## **Co-Editor Decision: Publish subject to technical corrections (10 Jul 2017) by Jennifer G. Murphy** Comments to the Author:

I encourage the authors to consider the comments from Reviewer 1 regarding the revised version of the manuscript. In particular, it would be beneficial to have more clarity about which version of the  $NH_3$  dataset was used and to what extent revisions might impact the trend analysis. Once this issue and the other details have been addressed, the manuscript is ready for publication.

#### Dear Dr. Murphy:

We have addressed the comments raised by Referee 1, and incorporated the comments/suggestions in the revised manuscript. In addition, we have also added detailed descriptions in Sect. 2.2 to clarity about which version of the NH<sub>3</sub> dataset was used and to what extent revisions might impact the trend analysis (lines 191-194).

Thank you very much for your consideration.

Sincerely,

Xiuying Zhang

On behalf of all co-authors

#### Referee #1

The authors have made an effort to address reviewer comments. Most technical issues have been addressed, but I have several suggestions for further revision.

We are grateful to the reviewer for the time and energy in providing helpful comments and guidance that have improved the manuscript. In this document, we describe how we have addressed the reviewer's comments. Detailed responses to each comment are given below (in blue).

1) The authors state in their response that this analysis is important because of the discussion on "the possible interactions between  $NO_2$  and  $NH_3$ ". However, this really isn't included or addressed in the manuscript. What exactly have the authors learned about this interaction, given the current results of their work?

Please refer to lines 276-286 in the main text.  $NH_3$  is an important alkaline gas that is abundant in the atmosphere, and is able to neutralize acidic components including  $HNO_3$  and  $H_2SO_4$  through the oxidation of  $NO_x$  and  $SO_2$ . We here just show a possible reason for an increase in  $NH_3$  columns in recent years due to decreased acidic gases ( $NO_2$  and  $SO_2$ ) under strong control policy in 12-th FYP over China.

2) The authors mention the update in the NH3 retrieval in September 2014, but it's unclear whether they are actually using this updated retrieval. Are they? If so, please explicitly state this. If they aren't, they must include a comment (in response to the comment by Van Damme in the open discussion) on how they expect this update could affect their conclusions about NH3 trends, if at all.

We have added the following text in Sect. 2.2 for clarifications:

"We did not use the IASI NH<sub>3</sub> after September 30 in 2014 for the trend analysis because an update of the input meteorological data on 30 September 2014 has caused a substantial increase of the retrieved

atmospheric NH3 columns."

#### Specific suggestions:

The trends are now included in %/yr. However, these are linear trends. So the authors must explain what this is relative to... Is it a % based on the first year of data? Or of the long term mean? First, NO2 increases by 4% per year to 2011, then decreases by 3.6 % per year to 2015... These % increases/decreases are very hard to compare when we don't know what the % is relative to. The authors must be more careful to explicitly explain how trends were calculated in their methods.

We have added the following text in Fig. 3 captions for clarifications:

"The percent increase or decrease rate (%) was the long term mean calculated by  $[100 \times (\frac{Y_2 - Y_1}{Y_1} + \frac{Y_3 - Y_2}{Y_2} + \dots + \frac{Y_{n+1} - Y_1}{Y_n})] \times \frac{1}{n}$ ."

 Section 2.3, the authors write "...although their thread values of 0.24 kg N ha-1 yr-1...". What is a "thread" value? This wording is not clear to me.

#### We have added "(the slope in Fig. 2)" after "thread" for clarification.

2) I think the trend lines in the top panels of Figure 3a and 3b should be removed, since they are no longer commenting on the linear trend using monthly means.

#### We have removed them as suggested.

As a further suggestion, it's strange having two different ranges on the same plot for the same species (bottom panels) For example, NO2 ranges from 1.5-3 on the left side, but from 1-7 on the right side). I suggest two separate narrow panels so that we can actually distinguish the lines.

The different ranges on the same plot are mainly because the different scales of NO<sub>2</sub> ranges in warm (April-September) and cold (October-March) months.

To distinguish these two lines, we have changed the range of right Y-axis to "from 1 to 5".

Temporal characteristics of atmospheric ammonia and nitrogen dioxide over China based on emission data, satellite observations and atmospheric transport modeling since 1980

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#### Abstract

China is experiencing intense air pollution caused in large part by anthropogenic emissions of reactive nitrogen (Nr). Atmospheric ammonia (NH<sub>3</sub>) and nitrogen dioxide (NO<sub>2</sub>) are the most important precursors for Nr compounds (including N<sub>2</sub>O<sub>5</sub>, HNO<sub>3</sub>, HONO and particulate NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup>) in the atmosphere. Understanding the changes of NH<sub>3</sub> and NO<sub>2</sub> has important implications for the regulation of anthropogenic Nr emissions, and is a requirement for assessing the consequence of environmental impacts. We conducted the temporal trend analysis of atmospheric NH<sub>3</sub> and NO<sub>2</sub> on a national scale

since 1980 based on emission data (during 1980-2010), satellite observations (for NH<sub>3</sub> since 2008 and for NO<sub>2</sub> since 2005) and atmospheric chemistry transport modeling (during 2008-2015).

