 1997; Finlayson-Pitts et al., 1989). Briefly, N2O5 is formed from the reversible reaction of nitrogen dioxide (NO2) with the photolabile nitrate radical (NO3; Reaction R1), which in turn is formed from reaction of NO2 with O3 (Reaction R2).

\[
\text{NO}_2 + \text{O}_3 \rightarrow \text{NO}_3 + \text{O}_2 \quad \text{(R1)}
\]

\[
\text{NO}_3 + \text{NO}_2 \rightarrow \text{N}_2\text{O}_5 \quad \text{(R2)}
\]

In ambient air, N2O5, NO3, and NO2 are usually in equilibrium; the equilibrium constant, K2, is temperature dependent, favouring NO3 and NO2 at higher temperatures (Osthoff et al., 2007). During daytime, NO3 (and, indirectly, N2O5) is removed primarily via its Reaction (R3) with NO (which is generated from NO2 photolysis and directly emitted, for example, by automobiles) and by NO3 photolysis (Reaction R4) (Wayne et al., 1991).

\[
\text{NO}_3 + \text{NO} \rightarrow 2\text{NO}_2 \quad \text{(R3)}
\]

\[
\text{NO}_3 + \text{h} \nu \rightarrow 0.9\text{NO}_2 + 0.1\text{NO} \quad \text{(R4)}
\]

The heterogeneous hydrolysis of N2O5 to nitric acid (HNO3) is an important nocturnal NO3 and odd oxygen (O3 = NO2 + O3) removal pathway (Chang et al., 2011; Brown et al., 2006a). On chloride containing aerosol, however, uptake of N2O5 yields up to a stoichiometric amount of ClNO2 (Reaction R5) (Behnke et al., 1997; Finlayson-Pitts et al., 1989):

\[
\text{N}_2\text{O}_5 + \text{H}_2\text{O}(\text{het}) + \varphi\text{Cl}^- (\text{het}) \rightarrow (2 - \varphi)
\]

\[
\text{HNO}_3(\text{het}) + \varphi\text{ClNO}_2, \quad 0 \leq \varphi \leq 1. 
\]  

(R5)

The ClNO2 yield, \( \varphi \), is primarily a function of aerosol chloride and water content (Behnke et al., 1997; Bertram and Thornton, 2009; Roberts et al., 2009; Ryder et al., 2014, 2015). Formation of ClNO2 impacts air quality in the following ways: since ClNO2 is long-lived at night, its primary fate is photodissociation (to Cl and NO2) in the morning hours after sunrise (Reaction R6) (Osthoff et al., 2008).

\[
\text{ClNO}_2 + \text{h} \nu \rightarrow \text{NO}_2 + \text{Cl} 
\]

(R6)

This reaction increases the morning abundance of O3, leading to greater net photochemical O3 production throughout the day. The other photo-fragment, the Cl atom, is highly reactive towards hydrocarbons and will initiate radical chain reactions that produce O3 and secondary aerosol (Behnke et al., 1997; Young et al., 2014). The fate and impact of ClNO2 is thus similar to that of nitrous acid (HONO), which also accumulates during the night and photodissociates in the morning to release NO and the hydroxyl radical (OH) that go on to produce O3 (Alicke et al., 2003).

Data collected during the 2006 Texas Air Quality Study – Gulf of Mexico Atmospheric Composition and Climate Study (TEXAQS-GOMACCS) have shown that ClNO2 production is efficient in the nocturnal polluted marine boundary layer even on primarily non-sea-salt aerosol surfaces (Osthoff et al., 2008). As a result, up to 15% of total odd nitrogen (NOy) was present in the form ClNO2 at night (Osthoff et al., 2008). The high efficiency of ClNO2 formation on aerosol of medium-to-low total chloride content has been confirmed by several laboratory investigations (Bertram and Thornton, 2009; Raff et al., 2009; Roberts et al., 2009) and direct measurements of N2O5 uptake on ambient particles (Riedel et al., 2012b). Some ambiguity remains as to the detailed mechanism of Reaction (R5), but there is agreement that acid displacement of HCl from supermicron (predominantly sea salt aerosol) to submicron (predominantly non-sea-salt aerosol) is a key step in the efficient production of ClNO2. These results suggested that this chemistry is active anywhere that pollution in the form of NO3 and O3 comes in contact with marine air, including the LFV.

However, while the yield of ClNO2 in Reaction (R5) is high in polluted coastal regions, the ClNO2 yield relative to
the amount of NO₃ produced via Reaction (R1) cannot be easily predicted because NO₃ is consumed by reactions with VOCs (Reaction R7), e.g., with biogenic VOCs such as isoprene and monoterpenes as well as aldehydes, and dimethyl sulfide (Wayne et al., 1991).

\[
\text{NO}_3 + \text{VOC} \rightarrow \text{products} \quad \text{(R7)}
\]

Previous studies in the LFV have shown high biogenic VOC concentrations (Biesenthal et al., 1997; Gurren et al., 1998; Drewitt et al., 1998) yet there was active nighttime nitrogen oxide chemistry and aerosol chloride present mainly as sea-salt-derived aerosol in > 1 µm diameter aerosol (Anlauf et al., 2006). During the Pacific 2001 study, measurements of the mixing ratios of NO, NO₂, peroxynitric anhydride (CH₃C(O)O₂NO₂, PAN), HONO, HNO₃, and NO₃ at three ground sites in the LFV indicated deficits of up to 15 % in the nocturnal NO₂ budget (Hayden et al., 2004) attributable to unquantified species such as alkyl nitrates, N₂O₅, and ClNO₂. McLaren and coworkers quantified mixing ratios of NO₂ and NO₃ by differential optical absorption spectroscopy (DOAS) at the Sumas Eagle Ridge site (~250 m above the floor of the LFV) as part of Pacific 2001 (McLaren et al., 2004) and off-shore on Saturna Island (Fig. 1) in the Strait of Georgia in 2005 (McLaren et al., 2010). The LFV data showed occasional episodes of active nocturnal nitrogen oxide chemistry in the residual layer with N₂O₅ contributing up to 9 % of NO₃, while the Saturna Island data showed NO₃ mixing ratios of > 20 parts-per-trillion by volume (10⁻¹² pptv) every night of measurement. McLaren et al. (2010) estimated that between 0.3 and 1.9 ppbv of ClNO₂ would be produced under these conditions. Efficient formation of ClNO₂ would be consistent with the unidentified O₃ precursor proposed by Ainslie and Steyn and is also a plausible explanation for part of the deficit in the NO₂ budget observed by Hayden et al. (2004).

Another feature of the LFV are somewhat unusual diurnal profiles arising from the vertical structure in pollutant concentrations. Measurements of O₃ and NO₂ using tethered balloons by Pisano et al. (1997) during Pacific 93 at the Harris Road site (located ~ 38 km NW of Abbotsford International Airport) revealed a highly stratified boundary layer with a shallow, 50 m deep isothermal surface layer (also called a nocturnal boundary layer, or NBL) and low surface O₃ concentrations at night. Nocturnal loss of surface O₃ is known to occur by several pathways, including dry deposition, titration with NO (Reaction R8), and reaction with unsaturated biogenic hydrocarbons (Neu et al., 1994; Kleinman et al., 1994; Trainer et al., 1987; Logan, 1989; Talbot et al., 2005). Titration of O₃ with NO is readily measured as the concentration of a product of Reaction (R8), NO₂, can be used to measure directly and conserves O₃.

\[
\text{O}_3 + \text{NO} \rightarrow \text{O}_2 + \text{NO}_2 \quad \text{(R8)}
\]

Usually, the major nocturnal sink of O₃ is dry deposition of O₃ and NO₂ (Lin et al., 2010).

![Figure 1. Map of the Lower Fraser Valley. YXX is the Abbotsford International Airport (measurement location for this study). YVR is the Vancouver International Airport. YYJ is the Victoria International Airport. BLI is the Bellingham International Airport. SAT is Saturna Island.](Image)

The balloon data also showed pools of NO₂ and O₃ in a ~100 m deep nocturnal residual layer (NRL) located 200 to 350 m above ground. Following the breakup of the nocturnal layers in the early morning, vertical down-mixing events of O₃ pollution were observed (McKendry et al., 1997). In this process, pollutants are entrained into the growing mixed layer from the NRL, i.e., the growing mixed layer in the hours after sunrise erodes the somewhat deeper NRL, and pollutants are mixed to the surface (Neu et al., 1994; Kleinman et al., 1994).

In this paper, we present the first measurements of ClNO₂ and N₂O₅ mixing ratios in the LFV. The data were collected at a surface site east of the Abbotsford International Airport (International Air Transport Association (IATA) airport code YXX) located approximately 35 km from the Pacific Ocean from 20 July to 5 August, 2012. Auxiliary measurements included NO, NO₂, NO₃, O₃, photolysis frequencies, and non-refractory PM₁ aerosol composition and size distributions. An analysis of nocturnal nitrogen oxide chemistry including the formation of ClNO₂ and its potential impact on nocturnal O₃ and NO₂ loss and radical budgets in the LFV is presented.

2 Experimental

2.1 Location

The map shown in Fig. 1 indicates the location of the study. Ambient air measurements were conducted at the T45 routine monitoring site located to the east YXX at latitude 49.0212 (N) and longitude −122.3267 (W) and ~ 60 m
above sea level (a.s.l.) and ∼ 30 km from the Pacific Ocean.
A raspberry field was located immediately to the W between
the end of the airport runway and the measurement site.
Nearby local sources included agricultural operations (such
as poultry farms) and emissions from motor vehicle traffic on
secondary roads and highways. YXX is located ∼ 60 km ESE
of the Vancouver International Airport (YVR) and the City of
Vancouver. Abbotsford is in the heart of the so-called “lower
mainland”, the low-lying region stretching from the Pacific
Ocean at Vancouver to the NW and the Canada–USA border
to the S (north of Bellingham, BLI) to the eastern end of the
Fraser Valley, with a total population in excess of 2 500 000.

