Review 1 of “NH3 Spatio-temporal variability over Paris, Mexico and Toronto and its link to PM2.5 during pollution events”

Referee: General comments:

The manuscript of egusphere-2022-413 presented the result of a unique project of the AmmonAQ that targeted different three areas of Paris, Toronto, and Mexico City. Towards better air quality by mitigating NH3 emissions, the finding of this study will contribute to improving the atmospheric environment. I generally agree with this manuscript being published; however, some points are concerned. Especially, I am wondering about the uncertainty of the satellite NH3 measurement dataset analyzed in this study. Please address the following specific points.

Authors: We would like to thank the referee for this positive review and all the relevant comments that we have addressed in the following document.

Specific comments:

P3, L52: Please specify these five countries.

The five countries are United-States, China, Netherlands, United-Kingdom, and Canada. We have included this list in the revised manuscript.

P3, L65: Are these increasing trends explained by the expansion of NH3 emissions, or meteorological variability (e.g., temperature)?

In Europe, NH3 emissions are relatively stable (with a decrease of 4% between 2008 and 2012 followed by an increase of 3% between 2013 and 2018). However, IASI observations reveal an increase that could be associated with meteorological conditions, with high temperature and drought recorded in 2018.

In Canada, emissions are also relatively constant between 2008 and 2018, but biomass burning sustains increased NH3 emissions in the northern hemisphere.

In Mexico, the EDGAR inventory shows an increase in NH3 emissions.

These conclusions must be tempered by the fact that the lifetime of NH3 in the atmosphere increases with decreasing concentrations of nitrogen oxides and sulfur.

We have added a sentence in the revised manuscript to explain the trends: “These trends are likely explained by increasing emissions, partly due to increased temperature (Europe) and biomass burning (Canada). However, decreasing concentrations of nitrogen and sulfur oxides e.g. in Europe and China also increase the ammonia atmospheric lifetime and plays a role in the reported upward trends.”

P3, L77: Does “the standard” indicate the standard in Mexico? Because this study conducted the comparison over three regions, it will be better to explicitly state it.

We have included ‘national’ standard to explicitly state it. In addition, we have modified the standard value for Mexico because it has recently changed in October 2021. Therefore, we have modified the sentence as follows: “In Mexico, PM2.5 concentrations often exceed the national standard of 41 µg/m³ for the 24-hour mean [NOM-025-SSA1-2021, 2021]”.

Because of the recent progress in satellite NH$_3$ measurement, I would like to strongly suggest including the discussion of the uncertainty of satellite data, such as the detection limit (https://doi.org/10.5194/acp-19-12261-2019).

In this section of the revised manuscript, we have added description about the detection limit: “The detection limit depends both on the atmospheric state (mainly thermal contrast and NH$_3$ abundance) and the instrument characteristics. For IASI, the minimum detection limit is found to be 4-6x10$^{15}$ molecules/cm$^2$ [Clarisse et al., 2010].”

What is the mottled pattern found over Canada during winter in Figure 3?

The sensitivity of IASI measurements is intimately related to the thermal contrast between the surface and the first layers of the atmosphere [Clerbaux et al., 2009]. When the detection is possible (with good thermal contrast), the peak sensitivity for NH$_3$ is in the boundary layer [Clarisse et al., 2010]. The high value over Canada and the Arctic in winter can be associated with high uncertainties in the NH$_3$ retrievals due to low thermal contrast and high emissivity from snow. We have added this sentence in the revised manuscript to clarify: “The high value over Canada and the Arctic in winter can be associated with high uncertainties in the NH$_3$ retrievals due to low thermal contrast”.

Can all satellite measured NH$_3$ close to zero be used in Figure 4?

Yes, removing or keeping these values won’t change the relationship between temperature and the ammonia averages, therefore the discussion of the following figure will not be affected. In Figure R1, we show the same plots as Figure 4 (in the manuscript) but with the values of NH$_3$ ≤ 0.25 × 10$^{16}$ molecules cm$^{-2}$ removed. Note that the $r^2$ did not change much, in the presence and absence of these values.
Figure R1. Same as Figure 4 in the revised manuscript but with NH$_3$ average $\leq 0.25 \times 10^{16}$ molecules cm$^{-2}$ removed.