Based on the emission data, during 1980-2010, both significant continuous increasing trend of NH<sub>3</sub> and  $NO_x$  were observed from REAS (Regional Emission inventory in Asia, for NH<sub>3</sub> 0.17 kg N ha<sup>-1</sup> y<sup>-2</sup> and for NO<sub>x</sub> 0.16 kg N ha<sup>-1</sup> y<sup>-2</sup>) and EDGAR (Emissions Database for Global Atmospheric Research, for NH<sub>3</sub> 0.24 kg N ha<sup>-1</sup> y<sup>-2</sup> and for NO<sub>x</sub> 0.17 kg N ha<sup>-1</sup> y<sup>-2</sup>) over China. Based on the satellite data and atmospheric chemistry transport modeling named as the Model for Ozone and Related chemical Tracers, version 4 (MOZART-4), the NO<sub>2</sub> columns over China increased significantly from 2005 to 2011 and then decreased significantly from 2011 to 2015; the satellite-retrieved  $NH_3$  columns from 2008 to 2014 increased at a rate of 2.37% y<sup>-1</sup>. The decrease in NO<sub>2</sub> columns since 2011 may result from more stringent strategies taken to control NOx emissions during the 12th Five-Year-Plan, while no control policy focused on NH<sub>3</sub> emissions. Our findings provided an overall insight on the temporal trends of both NO<sub>2</sub> and NH<sub>3</sub> since 1980 based on emission data, satellite observations and atmospheric transport modeling. These findings can provide a scientific background for policy-makers that are attempting to control atmospheric pollution in China. Moreover, the multiple datasets used in this study have implications for estimating long-term Nr deposition datasets to assess its impact on soil, forest, water and greenhouse balance.

Keywords: trends, seasonal cycle, ammonia

#### 1. Introduction

Reactive nitrogen (Nr) emissions have increased significantly in China due to anthropogenic activities such as increased combustion of fossil fuels, over-fertilization and high stocking rates of farm animals (Canfield et al., 2010;Galloway et al., 2008;Liu et al., 2013). Elevated Nr in the environment has led to

a series of effects on climate change and ecosystems, e.g. biodiversity loss, stratospheric ozone depletion, air pollution, freshwater eutrophication, the potential alteration of global temperature, drinking water contamination, dead zones in coastal ecosystems and grassland seed bank depletion (Basto et al., 2015;Lan et al., 2015;Shi et al., 2015). Atmospheric reactive N emissions are dominated by nitrogen oxides (NO<sub>x</sub> = NO + NO<sub>2</sub>) and ammonia (NH<sub>3</sub>) (Li et al., 2016a;Galloway et al., 2004). Atmospheric NO<sub>2</sub> and NH<sub>3</sub> are the most important precursors for Nr compounds including N<sub>2</sub>O<sub>5</sub>, HNO<sub>3</sub>, HONO and particulate NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup> in the atmosphere (Xu et al., 2015;Pan et al., 2012). Therefore, an understanding of both the spatial and temporal patterns of NO<sub>2</sub> and NH<sub>3</sub> is essential for evaluating N-enriched environmental effects, and can provide the scientific background for N pollution mitigation.

