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1 Introduction

Tropospheric ozone (O\textsubscript{3}) is important as it affects air quality and is a greenhouse gas. The Arctic has been warming at twice the global average rate over the past century (IPCC, 2007). While increases in long-lived greenhouse gases dominate Arctic warming, O\textsubscript{3} and other short-lived pollutants (e.g., aerosols) could also play an important role (Law and Stohl, 2007; Shindell, 2007; Quinn et al., 2008). Changes in local tropospheric O\textsubscript{3} affect the Arctic climate by altering local radiation fluxes with maximum impact near the tropopause (Hansen et al., 1997). A recent modeling study suggests that an increase in tropospheric O\textsubscript{3}, caused by increases in anthropogenic emissions, could have contributed about 0.3 °C annually to the 20th-century Arctic surface warming and about 0.4 °C–0.5 °C during winter and spring (Shindell et al., 2006). The impact of possible increases in boreal forest fire emissions and changes in stratospheric O\textsubscript{3} flux to the troposphere on Arctic surface warming are not yet well quantified.

Ozone is produced locally in the Arctic troposphere from its precursors (i.e., carbon monoxide (CO), hydrocarbons, nitrogen oxides (NO\textsubscript{x})) emitted from anthropogenic and biomass burning sources in adjacent continents (e.g., Penkett and Brice, 1986; Wofsy et al., 1992; Beine et al., 1997). Additional potential sources of O\textsubscript{3} in the Arctic troposphere include transport of O\textsubscript{3} from lower latitudes (Shindell et al., 2008) as well as transport from the stratosphere (Dibb et al., 2003; Allen et al., 2003). Stratospheric air contains high NO\textsubscript{x} and nitric acid (HNO\textsubscript{3}) and is also an important source of NO\textsubscript{x} when injected into the Arctic troposphere (Wofsy et al., 1992; Levy et al., 1999; Law and Stohl, 2007; Liang et al., 2009). NO\textsubscript{x} of stratospheric origin is the driving mechanism that leads to enhanced O\textsubscript{3} production in the Arctic upper troposphere (Liang et al., 2009). A better quantification of the contribution of various anthropogenic and natural sources to O\textsubscript{3} in the Arctic is important for understanding the temporal variation and radiative impact of O\textsubscript{3}, and how Arctic O\textsubscript{3} may change as climate warms and the stratospheric O\textsubscript{3} layer recovers. The NASA Arctic Research of the Composition of the Troposphere from Aircraft and Satellites (ARCTAS) mission was conducted in April and June–July 2008 (Jacob et al., 2010). Its goal was to better understand the factors driving changes in Arctic atmospheric composition and climate. The extensive and detailed measurements of O\textsubscript{3} and reactive nitrogen (NO\textsubscript{y}) species provide a great opportunity to examine the photochemistry of O\textsubscript{3} and NO\textsubscript{x}, and their sources in the Arctic. In this paper, we will use observations obtained onboard the NASA DC-8 aircraft during ARCTAS to examine O\textsubscript{3} and NO\textsubscript{x} in the Arctic and sub-Arctic region and their source attributions. Section 2 describes the observations used in this study. We use a set of marker tracers to identify various air masses sampled during ARCTAS and examine their chemical composition, as described in Sect. 3. NO\textsubscript{x} plays a determinative role in O\textsubscript{3} production in the troposphere. Therefore to better understand which sources contribute to O\textsubscript{3} in the Arctic troposphere, it is important to understand sources of NO\textsubscript{x} and the long-lived reservoir species of NO\textsubscript{y}, i.e., HNO\textsubscript{3} and peroxyacetyl nitrate (PAN). We examine NO\textsubscript{x} and its partitioning in various air masses in Sect. 4, followed by an analysis of O\textsubscript{3}, O\textsubscript{3} production and its dependence on NO\textsubscript{x} and HO\textsubscript{x} (OH+HO\textsubscript{2}) levels within individual air masses sampled during ARCTAS (Sect. 5). Conclusions are presented in Sect. 6.

2 Observations and model

The NASA ARCTAS mission had two phases. The spring deployment (ARCTAS-A), based in Fairbanks Alaska, involved nine flights by the NASA DC-8 aircraft between 1 April and 21 April 2008. The summer deployment (ARCTAS-B) took place between 26 June and 14 July 2008 (nine flights) and was operated from a base in Cold Lake, Canada. Figure 1 shows the geographical distribution of flight tracks of the DC-8 aircraft during ARCTAS. Here we use measurements obtained north of 50° N. The color symbols indicate the location of various air masses sampled during ARCTAS. Tracks not marked with color symbols indicate background atmosphere.
Table 1. Summary of ARCTAS observations used in this study.

| Species | Instrument & Methods | Reference |
|---------|----------------------|-----------|
| CO, NO, NO₂, NOy | Tunable Diode Laser Absorption Spectroscopy (TDLAS) | Diskin et al. (2002) |
| O₃, NO, NO₂, NOy, NOy, HNO₃ | Chemiluminescence | Weinheimer et al. (1994) |
| PAN | Chemical Ionization Mass Spectrometry (CIMS) | Slusher et al. (2004) |
| Alkyl nitrates | Thermal-Dissociation Laser Induced Fluorescence (TD-LIF) | Cleary et al. (2002) |
| OH, HO₂ | Laser Induced Fluorescence (LIF) | Brune et al. (1999) |
| CH₃CN* | Proton Transfer Reaction – Mass Spectrometry (PTR-MS) | Wisthaler et al (2002) |
| CFC-113, C₂H₆ | Whole Air Sampler – Gas Chromatography (WAS-GC) | Apel et al. (2003) |
| | Gas Chromatography – Mass Spectrometry (GC-MS) | Apel et al. (2003) |

* Multiple sets of measurements were available for several species used in this study, i.e., NO₂, OH, HO₂, HNO₃. The different measurements broadly agree with each other and the choice of measurements does not affect the conclusion of this study.

** Two sets of measurements were available for CH₃CN: (i) the PTR-MS measurement of CH₃CN is available every 8-s and 60-s, respectively, and (ii) the GC-MS measurements available every 120-s. In general, we use CH₃CN measured by PTR-MS. During time periods when there are no available PTR-MS measurements, we use the GC-MS measurements whenever possible.

Observations obtained onboard the DC-8 aircraft include O₃, HOₓ, NOₓ, as well as NOy reservoir species, hydrocarbons, halocarbons, aerosols (Jacob et al., 2010). Segregation between various air masses relies on the availability of simultaneous measurements of the marker tracers, e.g., CO for combustion plumes, acetonitrile (CH₃CN) for biomass burning and chlorofluorocarbons (CFCs) for stratospheric air. A detailed list of the species used in this study and the associated instrument specifications is presented in Table 1. Multiple merge files (1-s, 10-s, 60-s) were created for the ARCTAS measurements. Here, we rely on the 60-s merge. Although many species are available at higher frequency, measurements crucial to this analysis, i.e., halocarbons, from the Whole Air Sampler – Gas Chromatography, were only obtained every 160-s.