Is it available AK when comparing GEOS-Chem? The information on AK and how to calculate it in the comparison with the model is not described.

The IASI retrieval algorithm for NH$_3$ does not provide averaging kernels. We have qualitatively compared IASI NH$_3$ columns with GEOS-Chem NH$_3$ columns using all the morning IASI measurements available, following the recommendation provided in Van Damme et al. [2017]. Previous studies have demonstrated good agreement with surface and FTIR measurements (e.g., Clarisse et al., 2010; Van Damme et al., 2015; Dammers et al., 2017; Viatte et al., 2021; Van Damme et al., 2021), demonstrating that IASI retrievals are sensitive to NH$_3$ located in the lower layers of the atmosphere.

P5, L115: What is the actual gridded data (e.g., Figure 2) analyzed in this study?

We have gridded the IASI data at 0.25° x 0.25° degrees. This information has been included in the revised manuscript as follow: "In this work, we use version 3 of the ANNI-NH$_3$ product [Van Damme et al., 2021; Guo et al., 2021] from IASI Metop-A/B morning overpasses over the period 2008 to 2017 gridded a spatial resolution of 0.25° x 0.25°."
Although we can find the reason to choose the model simulation period of 2011 in P12, L344-L349, it is better to be shortly explained here. We have added a short sentence in this section: “because all three cities experienced both separate and combined PM$_{2.5}$ and NH$_3$ pollution events during this period.”

Are these three panels shown with the same horizontal distance? If different, the scaler might be helpful.

The three panels are approximately shown at the same horizontal distance, but we have inserted a scaler in each panel to help the reader.

![NH$_3$ concentration maps](image)

Figure R2: Same as Figure 2 in the revised manuscript.

Too coarse reanalysis resolution to investigate air mass trajectories on 50 km radius-circle at each city?

We have run 24-hour HYSPLIT back-trajectories every day for the whole IASI dataset from 2008 to 2017. The idea is to determine the effect of long-range transport affecting air quality within the cities during this decade. Running 3652 daily back-trajectories (10 years) using a finer meteorological reanalysis would have been very time consuming. In addition, results derived from a finer resolution meteorological dataset are very similar. We have performed a test for a sub dataset (July 2008) running 24h-backtrajectories ending in Paris using NCEP at 2.5° resolution and GDAS at 1° resolution (Figure R1). Visual inspection of the back trajectories shows that using a 2.5° resolution meteorological dataset is similar to a finer meteorological dataset at 1° resolution. We have inserted in the revised manuscript a sentence to address this concern: “Note that visual inspection of the back trajectories shows that using a 2.5° resolution meteorological dataset is similar to using a finer meteorological dataset at 1° resolution (GDAS).”
Figure R3: 24-h HYSPLIT back-trajectories ending in Paris using the meteorological dataset of GDAS (at 1° resolution, left panel) and NCEP (at 2.5° resolution, right panel) for July 2008.

P7, L188: Same to Europe and southern North America, source information of “(Canada)” or “(U.S.A.)” can be useful in this Table 1.

We have added information about the sources countries in the revised Table 1 as follows:

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*Fertilizer industry ** Soda ash industry
¹ Van Damme et al., 2018; Clarisse et al., 2019
² Dammers et al., 2019

P13, L358: Should the denominator be “observations” when comparing observation and model? Why model is referred to as a criterion?

We agree with the referee that the model is not referred here as a criterion. Therefore, we have changed the denominator to be observations. The result doesn’t change much the MRD, so the discussion remains valid.

P13, L359: Are these values positive? If model underestimation, are these negative?
To calculate the Mean Relative Difference (MRD), we have revised the formula to be more intuitive as follows: \( MRD = \frac{(\text{model} - \text{observations})}{\text{observations}} \) as suggested above. Thus, when the MRD is positive, model data are higher than the observations. Conversely, when the MRD is negative, model data are smaller than the observations. In the latter case, a negative MRD means that the model underestimate compared to the observations.

P14, L379: From the spatial mapping over Europe, this seems to be simply led to model overestimation, and this is not consistent with the timeseries and relevant discussion in the main text. Please confirm this figure.