To investigate the spatial and temporal variations of atmospheric NO<sub>2</sub> and NH<sub>3</sub>, ground measurements are acknowledged to be an effective way in monitoring the accurate concentrations of NO<sub>2</sub> and NH<sub>3</sub> (Xu et al., 2015;Pan et al., 2012;Meng et al., 2010). Ground measurements of NO<sub>2</sub> concentrations in China, including about 500 stations in 74 cities, have been monitored and reported to the public since January 2013 (Xie et al., 2015). By the end of 2013, this network was extended with hourly NO<sub>2</sub> concentrations from more than 850 stations in 161 cities. However, there are fewer NH<sub>3</sub> measurements across China than NO<sub>2</sub> measurements. The China Agricultural University has organized a Nationwide Nitrogen Deposition Monitoring Network (NNDMN) since 2010, consisting of 43 monitoring sites covering urban, rural (cropland) and background (coastal, forest and grassland) areas across China (Xu et al., 2015;Liu et al., 2011). Xu et al. (2015) reported the ground NH<sub>3</sub> concentrations throughout China for the first time, providing great potential to understand the ground NH<sub>3</sub> concentrations on a national scale. Other networks include (1) the Chinese Ecosystem Research Network (CERN) which was established in 1988, including 40 field stations (Fu et al., 2010). However, to our knowledge, there are no detailed reports about ground NH<sub>3</sub> concentrations from CERN on a national scale. (2) Four Chinese cities (Xiamen, Xi-An, Chongqing and Zhuhai) have joined the Acid Deposition Monitoring Network in East Asia (EANET) since 1999. However, only one site (Hongwen, Xiamen) in EANET measured the ground NH<sub>3</sub> concentrations and that data is not continuous. Finally, ground NH<sub>3</sub> concentrations at ten sites in Northern China from 2007 to 2010 have been reported by Pan et al. (2013). All of the above ground measurements provide the potential to understand NH<sub>3</sub> and NO<sub>2</sub> concentrations on a regional scale. However, there is limited information on the spatial and temporal variations of NH<sub>3</sub> and NO<sub>2</sub> in the atmosphere across China. This is due to the limited observation sites and monitoring period, as well as given the uneven distribution of the monitoring sites. Importantly, atmospheric NH<sub>3</sub> and NO<sub>2</sub> monitoring based on ground-based local sites may have limited spatial representativeness of the regional scale as both NH<sub>3</sub> and NO<sub>2</sub> are highly variable in time and space (Clarisse et al., 2009;Wichink Kruit et al., 2012;Boersma et al., 2007).

In order to complement ground-based measurements, satellite observation of NH<sub>3</sub> and NO<sub>2</sub> is a welcome addition for analyzing the recent trends of NH<sub>3</sub> and NO<sub>2</sub> in the atmosphere. Satellite remote sensing offers an opportunity to monitor atmospheric NH<sub>3</sub> and NO<sub>2</sub> with high temporal and spatial resolutions (Warner et al., 2017;Li et al., 2016b). NO<sub>2</sub> was measured by multiple space-based instruments including the Global Ozone Monitoring Experiment (GOME), SCanning Imaging Absorption SpectroMeter for Atmospheric CHartographY (SCIAMACHY), Ozone Monitoring Instrument (OMI) and Global Ozone Monitoring Experiment-2 (GOME-2). The OMI NO<sub>2</sub> provides the best horizontal resolution  $(13 \times 24 \text{ km}^2)$  among instruments in its class and near-global daily coverage (Levelt et al., 2007). OMI observations have been widely applied in environmental-related studies and

for the support of emission control policy (Russell et al., 2012; Zhao and Wang, 2009; Castellanos et al., 2015;Lamsal et al., 2015;Liu et al., 2016a;Foy et al., 2016). First measurements of NH<sub>3</sub> from space were reported over Beijing and San Diego areas with the Tropospheric Emission Spectrometer (TES) (Beer et al., 2008) and in fire plumes in Greece with the Infrared Atmospheric Sounding Interferometer (IASI) (Coheur et al., 2009). The first global map of NH<sub>3</sub> was created from IASI measurements by correlating the observed brightness temperature differences to NH<sub>3</sub> columns using the averaged datasets in 2008 (Clarisse et al., 2009). Shortly after that, many studies focused on developing techniques to gain more reliable NH<sub>3</sub> columns (Whitburn et al., 2016a;Van Damme et al., 2014b), validating the retrieved NH<sub>3</sub> columns using the ground measurements (Van Damme et al., 2014a;Dammers et al., 2016) and comparing the data with the results of the atmospheric chemistry transport models (Van Damme et al., 2014c; Whitburn et al., 2016a), and the estimated NH<sub>3</sub> columns obtained from Fourier transform infrared spectroscopy (FTIR) (Dammers et al., 2016). The retrieval algorithm of obtaining IASI NH<sub>3</sub> columns was based on the method described in Whitburn et al. (2016). Two main steps were performed to derive the  $NH_3$  columns from the satellite measurements. First, derive the spectral hyperspectral range index (HRI) based on each IASI observations (Walker et al., 2011; Van Damme et al., 2014b). Second, convert HRI to NH<sub>3</sub> columns based on a constructed neural network with input parameters including vertical NH<sub>3</sub> profile, satellite viewing angel, surface temperature and so on (Whitburn et al., 2016a). The progresses made on the satellite techniques provide possibility for understanding both the spatial and temporal variations of NH<sub>3</sub> and NO<sub>2</sub> in the atmosphere.