We also use results calculated by the NASA Langley box model (Olson et al., 2004) constrained by chemical and physical parameters measured by the DC-8 aircraft. Observed O₃, CO, NO, temperature, J(NO₂) and J(O₃) from the 60-s merge were used as model input. In addition, model calculations have been constrained by observed values of many trace gases, including H₂O₂, CH₃OOH, HNO₃, PAN, acetone, MEK, methanol, and ethanol when possible.

3 Air masses observed during ARCTAS

3.1 Air sampled during ARCTAS

While airborne field missions provide an extensive set of trace gas measurements over vast spatial regions, the flight plans are usually designed to target pollution plumes and thus biased towards these plumes. Using CO, a commonly used tracer for combustion and atmospheric transport, we analyze the representativeness of the ARCTAS sampling to the general characteristics of the Arctic troposphere.

We analyze the probability density function (PDF) of CO observed during ARCTAS-A (Fig. 2). The PDF of CO displays a unimodal distribution in the lower and mid troposphere during spring with peaks at 160 ppbv and 145 ppbv, respectively, implying a relatively well-mixed Arctic atmosphere. In the upper troposphere/lower stratosphere (UT/LS), the distribution is bimodal, with one peak at 125 ppbv and a secondary peak at ~50 ppbv representing tropospheric and stratospheric air masses, respectively. The PDF during ARCTAS-B (Fig. 2) displays multiple peaks in the troposphere. The primary peak around 100 ppbv...
(90–120 ppbv) represents the background atmosphere and the two peaks between 120–160 ppbv (present in the upper and lower troposphere) and >160 ppbv (present in the mid- and upper troposphere) are associated with either anthropogenic and/or biomass burning pollution. Acetonitrile is typically used as a tracer for biomass burning plumes (Lobert et al., 1990; Holzinger et al., 2001). The peak with CO > 160 ppbv has mean CH$_3$CN of 520 pptv, indicating these are mostly biomass burning plumes. The other pollution peak with CO between 120–160 ppbv has relatively low level of CH$_3$CN (200 pptv), suggesting these measurements are mostly anthropogenic pollution plumes. The fact that the combustion peaks are well separated from the background suggests these are fresh pollution plumes that have not yet mixed into the background. The extended tails of combustion plumes during summer implies that, unlike spring, the summertime sampling is highly biased towards pollution plumes.

### 3.2 Air mass identification

We use a comprehensive set of tracers to characterize air masses sampled by the DC-8 aircraft during ARCTAS. The detailed criteria applied to define each type of air mass are listed in Table 2. Note that the thresholds of the marker gases chosen to segregate air masses of different origin are highly subjective and can vary significantly depending on season, location, and the question of interest. While we choose some criteria based on previous literature (O$_3$ > 100 ppbv for air of stratospheric origin) and the PDF distribution of CO (Sect. 3.1) for combustion plumes, we heavily rely on tracer-tracer correlations for optimal segregation between different air masses (see Supplement, Figs. S1 and S2). We found that the CO-NO$_x$, CO-CO$_2$ and CO-CH$_4$ correlations are particularly useful in determining the threshold levels of markers for distinguishing air in the stratosphere, air associated with recent stratosphere-troposphere-exchange (STE), biomass burning and anthropogenic plumes.

We use CO and CH$_3$CN to distinguish anthropogenic and biomass burning pollution plumes. Since pollution plumes are not well separated from the background during spring (Sect. 3.1), we use the highest quartile of CO (>160 ppbv) to define pollution plumes. Within the pollution plumes, air masses with CH$_3$CN > 145 pptv are identified as biomass burning plumes and the remaining as anthropogenic pollution plumes. During summer, air masses with CO > 120 ppbv are defined as combustion plumes (Table 2). We further use CO > 160 ppbv and CH$_3$CN > 320 pptv to separate biomass burning air masses from anthropogenic plumes. The thresholds of CH$_3$CN ～145 pptv for ARCTAS-A and ～320 pptv for ARCTAS-B are chosen for optimal segregation between the biomass burning and anthropogenic pollutants based on the CO/$\text{CO}_2$, CH$_4$/CO, and C$_2$H$_6$/CO ratios (Table 2), which differ in these two types of air masses (see Supplement, Figs. S1 and S2).

Air in the stratosphere is enriched in O$_3$ and depleted in surface emitted pollutants such as long-lived CFCs (lifetime ～45–100 yr) as well as short-lived CO (lifetime ～two months). Stratospheric air can enter the troposphere through
Table 3a. Mean observed chemical composition of air masses sampled during ARCTAS-A\(^a\).

|                      | Background | Anthropogenic | Biomass | Stratosphere | STE |
|----------------------|------------|---------------|---------|--------------|------|
|                      | 2370 min   | Pollution     | Burning |              |      |
|                      | (59 %)\(^b\) | (17 %)        | (4 %)   | (9 %)        | (3 %) |
| 0–12 km\(^c\)       |            |               |         |              |      |
| CO (ppbv)            | 144 ± 14   | 172 ± 14      | 220 ± 42| 48 ± 14      | 100 ± 14 |
| O\(_3\) (ppbv)       | 63 ± 16    | 78 ± 12       | 363 ± 122| 150 ± 31     |      |
| OH (pptv)            | 0.04 ± 0.04| 0.02 ± 0.03    | 0.04 ± 0.03| 0.05 ± 0.04 |      |
| HO\(_2\) (pptv)      | 3.5 ± 1.9  | 3.2 ± 1.5     | 6.4 ± 4.1| 1.0 ± 0.4    |      |
| HO\(_x\) (pptv)      | 3.5 ± 1.9  | 3.2 ± 1.5     | 6.7 ± 4.2| 1.1 ± 0.4    |      |
| NO (pptv)            | 11 ± 28    | 32 ± 372      | 18 ± 17 | 50 ± 24      |      |
| NO\(_2\) (pptv)      | 6 ± 38     | 24 ± 211      | 33 ± 22 | 75 ± 29      |      |
| NO\(_x\) (pptv)      | 25 ± 65    | 65 ± 630      | 50 ± 40 | 150 ± 55     |      |
| PAN (pptv)           | 205 ± 80   | 345 ± 145     | 910 ± 475| 70 ± 30      |      |
| HNO\(_3\) (pptv)     | 30 ± 35    | 25 ± 30       | 40 ± 40 | 1470 ± 575   |      |
| ANs (pptv)           | NA         | NA            | NA      | NA           |      |
| NO\(_y\) (pptv)      | 410 ± 170  | 650 ± 660     | 1725 ± 955| 2035 ± 660   |      |

