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1 Direct Measurements of the Convective Recycling of the Upper Troposphere

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We present a statistical representation of the aggregate effects of deep convection on the 1 2 chemistry and dynamics of the Upper Troposphere (UT) based on direct aircraft observations of the chemical composition of the UT over the Eastern United States and 3 4 Canada during summer. These measurements provide new and unique observational 5 constraints on the chemistry occurring downwind of convection and the rate at which air in the UT is recycled, previously only the province of model analyses. These results 6 7 provide quantitative measures that can be used to evaluate global climate and chemistry 8 models.

1 Deep convection is a highly efficient mechanism for the vertical transport of air from 2 near the Earth's surface (0-2 km) to the Upper Troposphere (UT) (6-12 km) (1-5). 3 Typical convective storms have spatial scales of tens of kilometers and vertical velocities as large as 15 m sec⁻¹ (6), making their local influence in the UT extremely strong. The 4 5 rapid upward flow is balanced by much slower descending flow that occurs over a larger 6 spatial scale (7). Convection is also associated with lightning a significant source of NO_x 7 $(NO_x \equiv NO + NO_2)$ in the UT (8, 9). The source strength and spatial distribution of lightning NO_x emissions are not well known, with estimates ranging from 2-20 Tg(N) yr⁻¹ 8 for the global average (10), compared to 25 Tg(N) yr⁻¹ from fossil fuel combustion (11). 9 10 Although there have been a number of case studies of the chemical effects of individual 11 storms (12), studies of the aggregate effects of convection on the chemical composition 12 and radiative forcing of the UT have been largely the province of modeling and theory 13 (13, 14). Here we describe measurements that provide a direct link between an 14 observable property and the ensemble of convective events.

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The chemical and radiative consequences of convection and lightning are known to be large (2, 3, 15). Upper tropospheric O_3 , either transported directly from the boundary layer via convection or formed in situ following detrainment of convectively lofted O_3 precursors (NO_x, odd hydrogen radicals (HO_x) and hydrocarbons) in the outflow region, directly impacts climate through a positive radiative forcing (15). Additionally, deep convection accounts for a significant fraction of the net flux of moisture from near the Earth's surface to the UT (16). Thus, the rate at which the UT is turned over by 1 (

convection has important implications on the hydrological cycle and the magnitude of the water vapor feedback on global temperature (*17*).

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2

4 In this study we describe a method for calculating the time air spends in the upper 5 troposphere following convection from in situ measurements of the chemical composition 6 of the UT and discuss the chemistry occurring in the outflow region as a function of time 7 since convection. We use measurements of NO₂ (18, 19) (NO_x is calculated from NO₂, 8 O₃, HO₂ and photolysis rates), HNO₃ (20, 21), OH and HO₂ (22), O₃ (23), aerosol number 9 density (24), actinic flux (25), CO (26) and CO_2 (27) obtained during the Intercontinental 10 Chemical Transport Experiment – North America (INTEX-NA) aboard the NASA DC-8 11 (28). Measurements were made at altitudes between the surface and 12.5 km, over a wide 12 area of the US and Canada, west of 40° W and between 30 and 50° N. There were a large 13 number of vertical profiles allowing a reasonably unbiased statistical sampling of air over 14 this region during July and August of 2004.

15

16 We use the deviation of the observed NO_x to HNO₃ ratio from steady-state as an indicator 17 of convective influence. The NO_x to HNO_3 ratio is reset to near infinity in moist 18 convection as a result of preferential wet scavenging of HNO_3 relative to NO_x (i.e., the Henry's Law Constant for HNO₃ is ~ 10^8 times larger than for NO_x) (29). Further, 19 lightning initiated NO_x production, often coincident with convection, dramatically 20 21 enhances NO_x in the outflow region. The coupling of these processes makes the NO_x to 22 HNO₃ ratio in the UT an effective indicator of convective influence, where NO_x/HNO₃ 23 >> 1 is indicative of recent cloud outflow (30, 31). In the days following convection, the 1 ratio decays toward steady-state providing a chemical clock that marks the time an air-2 mass has spent in the UT following convection (32). A number of previous studies have 3 used species which have no upper tropospheric source (e.g. CH_3I) (33), or alternative 4 chemical ratios to provide estimates of age of air in the UT (34, 35). Our study is unique 5 because of the availability of high time resolution NO₂, OH and HNO₃ measurements that 6 allow us to build a much more extensive data set than previous studies. After the initial 7 turbulent mixing in the near field of the convection, mixing is slow, thus the time 8 evolution of NO_x/HNO_3 following convection depends largely on the partitioning of NO_x 9 (between NO and NO₂), the concentration of OH and the actinic flux.