In addition to satellite observations, the emission data are also very important for investigating the temporal trends of NH<sub>3</sub> and NO<sub>2</sub> such as the IIASA inventory (Cofala et al., 2007), EDGAR (Emission

Database for Global Atmospheric Research, RAINS-Asia (Regional Air Pollution Information and Simulation) and Asia REAS (Regional Emission inventory in Asia). REAS is considered as the first inventory by integrating historical, current and future emissions data for Asia based on a consistent methodology (Ohara et al., 2007), and EDGAR is the global emission data with 0.1 by 0.1 grid, which has the highest spatial resolution among different datasets mentioned above. Thus, REAS and EDGAR are used to analyze the historical trends of NH<sub>3</sub> and NO<sub>2</sub> during 1980-2010 in this study. Based on the EDGAR emission data, a widely used atmospheric transport model named as the Model for Ozone and Related chemical Tracers, version 4 (MOZART-4) was also used to model the temporal trend of NH<sub>3</sub> and NO<sub>2</sub> columns during 2008-2015 in comparison with the temporal trends of NH<sub>3</sub> and NO<sub>2</sub> columns measured by satellite instruments.

We aim at getting an overall insight on the temporal trends of both  $NO_2$  and  $NH_3$  since 1980 based on the multiple datasets including the emission data, satellite observations and atmospheric transport modeling. We herein show the Chinese national trend of REAS and EDGAR  $NH_3$  and  $NO_x$  emission data during 1980-2010, satellite-retrieved  $NH_3$  during 2008-2015 and  $NO_2$  columns (2005-2015), and atmospheric transport chemistry modeling  $NH_3$  and  $NO_2$  columns (2008-2015). It should be noted here that the satellite  $NH_3$  columns were retrieved from the IASI, and can only be obtained since 2008. It is beneficial to analyze the temporal variations of both  $NH_3$  and  $NO_2$ , hence providing a scientific basis for policy makers to reduce N-enriched environmental pollution in China.

#### 2. Materials and methods

#### 2.1. NH<sub>3</sub> and NO<sub>2</sub> Emissions

We examined the emission inventory dataset for Asia REAS (Regional Emission inventory in Asia) with 0.5  $^{\circ}\times$  0.5  $^{\circ}$  resolution for the period 1980-2010, and analyzed the temporal trends of NO<sub>x</sub> and NH<sub>3</sub>

over China. REAS v1.1 is believed to be the first inventory of integrating past, present and future dataset in Asia based on a consistent methodology. The REAS datasets have been validated by several emissions, and denote agreement with the recent growth status in Chinese emissions (Ohara et al., 2007). We also collected NO<sub>x</sub> and NH<sub>3</sub> emission data from EDGAR (Emissions Database for Global Atmospheric Research) v4.3.1, which was developed by the Netherlands Environmental Assessment Agency and European Commission Joint Research Centre (Jgj et al., 2002). The EDGAR emissions are calculated on the basis of a point emissions inventory conducted by the International Energy Agency. EDGAR also has a long time period 1980-2010 with the highest spatial resolution globally (0.1 °×0.1 °) (http://edgar.jrc.ec.europa.eu/overview.php?v=431).

### 2.2. Satellite observations

IASI is a passive remote-sensing instrument operating in nadir mode and measures the infrared radiation emitted by the Earth's surface and the atmosphere (Clarisse et al., 2009). It covers the entire globe twice a day, crossing the equator at a mean solar local time of 9:30 A.M. and P.M. and has an elliptical footprint of 12 by 12 km up to 20 by 39 km depending on the satellite-viewing angle. In this study we use daytime satellite observations as these are more sensitive to NH<sub>3</sub> and are associated with a large positive thermal contrast and a significant amount of NH<sub>3</sub> (Van Damme et al., 2014b;Whitburn et al., 2016a). The availability of measurements is mainly driven by the cloud coverage as only observations with cloud coverage lower than 25% are processed to be a good compromise between the number of data kept for the analysis and the bias due to the effect of clouds. As the amount of daily data is not always sufficient to obtain meaningful distributions (due to cloud cover or the availability of the temperature profiles from the EUMETSAT operational processing chain) (Van Damme et al., 2014b), it is more appropriate to consider monthly or yearly averages for this trend analysis. We

consider IASI observations with a relative error below 100% or an absolute error below  $5 \times 10^{15}$  molec. cm<sup>-2</sup> for analysis over China. For the error, the filtering depends on the use of the data. Doing this, low columns typical for background conditions with a large relative error but a small absolute error are also taken into account. For other applications, such as comparing with ground measurements, we would recommend to use a threshold of 75% or even 100% relative error. We gained the data upon request from the Atmospheric Spectroscopy Group at Université Libre De Bruxelles (http://www.ulb.ac.be/cpm/atmosphere.html). This data can be gridded on 0.1 °latitude×0.1 °longitude (Dammers et al., 2016), 0.25 °latitude×0.25 °longitude (Whitburn et al., 2016a) and 0.5 °latitude×0.5 ° longitude (Whitburn et al., 2016b) or even coarser resolutions depending on the usage of the data. For IASI NH<sub>3</sub>, we firstly divided China into  $0.5^{\circ}$  latitude× $0.5^{\circ}$  longitude grid. For each grid cell, we calculated the monthly arithmetic mean by averaging the daily values with observations points within the grid cell. Similarly, we calculated the annual arithmetic mean by averaging the daily values with observations points within the grid cell over the whole year.