\(^a\) For each type of air mass we include the observed mean ± one standard deviation. Chemical species that are significantly enhanced (> mean + one standard deviation) with respect to background at the corresponding altitude are highlighted in bold. \(^b\) The percentage sum of all identified air masses equals to 93 % and the remaining 7 % are ozone depleting events. \(^c\) The altitude span of individual air masses.

rapid synoptic eddy exchange activities, e.g., tropopause folds, or slow global-scale diabatic descent (Holton et al., 1995). The stratosphere-to-troposphere transport time ranges between a few days during rapid tropopause folding events that intrude deeply into the troposphere to the order of a month for shallow STE intrusions followed by subsequent slow diabatic descent. The difference in transport time can lead to significantly different levels of trace gases, in particular the short-lived species, such as O\(_3\), HNO\(_3\), and Be-7 (Liang et al., 2009). We use the combination of a short-lived tracer, O\(_3\) (>100 ppbv), and a long-lived tracer, CFC-113 (lowest quartile, <78 pptv) to identify air of stratospheric origin. We choose CFC-113 over the other two more common CFCs, CFC-11 and CFC-12. This is because emissions of CFC-113 have significantly decreased since 2000 (Liang et al., 2008), due to the phase-out required by the Montreal Protocol. The variation in CFC-113 is less tempered by variability due to tropospheric transport of fresh surface emissions, which can be significant, and reflects mainly the extent of mixing with stratospheric air. Therefore low CFC-113, together with high O\(_3\), is a better marker to distinguish air transported from the stratosphere. We apply an additional criterion, CO < 160 ppbv in spring (<120 ppbv in summer), to exclude any samples that have mixed to some extent with fresh combustion plumes. We also use CO levels to distinguish air that is of stratospheric origin, but has already penetrated into the troposphere through STE events (CO > 80 ppbv and >50 ppbv in spring and summer, respectively), from the air that still resides in the lowermost stratosphere (Table 2). This is because air of stratospheric origin can have very different NO\(_x\) partitioning and photochemical properties, e.g., O\(_3\) production rates, when it enters the troposphere and mixes with the tropospheric background, as compared to air that remains in the stratosphere. Note that the use of O\(_3\) > 100 ppbv for stratosphere-troposphere mixed air masses is a stringent criterion that distinguishes only the relatively fresh STE events from the background atmosphere.

The DC-8 aircraft also encountered a few deep convective events during ARCTAS-B. Air masses that have recently experienced deep convection contain enhanced levels of NO\(_x\) associated with freshly-ventilated air from the boundary layer and/or lightning and are depleted in HNO\(_3\) due to scavenging (e.g., Thompson et al., 1999; Liang et al., 2007). Thus we define air as being influenced by convection/lightning when NO\(_x\) exceeds 200 pptv and the NO\(_x\)/HNO\(_3\) ratio exceeds >1.2 pptv pptv\(^{-1}\). During ARCTAS-A, six minutes (<0.1 % of a total ~4200 min) of the data contain elevated NO\(_x\) (>100 pptv), which were of neither anthropogenic/biomass burning nor stratospheric origin. Since deep convection is not common during the high latitude spring, these measurements are most likely tied to fresh aircraft exhaust. We therefore exclude these air samples.

The remaining air masses are defined as background. Note that the DC-8 measurements in the Arctic marine boundary layer also include a few O\(_3\) depletion events (O\(_3\) < 30 ppbv) during spring (Neuman et al., 2010) as well as local plumes with high NO\(_x\) from coastal ship emissions in spring and Canadian power plants near Edmonton and Ft. McMurray in summer. We exclude these data in this analysis.
3.3 Air mass composition

A summary of the air mass composition sampled by the DC-8 aircraft is shown in Table 3a (for ARCTAS-A) and Table 3b (for ARCTAS-B). About 59% of the spring measurements are from the background troposphere. Pollution plumes account for 21% of the observations, 17% for anthropogenic pollution and 4% for biomass burning plumes. About 9% and 3% of the spring measurements, respectively, are of lowermost stratospheric air and air influenced by recent STE events. During ARCTAS-B, about 38% of the data are identified as fresh anthropogenic pollution and about 11% are attributed to fresh biomass burning plumes. However, as we discussed in Sect. 3.1, the ARCTAS-B measurements are biased towards combustion plumes and thus the above fractions are not representative of the general Arctic troposphere. Stratospheric air and STE together account for ~5% of the measurements. About 2% of the air sampled during ARCTAS-B was recently influenced by convection and/or lightning. Geographically, the majority of the convective and biomass burning plumes are located in the sub-Arctic between 50–70° N while anthropogenic and stratosphere-troposphere mixed air masses are found throughout the Arctic and sub-Arctic (Fig. 1).

The background Arctic troposphere during spring has mean CO concentrations of ~145 ppbv, O₃ of ~60 ppbv, and NOₓ of ~25 pptv (Table 3a). Background CO decreases with altitude (Fig. 3a, Table 3a), suggesting that pollution is mainly mixed into the background and trapped at low altitudes. Background O₃ and NOₓ remain relatively the same from spring to summer, but CO levels decrease to ~100 ppbv due to increased destruction by OH (Table 3b). Unlike in spring, CO in summer shows little dependence on altitude (Fig. 3b), indicating efficient vertical mixing.

Extensive aircraft measurements of the Arctic troposphere were available from the earlier Tropospheric O₃ Production about the Spring Equinox (TOPSE) campaign in spring 2002 (Atlas et al., 2003) and the Arctic Boundary Layer Expedition (ABLE 3A) during summer 1988 (Harriss et al., 1992). Measurements from these previous missions show springtime mean CO ~154 ppbv, O₃ ~67 ppbv, and NOₓ ~17 pptv (TOPSE) (Stroud et al., 2003) and summertime mean CO ~100 ppbv, O₃ ~70 ppbv, and NOₓ ~10–50 pptv (ABLE 3A) (Jacob et al., 1992) at 3–6 km in the Arctic mid-troposphere. Considering the likely variations associated with differences in air mass sampling and interannual variability, the ARCTAS measurements indicate that these important tropospheric trace gases, CO, NOₓ and O₃, have remained relatively unchanged in the Arctic mid-troposphere in the past two decades, despite the significant changes in processes that could have had a notable impact on the Arctic atmospheric composition, e.g., emissions regulation in Europe and N. America, rapid industrialization in East Asia, and destruction of the stratospheric O₃ layer.