10

11 Reactive Nitrogen Partitioning in the UT

12 The only significant chemical sinks of UT NO_x are reaction with OH to produce HNO₃ 13 (Equation 1) and nighttime loss through NO₃ (Equation 2a-b) followed by hydrolysis of 14 N₂O₅ to produce HNO₃ (*36*). NO_x is regenerated by nitric acid photolysis (and 15 subsequent NO₃ photolysis to NO₂) and reaction of OH with HNO₃ (Equations 3 and 4).

$$16 \qquad NO_2 + OH \to HNO_3 \tag{1}$$

$$NO_2 + O_3 \rightarrow NO_3 + O_2 \tag{2a}$$

$$NO_3 + NO_2 \rightarrow N_2O_5 \tag{2b}$$

$$19 \qquad HNO_3 \xrightarrow{h\nu} OH + NO_2 \tag{3}$$

 $20 HNO_3 + OH \to NO_3 + H_2O (4)$

Including the altitude dependent rain-out rate for HNO₃ (k_{rain-out}) as derived by Giorgi and
Chameides (*37*), the expected steady-state NO_x/HNO₃ is:

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$$1 \qquad \left(\frac{\left[NO_{x}\right]}{\left[HNO_{3}\right]}\right)_{Steady-State} = \frac{J_{HNO_{3}} + k_{HNO_{3}+OH}\left[OH\right] + k_{HNO_{3} rainout}}{\left(k_{NO_{2}+OH}\left[OH\right] + 2k_{N_{2}O_{5}+H_{2}O}\left[\frac{\left[N_{2}O_{5}\right]}{\left[NO_{2}\right]}\right]x\left(\frac{\left[NO_{2}\right]}{\left[NO_{x}\right]}\right)}$$
(3)

2

3 Our observations show the NO_x to HNO_3 ratio to be much higher than the ratio described 4 by equation 3 at altitudes greater than 6 km (c.f. Fig. 2). The difference between the 5 observed ratio and that predicted by equation 3 grows with altitude, reaching a maximum Previous observations of NOx and HNO3 (either measured directly or 6 at 10 km. 7 calculated from observations of NO_x, PAN and NO_y) have shown the NO_x/HNO₃ ratio to 8 be significantly larger than the steady-state prediction in the UT (30, 31, 38-42). This has 9 been shown to be primarily a result of convection and lightning reinitializing the system 10 before steady-state is achieved (30, 31). Although a series of other hypotheses have been 11 put forth (40-42), we (like Jaegle et al.) find no evidence for a mechanism other than 12 convection responsible for holding NO_x/HNO₃ out of steady-state in the UT.

13

14 Chemical Signatures of Convection

15 Fig. 3 depicts one of many convectively influenced air-masses sampled in the UT during 16 Three distinct convective events (40-80 km wide) are identified by INTEX-NA. 17 enhancements in NO_x/HNO₃ in Fig. 3a. Coincident enhancements are present in SO₂, an indicator of a recent boundary layer source for this air, and Ultra-fine Cloud 18 19 Condensation Nuclei (UCN) ($3 \le D_p \le 10$ nm), an indicator of cloud detrainment (Fig. 20 3b) (43, 44). Sharp decreases in CO_2 also indicate the convective lofting of boundary layer air depleted in CO_2 , a result of photosynthetic activity (Fig. 3c) (44). 21 22 Enhancements in CO, CH₂O and various hydrocarbons, relative to the surrounding UT

air, were also observed in these plumes, further indicating that these parcels originate
from the Planetary Boundary Layer (PBL). Backward air trajectories, initialized along
the flight track, coupled to the spatial and temporal distribution of cloud-to-ground
lightning strikes, indicate that the sampled air-mass was recently influenced by lightning
approximately one day prior to DC-8 sampling (c.f. Fig. 3 bottom panel) (45). Such
features with high NO_x/HNO₃ were observed throughout the UT during INTEX-NA.

7

8 To assess the extent to which the UT over the Eastern U.S. and Canada during the 9 summer of 2004 was influenced by convection and describe the chemical evolution of 10 convective outflow, we use a constrained time-dependent photo-chemical box model to 11 map the observed NO_x/HNO_3 to the time since the ratio was last reinitialized. The model 12 is described in the supplemental information included with this article. It is initialized 13 with observations at 1km vertical intervals from 6 to 12 km. The derived timing indicator 14 for the convectively influenced air sampled on 11 August 2006 is shown in Fig. 3d. The 15 properties of the ensemble of our measurements are shown in Figs. 4-6.

16

The aerosol size distribution provides an independent indicator of air recently detrained from clouds. Cloud-processed air is depleted of aerosol surface area permitting new particle formation in the outflow region (*43*, *44*). Fig. 4a depicts the fraction of condensation nuclei found in the 3-10 nm bin as a function of time since convective influence. The fraction of particles in this ultra-fine mode is largest during the first few days confirming that the NO_x to HNO₃ ratio, and the timing indicator derived from it, is reinitialized in the UT by cloud processing. Strong enhancements in CH₃OOH/H₂O₂ (not shown), also an indicator of recent cloud processing (*33, 46*), were observed during the
 first two days after cloud processing.

3

As expected, both elevated NO_x and suppressed HNO₃ are observed at short times (c.f. Figs. 4b-c). Enhancements in NO_x during the first few days is indicative of convection of boundary layer and/or lightning NO_x (47, 48). The suppression of HNO₃ at short times is clear indication of HNO₃ scavenging during convection. Fig. 4d confirms that reactive nitrogen (NO_y \equiv NO_x + total peroxy nitrates + HNO₃) is conserved during the chemical processing following convection, a fact which provides further support for the use of NO_x/HNO₃ as a marker representing time since convection.