The NO<sub>2</sub> columns are obtained from the OMI instrument on NASA's EOS Aura satellite globally everyday. We used the generated products by the project "Derivation of Ozone Monitoring Instrument tropospheric NO<sub>2</sub> in near-real time" (DOMINO) to analyze the temporal trends of NO<sub>2</sub> columns over China. In DOMINO products, only the observations with a cloud radiance fraction below 0.5 were processed for analysis. The retrieval algorithm is described in detail in the previous work (Boersma et al., 2007) and recent updates can be found in the DOMINO Product Specification Document (<u>http://www.temis.nl/docs/OMI NO2 HE5 1.0.2.pdf</u>). We used tropospheric NO<sub>2</sub> retrievals from the DOMINO algorithm v2.0. The retrieval quality of NO<sub>2</sub> products is strongly dependent on different aspects of air mass factors, such as radiative transfer calculations, terrain heights and surface albedo.

The OMI v2.0 data were mainly improved by more realistic atmospheric profile parameters, and include more surface albedo and surface pressure reference points than before (Boersma et al., 2011:Boersma 2016). The DOMINO al.,  $NO_2$ datasets available et are from http://www.temis.nl/airpollution/no2.html. We should state in particular that we used directly the DOMINO v2.0 products of monthly means from 2005 to 2015 over China for the trend analysis. The DOMINO NO<sub>2</sub> columns were gridded at a resolution of 0.125 °latitude×0.125 °longitude grid globally, which has been widely used for scientific applications (Ma et al., 2013;Ialongo et al., 2016;Castellanos et al., 2015).

To illustrate measurement availability, we presented here some measurement statistics. A total number of cloud-free daytime observations as characterized by the operational IASI processor by year were retrieved in China during 2008-2015 for NH<sub>3</sub> (Fig. 1b). We retrieved more observation numbers after 2010 than those during 2008-2009. In 2010, the update of the improved air temperature profiles, cloud properties products and cloud detection, which are important for calculating the thermal contrast, increased the quality of retrieval (Van Damme et al., 2014b;Van Damme et al., 2014c). In September 2014, there was another update of the air temperature profiles, cloud properties products and cloud detection for calculating the thermal contrast. The version of IASI NH<sub>3</sub> columns used in the present work was based on the method described in Whitburn et al. (2016). We did not use the IASI NH<sub>3</sub> after September 30 in 2014 for the trend analysis because an update of the input meteorological data on 30 September 2014 has caused a substantial increase of the retrieved atmospheric NH<sub>3</sub> columns. For the updates of the IASI-NH<sub>3</sub> data, you can refer to Van Damme et al. (2014b), Van Damme et al. (2014c) and Whitburn et al. (2016). The monthly observation numbers are also presented in Fig. 1a, showing that spring (Mar, Apr and May), summer (Jun, Jul and Aug), autumn (Sep, Oct and Nov) and winter (Dec, Jan and Feb) months represent 29%, 26%, 23% and 21%, respectively. Compared with large variations of observation numbers for  $NH_3$ , the observation numbers for  $NO_2$  varied less by year; winter season had the least, while other seasons varied little.

#### 2.3. Atmospheric transport chemistry model

Atmospheric transport chemistry model is also of central importance in modeling the tropospheric NO<sub>2</sub> and NH<sub>3</sub>. We applied a widely used atmospheric global atmospheric transport chemistry model named as the Model for Ozone and Related chemical Tracers, version 4 (MOZART-4) to simulate the tropospheric NO<sub>2</sub> and NH<sub>3</sub> columns during 2008-2015 in accordance with the time period of IASI NH<sub>3</sub> measurements.