The lowermost stratosphere, with low CO, can reach as low as 6 km during spring, likely during low tropopause events (Fig. 3a). Significantly fewer samples of the lowermost stratospheric air (1%) were sampled during summer at >10 km (Fig. 3b). This is consistent with the seasonal growth of tropopause height from spring to summer. Frequent STE events have been observed throughout spring and summer. Air masses associated with fresh STE events are present at altitudes >5 km (Fig. 3). Stratosphere-troposphere mixed air masses have higher CO, compared to air in the lowermost

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**Table 3b.** Same as Table 3a but for ARCTAS-B.

| Background | Anthropogenic Pollution | Biomass Burning | Stratosphere | STE | Convection |
|------------|-------------------------|-----------------|--------------|-----|------------|
| 1417 min   | 1232 min                | 342 min         | 32 min       | 131 min | 59 min     |
| (44 %)     | (38 %)                  | (11 %)          | (1 %)        | (4 %) | (2 %)      |
| 0–12 km    | 103 ± 11                | 103 ± 10        | 104 ± 10     | 102 ± 12 | 153 ± 39   | 415 ± 280 | 30 ± 6 | 91 ± 17 | 143 ± 28 |
| 0–3 km     | 57 ± 21                 | 34 ± 6          | 60 ± 15      | 70 ± 22 | 58 ± 18    | 49 ± 17   | 448 ± 48 | 164 ± 51 | 79 ± 11 |
| 0–6 km     | 0.13 ± 0.08             | 0.07 ± 0.06     | 0.16 ± 0.08  | 0.14 ± 0.06 | 0.11 ± 0.08 | 0.09 ± 0.07 | 0.09 ± 0.03 | 0.18 ± 0.07 | 0.23 ± 0.18 |
| 0–12 kmb   | 25 ± 30                 | 25 ± 35         | 20 ± 20      | 30 ± 25 | 95 ± 355   | 635 ± 1490 | 385 ± 50 | 110 ± 60 | 505 ± 350 |
| 0–10 kmb   | 210 ± 100               | 105 ± 55        | 230 ± 80     | 245 ± 95 | 350 ± 150  | 950 ± 660 | 70 ± 50 | 320 ± 45 | 415 ± 60 |
| 10–12 kmb  | 70 ± 85                 | 90 ± 105        | 80 ± 85      | 45 ± 60 | 85 ± 90    | 70 ± 70   | 1740 ± 330 | 515 ± 355 | 25 ± 20 |
| 6–12 kmb   | 20 ± 50                 | 40 ± 70         | 15 ± 30      | 10 ± 40 | 55 ± 90    | 195 ± 255 | NA     | 50 ± 65 | 105 ± 80 |
| 6–12 kmb   | 320 ± 165               | 245 ± 200       | 310 ± 140    | 370 ± 150 | 585 ± 445  | 2020 ± 2175 | 2210 ± 260 | 955 ± 355 | 1095 ± 355 |

*For each type of air mass we include the observed mean ± one standard deviation. Chemical species that are significantly enhanced (>mean ± one standard deviation) with respect to background at the corresponding altitude are highlighted in bold. b The altitude span of individual air masses.
stratosphere, reflecting mixing with tropospheric background air during stratosphere-to-troposphere transport.

The convective air masses observed during summer contain elevated CO (50% enhancement as compared to background) (Table 3b), indicating freshly ventilated surface pollution. Anthropogenic pollution plumes are present from the surface to the upper troposphere throughout spring and summer and contain elevated CO (∼170 ppbv in spring and ∼150 ppbv in summer) (Fig. 3). Biomass burning plumes are confined in the mid troposphere during spring with a moderate increase in CO (∼220 ppbv) (Fig. 3a, Table 3a). The majority of the biomass burning air masses sampled during summer are fresh fire plumes in the lower troposphere with marked high CO (∼415 ppbv) (Fig. 3a, Table 3b). More detailed analysis on how anthropogenic pollution and Siberian (spring phase) and Canadian (summer phase) fire emissions impact atmospheric gas and aerosol composition and O3 production can be found in Singh et al. (2010) and Alvarado et al. (2010).

4 Reactive nitrogen in the Arctic troposphere

The abundance of NOx plays a determinative role in O3 production in the background troposphere (Lin et al., 1988; Stillman et al., 1990; Jaeglé et al., 1998; Wennberg et al., 1998). While NOx is present in the background atmosphere at low levels, it can be recycled between the radical forms and its long-lived reservoir species, which adds complexity to an accurate understanding of the NOx budget in the atmosphere. We analyze NOx (NOx+PAN+ HNO3+nitrates) and its partitioning during ARCTAS to investigate the budget and source attribution of NOx in the Arctic and sub-Arctic troposphere. It is difficult to quantify the actual contribution of a certain source to reactive nitrogen species (same for O3 in Sect. 5) just based on observations. Therefore we examine the concentration of nitrogen species in individual air masses with respect to the background since the level of elevated concentration (shown in below as Δ values relative to the background concentrations) in an individual air mass reflects its potential as a source of nitrogen species.

4.1 Reactive nitrogen in various air masses

Reactive nitrogen in the background troposphere remains relatively constant from spring to summer (∼300–400 pptv) (Table 3 and Figs. 4 and 5). Nitrogen oxides (∼25 pptv) on average account for 5–10% of NOx. PAN is the largest reservoir species (∼200 pptv), accounting for 50% of NOy in spring and ∼70% in summer. The level of HNO3 is significantly lower than that of PAN, ∼30 pptv in spring and ∼70 pptv in summer. A small fraction of NOy (∼6%) is present as alkyl nitrates during summer.

The main sources of NOy in the troposphere at high latitudes are STE, and anthropogenic and biomass burning emissions (Fig. 4 and Table 3a). Combustion plumes are the major contributors of NOy in the middle troposphere mainly in the form of PAN and NOx, but little HNO3. Air influenced by the stratosphere on average contains the highest level of NOy above 6 km. Air masses associated with STE contain elevated levels of NOx and HNO3. Compared to air in the lowermost stratosphere, they contain much less NOy (40% of that in the lowermost stratosphere) and different NOy partitioning (less HNO3 and more PAN).

All sources, including anthropogenic and biomass burning emissions, convection, and STE contribute to NOy in the Arctic/sub-Arctic troposphere during summer (Fig. 5 and Table 3b). The NOy vs. CO relationship is more dispersed
during summer, compared to a clear and compact correlation in individual air masses in spring, implying more mixing among air masses of different origin. Biomass burning air masses contain high concentrations of NO\textsubscript{x}, PAN, and alkyl nitrates (AN) (\(\Delta\text{NO}_x\sim 600\) pptv, \(\Delta\text{PAN}\sim 750\) pptv, \(\Delta\text{ANs}\sim 200\) pptv) and is the dominant contributor to NO\textsubscript{y} (\(\Delta\text{NO}_y\sim 1700\) pptv) in the mid and lower troposphere. Anthropogenic emissions also contribute, but their impacts are much less pronounced (\(\Delta\text{NO}_y\sim 250\) pptv, \(\Delta\text{NO}_3\sim 70\) pptv, \(\Delta\text{PAN}\sim 150\) pptv, and no elevated ANs and HNO\textsubscript{3}). In the upper troposphere (>6 km), convection, STE, and biomass burning all contribute significantly to NO\textsubscript{y}. Convection is the dominant source of NO\textsubscript{x} (\(\Delta\text{NO}_x\sim 600\) pptv) while biomass burning pollution is the dominant contributor to PAN. Air masses influenced by STE contain high NO\textsubscript{y}, comparable to that in convective air masses. They display high NO\textsubscript{x} (\(\Delta\text{NO}_x\sim 100\) pptv) and HNO\textsubscript{3}(\(\Delta\text{HNO}_3\sim 400\) pptv) as stratospheric air is commonly enriched with NO\textsubscript{x} and HNO\textsubscript{3}. They also have significantly elevated in PAN (50% more than the background), with a mean concentration (320 pptv) almost comparable to that in anthropogenic plumes (350 pptv).