In the revised figure (see figure R4), we choose to be consistent and show (GEOSChem-IASI) \( \text{NH}_3 \) columns. Therefore, if this difference is negative (blue colour in the figure), then IASI \( \text{NH}_3 \) columns are higher than the model ones, leading to an underestimation of the model. To clarify, we have added “(model-observations)” in the caption of the revised manuscript.

![Figure R4: same as Figure 8 in which we choose to show the difference (model-observations) to be consistent throughout of the revised manuscript.](image)

P15, L413-418: From this comparison on PM2.5 component, I am suspicious about the result in other cities of Paris and Mexico City. When we considered this poor performance for PM2.5 components, the result in Figure 9 and the relevant discussion seems to be meaningless. Is this performance for PM2.5 component useful (worse or better than other studies)? If not, I would like to request to reconsider this final section in P14, L391-P16, L425.

The component comparison is important in Toronto because it helps explain why the model is performing so poorly there.
Unfortunately, to our knowledge, there are no \( \text{PM}_{2.5} \) chemical composition observations available to compare individual components in Paris and Mexico City in March 2011. However, we found in the literature relevant results that could strengthen the final discussion.

In Paris, the March 2011 pollution episode has indeed been analyzed in terms of geographical origins and aerosols properties [Chazette et al., 2017] but not in term of aerosol speciation.

Comparing the GEOS-Chem outputs used in this study with two years of observations of aerosol chemical composition in Paris (2011-2013) [Petit et al., 2015], we found that sulfate component is slightly higher in the GEOS-Chem model than in the springtime observations (21% compared to 11%) whereas modelled organic carbon is lower than the observations (8% compared to 33%). This springtime underestimation of organics in atmospheric models has previously been reported in Paris [Sciare et al., 2010; Petit et al., 2015; Lanzafame et al., 2021] and could be associated with an underestimation of the organic matter emissions from residential contribution [Van der Gon et al., 2015].

Regarding the secondary aerosol, observations in Paris during the March 2015 pollution event show that it accounts for more than 50% of the PM concentration [Petit et al., 2017], which is in agreement with the SNA partition simulated in our GEOS-Chem model.

In Mexico City, a very recent paper [Retama et al., 2022] shows that during the dry-warm season of 2019 (between March and June), SNA correspond to 30% of the aerosol mass concentration, which is consistent with what has been reported before [Cheng et al., 2016] and the chemical composition modelled in our study. The organic fraction is found to be dominant in the observations [Retama et al., 2022] as suggested in the GEOS-Chem model over Mexico City.

We have inserted part of this discussion in the revised manuscript as follows:

“...In Paris, the March 2011 pollution episode has been analyzed in terms of geographical origins and aerosols properties [Chazette et al., 2017] but not in terms of aerosol speciation.

Comparing the GEOS-Chem outputs used in this study with two-years observations of aerosol chemical composition in Paris (2011-2013) [Petit et al., 2015], we found that the sulfate component is slightly higher in the GEOS-Chem model than in the springtime observations (21% compared to 11%) whereas modelled organic carbon is lower than the observations (8% compared to 33%). This springtime underestimation of organics in atmospheric models has previously been reported in Paris [Sciare et al., 2010; Petit et al., 2015; Lanzafame et al., 2021] and could be associated with an underestimation of the organic matter emissions from residential contribution [Van der Gon et al., 2015]. Regarding the secondary aerosol, observations in Paris during the March 2015 pollution event show that it accounts for more than 50% of the PM concentration [Petit et al., 2017], which is in agreement with the SNA partition in our GEOS-Chem model simulation.

In Toronto, \( \text{PM}_{2.5} \) speciation is monitored by the National Air Pollution Surveillance Program (NAPS, https://www.canada.ca/en/environment-climate-change/services/air-pollution/monitoring-networks-data/national-air-pollution-program.html) network. Observations in March 2011 reveal that inorganic nitrate burden is overestimated by a factor 2 in the GEOS-Chem run (41% in the model compared to 20% in the observations), whereas sulfate and black carbon abundances are underestimated by a factor 2 (15 and 6% in the model compared to 27 and 12% in the observations).

