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Evaluation of nitrous acid sources and sinks in urban outflow

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HIGHLIGHTS
- A two-layer box model evaluates HONO sources, sinks in outflow of Dallas–Fort Worth.
- Monte Carlo simulation is applied to scenarios with 3 recently identified sources.
- Improved model outcomes result from inclusion of 2 of 3 recently identified sources.
- A substantial unknown source is still required for agreement with observation.
- Missing HONO source is moderately correlated with jNO2, weakly correlated with NO2.

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ABSTRACT
Intensive air quality measurements made from June 22–25, 2011 in the outflow of the Dallas–Fort Worth (DFW) metropolitan area are used to evaluate nitrous acid (HONO) sources and sinks. A two-layer box model was developed to assess the ability of established and recently identified HONO sources and sinks to reproduce observations of HONO mixing ratios. A baseline model scenario includes sources and sinks established in the literature and is compared to scenarios including three recently identified sources: volatile organic compound-mediated conversion of nitric acid to HONO (S1), biotic emission from the ground (S2), and re-emission from a surface nitrite reservoir (S3). For all mechanisms, ranges of parametric values span lower- and upper-limit values. Model outcomes for ‘likely’ estimates of sources and sinks generally show under-prediction of HONO observations, implying the need to evaluate additional sources and variability in estimates of parameterizations, particularly during daylight hours. Monte Carlo simulation is applied to model scenarios constructed with sources S1–S3 added independently and in combination, generally showing improved model outcomes. Adding sources S2 and S3 (scenario S2/S3) appears to best replicate observed HONO, as determined by the model coefficient of determination and residual sum of squared errors ($r^2 = 0.55 \pm 0.03, SSE = 4.6 \times 10^5 \pm 7.6 \times 10^4$ ppt$^2$). In scenario S2/S3, source S2 is shown to account for 25% and 6.7% of the nighttime and daytime budget, respectively, while source S3 accounts for 19% and 11% of the nighttime and daytime budget, respectively. However, despite improved model fit, there remains significant underestimation of daytime HONO; on average, a 0.15 ppt/s unknown daytime
1. Introduction

Atmospheric nitrous acid (HONO) is important due to the role of HONO in generation of the hydroxyl radical (OH). There are a number of known sources of OH in the troposphere; however, OH production from HONO is of interest because the sources, fate, and diurnal cycling of HONO in the atmosphere have only recently begun to be elucidated. Models of atmospheric HONO generally employ a mass balance approach that allows evaluation of the HONO budget, often with a potentially limiting photostationary state assumption. As summarized by Spataro and Ianniello (2014), models generally include sources, sinks, and transport, the last relevant as formation processes hypothesized to occur at the ground result in vertical gradients of HONO.

Homogeneous and heterogeneous reactions, as well as direct emission of HONO from combustion sources, contribute to the presence of HONO in the troposphere (Finlayson-Pitts and Pitts, 1999). Nitrous acid strongly absorbs sunlight at wavelengths shorter than 390 nm resulting in photolytic degradation to OH and nitric oxide (NO). This results in suppressed, but non-zero, mixing ratios of daytime HONO due to the presence of daytime sources (Kleffmann, 2007). At night, the absence of this photolytic loss mechanism results in HONO accumulation, generally on the order of 0.1 ppb—10 ppb (Kleffmann et al., 2003; Su et al., 2008; Young et al., 2012). The resumption of HONO photolysis after sunrise can lead to substantial formation of OH in the early morning. Aicke et al. (2003) report that during the BERLIOZ investigation at a rural, lightly trafficked site with low anthropogenic emissions during the summer months, photolysis of HONO was the dominant source of OH in the morning, and contributed as much as 20% of 24-h integrated OH production.

Modeling studies generally show the need for an unknown daytime source to close the HONO budget (Stoffelbach et al., 1997; Lee et al., 2015). A number of photochemically driven homogeneous reactions have been identified or considered: e.g., the known reaction of OH and NO and the hypothesized reaction of photolytically excited nitrogen dioxide (NO₂) and water (Li et al., 2008). The latter, however, may not proceed sufficiently rapidly or at adequate yields to affect HONO mixing ratios in the atmosphere (Carr et al., 2009). Other potential homogeneous sources are under discussion and review. For example, Li et al. (2014) proposed an internal source of HONO that consumed nitrogen oxides, although follow up discussion and further experiments indicate the source was likely strongly overestimated (Li et al., 2015; Ye et al., 2015).

Nitrous acid formation mediated by aerosol surface area (SA) is a topic of ongoing research, largely because the complexity of aerosol sources results in substantial uncertainty regarding their ultimate role in HONO formation. Static surfaces such as the ground (Stemmler et al., 2006) also may enhance HONO formation. Other hypothesized daytime sources include emissions resulting from acid/base chemistry in soils (Su et al., 2011) and photolysis of nitric acid (HNO₃) on forest canopy surfaces (Zhou et al., 2011). Photo-enhanced conversion of NO₂ on organic surfaces, including the ground and aerosols, are also thought to contribute to the daytime HONO budget (George et al., 2005; Stemmler et al., 2006, 2007).

Given the many identified and proposed HONO source and sink mechanisms, single value estimates of parameterizations of HONO sources and sinks limit the ability to understand the impact of variability in multiple input parameters on models of HONO dynamics in the atmosphere. Monte Carlo simulation (MCS) provides a tool to observe the combined effects of ranges of input parameters and the resulting impact on the agreement between model output and measurements. In this work, we identify fourteen HONO sources or sinks established in the literature, including three sources that have recently (2013—2014) been identified. We evaluate these recently identified sources through incorporation into a baseline model with a full-factorial, deterministic screening analysis. We then identify scenarios for which we stochastically parameterize source and sink mechanisms with MCS to determine probability distributions of modeled HONO mixing ratios.

