Response to the Anonymous Referee # 1

We would like to thank the Referee for time and effort put into reviewing this manuscript. Please see below our responses to your comments.

One of the major issues here is the lack of continuous or sufficient HONO measurements. I can hardly count 6 diurnal cycles of HONO measurements in Fig. 6, which is properly not adequate to discuss model-measurements comparison. From Fig. 6, HONO simulations seem to be improved only during the early morning and most apparently during the Sep. 12th. Figs 4 and 5 are shown only for Sep 12th, what about the other days?

--- HONO is usually not measured routinely, which is a shortcoming of this as well as other studies (e.g., Wang et al. (2011)). Limited availability of measured data is already mentioned in the manuscript (see page 21324 lines 16-19). We hope that the new emission ratio would be tested in other areas along with different HONO measurements that would provide additional validation of HONO emissions. For our study we will add more detailed analysis of the dataset that is available to us as well as more analysis of the modeling results itself, especially on the potential impact of these higher emissions on modeled mixing ratios (see).

--- Since only HONO emissions from mobile sources were increased we expect to see the largest differences in mixing ratios during early morning times when the traffic emissions are the highest, the mixing layer height low allowing for accumulation of HONO, and photochemistry not very active. We will incorporate this statement into the manuscript.

Also from Fig. 6 (since it is the only figure that show several days of HONO diurnal cycles), it seems that HONO was much better simulated with the (N) scenario on Sept. 11th, 19th and 26th, which are significantly overestimated by the new (NH) scenarios. HONO simulations seem to be improved only on the Sep. 12th, 18th.

--- Variations of simulated HONO mixing ratios from day to day are influenced not only by emissions but also by other parameters, for example, the model capability to predict grow of the mixing layer as well as clouds that influence photolysis rates. To more clearly present differences between the simulations cased we prepared the average diurnal profile of measured HONO and compared it with simulated N and NH cases (see Fig. 7 below). It can be seen that NH case improves HONO morning peaks. As mentioned above, the increase in the morning can be explained by high traffic emission during morning times, low mixing layer height and accumulation of HONO since photochemistry is not very active.

--- HONO simulations with the new ratio are improved on Sep. 12, 18 as well as on Sep. 23, 25, and 30.
The authors should also plot the Measured vs Simulated HONO for both scenarios (N and NH) and for each complete diurnal cycle and for the mean simulated period (though statistical mean is shown in table 3), so that we can get a clearer picture if the new ER (NH scenario) would consistently improve HONO or only under certain conditions. Why it is only improved on the 12th?

--- Diurnal profiles of the measured and simulated HONO from both scenarios (N and NH) are already presented in Figure 6. As mentioned above, based on the comparison of measured NO\textsubscript{x} from very representative dataset at many stations around Houston taken during the whole month of September, we believe that N case better reflects observed NO\textsubscript{x} and since HONO is derived directly from NO\textsubscript{x}, reducing emissions of NO\textsubscript{x} resulted in HONO reduction.

--- The additional figure with an average HONO profiles (see below) will be added to the manuscript as Fig. 7 (current Fig. 7 will become Fig. 8) along with the following description:

Figure 7 shows the average diurnal profiles of measured and simulated HONO mixing ratios. Since only HONO emissions from mobile sources were increased the largest differences in mixing ratios occur during early morning times when the traffic emissions are the highest, the mixing layer height low allowing for accumulation of HONO, and photochemistry not very active. The model underprediction during daytime can be explained by the fact the default model version that we used in this study does not account for the photochemical HONO sources. Also, too low modeled average profile during daytime is caused by underpredictions of HONO on Sep. 23-25 which can be attributed to stronger modeled winds in comparison to weak observed winds causing HONO to be removed from the observational site. It is worth to note that all available measured data for HONO for the September 2013 are from weekdays and the higher HONO/NO\textsubscript{x} ratio measured in Houston was calculated based on measurements taken during weekdays.”

![Graph showing diurnal variation of HONO](image)
The statistical mean in table 3 is misleading because the overestimated and underestimated HONO cancel each other resulting in slightly improved mean simulated HONO. So here also, the authors should show the results for each diurnal cycle (not the mean).

--- We agree that the mean values might be misleading but in addition to the mean value we also calculated and presented in table 3 the absolute mean error. The underestimated and overestimated values do not cancel each other in calculation of the absolute mean error because the absolute values are taken for the calculation (please refer to equation (1)).

On the days 23-25th, HONO measurements are still significantly underestimated, especially during the afternoon time. This underestimation should also be discussed in more details. In fact, most of the HONO unknown sources are reported during afternoon hours (e.g., Kleffmann et al., 2005; Elshorbany et al., 2012). During the early morning, the so called [HONO]pss (zero net OH source), which account for the known gas phase HONO formation from OH+NO and loss through photolysis and reaction with OH, accounts for most of the early morning peak.

--- We agree that part of underprediction of HONO on those days may be due to photochemical HONO formation that is not accounted in the model. Also, on September 21-25 a cold front was passing through the Houston area, with high pressure system. On Sep. 23 – 25 the model shows stronger easterly winds than the observation that contribute to faster transport and removal of HONO from the observation site.

Why these high emission ratios, Could the authors try to shed some light on the type of fleet in Huston Metropolitan Area, compared to other cities in the US or to the fleet in Europe, does the fleet type and quality changed over time (Benzene, diesel, natural gas, hyprid cars, ::: .etc), why are ER are different that reported before?

--- We will add the following discussion about that:

“The HONO/NOx ratio reported by Kurtenbach et al. (2001) is based on measurements performed between 6 am and 2 pm, for both weekdays and weekends where 22 200 ± 400 vehicles were passing on weekdays and 13 300 ± 1 400 cars passing on weekends. The vehicle fleet was composed of 6.0% heavy-duty trucks, 6.0% commercial vans, 12% diesel and 75% gasoline powered passenger cars, and 1.0% motorcycles. The measurements made by Rappenglueck et al. (2013) reflect high traffic, early morning conditions (4-8 am) on weekdays. The measurements were performed at highway junction in Houston with very high traffic load (about 400 000 vehicles passing daily), which is much larger than that in the tunnel study. The vehicle fleet was represented by 93-95% of gasoline fueled vehicles and 5-7% by diesels during the morning hours. Another difference between these two studies is in vehicle speed, with a typical speed of 50-90 km/h in the tunnel studies and much lower speed during the morning peak traffic hours in Houston.”
“Since the newly reported ratio reflects high traffic conditions during the morning rush hours on weekdays our model sensitivity study provides estimate of the upper bound of the impact of HONO emissions on pollutant levels in urban areas.”