2.2 Measurement techniques

The measurement techniques used for this study are listed in
Table 1. Data were averaged to 5 min prior to presentation.

The instruments measuring O3 and nitrogen oxides were
housed in an air-conditioned trailer and sampled from a com-
mon 0.635 cm (1/4 in.) outer diameter (OD) and 0.476 cm
(3/16 in.) inner diameter (ID) Teflon™ inlet at a height of
4 m above ground; the setup is depicted in Fig. 3 of Tokarek
et al. (2014). A scroll pump whose flow rate was throttled
using a 50 standard litres per minute (slpm) capacity mass
flow controller was connected to the end of the common in-
let to minimize the residence time of the sampled air and
to reduce inlet “aging”, i.e., accumulation of aerosol on fil-
ters of individual instruments, whose inlets tapped into the
main inlet line at 90°. The total inlet flow was in the range of
18 to 20 slpm.

Measurements of PM1 aerosol composition and size dis-
tributions (Sct. 2.3) and of meteorological data were made
from the research trailer housing the routine measurements at
the site. The Agilent VOC measurements were made from a
research trailer owned by Environment and Climate Change
Canada (ECCC).

2.2.1 Quantification of CINO2 by iodide chemical
ionization mass spectrometry

Mixing ratios of CINO2 were quantified as iodide cluster
ions at m/z 208 using the THS Instruments iodide chemical
ionization mass spectrometer (iCIMS) described by Mielke
et al. (2011) and calibrated using the scheme by Thaler et
al. (2011). In this method, a gas stream containing CINO2 is
generated from reaction of Cl2 (Praxair, 10 ppmv in N2) with
an aqueous slurry saturated with NaNNO2 (Sigma-Aldrich)
(Reaction R9):

\[
\text{Cl}_2(g) + \text{NO}_2^-(aq) \rightleftharpoons \text{CINO}_2(g) + \text{Cl}^-(aq). \quad (R9)
\]

This gas stream was periodically added to the main inlet with
the aid of a normally open two-way valve connected to a vac-
uum pump in a similar fashion as described earlier for “N2O5
and PAN (Tokarek et al., 2014; Odame-Ankrah and Osthoff,
2011). The CINO2 content of the calibration gas stream was
quantified by thermal dissociation cavity ring-down spec-
troscopy (TD-CRDS) as described in Sect. 2.2.2. In total, 31
calibrations for CINO2 were carried out, spread out evenly
over the measurement period. The iCIMS response factor
at m/z 208 was (0.40 ± 0.06) Hz pptv⁻¹ (where the error
represents the standard deviation of repeated calibrations),
normalized to 10⁶ counts of reagent ion at m/z 127. The
37ClNO2I⁻ ion at m/z 210 was also monitored and found to be
(0.298 ± 0.004) times the signal at m/z 208 (r² = 0.944),
slightly lower than Standard Mean Ocean Chloride 37Cl mole
fraction in seawater of ∼ 0.319 (Wieser and Berglund, 2009)
and our previously observed ratios of 0.315 ± 0.003 in Cal-
gary (Mielke et al., 2011) and 0.3065 ± 0.0002 in Pasadena
(Mielke et al., 2013). The reason(s) for these differences is
unclear but they may be a result of fractionation processes
(Koecher and Wassenaar, 2010; Volpe et al., 1998), a topic
outside the scope of this paper.

The iCIMS was also used to quantify mixing ratios of PAN
at m/z 59 and PPN at m/z 73 (Slusher et al., 2004; Mielke
et al., 2011; Mielke and Osthoff, 2012). For this reason, part
of the instrument’s inlet prior to the ion–molecule reaction
region was heated to 190°C to dissociate PANs into their
respective peroxyacyl radicals and NO2. Further, the collis-
sional dissociation chamber (CDC) was operated in declus-
tering mode (∼ 22.7 V) to break up ion clusters. Calibrations
and matrix effect correction procedures and a time series
of the PAN and PPN data were presented by Tokarek et
al. (2014).

2.2.2 Quantification of NO2 and N2O5 by cavity
ring-down spectroscopy

The CRDS used in this work was an amalgamated ver-
sion of two instruments described earlier (Paul and Osthoff,
2010; Odame-Ankrah and Osthoff, 2011), called Improved
Detection Instrument for Nitrogen Oxide Species (iDinos)
(Odame-Ankrah, 2015). A schematic of the optical layout is
shown in Fig. 2. The optical bread board, instrument frame,
electronic, and data acquisition components were as de-
scribed by Paul and Osthoff (2010). The new instrument was
set up with up to six parallel detection channels: four 405 nm
“blue” diode laser CRDS cells for quantification at NO2 via
its absorption at 405 nm with a distance between the pairs
of high-reflectivity (HR) mirrors (Advanced Thin Films) of
112.5 cm, of which 92.0 cm were filled with sample air, and
two newly constructed 662 nm “red” diode laser CRDS cells
for quantification at NO1 via its absorption at 662 nm with a
distance between the HR mirrors (Los Gatos) of 93.0 cm
of which 73.0 cm were filled with sample air. Light exiting
the far ends of the CRDS cells was collected using fixed-
focus collimating lenses and multi-mode optical fibers (Thor-
labs) connected to photomultiplier tubes (PMTs; Hamamatsu
H9433-03MOD) with 10 MHz bandwidth. Bandpass filters
(Thorlabs FB405-10 and FB660-10) were placed between the
PMTs and the end of the optical fibers.
The two laser diodes were simultaneously square-wave modulated by a function generator (SRS DS335). The PMT voltages were digitized using an eight-channel 14-bit data acquisition card (National Instruments PCI-6133; 2.5 MS s\(^{-1}\)) simultaneous sampling sample rate) connected to a laptop computer via a PCMCIA-to-PCI expansion unit (Magma CB4DRQ) and controlled by software written in LabVIEW\textsuperscript{TM} (National Instruments).

Ring-down time constants (\(\tau\)) were determined from a linear fit to the logarithm of the digitized PMT voltage as described by Brown et al. (2002) immediately after acquisition of the ring-down traces (which were co-added to a user-selectable averaging time prior to the fit). The fitting algorithm requires the subtraction of the PMT voltage offset prior to taking the logarithm; this offset was measured between ring-down events after the signal had returned to baseline, which limited the repetition rate of the diode lasers and the number of traces averaged per second to a frequency of 300 Hz.

Ring-down time constants in the absence of the target absorber (\(\tau_0\)) were determined by flooding the inlet (each once per hour) with ultra-pure, or “zero”, air (Praxair) for the 405 nm channels and by titration with NO for the 662 nm channel (Brown et al., 2001; Simpson, 2003). Typical values of \(\tau_0\) were in the range of 63 to 67 \(\mu\)s and between 198 and 210 \(\mu\)s for the blue and red channels, respectively. The baseline precision (i.e., standard deviation, \(\sigma\)) of the NO\(_2\) and NO\(_3\) measurements were \(\pm 80\) and \(\pm 3\) pptv (1 s data), respectively. For the NO\(_3\) channels, additional noise was introduced by variable background absorption of NO\(_2\), O\(_3\), and water vapour which produce small, spurious structure in the 662 nm absorption signal (Dubé et al., 2006) and were not tracked well by the interpolation of the baseline from the hourly \(\tau_0\) determinations.

During the Abbotsford campaign, only five (four blue and one red) CRDS channels were operated because of delays in the fabrication of the final set of CRDS mirror holders. The 662 nm CRDS cell sampled from a Teflon\textsuperscript{TM} inlet heated to 130 \(^\circ\)C for quantification of NO\(_x\) plus the NO\(_3\) generated from thermal dissociation N\(_2\)O\(_5\) (Brown et al., 2001; Simpson, 2003; Dubé et al., 2006). Under the high-NO\(_x\) conditions of this study, equilibrium (2) was sufficiently far to the right (see Sect. 3.3) such that [NO\(_3\)] + [N\(_2\)O\(_5\)] \(\approx\) [N\(_2\)O\(_3\)]; i.e., the concentration measured could be equated with [N\(_2\)O\(_3\)] without introducing a large error (i.e., < 5\%). The four 405 nm CRDS cells were operated as follows: the first sampled from an ambient temperature inlet and was used to quantify NO\(_2\). The second sampled from a quartz inlet heated to 250 \(^\circ\)C and was used to quantify NO\(_2\) plus total peroxyacyl nitrate (\(\Sigma\)PAN) (Paul et al., 2009; Paul and Osthoff, 2010). Data from this channel will be presented in a future publication. The third was operated with a quartz inlet heated to 450 \(^\circ\)C to enable ClNO\(_2\) calibrations (Thaler et al., 2011). Quantification of total alkyl nitrates (\(\Sigma\)AN) in ambient air was not attempted because of the high NO\(_3\) levels and resulting large subtraction errors (Thieser et al., 2016). The
Fourth 405 nm CRDS cell was connected with polycarbonate tubing (3/8 in. OD and 1/4 in. ID) in series to the 662 nm channel and was used to calibrate the response of the N$_2$O$_5$ channel, which is a function of the transmission efficiency of N$_2$O$_5$ through the inlet and the overlap of the diode laser spectrum with the NO$_3$ absorption line (Odame-Ankrah and Osthoff, 2011). The role of the polycarbonate tube was to scrub NO$_3$ exiting the N$_2$O$_5$ channel, allowing detection of only the NO$_2$ generated from thermal dissociation of N$_2$O$_5$ and to prevent recombination of NO$_3$ and NO$_2$ in the blue calibration channel (Wagner et al., 2011).

N$_2$O$_5$ was generated in situ by adding an excess of O$_3$ (generated by passing O$_2$ past a 254 nm Hg lamp) to nitric oxide (NO) in a 0.635 cm (1/4 in.) OD and 0.476 cm (3/16 in.) ID Teflon™ calibration line and allowed to equilibrate (i.e., until the output was constant) offline before being switched inline on demand. The N$_2$O$_5$ response (which accounted for N$_2$O$_5$ loss in the sampling line and slight mismatches of the laser wavelengths with the NO$_3$ absorption line) varied between 65 and 100 % and depended on inlet “age”; the Teflon™ inlet and aerosol inlet filter were changed every 2–3 days. The accuracy of the NO$_2$ and N$_2$O$_5$ data were ±10 and ±25 %, respectively, driven mainly by the systematic uncertainty of the NO$_2$ absorption cross section and of the N$_2$O$_5$ inlet transmission efficiency (Odame-Ankrah, 2015).