In Mexico City, the organic matter represents the most abundant fraction of the aerosol, which is consistent with measurements made during several campaigns performed in the dry season of 2006 during the Megacity Initiative: Local And Global Research Observations (MILAGRO, [Molina et al.,
2010]) and Aerosoles en Ciudad Universitaria (ACU) in 2015 [Salcedo et al., 2018]. Observations performed during the dry-warm season of 2019 reported that SNA correspond to 30% of the aerosol mass concentration, which is consistent with what has been reported before [Cheng et al., 2016] and the chemical composition modelled in our study. The organic fraction is found to be dominant in the observations [Retama et al., 2022] as suggested in the GEOS-Chem model over Mexico-City. Daily cycles appear overexaggerated in the model with maxima well represented and minima greatly underestimated. This could suggest model issues in term of atmospheric dynamics (removal/transport or planetary boundary layer dynamics) due to coarseness of grid.”

**Technical corrections:**

Authors: All technical corrections have been addressed in the revised manuscript.

P10, L281: 4.71 “×” 1015?

P12, L324 and L325: “m.s−1”? Does it need the period?

References:


Sciare, J., d’Argouges, O., Zhang, Q. J., Sarda-Estève, R., Gaimoz, C., Gros, V., Beekmann, M. and Sanchez, O.: Comparison between simulated and observed chemical composition of fine aerosols in Paris (France) during


Review 2 of “NH3 Spatio-temporal variability over Paris, Mexico and Toronto and its link to PM2.5 during pollution events”

Referee:

Summary

This paper analyses 10 years of IASI NH3 data over three large domains, each of which encompasses a major metropolitan area (Paris, Toronto and Mexico City). The ten year average and seasonal means of the NH3 total columns from the IASI instruments deployed on Metop-A and Metop-B are presented and the relationships between NH3 amounts and temperature and precipitation are evaluated; the authors find a strong correlation between temperature and NH3 amounts in the Paris and Toronto domains, but only a weak one in the Mexico City domain. This analysis is extended to sub-domain scales, focusing on a number of previously identified source regions, and expanded to include the effect of relative humidity. Many of the smaller domains show an interesting local maxima in the NH3 vs temperature plot that coincides with fertilization activities. The impact of wind direction on air quality is also examined, using back-trajectories from HYSPLIT and a cluster analysis, along with in situ PM2.5 data from local networks. The PM 2.5 data is also used in conjunction with the NH3 columns to identify and count pollution events; wind roses are then constructed to determine the wind patterns on days with high pollution; interestingly PM2.5 and NH3 are not always high on the same days. Finally IASI NH3 and the local PM2.5 data are compared against GEOS-Chem output for one month. This last analysis is interesting but too limited to provide really useful information on GEOS-Chem performance. The paper is well laid out and clearly written. The plots are of high quality and in general easily understood, though a few require more detailed captions, described below. It requires only some minor edits and clarifications to be acceptable for publication. Overall, a good illustration of how to apply a number of different techniques and data sources to the problem of understanding the drivers of pollution events.

Authors: We would like to thank the referee for this positive review and all the relevant comments that we have addressed in the following document.

Technical comments

Section 3.1: How are the IASI averaged and what is the grid resolution? How are the boxes doe each source region defined?

We have gridded the IASI data at 0.25° x 0.25° degrees. This information has been included in the revised manuscript. The sources regions are defined by looking at local enhancement in the 10-year average of the IASI total column.

Figure 3: Please comment on the high NH3 values over the Arctic.

The sensitivity of IASI measurements is intimately related to the thermal contrast between the surface and the first layers of the atmosphere [Clerbaux et al., 2009]. When the detection is possible (with good thermal contrast), the peak sensitivity for NH3 is in the boundary layer [Clarisse et al., 2010]. The high value over Canada and the Arctic in winter can be associated with high uncertainties
in the NH$_3$ retrievals due to low thermal contrast and high emissivity from snow. We have added in the revised manuscript this sentence to clarify: «The high value over Canada and the Arctic in winter can be associated with high uncertainties in the NH$_3$ retrievals due to low thermal contrast».

Lines 266-274: Is each back-trajectory associated with the 50 km NH$_3$ mean for that day? Could the authors please briefly describe the clustering approach? Are the NH$_3$ means clustered according to the corresponding back-trajectories?

We have explained the method following the different steps (Figure R1) and added these descriptions in the revised supplementary information Figure S7.