11

12 Chemical Processing in Convective Outflow

Mapping the ensemble of observations made throughout the UT onto the coordinate of time since convection allows us to assess the chemical and dynamical processes occurring following convection, without attempting a Lagrangian convection study. In this analysis we concentrate on the time evolution of CO and O₃.

17

The time evolution of CO, following detrainment into the UT, is set by the abundance of OH and the rate at which the convective plume entrains air from the background UT (c.f. Fig. 5a). Due to the direct dependence of the chemical clock on HO_x , we constrained both OH and HO_2 to the observations as a function of NO_x and pressure in the timedependent model used to generate time. As a result, we can iterate the model to determine the proper mixing rate of the convective plume by matching the modeled and

1 observed time evolution of CO following convection. Using this approach for a series of 2 long lived species (e.g. CO, CH₄, CH₃OH and others), we calculate an average mixing rate of 0.05 ± 0.02 day⁻¹ following detrainment into the UT. This is in good agreement 3 with the upper limit of 0.06 - 0.1 day⁻¹ determined by Ray et al. from observations of 4 5 convective plumes observed in the stratosphere during the CRYSTAL-FACE mission 6 (49), however it is slower than the 2 day dilution time-scale determined by Wang et al. 7 from observations during the SONEX Experiment (32). Since the DC-8 did not routinely 8 sample in the turbulent environment directly surrounding convective outflow, this mixing 9 rate likely reflects diffusive and shear induced mixing subsequent to the initial turbulent 10 mixing occurring during detrainment from the convective system. Entrainment of UT air 11 either during convective lofting or cloud detrainment is discussed in the next section.

12

13 The O₃ mixing ratio as a function of time since convection is shown in Fig. 5b. We find 14 that on average, convectively lofted air-masses contain less O₃ than the background UT. 15 This result is consistent with the observed vertical gradient in O₃ observed over the 16 continental US during INTEX-NA, with lower O_3 in the PBL than above (50). Rapid 17 changes in the O₃ mixing ratio are observed during the first two days following detrainment, with the observed O_3 15 nmol mol⁻¹ above the initial value by the end of day 18 19 two. The observed rate of increase slows exponentially with an asymptote at long time approaching zero and the O_3 mixing ratio approaching a constant value of 85 nmol mol⁻¹. 20 21 This is a surprising result, as our model of the O_3 rate of change never approaches zero, but continues to predict a net increase of 3 nmol mol⁻¹ O₃ day⁻¹ at the end of day five 22 23 (51).

2 Constraints on the Convective Turnover Rate of the UT

3 The convective turnover rate of the upper troposphere is critical for accurately describing 4 NO_x , HO_x and O_3 chemistry in the UT (52). However, at present there is a paucity of 5 observation based constraints available (either meteorological or chemical) to test the 6 aggregate effects of convection in the current generation of global chemistry and climate 7 models. To determine the convective turnover rate of the UT from the observations 8 presented here, both the extent to which the UT is influenced by convection and the 9 fraction of PBL air in the convectively influenced air-masses must be known with high 10 confidence.

11

To determine the fraction of PBL air contained in fresh convective outflow, we use observations of insoluble long-lived species made throughout the INTEX-NA campaign over the continent. Assuming that we conducted a statistically unbiased sampling of both the boundary layer and free troposphere during INTEX-NA, we can calculate the fraction of PBL air present in fresh convection (f) through the following equation:

17

18
$$[X]_{UT(t=0)} = f [X]_{surface} + (1-f) [X]_{UT}$$
 (4)

19

where [X]_{UT (t=0)} is the mean mixing ratio of species X in fresh convective outflow (as
identified using our timing indicator), [X]_{UT} is the mean mixing ratio of species X in the
UT (7.5-11.5 km) and [X]_{surface} is the mean mixing ratio of species X between 0-1.5 km.
Using observations of CO, CO₂, CH₃OH, CH₄ and C₂H₆ we calculate the fraction of PBL

1 air in fresh convection to be 0.19 ± 0.05 , 0.11 ± 0.03 , 0.26 ± 0.05 , 0.15 ± 0.05 , and 0.34 ± 0.05 2 0.09, respectively. We calculate a weighted average for the fraction of PBL air in convective outflow of 0.17 ± 0.02 by weighting each value by the inverse square of its 3 4 uncertainty. This implies that convectively lofted PBL air rapidly entrains the 5 surrounding air either during ascent or in the turbulent environment of the detraining 6 flow. These results are consistent with: i.) the observations of Ray et al., who determined 7 the fraction of tropospheric air in convective plumes sampled in the stratosphere to be 8 between 0.1 and 0.4 (49), ii.) the observations of Cohan et al., who calculate the fraction 9 of BL air in fresh convection outflow to be between 0.32 and 0.64 from observations of 10 CHBr₃, CH₃OOH and CH₃I in fresh convection (33) and iii.) the modeling studies of 11 Mullendore et al., who calculate the fraction of PBL air present in the convective outflow 12 region of a supercell storm to be 0.26, 10 hours after storm initialization (53).

13

14 Fig. 6a shows the normalized frequency distribution of the observed time since 15 convection based on the ratio of NO_x to HNO₃. We find that 54% of the air sampled 16 between 7.5 and 11.5km had been influenced by convection during the past two days. 17 The convective outflow was strongest between 9.5 and 10.5 km, where the fraction of 18 sampled air that is less than two days old exceeds 69%. The vertical distribution 19 presented here is consistent with previous observations and model analyses of convective 20 outflow to the UT from individual storms (4, 54) and the vertical distribution of 21 convectively influenced laminae observed in O_3 sonde data from the summer of 2004 22 over the northeastern U.S. (55). The shift toward longer times between 10.5 and 11.5 km 23 suggests that either convective cloud tops on average do not extend higher than 10.5 km over the mid-latitude during the summer (56) or that transport of stratospheric air, rich in
 HNO₃, contributes to keeping the NO_x to HNO₃ ratio low at altitudes greater than 10.5
 km (57).

4

5 To constrain the turnover rate of the UT from the ensemble statistics generated from our 6 calculated time since convective influence (Fig. 6a), we constructed a two dimensional 7 model of the UT. We assume that it takes 4 days for any individual box to pass through 8 the sampling region and that each box has not been influenced by convection upon 9 entering the sampling window. Every six hours we: i.) represent convection by 10 randomly reinitializing the age of x% of the boxes in the sampling domain to 0 (the value of x is determined by the turnover rate (varied between 0.05 and 0.2 day⁻¹) and the 11 12 fraction of PBL air contained in fresh convection (assumed to be a constant at 0.17)) and ii.) dilute each box with the mean value of the adjacent 8 boxes at the rate of 0.05 day⁻¹. 13

14

15 Fig. 6b depicts the observed and three calculated normalized frequency distribution of 16 time since convective influence between 7.5 and 11.5 km. The shape of the distribution 17 suggests that UT air sampled during INTEX-NA was largely influenced by convection, 18 and that convectively lofted plumes did not have sufficient time to either mix or age prior 19 to sampling by the DC-8, but instead were transported to the East out of the domain. 20 Frequency distributions of time since convection, using three different convective 21 turnover rates, in the Eastern half of the 2-D UT model analysis (where we sampled most 22 frequently) are also shown in Fig. 6b. Assuming the DC-8 made a statistically unbiased 23 sampling of the continental UT during summer, the best match between the model and observations would imply a convective turnover rate between 0.1 - 0.2 day⁻¹. However, if
we assume the DC-8 had a positive bias toward sampling fresh convection in accordance
with Fuelberg et al., our observed frequency distributions are consistent with a convective
turnover rate closer to 0.1 day⁻¹ (45, 58).

5

6 For comparison, the model detrainment cloud mass flux between 400 and 200 hPa 7 (approximately 7.2 - 11.8 km based on standard atmosphere) for the INTEX-NA sampling domain (80° - 100°W for 30° - 35°N and 70° - 100° W for 35° - 50° N) between 8 July 1st and August 15th 2004 was 0.0085 kg m⁻² sec⁻¹. This corresponds to a turnover 9 rate of 0.37 day⁻¹ (using a column mass of 1.9×10^3 kg m⁻² between 7.2 - 11.8 km). As 10 11 expected, the observed and modeled detrainment rates for the extratropics are slower than 12 in the tropics. For example, Folkins and Martin determine a maximum convective detrainment rate of 0.4 day⁻¹ at 12.5 km from calculation of the clear-sky radiative and 13 14 precipitation induced evaporative cooling rates using a one-dimensional model 15 constrained by observations of temperature and water vapor (59).

16

17 Conclusions

We present a statistical representation of the aggregate effects of convection on the chemistry and dynamics of the upper troposphere using in situ measurements taken aboard the NASA DC-8 during the summer of 2004 over the Eastern U.S. and Canada. These observations provide a new and unique constraint on: i.) the extent to which convection perturbs the continental UT during summer, ii.) the fraction of boundary layer air present in convective outflow, and iii.) the convective overturn rate of the upper

troposphere. In addition, the chemical clock described here defines a coordinate that can be used to assess the chemistry occurring down-wind of convective injection. These direct measures of atmospheric rates present a new opportunity for quantitative tests of model representations of processes governing UT ozone, convection, lightning and their impact on climate.



1

4 Figure 1: In moist convection, air from near the Earth's Surface is rapidly transported 5 upwards and detrained into the Upper Troposphere. In this process, Nitric Acid (highly soluble) is efficiently scavenged, while NO_x (insoluble) remains. NO_x is dramatically 6 elevated by concurrent lightning NO production, resulting in high NO_x to HNO₃ ratios in 7 the convective outflow region. Following detrainment into the UT, NO_x is converted to 8 9 HNO₃ by OH during the day and via NO_3/N_2O_5 at night. The chemical evolution of the 10 NO_x/HNO₃ ratio provides a unique indicator of the time a sampled air-mass has been in the UT following convection. 11



Figure 2: The observed deviation of the NO_x to HNO_3 ratio from steady-state as a

5 function of altitude in the UT. The mean values within 500 m vertical bins are shown 6 with circles (\circ). The steady-state NO_x to HNO₃ ratio was calculated from measured NO_x, 7 OH and J_{HNO3} and includes the rain out parameterization of Giorgi and Chameides 8 (1985).



2 Figure 3: top panel Time series of measurements taken in the vicinity of recent 3 convective activity on 11 August 2004 between 5 and 9 km. Panel A suggests the 4 sampling of a series of fresh convective plumes, indicated by a sharp increase in the 5 NO_x/HNO₃ ratio. Panels B & C depict coincident enhancements in SO₂ and UCN (3nm > 6 $D_p > 10$ nm) and coincident sharp drops in CO₂, indicative of the convective lofting of 7 boundary layer air depleted in CO₂. The derived time since the sampled air-mass had been influenced by convection is shown in Panel D. bottom panel NLDN lightning hits 8 (small dots) on the 10th and 11th of August. The color-code represents the time of the hit 9 10 (hours) prior to aircraft sampling. The DC-8 sampling location corresponding to measurements shown in Figure 1 is located on the Maine – New Brunswick border [46°N 11 12 67°W]. The two day back trajectory [•] (initialized at the point of the second convective 13 plume shown in Panel A) is also color-coded by time prior to DC-8 sampling (dots with 14 black edges).





0.6

3 Figure 4: Observations of the fraction of ultra-fine condensation nuclei [number density 4 of aerosol (3-10 nm) / total aerosol number density] (Panel A), NO_x (Panel B), HNO₃ 5 (Panel C) and NO_v (Panel D) as a function of modeled time since convective influence. 6 The mean $(-\Box)$ and median $(-\odot)$ of the observations, within 8 hour bins, is shown along 7 with interquartile range (shaded region). Results from the time-dependent box model, 8 initialized at 10 km and 12PM, are shown with dashed lines for the gas phase species (B-9 D).



Figure 5: Observations of CO (Panel A) and Ozone (Panel B) as a function of modeled time since cloud processing in the UT. The mean $(-\Box)$ and median $(-\circ)$ of the observations, within 8 hour bins, is shown along with the interquartile range (shaded region). Results from the time-dependent box model, initialized at 10 km and 12PM, are shown with dashed lines.





Figure 6: *left panel* Normalized frequency distribution in the time since convective influence, as calculated from observations of the NO_x to HNO_3 ratio made during the summer of 2004. Calculations are separated into 1 km altitude bins (ranging from 7.5-11.5 km). The fraction of air that had been influenced by convection within the past two days (f < 2 days) is included in the figure legend. *right panel* Comparison of observed frequency distribution (7.5-11.5 km) with various modeled representations of the convective turnover rate.

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past 2 days. In Fuelberg et al., the authors determine that when considering the entire INTEX-NA sampling domain (both in space and time), convection was present in 12.5% of the grid points. This is substantially smaller than the percent of observations within 6 hours of convection (21.4%), suggesting that the DC-8 had a positive bias toward sampling fresh convection. This bias is reflected in the sharp drop in population between day 1 and 2 as shown in Fig. 6a. Correcting for this bias has little effect on our assessment of the fraction of air less than 2 days old, lowering our results from 0.43, 0.56, 0.69, and 0.43 to 0.38, 0.50, 0.62, and 0.39 at 8, 9, 10, and 11 km respectively.

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1 Supplemental Online Information

2 **1. INTEX-NA Experiment Description and Instrument Descriptions**

The Intercontinental Chemical Transport Experiment – North America (INTEX-NA) 3 4 took place between 1 July and 14 August 2004. Research flights were conducted out of 5 Dryden Flight Research Center (Edwards AFB, CA), Mid-America Airfield (Mascoutah, 6 IL); and PEASE International Trade-Port (Portsmouth, NH). Figure S1 depicts the 7 vertical and horizontal extent of research flights conducted aboard the NASA DC-8 8 during INTEX-NA (1). DC-8 flight tracks are shown in the left panel of Figure 1 and the 9 number of samples (10 second averaging time) in 1km vertical bins are shown in the right 10 panel. In situ observations relevant to this study include; NO₂, HNO₃, OH, O₃, CO, CO₂, 11 SO₂ and Ultra-fine Condensation Nuclei (UCN). Table S1 describes the detection 12 threshold, uncertainty and time response for each measurement used in this analysis.

13

14 NO₂ LIF Instrument The NO₂ instrument flown aboard the DC-8 was described in 15 detail by Thornton et al. (2), with specifics of the jet-expansion described by Cleary et al. 16 (3). Briefly, NO₂ fluorescence is detected at 1Hz following excitation of a specific jet-17 cooled rovibronic transition in NO₂ at 585 nm. Red-shifted fluorescence is imaged at 90° 18 onto an air cooled photomultiplier tube that is both optically and temporally filtered to 19 remove laser scatter. NO₂ mixing ratios are calculated directly from fluorescence counts 20 following calibration to NO₂ gas standards and measurements of the instrument zero from 21 compressed air mixtures containing zero NO₂. Selectivity to NO₂ is demonstrated by 22 tuning on and off of a specific NO₂ resonance, where the difference in observed 23 fluorescence at the two different frequencies is attributed solely to NO₂. We calculate a NO₂ detection threshold of 8 pptv in 10 seconds at the surface and 25 pptv in 10 seconds
at the aircraft ceiling (12.5 km).