4.2 PAN in air masses influenced by STE

Using CO as a proxy for transport and air mass inter-mixing, we examine how mixing ratios of reactive nitrogen species change as air of stratospheric origin mixes with tropospheric background air during STE events (Parrish et al., 1998). As it mixes with tropospheric air, an air parcel of stratospheric origin moves along the mixing line (thick green dashed lines) in a scatter plot (Fig. 6). (We refer to tropospheric air masses influenced by stratospheric air as “stratosphere-troposphere mixed air masses” hereafter.) When active chemical production and loss of NO\textsubscript{y} species occurs, the air parcel deviates from the mixing line. Wet scavenging adds some complexity as it is a significant loss of HNO\textsubscript{3} (therefore NO\textsubscript{y}) in the troposphere in summer and thus lowering the mean concentrations of HNO\textsubscript{3} and NO\textsubscript{y} in the tropospheric background air. The majority of the observed mixing ratios of NO\textsubscript{x}, HNO\textsubscript{3}, NO\textsubscript{y}, and the springtime PAN in the stratosphere-troposphere mixed air is within the envelope of variations that can be explained by mixing alone (thin gray dashed lines). The observed mean PAN in the summertime stratosphere-troposphere mixed air is 320 pptv.
Assuming that the mean CO of ∼90 ppbv in the stratosphere-troposphere mixed air masses results from mixing stratospheric air with tropospheric background, PAN can increase to ∼220 pptv assuming PAN and CO increase proportionally. Although the air masses that we identified as being influenced by STE display low levels of CO (<160 ppbv for spring and <120 ppbv for summer), it is possible that these air masses might experience some mixing with combustion plumes, which are considered the main sources of PAN in the atmosphere (e.g., Singh et al., 1992; Alvarado et al., 2010). Air masses influenced by convection sampled during ARCTAS-B also display elevated levels of PAN (Table 3b). Mixing with these tropospheric plumes can potentially increase the level of PAN. However, the mixing lines between stratospheric air and anthropogenic, biomass burning, and convective air masses between 6-12 km fall closely to that of the background air and do not help much in explaining the observed high level of PAN in at least half of the stratosphere-troposphere mixed air masses.

What could have contributed to the high PAN observed in air influenced by STE during ARCTAS in summer? One possible explanation is active photochemical production and conversion of nitrogen from HNO$_3$ to PAN. In our earlier modeling study, Liang et al. (2009), we conducted a detailed budget analysis of NO$_x$ in the Arctic using the GMI CTM which contains a fully-coupled tropospheric and stratospheric chemistry scheme. The model results suggest that significant conversion occurs of nitrogen from HNO$_3$ to NO$_x$ and then to PAN as air of stratospheric origin mixes into air in the upper troposphere in the Arctic during summer.

We now analyze the ARCTAS-B measurements together with box model results to examine if indeed there exists active conversion of nitrogen from HNO$_3$ to PAN in the stratosphere-troposphere mixed air masses. Fig. 7 shows the concentrations of several trace gases in a stratosphere-troposphere mixed plume (low CO and high O$_3$) sampled by the DC-8 aircraft on 9 July 2008. This is a relatively fresh stratosphere-troposphere mixed plume with high NO$_x$, HNO$_3$, and medium PAN (∼300 pptv). The DC-8 aircraft sampled another stratosphere-troposphere mixed plume on 8 July 2008 (Fig. 8). This plume contains two subplumes: A and B. Subplume A is a fresh stratosphere-troposphere mixed plume with chemical composition similar to that of the 9 July plume (high NO$_x$, HNO$_3$, and medium PAN). Subplume B is an aged stratosphere-troposphere mixed plume with relatively low NO$_x$, HNO$_3$, and high PAN (maximum
Fig. 6. Scatter plots of NO$_x$, PAN, HNO$_3$, and NO$_y$ vs. CO in the background air between 6-12 km (black dots), stratospheric air (purple squares), and stratosphere-troposphere mixed air (lilac crosses) for ARCTAS-A (top panels) and ARCTAS-B (bottom row). The mean mixing ratios within each air mass is plotted (red squares) with horizontal bars indicating one standard deviation of CO and vertical bars for one standard deviation of the corresponding reactive nitrogen species. This figure demonstrates how trace gas mixing ratios change as air of stratospheric-origin mixes with background air during STE. Note in panel (h) we have added the biomass burning, anthropogenic and convective air masses between 6–12 km as well to demonstrate the impact on PAN in the stratosphere-troposphere mixed air if mixing with these tropospheric air masses occurs. For air that do not experience active chemical production and loss of reactive nitrogen species, as well as wet scavenging (important for HNO$_3$, and therefore NO$_y$), an air parcel should in general follow the mixing line (thick gray dashed lines) as it descends into the troposphere. The thin gray dashed lines outline the envelope of mixing lines when allowing one sigma deviation from the mean concentration in both the stratospheric air and tropospheric background.

\[ \sim 400 \text{ pptv} \]. The differences in NO$_y$ partitioning between the fresh and aged stratosphere-troposphere mixed plumes are likely linked to photochemical process which occur when air of stratospheric origin is mixed with tropospheric background: stratospheric NO$_x$ and HNO$_3$ are converted to PAN in the presence of tropospheric VOCs.

We examine the photochemical equilibrium state within three types of air masses using the NASA Langley box model (Olson et al., 2004): the mean tropospheric background between 6–12 km, the lowermost stratosphere, and the stratosphere-troposphere mixed air (Fig. 9). We conduct two base simulations initiated with mean observed chemical composition (Table 3b) in the upper tropospheric background (BCG) (Fig. 9a) and the lower stratosphere (STR) (Fig. 9b), respectively, and ran the box model for 20 days. The BCG case shows that in general the NO$_y$ partitioning in the background air masses sampled during ARCTAS-B is not at equilibrium and PAN is steadily converted to HNO$_3$ at a rate of \( \sim 2 \text{ pptv day}^{-1} \). The NO$_y$ partitioning in the sampled lower stratospheric air is approximately at equilibrium (Fig. 9b).