**2.2.3 Measurements of O$_3$, NO, and NO$_y$**

Mixing ratios of O$_3$ were monitored by UV absorption in a commercial instrument (Thermo 49) and were accurate within ±2 % and ±1 ppbv. An NO-O$_3$ chemiluminescence instrument (Thermo 42i) was used to monitor mixing ratios of NO and NO$_y$, which was reduced to NO in a Mo converter heated to ∼ 320 °C placed outside a short distance (~1 m) from the sample inlet. This instrument sampled from the main inlet via a Teflon™ filter and filter holder and was calibrated daily against CRDS as described by Tokarek et al. (2014). The slope uncertainty for each multipoint calibration was ±15 %. Interpolation between calibration runs gave an overall uncertainty of ±30 %. The NO zero offset uncertainty (needed for calculating the NO$_3$ loss rate with respect to reaction with NO, Reaction R9) was ±10 ppbv.

**2.2.4 VOC measurements**

Volatile organic compounds were monitored with a commercial gas chromatograph–mass spectrometer (GC-MS; Agilent model 7890A and 5975C) equipped with an FID detector and a Markes Unity 2 pre-concentrator with an ozone precursor trap cooled to −25 °C.

In a typical sampling sequence, a 500 mL air sample was collected at a flow rate of 25 mL min$^{-1}$, taken from the centre flow of a 1.27 cm (1/2 in.) stainless steel inlet line which was continuously sampling ambient air at 5 L min$^{-1}$. The sampled air flowed through a 0.318 cm (1/8 in.) stainless steel line and particles were removed using a 1 µm pore size fritted filter. Once 500 mL of air were collected, the pre-concentrator was flushed with helium to remove air while awaiting injection. At the start of a GC run, the sample in the pre-concentrator was flash heated to 300 °C and held for 3 min. The sample was separated on two columns with the entire sample going through the Agilent VRX column with a Dean switch directing the first gases emitted to a second GasPro column and then to the FID detector (~ < C4), while the heavier compounds were detected using the MS detector in scan mode.

The cycle time for the GC analysis was 1 h with the sample being collected during the previous runs analyses. The 20 min sample was taken at the start of a 1 h time period.

Due to the low temperature of the trap, the air was dried using a trap at −30 °C. The trap was heated and dried between each sample and reconditioned for 10 min prior to sample collection. All sample lines were stainless steel with a Restek Sulfinert™ coating to minimize sample loss on the lines. Calibrations were performed once per day for 105 species using a 100 ppbv US Environmental Protection Agency (EPA) photochemical assessment monitoring system (PAMS) and a 100 ppbv EPA air method, toxic organics – 15 (TO15) standard at an approximate concentration. The terpenes were semi-quantitatively measured as a calibration source was not available at the time and only the changes in concentration...
strength with time of day were used. The accuracy of the measurements varied depending on the species but was better than ±30 % throughout. Peaks were manually reintegrated using Chemstation software from Agilent. Table S1 in the Supplement summarizes the VOCs quantified.

### 2.3 Aerosol measurements

The chemical composition of non-refractory PM1 was monitored using an Aerosol Chemical Speciation Monitor (ACSM, Aerodyne), which reported concentrations of NO3⁻, SO4²⁻, Cl⁻, NH4⁺, and total organics. A general description of this instrument designed for routine monitoring has been given by Ng et al. (2011). The composition of the refractory aerosol (i.e., sea salt) was not quantified.

Submicron aerosol size distributions were quantified by a scanning mobility particle sizer (SMPS, TSI 3034). This instrument measured aerosol particles in the range from 10 to 487 nm using 54 size channels (32 channels per decade). Both of these instruments were housed in a trailer operated by Metro Vancouver. The ACSM and the SMPS sampled air off a shared stainless steel inlet that had a total flow of 5 L min⁻¹ and contained a PM2.5 sharp cut filter at the inlet and was operated at ambient relative humidity.

### 2.4 Photolysis frequencies

Photolysis frequencies were determined by solar actinic flux spectroradiometry (Hofzumahaus et al., 1999) using a commercial radiometer with 2π receptor optics and photodiode array (PDA) detector (Metcon; 512 pixels, wavelength range 285–690 nm) calibrated by the manufacturer. The spectrometer was mounted facing up (zenith view) and hence measured the down-welling radiation. On several days, the spectrometer was inverted hourly to determine the up-welling radiation, which was added to the down-welling flux. Photolysis frequencies including \( j(\text{NO}_3) \), \( j(\text{NO}_2) \), \( j(\text{O}_1\text{D}) \), and \( j(\text{ClNO}_2) \) were calculated using reference spectra and quantum yields from Sander et al. (2010) and Ghosh et al. (2012). Table 2 gives the ratio of observed up-welling to down-welling for selected photolysis frequencies. For 3 August (a cloud-free day), the measurements were compared to (hourly) predictions with the online Tropospheric Ultraviolet and Visible (TUV) radiation model V5.0 (Madronich and Flocke, 1997); with default settings, the model reproduced the measured \( j(\text{NO}_2) \) and \( j(\text{O}_1\text{D}) \) quite well: a scatter plot of observed against TUV rate constants had correlation coefficients (\( r \)) of 0.997 and 0.998, slopes of 1.06 ± 0.02 and 1.10 ± 0.02, and offsets of \( (3 \pm 1) \times 10^{-4} \text{s}^{-1} \) and \( (5 \pm 3) \times 10^{-7} \text{s}^{-1} \).

### 2.5 Box model simulations of the nocturnal \( \text{O}_3 \) and \( \text{O}_x \) loss in the NBL

A box model was set up to reconcile the median nocturnal decays of \( \text{O}_3 \) and \( \text{O}_x \). These simulations are intended as back-of-the-envelope type estimates of major processes only since an accurate description of the nocturnal boundary layer chemistry would require modelling of horizontal and vertical transport, i.e., altitude-resolved information not available in this study (Geyer and Stutz, 2004). The model’s assumptions are a well-mixed NBL that is decoupled from the NRL above it as observed by earlier balloon vertical profiling (Pisano et al., 1997). \( \text{O}_3 \) and \( \text{NO}_2 \) dry deposition velocities of \( \nu_d(\text{O}_3) = 0.2 \text{cm s}^{-1} \) and \( \nu_d(\text{NO}_2) = \alpha \times \nu_d(\text{O}_3) \) with \( \alpha = 0.65 \) (Lin et al., 2010), and negligible chemical loss other than titration of \( \text{O}_3 \) by NO (Reaction R8) and by reaction with a generic biogenic hydrocarbon (assumed to react with \( \text{O}_3 \) with a rate coefficient of \( 5 \times 10^{-11} \text{cm}^3 \text{molec}^{-1} \text{s}^{-1} \), i.e., the rate coefficient for reaction of \( \alpha \)-pinene with \( \text{O}_3 \); Seinfeld and Pandis, 2006). Simulations were initiated with the median \( \text{NO}_2 \) and \( \text{O}_3 \) concentrations observed at sunset. More details are given in the Supplement.

### 3 Results

#### 3.1 Overview of data set

#### 3.1.1 Meteorology

A time series of local wind direction and speed are displayed in Fig. 3d. During the 2-week-long measurement period, the air flow to the site was from the Pacific Ocean to the SW and WSW with a moderate wind speed of 8.7 km h⁻¹ (median value). On most nights, local wind speeds were calm, i.e., < 5 km h⁻¹ (median speed 3.6 km h⁻¹) and from variable directions, though predominantly from the W and N. The two exceptions were the nights of 22–23 July and 1–2 August when stronger winds (> 5 km h⁻¹) from the W and SW persisted. These nights saw relatively high \( \text{ClNO}_2 \) mixing ratios (see Sect. 3.1.4).

The air temperatures were quite mild and ranged from a minimum of 11.0 °C to a maximum of 31.9 °C. The warm temperatures shifted equilibrium \( K_2 \) from \( \text{N}_2\text{O}_5 \) towards \( \text{NO}_3 \) and \( \text{NO}_2 \) (further analyzed in Sect. 3.2.2). At night, temperatures frequently dropped to the dew point, resulting in occasional fog formation (shown as grey rectangles in Figure 3D), sometimes after sunrise. Fog droplets are strong sinks for \( \text{N}_2\text{O}_5 \) (Osthoff et al., 2006). In total, the impact of fog was minor, affecting 5 % of the data. In addition, there were two periods with precipitation: the first occurred inter-
mittently on 20 July until the morning of 21 July. The second rainfall event was a 24 h period from mid-day 22 July to the afternoon of 23 July (shown as blue dots in Fig. 3d). 23 July also exhibited the highest wind speeds of the campaign (Fig. 3c) and lowest daytime photolysis frequencies. The time series of \( j(\text{ClNO}_2) \) is shown as a representative example in Fig. 3a. The photolysis data indicate that it was sunny on 6 days (25, 26, 29 July and 1, 4, 5 August) and that the remaining days had variable cloud cover, consistent with hourly meteorological logs that showed 10% of the measurement period affected by precipitation.

### 3.1.2 NO and NO\(_2\)

The rates of \( \text{N}_2\text{O}_3 \) and ClNO\(_2\) formation depend on the rate of NO\(_3\) production, \( P(\text{NO}_3) = k_1[\text{NO}_2][\text{O}_3] \) (analyzed further in Sect. 3.2.2); therefore, it is informative to first examine the mixing ratios of NO\(_2\) and O\(_3\) (see Sect. 3.1.3). The time series of NO, NO\(_2\), O\(_3\), and O\(_3\) (\( \text{O}_3 + \text{NO}_2 \)) mixing ratios are shown in Fig. 3c, and their diurnal averages are shown as 10th, 25th, 50th, 75th, and 90th percentiles in Fig. 4b and c.