At the end, more scientific discussion of the results is still required. For example, why OH is only enhanced by ~5% though HONO is enhanced by 35% (Table 3) on Sep. 12th. What is the contribution of HONOprod to the total simulated HONO? Here also Fig. 7 should include all other simulated cycles, i.e. not only one single event.

Based on the 1 month of simulated surface concentrations the average increase in the morning OH (between 6 – 8 a.m. LT) is 14% at the location of the Moody Tower and 3% when averaged over the urban area. The ozone increase is below 1% for both the Moody Tower and the urban area. The average increase in OH during daytime (6 a.m. – 8 p.m. LT) is 7% for the Moody Tower and 1% for the urban area. The increase in ozone is again below 1%. To obtain more insights on the fate of HONO we performed additional model simulations and analysis for the Moody Tower site for Sep. 10-13, 2013. At the surface at the location of the Moody Tower the average contribution of vertical transport to the loss of HONO is 77%, horizontal transport contributes 8%, chemical removal 11% and dry deposition 5%. At the second model layer, which corresponds to the altitude of measurements, transport (horizontal and vertical) continue to be a dominant loss process contributing on average 77% to the total HONO loss while chemical loss contributes only 23% to the total loss. The chemical loss of HONO is dominant only during couple of morning hours. Figure 9 shows hourly values of HONO mixing ratios for Sep. 10-13, 2013 along with process contributing to changes in the mixing ratios at the grid cell corresponding to measurements taken at the Moody Tower (simulated data extracted from the second model layer). This explains the fact that even though HONO mixing ratios significantly increased upon additional emissions, HONO was removed mainly by transport with only small portions taking part in chemical reactions converting it to OH and furthermore to O3. Also, the main impact of chemistry is during early morning hours following the peak in HONO.
Figure 9. HONO mixing ratios (black line) and processes contributing to changes in HONO mixing ratio at the Moody Tower site where the measurements were taken, which corresponds to the second model layer. VTRAN is vertical transport, HTRAN is transport in horizontal direction, and CHEM_HONO correspond to changes due to chemical reactions.

--- We believe that the above presented analysis of the 1 month dataset provide sufficient information on OH increases and do not see a need of adding more graphics in Figure 7.

--- Instead of showing

Technical corrections
Page 21317, line 9: HONO photolysis during the early morning was first reported by Perner and Platt (1979) and Harris et al. (1982). Add these references before Czader et al., (2012) and write (e.g.,) at the statement’s beginning.

--- We will certainly add the above mentioned references and modify the manuscript according to your suggestion.

Page 21325, line 18: 12 September.
Page 21325, line 22: 12 September.
Response to the Anonymous Referee # 2

We would like to thank the Referee for time and effort put into reviewing this manuscript. Please see below our responses to your comments.

Please check the word “sheds”. Should it be “shed”?

--- yes, it should be “shed”, we will modify it.

Section 2 – Methodology
Indeed several studies, mentioned in the article, have suggested that 2005-2008 NEI overestimates NOx emissions in Houston. The authors simulated air quality for 2013 using the revised 2008 NEI. In theory, the revision of the 2008 NEI accounts for the NOx emissions reduction that occurred between 2008 and 2013. Since the base 2008 NEI contains higher NOx estimates, the revised NEI for 2013 that the authors used in the study still likely to over-estimate NOx emissions in Houston. Thus, some discussions are needed to indicate such possibility and relate to the over-predictions of NOx mixing ratios shown in Table 2 and Figure 2-3.

--- We will add the following discussion to the manuscript on page 21323, line9:

“Even though in our study we adjusted NOx emissions to reflect emission reduction between the year 2008 and 2013 some overpredictions may occur since, as pointed by Choi (2014), NOx rates in the base 2008 inventory might be too high.”

Section 2.1 – Adjusting NOx and HONO emissions
The authors used a newly reported HONO speciation factor. Should the new speciation factor be used for all urban areas or be limited only to Houston? Some discussion will be helpful to air quality modelers.

--- We will add the following discussion about that on page 21320 in line 24 after sentence “..tunnel measurements in 2001.”:

“The HONO/NOx ratio reported by Kurtenbach et al. (2001) is based on measurements performed between 6 am and 2 pm, for both weekdays and weekends where 22 200 ± 400 vehicles were passing on weekdays and 13 300 ± 1 400 cars passing on weekends. The vehicle fleet was composed of 6.0% heavy-duty trucks, 6.0% commercial vans, 12% diesel and 75% gasoline powered passenger cars, and 1.0% motorcycles. The measurements made by Rappenglueck et al. (2013) reflect high traffic, early morning conditions (4-8 am) on weekdays. The measurements were performed at highway junction in Houston with very high traffic load (about 400 000 vehicles passing daily), which is much larger than that in the tunnel study. The vehicle fleet was represented by 93-95% of gasoline fueled vehicles and 5-7% by diesels during
the morning hours. Another difference between these two studies is in vehicle speed, with a
typical speed of 50-90 km/h in the tunnel studies and much lower speed during the morning peak
traffic hours in Houston.”

--- Also, the following will be added at the end of paragraph in line 27 on page 21320:

“Since the newly reported ratio reflects high traffic conditions during the morning rush hours on
weekdays our model sensitivity study provides estimate of the upper bound of the impact of
HONO emissions on pollutant levels in urban areas.”

Section 3.2 – HONO Modeling
What is average increase in morning OH for the entire simulation? Similarly, what is its impact
on average morning ozone for the entire simulation period?

--- We will add the following discussion:

“Based on the 1 month of simulated data the average increase in the morning OH (between 6 – 8
a.m. LT) is 14% at the location of the Moody Tower and 3% when averaged over the urban area.
The ozone increase is below 1% for both the Moody Tower and the urban area. The average
increase in OH during daytime (6 a.m. – 8 p.m. LT) is 7% for the Moody Tower and 1% for the
urban area. The increase in ozone is again below 1%”.

Section 4 - Summary
OH predictions have not been compared to any observed data. Thus, it cannot be concluded that
model under-predicts OH.

--- The sentence on page 21326 in lines 22-24 will be re-written as follow:

“This study results could shed light on the underestimated HONO in the morning from
global/regional chemical transport model with the typical emission ratio of 0.8% HONO
emission out of the total NOx emissions. In addition, since HONO is the major radical source in
the morning (e.g., Perner and Platt, 1979; Harris et al., 1982; Czader et al., 2013),
derunderpredictions of HONO would lead to underprediction of OH radical.”

Need to clarify that total NOx emissions are not used for speciating HONO emissions; only
mobile source NOx emissions have been used.

--- This information is provided in the Methodology section on page 21320, lines 19-21:

“NEI provides emission rates for nitrogen oxides, during the processing with SMOKE NOx
emissions for mobile sources are separated into 90% NO, 9.2% NO2, and 0.8% HONO.”
--- We will also modify lines 3-4 in the summary as follow:

“In addition, HONO/NOx emission ratio from mobile sources was increased and its impact on HONO mixing ratios was evaluated.”

Table 1 and 2
Units are not included in the tables.

--- we will correct that and add units (ppbv) next to the mean, max. value, bias, and absolute mean error (AME) headers in tables. R and IOA are unitless.

Table 3
It shows “Sim. H”; it will probably be “Sim. NH”.

--- yes, it should be “Sim NH”, we will correct that.

Figure 4
Need to specify date and local time in the figure caption.

--- we will replace the caption with the following:

“Snapshot of differences in HONO emissions between a case with emission ratio of HONO/NOx =0.016 (NH) and default emissions of HONO/NOx=0.008 (N) at 7 a.m. LT on September 12, 2013.”

Figure 5
Need to specify date and local time in the figure caption. Figure caption states base HONO emissions but parenthesis shows (N).

--- we will modify the caption as follow:

“Differences in HONO mixing ratios between a case with 0.016 HONO/NOx emission ratio (NH) and 0.008 HONO/NOx emissions (N) for the surface (left) and the second model layer (right) at 7 a.m. LT on September 12, 2013.”

Figure 7
Need to specify date and local time in the figure caption. Figure caption states differences between the base and increased HONO emissions case. I think case N is used, not the base case.

--- we will modify the caption as follow:
“OH mixing ratios (left) and differences in OH mixing ratios (right) between the case with 0.008 HONO/NO$_x$ emission ratio (N) and 0.016 NO$_x$/HONO emission ratio (NH) at noon local time on September 12, 2013.”
Impact of updated traffic emissions on HONO mixing ratios simulated for urban site in Houston, Texas

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Abstract

Recent measurements in Houston show that HONO traffic emissions are 1.7% of NO\textsubscript{x} emissions which is about twice the previously estimated value of 0.8% based on tunnel measurements in 2001. The 0.8% value is widely used to estimate mobile emissions of HONO for air quality modeling applications. This study applies the newly estimated HONO/NO\textsubscript{x} ratio in the WRF-SMOKE-CMAQ modeling system and estimates the impact of higher HONO traffic emissions on its mixing ratios. Since applied emission inventory resulted in overestimates of NO\textsubscript{x} mixing ratios and because HONO emissions and chemical formation depends on the magnitude of NO\textsubscript{x}, thus, before proceeding with HONO emission modifications emissions of NO\textsubscript{x} were adjusted to reflect current emission trends. The modeled mixing ratios of NO\textsubscript{x} were evaluated against measured data from a number of sites in the Houston area. Overall, the NO\textsubscript{x} mean value dropped from 11.11 ppbv in the base case to 7.59 ppbv in the NO\textsubscript{x} adjusted case becoming much closer to the observed mean of 7.76 ppbv. The Index of Agreement (IOA) is improved in the reduced NO\textsubscript{x} case (0.71 vs. 0.75) and the Absolute Mean Error (AME) is lowered from 6.76 to 4.94. The modeled mixing ratios of HONO were evaluated against the actual observed values attained at the Moody Tower in Houston. The model could not reproduce the morning HONO peaks when the low HONO/NO\textsubscript{x} ratio of 0.008 was used to estimate HONO emissions. Doubling HONO emissions from mobile sources resulted in higher mixing ratios, the mean value increased from 0.30 ppbv to 0.41 ppbv becoming closer to the observed mean concentrations of 0.69 but still
low; AME was slightly reduced from 0.46 to 0.43. IOA for simulation that used the 2001 emission values is 0.63 while for simulation with higher HONO emission it increased to 0.70. Increased HONO emissions from mobile sources resulted in 14% increase in OH during morning time at the location of the Moody Tower and 3% when averaged over urban area. The increase calculated for daytime was 7% and 1% for the Moody Tower and the urban area, respectively. Increased HONO emissions impacted OH mixing ratio, up to about 6% increase was found during morning and mid-day hours. The impact on ozone is was found to be marginal. This study results sheds light on the underestimated HONO and OH in the morning from global/regional chemical transport models with the typical emission of 0.8% HONO emission out of the total NOx emissions.

1. Introduction

Photolysis of nitrous acid (HONO) is an important source of hydroxyl radical (OH). OH plays a crucial role in the oxidation of volatile organic compounds (VOCs) leading to the formation of ozone and secondary organic particulate matter. Main sources of OH are photolysis of ozone, formaldehyde, alkenes, and nitrous acid (Elshorbany et al., 2009; Mao et al., 2010; Kim et al., 2014). Photolysis of ozone and formaldehyde are the most important sources of OH during mid-day and afternoon hours; however, the highest contribution to radical production during early morning hours comes from photolysis of HONO (e.g. Perner and Platt, 1979; Harris et al., 1982; Czader et al., 2012, 2013).

HONO can be either formed through chemical reactions or emitted to the atmosphere from combustion processes. Among the most known chemical sources of HONO is the gas-phase formation from the reaction between OH and nitric oxide (NO) (Pagsberg et al., 1997) and the heterogeneous formation on surfaces from the hydrolysis of nitrogen dioxide (NO2) (Kleffmann et al., 1998; Finlayson-Pitts et al., 2003). Other chemical sources of HONO are described elsewhere (Kleffmann et al., 2005, 2007; George et al. 2005; Stemmler et al. 2006, 2007; Crowley and Carl, 1997; Li et al., 2008, 2009; Carr et al., 2009; Amedro et al., 2011). Emissions of HONO from traffic were estimated by Kirchstetter et al. (1996) and Kurtenbach et al. (2001)
who performed tunnel studies and reported exhaust emission ratio of HONO to NOx in a range of 0.003-0.008. The value of 0.008 is used in the Community Multiscale Air Quality (CMAQ) model to calculate HONO emissions from mobile sources (Foley et al., 2010) as well as in other models, for example, in a box model employed to study HONO sources in Houston (Wong et al. 2013). The relative contribution of HONO emissions from traffic to other sources when using the HONO to NOx ratio of 0.008 is about 9% based on simulations for eastern U.S. (Sarwar et al. 2008). For high NOx areas in China Li et al. (2011) calculated as high as 26% contribution of HONO emissions to its total sources but they could not reproduce the high morning peak values of HONO associated with traffic emissions. Czader et al. (2012) studied HONO formation for Houston conditions and also applied the 0.008 HONO/NOx ratio to estimate HONO emissions. In addition to default sources of HONO present in CMAQ they implemented photolytic HONO formation; however, on many occasions the peak morning values continued to be underpredicted by the model. Recent measurements performed in Houston in 2009 show that the observed HONO/NOx emission ratio is 0.017 (Rappenglueck et al., 2013), which is about twice as high as previously reported and implemented in CMAQ modeling system. The impact of using higher HONO emissions in air quality modeling applications has not been evaluated. Therefore, in this work HONO emissions from mobile sources will be doubled to reflect the newly reported HONO/NOx emission ratio and the impact of higher HONO traffic emissions on its mixing ratios will be estimated in the WRF-CMAQ modeling system. The impact of increased HONO on the OH and O3 will also be investigated in this study.

Because in air quality applications HONO is derived from the total NOx reported in an emission inventory and chemical formation of HONO is directly related to NO and NO2 mixing ratios; therefore, HONO predictions by air quality models depend on how well the model reflects captures emissions of NOx. Czader et al. (2012) pointed out that the correlation between measured and simulated HONO values increased significantly when data points with wrong NO2 prediction were ignored and only data for which NO2 values were simulated within 70% of the measured value were considered. Therefore, accurate estimation of NOx in air quality models is crucial to properly simulate HONO mixing ratios. Previous studies used remote sensing and in-situ surface observations to analyze accuracy of NOx emissions and indicated that the National
Emission Inventory (NEI) has large uncertainty in emissions in urban areas (Choi et al., 2012; Choi, 2014). Of particular, Choi (2014) issued that both NEI2005 and NEI2008 have significant NOx overestimates in Houston. Thus, in this study, before proceeding with modifications of HONO emissions, NOx emissions will be adjusted using the U.S. Environmental Protection Agency (EPA) annual trend values and the absolute amounts of simulated surface NOx concentrations will be evaluated.

2. Methodology

Meteorological parameters were derived with the Weather Research and Forecasting (WRF) model version 3.5 (Skamarock et al., 2008). NCEP North American Regional Reanalysis (NARR) data provided by the NOAA/OAR/ESRL PSD (available at http://www.esrl.noaa.gov/psd/) were utilized to initialize WRF simulations. The 2008 National Emission Inventory (NEI2008) generated by the Environmental Protection Agency (EPA) was processed with the Sparse Matrix Operator Kernel Emissions (SMOKE) system to obtain gridded, chemically and temporally resolved emission files ready to use in an air quality model. The air quality simulations were performed with the three-dimensional Community Multiscale Air Quality (CMAQ) model (Byun and Schere, 2006) version 5.0.1 with the Carbon Bond 05 chemical mechanism and aerosol 5 module (cb05tucl_ae5_aq).

Simulations were performed for a domain with 4 km grid resolution covering southeast Texas, with 84 grid cells in east-west direction, 66 grid cells in south-north direction, and 27 vertical layers. The boundary conditions were obtained from the University of Houston air quality forecasting system (http://spock.geosc.uh.edu) from a larger domain with 12 km grid resolution, 150 grid cells in east-west direction and 134 grid cells in south-north direction. Initial conditions were also obtained from the air quality forecasting results from the nested south-east Texas domain. Simulations were performed for the month of September 2013 during which the DISCOVER-AQ campaign took place in Houston providing many different meteorological and chemical measurements that can be utilized for model evaluation.
2.1 Adjusting NO\textsubscript{x} and HONO emissions

Previous studies used remote sensing and in-situ surface observations to analyze accuracy of NO\textsubscript{x} emissions and pointed to the fact that the National Emission Inventory (NEI) has large uncertainties in emissions for urban areas (Choi et al., 2012; Choi, 2014). Of particular, Choi (2014) issued that both NEI2005 and NEI2008 might have significant overestimates of NO\textsubscript{x} emissions in Houston even with the consideration of the uncertainties caused from other chemical and physical processes. Adequate estimation of NO\textsubscript{x} emissions is critical for properly predicting HONO mixing ratios.

Since our simulations employed NEI2008 there was a need of adjusting emissions to reflect conditions of 2013. In this study, instead of relying on the remote-sensing-derived data or surface-measured data to adjust an emission inventory (e.g., Kim et al., 2009; Kim et al., 2011; Choi et al., 2012; Choi, 2014) we use the long-term trends of anthropogenic NO\textsubscript{x} emission reported by U.S. EPA. Then the impact of the adjusted NO\textsubscript{x} emissions on surface NO\textsubscript{x} concentrations is evaluated by comparing the simulated and observed NO\textsubscript{x} concentrations. According to EPA, emissions of nitrogen oxides from anthropogenic sources were reduced between 2008 and 2013. Table 1 shows emission values based on the EPA trends (available at: http://www.epa.gov/ttn/chief/trends/index.html#tables) for on-road mobile sources and other anthropogenic sources excluding wildfires. Relatively to values for the year 2008 there was 28% reduction in on-road mobile NO\textsubscript{x} emissions on a nationwide scale and 20% reduction in other anthropogenic NO\textsubscript{x} emissions in year 2013. To follow the emissions trends we created a sensitivity case in which on-road NO\textsubscript{x} emissions were reduced by 30% and anthropogenic point source emissions were reduced by 20%.

NEI provides emission rates for nitrogen oxides, during the processing with SMOKE NO\textsubscript{x} emissions for mobile sources are separated into 90% NO, 9.2% NO\textsubscript{2}, and 0.8% HONO. However, Rappenglueck et al. (2013) reports much higher HONO contribution from mobile sources in Houston; based on all measurements HONO traffic emissions are 1.7% of NO\textsubscript{x} emissions which is about twice the previously estimated value of 0.8% based on tunnel...
measurements in 2001. The HONO/NO\textsubscript{x} ratio reported by Kurtenbach et al. (2001) is based on
measurements performed between 6 am and 2 pm, for both weekdays and weekends where 22
200 ± 400 vehicles were passing on weekdays and 13 300 ± 1 400 cars passing on weekends,
The vehicle fleet was composed of 6.0% heavy-duty trucks, 6.0% commercial vans, 12% diesel
and 75% gasoline powered passenger cars, and 1.0% motorcycles. The ratio calculated by
Rappenglueck et al. (2013) is based on measurements performed during weekdays reflecting
high traffic, early morning conditions (4-8 am). The measurements were performed at highway
junction in Houston with very high traffic load (about 400 000 vehicles passing daily), which is
much larger than that in the tunnel study. The vehicle fleet was represented by 93-95% of
gasoline fueled vehicles and 5-7% by diesels during the morning hours. Another difference
between these two studies is in vehicle speed, with a typical speed of 50-90 km/h in the tunnel
studies and much lower speed during the morning peak traffic hours in Houston. To reflect the
latest values of HONO emissions measured in Houston in air quality modeling
additional sensitivity case was created in which contribution of HONO from mobile sources was
doubled at the cost of NO\textsubscript{2}. The following speciation was used for the sensitivity case: 90% NO,
8.4% NO\textsubscript{2}, and 1.6% HONO. It is worth to note that since the newly reported ratio reflects high
traffic conditions during morning rush hours on weekdays our model sensitivity study provides
estimate of the upper bound of the impact of HONO emissions on pollutant levels in urban areas.

The following three simulations cases are performed and analyzed in this study:

- **B** – base case, with NO\textsubscript{x} emissions rates obtained from NEI2008 and HONO/NO\textsubscript{x} = 0.008;
- **N** – reduced emissions of NO\textsubscript{x} case: mobile sources * 0.7, point sources * 0.8; **HONO/NO\textsubscript{x} = 0.008**;
- **NH** – similar as N but with doubled HONO emissions from mobile sources, this is
  HONO/NO\textsubscript{x}=0.016.

### 2.2 Measurements

Measured values from the Continuous Ambient Monitoring Stations (CAMS) system, operated
by the Texas Commission on Environmental Quality (TCEQ), were utilized for evaluating NO\textsubscript{x}
emission inventory. During the time period of interest 30 stations inside our 4 km modeling domain reported NOx measurements. Figure 1 show location of sites in the Houston – Galveston metropolitan areas, where color of the symbol indicates the measured mean NOx mixing ratios during the month of September 2013. Several sites, such as 78, 84, 618, 619, and 1016 have low mean values; those sites reflect regional and/or suburban conditions. Couple sites, such as 26 and 53, have medium range NOx values reflecting urban air mixture dominated by traffic emissions. Many sites close to highways or in downtown Houston and east of downtown are exposed to heavy traffic as well as a combination of traffic and industrial emissions. They have very high NOx mean values; those are CAMS sites 1, 8, 114, 403, 408, 411 and the Moody Tower (MT) site described below.

The Moody Tower, located east of downtown, was designated as a “super” site during air quality study campaigns in Houston in years 2006 (Lefer and Rappenglück, 2010) and 2009 (Olaguer et al., 2013) during which many chemical and meteorological measurements were taken. During September 2013 measurements at the Moody Tower complimented the DISCOVER-AQ campaign. The measurements were taken at 60 m a.g.l. In addition to NOx and ozone, HONO was also measured on several days during the month of September 2013.

3. Results

3.1 Evaluation of NOx modeling

Table 2 shows summary of statistical parameters for modeling NOx mixing ratios for the base case (B) and the reduced NOx case (N) as compared to measured values at CAMS sites, where R is the Pearson coefficient, AME – absolute mean error calculated as:

\[
AME = \left( \frac{1}{n} \right) \sum_{i=1}^{n} |C_m - C_o|
\]
"n" is the number of data points, “m” corresponds to modeled values and “o” to observed ones;

IOA – index of agreement, calculated according the following equation:

\[ IOA = 1 - \frac{\sum^n (C_m - C_o)^2}{\sum^n (|C_o - \bar{\delta}| + |C_m - \bar{\delta}|)^2} \]

“\(\bar{\delta}\)” corresponds to observed mean value. Compared to a Pearson coefficient the index of agreement is a more comprehensive measure of how well the concentrations are predicted since it takes into account not only scattering of data but also biases (Willmott, 1981).

Statistical parameters were calculated for all available data pairs from CAMS sites inside the modeling domain. The measured mean value from all sites is 7.76 ppbv, the simulated mean value dropped from 11.11 ppbv in the base case to 7.59 ppbv in the reduced NO\textsubscript{x} case becoming closer to the observed mean. Both, R and IOA are improved in the reduced NO\textsubscript{x} case (R=0.58, IOA = 0.71 in the base case, R=0.59, IOA = 0.75 in the reduced NO\textsubscript{x} case) and AME is lowered from 6.76 ppbv to 4.94 ppbv. Overall, the reduced NO\textsubscript{x} simulation case gives better NO\textsubscript{x} prediction in comparison to the base case. When looking at individual stations affected by emissions from different sources the improvement from NO\textsubscript{x} reductions is beneficial for most of sites, but leads to underpredictions at several sites. Many stations with medium range NO\textsubscript{x} mixing ratios, such as CAMS 35 and 53 show improvement from NO\textsubscript{x} reduction. There are also cases when NO\textsubscript{x} continue to be too high even after reduction of emissions. This is the case for CAMS sites 26 and 78 that represent sub-urban conditions with low measured NO\textsubscript{x} mixing ratios (usually below 10 ppb) and low mean values of 5.61 and 3.29, respectively. The model represents them as urban sites with significant traffic signature and therefore with much higher than measured mixing ratios. Even though in our study we adjusted NO\textsubscript{x} emissions to reflect emission reduction between the year 2008 and 2013 some overpredictions may occur since, as pointed by Choi (2014), NO\textsubscript{x} rates in the base 2008 inventory might be too high. Very high NO\textsubscript{x} mixing ratios are recorded in areas with heavy traffic and close to industrial facilities in the
eastern part of Houston; these are such as at CAMS stations 1, 403, 411, and 416. NOx mixing ratios at those stations were heavily overpredicted and consequently those stations benefit the most from NOx reductions as presented in Figure 2. Our results are similar to the previous study by Choi (2014) who issued that NOx mixing ratios at urban regions are overpredicted by air quality models, but NOx at the rural regions are underpredicted.

The Moody Tower site served as a super site for couple of measurements campaigns in Houston and many different chemical and meteorological parameters were measured there, including NO, NO2, and HONO. It is located in close proximity to downtown and major highways and is affected by quite high NOx emissions. Figure 3 shows comparison of measured at the Moody Tower and simulated mixing ratios of NO (top) and NO2 (bottom). Again, two simulation cases are compared: the case with regular emissions as included in NEI2008 (B) and the reduced NOx emissions case (N). It can be seen that for both compounds the peak values were overpredicted by the base case while reduced NOx case resulted in lower mixing ratios making them closer to the observed values. In particular, NO mixing ratios are much better predicted by reduced NOx emissions case. Both, NO2 morning peaks and low range day and nighttime NO2 values, although lowered, continue to be overpredicted most of the time.

3.2 HONO modeling

Since reduction of NOx emissions resulted in better prediction of NOx mixing ratios at the Moody Tower and nearby areas this case was used as a base for testing impact of increased HONO emissions. Figure 4 shows changes in HONO emissions rates between the sensitivity case in which HONO/NOx=0.016 (indicated as NH) and the base case that used HONO/NOx=0.008 (indicated as N). Doubling HONO emissions resulted in up to 0.01 mole/s increase in emission rates from mobile sources along highways. Figure 5 show differences in simulated mixing ratios of HONO for morning conditions at 7 a.m. LT that corresponds to the time of the highest HONO emissions from traffic and the highest HONO mixing ratios. The left panel shows results for the surface layer. It can be seen that changes of in HONO mixing ratio at the surface occur along highways following the pattern of emission changes presented in Figure
4. Differences in HONO mixing ratios at the second modeled layer, which corresponds to measurements taken at the Moody Tower, are shown in the right panel of Figure 5. At this level the air is mixed and the spatial signature of mobile emissions diminishes.

HONO is not routinely measured in Houston; in spite of that, during September 2013 HONO was measured at the Moody Tower to compliment measurements during DISCOVER-AQ campaign. However, the measurements were not continuous and the data are limited to several days. Figure 6 shows timeseries of measured and simulated HONO mixing ratios at the Moody Tower. The mixing ratios obtained from the reduced NOx simulation case (N), for which the HONO/NOx emission ratio of 0.008 was used, are much lower than observed HONO values. The values from the increased HONO case (NH), with the HONO/NOx emission ratio of 0.016, are higher, especially the morning peaks, and closer to the observations. The statistical parameters for HONO modeling at the Moody Tower are presented in Table 3. The mean value increased from 0.30 in the base case to 0.41 ppbv in the increased HONO emissions case but continue to be lower than the observed mean of 0.69 ppbv. The index of agreement increased from 0.63 to 0.70 indicating benefits of increased HONO emissions. Clearly, improvement in HONO peak values can be seen on September 12, 18, 23, 24, 25 and 30, especially on September 12 the model with increased HONO emissions nicely follow HONO peak while the case with low HONO/NOx emission rates resulted in underprediction of the peak value. However; as pointed by Czader et al. (2012) HONO predictions depends on how well the model captures NOx concentrations, especially NO2, since heterogeneous HONO formation is directly related to NO2 concentrations and greatly influences morning HONO mixing ratios. It can be seen that overprediction of NO and NO2 on September 11, 19, and 24 leads to overprediction of HONO. We can conclude that misprediction of precursors is responsible for HONO misprediction and expect that if NOx mixing ratios for those days are accurately simulated also HONO values would be close to observation. This is not a case on September 18 when, despite the fact that NO is well predicted and NO2 overpredicted, HONO peak is underpredicted. The reasoning for that is unknown, but it is probably due to the uncertainties in other HONO sources. Also, variations of simulated HONO mixing ratios from day to day are influenced not only by emissions but also by other parameters, for example, the model capabilities to predict grow of the mixing layer and wind fields as well as
clouds that influence photolysis rates. To more clearly present differences between the two simulated cases (N and NH) and measured data we calculated the average diurnal profiles of HONO and presented them in Figure 7. The modeled profiles follow the measured one showing high peak in the morning and low values during a daytime. It can be seen that the NH scenario, in which higher emission ratio was utilized, improves HONO morning peaks. Since only HONO emissions from mobile sources were increased it is expected to see the largest differences in mixing ratios during early morning times when the traffic emissions are the highest, the mixing layer height low allowing for accumulation of HONO, and photochemistry not very active. It is worth to note that all available measured data for HONO for September 2013 are from weekdays and the higher HONO/NO\textsubscript{x} ratio measured in Houston was also calculated based on measurements taken during weekdays. The model underprediction during daytime can be explained by the fact that the default model version that we used in this study does not account for the photochemical HONO sources. Also, too low modeled average profile during daytime is caused by underpredictions of HONO on Sep. 23-25 which can be attributed to stronger modeled winds in comparison to weak observed winds causing modeled HONO to be removed from the observational site.

The photolysis of HONO is a source of hydroxyl radical. Figure 7 shows a snapshot of spatial pattern of OH mixing ratios (left) and differences in OH mixing ratios (right) between simulations with increased HONO emissions (NH) and regular emissions with 0.008 HONO/NO\textsubscript{x} emissions ratio (N) for September 13\textsuperscript{2}, which is a day with nicely predicted HONO mixing ratios. An increase in OH occurs along highways corresponding to increased HONO mobile emissions. Based on the 1 month of simulated surface concentrations the average increase in HONO due to doubling its emissions from mobile sources is 36% at the location of the Moody Tower and 10% when averaged over the urban area. The average increase in the morning OH (between 6 – 8 a.m. LT) is 14% at the location of the Moody Tower and 3% when averaged over the urban area. The ozone increase is below 1% for both the Moody Tower and the urban area. The average increase in OH during daytime (6 a.m. – 8 p.m. LT) is 7% for the Moody Tower and 1% for the urban area. The increase in ozone is again below 1%. Since HONO emissions from mobile sources that peak in the morning were modified therefore, it is understandable that the
impact of these additional HONO emissions on OH and ozone is higher during morning time than afternoon hours. To obtain more insights on the fate of HONO we performed additional model simulations in which we utilized the process analysis that provides information on chemical and physical processes influencing pollutant mixing ratios. The analysis was performed for the Moody Tower site for Sep. 10-13, 2013. At the surface, at the location of the Moody Tower the average contribution of vertical transport to the loss of HONO is 77%, horizontal transport contributes 8%, chemical removal 11% and dry deposition 5%. HONO mixing ratios along with process affecting changes in mixing ratios for the second model layer, which corresponds to the altitude of measurements, are presented in Figure 9. It can be seen that transport (horizontal and vertical) continue to be a dominant loss process at this altitude contributing on average 77% to the total HONO loss while chemical loss contributes only 23% to the total loss. The chemical loss of HONO is dominant only during couple of morning hours. This explains the fact that even though HONO mixing ratios significantly increased upon additional emissions, HONO was removed mainly by transport with only small portions taking part in chemical reactions converting it to OH and furthermore to $O_3$. Doubling HONO emissions resulted in up to 6% of OH increase. The impact of increasing HONO emissions on ozone mixing ratios is smaller. For example, for September 13, the maximum change in ozone is 0.45 ppbv at 11 am L.T., the impact of increased HONO emissions on the afternoon peak ozone value is even smaller, at the 1 ppt level (not shown).

4. Summary

The WRF - SMOKE - CMAQ modeling system was used for evaluation and adjustment of NOx emissions. In particular, HONO/NOx emission ratio from mobile sources was increased and its impact on HONO mixing ratios as well as on OH and $O_3$ was evaluated. Effects of applying increased HONO/NOx emission ratio from mobile sources on HONO mixing ratios were evaluated.

First, NOx emissions were adjusted to reflect emission trends. Simulations with adjusted NOx emissions resulted in overall better NOx prediction as mixing ratios become closer to measured
values. The average NOx mean value from all analyzed sites dropped from 11.11 ppbv to 7.59 ppbv and is much closer to the observed mean of 7.76 ppbv, IOA is improved in the reduced NOx case (0.71 vs. 0.75) and the AME is lowered from 6.76 to 4.94. Therefore, the reduced NOx case was taken as a base for adjusting HONO emissions according to values measured in Houston.

Doubling HONO emission from mobile sources and therefore making them closer to the newly reported HONO/NOx ratio of 0.017 resulted in increased HONO mixing ratios especially during morning peak values. Based on 1 month of simulated data 36% increase in HONO mixing ratio at the location of the Moody Tower was obtained from the case with higher emission ratios utilized in simulation. The increase in HONO values averaged over the urban area was 10%. Simulated HONO mixing ratios were compared to values measured at the Moody Tower. The mean value increased from 0.30 ppbv in the base HONO emission case to 0.41 ppbv in the increased HONO emission case and become closer to the observed mean of 0.69, but still low. The index of agreement for simulation that used the 2001 HONO/NOx emission ratio of 0.008 is 0.63 while for the simulation with doubled HONO emissions IOA increased to 0.70. Increased HONO emissions from mobile sources resulted in up to 6%–14% increase in OH during morning time at the location of the Moody Tower and 3% when averaged over urban area. The increase calculated for daytime was 7% and 1% for the Moody Tower and the urban area, respectively. The impact on ozone was found to be marginal (below 1%).

This study results could shed light on the underestimated HONO in the morning from global/regional chemical transport model with the typical emission ratio of 0.8% HONO emission out of the total NOx emissions. In addition, since HONO is the major radical source in the morning (e.g., Perner and Platt, 1979; Harris et al., 1982; Czader et al., 2013), underpredictions of HONO would lead to underprediction of OH radical. This study results could shed light on the underestimated HONO and OH in the morning from global/regional chemical transport model with the typical emission ratio of 0.8% HONO emission out of the total NOx emissions.
Acknowledgements. The authors would like to thank the Texas Air Research Center (TARC) for supporting this work. They are also thankful to Lijun Diao for help in setting up WRF and to Hyuncheol Kim for helping with CAMS dataset.

References


Choi, Y. 2014: The impact of satellite-adjusted NOx emissions on simulated NOx and O3 discrepancies in the urban and outflow areas of the Pacific and Lower Middle US, Atmospheric Chemistry and Physics, 14, 675-690.

Choi, Y., Kim, H., Tong, D., Lee, P. 2012: Summertime weekly cycles of observed and modeled NOx and O3 concentrations as a function of land use type and ozone production sensitivity over the Continental United States, Atmospheric Chemistry and Physics, 12, 6291-6307.


doi:10.1016/S1352-2310(01)00138-8

Lefer, B., and B. Rappenglück (2010), The TexAQS-II radical and aerosol measurement project
(TRAMP), Atmos. Environ., 44, 3997-4004.

Li, S., Matthews, J., and Sinha, A.: Response to Comment on “Atmospheric Hydroxyl Radical
Production from Electronically Excited NO2 and H2O”, Science, 324, 336,

Li, S., Matthews, J., and Sinha, A.: Atmospheric Hydroxyl Radical Production from

Li, Y., An, J., Min, M., Zhang, W., Wang, F., Xie, P., 2011: Impacts of HONO sources on the air
quality in Beijing, Tianjin, and Hebei Province of China. Atmospheric Environment, 45, 4735-
4744.

Mao, J., X. Ren, S. Chen, W. H. Brune, Z. Chen, M. Martinez, H. Harder, B. Lefer, B.
Rappenglück, J. Flynn, and M. Leuchner. 2010. Atmospheric oxidation capacity in the summer
of Houston 2006: Comparison with summer measurements in other metropolitan studies. Atmos.

Olaguer, E. P., C. E. Kolb, B. Lefer, B. Rappenglück, R. Zhang, and J. P. Pinto (2013),
Overview of the SHARP campaign: motivation, design, and major outcomes, J. Geophys. Res.

OH + NO(+M)→HONO(+M) and the determination of the UV absorption cross section of

Perner, D., U. Platt: Detection of nitrous acid in the atmosphere by differential optical

precursors and related species from traffic as observed and modeled at an urban highway
junction, Journal of the Air & Waste Management Association, 63:11, 1270-1286, DOI:

Sarwar, G., Roselle, S. J., Mathur, R., Appel, W., Dennis, R. L., and Vogel, B.: A comparison of
CMAQ HONO predictions with observations from the Northeast Oxidant and Particle Study,

Skamarock, W. C., and Klemp, J. B.: A time-split non-hydrostatic atmospheric model for

reduction of nitrogen dioxide on humic acid as a source of nitrous acid, Nature, 440, 195–198,
2006.

Stemmler, K., Ndour, M., Elshorbany, Y., Kleffmann, J., D’Anna, B., George, C., Bohn, B., and
Ammann, M.: Light induced conversion of nitrogen dioxide into nitrous acid on submicron
Table 1. EPA emission trends for NOx (values reported in thousands of tons).

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<th>2010</th>
<th>2011</th>
<th>2012</th>
<th>2013</th>
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<td>6,241</td>
<td>5,734</td>
<td>5,786</td>
<td>5,398</td>
<td>5,010</td>
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<tr>
<td>other</td>
<td>9,872</td>
<td>9,540</td>
<td>9,144</td>
<td>8,594</td>
<td>8,114</td>
<td>7,914</td>
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<tr>
<td>total</td>
<td>16,813</td>
<td>15,781</td>
<td>14,878</td>
<td>14,380</td>
<td>13,512</td>
<td>12,924</td>
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Table 2. Summary of statistical parameters for the base case simulation (B) and reduced NOx case (N).

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<th>AME (ppb)</th>
<th>IOA</th>
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<td>2</td>
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<td>5.42</td>
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Table 3. Statistical parameters for modeling HONO mixing ratios for the Moody Tower site.

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<td>Mean Bias</td>
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<td>Sim. Red-NO$_{x}$ (N)</td>
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<td>Sim. (NH)</td>
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<td>Sim. (NH)</td>
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Figure 1. Locations of stations performing NO\textsubscript{x} measurements in the Houston-Galveston-Brazoria area during September 2013.
Figure 2. Timeseries comparing measured NOx against values simulated with the base case and the reduced NOx case at CAMS sites 1 and 411.
Figure 3. NO and NO$_2$ mixing ratio measured at the Moody Tower site and modeled with the base case emissions as well as with reduced NO$_x$ emissions.
Figure 4. Snapshot of differences in HONO emissions between a case with emission ratio of HONO/NO$_x$=0.016 (NH) and default emissions of HONO/NO$_x$=0.008 (N) at 7 a.m. LT on September 12, 2013. Difference in HONO emissions between increased HONO case (NH) and default HONO emissions (N).

Figure 5. Differences in HONO mixing ratios between a case with 0.016 HONO/NO$_x$ emission ratio (NH) and 0.008 HONO/NO$_x$ emissions (N) for the surface (left) and the second model layer (right) at 7 a.m. LT on September 12, 2013. Differences in HONO mixing ratios between the increased HONO emission case (NH) and the base HONO emissions (N) for the surface (left) and the second model layer (right).
Figure 6. HONO mixing ratios measured at the Moody Tower site and modeled with and the regular HONO emissions (N) for which the HONO/NO$_x$ emission ratio of 0.008 was used, and the increased HONO case (NH) for which the HONO/NO$_x$ emission ratio of 0.016 was used.
Figure 7. Average diurnal variation of HONO at the Moody Tower measurement site.
Figure 7. OH mixing ratios (left) and differences in OH mixing ratios (right) between the case with 0.008 HONO/NO\textsubscript{x} emission ratio (N) and 0.016 HONO/NO\textsubscript{x} emission ratio (NH) at noon local time on September 12, 2013. OH mixing ratios (left) and differences in OH mixing ratios between the base case and increased HONO emission case (right).
Figure 9. HONO mixing ratios (black line) and processes contributing to changes in HONO mixing ratio at the Moody Tower site where the measurements were taken, which corresponds to the second model layer, where VTRAN is vertical transport, HTRAN is transport in horizontal direction, and CHEM_HONO correspond to changes due to chemical reactions.