We now illustrate what happens when lower stratospheric air mixes with tropospheric background air. We initialize the box model with trace gas concentrations assuming a typical BCG point sampled by the 8 July flight mixes with a typical STR point sampled by the same flight (Fig. 9c). A mixing ratio of 70% BCG and 30% STR air sampled near the STR point by the 8 July flight is used which yields CO and CH$_4$ levels approximately close to those sampled in the stratosphere-troposphere mixed subplume A on 8 July (Fig. 8). Within the simulated stratosphere-troposphere mixture, there is active conversion from HNO$_3$ to PAN at \( \sim 3.7 \text{ pptv day}^{-1} \) (net \( \sim 75 \text{ pptv production of PAN in 20 days} \)) and the mixture is not yet reaching equilibrium after the 20 days’ simulation period. A more generalized case initialized with a mixture of mean BCG concentration (70%) and mean STR air (30%) shows similar net conversion
from HNO$_3$ to PAN but at a slower rate, $\sim 2$ pptv day$^{-1}$ (net $\sim 40$ pptv formation of PAN in 20 days) (Fig. 9d).

While the lower stratospheric or upper tropospheric background alone displays zero production or net destruction of PAN, the mixture of the two during STE events provides a unique environment, where tropospheric air which has high VOC and H$_2$O meets stratospheric air which contains high NO$_x$ and O$_3$, resulting in the active photochemical production of PAN. This occurs for two reasons: (i) increased degradation of ethane due to increases in OH (Table 3b) (the increase in OH in stratosphere-troposphere mixed air is due to the mix of O$_3$-rich stratospheric air with moist tropospheric air), (ii) increased PAN formation mainly through the following reactions:

$$
C_2H_6 + OH \rightarrow C_2H_5O_2 + H_2O
$$

$$
C_2H_5O_2 + NO \rightarrow CH_3CHO + NO_2 + HO_2
$$

$$
CH_3CHO + OH \rightarrow CH_3C(O)O_2
$$

$$
CH_3C(O)O_2 + NO_2 \leftrightarrow PAN
$$

A simplified version of the box model, which only includes C$_2$H$_6$, CH$_3$CHO and their intermediate products, CH$_3$C(O)O$_2$, NO$_x$, PAN, HNO$_3$, and constrained with the observed OH, HO$_2$, and photolysis rates, indicates the above chemistry alone is able to explain the $\sim 2$ pptv day$^{-1}$ PAN production calculated by the Langley box model (see Supplement, Fig. S3). Additional sensitivity runs suggest that the PAN production rate is most sensitive to the OH as well as NO$_x$ concentrations. Although we successfully demonstrated that the mixing during STE events leads to the photochemical production of PAN, the calculated box model results were only able to explain approximately 40% of the observed 100 pptv increase in PAN that could not be explained by mixing. It is important to note that our box model calculation is an idealized simplification of the real atmospheric chemical processes. The calculated OH and NO, NO$_2$ levels also differ slightly from the observed values, which can impact the calculated PAN production rate. It is desirable to conduct a comprehensive analysis using 3-dimensional chemical transport model for a better quantification of the impact of STE on tropospheric PAN production.

5 Ozone and ozone production in the Arctic troposphere

The photochemical balance between O$_3$ production and loss is an intricate play between NO$_x$, HO$_x$, and O$_3$ (Jaegle et al., 1998; Wennberg et al., 1998). In this section, we examine O$_3$, the O$_3$ production rate, and ozone’s dependence on NO$_x$ and HO$_x$ to understand the sources of O$_3$ in the Arctic and sub-Arctic troposphere.
Fig. 9. Time-dependent box-model calculation of NO\textsubscript{x}, PAN, and HNO\textsubscript{3} partitioning in (a) mean upper tropospheric background and (b) mean lower stratospheric air. We also show results from a third box-model calculation aiming to simulate the stratosphere-troposphere mixed subplume A sampled by the July 8 flight (c). We assume injection of the lower stratospheric air into the background air mass with a mixture ratio of 30% stratospheric air vs. 70% background air which yields approximately the measured CO and CH\textsubscript{4} in the stratosphere-troposphere mixed plume. (d) shows what happens in a more generalized stratosphere-troposphere mixed case assuming 30% stratospheric air mixes with 70% tropospheric air using the measured mean concentrations in the two air masses during ARCTAS-B.

5.1 Dependence of ozone production on NO\textsubscript{x} and HO\textsubscript{x}

Ozone is produced in the troposphere mainly through the following chemical reactions,

\begin{align*}
\text{OH} + \text{CO} & \rightarrow \text{HO}_2 + \text{CO}_2 \\
\text{HO}_2 + \text{NO} & \rightarrow \text{OH} + \text{NO}_2 \\
\text{NO}_2 + \text{hv} & \rightarrow \text{NO} + \text{O} \\
\text{O} + \text{O}_2 & \rightarrow \text{O}_3
\end{align*}

Net: \( \text{CO} + 2\text{O}_2 \rightarrow \text{CO}_2 + \text{O}_3 \) with Reaction (2) being the rate-limiting step. Ozone can also be produced from reaction of peroxy radicals (RO\textsubscript{2}) with NO:

\begin{align*}
\text{RO}_2 + \text{NO} & \rightarrow \text{CH}_2\text{O} + \text{HO}_2 + \text{NO}_2
\end{align*}

Ozone is destroyed in the troposphere through photolysis and reaction with HO\textsubscript{x} (OH + HO\textsubscript{2}):

\begin{align*}
\text{O}_3 + \text{hv} & \rightarrow \text{O}(^{1}\text{D}) + \text{O}_2 \\
\text{O}(^{1}\text{D}) + \text{H}_2\text{O} & \rightarrow 2\text{OH} \\
\text{HO}_2 + \text{O}_3 & \rightarrow \text{OH} + 2\text{O}_2 \\
\text{OH} + \text{O}_3 & \rightarrow \text{HO}_2 + \text{O}_2
\end{align*}

Figure 10 shows the dependence of the calculated instantaneous NP(O\textsubscript{3}) on levels of NO\textsubscript{x} during ARCTAS for high and low HO\textsubscript{x} conditions and vice versa. While NO\textsubscript{x} remains relatively unchanged from spring to summer, the O\textsubscript{3} production rate during summer is \( \sim 10 \) times higher than that calculated for spring, due to active photochemistry with increasing insolation and humidity. The level of NO\textsubscript{x} plays a determinative role in the photochemical production of O\textsubscript{3}. The NP(O\textsubscript{3}) increases rapidly with increasing levels of NO\textsubscript{x},

We use the O\textsubscript{3} production and loss rates calculated by the NASA Langley box model (Olson et al., 2004) constrained by chemical and physical parameters measured by the DC-8 aircraft. For this study, we use mostly the instantaneous production and loss rates from the box model calculation. Note that the catalytic destruction of O\textsubscript{3} by bromine radicals is also included in the Langley box model to account for the O\textsubscript{3} depletion events sampled during ARCTAS. For simplicity and clarity, we exclude data that contain elevated bromine (BrO > 1.5 pptv). The calculated net O\textsubscript{3} production (formation-destruction) rates, NP(O\textsubscript{3}), can be approximated as:

\[ \text{NP}(\text{O}_3) = \text{P}(\text{O}_3) - \text{L}(\text{O}_3) = K_2[\text{HO}_2][\text{NO}] + K_5[\text{RO}_2][\text{NO}] - K_6[\text{H}_2\text{O}][\text{O}(^{1}\text{D})] - K_7[\text{HO}_2][\text{O}_3] - K_8[\text{OH}][\text{O}_3] \]
suggesting that the Arctic troposphere is in the NO\textsubscript{x}-limited regime. The rate NP(O\textsubscript{3}) increases with increasing NO\textsubscript{x} is dependent on the abundance of HO\textsubscript{x} radicals. When HO\textsubscript{x} is high (>4 pptv in spring and >10 pptv in summer which occur mostly in combustion plumes), the NP(O\textsubscript{3}) increases drastically as NO\textsubscript{x} increases. At low HO\textsubscript{x} concentrations (background and air of stratospheric origin), the NP(O\textsubscript{3}) displays a weak increase with increasing NO\textsubscript{x} as both production (Reaction 2) and loss (Reactions 7 and 8) are slow. The dependence of NP(O\textsubscript{3}) on HO\textsubscript{x} is rather complex, impacted by levels of NO\textsubscript{x}. On the one hand, HO\textsubscript{x} can enhance O\textsubscript{3} production through Reaction (2). On the other hand, it provides a reaction partner for O\textsubscript{3} destruction in Reactions (7) and (8). At high NO\textsubscript{x} levels (e.g., fresh combustion plumes, STE events, and convection), the NP(O\textsubscript{3}) shows a positive dependence on HO\textsubscript{x}. When NO\textsubscript{x} is low (<40 pptv, i.e., background and aged pollution plumes), the NP(O\textsubscript{3}) is either insensitive to HO\textsubscript{x} (spring) or decreases with increasing HO\textsubscript{x} and becomes negative when HO\textsubscript{x} exceeds 20 pptv (summer).

5.2 O\textsubscript{3} and O\textsubscript{3} production in various air masses

The mean background O\textsubscript{3} in the Arctic and sub-Arctic troposphere remains relatively constant from spring to summer, increasing with altitude from 30–40 ppbv at the surface to 60–70 ppbv in the middle and upper troposphere (Fig. 11b and e and Table 3). The middle troposphere (3–8 km) shows net O\textsubscript{3} destruction while the lower (<3 km) and upper troposphere (>8 km) has NP(O\textsubscript{3}) greater than zero (net O\textsubscript{3} formation), regulated by the mean NO\textsubscript{x} concentration. NO\textsubscript{x} ∼ 20 pptv is a critical level (Klonecki and Levy, 1997) in the Arctic troposphere that separates the middle troposphere (NO\textsubscript{x} < 20 pptv, net O\textsubscript{3} destruction regime, Fig. 11c and f) from the lower and upper troposphere where NO\textsubscript{x} exceeds 20 pptv, hence, net O\textsubscript{3} formation (Fig. 11c and f).

Air masses associated with recent STE intrusions have a mean O\textsubscript{3} mixing ratio of ∼150 ppbv (vs. ∼70 ppbv in background air between 6–12 km), implying that the direct transport of high O\textsubscript{3} from the stratosphere to the troposphere during STE events are a significant source of O\textsubscript{3} in the Arctic during spring and summer, particularly in the upper troposphere (Fig. 11). In addition, air masses influenced
by stratospheric air show net O₃ formation with instantaneous NP(O₃) as high as ~2 ppbv day⁻¹ in spring and ~5 ppbv day⁻¹ in summer. The active photochemical production to some extent offsets the dilution by tropospheric background air which has relatively low O₃ and helps in maintaining the high O₃ level during STE transport. We are able to calculate the reaction rates offline for Reactions (2), (6), (7), and (8) using observed O₃, temperature, and the instantaneous OH, HO₂, NO, O(¹D) calculated by the NASA Langley box model. Figure 12 shows the calculated mean instantaneous rates for these reactions in the stratospheric air and stratosphere-troposphere mixed air, in contrast to the background between 6–12 km. As stratospheric air is entrained into the troposphere during STE, it mixes with the surrounding moist background air, therefore increasing HOₓ through increasing K₆[O(¹D)][H₂O] (e.g., Esler et al., 2001). This increases ozone production by increasing K₇[HO₂][O₃] + K₈[OH][O₃]. Note that although ozone loss through Reaction (6) (K₆[O(¹D)][H₂O]) also increases significantly, particularly in summer, its rate is much smaller than those of Reactions (2), (7), and (8). The increase in the O₃ production term outweighs the loss terms and results in an increase in NP(O₃) from near zero to net O₃ formation.

Convection/lightning is an important source of O₃ during ARCTAS-B. The summertime convective/lightning air mass is elevated in O₃ (ΔO₃ ~ 10 ppbv, Table 3b) and displays high net O₃ production. This O₃ enhancement value is similar to that measured in the convective/lightning air masses (ΔO₃ ~ 5 ppbv) during the Intercontinental Chemical Transport Experiment-Phase A (INTEX-A) mission in North America during summer 2004 (Liang et al., 2007). The anthropogenic air mass in general contains relatively low NOₓ (Figs. 4 and 5) and therefore weaker O₃ production (Fig. 11). Despite the positive NP(O₃) rates, the anthropogenic air mass sampled during both ARCTAS-A and ARCTAS-B show no elevated O₃, compared to the background, throughout most of the troposphere. The springtime biomass burning air mass shows active O₃ production (mean instantaneous NP(O₃) ~ 2 ppbv day⁻¹ and ΔO₃/ΔCO = 0.22) and an average ΔO₃ of 16 ppbv. The majority of these plumes are aged agricultural biomass burning plumes from Siberia (Fisher et al., 2010). These values are comparable to those reported in the biomass burning plumes in East Asia during the Transport and Chemical Evolution over the Pacific (TRACE-P)
mission ($\Delta O_3 \sim 2–15$ ppb) (Tang et al., 2003), the contribution of Indonesia biomass burning calculated by Duncan et al. (2003) ($\Delta O_3 \sim 10–20$ ppb) as well as the $\Delta O_3/\Delta CO$ ratio in the North American boreal fire plumes during INTEX-A ($\Delta O_3/\Delta CO = 0.25$) (Pfister et al., 2006). The $\Delta O_3/\Delta CO$ in the aged biomass burning plumes sampled in the equatorial Atlantic during fall 1992 were significantly higher, with mean $\Delta O_3/\Delta CO = 0.7$ (Mauzerall et al., 1998). In contrast, biomass burning air masses sampled in summer, although showing very high positive NP($O_3$) values, display no elevated $O_3$. Alvarado et al. (2010) conducted a detailed analysis of NOx and PAN photochemistry in fresh boreal fire plumes during ARCTAS-B and their impact on $O_3$ and found little evidence of $O_3$ formation in the smoke plumes in the aircraft, satellite, or model results, in accordance with our findings. Paris et al. (2009) also found negative $O_3$/CO ratio ($-0.04$ ppbv ppbv$^{-1}$), indicating photochemical destruction of $O_3$ in the summertime Siberian fire plumes observed during the YAK-AEROSIB/POLARCAT experiment. While the NP($O_3$) in the summer fire plumes are high, this high production is not sustainable as $O_3$ production efficiency is low and NOx are rapidly converted to PAN in $\sim 2$ h (Alvarado et al., 2010). However, a study by Real et al. (2007) demonstrated that thermal decomposition of PAN in a biomass burning plume can lead to a 17 ppbv increase of $O_3$ in 5 days during its long-range transport from Alaska to Europe.

As we have discussed in Sect. 3.1, measurements obtained during ARCTAS-B are concentrated in the sub-Arctic between $50^\circ$N–$70^\circ$N. Our conclusions may not represent the true Arctic troposphere as more convection occurs in the continental sub-Arctic and is closer to pollution sources. We narrow our investigation to focus only on the measurements obtained north of $70^\circ$N during ARCTAS-B (Fig. 13). While there is a significant fraction of convection/lightning air masses identified during ARCTAS-B, none is sampled in the deep Arctic. Air masses associated with STE events are the only notable contributors to NOx and $O_3$ north of $70^\circ$N. In addition, the stratosphere-troposphere mixed air mass is
the only air mass that displays net $O_3$ formation above 2 km. The mean 24-h averaged NP($O_3$) (calculated by the Langley box model) in the fresh stratosphere-troposphere mixed air mass sampled during ARCTAS-B is $\sim 0.9$ ppbv day$^{-1}$. These findings confirm our previous proposition that STE, in addition to its direct contribution to $O_3$, is the driving mechanism of net $O_3$ formation in the Arctic upper troposphere as calculated in the GMI Combo CTM (0–10 ppbv month$^{-1}$ in July) (Liang et al., 2009).

### 6 Conclusions

The NASA ARCTAS mission presents a unique opportunity with its extensive suite of measurements to examine $O_3$ photochemistry in the Arctic troposphere. The PDF distribution of the observed CO suggests that the DC-8 aircraft measurements obtained during spring (ARCTAS-A) are representative of the mean Arctic troposphere. Measurements obtained during the summer deployment (ARCTAS-B) are highly biased towards pollution plumes, a fact to consider in understanding the mean chemical composition of the summertime Arctic troposphere and the impact of pollution plumes.

We conducted a detailed analysis to examine $O_3$ and NO$_x$ in the Arctic and sub-Arctic region and their source attribution. Using a number of marker tracers, we were able to identify various air masses from the background, including anthropogenic pollution, biomass burning emissions, air masses associated with fresh stratosphere-troposphere exchange, and convection and/or lightning influences.

The background Arctic troposphere has mean $O_3$ of $\sim 60$ ppbv and NO$_x$ of $\sim 25$ pptv throughout spring and summer. The mean CO mixing ratio decreases from $\sim 145$ ppbv during spring to $\sim 100$ ppbv in summer. The Arctic troposphere is in the NO$_x$-limited regime with much of the free troposphere showing net $O_3$ destruction due to its low NO$_x$ concentration. These observed CO, NO$_x$, and $O_3$ mixing ratios are not notably different from the values measured during the 1988 ABLE-3A and the 2002 TOPSE field campaigns.

Air masses associated with recent stratosphere-troposphere exchange are present at $>5$ km during spring and summer. These air masses with mean $O_3$ concentrations of 140–160 ppbv are significant direct sources of $O_3$ in the Arctic ($>70^\circ$ N) troposphere. Air of stratospheric origin is also significantly elevated in NO$_x$ (mean $\sim 75$ pptv in spring and 110 pptv in summer) and HNO$_3$ (mean $\sim 290$ pptv in spring and 500 pptv in summer), which will further release NO$_x$ through photochemical destruction. Driven by high NO$_x$, these air masses display active net $O_3$ formation with instantaneous production rates as high as $\sim 2$ ppbv day$^{-1}$ in spring and $\sim 5$ ppbv day$^{-1}$ in summer. During ARCTAS-B, several plumes that were influence by stratospheric air also show conversion of HNO$_3$ to PAN. This active production of PAN is the result of increased degradation of ethane in the stratosphere-troposphere mixed air to form CH$_3$CHO, followed by subsequent formation of PAN under high NO$_x$ conditions. This implies that the impact of NO$_y$-enriched stratospheric air on tropospheric NO$_x$, and therefore $O_3$ production, can be extended much further as the resulting PAN is transported to the lower altitudes and releases NO$_x$ downwind through thermal decomposition (e.g., Moxim et al., 1996; Honrath et al., 1996). Although a quantitative estimate of the impact of the influx of NO$_y$ from the stratosphere on tropospheric NO$_x$, PAN, and, subsequently, $O_3$ production is yet to be determined through more comprehensive 3-dimensional global chemistry modeling studies, the findings from the ARCTAS measurements suggest that an accurate representation of this influx, in addition to stratospheric influx of $O_3$, is essential in tropospheric chemistry transport models of the Arctic to accurately simulate $O_3$, NO$_x$, and PAN in the Arctic troposphere.

Although anthropogenic and biomass burning pollution plumes show highly elevated hydrocarbons and NO$_y$ (mostly in the form of NO$_x$ and PAN). Except some aged Siberia biomass burning plumes during spring, the majority of the pollution plumes has $O_3$ levels comparable to that in the Arctic troposphere, and thus unlikely to further increase the background $O_3$. However, it is important to point out that anthropogenic and biomass burning emissions can still exert an impact on $O_3$ in the Arctic through increasing the background $O_3$ in the mid-latitudes which then enters the polar troposphere through long-range transport, as demonstrated by Shindell et al. (2008). Convection and/or lightning influences are of negligible importance as a source of $O_3$ in the Arctic but can have significant impacts in the upper troposphere in the continental sub-Arctic during summer.

**Supplementary material related to this article is available online at:**
http://www.atmos-chem-phys.net/11/13181/2011/acp-11-13181-2011-supplement.pdf.

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