2. Methods

2.1. Measurements

Measurements of gas- and particle-phase constituents were made from May 30 to July 1, 2011 in a semi-urban area approximately 68 km northwest of the Dallas—Fort Worth (DFW) metropolitan area. The monitoring site was co-located with the Texas Commission on Environmental Quality Eagle Mountain Lake (EML) continuous ambient monitoring station (CAMS 75). Further details regarding the geography, surrounding industrial and biogenic activities, and site conditions have been outlined previously (Rutter et al., 2015).

Temperature, humidity (Vaisala, HMP-45C in a RM Young 10-plate solar radiation shield), and planetary boundary layer (PBL) height (Vaisala, CL31) were measured throughout the duration of the campaign. Mixing ratios of HONO and HNO₃ were measured every 5 min using a method that coupled a mist chamber with ion chromatography (Dionex, CD20-1), described in greater detail elsewhere (Dibb et al., 2004). First-order photolysis rate constants (k-values) were determined with radiometric measurements of actinic flux determined with a 2-pi double monochrometer with photomultiplier and subsequent calculations following IUPAC recommendations. Nitrogen oxides were recorded every minute using a chemiluminescence trace level NO–NO₂–NOx analyzer (Thermo Electron Corp., Model 42C) equipped with a Blue Light Converter (Air Quality Design, Inc.) for NO₂ quantification. Hydroxyl radical was observed using atmospheric pressure chemical ionization mass spectrometry (Kim et al., 2013). One-hour averaged mixing ratios of volatile organic compounds (VOCs) were measured using a thermal desorption gas chromatograph with flame ionization detection (Perkin–Elmer O₂ Precursor Analyzer System). Continuous measurements of number-based particle size distributions (diameter range of 20 nm–500 nm) were made every 10 min with a scanning electrical mobility sizer (SEMS, Brechtel Inc. Model 2002) and were converted to SA distributions assuming spherical particles. Concentrations of particulate phase nitrate were determined with an Aerodyne high-resolution time-of-flight aerosol mass spectrometer, as described by Rutter et al. (2015). Black carbon concentrations were measured using an aethalometer.
2.2. Baseline model

A two-layer box model describing HONO mixing ratios was developed, with the height of the first layer set to 36 m to represent a surface layer and the height of layer 2 set to 72 m to facilitate use of HONO observations above the surface layer that are available in the literature. Established source (labeled as ‘B1–B8’ in Table 1) and sink mechanisms (labeled ‘L1–L3’ in Table 1) are described in full in the Supporting Information (SI) (including Figs. S1–S5 and Eqs. S1–S20). The timeframe selected for continuous modeling was 22 June 01:00 to 25 June 14:00 (all times local) based on the longest uninterrupted period during the campaign with observations of HNO3, HONO, aerosol SA, NO2, NO, gas-phase chloride (assumed to be hydrochloric acid, HCl), and HONO. Mixing ratios of constituents during this period were generally typical of the broader study period, Eq. (1) describes baseline sources and sinks modeled with a transient approach:

\[
\frac{d[HONO]_{\text{trans}}}{dt} = F_{B1} + F_{B2} + F_{B3} + F_{B5} + F_{B6} + F_{B7} + F_{B8} - (F_{L1} + F_{L2} + F_{L3}) - \Psi_{\text{trans}}
\]  

(1)

where \([HONO]_{\text{trans}}\) is the mixing ratio of HONO from modeled transient sources and sinks (ppt), \(dt\) is the time step (s) between measurements for which observations of all constituents present in Eq. (1) were made, \(F\) represents the source or sink strength of the indicated mechanism (ppt/s), and \(\Psi_{\text{trans}}\) is the loss (or source) of HONO from layer 1 to (or from) layer 2 due to vertical transport (ppt/s).

Eq. (1) describes the transient processes occurring in the model; source B4 was incorporated into the model after accounting for transient processes as shown in Eq. (2):

\[
[HONO]_{\text{total}} = [HONO]_{\text{trans}} + f_{\text{emiss}}\Delta[NOx]
\]  

(2)

where \([HONO]_{\text{total}}\) is the mixing ratio of HONO at a time step resulting from transient and instantaneous processes (ppt) and \(f_{\text{emiss}}\) is the direct HONO emission factor described in Table 1. Eq. (2) may overestimate the contribution of B4 in a box-model, as during the daytime, HONO will rapidly photolyze prior to the measurement of emitted NOx.

Vertical transport, \(\Psi_{\text{trans}}\) (ppt/s), is calculated using a first-order flux-gradient relationship simulated with the 1D CACHE model (Bryan et al., 2012) where mass is transported by eddy diffusion at a magnitude proportional to the eddy diffusivity for heat (\(K_h\)), shown in Eq. (3):

\[
\Psi_{\text{trans}} = -K_h(z,t) \frac{\partial[C(z,t)]}{\partial z} \frac{1}{h}
\]  

(3)

where \(K_h(z,t)\) is the eddy diffusivity (m²/s) at height \(z\) (m) and time \(t\). As shown in Eq. (3), estimates of flux are divided by \(h\), the height of the second layer in the model (m), prior to inclusion in Eq. (1).

Two 1D simulations during the campaign were used to derive \(K_h\), including one simulation for 7–9 June and one for 10–12 June. For the layers corresponding to the upper boundary that are used in the results here, \(K_h\) is derived based on a length scale, vertical wind shear, and a stability parameter (Forkel et al., 1990). It is calculated at each time step within the model, providing a diurnal cycle that is based on meteorological conditions during the campaign.