3

4 **2. 0-D Time Dependent Model**

5 The chemical evolution of convective outflow was modeled using a 0-D time dependent 6 model. The model was initialized with chemical conditions, altitudes and detrainment 7 times consistent with observations of fresh convection made during INTEX-NA. As time propagates in the model, we calculate the production and loss of O₃, CO, NO, NO₂, NO₃, 8 9 N₂O₅, PAN, HO₂NO₂, HNO₃, OH, HO₂, RO₂, H₂O₂, CH₃OOH, H₂CO and C1-C6 10 Hydrocarbons for 20 days following cloud detrainment. The conversion of NO_x to HNO₃ 11 in the outflow region is used as an indicator of time since convection. Figure S2 depicts 12 the results of a single run initialized at 10km with a noon detrainment time. Initial 13 conditions correspond to $[NO_x]_i = 800$ pptv, $[O_3]_i = 65$ ppbv and $[CO]_i = 105$ ppbv. Rapid 14 conversion of NO_x to HNO_3 is observed during the first few days as the system 15 approaches steady-state. In this analysis we assume: i.) HNO₃ is scavenged with unit efficiency in deep convection, ii.) $\gamma_{N205} = 0.01$ and iii.) HNO₃ is not scavenged by 16 17 aerosols (or rain) following injection into the UT. All kinetic rates used in this analysis 18 were taken from the NASA JPL Chemical Kinetics and Photochemical Data for Use in 19 Atmospheric Studies, Evaluation Number 14 (4).

20

21 2.1 Treatment of OH and HO₂

The calculated time since convective detrainment is directly coupled to the HO_x budget through the daytime NO_x sink to HNO_3 via reaction with OH. As in other model

1 descriptions of the UT during INTEX-NA (5, 6), our unconstrained model over-estimates 2 OH by nearly a factor of two in the UT and under-estimates HO_2 by a similar amount. Due to the direct dependence of our timing indicator on HO_x , we constrain the mixing 3 4 ratios of OH and HO_2 to the observed values as a function of NO_x and altitude. Figure S3 depicts the modeled mixing ratios of OH and HO₂ (lines), constrained to the 5 6 observations (dots), as a function of NO_x and SZA at 10km. The observed OH is a strong 7 function of NO_x, while observations of HO₂ remain insensitive to NO_x. Constraints for 8 OH and HO₂ were derived independently for each 1km altitude bin. Constraining OH 9 and HO₂ to the observations increases the time required for the NO_x-HNO₃ system to 10 reach steady-state (by slowing the rate of $OH + NO_2$) and enhances the modeled O_3 11 production in the outflow region (by speeding up the rate of HO_2+NO).

12

13 **2.2 Calculation of Time since Convection**

The time since a sampled air-mass had been cloud processed is calculated by applying the mapping of time to NO_x/HNO_3 derived in the box model to the observed NO_x to HNO_3 ratio. Figure S4 depicts the best-fit relation between the modeled NO_x to HNO_3 ratio and time since cloud processing at 10km. This function is calculated at 1km increments from 6-12km and applied to the measured NO_x to HNO_3 ratio.

19

20 **2.3 Model Assumptions and Uncertainty**

In order to access the uncertainty in the calculated time, we ran the time-dependent model under various different conditions encountered during INTEX-NA (e.g. $[NO_x]_i$ (0.2-3.0 ppbv), $[O_3]_i$ (40-100 ppbv), $[CO]_i$ (80-150 ppbv), detrainment time (noon, 4PM,

midnight), altitude (6-12km) and time of year (June-September). As illustrated in figure 1 2 S4, the NO_x to HNO₃ ratio has good resolution (i.e. large rate of change per unit time) during the first five days following convection. Beyond five days small changes in 3 4 NO_x/HNO_3 correspond to larger changes in the derived time. From the variance in the 5 calculated time of individual model runs, we estimate the uncertainty in our modeled time 6 to be ± 6 hours at 1 day, ± 12 hours at 2 days and ± 1 day at 4 days. In addition, the 7 INTEX-NA sampling domain did not permit frequent measurement of aged convection 8 (>5 days). For these reasons we limit our analysis to the first five days following 9 convection.

10

11 **2.4 Treatment of Mixing**

The mixing rate was determined by iterating the model until we had closure between the observed and modeled time evolution of a suite of long-lived species (e.g. CO, CH₄, CH₃OH and others). The modeled time rate of change of species X is determined as:

15
$$\frac{d[X]}{dt} = P(X) - L(X) - k_{dilution} ([X] - [X]_{Background})$$

where P(X) represents the chemical production of species X, L(X) represents the chemical loss of species X and $k_{dilution}$ is the mixing rate of the convective plume with background UT. We find this mixing term to be on average 0.05 ± 0.02 day⁻¹. That is, after 5 days, the plume still has 75% of its original contents. While individual convective plumes may mix faster (or slower) that this, the aggregate mixing rate of all the sampled plumes can be described by this rate. This rate supports the conclusion that over the course of 5 days, the convective plume remains relatively isolated from the background

- UT. Background mixing ratios used in the dilution calculation were taken as the mean
 observed UT mixing ratios outside of fresh convective plumes.
- 3

4 Due to subsidence of convectively lofted air parcels following injection, our calculated 5 time represents a lower bound for age as the chemical clock speeds up (due to NO_x 6 repartitioning) as the parcel descends in altitude. However, this is a relatively small 7 effect as calculated subsidence rates are approximately 35 hPa day⁻¹ (7).