The median NO and NO\(_2\) mixing ratios for the entire campaign were 0.9 and 5.9 ppbv, respectively. The average NO\(_3\)/NO\(_2\) ratio for the entire campaign was 0.9 ± 0.4. These concentration levels are characteristic of an urban air mass impacted by relatively fresh emissions from combustion engines in automobiles.

At night, mixing ratios of NO were generally lower than during the day though not negligible (median 0.3 ppbv, Fig. 4b) as NO was oxidized by O\(_3\) to NO\(_2\) (Reaction R8) and was not replenished by NO photolysis. However, mixing ratios of NO increased throughout the night, often coinciding with complete nocturnal removal of O\(_3\) (see Sect. 3.1.3), which indicates the presence of nearby combustion sources of NO\(_3\) (most likely automobile exhaust). The presence of NO titrates NO\(_3\) (Reaction R3) and effectively shut down \( \text{N}_2\text{O}_3 \) and ClNO\(_2\) production for most of the study: 68% of the measurement period had NO mixing ratios > 100 ppbv and NO\(_3\) lifetimes (with respect to its reaction with NO\(_3\)) of < 15 s. In contrast, NO\(_2\) mixing ratios were highest at night (median 7.3 ppbv), amplified further by NO\(_3\) emissions that continued throughout the night and likely by low nocturnal mixing heights (see discussion).

Mixing ratios of NO and NO\(_2\) were highest in the morning hours. Concentration changes at this time of day are difficult to interpret since the NBL breaks up during this time, resulting in vertical mixing of air masses, photolabile species (e.g., ClNO\(_2\), HONO, \( \text{N}_2\text{O}_5 \)) that accumulated overnight begin to photodissociate, and local emissions change with the onset of rush hour.

In contrast to the morning increase in NO, an afternoon/early evening maximum in NO was absent. This can be rationalized by a greater mixing height and abundance of oxidants that oxidize NO to NO\(_2\), i.e., O\(_3\) (see Figs. 3 and 4 and Sect. 3.1.3) and organic peroxy radicals in the afternoon, a topic outside the scope of this paper.

### 3.1.3 \( \text{O}_3 \) and O\(_3\)

The time series of \( \text{O}_3 \) mixing ratios and its diurnal profile are shown in Figs. 3c and 4c, respectively. \( \text{O}_3 \) mixing ratios were small (average ±1 standard deviation of 16 ± 12 ppbv) and peaked at ∼ 17:00 P PST in the afternoon. The highest concentrations were observed on August 4 from 13:55 to 15:30, when mixing ratios were 64 ± 1 ppbv (the 8 h running average was 52 ppbv). These levels were well below the CAAQS 8 h standard of 63 ppbv and the 1 h National Ambient Air Quality Objective of 82 ppbv, smaller than the pre-2003 data analyzed by Ainslie and Steyn (2007), who reported between 10 and 20 \( \text{O}_3 \) 1 h exceedances of 82 ppbv in the 1980s, and of similar magnitude as observed by a high-density monitoring network in the region in 2012 (Bart et al., 2014), which observed peak \( \text{O}_3 \) levels of 74 and 83 ppbv at Abbotsford on 8 July and 17 August, respectively.

A recurring feature of this data set was the rapid and often complete loss of \( \text{O}_3 \) at night (Fig. 4c). This was accompanied by an increase in the NO\(_2\) mixing ratios, though by less (+6 ppbv on average) than the amount of \( \text{O}_3 \) that was lost (∼26 ppbv on average), showing that NO to NO\(_2\) conversion (Reaction R8) was a contributor, though minor (∼25%) to the nocturnal \( \text{O}_3 \) loss.

The diurnal profile of O\(_3\) is similar to that of \( \text{O}_3 \), in that the highest concentrations occurred in the afternoon (at ∼ 18:00) and a considerable fraction of O\(_3\) was removed at night. At sunset, a median amount of 26 ppbv of O\(_3\) were present, which decreased to 12 ppbv at sunrise (Fig. 4c). The pathways contributing to nocturnal \( \text{O}_3 \) and O\(_3\) loss are probed using box model simulations in Sect. 3.2.1.

There were two (out of 16 total) nights when \( \text{O}_3 \) was not completely removed. On 22–23 July and 1–2 August, \( \text{O}_3 \) mixing ratios dropped from a daytime maxima of ∼ 33 ppbv to non-zero nocturnal minima of ∼ 16 ppbv. On both of these nights, ClNO\(_2\) and \( \text{N}_2\text{O}_3 \) mixing ratios were elevated (Fig. 3a), and the two largest ClNO\(_2\) to NO\(_3\) ratios were observed (Fig. 3b). The local wind speeds were > 6 km h\(^{-1}\), whereas on other nights, local winds were calmer (Fig. 3c). The greater local wind speeds likely induced more turbulence and a higher vertical mixing height.

### 3.1.4 \( \text{N}_2\text{O}_5 \) and ClNO\(_2\)

Time series of ClNO\(_2\) and \( \text{N}_2\text{O}_5 \) mixing ratios and ClNO\(_2\) photolysis frequencies are shown in Fig. 3a. Mixing ratios of ClNO\(_2\) and \( \text{N}_2\text{O}_5 \) were small (campaign averages at night of 4.0 and 1.4 pptv, respectively). The mixing ratios peaked prior to sunrise at median values of 7.9 and 7.8 pptv for ClNO\(_2\) and \( \text{N}_2\text{O}_5 \), respectively. The highest mixing ratios of this campaign were 97 pptv for ClNO\(_2\) and 23 pptv for \( \text{N}_2\text{O}_5 \), both observed on the night of 1–2 August. This night was
Figure 3. (a) Time series of \( N_2O_5 \) and ClNO\(_2\) mixing ratios (left axis) and ClNO\(_2\) photolysis frequency (right axis) observed at T45 near the Abbotsford International Airport. (b) Time series of the ratios of ClNO\(_2\) and \( N_2O_5 \) to NO\(_y\) (left axis) and of NO\(_y\) (right axis). (c) Time series of NO, NO\(_2\), O\(_3\), and O\(_x\) (= NO\(_2\) + O\(_3\)) mixing ratios. (d) Time series of local wind direction (left axis) and speed (right axis). The blue and grey dots above the time series indicates periods of precipitation (drizzle or rain) and fog, respectively, as identified in hourly meteorological logs.

also the only time when nocturnal ClNO\(_2\) mixing ratios exceeded 20 pptv and is analyzed in greater detail in Sect. 3.2.3.

Consistent with their low mixing ratios, neither ClNO\(_2\) nor N\(_2\)O\(_5\) were significant components of NO\(_y\) (Fig. 3b): on average, they contributed 0.1 % to the nocturnal NO\(_y\) budget, though NO\(_y\) mixing ratios were large (median 6.3 ppbv at night), typical for a site impacted by urban emissions. The only exception was the night of 1–2 August, when ClNO\(_2\) and N\(_2\)O\(_5\) constituted 2.6 and 1.6 % of NO\(_y\), respectively, and NO\(_y\) mixing ratios were 4.4 ppbv on average (Fig. 3b).

The ClNO\(_2\) and N\(_2\)O\(_5\) mixing ratios are displayed as functions of time of day in Fig. 4a. Before midnight local time, N\(_2\)O\(_5\) mixing ratios were slightly larger (median value of 1.8 pptv on average) than those of ClNO\(_2\) (median value of 1.4 pptv on average), whereas after midnight ClNO\(_2\) mixing ratios were larger than those of N\(_2\)O\(_5\) (2.0 vs. 0.6 pptv). The latter is consistent with observations at other ground sites, which generally showed higher concentrations of the longer-lived ClNO\(_2\) prior to sunset (Thornton et al., 2010; Mielke et al., 2013). The higher N\(_2\)O\(_5\) than ClNO\(_2\) abundances at the beginning of the nights suggest that the N\(_2\)O\(_5\) production rate at that time exceeded its ability to react heterogeneously and convert to ClNO\(_2\), potentially due to a lack of available aerosol chloride or otherwise reduced N\(_2\)O\(_5\) heterogeneous uptake parameters (Thornton et al., 2010).

Production of ClNO\(_2\) from N\(_2\)O\(_5\) uptake on aerosol ceases after sunrise because of the rapid removal of N\(_2\)O\(_5\) and NO\(_3\) as the latter is titrated by NO and destroyed by photolysis (Reactions R3, R4) (Wayne et al., 1991). In spite of this, ClNO\(_2\) mixing ratios frequently (on 12 out of 15 measurement days) continued to increase after sunrise (Figs. 3a and 4), peaking on average at \(~07:45\) in the morning approximately 2 h after sunrise. The median mixing ratio at that time was 6.7 pptv larger than the median value of 5.3 pptv observed at sunrise. The most prominent example of this phenomenon occurred on the morning of 26 July. For a 2-hour...
period leading up to sunrise, there was fog (virtually ensuring the absence of N\textsubscript{2}O\textsubscript{5}), and ClNO\textsubscript{2} mixing ratios were < 5 pptv. The fog then dissipated at sunrise. One hour later, ClNO\textsubscript{2} mixing ratios increased to > 40 pptv. Similar events (though with more modest ClNO\textsubscript{2} increases) were observed on the mornings of 22, 23, 25, 27, 28, 30, 31 July and 1 August. Two of these (23 and 27 July) overlapped with brief fog events.

Qualitatively similar ClNO\textsubscript{2} morning peaks have been observed at other ground sites and were rationalized by vertical mixing (Tham et al., 2016; Bannan et al., 2015; Faxon et al., 2015).

In the period after the ClNO\textsubscript{2} morning peak after ~ 09:00, ClNO\textsubscript{2} mixing ratios decreased, coinciding with the increasing ClNO\textsubscript{2} photolysis rate. Box model simulations (see Supplement) indicate that the decay of ClNO\textsubscript{2} (after 09:00) was consistent with its destruction by photolysis.

There were two exceptions: the mornings of 27 July and 2 August, when the decay of ClNO\textsubscript{2} concentration occurred at a rate faster than its photolysis. On 27 July, fog was not observed until 08:00, at which time the ClNO\textsubscript{2} mixing ratio rapidly decreased because of dissolution and/or an air mass shift to one with a different chemical history. On 2 August, the campaign maximum of 97 pptv was observed at 04:40 prior to sunrise, followed by a sharp decline. Hourly logs indicated scattered showers at 06:00.

### 3.1.5 PM\textsubscript{1} size distribution and composition measurements

The time series of PM\textsubscript{1} surface area density (S\textsubscript{A}) observed by the SMPS is shown in Fig. 5a. The aerosol loadings were modest: the average (median) surface area density was 128 (104) \( \mu \text{m}^2 \text{ cm}^{-3} \) and ranged from extremes of 26 to 618 \( \mu \text{m}^2 \text{ cm}^{-3} \). The size distribution data show that bulk of the surface area (i.e., the mean diameter (\( \bar{D}_s \))) is in the range of 200 to 300 nm, such that most of the area of the accumulation mode was captured. However, the surface area calculations do not include contributions from larger diameter particles which were not quantified. Shown on the right-hand side of Fig. 5a is the rate coefficient for heterogeneous uptake of N\textsubscript{2}O\textsubscript{5}, \( k_{\text{N}_2\text{O}_5} \), calculated using Eq. (1).

\[
k_{\text{N}_2\text{O}_5} = \frac{1}{4} \gamma \tau S_A
\]

Here, \( \gamma \) and \( \tau \) are the uptake probability and the mean molecular speed of N\textsubscript{2}O\textsubscript{5}, respectively. Equation (1) is valid for uptake on small, submicron aerosol as it neglects gas-phase diffusion limitations (Davidovits et al., 2006). For this calculation, a \( \gamma \) value of 0.025 was assumed. The average (\( \pm 1 \) standard deviation) of \( k_{\text{N}_2\text{O}_5} \) was \( (2 \pm 1) \times 10^{-4} \text{ s}^{-1} \).

The ACSM submicron aerosol composition data are shown as a time series in Fig. 5b and as a function of time of day in Fig. 6. Consistent with the size distributions, mass loadings were also modest overall (average 2.3 \( \mu \text{g m}^{-3} \)). The ACSM factor analysis identified oxygenated organic aerosol (OOA) as the largest mass fraction of the non-refractory aerosol (average \( \pm \) standard deviation 1.4 \( \pm 1.2 \mu \text{g m}^{-3} \), 63.3\% of the total aerosol mass measured by the ACSM). Hydrocarbon-like organic aerosol (HOA) associated with primary emissions was a minor component (average 0.03 \( \mu \text{g m}^{-3} \), 1.1\%) but occasionally enhanced in plumes (maximum 8.3 \( \mu \text{g m}^{-3} \)). The OOA did not exhibit a discernible diurnal profile (Fig. 6a), which is consistent with the modest photochemistry at this site as judged from the modest peak O\textsubscript{3} levels observed. The inorganic mass fraction was dominated by nitrate (0.47 \( \pm 0.40 \mu \text{g m}^{-3} \), 20.7\%). The second most abundant inorganic component was ammonium (0.21 \( \pm 1.4 \mu \text{g m}^{-3} \), 8.8\%) followed by sulfate (0.15 \( \pm 0.15 \mu \text{g m}^{-3} \), 6.8\%). The data are of similar magnitude as aerosol mass spectrometry (AMS) data collected at nearby Langley as part of Pacific 2001 (Boudries...
et al., 2004); then, organics were also the largest component (average of 1.6 µg m$^{-3}$, 49 %), though sulfate and ammonium mass loadings were larger (0.88 and 0.44 µg m$^{-3}$; 25 and 14 %, respectively) and nitrate mass loadings smaller (0.38 µg m$^{-3}$, 12 %).

The neutralization ratio, \( NR \approx \frac{[\text{NH}_4^+]}{([\text{NO}_3^-] + 2[\text{SO}_4^{2-}])} \) (Zhang et al., 2007), where the square brackets denote molar concentrations (calculated from the mass concentrations reported by the ACSM by dividing by the appropriate molecular weights), was 1.19 (median value). The high NH$_3$ content is qualitatively consistent with the non-quantitative data collected by Metro Vancouver (using a Thermo Scientific 17i NH$_3$/NO/NO$_2$/NOx analyzer), which showed large concentrations of gas-phase NH$_3$ (Fig. S1).

The ACSM software also reported non-refractory chloride with an average ($\pm$1 standard deviation) concentration of 0.01 $\pm$ 0.03 µg m$^{-3}$, though it is unclear whether this signal was real as it did not vary over the course of the campaign and was below the stated ACSM detection limit of 0.2 µg m$^{-3}$ (Ng et al., 2011).

Aerosol nitrate exhibited a clear diurnal profile with higher concentrations at night (Fig. 6b). In particular, the amount of aerosol nitrate increased at the beginning of the night, when the nocturnal NO$_3$ production rates were greatest.

Previous AMS measurements in Vancouver during the month of August as part of Pacific 2001 reported a slightly higher total mass loadings of 7.0 µg m$^{-3}$ that included a greater HOA component (2.4 µg m$^{-3}$, 34 %) and a smaller nitrate fraction (0.6 µg m$^{-3}$, 8.5 %) (Alfarra et al., 2004; Jimenez et al., 2009) than observed here. The lower HOA in this data set is likely a result of tighter emission controls implemented since the earlier study, a topic outside the scope of this paper.

### 3.1.6 Hydrocarbon measurements

Mixing ratios of hydrocarbons were quantified during daytime and during the nights of 2–3 and 3–4 August. A portion of the hydrocarbon data is shown in Fig. 7a. Mixing ratios were generally smaller during the day than during night, due to the larger daytime mixing heights. On the nights of 2/3 and 3/4 August, NO$_3$ was not detected, consistent with low \( P(\text{NO}_3) \) values as \( \text{O}_3 \) mixing ratios approached zero (Fig. 3). At the same time, there were strong NO$_3$ sinks present: mixing ratios of \( \alpha \)-pinene and limonene (left-hand axis) increased throughout the night, as thermal emissions continued into the shallow NBL. In contrast, mixing ratios of isoprene, whose emissions are driven by photosynthesis (Hewitt et al., 2011; Guenther et al., 1995), increased at the beginning of the nights and then decreased as isoprene was removed by oxidation with \( \text{O}_3 \) and NO$_3$ and by transport. Throughout both nights, the site was also influenced by anthropogenic hydrocarbons (e.g., isooctane and toluene, right-hand axis). Because synoptic conditions as judged from local wind speed and direction (Fig. 3d) were similar on most of the other nights when hydrocarbons were not quantified, the data shown in Fig. 7a were likely representative for much of the campaign.

The VOC data were not sufficiently comprehensive to allow an accurate determination of the NO$_3$ loss frequency to hydrocarbons, given by \( k_{\text{NO}_3^+ + \text{VOC},i} / [\text{VOC}] \). Shown in Fig. 7b is the loss frequency of NO$_3$ to isoprene, calculated by multiplying its concentration with the NO$_3$ rate coefficient taken from Seinfeld and Pandis (2006). Loss of NO$_3$ to isoprene was a small sink compared to its loss to NO via Reaction (R3) and NO$_3$ photolysis (Reaction R4) but was approximately on par with its indirect loss, i.e., the heterogeneous uptake of N$_2$O$_5$. 
Figure 6. Diurnal averages of submicron (PM$_{1}$) ACSM data. (a) Organic aerosol displayed as hydrocarbon-like organic aerosol (HOA) and oxygenated organic aerosol (OOA) factors. (b) Inorganic aerosol fractions. (c) Neutralization ratio (NR).

3.2 Analysis

3.2.1 Box model simulations of the nocturnal O$_3$ and O$_x$ loss in the NBL

In initial simulations, the O$_3$ and NO$_2$ deposition rates were tuned until the median nocturnal O$_3$ loss was reproduced. An O$_3$ dry deposition rate of $4 \times 10^{-5}$ s$^{-1}$ produced a simulation that reasonably matched the observations (Fig. S2). The magnitude of this rate corresponds to a NBL height of 50 m, the same mixing height that was frequently observed in balloon vertical profiles reported by Pisano et al. (1997). However, since wind speeds at night were low during the study (median 3.6 km h$^{-1}$), the aerodynamic resistance to vertical transport was likely elevated due to reduced turbulence. It is therefore conceivable that the O$_3$ dry deposition velocity was in actuality smaller than the values taken from Lin et al. (2010) and the mixing height was greater than 50 m.

Modelling studies have assumed N$_2$O$_5$ and NO$_3$ deposition velocities of up to 2 cm s$^{-1}$ in urban areas (Sander and Crutzen, 1996); adopting this value allows the dry deposition rate constants of N$_2$O$_5$ and NO$_3$ to be estimated at $\sim 4 \times 10^{-4}$ s$^{-1}$, which is on par with the estimated heterogeneous uptake rate constant of N$_2$O$_5$ on submicron aerosol.

Next, the generic biogenic VOC was added. For this, a biogenic hydrocarbon abundance of 1 ppbv at sunset (mostly isoprene – see Fig. 7) and a (monoterpene) emission rate of $3 \times 10^5$ molecules cm$^{-3}$ s$^{-1}$ based on the crop emission factor given by Guenther et al. (2012) into a 50 m deep NBL were assumed. This assumed flux gives a similar emission rate as the 0.3 ppbv increase over a 6 h period observed on 3–4 August (Fig. 7). The addition of this biogenic VOC only had a marginal effect on O$_x$ (Fig. S3).

The simulations presented in Fig. S2 underpredict the observed loss of O$_3$, necessitating the addition of an NO source that results in selective removal of O$_3$ while preserving O$_x$. Since automobiles are the largest NO$_x$ source in the region, a constant emission source of 95 % NO and 5 % NO$_2$ (Wild et al., 2017) was added and its magnitude varied. The NO$_x$ source strength necessary to reproduce the median O$_3$ loss was $\sim 1.1$ ppbv h$^{-1}$. The simulation results using these pa-
rameters are superimposed (in red) in Fig. 4c. There is reasonable agreement between the simulations and observations of O$_3$ and O$_2$ until $\sim$03:00 (and between simulation and observation of NO, Fig. S4). This shows that the nocturnal O$_3$ and O$_2$ loss can be rationalized without active NO$_3$ and N$_2$O$_5$ chemistry and suggests that NO$_3$, N$_2$O$_5$, and ClNO$_2$ did not contribute significantly to O$_2$ and O$_3$ loss in the NBL.

### 3.2.2 Metrics of nocturnal nitrogen oxide chemistry: $P$(NO$_3$), $\phi$(ClNO$_2$), and $\tau$(N$_2$O$_5$)

Nocturnal N$_2$O$_5$ chemistry was analyzed using several common metrics: the rate of NO$_3$ production by Reaction (R1), $P$(NO$_3$) = $k_1$[NO$_2$][O$_3$], the yield of ClNO$_2$ relative to the total amount of NO$_3$ formed at night, $\phi$(ClNO$_2$), and the steady-state lifetime of N$_2$O$_5$, $\tau$(N$_2$O$_5$).

The time-of-day dependence of $P$(NO$_3$) is shown in Fig. 8a. The NO$_3$ production rates were small (median values < 0.3 ppbv h$^{-1}$) and were larger during the day than at night due to the low O$_3$ mixing ratios. After midnight, for example, the median $P$(NO$_3$) was (55 ± 23) pptv h$^{-1}$. These are very modest NO$_3$ production rates for a site influenced by urban emissions. In a recent study on a mountain top in Hong Kong, for instance, $P$(NO$_3$) in excess of 1 ppbv h$^{-1}$ was observed in polluted air (Brown et al., 2016).

The median integrated nocturnal NO$_3$ production over the course of the night was 940 pptv (Fig. 8a, right-hand axis), of which 600 pptv was produced before midnight. The amount of ClNO$_2$ produced relative to this amount, $\phi$(ClNO$_2$), was very small (median 0.17 %, maximum 5.4 % on the morning of 2 August) and considerably less than reported by our group for Calgary (median 1.0 %) (Mielke et al., 2016) and Pasadena, CA (median 12 %) (Mielke et al., 2013).

A frequently calculated metric of nighttime nitrogen oxide chemistry is the steady-state lifetimes of NO$_3$ and N$_2$O$_5$, $\tau$(NO$_3$), and $\tau$(N$_2$O$_5$) (Alden et al., 2006; Heintz et al., 1996). The latter is calculated from (Brown et al., 2003; Brown and Stutz, 2012)

$$\tau(\text{N}_2\text{O}_5) = \frac{[\text{N}_2\text{O}_5]}{P(\text{NO}_3)} = \frac{[\text{N}_2\text{O}_5]}{k_1[\text{NO}_2][\text{O}_3]} \approx \left(\frac{k_{N_2O_5}}{k_{\text{N}_2\text{O}_5}} + \frac{k_{\text{NO}_3}}{K_2[\text{NO}_2]}\right)^{-1}. \tag{2}$$

Here, $k_{N_2O_5}$ and $k_{\text{NO}_3}$ are the pseudo-first-order loss-rate coefficients of N$_2$O$_5$ and NO$_3$, respectively, and $K_2$ is the equilibrium constant for equilibrium (Reaction R2).

A central assumption in Reaction (R2) is that NO$_3$, NO$_2$, and N$_2$O$_5$ more rapidly equilibrate than NO$_3$ is formed and either NO$_3$ or N$_2$O$_5$ is destroyed; i.e., NO$_3$ + N$_2$O$_5$ are assumed to be in steady state with respect to production and loss. Brown et al. (2003) outlined potential pitfalls concerning the validity of the steady-state approximation and recommended that box model simulations are carried out to evaluate if a steady state in N$_2$O$_5$ can be assumed. Using the median nocturnal NO$_3$ and O$_3$ mixing ratios of 7.5 ppbv and 18 to 5.0 ppbv, respectively, a temperature of 286 K, and assumed N$_2$O$_5$ and NO$_3$ pseudo-first-order loss frequencies of $1 \times 10^{-3}$ s$^{-1}$ and between $1 \times 10^{-2}$ s$^{-1}$ and 0 s$^{-1}$, the time to achieve steady state in N$_2$O$_5$ is 70 min or less (see Supplement). Thus, the steady-state assumption is reasonable for this data set.

A key parameter in Eq. (2) is the strongly temperature-dependent equilibrium constant $K_2$ (Osthoff et al., 2007). At night, the air temperatures during this study were quite warm (median nocturnal minimum of +13 °C) and did not vary a lot between nights (Fig. 8b). The warm temperatures shift equilibrium (Reaction R2) away from N$_2$O$_5$ and towards NO$_3$ and NO$_2$, making losses via NO$_3$ (Reactions R3–R4, R7) more competitive with the losses of N$_2$O$_5$ (that produce ClNO$_2$; $R$), i.e., the $k_{\text{NO}_3}$ term in Eq. (11) becomes large relative to $k_{\text{N}_2\text{O}_5}$. In contrast, the relatively high NO$_2$ mixing ratios (median value 7.5 ± 0.8 ppbv) shift the equilibrium towards N$_2$O$_5$. Thus, in spite of the relatively warm temperatures, the N$_2$O$_5$:NO$_3$ equilibrium ratios were large on aggregate (> 15; Fig. 8b), enabling ClNO$_2$ formation via Reaction (R5).

The steady-state lifetime of N$_2$O$_5$, $\tau$(N$_2$O$_5$), is shown as a diurnal average in Fig. 8c. The median $\tau$(N$_2$O$_5$) at night was short (~1 min), and the 90th percentile peaked at a modest 7.6 min at sunrise, considerably shorter than observed above the NBL (Brown et al., 2006b) and at other ground sites (Wood et al., 2005; Crowley et al., 2010; Brown et al., 2016).

Superimposed on the right-hand side of Figure 8C are upper limits to the steady-state lifetime of N$_2$O$_5$, calculated using the sum of pseudo-first-order rate coefficients for the titration of NO$_3$ by NO ($k_3$[NO], Reaction R3), NO$_3$ photolysis ($j$[NO$_3$], Reaction R4), and NO$_2$ dry deposition ($k_{\text{dep}}$(NO$_2$)), all divided by the N$_2$O$_5$ over NO$_3$ ratio at equilibrium given by $K_2$NO$_2$ (Fig. 8b), plus the pseudo-first-order rate coefficient for N$_2$O$_5$ heterogeneous uptake ($k_{\text{het}}$(N$_2$O$_5$), Eq. 1) plus N$_2$O$_5$ dry deposition ($k_{\text{dep}}$(N$_2$O$_5$)).

$$\tau(\text{N}_2\text{O}_5) = \left(\frac{k_{\text{NO}_3}}{K_2[\text{NO}_2]} + k_{\text{N}_2\text{O}_5}\right)^{-1} < \left(\frac{k_3[\text{NO}] + j[\text{NO}_3] + k_{\text{dep}}(\text{NO}_2)}{K_2[\text{NO}_2]}\right)^{-1} + k_{\text{het}}(\text{N}_2\text{O}_5) + k_{\text{dep}}(\text{N}_2\text{O}_5)\right)^{-1} \tag{3}$$

The dry deposition rate constants were set to $4 \times 10^{-4} \text{ cm}^2 \text{s}^{-1}$ (see Sect. 3.2.1), which likely overestimates dry deposition during the day due to higher mixing heights; however, the error this introduces is negligible compared to the large daytime sinks such as NO$_3$ photolysis and its reaction with NO. Missing from Eq. (3) are losses of NO$_3$ to hydrocarbons (which were omitted because of the poor VOC data coverage) and terms for NO$_3$ and N$_2$O$_5$ wet (i.e., on cloud and rain droplets) deposition. Periods affected by precipitation or fog (shown in Fig. 3d) were hence excluded from the calcu-
Figure 8. (a) NO$_3$ production rate $P(\text{NO}_3) = k_1\text{[NO]}[\text{O}_3]$ as a function of time of day. The red line is the total amount NO$_3$ generated since sunset, $P(\text{NO}_3)\,dt$. (b) Equilibrium ratio of N$_2$O$_5$ to NO$_3$ calculated by multiplying the temperature-dependent equilibrium constant, $K_2$, with the NO$_3$ concentration, [NO$_3$] (left axis), and air temperature (right axis). (c) Steady-state lifetime of N$_2$O$_5$ (left axis) and upper limits calculated using Eq. (3) (right axis) as functions of time of day.

Figure 9. Scatter plot of ClNO$_2$ mixing ratios with submicron (PM$_1$) ACSM NO$_3$ data. The slopes were calculated for three periods: 2 August, 01:25–04:55 (red dots; slope $= 219 \pm 103$; $\varphi = 0.72$), 23 July, 03:00–04:25 (blue dots slope $= 44 \pm 97$; $\varphi = 0.21$), and 21 July, 02:25–05:20 (purple dots slope $= 12 \pm 17$; $\varphi = 0.06$).

3.2.3 Heterogeneous conversion of N$_2$O$_5$ to ClNO$_2$ on the night of 1/2 August

Phillips et al. (2016) recently applied several methods to estimate the N$_2$O$_5$ uptake parameter ($\gamma$) and yield of ClNO$_2$ ($\varphi$) from ambient measurements of NO$_3$, N$_2$O$_5$, ClNO$_2$, and aerosol nitrate. One of these methods uses the covariance of ClNO$_2$ and aerosol nitrate production rates, $P(\text{NO}_3^-)$ and $P(\text{ClNO}_2)$:

$$\varphi = 2(\frac{P(\text{NO}_3^-)}{P(\text{ClNO}_2) + 1})^{-1}, \quad (4)$$

$$\gamma = 2\left(\frac{P(\text{NO}_3^-)}{P(\text{ClNO}_2)}\right)/(cS_A[N_2O_5]). \quad (5)$$

In the above equations, $c$ is the mean molecular speed of N$_2$O$_5$ ($\approx 237$ m s$^{-1}$). The use of Eqs. (4)–(5) assumes that the relevant properties of the air mass are conserved (i.e., identical upwind of and at the measurement location and affected identically by air masses mixing), that losses of measured species are not significant, that the efficiency of N$_2$O$_5$ uptake and production of ClNO$_2$ and NO$_3$ is independent of particle size, and that partitioning of HNO$_3(g)$ and aerosol nitrate between the gas and particle phases does not occur (Phillips et al., 2016). It is assumed further that production of nitrate from N$_2$O$_5$ uptake on refractory aerosol (that the ACSM does not quantify) is minimal.

In this data set, ClNO$_2$ and non-refractory PM$_1$ nitrate rarely covaried (Fig. 9); the only instance showing a modest correlation ($r = 0.66$) is the time period prior to sunrise of 2 August (shown as red dots in Fig. 9).

The night of 1–2 August exhibited the highest nocturnal nitrogen oxide concentrations for the entire campaign. Winds were initially from the NW and relatively light ($4.8 \pm 0.7$ km h$^{-1}$) and after 01:00 picked up in speed (to $8 \pm 1$ km h$^{-1}$) and shifted to the W. Judging from the HYbrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) back trajectories (Stein et al., 2015), the upwind air
had moved in from the coast, roughly from the direction of the city of Victoria, BC (Odam-ANKrah, 2015).

After sunset at ~21:00 local time, N$_2$O$_5$ levels started increasing and continued to increase until about 01:30 (Fig. 3a). The steady-state N$_2$O$_5$ lifetime at this time was the highest of the campaign, ~10 min. At 01:20, ClONO$_2$ mixing ratio increased from 20.4 pptv at 01:25 to 93.7 pptv at 04:55 and the PM$_1$ content from 0.10 to 0.34 µg m$^{-3}$ (40 to 127 pptv). During this time, N$_2$O$_5$ mixing ratios and PM$_1$ surface area density were relatively constant, 11 ± 6 pptv and 67 ± 4 µg m$^{-3}$ (average ± standard deviation), respectively. The combined amount of N$_2$O$_5$, ClONO$_2$ and PM$_1$ NO$_3$ produced (172 pptv) is less than the amount of NO$_3$ produced from Reaction (R1) which was 519 pptv during this period.

From Eqs. (4) and (5), a ClONO$_2$ yield of $\psi = 0.7 \pm 0.3$ and an N$_2$O$_5$ uptake probability of $\gamma = 0.15 \pm 0.07$ were calculated for this period. Both of these values are upper limits because production of ClONO$_2$ from uptake of N$_2$O$_5$ on unquantified supermicron (i.e., > 0.5 µm) or refractory aerosol (which takes place simultaneously) is not accounted for.

A $\gamma$ value of >0.05 is greater than can be rationalized from laboratory and field studies (Chang et al., 2011) and is hence unrealistic. This suggests that ClONO$_2$ production took place predominantly on supermicron or refractory aerosol, which likely was comprised of mainly sea-salt-derived aerosol (Anlauf et al., 2006). However, if one assumes that “all” of the ClONO$_2$ is produced on supermicron or refractory aerosol such that $P$(ClONO$_2$) on submicron aerosol equals 0 pptv s$^{-1}$ (which is not unreasonable considering the absence of measurable amounts of aerosol chloride in this size fraction; see Sect. 3.1.5), a $\gamma$ value of 0.08 ± 0.04 is calculated. This large value suggests very efficient N$_2$O$_5$ uptake (and conversion to aerosol nitrate) on the non-refractory submicron aerosol that night.

3.3 Impacts of ClONO$_2$ on radical production

Photolysis of ClONO$_2$ increases the rates of photochemical O$_3$ production (and hence worsen air quality) by producing NO$_2$ and reactive Cl atoms (Reaction R6). The amounts of ClONO$_2$ available for photolysis in the morning (median 3.5 pptv at sunrise and 6.8 pptv at 08:00 local time) were too small to have had a measurable impact on local NO$_3$ concentrations (Fig. 3c) but were sufficiently large to, at least occasionally, impact radical budgets.

Figure 10 shows the instantaneous radical production rates of Cl and OH, $P$(Cl) = $j$(ClONO$_2$) $\times$ [ClONO$_2$] and $P$(OH) from reaction of O($^1$D)+H$_2$O. The latter was calculated from an assumed steady state in O($^1$D) with respect to its production from O$_3$ photolysis and reactions with N$_2$, O$_2$, and H$_2$O as described by Mielke et al. (2016). This analysis does not account for OH radical production from photolysis of nitrous acid or aldehydes and, hence, overestimates the importance of Cl radicals.

The largest $P$(Cl) values were observed on 26 July, 07:45 local time (9.5 x 10$^4$ atoms cm$^{-3}$ s$^{-1}$), accounting for 40 % of the total radical production. The largest fraction of radicals produced from ClONO$_2$ photolysis was observed on the same day at 06:35 local time (74 %, 7.8 x 10$^3$ atoms cm$^{-3}$ s$^{-1}$). The photolysis of ClONO$_2$ produces a median value of 6.5 x 10$^3$ atoms cm$^{-3}$ s$^{-1}$ during daytime, which is negligibly small compared to the median $P$(OH) of 3.8 x 10$^6$ molecules cm$^{-3}$ s$^{-1}$ at noon.

4 Discussion

It is now well-established that ClONO$_2$ is an abundant nitrogen oxide in many regions of the troposphere (Table 3). The results presented in this paper are atypical in that they show consistently small ClONO$_2$ mixing ratios in spite of close proximity to sources, i.e., in a region where nearby oceanic emissions of sea salt aerosol and NO$_3$ emissions from a megacity combine. In the following, factors contributing to the low ClONO$_2$ mixing ratios observed in this study and broader implications of ClONO$_2$ in the LFV are discussed.

The main reason for the low ClONO$_2$ mixing ratios observed in this work are the low nocturnal mixing ratios of O$_3$ and small NO$_3$ production rate, $P$(NO$_3$), resulting from the stratification of the boundary layer at night and decoupling of the shallow NBL from the NRL. In the following, it is assumed that a boundary layer structure similar to those observed during PACIFIC 93 (Pisano et al., 1997; McKendry et al., 1997; Hayden et al., 1997) also existed on most measurement nights of this study. Once the nocturnal boundary layer formed at sunset, O$_3$ and O$_x$ in the NBL were rapidly (lifetime of ~4 h) removed. The box model simulations presented in Sect. 3.2.1 show that this removal can be rationalized by dry deposition and titration of O$_3$ with NO and
biogenic VOCs alone, leaving little room for nitrogen oxide chemistry to destroy O$_3$ or NO$_2$, for example, via heterogeneous formation of HONO, which destroys NO$_2$ (Stutz et al., 2004a; Indarto, 2012), or formation of N$_2$O$_5$ and subsequent heterogeneous hydrolysis, which consumes two molecules of NO$_2$ and 1 molecule of O$_3$ (Brown et al., 2006a). It is the often complete absence of O$_3$ at night which distinguishes this data set from the other measurement locations for which CINO$_2$ data have been reported, including continental sites where aerosol chloride is likely less abundant (Table 3).

A compounding factor in this study was the occasional formation of fog and occasional precipitation events. Fog droplets act as a very rapid sink for NO$_3$ and N$_2$O$_5$ (Osthoff et al., 2006), which shuts down CINO$_2$ production, and may have also directly contributed episodically to CINO$_2$ losses, for example on the morning of 27 July. Overall, though, the contribution of fog to CINO$_2$ losses in this data set was minor, as only 5% of the measurement period was impacted by fog. However, this potential CINO$_2$ loss mechanism should be investigated further in future lab studies.

The rapid drop of CINO$_2$ mixing ratio at around 06:00 of 2 August is interesting in that it coincided with a very brief precipitation event. Though an air mass shift cannot be ruled out, this coincidence suggests the possibility that scavenging of CINO$_2$ by rain droplets followed by hydrolysis may be a possible loss pathway. Scavenging of NO$_3$, N$_2$O$_5$, and CINO$_2$ by rain droplets is currently not constrained by laboratory investigations (unlike other gases, such as SO$_2$ or NH$_3$; Hannemann et al., 1995). Similarly to fog, precipitation was not a major factor in this data set as it affected only 10% but may be in other locations or seasons that experience higher rainfall amounts.

An important observation is the lack of non-refractory PM$_1$ chloride (Fig. 5b). This suggests that there was limited redistribution of chloride from acidification of sea salt aerosol onto other aerosol surfaces in this data set. Such a redistribution was observed, for example, during the CalNEX-LA campaign, where the AMS measured a median chloride concentration of ∼0.1 μg m$^{-3}$ on non-refractory aerosol (Mielke et al., 2013). This in turn implies that the submicron aerosol surface did not significantly participate in the production of CINO$_2$ from N$_2$O$_5$ uptake in the NBL, broadly consistent with the conclusions in Sect. 3.2.3 and consistent with measurements of water-soluble aerosol components in the LVF during Pacific 2001 (Anlauf et al., 2006) that showed no evidence for chloride redistribution to PM$_1$ from larger particles where aerosol chloride was present.

The low observed τ(N$_2$O$_5$) is consistent with earlier studies that reported strong vertical gradients in τ(N$_2$O$_5$) due to elevated near-surface sinks from emissions by plants (i.e., monoterpenes) and automobiles (i.e., NO and butadiene; Curren et al., 2006) that titrate NO$_3$ (Stutz et al., 2004b; Wang et al., 2006; Brown et al., 2007; Young et al., 2012). An emblematic example is the study by Wood et al. (2005) at a ground site east of the San Francisco Bay Area in January 2004: They observed relatively modest N$_2$O$_5$ mixing ratios of up to 200 pptv, corresponding to τ(N$_2$O$_5$) < 5 min for the entire study period. Studies for which vertically resolved data were available (e.g., Stutz et al., 2004b; Wang et al., 2006; Brown et al., 2007; Young et al., 2012; Tsai et

Table 3. Maximum CINO$_2$ mixing ratios observed to date.

<table>
<thead>
<tr>
<th>Location</th>
<th>Type</th>
<th>Maximum mixing ratio</th>
<th>Reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Houston, TX</td>
<td>Off-shore, coastal, and inland</td>
<td>1.2 ppbv</td>
<td>Osthoff et al. (2008)</td>
</tr>
<tr>
<td>New England</td>
<td>Off-shore</td>
<td>90 pptv</td>
<td>Kercher et al. (2009)</td>
</tr>
<tr>
<td>Pasadena, CA</td>
<td>Off-shore</td>
<td>2.15 ppbv</td>
<td>Riedel et al. (2012a)</td>
</tr>
<tr>
<td>La Jolla, CA</td>
<td>Coastal</td>
<td>30 pptv</td>
<td>Kim et al. (2014)</td>
</tr>
<tr>
<td>Boulder, CO</td>
<td>Continental</td>
<td>425 pptv</td>
<td>Thornton et al. (2010)</td>
</tr>
<tr>
<td>Calgary, AB</td>
<td>Continental</td>
<td>330 pptv</td>
<td>Mielke et al. (2016, 2011)</td>
</tr>
<tr>
<td>Erie, CO</td>
<td>Continental</td>
<td>1.3 ppbv</td>
<td>Riedel et al. (2013), Brown et al. (2013)</td>
</tr>
<tr>
<td>Feldberg, GER</td>
<td>Continental</td>
<td>800 pptv</td>
<td>Phillips et al. (2012), Phillips et al. (2016)</td>
</tr>
<tr>
<td>Horsepool, UT</td>
<td>Continental</td>
<td>500 pptv</td>
<td>Edwards et al. (2014)</td>
</tr>
<tr>
<td>Pasadena, CA</td>
<td>Coastal, inland</td>
<td>3.5 ppbv</td>
<td>Mielke et al. (2013)</td>
</tr>
<tr>
<td>London, UK</td>
<td>Coastal, inland</td>
<td>724 pptv</td>
<td>Bannan et al. (2015)</td>
</tr>
<tr>
<td>Hongkong, PRC</td>
<td>Coastal, inland</td>
<td>2.0 ppbv</td>
<td>Tham et al. (2014)</td>
</tr>
<tr>
<td>Southeast TX</td>
<td>Coastal, inland</td>
<td>280 pptv</td>
<td>Faxon et al. (2015)</td>
</tr>
<tr>
<td>Hongkong, PRC</td>
<td>Coastal, inland</td>
<td>4.7 ppbv</td>
<td>Wang et al. (2016)</td>
</tr>
<tr>
<td>North China Plain</td>
<td>Continental</td>
<td>2.1 ppbv</td>
<td>Tham et al. (2016)</td>
</tr>
<tr>
<td>North China Plain</td>
<td>Continental</td>
<td>776 pptv</td>
<td>Wang et al. (2017)</td>
</tr>
<tr>
<td>Abbotsford, BC</td>
<td>Coastal, inland</td>
<td>97 pptv</td>
<td>This work</td>
</tr>
</tbody>
</table>
al., 2014) generally showed higher N₂O₅ concentrations and hence larger r(N₂O₅) aloft in the NRL than at the surface. A different scenario likely played out aloft in the NRL, which would exhibit higher NO₃ production rates (via Reactions R1) than the surface layer. Assuming levels of 20 ppbv of O₃ and NO₂ in the NRL (Pisano et al., 1997; McKendry et al., 1997), the NO₃ production rate would equal ~1.1 ppbv h⁻¹ in the NRL, roughly on par with values recently reported for Hong Kong, the current record holder for CINO₂ mixing ratios (Brown et al., 2016; Wang et al., 2016). Recent aircraft and tower studies have shown high rates of production of CINO₂ aloft (Riedel et al., 2013; Young et al., 2012), which likely also occurred in this work.

In contrast, the low mixing height of the NBL is conducive to high levels of biogenic hydrocarbons (Sect. 3.1.6). The nocturnal temperatures during this study were quite warm and did not vary a lot between nights (Fig. 8b). Emissions of monoterpenes, which are reactive towards NO₃, are driven by a temperature-dependent process from storage tissue within the plants at night (Guenther et al., 1995) and, hence, were likely substantial. Their presence is likely responsible for the difference between the “observed” N₂O₅ steady lifetimes, τ(N₂O₅), and upper limit calculated using equation (3) before midnight (Figs. 8c and 8d). Even if one assumes a relatively large uptake probability of γ = 0.025 and accounts for the large ratios of N₂O₅ : NO₃, the loss rate of N₂O₅ on submicron aerosol was likely small in comparison to losses via NO₃ for most of this data set (Fig. 7b). Hence, only a small fraction of the integrated nocturnal NO₃ production of 940 pptv resulted in CINO₂ formation at the surface.

Because of the relatively long lifetime of CINO₂, the breakdown of the surface layer and merging of the surface air with the NRL constituted itself as a CINO₂ “morning peak” in a similar manner as what has recently been reported at other locations (Tham et al., 2016; Bannan et al., 2015; Faxon et al., 2015). This morning peak is rationalized by higher net CINO₂ production in the NRL; the breakup of this layer ~2 h after sunrise then mixes CINO₂ down to the surface. Such a vertical mixing process was not seen during CalNex-LA (Young et al., 2012; Tsai et al., 2014) where the NBL was sufficiently deep to prevent complete O₃ removal and the CINO₂ produced mixed down to the surface at night.

Assuming a 100 m deep NRL where CINO₂ production takes place, a mixed layer height of 500 m by 08:00 (Pisano et al., 1997) and negligible destruction of CINO₂ by photolysis (which is reasonable as the lifetime of CINO₂ with respect to photolysis is >4.6 h at that time of day), a morning increase in CINO₂ mixing ratio by 40 pptv at the surface as seen on the morning of 26 July suggests a pool of CINO₂ in the NRL at sunrise of ~200 pptv, likely a modest value considering that the (assumed) NO₃ production rate may have integrated to ~9 ppbv over the course of the night.

The largest nocturnal CINO₂ mixing ratios were observed on 22–23 July and 1–2 August. Both of these nights exhibited high wind speeds and are counterexamples to what was observed on other nights. We speculate that the higher levels of wind shear and turbulence altered the nocturnal boundary layer structure which exhibited a greater degree of vertical mixing and higher O₃ concentrations at the surface. Consistent with this interpretation and the notion that an isolated NRL with higher net CINO₂ production was absent on those nights, the mornings of 23 July and 2 August did not show a “morning peak”. In contrast, low surface wind speeds were observed on the other nights, facilitating a stable and shallow nocturnal surface layer.

It is conceivable that a land–sea breeze effect transported air from a region closer to the coast that saw higher CINO₂ production than at Abbotsford, i.e., that the CINO₂ morning peaks are generated by horizontal as opposed to vertical transport. Large NO₃ mixing ratios have been reported at Saturna Island (McLaren et al., 2010), which strongly suggest that sizeable reservoirs of CINO₂ form offshore at night. However, it is not known how far inland these reservoirs extend. Considering the average wind speed in the morning (6 km h⁻¹), distance to the coast (35 km), and close proximity (200 m) of the site to the bottom of the polluted NRL with documented high nocturnal pollution levels and early morning down-mixing events, the vertical transport explanation is much more likely correct. Nevertheless, measurements of CINO₂ at a site closer to the coast (e.g., at White Rock) would be beneficial.

Formation of CINO₂ affects air quality through its photolysis which generates O₃, NOₓ, and reactive Cl radicals in the morning, leading to higher net photochemical O₃ production (Sarwar et al., 2014). In spite of the low levels of ClNO₂ observed in this work, the production of radicals from its photodissociation was not always negligible (Fig. 10). Conditions leading to O₃ exceedances did not develop during this study. If such conditions had developed, it is highly likely that this radical generation pathway would have played a much greater role.

The data presented here suggest that higher rates of ClNO₂ and subsequent radical generation take place routinely in layers aloft, processes that are not directly observable at the surface but whose implications are felt as the ultimate product, O₃, is sufficiently long-lived to mix down to the surface (McKendry et al., 1997). Future studies should therefore target the NRL, for example through missed-approaches by aircraft, a blimp, or from a tall tower, especially during episodes of a developing O₃ exceedance event and also include composition measurements of refractory aerosol.

5 Summary and conclusions

In this paper, we have presented the first measurements of CINO₂ and N₂O₅ mixing ratios in the LFV. In spite of the close proximity to NOₓ (Greater Vancouver) and sea salt aerosol (the Pacific Ocean) sources, CINO₂ and N₂O₅ mixing ratios were small (maximum of 97 and 27 pptv, respec-
tively) and smaller than observed at other measurement locations for which CINO$_2$ abundances were reported. The low mixing ratios are explained through the removal of O$_3$ by deposition and titration with NO in a shallow nocturnal surface layer. Measurements of submicron aerosol composition by ACSM showed no enhancements of particle-phase chloride, which is in contrast to locations where high CINO$_2$ mixing ratios were observed (such as Pasadena; Mielke et al., 2013) and indicates that there was little processing and redistribution of sea-salt-derived chloride at this location. There is indirect evidence that higher production of CINO$_2$ took place above the measurement site in the NRL, observed via down-mixing after the breakup of the NBL in the morning, and highlights the need for future vertically resolved measurements (e.g., from an aircraft platform) of CINO$_2$ and N$_2$O$_5$ mixing ratios in the LFV. Conditions leading to O$_3$ exceedances did not develop during the relatively short measurement period of 2 weeks, such that the full impact that nocturnal formation of CINO$_2$ could have on radical production and NO$_2$ recycling remains unquantified.

**Data availability.** The data used in this study are available from the corresponding author upon request (hosthoff@ucalgary.ca).

**Supplement.** The supplement related to this article is available online at: https://doi.org/10.5194/acp-18-6293-2018-supplement.

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

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