Observations of HONO were made at one elevation, approximately 10 m above surface, and were used to represent the HONO mixing ratio in layer 1 of the model. Eq. (3) requires an estimate of the HONO mixing ratio in layer 2 to estimate the HONO gradient. Three scenarios were considered: 1) no gradient (i.e., [HONO] in layer 1 equals that in layer 2 at all times); 2) a gradient created using fractions of [HONO] presented in VandenBoer et al. (2013), representative of a stronger nighttime gradient and a weaker daytime gradient (GrN); and 3) a gradient created from fractions of [HONO] presented in Villena et al. (2011) that is representative of a stronger daytime gradient and weaker nighttime gradient (GrD). Diurnal profiles of the three gradient conditions are shown in Fig. S6 of the SI and implications of this limitation are discussed in Section 3.2.

2.3. Parameterization and evaluation of newly identified HONO sources

Three recently identified HONO source mechanisms were parameterized to assess the potential of these mechanisms (in conjunction with B1–B8 and L1-L3) to independently or jointly account for HONO mixing ratios observed in DFW. The three mechanisms, listed in Table 1 as S1, S2, S3 are incorporated into Eq.

Table 1

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>ID Parameter</th>
<th>Lower-limit</th>
<th>Likely</th>
<th>Upper-limit</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aerosol uptake of NO2</td>
<td>B1 ( \gamma_{NO2} ) (–)</td>
<td>( 2.0 \times 10^{-7} )</td>
<td>( 1.0 \times 10^{-6} )</td>
<td>( 5.0 \times 10^{-6} )</td>
<td>Kleffmann et al. (1998); Aumont et al. (2003)</td>
</tr>
<tr>
<td>Photoenhanced aerosol uptake of NO2</td>
<td>B2 ( \gamma_{NO2_{hv}} ) (–)</td>
<td>( 4.0 \times 10^{-6} )</td>
<td>( 1.0 \times 10^{-5} )</td>
<td>( 1.0 \times 10^{-5} )</td>
<td>Stemmler et al. (2007); Wong et al. (2013)</td>
</tr>
<tr>
<td>Photoenhanced conversion of NO2 soot</td>
<td>B3 ( \gamma_{NO2_{soot}} ) (–)</td>
<td>( 4.0 \times 10^{-7} )</td>
<td>( 5.0 \times 10^{-7} )</td>
<td>( 6.0 \times 10^{-7} )</td>
<td>Monge et al. (2010)</td>
</tr>
<tr>
<td>Direct HONO emission</td>
<td>B4 ( f_{\text{trans}} ) (3V, ( \Delta[\text{HONO}] ))</td>
<td>( 0.0029 )</td>
<td>( 0.0055 )</td>
<td>( 0.0080 )</td>
<td>Kirchstetter et al. (1996); Kurtenbach et al. (2001)</td>
</tr>
<tr>
<td>OH + NO</td>
<td>B5 ( k_{\text{NO}}(T) ) (cm³ molec⁻¹ s⁻¹)</td>
<td>( 3.0 \times 10^{11} )</td>
<td>( 3.6 \times 10^{11} )</td>
<td>( 4.3 \times 10^{11} )</td>
<td>NASA (2011)</td>
</tr>
<tr>
<td>HONO from surface HNO3 photoysis</td>
<td>B6 ( f_{\text{HONO, HNO3}} ) (s⁻¹)</td>
<td>( 1.0 \times 10^{-3} )</td>
<td>( 1.2 \times 10^{-3} )</td>
<td>( 1.4 \times 10^{-3} )</td>
<td>Zhou et al. (2003)</td>
</tr>
<tr>
<td>HONO from NO2 conversion at ground</td>
<td>B7 ( \gamma_{NO2_{2,NO}} ) (–)</td>
<td>( 1.0 \times 10^{-6} )</td>
<td>( 5.0 \times 10^{-6} )</td>
<td>( 1.0 \times 10^{-5} )</td>
<td>Kleffmann et al. (1998); Kurtenbach et al. (2001)</td>
</tr>
<tr>
<td>Photoenhanced NO2 conversion at ground</td>
<td>B8 ( \gamma_{NO2_{2,NO}} ) (–)</td>
<td>( 1.7 \times 10^{-5} )</td>
<td>( 2.0 \times 10^{-5} )</td>
<td>( 6.0 \times 10^{-5} )</td>
<td>Stemmler et al. (2006); Wong et al. (2013)</td>
</tr>
<tr>
<td>HONO photolysis &amp;</td>
<td>( f_{\text{HONO, HNO3}} ) (s⁻¹)</td>
<td>( 3.6 \times 10^{-2} )</td>
<td>( 5.8 \times 10^{-2} )</td>
<td>( 8.3 \times 10^{-2} )</td>
<td>Rutter et al. (2014)</td>
</tr>
<tr>
<td>Biotic release, ground</td>
<td>S1 ( f_{\text{Biotic, NO2}} ) (ppt/s)</td>
<td>( 3.6 \times 10^{-2} )</td>
<td>( 5.8 \times 10^{-2} )</td>
<td>( 8.3 \times 10^{-2} )</td>
<td>Osvald et al. (2013)</td>
</tr>
<tr>
<td>Re-emission from NO2(p) reservoir</td>
<td>S2 ( f_{\text{NO2(p)}} ) (molec cm⁻² s⁻¹)</td>
<td>( 1.0 \times 10^{-2} )</td>
<td>( 9.0 \times 10^{-2} )</td>
<td>( 2.0 \times 10^{-1} )</td>
<td>VandenBoer et al. (2014)</td>
</tr>
<tr>
<td>HONO uptake at ground</td>
<td>L1 ( \gamma_{\text{HONO, gr}} ) (–)</td>
<td>( 1.0 \times 10^{-6} )</td>
<td>( 2.0 \times 10^{-6} )</td>
<td>( 1.8 \times 10^{-5} )</td>
<td>VandenBoer et al. (2013); Wong et al. (2013); Trick (2004)</td>
</tr>
<tr>
<td>HONO + OH</td>
<td>L2 ( \gamma_{\text{HONO-OH}} ) (cm³ molec⁻¹ s⁻¹)</td>
<td>( 6.75 \times 10^{-12} )</td>
<td>( 4.5 \times 10^{-10} )</td>
<td>( 3.0 \times 10^{-12} )</td>
<td>NASA (2011)</td>
</tr>
<tr>
<td>HONO photolysis</td>
<td>L3 ( \gamma_{\text{HONO}} ) (s⁻¹)</td>
<td>( 1.8 \times 10^{-7} )–( 3.9 \times 10^{-5} )</td>
<td></td>
<td></td>
<td>This investigation</td>
</tr>
</tbody>
</table>

* Maximum–minimum range of the experimentally determined time-series values of \( \gamma_{\text{HONO}} \) input to the model (not varied).
Source S1 is the formation of HONO from the reduction of HNO₃ to HONO mediated by VOCs emitted from motor vehicles (Rutter et al., 2014). The source strength (Fₛ₁, ppt/s) was parameterized using HONO source strength and reactant mixing ratios presented in Table 1 of Rutter et al. (2014) and is shown in Eq. (4):

$$F_{S1} = \frac{f_{HONO\text{,VOC}}}{f_{HNO_3\text{,EML}}} \left[ \frac{[\text{Propylene}]_{\text{baseline}}}{[\text{Propylene}]_{\text{Max.DFW}}} \right] \left[ \frac{[\text{HNO_3}]_{\text{EML}}}{[\text{HNO_3}]_{\text{Rutter}}} \right]$$

where $f_{HONO\text{,VOC}}$ is the observed HONO formation rate (ppt/s) in Rutter et al. (2014), and normalizing ratios are further described in the SI. Estimates of ‘likely’ $f_{HONO\text{,VOC}}$ were taken for experiments conducted at 50% RH while ‘lower-limit’ and ‘upper-limit’ estimates were taken as the minimum and average across experiments shown in Table 1 of Rutter et al. (2014). Normalizing assumptions shown in Eq. (4) resulted in, on average, -95% reduction of $f_{HONO\text{,VOC}}$ when calculating $F_{S1}$. The form of the parameterization in Eq. (4) is speculative; propylene is chosen as a proxy for reactive VOCs while benzene is chosen to account for dilution that may occur as air masses move from DFW to EML (see Fig. S7 in the SI for a diurnal profile of propylene/benzene). Identification of specific reactive species participating in the HONO formation process identified in Rutter et al. (2014) would enable improvements in development and assessment of parameterizations of VOC-mediated conversion of HNO₃ to HONO.

Source S2 is HONO emissions from soil bacteria as described by Oswald et al. (2013). Emission from the soil ($F_{S2}$, ppt/s) was assumed to mix instantaneously through the first model layer as shown in Eq. (5):

$$F_{S2} = \frac{f_{\text{soil}}}{h} I_{S2}$$

where $f_{\text{soil}}$ is the “optimum” HONO flux from a soil type (molec cm⁻² s⁻¹), $h$ is the height of the model layer, and $I_{S2}$ represents the conversion factor to ppt/s prior to inclusion in Eq. (1) (see the SI Eqs. S21–S24 for an example calculation). The ‘lower-limit’ value of $f_{\text{soil}}$ was taken as the value of HONO flux for pasture, and the ‘upper-limit’ value was taken as that for grassland. No ‘likely’ value of $f_{\text{soil}}$ was selected, as pasture and grassland were the only two relevant soil types for the DFW region. Despite specifying a ‘lower-limit’ value, this investigation may be effectively considering the high end of contribution of soil bacteria to HONO because “optimum” values of flux are used for both soil types.

Source S3 is the re-emission of HONO from a surface nitrite reservoir by displacement from HNO₂ and HCl, as in VandenBoer et al. (2014, 2015) and shown in Eq. (6):

$$F_{S3} = \frac{[\text{HNO}_2] + [\text{HCl}]}{h} v_d \eta$$

where $F_{S3}$ is the source strength of S3 (ppt/s), $v_d$ is the deposition velocity of HNO₂ and HCl, taken as 1 cm s⁻¹, and $\eta$ is the displacement efficiency, ranging from 1% to 20% for ‘lower-limit’, ‘likely’, and ‘upper-limit’ values, respectively (VandenBoer et al., 2014). This parameterization was constrained by the calculation of a ‘reservoir’ of nitrite from deposited HONO, approximated from a material balance on the ground where the source of nitrite is mechanism L1 and loss is due to displacement from mechanism S3. Mechanism S3 was set to 0 when the reservoir was equal to 0. As there may be additional sources of surface nitrite other than gas-phase HONO and surface nitrite accumulation over greater than diurnal time-scales, Eq. (6) likely represents a conservative estimate of the source strength of S3. Further description of the constraints on source S3 is given in the SI, and dynamics are depicted in Fig. S8, also in the SI.

2.4. Model calculation and assessment

Nitric acid mixing ratios were first modeled with the baseline scenario using the B and L parameterizations summarized in Table 1. The ‘likely’ parameterization incorporates HONO source and sink estimations most representative of each mechanism, while ‘upper-limit’ and ‘lower-limit’ are values that result in maximum or minimum HONO production, respectively, e.g. in the ‘upper-limit’, parameterizations of sources result in greater formation while those of sinks result in lower loss rates. Predictions of HONO mixing ratios were assessed through the residual sum of squared errors (SSE) and the coefficient of determination ($r^2$), both determined from differences between modeled and measured HONO mixing ratios.

Model scenarios were constructed to assess the three new mechanisms (mechanism ID = S1, S2, and S3 shown in Table 1) and gradient conditions (GrN or GrD); scenarios are named according to the gradient used and sources added, e.g., GrN S2/S3 refers to a model scenario with the stronger nighttime gradient as described previously and with sources S2 and S3 added to baseline sources B1–B8 and sinks L1–L3. Sources S1–S3 were added to the baseline model in a full-factorial deterministic screening analysis (using ‘likely’ estimates of parameterizations) to identify scenarios for further analysis. Monte Carlo simulation (Crystal Ball v. 11.1.2.3, Oracle) was used to evaluate the probability of model scenarios to account for observed HONO mixing ratios. Input distributions of source and sink parameterizations were assumed to be triangular probability distributions, bounded by ‘lower-limit’ and ‘upper-limit’ values with the ‘likely’ value as the most frequently occurring. Model sensitivity to the number of trial simulations was performed to ensure a trial-independent solution was achieved; all MCS were conducted with 5000 iterations. A bounded evolutionary solver was applied to the baseline model scenario and to the model scenario with the highest $r^2$ and lowest residual SSE in the deterministic screening analysis. The evolutionary solver used a genetic algorithm to estimate source and sink parameterizations with a minimum SSE across the range of ‘lower-limit’ to ‘upper-limit’ values for each source or sink mechanism.

3. Results and discussion

3.1. Ambient air monitoring in the outflow of DFW

Experimental observations of mixing ratios of ambient gases and particles input to the model are shown in Fig. 1; diurnal profiles of selected constituents across the full monitoring campaign are shown in Fig. S9 of the SI. Values of HONO/NO₂ are variable and elevated during the daytime, possibly indicative of a secondary daytime source of HONO. Mixing ratios of HNO₂ are suppressed in the morning and evenings and elevated during daytime hours, likely a result of strong daytime HNO₂ production from the reaction of NO₂ and OH (Anjea et al., 1994). The highest observed mixing ratios of HNO₂ across the full monitoring campaign are included in the model period shown in Fig. 1, exceeding 5000 ppt in the early evening of June 22, 2011. Mixing ratios of HCl exhibit similar trends to those observed for HNO₂. Mixing ratios of HONO show accumulation over the nighttime and suppression during the daytime, a result of the strong loss due to photolysis and convective dilution during the daytime hours. Aerosols and aerosol-phase constituents appear elevated during the nighttime hours of 6/23 and 6/24 compared to daytime concentrations, but are suppressed during
the nighttime of 6/25. Across the model period, the SA of particulate matter averages 125 μm² cm⁻³, consistent with typical values across the month-long monitoring campaign (Fig. S1), and ranges 22 μm² cm⁻³—392 μm² cm⁻³.

3.2. Baseline model

Mixing ratios of HONO are first calculated with the model under the baseline scenario for 'likely' estimates of parameterizations. Predicted and measured mixing ratios of HONO for the baseline scenario with three HONO gradient conditions described in Section 2.2 are shown in Fig. 2. The "no gradient" condition results in substantial over-estimation of nighttime HONO mixing ratios, logical given the role of the ground surface in HONO formation processes included in the baseline scenario and the first layer height of 36 m. Conversely, the GrN and GrD conditions both result in underestimation of nighttime HONO, with relatively small differences between the two conditions. A strong daytime sink, due to photolysis, results in suppression of modeled daytime mixing ratios below observation for all three gradient conditions, implying the need for daytime sources beyond those considered in the baseline scenario. The underestimation may also result from the limited vertical resolution in the two-layer box model used here and the measurement height in the lower portion of the first layer (10 m); it is likely that a continuous HONO gradient is present in the 36 m of the model first layer resulting in a lower modeled mixing ratio across the first model layer than the 10 m observation.

While relatively few studies report measurements of vertical gradients of HONO, available profiles generally show higher HONO mixing ratios in surface layers than aloft, indicative of ground surface HONO formation. Michoud et al. (2014) summarize several studies reporting vertical gradients, four of which show the
presence of a vertical gradient (Veitel, 2002; Zhang et al., 2009; Villena et al., 2011; Wong et al., 2012) and one that does not (Häseler et al., 2009). VandenBoer et al. (2013) report high-resolution vertical profiles measured from a tower in Boulder, CO, and show the presence of both daytime and nighttime HONO gradients. Veitel (2002) report that over 13 months of measurements, HONO mixing ratios were observed to decrease with height under nearly all atmospheric conditions. For the present investigation, we interpret the over-prediction of HONO mixing ratios in the nighttime for the “no gradient” condition, when convective mixing is most likely to be diminished, to indicate a HONO vertical gradient. Thus, conditions GrN or GrD better represent the vertical structure of HONO mixing ratios in the outflow of DFW. While this appears to be in agreement with the preponderance of available HONO vertical gradient measurements, a site-specific HONO gradient would clearly improve the present study. Nevertheless, parameterizations here allow an estimation of the source and sink processes in the outflow of DFW and exploration of two estimates of gradients to assess model sensitivity to the HONO vertical profile. The impact of the vertical gradient and of parameterizations of established and recently identified HONO sources and sinks are further explored in Sections 3.3–3.5.

3.3. Deterministic screening analysis

A deterministic screening analysis was employed to evaluate model outcomes when sources S1–S3, acting independently or in any combination, are incorporated into the model. This full-factorial analysis, consisting of 24 possible scenarios, is conducted for only the ‘likely’ parameterizations of the mechanisms, as shown in Table S1 of the SI. Full output of model runs across all gradient conditions and scenarios of parameterizations are provided in Figs. S10–S12.

Generally, ‘likely’ estimates of parameterizations showed improved model fit compared to ‘upper-limit’ estimates, implying additional sources of HONO, rather than increased production from baseline sources result in improved model outcomes. Subsequent discussion in this section reflects ‘likely’ parameterizations. Scenarios identified for further investigation are those with a combination of low SSE and high $r^2$. The baseline model generally is characterized by the highest model SSE, and the addition of source mechanisms S1–S3 generally lowers SSE and increases $r^2$. In cases, however, the SSE is lowered while the $r^2$ decreases (for example, from GrN Baseline to GrN S1). This is a result of improvement in model prediction for only a subset of times in the modeling period. The screening analysis identified scenario S2/S3 and scenario S1/ S2/S3 as having the lowest SSE and highest $r^2$ (SSE range: $4.3 \times 10^6–6.7 \times 10^6$; $r^2$ range: 0.42–0.58). These scenarios, along with baseline scenarios for comparison, are further explored with MCS and an evolutionary solver.

3.4. Monte Carlo simulation

Six model scenarios that vary the new sources and vertical gradient conditions were evaluated with MCS to incorporate uncertainty and variability in each mechanism into the model; model estimates of HONO are determined as probabilistic distributions at each model time step. Summarized output of MCS are shown in Figs. S10 as hourly-averaged diurnal profiles of measured and modeled distributions of HONO mixing ratios across the model period. The MCS reinforces the conclusions that ‘baseline’ source mechanisms cannot explain observed HONO mixing ratios; in the GrN Baseline condition, 90th percentile values of model output underestimate observed HONO mixing ratios in 23 of 24 reported hours, and 75th percentile values underestimate observed HONO mixing ratios all 24 reported hours.

The addition of source mechanisms S2 and S3 to the model (Fig. 3) results in improved agreement between the model and observations for nighttime mixing ratios of HONO for both GrN and GrD conditions. GrN S2/S3 shows 9 of the 10 h in the 21:00–07:00 nighttime period are between the 10th and 90th percentile values determined in the model. GrD S2/S3 shows improvement over the GrD Baseline condition; however, metrics of goodness of fit are lower than GrN S2/S3, and there is less improvement over baseline. This appears to be a result of sustained accumulation over the nighttime period, due to the smaller HONO nighttime vertical gradient in the GrD condition. Under both GrN and GrD conditions for scenario S2/S3, daytime mixing ratios of HONO remain substantially underpredicted as in the baseline condition.

The addition of all three sources (S1, S2, and S3) does not appear to resolve underprediction of the daytime HONO mixing ratio. In the GrN condition, the addition of source S1 results in a small increase in over-estimation of nighttime HONO mixing ratios, and metrics of model fit worsen. In the GrD condition, there is a limited impact from the combined effect of sources S1, S2 and S3, with a modest reduction in both SSE and correlation coefficient when comparing GrD S1/S2 to GrD S1/S2/S3. Fig. 3 shows GrN S2/S3 results in improved model fit compared to other scenarios, although daytime HONO remains substantially underestimated.

An estimation of average total and relative source and sink strength across both nighttime (21:00–07:00) and daytime (07:00–21:00) is shown in Fig. 4 for GrN S2/S3. Estimates of sources and sinks are reported for ‘likely’ values of parameterizations for the indicated time period. Considerable temporal differences in the contributions of various source and sinks to the HONO budget exist. At night, HONO from NO2 conversion at the ground (B7) is the major source, contributing 53% of the HONO budget. Biotic release from the ground (S2) and re-emission from the nitrite reservoir (S3) are the next two largest contributors at 25% and 19%, respectively. Nighttime HONO is slightly over-estimated; an ‘unknown’ nighttime sink of 0.0016 ppt/s, or 3% of the total, is required to bring the model into agreement with observations. Major nighttime sinks are vertical transport and deposition of HONO at the ground surface, contributing 73% and 21%, respectively. These nighttime sources and sinks are in general agreement with relative estimates of mechanisms reported by Czader et al. (2012), who report 71% of HONO production due to heterogeneous surface chemistry and losses due to transport and deposition of 77% and 23%, respectively, during the nighttime and pre-sunrise morning.

During the daytime, a missing HONO source dominates; however there are meaningful contributions to the daytime HONO budget from S3, S2, B8, B7 and B5. A missing daytime source of 0.15 ppt/s, or 67% of the total HONO source budget shown in Fig. 4, is needed to bring modeled and measured results into full agreement. This “missing” source is in the range of magnitudes identified in other investigations, ranging from 0.03 to 0.3 ppt/s (Su et al., 2008; Elshorbany et al., 2009; Sørgel et al., 2011; VandenBoer et al., 2013; Lee et al., 2015). Unless there is a positive artifact that depends on sunlight, a strong daytime source is needed to balance the substantial sink of HONO due to photolysis (89% of the total sink). In section 3.5, we explore the potential for ‘best fit’ estimates of parameterizations in GrN S2/S3 to close some portion of the HONO budget through optimization of parameterizations across the range of values presented in Table 1.

3.5. Evolutionary solver and sensitivity analysis

An evolutionary solver was employed to estimate the optimal combination of input values within ‘lower-limit’ to ‘upper-limit’ ranges of parameterizations and the resulting impact on the
estimate of the “missing” HONO source or sink. The evolutionary solver was applied to the GrN baseline scenario and GrN S2/S3. Model outcomes with optimal estimates for GrN baseline and GrN S2/S3 are shown in Fig. 5 and parameterizations are reported in Table 2.

Across optimization of both GrN Baseline and GrN S2/S3, the largest changes to the parameterizations relate to heterogeneous conversion of NO₂ on aerosol (B1 and B2) and on the ground (B7, B8), and HONO uptake to the ground (L1). Aerosol processes increase substantially as a result of a speculative upper-limit as described in the SI; B1 was allowed to vary over 1.5 orders of magnitude and B2 over 2.5 orders of magnitude based on prior modeling studies, rather than experimental estimates. However, contributions from B1 and B2 remain limited (<1% as can be determined from absence of B1 and B2 in Fig. 4), in part a result of the two layer box-model used here that emphasizes ground-level phenomena. In both GrN Baseline and GrN S2/S3, the optimization resulted in B8 at the upper-limit of the parameterization.
Source B7 increased by ~2× in GrN Baseline, but more moderately in GrN S2/S3, a result of the contribution of sources S2 and S3 in GrN S2/S3. In GrN S2/S3, deposition loss (L1) increased, a result of the need to balance increases in parameterizations of sources that act over both daytime and nighttime periods (e.g., S3) and contribute to reductions in the daytime “unknown” source but also nighttime accumulation.

Fig. 5 shows greater improvements in metrics of model goodness of fit for the optimal solution of GrN S2/S3 compared to the optimal solutions of the GrN Baseline. This indicates that baseline mechanisms are not able to similarly explain HONO observations under any combination of input parameters compared to the scenario with S2/S3 present. This appears to largely result from stronger parameterizations of S2/S3 resulting in improved estimates of daytime HONO mixing ratio, although levels are still lower than observed. Best-fit parameterizations of GrN S2/S3 result in a missing daytime source of 0.10 ppt/s, reduced from 0.15 ppt/s (Fig. 4), implying that a substantial missing HONO source remains even across a statistically optimized range of parameterizations.

The “best-fit” estimates of GrN S2/S3 reflect an improved statistical outcome for the model when parameterizations are allowed to vary across a range of values. Parameterizations in Table 2 with larger percentage changes imply a combination of model sensitivity to the parameter as well as uncertainty in the value of the parameterization. We conducted a sensitivity analysis to identify the most important parameterizations impacting the estimates of goodness-of-fit, the model $r^2$ and SSE. The sensitivity analysis for GrN S2/S3 is summarized in Table S2 of the SI, reported as the Spearman’s rank correlation coefficient ($\rho$) between each mechanism’s input parameter and the model output $r^2$ or SSE. Uptake of NO$_2$ at the ground (B7) is the parameter with the largest impact on both the model SSE and $r^2$, by a comparatively large margin. Given that there is a wide range of estimates of the uptake coefficient parameterizing B7 in the literature, this source represents a large source of uncertainty in the model. Sources S3, B8, and S2 are the next three strongest correlations with model SSE; interestingly, all four sources with highest sensitivity (B7, B8, S2, and S3) are ground-level phenomena. Source B7 was strongest correlated with

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**Fig. 4.** Relative contribution to HONO source or sink strength in GrN S2/S3 with ‘likely’ estimates of parameterizations. Contributions are averaged for the time period indicated above each pie chart across the modeling period (6/22/2011 01:00–6/25/2011 14:00 local time). Unknown source or sink is determined by stepwise addition of HONO source or sink such that modeled HONO equals measured HONO.

**Fig. 5.** Model performance with best-fit parameters for the nighttime gradient (GrN) scenario with sources S2 and S3, compared to the nighttime gradient scenario with only baseline sources included.
night-time (21:00–07:00) HONO mixing ratios while source S3 was strongest correlated with daytime HONO. This underscores the importance of characterizing the role of the ground surface mechanisms, including biotic release and ground-level chemical transformations. The presence of a substantial missing daytime source is further explored via estimation of correlation coefficients between measured constituents and products of constituents with the missing HONO source, similar to the analysis presented by Lee et al. (2015). This analysis employed time-series measurements for constituents and the estimate of missing HONO at each time step required for model agreement with observation. Outcomes are shown in Table S3 for ‘likely’ and ‘best-fit’ estimates of GrN S2/S3. Relatively strong correlation coefficients ($r^2 > 0.5$) were observed for $J_{NO2}$ and $J_{NO2} \times$ temperature with the missing HONO source, the latter in close agreement to the results of Lee et al. (2015). However, the correlation of $J_{NO2} \times NO2$ with the missing HONO source is weak ($r^2 = 0.09–0.17$), as is the correlation of $J_{NO2} \times$ SEMS $SA \times NO2$ ($r^2 = 0.08–0.16$) and with NO2 alone ($r^2 = 0.21–0.25$). The stronger correlation with $J_{NO2}$ and $J_{NO2} \times$ temperature may imply photo-sensitized conversion on organics, including humic acids, which are mainly ground surface sources (Stemmler et al., 2006, 2007), are underestimated. The weak correlation of the missing HONO source with NO2 and products containing NO2 mixing ratios appears aligned with a recent analysis of weekday–weekend HONO and NO2 relationships that shows HONO production rates do not increase with increases in NO2, implying daytime HONO production may not be rate-limited by NO2 (Pusede et al., 2015). Weakening correlations for products of gas- and particle-phase constituents and $J_{NO2}$ also may result from the two-layer model that lends greater emphasis to interactions at the ground level, consistent with the results of the sensitivity analysis in Table S2 and discussed previously.

### 3.6. Model limitations

The model described in this work is subject to a number of important limitations. Source S1 assumes the source strength determined in the laboratory is possible in the ambient environment, with several normalizing assumptions. However, as we did not observe meaningful formation of HONO from source S1, the impact of the speculative parameterization is therefore limited in this investigation. Future field efforts should further investigate the potential for VOC-mediated reduction of HNO3 to HONO in near-source environments. Source S2 was parameterized using a single value for a model simulation; there are likely to be diurnal variations in biological activity and soil water content that would impact the parameterization of source S2. Source S3 considered only gas-phase HONO as an input to the surface nitrite reservoir and that the reservoir was empty at the beginning of the model period. This may result in a conservative estimate of the contribution of source S3.

Input distributions in MCS were assumed to be triangular. This assumption may over-weight estimates of parameterizations at the ‘upper-limit’ and ‘lower-limit’ extents of the distribution as compared to a normal distribution. A triangular distribution was chosen, in part, to ensure parameterizations did not exceed upper or lower-limit estimates in MCS. The two-layer box model uses instantaneous and in-situ mixing ratios to constrain the model, with the assumption of instantaneous mixing up to the first layer height. Transport between layers was estimated using an approximation of HONO vertical gradients at similar heights taken from literature. We assume transport time for NO2 sources that exceeds the atmospheric age of HONO (Lee et al., 2013). During the daytime periods (07:00–21:00), the atmospheric age of HONO across the modeling period in this work averaged 19.4 min and ranged from 8.9 to 128 min. We assume NO2 sources input to the model originate from the metropolitan DFW area (~70 km away), while the wind speed averaged 19 km/h, resulting in a transport time of 220 min.

### 4. Conclusions

Model predictions of HONO that account for ranges in parameterizations of HONO source and sink mechanisms enable a statistical assessment of the likelihood of the model to match observation. Observations of HONO appear most accurately simulated for GrN S2/S3. Model output for GrN S2/S3 accounted for, on average, 33% of the daytime HONO budget and 103% of the nighttime HONO budget. Major nighttime sources included (in order) NO2 conversion at the ground (B7), biotic release from soil (S2), and re-emission from the nitrite reservoir (S3). Major daytime sources include S3, S2, photo-enhanced NO2 conversion at the ground (B8), B7, and the reaction of OH with NO (B5). Model fit improved after application of an

### Table 2

<table>
<thead>
<tr>
<th>ID</th>
<th>Parameter</th>
<th>Best-fit estimate (% difference from ‘likely’)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>GrN S2, S3</td>
</tr>
<tr>
<td>B1</td>
<td>$J_{NO2}$ (-)</td>
<td>$3.9 \times 10^{-6}$ (294%)</td>
</tr>
<tr>
<td>B2</td>
<td>$J_{NO2, hv}$ (-)</td>
<td>$8.5 \times 10^{-6}$ (5000%)</td>
</tr>
<tr>
<td>B3</td>
<td>$J_{NO2, hv, BET}$ (-)</td>
<td>$5.3 \times 10^{-7}$ (6%)</td>
</tr>
<tr>
<td>B4</td>
<td>$f_{max}$ (5V, HONO/NO2)</td>
<td>$1.1 \times 10^7$ (6.5%)</td>
</tr>
<tr>
<td>B5</td>
<td>$k_{NO2}$ (cm$^3$ molecule$^{-2}$ s$^{-1}$)</td>
<td>$1.0 \times 10^{-11}$ (4.4%)</td>
</tr>
<tr>
<td>B6</td>
<td>$J_{HONO, hv}$ (cm$^{-1}$ s$^{-1}$)</td>
<td>$7.6 \times 10^{-6}$ (9%)</td>
</tr>
<tr>
<td>B7</td>
<td>$f_{NO2}$ (mole cm$^{-2}$ s$^{-1}$)</td>
<td>$1.2 \times 10^{-5}$ (3%)</td>
</tr>
<tr>
<td>B8</td>
<td>$f_{NO2,vd}$ (ppt/s)</td>
<td>$1.8$ (4.6%)</td>
</tr>
<tr>
<td>S1</td>
<td>$J_{NO2, NO}$ (ppt/s)</td>
<td>$6.1 \times 10^{-6}$ (22%)</td>
</tr>
<tr>
<td>S2</td>
<td>$f_{max}$ (mole cm$^{-2}$ s$^{-1}$)</td>
<td>$6 \times 10^{-6}$ (200%)</td>
</tr>
<tr>
<td></td>
<td>$v_{g} - \eta$ (cm s$^{-1}$)</td>
<td>n/a</td>
</tr>
<tr>
<td>S3</td>
<td>$J_{NO2, NO}$ (cm$^{-1}$ s$^{-1}$)</td>
<td>$2.8 \times 10^{6}$ (66%)</td>
</tr>
<tr>
<td>L1</td>
<td>$J_{HONO, hv}$ (cm$^{-1}$ s$^{-1}$)</td>
<td>0.18 (105%)</td>
</tr>
<tr>
<td>L2</td>
<td>$k_{HONO, hv}$ (cm$^{-1}$ s$^{-1}$)</td>
<td>$5.7 \times 10^{-5}$ (185%)</td>
</tr>
<tr>
<td>L3</td>
<td>$J_{HONO}$ (s$^{-1}$)</td>
<td>$5.7 \times 10^{-4}$ (283%)</td>
</tr>
<tr>
<td></td>
<td>Missing source or sink: daytime, nighttime (ppt/s)</td>
<td>unchanged</td>
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evolutionary solver, resulting in a reduction of the estimate of the unknown daytime source for GrN S2/S3. However, the presence of a substantial unknown daytime source (on average 0.10 ppt/s) even with a statistically optimal fit for GrN S2/S3 implies sources of HONO other than those evaluated here must be included to reproduce accurately daytime HONO mixing ratios. Analyses of model sensitivity and correlations between the missing HONO source and constituents imply the presence of additional, or underestimation of considered, ground-level HONO sources in this investigation.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.atmosenv.2015.12.044.

References