8

9 **3.0** Comparison of Chemical and Meteorological Convective Influence Calculations

10 The results presented here provide a chemical constraint on the rate at which the UT over 11 the continental US is influenced by convection during summer. In addition to the 12 meteorological analysis of Fuelberg et al., presented in this manuscript, Thompson et al. 13 assessed the effects of convection on the O₃ budget in the UT, over eastern North 14 America, during the summer of 2004. Using results from the INTEX Ozone Sonde 15 Network Study (IONS), the authors conclude that 10-15% (lower-limit) of the below 16 tropopause O_3 can be attributed to the interaction of regional pollution with convection 17 and lightning (8).

18

The INTEX-NA sampling period (June-August) and region (Eastern North America) is characterized by intense lightning activity. Cooper et al. calculated that 13% of the global annual lightning NO_x emissions occurred between 108° W – 18 ° E and 18 ° -72° N the between June 21 and August 15, 2004 (9). In terms of lightning flash counts, Hudman et al., conclude that 2004 was typical (within 20% of the mean) of the past 5

years (6). Persistent frontal passages prevented the formation of stagnant high pressure
 systems, typically observed during the summer over the northeastern United States (10).
 These frequent passages led to both record low temperatures and number of O₃
 exceedances in the northeast (11). In contrast UT/LS O₃, as observed from the IONS
 network, was comparable to the climatology (11).

7 **4.0 Measurement Uncertainties**

8 In this analysis we calculate NO_x from observations of NO₂, O₃, HO₂ and photolysis rates 9 measured directly on the DC-8. NO was measured directly on the DC-8 via a 10 chemiluminescence commercial grade detector. The sensitivity of the 11 chemiluminescence instrument (detection threshold > 50 pptv) and long integration time 12 (1 minute) prohibited its use in these calculations. NO was calculated from steady-state 13 using the following equation:

14
$$[NO]_{Steady-State} = \frac{J_{NO_2} [NO_2]}{k_{NO+O_3} [O_3] + k_{NO+HO_2} [HO_2]}$$

We calculate the accuracy of the derived NO to be better than \pm 30% based on the propagation of the individual errors used on the calculation. Observations of Nitrogen Dioxide, Ozone and JNO₂ made during the INTEX-NA campaign were compared directly with measurements made aboard the NOAA WP-3D during a series of in-flight comparisons. During these experiments the principal individual components (NO₂, O₃ and JNO₂) showed agreement to within their stated instrumental uncertainty.

1 In Figure S6 we compare the measured NO_x (using the chemiluminescence and the Laser 2 Induced Fluorescence measurements) and NO_x calculated from steady-state for the entire 3 INTEX-NA campaign. The steady-state NO_x agrees with the measured NO_x to within the 4 calculated uncertainty when averaged to 1-minute and divided into 1km vertical bins. 5 The observed upper tropospheric NO_x concentrations during INTEX-NA are on average 6 much higher than previous reported on intensive aircraft field campaigns over North 7 America. During the SUCCESS campaign, Jaegle et al. report mean NO concentrations 8 of 0.030 ± 0.022 ppbv and 0.061 ± 0.045 ppbv, for the altitude ranges of 8-10 km and 10-9 12 km, respectively (12). However, it must be noted that the scientific objective of 10 SUCCESS was the sampling of aircraft exhaust and contrails, thus the values reported in 11 Jaegle et al., were filtered to exclude fresh aircraft exhaust ($CO_2 > 368$ ppmv and NO >12 300 pptv). Crawford et al., report mean NO concentrations of 0.1 ppbv for observations 13 made between 6-12 km during the SONEX campaign during the fall of 1997 over the 14 North Eastern United States and the North Atlantic (13). Neither of these studies 15 provides a direct comparison to the INTEX-NA data-set as SUCCESS was conducted in 16 the spring and SONEX in the fall, while peak lightning and convection occurs over the 17 continental North America during summer. The best comparison comes from NO_x 18 measured aboard a commercial passenger aircraft during the NOXAR program between 1995 and 1997. Brunner et al. report a mean UT NO_x concentration of 0.4 ppbv during 19 20 June, July and August over North America (14). As seen in Figure S6, our observations 21 during the summer of 2004 are consistent with this.

We use the Caltech CIMS HNO₃ due to its fast time response (5 seconds as compared to 105 seconds for the UNH Mist Chamber Technique) and the UNH MC results when the fast HNO₃ was unavailable. To account for the systematic bias between the two observations in the UT ([HNO₃]_{UNH} = $0.6 \times [HNO_3]_{Caltech}$), we scale both the CIMS and MC observations to split the difference between the two measurements (i.e. we increase [HNO₃]_{UNH} by 20% and decrease [HNO₃]_{CIT} by 20%).