1) For each day, we have run HYSPLIT back-trajectories ending in the cities at the overpass time of the IASI satellite (blue lines in Figure R1).

2) For each day, we have calculated the amount of NH$_3$ derived from IASI observations within a circle of 50km radius around the cities (orange cylinder in Figure R1).

3) We have run the cluster analysis to merge trajectories that are near each other (green lines in Figure R1). The cluster analysis computes the spatial variance and minimize differences between trajectories within a cluster while differences between clusters are maximized [Abdalmogith et al., 2005; https://www.ready.noaa.gov/documents/Tutorial/html/traj_cluster.html]. NH$_3$ mean concentrations measured inside the cities by IASI have been allocated to the different mean cluster trajectories according to the corresponding back-trajectories.

*Figure R1: method to analyze the impact of long-range transport on NH$_3$ concentrations measured over the cities*

Line 289: Which sources have increased near Toronto?

According Yao and Zhang (2019), NH$_3$ concentrations around Toronto (near Granby – box A in Figure 2 and Elmira-Kitchener-Guelph – box B in Figure 2) have increased in the last decade as opposed to the reported emissions. Yao and Zhang (2019) explained the NH$_3$ trends by the reduced chemical loss of atmospheric NH$_3$ to form particulate NH$_4^+$. We have modified the sentence in the revised manuscript to clarify this point: “9 to 17% of NH$_3$ concentrations are coming from the west and the east of Toronto (cluster 3, 4, and 5) where atmospheric NH$_3$ have increased in the last decade possibly due to the reduced chemical loss of atmospheric NH$_3$ to form particulate NH$_4^+$ (Boxes A and B in Figure 2, [Yao and Zhang, 2019])”

Line 303: What are the criteria used for defining a pollution event?
We have used the same method than the one conducted in Yamanouchi et al. (2021). Pollution events are defined when residuals of the Fourier fit are above 2 standard deviations. We have inserted “2 standard deviations” in the sentence of the revised manuscript.

**Line 313: It would be useful to add the spring plot here**

We have added the plot showing the seasonal occurrence of the pollution events in the revised manuscript.

**Line 319: What does the radial distance in the wind roses in Figure 7 indicate?**

The radial distance in the wind roses indicates the frequency of the wind direction occurrence. We have added this information in the revised manuscript.

**Line 350: Please state the coincidence criteria.**

The criteria were stated in the following sentence. To clarify, we have arranged this sentence: “Spatial and temporal coincidence criteria have been applied to GEOS-Chem outputs to compare with IASI morning observations, such as: model outputs between 8.30 and 11.30 AM coincident with IASI overpasses have been selected, and only collocated model outputs (at 0.5° × 0.625° horizontal resolution) have been selected coincident with IASI observations.”

**Lines 360-364: I don’t agree that the GEOS-Chem and IASI are in good agreement; the authors should state that the two datasets capture some of the same pollution events.**

We have modified the text accordingly.

**Figure S4: Are the bins for the RH plot also specified by temperature? This does not seem right.**

In Figure S4, lower panel, the NH$_3$ values are averaged per bins of relative humidity with a 1% RH between each consecutive bin. We modified the description of Figure S4 in order to make this clearer.

**Minor edits**

Authors: All minor edits have been addressed in the revised manuscript.

Lines 32-33: ... in Paris and Mexico City;

Line 65: However, NH$_3$ concentrations are increasing in many countries: France, Canada and Mexico reported increases of .......

Line 67: ... are composed of ...

Line 72: ... nitrate formed from ...

Line 127: ... are located within a 50-km

Line 134: ... the same month

Line 150: Note that

Line 153: ... daily 24-hour back-trajectories ...
Finally, all back-trajectories are combined. (not sure here what the authors mean to say)

We have modified this sentence as: “Finally, every back-trajectory that are near to each other are merged in clusters and associated with the corresponding local-scale IASI NH₃ concentrations”.

practices (which are dominant over Europe and North America)

in this region closer to the Equator

(with small contributions from industries)

concentrations. In order to analyze ...

associated with the highest ...

on average, are originating ...

concentrations are observed

numbers of coincident observations

sources are

surrounding regions:

These lead

and Mexico pollution is transported along the northeast-southwest line,

launched in

References:


