Evaluation of simulated photochemical partitioning of oxidized nitrogen in the upper troposphere

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Abstract. Regional and global chemical transport models underpredict NOₓ (NO + NO₂) in the upper troposphere where it is a precursor to the greenhouse gas ozone. The NOₓ bias has been shown in model evaluations using aircraft data (Singh et al., 2007) and total column NO₂ (molecules cm⁻²) from satellite observations (Napelenok et al., 2008). The causes of NOₓ underpredictions have yet to be fully understood due to the interconnected nature of simulated emission, transport, and chemistry processes. Recent observation-based studies, in the upper troposphere, identify chemical rate coefficients as a potential source of error (Olson et al., 2006; Ren et al., 2008). Since typical chemistry evaluation techniques are not available for upper tropospheric conditions, this study develops an evaluation platform from in situ observations, stochastic convection, and deterministic chemistry. We derive a stochastic convection model and optimize it using two simulated datasets of time since convection, one based on meteorology, and the other on chemistry. The chemistry surrogate for time since convection is calculated using seven different chemical mechanisms, all of which predict shorter time since convection than our meteorological analysis. We evaluate chemical simulations by inter-comparison and by pairing results with observations based on NOₓ:HNO₃, a photochemical aging indicator. Inter-comparison reveals individual chemical mechanism biases and recommended updates. Evaluation against observations shows that all chemical mechanisms overpredict NOₓ removal relative to long-lived methanol and carbon monoxide. All chemical mechanisms underpredict observed NOₓ by at least 30%, and further evaluation is necessary to refine simulation sensitivities to initial conditions and chemical rate uncertainties.

1 Introduction

Total oxidized nitrogen [NOₓ = NO + NO₂ + NO₃ + N₂O₅ + HNO₂ + HNO₃ + HO₂NO₂ + CH₃(CH₂)₆C(O)OOONO₂ + RNO₅] includes many compounds with a wide variety of physical properties and environmental roles. Nitrogen oxides (NOₓ = NO + NO₂) are water insoluble, chemically reactive in the atmosphere, and serve as precursors to ozone. Peroxy nitrates (PNs = HO₂NO₂ + CH₃(CH₂)₆C(O)OOONO₂) are insoluble, their chemical reactivity is temperature dependent, and they act primarily as a reservoir for NOₓ. Nitric acid, on the other hand, is highly water soluble, chemically stable, and is a primary component of acid rain. The partitioning of the NOᵧ between component compounds is controlled by a mix of physical (i.e., emissions and transport) and chemical (i.e., aqueous, particle, and gas-phase) processes and is critical to accurate simulation of environmental stress. The partitioning between NOᵧ compounds influences the efficiency of NOᵧ wet scavenging, the availability of HNO₃ for acid rain, and the amount of NOₓ for production of the greenhouse gas ozone. As a greenhouse gas, ozone is 10 times more efficient in the upper troposphere than in the lower troposphere (Lacis et al., 1990). The upper troposphere, with its high ozone mixing ratio and high radiative
forcing efficiency, is also where chemical transport models (CTMs) underpredicted the NO\textsubscript{x} precursor (Napelenok et al., 2008; Bertram et al., 2007; Singh et al., 2007; Napelenok et al., 2008).

Underprediction of upper tropospheric NO\textsubscript{x} could be caused by any of the interrelated chemical and physical processes in CTMs that affect NO\textsubscript{x} partitioning. Increasing simulated NO\textsubscript{x} from aircraft and lightning increase NO\textsubscript{x} mixing ratios, but does not resolve the bias. Pickering et al. (2009) found that lightning improved NO\textsubscript{x}, but most bias improvement was below 8 km. Hudman et al. (2007) concluded that lightning emissions improved simulated NO\textsubscript{x} mixing ratios, but the median simulated NO\textsubscript{x} mixing ratio was still 300 ppt low-biased and the primary chemical sink (HNO\textsubscript{3}) was now overpredicted. Other emission studies have quantified NO\textsubscript{x} emissions from aircraft (Eyers et al., 2004; Sutkus et al., 2003), which are generally small compared to lightning except perhaps directly in particular flight paths (Hudman et al., 2007). Zero dimensional modeling studies have suggested either missing observations or errors in chemical transformation of radical precursors in the upper troposphere (Olson et al., 2006; Ren et al., 2008). Emissions, physics, and chemistry both contribute to the NO\textsubscript{x} mixing ratios, requiring evaluation of each process in isolation.

This study develops and implements a new evaluation technique designed to isolate simulated chemistry in the upper troposphere. Chemistry evaluation, to date, uses either smog chamber experiments or quasi-Lagrangian measurements. Smog chamber experiments provide a direct evaluation in a controlled environment, but chamber experiments are carried out at surface level temperatures and pressures (T ≈ 298 K, P ≈ 1 atm) and typically high NO\textsubscript{x} mixing ratios (NO\textsubscript{x} > 50 ppb), which are significantly different from the upper troposphere (medians from this study: T = 240 K, P = 0.31 atm, NO\textsubscript{x} = 0.4 ppb). Quasi-Lagrangian aircraft measurements can provide temperature/pressure appropriate time-series case studies, but the Lagrangian nature of the sampling is often difficult to verify given uncertainty in meteorology (Real et al., 2008). Smog chamber evaluations do not have appropriate environmental conditions, and quasi-Lagrangian sampling does not provide enough high-quality samples for statistical evaluation. Any upper tropospheric evaluation must account for both environmental conditions and air parcel interaction with meteorology.

We propose a statistically robust chemical evaluation using in situ upper tropospheric aircraft observations from the Intertropical Chemical Transport Experiment (INTEX-NA; Singh et al., 2006). Although these aircraft measurements do not sample a single air parcel through space and time, the measurements can be grouped and sorted by photochemical age using a technique developed by Bertram et al. (2007). This technique assumes that the “youngest” air parcels are the result of deep convection events. Deep convection mixes air from the earth’s surface into the upper troposphere and is generally associated with precipitation that removes water soluble HNO\textsubscript{3}, but not less soluble NO\textsubscript{x} (Prather and Jacob, 1997; Jaegle et al., 1998). Thus air parcels immediately following convection have very high ratios of NO\textsubscript{x}:HNO\textsubscript{3}. After deep convection, air parcels undergo chemical processing that converts NO\textsubscript{x} to HNO\textsubscript{3}, reducing the NO\textsubscript{x}:HNO\textsubscript{3} ratio until the air parcel is removed from the upper troposphere by convective downdrafts or subsidence. Initial deep convection is identified by high NO\textsubscript{x}:HNO\textsubscript{3} and subsequent removal is modeled stochastically. Therefore, the observed NO\textsubscript{x}:HNO\textsubscript{3} ratio provides a relative metric of time since convection that can be used to create a time-series. This time-series is suitable for evaluating chemistry in the upper troposphere because it has appropriate environmental conditions and enough observations for statistical evaluation.

This study uses a relative time-series of observations to evaluate photochemical aging predicted by seven different chemistry representations. Each chemistry representation, called a chemical mechanism, uses reaction sets with varying degrees and methods of simplification (Dodge, 2000). We selected seven chemical mechanisms from chemical transport models with spatial scales ranging from point to global. The complexity of each chemical mechanism also ranges from near-explicit to condensed. Near-explicit chemical mechanisms represent all known chemical compounds and reactions. Although all known reactions are included, many reactions have large uncertainty in the rate coefficient and stoichiometric yield. Condensed mechanisms use abstractions to reduce the computational load, but often include empirical tuning for conditions that may limit the applicability of the mechanism to all environmental conditions.

We evaluate each chemical mechanism to test three main questions. First, is the rate of chemical aging consistent between chemical mechanisms and observations? Second, are biases consistent for all chemical mechanisms, and therefore, fundamental to the state of the science, or can mechanism differences identify misrepresentations? Third, to what extent can chemical mechanisms’ photochemical aging cause underprediction of NO\textsubscript{x}? Finally, we evaluate factors that contribute to partitioning biases for total oxidized nitrogen in an attempt to improve the individual chemical mechanisms.

\section{Methods}
\subsection{Modeling framework}

This study simulates photochemical aging and physical processing of air parcels following deep convection. Deep convection mixes lower tropospheric air with VOCs and NO\textsubscript{x} into upper tropospheric air (Bertram et al., 2007). Deep convection produces clouds that scavenge water soluble HNO\textsubscript{3} and lightning that produces NO\textsubscript{x}. These two processes result in high NO\textsubscript{x}:HNO\textsubscript{3} ratios that can identify air parcels transported by recent convection. After convection, the air parcel photochemically ages, converting NO\textsubscript{x} to HNO\textsubscript{3}, and
mixes with background upper tropospheric air until it is removed from the upper troposphere by convective downdrafts or subsidence. Particle chemistry is most likely of limited importance in our study due to low particle surface area. In an initial analysis, less than 1% of HNO$_3$ is in the particle phase at equilibrium. Aircraft observe air parcels at varying time since convection and, therefore, with varying extents of photochemical aging. We then developed a model to reproduce the observed distribution of photochemical age. To reproduce the distribution of air parcels, our model framework simulates gas-phase chemistry, photolysis, mixing into background air (i.e., dilution, dispersion, diffusion), and subsequent convection. Subsequent convection is caused by meteorological processes external to our box model, and we simulate this process stochastically using a distribution of time between convective influence. First, we simulate 10 d of chemical aging, or air parcel lifetime, for a variety of physical and chemical conditions representative of recently convected air parcels in the INTEX-NA observational database. In the real environment, we expect that air parcel lifetimes have a distribution that is governed by subsequent convection. We stochastically simulate subsequent convection by optimizing the distribution of air parcel lifetimes for consistency with observed chemical mixing ratios. The air parcel lifetimes can be evaluated against the empirical distribution, and the predicted distribution of chemical species during the air parcel lifetime can be compared to observed mixing ratios.

2.2 Observations

Aircraft observations provide box model initial conditions and photochemical age needed for model evaluation. We first sorted observations using NO$_3$:HNO$_3$ as a chemical indicator of photochemical age. The measurements with the highest NO$_3$:HNO$_3$ ratios provide physical conditions and initial chemical mixing ratios for model simulations using seven chemical mechanisms. The predictions are then evaluated against the observational time-series to assess the performance of simulated chemistry.

We use aircraft observations from the National Aeronautics and Space Administration (NASA) DC-8 aircraft flights during Intercontinental Chemical Transport Experiment – North America (INTEX-NA) campaign (Singh et al., 2006). We started with the 10-s averaged NASA DC-8 observational database (n = 56465). We then filtered the observation database to include only measurements of the upper troposphere (8 km < altitude < 10 km). We exclude air parcels with any fractional cloud presence that would have active wet scavenging, which would influence NO$_3$:HNO$_3$. We also removed air parcels that might have been influenced by stratospheric intrusion (³⁸Be:²¹⁰Pb > 1000) or biomass burning (CH$_3$CN > 200 ppt). The remaining observations fall into two distinct groups: those influenced by polluted air (CO ≥ 80 ppb) and those influenced by background air (CO < 80 ppb) (Singh et al., 2007). Our analysis has been performed with both polluted and background influenced observations and excluding background observations. Both analyses give similar results. In this study, we focused on the influence of polluted air and include only those air parcels with over 80 ppb CO (n = 861). These observations represent upper troposphere air parcels with varying photochemical age.

Observations of NO and HNO$_3$ in the INTEX-NA dataset have known uncertainties and limitations that require adjustment. The NO chemiluminescence measurement has a long 1-minute integration time, is most reliable for mixing ratios greater than 100 ppt (Singh et al., 2007) and, during the INTEX-NA campaign, has a strong bias compared to steady-state NO ($\text{NO}_{\text{SS}}=[\text{NO}_2]/(k_{\text{NO}_2}+k_{\text{NO}_2+\text{HO}_2}[\text{HO}_2])$). For this analysis, we require finer time resolution and detection of low NO (17.7% of [NO]$_{\text{SS}}$ observations are below 100 ppt), so we use the steady-state calculated mixing ratio. During the INTEX-NA study, the University of New Hampshire (UNH) and the California Institute of Technology (CIT) took two separate measurements of HNO$_3$. When observations are available simultaneously between 8 and 10 km, the linear least squares fit of UNH as a function of CIT is 61.7%. The reason for the discrepancy is currently unknown, so we and other researchers (Bertram et al., 2007) adjust UNH by a factor of 1.2 and CIT by a factor of 0.8. The CIT measurement has better time resolution, but less temporal coverage. We use the adjusted CIT measurement when available and fill in measurement gaps with the adjusted UNH measurement. The large discrepancy in the HNO$_3$ measurement could bias our chemical surrogate of age. As a result, we have performed this analysis using CIT, UNH, and the adjusted HNO$_3$ values (see Appendix). The conclusions of this study are robust to the choice of HNO$_3$ measurement. Since the cause of the discrepancy is unknown, we use the adjusted values in the rest of this analysis.

The upper troposphere observations are then divided into age groups according to photochemical age as assessed by NO$_3$:HNO$_3$. The observed NO$_3$:HNO$_3$ ratio in our filtered dataset is log-normally distributed, and we split observations into 4 age groups that are non-overlapping, have comparable sample sizes, and capture the range of air parcel aging. The age groups, which represent relative photochemical age, will be referred to as fresh, young, midage, or old. Each category has a minimum of 215 observations (fresh: 216, young: 215, midage: 215, old: 215). An additional classification, “initial”, was added to capture immediate convection for model initialization. The initial age group includes the youngest 50% of the fresh observations and represents air parcels that have been convected most recently. Figure 1 shows the total oxidized nitrogen (NO$_x$) partitioning of each age group and shows that pernitric acid, PANS, and organic nitrates did not show strong age-dependent mixing ratios.
Fig. 1. Nitrogen partitioning of fresh, young, midage and old age categories demonstrates influence of chemical aging. Each age category has been tested for statistical difference (p < 0.0001) from the preceding age category (∗) and fresh (#).

2.3 Box model

We use a common box model framework for all simulations to remove artifacts of multiple modeling systems and isolate differences between seven chemical mechanisms. The use of a common box model removed variability in ordinary differential equation solvers and physical representations. The Dynamically Simple Model of Atmospheric Chemical Complexity (DSMACC) provided the flexibility and power necessary to model all our chemical mechanisms. The DSMACC model (Emmerson and Evans, 2009) is based on the Kinetic Pre-Processor (KPP) (Sandu and Sander, 2005), which has a flexible rate coefficient representation. The flexible rate representation allowed all seven chemical mechanisms to use their native reaction rate coefficient forms. We have added a mixing process to the DSMACC model to account for dilution, dispersion, and diffusion. Air parcels mix in “background” air where each chemical species mixing ratio is the mean of observations described above. The “background” air includes air parcels influenced by both polluted and background air. The rate of mixing is assumed to be constant and set to 5% per day (Bertram et al., 2007). Sensitivity analysis using up to 10 times the mixing rate, and/or more complex dynamic background and dynamic mixing rates did not yield meaningfully different results (see Fig. A3). This box model represents only gas-phase chemical reactions and mixing with background air; there is no particle or aqueous chemistry.

2.4 Gas-phase chemistry

This study evaluates gas-phase chemistry from seven chemical mechanisms that each have different research goals. Carbon Bond version 2005 (Yarwood et al., 2005) and the State Air Pollution Research Center ’99 (SAPRC99) (Carter, 2000) are typically used for urban to continental simulation. In addition, SAPRC ’07 (Carter, 2009) and Regional Atmospheric Chemical Mechanism version 2 (RACM2) (Stockwell et al., 2008; Goliff and Stockwell, 2008; Goliff et al., 2011) are mechanisms that are planned to be included in the Environmental Protection Agency’s Community Multi-scale Air Quality model. The Goddard Earth Observing System-Chemistry (GEOS-Chem) (Mao et al., 2009) and Model for OZone And Related chemical Tracers (MOZART-4) (Emmons et al., 2010) are typically used for global simulation. The near-explicit LEEDS Master Chemical Mechanism (MCM) (Saunders et al., 1997) is typically used in box model or trajectory simulations. For MCM, we extracted only those chemical reactions that would be active given our initial conditions and subsequent chemical products. The seven chemical mechanisms we evaluate are used for a range of research goals and have a range of computational complexity (see reactions and species in Table 1).

2.5 Photolysis

Gas-phase chemistry simulations typically use different photo-dissociation models that strongly influence radical initiation and photochemical cycling. Each chemical mechanism evaluated in this study is typically used in a host chemical transport model (i.e., CMAQ, GEOS-Chem, MOZART4, and SBOX) with specific photolysis models to calculate photo-dissociation rates. For example, Carbon Bond and SAPRC chemical mechanisms both used the CMAQ photolysis preprocessor (JPROC), GEOS-Chem used FAST-J photolysis (Wild et al., 2000), MCM used the Tropospheric Ultraviolet model (TUV) version 4.2 (Madronich, 2002), RACM2 used a predecessor of TUV, and MZ4 used TUV version 4.6. Not all photolysis models have implemented pressure/temperature sensitivities and the resulting difference in photolysis rates dominated the nitrogen partitioning in our initial tests. Particularly, representation of carbonyl photolysis temperature/pressure dependence led to differences in PAN predictions and the representation of near-IR photolysis (0.00001 s⁻¹; Murphy et al., 2004) of pernitric acid led to diverse predictions. The different photolysis rates were not a function of chemical mechanism, but rather of the photolysis model calculation. To truly focus on chemical mechanism differences, the photolysis rates were standardized using TUV v4.6 with modifications consistent with those used in DSMACC for MCM with one exception. All mechanisms except MCM had photolysis reactions for PAN and HO2NO2 and, for this analysis, PAN and HO2NO2 photolysis has been added to MCM.

2.6 Base simulations

Each chemical mechanism simulates chemical aging for each initial observed air parcels (n₁ = 108). The initial air parcels were used as the initialization of all simulated physical and chemical conditions (see Table 2). The initial observations of chemical species were mapped to their appropriate chemical mechanism species. Where particular chemical compound measurement was not concurrently available, the median of all initial values for that compound was used. An additional
Table 1. Overview of chemical mechanisms in this study.

<table>
<thead>
<tr>
<th>Chemical Mechanism (abbreviation)</th>
<th># Rxns</th>
<th># Spcs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon Bond ‘05 (CB05)</td>
<td>176</td>
<td>62</td>
</tr>
<tr>
<td>State Air Pollution Research Center ‘99 (SAPRC99)</td>
<td>222</td>
<td>77</td>
</tr>
<tr>
<td>State Air Pollution Research Center ‘07 (SAPRC07)</td>
<td>691</td>
<td>153</td>
</tr>
<tr>
<td>Model for Ozon e And Related chemical Tracers “Standard” (MZ4)</td>
<td>196</td>
<td>86</td>
</tr>
<tr>
<td>GEOS-Chem “full” (GEOS-Chem)</td>
<td>286</td>
<td>88</td>
</tr>
<tr>
<td>Regional Atmospheric Chemistry Mech v.2 (RACM2)</td>
<td>349</td>
<td>117</td>
</tr>
<tr>
<td>Master Chemical Mechanism Active Subset (MCM)</td>
<td>4685</td>
<td>1610</td>
</tr>
</tbody>
</table>

Table 2. Median observed values for filtered initial (n_i = 108) and background (n_bkg = 1006) selected chemical compounds and physical conditions.

<table>
<thead>
<tr>
<th>Measured</th>
<th>Background</th>
<th>Initial</th>
<th>Principal Investigator</th>
</tr>
</thead>
<tbody>
<tr>
<td>Altitude</td>
<td>8841 m</td>
<td>9149 m</td>
<td>J. Barrick, NASA LaRC</td>
</tr>
<tr>
<td>Pressure</td>
<td>314.7 hPa</td>
<td>300.6 hPa</td>
<td>M. Avery, NASA LaRC</td>
</tr>
<tr>
<td>Temperature</td>
<td>241.1 K</td>
<td>233.7 K</td>
<td>Derived from NO_2, O_3, and HO_2</td>
</tr>
<tr>
<td>HO</td>
<td>0.5396 pptv</td>
<td>0.6101 pptv</td>
<td>W. Brune, Pennsylvania State University; Adjusted according to Ren et al. (2008)</td>
</tr>
<tr>
<td>HO_2</td>
<td>13.16 pptv</td>
<td>11.24 pptv</td>
<td>G. Huey, Georgia Institute of Technology</td>
</tr>
<tr>
<td>O_3</td>
<td>77.76 ppbv</td>
<td>70.61 ppbv</td>
<td>D. Blake, UC Irvine, and E. Atlas, Univ. of Miami</td>
</tr>
<tr>
<td>NO_2</td>
<td>95.52 pptv</td>
<td>153.6 pptv</td>
<td>P. Wennberg, California Institute of Technology; R. Talbot, Univ. of New Hampshire; Adjusted following Bertram et al. (2007)</td>
</tr>
<tr>
<td>NO</td>
<td>203.3 pptv</td>
<td>411.8 pptv</td>
<td>G. Sachse, NASA LaRC</td>
</tr>
<tr>
<td>HNO_3</td>
<td>280.1 pptv</td>
<td>125.9 pptv</td>
<td>P. Wennberg, California Institute of Technology; B. Heikes, Univ. of Rhode Island; Adjusted following Bertram et al. (2007)</td>
</tr>
<tr>
<td>HNO_2NO_2</td>
<td>82.00 pptv</td>
<td>67.80 pptv</td>
<td>G. Sachse, NASA LaRC</td>
</tr>
<tr>
<td>H_2O_2</td>
<td>234.2 pptv</td>
<td>195.9 pptv</td>
<td>D. Blake, UC Irvine, and E. Atlas, Univ. of Miami</td>
</tr>
<tr>
<td>CO</td>
<td>98.36 ppbv</td>
<td>108.0 ppbv</td>
<td>G. Sachse, NASA LaRC</td>
</tr>
<tr>
<td>CH_4</td>
<td>1.789 ppmv</td>
<td>1.784 ppmv</td>
<td>D. Blake, UC Irvine, and E. Atlas, Univ. of Miami</td>
</tr>
<tr>
<td>C_2H_6</td>
<td>790.0 pptv</td>
<td>800.0 pptv</td>
<td>P. Wennberg, California Institute of Technology; B. Heikes, Univ. of Rhode Island; Adjusted following Bertram et al. (2007)</td>
</tr>
<tr>
<td>C_3H_8</td>
<td>146.0 pptv</td>
<td>153.5 pptv</td>
<td>P. Wennberg, California Institute of Technology; B. Heikes, Univ. of Rhode Island; Adjusted following Bertram et al. (2007)</td>
</tr>
<tr>
<td>C_2H_4</td>
<td>1.500 pptv</td>
<td>1.500 pptv</td>
<td>P. Wennberg, California Institute of Technology; B. Heikes, Univ. of Rhode Island; Adjusted following Bertram et al. (2007)</td>
</tr>
<tr>
<td>Speciated alkyl nitrates (RNO_3)</td>
<td>8.630 pptv</td>
<td>8.630 pptv</td>
<td>P. Wennberg, California Institute of Technology; B. Heikes, Univ. of Rhode Island; Adjusted following Bertram et al. (2007)</td>
</tr>
<tr>
<td>CH_3O</td>
<td>174.5 pptv</td>
<td>437.0 pptv</td>
<td>A. Fried, NCAR; B. Heikes, Univ. of Rhode Island</td>
</tr>
<tr>
<td>CH_3C(O)H</td>
<td>83.80 pptv</td>
<td>117.5 pptv</td>
<td>H. Singh, NASA ARC</td>
</tr>
<tr>
<td>CH_3C(O)CH_3</td>
<td>1475. pptv</td>
<td>1375. pptv</td>
<td>A. Fried, NCAR; B. Heikes, Univ. of Rhode Island</td>
</tr>
<tr>
<td>CH_3C(O)C_2H_5</td>
<td>71.25 pptv</td>
<td>95.00 pptv</td>
<td>H. Singh, NASA ARC</td>
</tr>
<tr>
<td>PAN</td>
<td>374.9 pptv</td>
<td>370.6 pptv</td>
<td>H. Singh, NASA ARC</td>
</tr>
<tr>
<td>CH_3C(O)OOH</td>
<td>172.8 pptv</td>
<td>226.1 pptv</td>
<td>P. Wennberg, California Institute of Technology; B. Heikes, Univ. of Rhode Island; Adjusted following Bertram et al. (2007)</td>
</tr>
</tbody>
</table>

Simulation was generated using the initial age group median value of every chemical compound. Each chemical mechanism was used to simulate 10 d of chemical processing for each initial air parcel and the median air parcel (n_s = n_i + 1). Nighttime simulation results (i.e., solar zenith angles, \( \theta \), higher than 75°) are ignored to be consistent with exclusively daytime observations (i.e., 8 < \( \theta \) < 75°).

2.7 Stochastic convection model description

Our simulations must take into account the frequent exchange between the upper and lower troposphere. Convective updrafts loft air parcels into the upper troposphere which are later removed from the upper troposphere by convective downdrafts or subsidence. The time between convective lofting and removal, hereafter air parcel lifetime \( \tau_{air} \), can be used to calculate the instantaneous or sampled distribution
of time since convection $Pr(t)$ in the upper troposphere. To accurately represent observed upper tropospheric air parcels, we must derive the distribution of time since convection $Pr(t)$ and subset our simulation results accordingly. We estimate the distribution of time since convection using a maximum likelihood technique with one stochastic model and two observationally-derived datasets of time since convection.

The stochastic model for the distribution of air parcel lifetimes and of time since convection $Pr(t)$ is both exponential. From an air parcel’s perspective, encountering a downdraft is a random and time independent event that will have an exponential distribution (Gallager, 1996). If the INTEX-NA observations were an unbiased random sample, Eq. (1) would describe the distribution of observed time since convection, where $\tau_{air}$ is the mean air parcel lifetime. The INTEX-NA observations, however, preferentially sampled freshly convective air parcels. Bertram et al. (2007) showed that 21.4% of air parcels had time since convection less than 6 h, which is far greater than 12.5% of the sampling domain with convective activity during the INTEX-NA campaign. To approximately correct the sample bias, we increase the relative probability of sampling time since convection less than 6 h by a factor of 2. Equation (2) doubles the probability of sampling young ($t < 6$ h) air parcels (real or simulated), and still has only one fitting parameter $\tau_{air}$.

$$Pr(t) = \frac{1}{\tau_{air}} \exp\left(-\frac{-t}{\tau_{air}}\right)$$  \hspace{1cm} (1)

$$Pr(t) = \begin{cases} 
\frac{1}{\kappa(2-\exp\left(\frac{-t}{\tau_{air}}\right))} \exp\left(\frac{-t}{\tau_{air}}\right) & \text{if } t \leq 6 \\
\frac{1}{\tau_{air}(2-\exp\left(\frac{-t}{\tau_{air}}\right))} \exp\left(\frac{-t}{\tau_{air}}\right) & \text{if } t > 6 
\end{cases} \hspace{1cm} (2)$$

where $\kappa = \frac{-6}{\log\left(2\exp\left(\frac{-6}{\tau_{air}}\right)-1\right)}$ and $\tau_{air} \geq 9$. 

We estimate $\tau_{air}$ by fitting our statistical models (Eqs. 1 and 2) to two empirical estimates of INTEX-NA observed time since convection. First, we use back trajectory encounters with convection systems calculated by Fuelberg et al. (2007). Second, we use our statistical model with chemical mechanisms to reproduce NO$_x$:HNO$_3$, a chemical indicator of time since convection. These two approaches, described in detail below, require different assumptions, rely on different models, and provide independent estimates of estimate $\tau_{air}$.

Fuelberg et al. (2007) simulated back trajectories and estimated time since convection, which we use to optimize our statistical model. Fuelberg et al. (2007, Table 3) reported the cumulative distribution function (CDF) of time since convection event at intervals starting at 6 h, and ending at 240 h. At 240 h, 91.8% of observations had encountered convection, which leaves 8.2% of observations with unknown time since convection. As a conservative approach, we fit our time since convection model to both the reported and renormalized CDF and provide the range of results as the back trajectory estimate of $\tau_{air}$. For both the original and renormalized dataset, we find the $\tau_{air}$ (between 1 and 240 h) that minimizes the sum of squared prediction error.

Chemical indicators of time since convection, such as NO$_x$:HNO$_3$, provide a second dataset for determining $\tau_{air}$. The chemical evolution of NO$_x$:HNO$_3$ is reproducible by chemical simulations, using chemical mechanisms, and then subsetting results proportional to Eq. (2). We iteratively subset our base simulations according to the probability of time since convection for each possible $\tau_{air}$ (1–240 h). To maximize the size of each subset, we normalize the probability of time since convection to a percentage (exponential: Eq. 3, bias-corrected: Eq. 4) of simulations at each model output time. Each result subset is an ensemble of simulated NO$_x$:HNO$_3$ with varying initial conditions and time since convection. We then selected the optimal $\tau_{air}$ based on the agreement of the simulation ensemble NO$_x$:HNO$_3$ with observed NO$_x$:HNO$_3$.

$$p(t) = \exp\left(-\frac{-t}{\tau_{air}}\right)$$  \hspace{1cm} (3)

$$p(t) = \begin{cases} 
\exp\left(\frac{-t}{\kappa}\right) & \text{if } t \leq 6 \\
\frac{1}{\tau_{air}} \exp\left(-\frac{-t}{\tau_{air}}\right) & \text{if } t > 6 
\end{cases} \hspace{1cm} (4)$$

where $\kappa = \frac{-6}{\log\left(2\exp\left(\frac{-6}{\tau_{air}}\right)-1\right)}$ and $\tau_{air} \geq 9$.

For each $\tau_{air}$, we then evaluate the agreement of the simulation ensemble with observed NO$_x$:HNO$_3$ using the non-parametric Anderson-Darling K-sample goodness-of-fit statistic (Scholz and Stephens, 1987). The Anderson-Darling test makes no assumptions about data distribution (i.e., skew, kurtosis, etc.), and is particularly sensitive on tails of data distributions. Further, the fit criterion ($A_{LKB}^2$) is inversely proportional to goodness-of-fit, which makes it ideal for optimization. For each chemical mechanism, we minimize the fit criterion to identify the optimal $\tau_{air}$.

### 2.8 Evaluation approach

We derive $\tau_{air}$ using one approach that relies on back trajectory simulation and another that depends on chemical simulation. Using the back trajectory dataset provides an estimate of $\tau_{air}$ that depends on the accuracy of a meteorology model. Using the chemical mechanism approach provides an estimate of $\tau_{air}$ that depends on the modeled NO$_x$ to HNO$_3$ conversion. If these two approaches confirm each other, we gain confidence that the chemical mechanisms are photochemically aging at the same rate as observations. If these two approaches conflict, we further evaluate chemical simulation results for evidence that the chemical aging rate is consistent or inconsistent with observed mixing ratios.
We evaluate simulation results, sampled by optimal $\tau_{\text{air}}$, to test the consistency of chemical aging precursors and products. Chemical aging, here assessed by NO$_x$:HNO$_3$, includes the net production or loss of all oxidation precursors and products. If the chemical aging is consistent with observations, other oxidation precursors and products should also be correctly predicted. Our null hypothesis is that, given the same amount of nitrogen oxidation, simulated and observed mixing ratios will be statistically similar for chemical species that were not used to optimize $\tau_{\text{air}}$. The predicted distributions of mixing ratios for simulated and observed chemical species are statistically compared using a Mann-Whitney-Wilcoxon rank sum test (hereafter rank sum test) (Mann and Whitney, 1947). The rank sum test compares the entire distribution (i.e., not just the mean, median or mode) to test if one is statistically greater than the other. The rank sum test is a non-parametric test and, as such, makes no assumptions about data distribution (i.e., skew, kurtosis, etc). There is no perfect comparison between simulated and measured chemical mixing ratios. For instance, the aircraft observations are time (10 s) and space (1.5 to 3 km) averaged while predictions are instantaneous. The averaging of observations could smooth out some extremes; this is especially true for fast reacting radical species (Olson et al., 2006). To account for some anticipated variation, this study requires a very high degree of confidence to conclude that observations are distinct from model mixing ratios. We only reject the null hypothesis if the probability of the difference in distributions is less than 0.01% ($p < 0.0001$).

3 Results

3.1 Stochastic convection: back trajectory results

The back trajectory estimation technique has four discrete estimates of mean air parcel lifetime ($\tau_{\text{air}}$). Each estimate comes from combining a time since convection dataset, either the unadjusted or renormalized, and a statistical model, either the exponential (Eq. 1) or bias-corrected (Eq. 2) as described in the Stochastic Model Description. Figure 2 shows that renormalizing the back trajectory dataset shortens the $\tau_{\text{air}}$ estimate, while using the bias-corrected statistical model lengthens the $\tau_{\text{air}}$ estimate. Both the renormalized dataset and the bias-corrected statistical model incrementally improve the coefficient of correlation ($R^2$). Using the unadjusted back trajectory results, the exponential model (a) predicts $\tau_{\text{air}}=40 \text{ h}$ and our bias-corrected model (b) predicts $\tau_{\text{air}}=51 \text{ h}$. With renormalized back trajectory results, the exponential model (c) predicts $\tau_{\text{air}}=47 \text{ h}$ and our bias-corrected model (d) predicts $\tau_{\text{air}}=58 \text{ h}$.

3.2 Stochastic convection: NO$_x$:HNO$_3$ results

The chemical mechanism technique of estimating of $\tau_{\text{air}}$ consistently yielded shorter $\tau_{\text{air}}$ values than the back trajectory approach. Figure 3 compares the back trajectory and chemical mechanism $\tau_{\text{air}}$. The shortest $\tau_{\text{air}}$ estimates for all chemical mechanisms was derived using the exponential model (18–23 h). When using the optimized exponential model (Eq. 3), all chemical mechanisms, except SAPRC99 and RAC2, predicted NO$_x$:HNO$_3$ ratios that are statistically different from observations. When the convection model is corrected for sampling bias (Eq. 4), estimated air parcel lifetimes are longer (28–34 h) and NO$_x$:HNO$_3$ compares better with observations. When correcting for sampling bias, Fig. 4 shows that all the chemical mechanisms capture the general shape of the observed NO$_x$:HNO$_3$. As a result, the Anderson-Darling goodness-of-fit test cannot reject the null hypothesis that NO$_x$:HNO$_3$ is consistent with observations ($\alpha < 0.01$). Even though the chemical mechanisms capture the distribution of observed NO$_x$:HNO$_3$, the highest $\tau_{\text{air}}$ estimate is 6 h shorter than the shortest back trajectory estimate. The back trajectory estimates of time since convection are all longer than any estimate by chemical mechanisms. If any of the back trajectory $\tau_{\text{air}}$ estimates are correct, all of the chemical mechanisms too rapidly remove NO$_x$. Because NO$_x$ components NO and NO$_2$ are in steady state, this leads to an underprediction of NO$_2$. We estimate the NO$_2$ low-bias by sampling simulated results using our statistical model of convection optimized with back trajectory time since convection. Even when we sample the simulation results using the lowest $\tau_{\text{air}}$ estimate (40 h), we underpredict NO$_2$ by at least 30%.

3.3 Chemical mixing ratio evaluation

The chemical mechanism and back trajectory $\tau_{\text{air}}$ estimates disagree, suggesting a need to further evaluate predicted oxidation precursors and products. We evaluate oxidation precursors and products to assess our confidence in the chemical mechanism estimate and to understand chemical mechanism
Given the same amount of nitrogen aging, we also expect oxidation precursors to compare well. Chemical mechanisms, however, tended to underpredict quickly-reacting carbonyls acetaldehyde (CH$_3$CHO) and peroxy acetic acid (CH$_2$C(O)OOH) while overpredicting longer-lived species carbon monoxide (CO) and methanol (CH$_3$OH). Acetaldehyde observations, for example, showed no statistical trend, but the predicted mixing ratios decrease with time. All chemical mechanisms underpredict the acetaldehyde magnitude and inter-quartile range almost immediately. Peroxy acetic acid observations also had no statistically significant decrease with time, but predictions bias depended on the chemical mechanism. For peroxy acetic acid mixing ratios, the SAPRC99 mechanism overpredicted, SAPRC07, RACM2 and GEOS-Chem underpredicted, while MZ4 and CB05 performed statistically well. For longer lived carbon monoxide and methanol, all chemical mechanisms overpredicted as early as the young age group.

These long-lived species, particularly methanol (CH$_3$OH) and carbon monoxide (CO), are important because they are alternative indicators of time. Methanol and carbon monoxide are lost exclusively by slow, well-known OH· reactions and have relatively little secondary chemical production in the upper troposphere. The bias in predicted carbon monoxide, when NO$_x$:HNO$_3$ is used as a surrogate for time, is a clear discrepancy. The chemical mechanisms incrementally remove long-lived carbon as a function of integration time, but as a function of NO$_x$:HNO$_3$ there is little integration time difference between age groups. As a result, long-lived carbon is relatively constant between age categories until the parcel is old.

### 3.4 Chemical mechanism biases and recommendations

In several cases, chemical mechanisms had striking biases that can be explained by either modeling assumptions or updates to the kinetic literature. The CB05 mechanism had by far the highest bias for organic nitrates (RNO$_3$), which can be explained by its representation of acetone. Both GEOS-Chem and RACM2 oxidized peroxy acetic acid much faster than the other chemical mechanisms, which can be explained by the choice of kinetic surrogate. All mechanisms overpredict peroxy nitric acid during the young age group, which can be improved by updating the OH· rate constant. Each of these issues is explored in detail below, and implemented to see the change in estimated air parcel lifetime ($\tau_{\text{air}}$).

The CB05 simulations partition up to 25% of total nitrogen into RNO$_3$, but all chemical mechanisms that explicitly represent acetone predict less than 3%. The RNO$_3$ production is a sink for both HO$_x$ and NO$_x$, decreasing availability of OH· and NO$_2$, which leads to CB05 predicting the lowest HNO$_3$. The overprediction of RNO$_3$ by CB05 is a result of structural lumping that combines acetone into the model species PAR. The CB05 PAR species holds all singly bonded carbon, but also holds all carbon from acetone (Yarwood et al.,...
Fig. 4. Observed NO$_x$:HNO$_3$ (bars) compared to simulated (lines) from each chemical mechanism using the optimized, bias-corrected statistical model.

Fig. 5. Simulation results (circle: median; bars: 0, 25, 75, 100 percentiles) and observations (box and whisker) binned by NO$_x$:HNO$_3$. For observations, each age category is superscripted for statistical difference ($p < 0.0001$) from the preceding ($*$) and fresh (#) age group. For model predictions, the median for each chemical mechanism is left hollow when statistically different ($p < 0.0001$) from the observations.
Acetone has a long lifetime and high mixing ratios in the upper troposphere, so it can dominate the carbon in PAR (see Fig. 6). The PAR + OH· organic nitrate yield, however, is based on urban, surface PAR reactivity (i.e., primarily alkanes). In CB05, the PAR + OH· reaction creates an operator species (XO2N), directly (13%) and indirectly (3%), that yield >10% organic nitrates production. In contrast, explicit representation of acetone in GEOS-Chem yields 3.6% organic nitrates. In the upper troposphere where acetone is the dominant PAR contributor, the organic nitrate fraction would have to be adjusted or acetone would need to be handled explicitly). A simple adjustment in CB05 of organic nitrate yield to 3% (as in GEOS-Chem) improves organic nitrate yield significantly and increases the \( \tau_{\text{air}} \) to 40 h, which is also the lower bound back trajectory \( \tau_{\text{air}} \) estimate.

GEOS-Chem and RACM2 predict a median peroxy acetic acid (CH3C(O)OOH) mixing ratio less than the observed 25th percentile by the young age group. Peroxy acetic acid is the second largest acyl peroxy radical source (i.e., PAN precursor) in the first 6 h of simulated aging. The primary loss pathway for peroxy acetic acid is reaction with OH·, but the OH· rate coefficient is not available in the literature. Both chemical mechanisms that underpredict peroxy acetic acid choose methyl peroxide as a surrogate compound for the OH· rate coefficient. The chemical mechanisms that perform better, however, use the acetic acid OH· rate coefficient. At upper tropospheric temperature and pressure, the acetic acid rate coefficient reported by Sander et al. (2006) (not updated from 2003 report see errata) and Atkinson et al. (2006) are both roughly ten times lower than the methyl peroxide OH· rate. Preliminary peroxy acetic acid OH· rate studies confirm the \( k_{\text{OH}} \) similarity to CH3C(O)OH (Orlando and Tyndall, 2002).

Pernitric acid is overpredicted by all chemical mechanisms and acts as an important radical sink in the upper troposphere. In the upper troposphere, pernitric acid that is formed (\( \text{HO}_3 + \text{NO}_2 \rightarrow \text{HO}_2\text{NO}_2 \)) is thermally stable, and the primary loss is OH· + \( \text{HO}_2\text{NO}_2 \) (see Fig. 7). The net pernitric acid reaction consumes two \( \text{NO}_3 \) radicals (Wennberg et al., 1998) and, in this study, this net reaction accounts for 29% of the radicals terminated in the first 6 h. We recommend using the latest \( k_{\text{OH} + \text{HO}_2\text{NO}_2} \) (Jimenez et al., 2004) which improves \( \text{HO}_2\text{NO}_2 \) agreement with observations and increases competition of pernitric acid with \( \text{NO}_2 \) for OH· radicals. Even with this recommendation, the pernitric acid reaction rates have large uncertainties at low temperatures and laboratory studies are restricted to temperatures above those typical in the upper troposphere (Atkinson et al., 2004; Gierczak et al., 2005; Sander et al., 2006).

The peroxy acetic acid and peroxy nitrate recommendations implemented together into our working version of GEOS-Chem. These changes improve peroxy acetic acid and pernitric acid predictions, and increase the \( \tau_{\text{air}} \) estimate from 32 to 34 h. The new predicted \( \text{NO}_3\cdot\text{HNO}_3 \) is now statistically consistent with observations at the \( p < 0.01 \) level. Despite the improved \( \text{NO}_3\cdot\text{HNO}_3 \), the \( \tau_{\text{air}} \) estimate is still 6 h shorter than the lowest back trajectory estimate, and the marginally longer \( \tau_{\text{air}} \) has little affect on long-lived carbon.

4 Discussion

The evidence gathered here suggests that the chemical mechanisms photochemically age \( \text{NO}_x \) too quickly. First, all chemical mechanism estimates of air parcel lifetime, the time necessary to age \( \text{NO}_x \), are at least 15% shorter than the shortest back trajectory estimate. Second, the chemical mechanism air parcel lifetime estimates are insufficient to remove long-lived carbon, as seen in observations. Given these discrepancies, we conclude that chemical mechanisms will be low-biased for \( \text{NO}_x \) in the upper troposphere at any given time since convection.

We investigated individual chemical mechanism biases to develop and test recommendations. Peroxy acetic acid, a peroxy acetyl nitrate precursor, is removed too quickly by chemical mechanisms that use methyl peroxide; we recommend all mechanisms use acetic acid as a surrogate until a specific rate is available. Peroxy nitrates are an important radical sink in the upper troposphere, and we recommend several updates. The primary peroxy nitrate loss reactions in the upper troposphere are photolysis and hydroxyl attack. For photolysis, we recommend that all chemical mechanisms include photolysis for PANs and pernitric acid, and that pernitric acid near IR photolysis be included. For hydroxyl attack, we recommend updating the OH· reaction rate (Jimenez et al., 2004). Finally, explicit or targeted parameterization of acetone is necessary to properly model radical cycling in the upper troposphere. Improved representation of acetone will decrease overpredictions of alkyl nitrates, which will alter radical cycling and total oxidation. The recommendations improved target species predictions and increased air parcel lifetime, but did not solve overpredictions of long-lived carbon.

This analysis included only gas-phase chemistry and ignores heterogeneous processing that also affects the \( \text{NO}_3\cdot\text{HNO}_3 \) ratio. Including \( \text{N}_2\text{O}_5 \) heterogeneous hydrolysis
would exacerbate the rate of NO$_x$ to HNO$_3$ conversion (Jaeglé et al., 1998; Olson et al., 2001; Evans and Jacob, 2005). Mineral dust and ice particle uptake of HNO$_3$ would buffer or counteract the effect of N$_2$O$_5$ hydrolysis. Our initial analysis of HNO$_3$ uptake suggests that this rate would be small compared to nitric acid production NO$_2$ + OH$. We intend to evaluate heterogeneous processing in more depth along chemical rate analysis there.

During this study, several best practices for atmospheric chemical modeling became apparent. The chemical system is very sensitive to the photolysis rates, and so it is critical to simulate photolysis in a detailed way when evaluating the chemical mechanisms. Photolysis simulations need to represent up-to-date pressure/temperature sensitivities. For instance, two models evaluated for use in this study did not include temperature/pressure sensitivities, which are critical in the upper troposphere. To accurately simulate temperature/pressure sensitivities, photolysis rates need to be calculated at the chemical transport model vertical resolution. Photolysis rates of many species (e.g. ozone) exhibit complex shape throughout the troposphere and linear interpolation can drastically underpredict local minima and maxima. Coarse resolution in some photolysis preprocessors is most likely a hold over from historically coarser CTM vertical resolutions. Also, ensure that the chemical mechanism used accounts for PAN photolysis and near-IR HO$_2$NO$_2$ photolysis. Photolysis is the dominant PAN chemical loss process in the upper troposphere, where many have reported PAN overprediction (Pickering et al., 2009; Yu et al., 2010; Fang et al., 2010).

5 Conclusions

This study uses a new probabilistic approach to isolate simulated chemistry for evaluation in the upper troposphere. This approach uses a large number of observations for statistical power and parameterizes processes whose stochastic nature precludes box model simulation. Parameterizing all other processes isolates gas-phase chemistry and produces an ideal modeling system for evaluation in the upper troposphere. Other upper troposphere gas-phase evaluations rely on steady-state assumptions or quasi-Lagrangian measurements. Steady-state assumptions may not be valid in the upper troposphere because convective mixing constantly perturbs NO$_x$ and radical mixing ratios (Prather and Jacob, 1997). Quasi-Lagrangian analysis provides a direct evaluation approach when sufficient observations are available and their Lagrangian nature can be confirmed. Both the quasi-Lagrangian approach and our probabilistic approach have benefits that can complement each other to strengthen our body of knowledge where time-series observations from a single air parcel (e.g. smog chamber experiments) are not available.

One specific goal of this study was to characterize the contribution of chemistry to upper troposphere underprediction of NO$_3$. The results presented here confirm previously reported NO$_2$ underpredictions, and do so in an isolated chemistry model. All evaluated chemical mechanisms converted NO$_3$ to HNO$_3$ too rapidly and, consequently, underpredicted NO$_2$ by at least 30%. Even if all emissions, physical transport, and aqueous-phase chemistry were accurately simulated by a chemical transport model, gas-phase chemistry would cause model underpredictions of NO$_2$.

This paper isolates chemistry and establishes NO$_2$ bias caused by chemistry. While this work does not resolve the problem, the modeling framework described provides a test.
Table A1. Optimization results for stochastic convection using the pure exponential model and the model with correction for preferential sampling. Table includes optimal air parcel lifetime ($\tau_{\text{kaN}}$) and Anderson Darling goodness-of-fit test value ($T_{\text{kaN}}$) for alternative background mixing rate sensitivities. The predicted NO$_x$:HNO$_3$ is statistically different than observations when $T_{\text{kaN}}$ is greater than 3.752 ($\alpha = 0.01$).

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>exponential $\tau_{\text{air}}$</th>
<th>$T_{\text{kaN}}$</th>
<th>corrected $\tau_{\text{air}}$</th>
<th>$T_{\text{kaN}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>GEOS-Chem</td>
<td>23</td>
<td>9.9</td>
<td>33</td>
<td>2.77</td>
</tr>
<tr>
<td>2 \times Mix</td>
<td>21</td>
<td>10.6</td>
<td>32</td>
<td>3.62</td>
</tr>
<tr>
<td>4 \times Mix</td>
<td>20</td>
<td>11.6</td>
<td>31</td>
<td>5.59</td>
</tr>
<tr>
<td>10 \times Mix</td>
<td>18</td>
<td>15.9</td>
<td>28</td>
<td>10.2</td>
</tr>
<tr>
<td>GC*</td>
<td>24</td>
<td>8.32</td>
<td>26</td>
<td>2.28</td>
</tr>
<tr>
<td>M10 \times Init</td>
<td>28</td>
<td>65.9</td>
<td>43</td>
<td>51</td>
</tr>
<tr>
<td>DynMix</td>
<td>24</td>
<td>7.42</td>
<td>36</td>
<td>2.28</td>
</tr>
</tbody>
</table>

environment for further analysis. Initial analyses demonstrate that results presented here are robust to uncertainty in initial conditions, but that rate expression uncertainty can meaningfully slow chemical mechanism NO$_x$ aging. Future research will conduct sensitivity tests to identify key rate expressions. The results from subsequent sensitivity tests should be used to direct gas-phase rate research that will improve state of the science chemical mechanisms.

A1 Observation

The spatial distribution of all age groups (defined in the paper) are shown in Fig. A1.

A2 Subsequent removal

Upper troposphere air parcel lifetimes are limited by subsequently encountered convection and large scale subsidence. Air parcels in the UT subside along isentropic surfaces, but not as rapidly as they are removed by convection related processes (Prather and Jacob, 1997; Jaegle et al., 1998). The importance of convection is most clear in the tropics where convection is very frequent. To confirm the importance of convection during the INTEX-NA, we simulated back trajectories a 12 locations forming a grid over the Northeastern United States using the Hybrid Single Particle Lagrangian Integrated Trajectory Model (HYSPLIT Draxler and Hess, 1997). During a 84 h (70% of the time between INTEX-NA convective events Fuelberg et al., 2007) back trajectory with only isentropic vertical motion, Fig. A2 shows that only 3 of the 12 simulations originated below 8 km or above 10 km. This confirms our conceptual model of convection as the dominant removal process of air parcels from the upper troposphere.

A3 Alternate background mixing scenarios

Our analysis uses background mixing calculated by Bertram et al. (2007), but there are significantly higher literature values. Figure A3 shows 5 mixing sensitivity analysis. We test the sensitivity of our analysis by scaling our mixing parameter by 2, 4, and 10. Standard mixing is 5% per day, so these scaling values evaluate to 10%, 20%, and 50%. Twenty percent is the upper bound of values found in the literature (Bertram et al., 2007, and references therein) and 50% is used to demonstrate the influence of drastically increasing mixing. We also test the possibility of variable mixing efficiency and variable boundary conditions. This dynamic mixing test (DynMix) has mixing efficiency of 50% per day in the initial age group, 25% per day in the fresh age group, and 5% in midage and old age groups. These initial and fresh air parcels vigorously mix in chemical mixing ratios set by the air parcel’s initial conditions. When the air parcel transitions to young age status, I decrease the mixing to 5 \times (also tried 6 \times) and start mix (1:1) of initial and background air, where background air is the average mixing ratio of all observations. When the air parcel transitions to midage, I return to the standard mixing rate and mix in “background” air. This is an extreme assumption because surrounding parcels should also be aging during the fresh time period. The air parcel lifetimes and the predicted NO$_x$:HNO$_3$ evaluation are shown in Table A1.

A4 Implications of measurement discrepancy for nitric acid

The disagreement between HNO$_3$ measured by CIT and UNH has the potential to influence our mean air parcel lifetime. The CIT and UNH measurements are not always coincident in time, so analysis using either has fewer total observations. When using just the CIT measurements, there are only 507 total observations and only 65 initial conditions. When using the UNH measurements, there are 842 total observations and 107 initial conditions. The CIT and UNH measurements both have log-normal distributions. The CIT distribution is broader and less smooth than the UNH distribution (see Fig. A4).

Our modeling framework is able to capture NO$_x$:HNO$_3$ distribution using the UNH and CIT measurements. Both the UNH and CIT mean air parcel lifetimes ($\tau_{\text{UNH}}$ and $\tau_{\text{CIT}}$) are shorter than predicted with adjusted values (see Fig. A5). Using the lower UNH measurements increases the NO$_x$:HNO$_3$ values and shifts the NO$_x$:HNO$_3$ age bin cut-points to higher values. Because the chemistry model tends to overpredict conversion of NO$_x$ to HNO$_3$, the higher NO$_x$:HNO$_3$ observations and higher cut-points cause $\tau_{\text{UNH}}$ to be shorter than the standard $\tau_{\text{air}}$. Using the higher CIT measurements increases NO$_x$:HNO$_3$ values and shifts the NO$_x$:HNO$_3$ age bin cut-points to lower values. With lower cut-points, we might expect an increased $\tau_{\text{CIT}}$ relative to the adjusted values. The
Fig. A1. Map (a) and distribution (b, c) of spatial locations of aircraft observations categorized by age groups (initial, fresh, young, mid-age, and old). Age group definitions are shown in Fig. 4.

Fig. A2. HYSPLIT back trajectories for 12 northeast locations at 9 km altitude with only more than half of the air parcels originating between 8 and 10 km.
Fig. A3. Same as Fig. 5, but for GEOS-Chem with standard and alternate background mixing.

Fig. A4. Observed NO$_x$:HNO$_3$ (bars; left: CIT, right: UNH) compared to simulated (lines) from the GEOS-Chem chemical mechanism using the optimized, bias-corrected statistical model.

Fig. A5. Estimated mean air parcel lifetimes ($\tau_{\text{air}}$) derived from back trajectory and chemical simulation. Chemical simulations use our updated GEOS-Chem mechanism with the adjusted HNO$_3$, the unadjusted CIT measurement, or the UNH unadjusted measurement. Asterisks indicate whether chemically simulated NO$_x$:HNO$_3$ is statistically consistent with observations ($\alpha < 0.01$) when using the exponential (left, Eq. 3) and bias-corrected (right, Eq. 4) statistical models.
CIT measurements, however, extend the distributions low-value tail that the model cannot capture without over predicting values from 0.2 to 0.5. Because our goodness-of-fit statistic is sensitive to the whole distribution, $\tau_{CTP}$ is 1 hour shorter than when using adjusted HNO$_3$ values.

Using either measurement by itself increases the discrepancy between $\overline{\tau_{air}}$ inferred from back trajectories and $\overline{\tau_{air}}$ inferred from chemical simulation. By increasing the discrepancy, using either measurement by itself increases our estimation of the chemistry-based NO$_2$ low-bias.

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