8 Due to the observed systematic bias, the choice of which HNO_3 measurement to use in 9 the analysis has the potential to complicate our conclusions. To address these effects we 10 have conducted the analysis using Nitric Acid as measured from: i.) the Mist Chamber -11 Ion Chromatography Instrument, ii.) the Chemical Ionization Mass Spectrometer and iii.) 12 the scaled difference between the two techniques. The results are compared in Figures 13 S7 and S8. Figure S7 shows the normalized frequency distribution in the time since 14 convective influence, as calculated from observations of the NO_x to HNO₃ ratio made 15 during the summer of 2004. Figure S8 depicts the fraction of air that had been influenced by convection within the past two days ($f_{< 2 \text{ days}}$) as a function of altitude. Calculations 16 17 derived from the Mist Chamber - Ion Chromatography Instrument result in a higher 18 fraction of convectively influenced air when compared with calculations made using 19 measurements from the CIMS instrument. When compared with the results shown in 20 Figure 6B of the manuscript, on which our conclusions regarding the convective overturn 21 rate are drawn from, calculations using either the CIMS, MC-IC or the scaled difference result in a convective overturn rate between 0.1 and 0.2 day⁻¹. 22



1 **2. Supplemental Figures**

- 3 Figure S1: *left panel* INTEX-NA flight tracks made between 1 July 2004 and 14 August
- 4 2004 aboard the NASA DC-8. right panel Number of samples (using 10-sec averaged
- 5 data) within 1km altitude bins between 0-12 km during the entire campaign.





was initialized at 12PM local time at 30°N in August using $[NO_x]_i = 800$ pptv, $[CO]_i =$

^{6 105} ppbv at $[O_3]_i = 65$ ppbv.



Figure S3: Model representation of OH (left panel) and HO₂ (right panel) as a function of SZA and $[NO_x]$. Model results (solid lines) are shown on top of the *in situ* observations (dots). The model was initialized at noon at 10km with $[NO_x]_i = 800$ pptv, $[CO]_i = 105$ ppbv at $[O_3]_i = 65$ ppbv. Observations shown were taken aboard the DC-8 between 9 and 11 km.



1

Figure S4: Observed NO_x to HNO₃ ratios are converted to a time since last convective influence using the best fit equation relating the NO_x/HNO₃ ratio to time as calculated 7 using the time-dependent model in 1km altitude bins from 7.5-11.5 km. The above

8 equation is valid for pressure altitudes between 9.5 and 10.5 km.



Figure S5: Comparison of chemical (grey bars) and meteorological constraints (-o-, -□-)
on convective influence during INTEX-NA. Convective influence on air *sampled* by the
DC-8 is shown with blue circles (-o-), while convective influence on the entire INTEXNA domain is shown with red squares (-□-).



3 Figure S6: Comparison of observed (grey lines) and steady-state (black lines) NO_x for

- 4 the entire INTEX-NA field campaign. The shaded region represents the interquartile 5 range of the calculated NO_x .
- 6



- 4 **Figure S7:** Normalized frequency distribution in the time since convective influence, as
- 5 calculated from observations of the NO_x to HNO_3 ratio made during the summer of 2004.
- 6 Calculations were conducted using Nitric Acid as measured from: i.) the Mist Chamber –
- 7 Ion Chromatography Instrument (-□-), the Chemical Ionization Mass Spectrometer (-○-)
- 8 and the scaled difference (--).
- 9





1 2 3 4 5 6 Figure S8: Fraction of air that had been influenced by convection within the past two days ($f_{< 2 \text{ days}}$) as a function of altitude. Calculations were conducted using Nitric Acid as measured from: i.) the Mist Chamber - Ion Chromatography Instrument (-u-), the Chemical Ionization Mass Spectrometer (-0-) and the scaled difference (--).

Species	Measurement Technique	Detection Threshold	Accuracy	Time Response	Reference
NO ₂	LIF^{1}	8 pptv / 10 sec	± 10% 1σ	1 Hz	(2, 3)
HNO ₃	CIMS ³	10 pptv / 0.5s	± 30%	0.5s	(15)
				sample	
				every 5s	
	Mist Chamber – IC ⁴	5 pptv / 105 sec		105 sec	(16)
OH	LIF^1	0.01 pptv	$\pm 32\%$	20 sec	(17)
			2σ - 1 min		
O ₃	Chemiluminescence	Precision =	± 2 ppbv,	1 Hz	(18)
		± 0.8 pptv, $\pm 1\%$	±3%		
CO	IR-Absorption	Precision =	± 1.4 ppby	1 Hz	(19)
	in noonpuon	± 1 ppbv, $\pm 1.5\%$	$\pm 2.6\% 2\sigma$		()
		of reading	,		
CO ₂	IR-Absorption	Precision <	± 0.25 ppmv	1 Hz	(20)
		0.07 ppmv			
UCN	TSI CN counter ⁵	± 10%	± 10%	1 Hz	(21)
J _{NO2}	Actnic Flux	4.1 x 10 ⁻⁷	$\pm 8\%$	1 Hz	(22)
	Spectroradiometer	Precision =			
		±4.5%			

¹LIF – Laser Induced Fluorescence

NO₂ detection threshold is 8 pptv / 10 sec at 760 Torr (ground) and 20 pptv / 10 sec at 200 Torr (10 km)

³CIMS – Chemical Ionization Mass Spectrometry

⁴IC – Ion Chromatography

234 567 ⁵UCN (Ultra-fine Condensation Nuclei) was obtained by the difference of the UCN (D_p >3nm, TSI 3025) and CN (D_P>10nm, TSI 3010) Condensation Nuclei (CN) instruments.

8

9 Table S1: Detection thresholds, measurement uncertainty and time response of the in

situ measurements used in this study. (Note: $ppmv = \mu mol mol^{-1}$, $ppbv = nmol mol^{-1}$ and 10

 $pptv = pmol mol^{-1}$) 11

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