The MOZART-4 model is driven by the meteorological data from the NASA Goddard Earth Observing System Model, Version 5 (GEOS-5) at a resolution of  $1.9^{\circ}$  latitude  $\times 2.5^{\circ}$  longitude spatially. The emission data applied for driving the simulations are based on the updated EDGAR emission inventories. 12 bulk aerosol compounds, 39 photolysis, 85 gas species as well as 157 gas-phase reactions were integrated in MOZART-4. The chemical mechanism on N compounds including the NO<sub>2</sub>, NH<sub>3</sub> and aerosols are detailedly integrated to MOZART-4, which is considered to be suitable for tropospheric chemical compositions (Emmons et al., 2010;Pfister et al., 2008;Sahu et al., 2013). The output data used in the current work are temporally varying six hours every day, which were upon request by Louisa Emmons at National Center for Atmospheric Research (NCAR). The monthly means of NO<sub>2</sub> and NH<sub>3</sub> columns were averaged by the daily data, and then used for the trend analysis over China. For more details about MOZART-4, the reader should refer to previous studies (Emmons et al., 2010;Brasseur et al., 1998;Beig and Singh, 2007).

#### 3. Results and discussions

#### 3.1. NH<sub>3</sub> and NO<sub>2</sub> emissions during 1980-2010

We conducted the temporal analysis of NH<sub>3</sub> and NO<sub>x</sub> emissions since 1980 based on REAS and EDGAR. Both significant continuous increasing trends of NH<sub>3</sub> and NO<sub>x</sub> were observed from REAS (for NH<sub>3</sub> 0.17 kg N ha<sup>-1</sup> y<sup>-2</sup> and for NO<sub>x</sub> 0.16 kg N ha<sup>-1</sup> y<sup>-2</sup>) and EDGAR (for NH<sub>3</sub> 0.24 kg N ha<sup>-1</sup> y<sup>-2</sup> and for NO<sub>x</sub> 0.17 kg N ha<sup>-1</sup> y<sup>-2</sup>) over China (Fig. 2). We found a relatively consistent increase in NO<sub>x</sub> emission from EDGAR and REAS over China, i.e. 0.17 kg N ha<sup>-1</sup> y<sup>-2</sup> vs 0.16 kg N ha<sup>-1</sup> y<sup>-2</sup>, but inconsistency in the magnitude of NH<sub>3</sub> emissions from EDGAR and REAS over China, i.e. 0.17 kg N ha<sup>-1</sup> y<sup>-2</sup> vs 0.16 kg N ha<sup>-1</sup> y<sup>-2</sup>. The increase rate in NH<sub>3</sub> emissions over China from EDGAR was much higher than that from REAS, indicating the magnitude of increase trend in NH<sub>3</sub> over China remains a debate, although their thread values (the slope in Fig. 2) of 0.24 kg N ha<sup>-1</sup> y<sup>-2</sup> (EDGAR) vs 0.17 kg N ha<sup>-1</sup> y<sup>-2</sup> (REAS) both reflected a continuous increasing trend (in this regard they are consistent). It implies that, at least, the NH<sub>3</sub> emissions are indeed increasing during 1980-2010. We also conducted a simple correlation analysis of the NH<sub>3</sub> (Fig. 2a) and NO<sub>x</sub> (Fig. 2b) from REAS and EDGAR, showing agreement in the magnitude (slope=1.06) and temporal trend (R<sup>2</sup>=0.96) for NO<sub>x</sub>, but some inconsistency in the increase rate (slope=1.33) for NH<sub>3</sub>.

The discrepancy in the magnitude of  $NH_3$  increase rate from REAS and EDGAR (0.24 kg N ha<sup>-1</sup> y<sup>-2</sup> vs 0.17 kg N ha<sup>-1</sup> y<sup>-2</sup>) in China since 1980 may be caused by the different emission factors considered for estimating  $NH_3$  emissions. The EDGAR v4.3.1  $NH_3$  emissions were calculated based on a variety of sectors including agriculture, shipping, waste solid and wastewater, energy for buildings, process emissions during production and application, power industry, oil refineries, transformation industry, combustion for manufacturing, road transportation, railways, pipelines and off-road transport, while the

REAS v1.1 NH<sub>3</sub> emissions focused mainly on the agriculture source (i.e., manure management of livestock and fertilizer application) (Crippa et al., 2015;Ohara et al., 2007). Moreover, the fundamental methodology on estimating the REAS v1.1 NH<sub>3</sub> emissions did not consider the seasonal agricultural variations compared with that of EDGAR v4.3.1 NH<sub>3</sub> emissions (Kurokawa et al., 2013), and the removal efficiency (as a key element to estimate NH<sub>3</sub> emissions) was also reported to be much higher in REAS v1.1 than in EDGAR v4.3.1 (Kurokawa et al., 2013).

A previous study (Liu et al., 2013) summarized published data on the national anthropogenic NH<sub>3</sub> and NO<sub>x</sub> emissions with multi-periods in China (Wang et al., 2009;Wang et al., 1997;Streets et al., 2003;Klimont et al., 2001;Sun and Wang, 1997;Olivier et al., 1998;FRCGC, 2007), and also analyzed the temporal pattern of NH<sub>3</sub> emissions. Their results showed that the NH<sub>3</sub> emissions had increased at an annual average rate of 0.32 Tg N y<sup>-2</sup> (about 0.33 kg N ha<sup>-1</sup> y<sup>-2</sup>). The increase rate of NH<sub>3</sub> emissions (0.33 kg N ha<sup>-1</sup> y<sup>-2</sup>) by Liu et al. (2013) was double that in REAS (0.17 kg N ha<sup>-1</sup> y<sup>-2</sup>), implying that the NH<sub>3</sub> increase rate in China is still an open question, and should be further studied.

#### 3.2. Satellite NH<sub>3</sub> and NO<sub>2</sub> over China in the recent decade

#### 3.2.1. Temporal trends

We referred to the method of a previous study (Russell et al., 2012) to conduct the temporal trend analysis by calculating the average values during cold months (October-March) and warm months (April-September) respectively. We herein concentrated more on the temporal analysis of satellite observations during warm months because of the relatively lower uncertainty in comparison with that during cold months. Fig. 3 shows the temporal trend of NO<sub>2</sub> columns during warm and cold months between 2005 and 2015 as well as monthly average values. From satellite observations, the NO<sub>2</sub> columns over China increased with a slope of  $0.063 \times 10^{15}$  molec. cm<sup>-2</sup> y<sup>-1</sup> (4.07% y<sup>-1</sup>) in warm months from 2005 to 2011 and then decreased with a slope of -0.072 molec. cm<sup>-2</sup> in warm months (-3.62% y<sup>-1</sup>) from 2011 to 2015 (Fig. 3 ). The decreasing trends were consistent with NO<sub>x</sub> emissions since 2011 over China (decreasing from 24.04×10<sup>6</sup> ton in 2011 to 20.78 ×10<sup>6</sup> ton in 2014, China Statistical Yearbook, http://www.stats.gov.cn/). During the Chinese 11th Five-Year-Plan (FYP) period (2006-2010), Chinese government undertook a series of strategies to increase energy efficiency and to reduce NO<sub>x</sub> emissions, but NO<sub>x</sub> emissions were not successfully restrained, which created a big challenge for improving air quality over the country (Xia et al., 2016). During the 12th FYP period (2011-2015), more stringent strategies were implemented to control NO<sub>x</sub> emissions, including the application of selective catalytic/non-catalytic reduction (SCR/SNCR) systems in the power sector, staged implementation of tighter vehicle emission standards and a series of standards with aggressive emission limits for power, cement, and the iron and steel industries. These strategies are believed to have helped achieve national targets of NO<sub>x</sub> emission abatement (Xia et al., 2016).

However, the satellite-retrieved NH<sub>3</sub> columns increased with a slope of  $0.118 \times 10^{15}$  molec. cm<sup>-2</sup> y<sup>-1</sup> (2.37% y<sup>-1</sup>) in warm months from 2008 to 2014 (Fig. 3). The percent increase rate for NH<sub>3</sub> by year (2.37% y<sup>-1</sup>) from 2008 to 2014 is lower than that for NO<sub>2</sub> (4.07% y<sup>-1</sup>) from 2005 to 2011, although the absolute NH<sub>3</sub> increase rate of  $0.118 \times 10^{15}$  molec. cm<sup>-2</sup> y<sup>-1</sup> from 2008 to 2014 was higher than absolute NO<sub>2</sub> increase rate of  $0.063 \times 10^{15}$  molec. cm<sup>-2</sup> y<sup>-1</sup> from 2005 to 2011. An increase in NH<sub>3</sub> columns from IASI may be due to decreased NH<sub>3</sub> removal leading to a larger fraction maintaining in gaseous state for a long time rather than changing to the condensed phase. Specifically, NH<sub>3</sub> is considered as an important alkaline gas that is abundant in the atmosphere, and is able to neutralize acidic components including HNO<sub>3</sub> and H<sub>2</sub>SO<sub>4</sub> through the oxidation of NO<sub>x</sub> and SO<sub>2</sub>, respectively (Li et al., 2014;Liu et al., 2017c;Xu et al., 2015). The decreased NH<sub>3</sub> removal to some degree can be

attributed to continuous decreased acidic gases including the NO<sub>2</sub> and SO<sub>2</sub> over China under strong control policy in 12-th FYP, which can largely decrease the fraction of the chemical conversion to (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> and NH<sub>4</sub>NO<sub>3</sub> in the atmosphere. Increasing trend in NH<sub>3</sub> columns may be associated with continuous N fertilizer use for guaranteeing increase of crop productions (Erisman et al., 2008). Although there was no strong NH<sub>3</sub> emission control regulation, N fertilizer efficiency should be further improved over China. In 2015, the Ministry of Agriculture formally announced a "Zero Increase Action Plan" for national fertilizer use by 2020, which requires the annual increase in total fertilizer use will be less than 1% from 2015 to 2019, with no further increment from 2020 (Liu et al., 2015).

If the "Zero Increase Action Plan" for N fertilizer can be effective, future NH<sub>3</sub> emissions should be consistent with the current NH<sub>3</sub> emissions. In addition, due to strong emission control of NO<sub>x</sub>, the NO<sub>x</sub> emissions were believed to decrease significantly from 2011 to 2015. We can reasonably make two major conclusions. First, the atmospheric NO<sub>2</sub>, as a key indicator of oxidized N compounds (NO<sub>2</sub>, HNO<sub>3</sub> and NO<sub>3</sub><sup>-</sup>), decreased since 2011, and will continue to decrease under the current policy. Second, the atmospheric NH<sub>3</sub>, as a key indicator of reduced N (NH<sub>3</sub> and particulate NH<sub>4</sub><sup>+</sup>), will slightly increase or stay at the current level in the future with the "Zero Increase Action Plan". Thus, due to a decreasing trend of oxidized N (NO<sub>x</sub>-N), ammonia N (NH<sub>x</sub>-N) should still dominate Nr deposition (oxidized N plus reduced N) in China, and is expected to play a more significant role in Nr deposition. Therefore, monitoring the reduced N on a regional scale is encouraged to assist in enacting effective measures to protect the environments and public health, with respect to air, soil and water quality.

#### **3.2.2. Spatial pattern**

High NH<sub>3</sub> columns were found in Beijing, Hebei, Henan, Shandong, Hubei and Jiangsu provinces and in Eastern Sichuan province (Fig. 4a), which were consistent with their high NH<sub>3</sub> emissions due to intensive fertilizer application and livestock (Huang et al., 2012). Guangdong, Guangxi, Hunan and Jiangxi provinces also showed high NH<sub>3</sub> columns, due to high volatilization from paddy fields in these regions, with rice being the dominant crop and contributing the most emissions. High NH<sub>3</sub> columns in southern China are in agreement with the high percent paddy farmland area (Fig. S1a) and the high NH<sub>3</sub> columns in northern China are in agreement with the high percent dry farmland area (Fig. S1b). In addition, the NH<sub>3</sub> emissions from vehicles in urban areas could also contribute to the observed high NH<sub>3</sub> columns. For example, in Beijing, the contribution of vehicles equipped with catalytic converters, particularly since the introduction of three-way-catalysts, to non-agricultural NH<sub>3</sub> emissions has recently been considered and might be the most important factor influencing NH<sub>3</sub> concentrations in urban cities (Meng et al., 2011;Xu et al., 2017). In addition, Xinjiang province also emits remarkable NH<sub>3</sub> emissions related to sheep manure management (Huang et al., 2012;Kang et al., 2016;Zhou et al., 2015;Liu et al., 2017a). The lower NH<sub>3</sub> columns are located mostly in the Tibet Plateau area, where there is a minimal amount of arable land and low use of synthetic nitrogenous fertilizers.

NO<sub>2</sub> columns (Fig. 4b) show significantly higher values over vast areas covering North China, East China, and the Sichuan Basin. The NO<sub>2</sub> columns also show high values over the Pearl River Delta, the southern part of Northeast China, and some areas in Northwest China. High NO<sub>2</sub> columns are mostly distributed in populated areas (Fig. S2), where there is a mix of various anthropogenic NO<sub>x</sub> sources, such as vehicles and industrial complexes (Wang et al., 2012;Xu et al., 2015;Meng et al., 2010). It should be noted that an enhanced emission intensity from transportation is confirmed since 2005, even with staged implementation of tightened emission standards for on-road vehicles (Wang et al., 2012). For example, NO<sub>x</sub> emissions from transportation grew to 30% for the whole country in 2014, and the values reached 44%, 55%, and 33% for Beijing, Shanghai, and Guangdong, respectively (Xia et al., 2016). Therefore, transportation is believed to play an increasingly important role in regional  $NO_2$  pollution, especially when emissions from stationary sources are gradually controlled through increased penetration of selective catalytic/non-catalytic reduction (SCR/SNCR) systems.

#### 3.2.3. Limitations of satellite observations

It is difficult to gain whole coverage over China based on the daily data for both IASI NH<sub>3</sub> and OMI NO<sub>2</sub>. For daily NO<sub>2</sub>, the spatial coverage gained by OMI were influenced by cloud radiance fractions, surface albedo, solar zenith angles, row anomaly and so on (Russell et al., 2011;De Smedt et al., 2015). "Row anomaly" issue resulting from the OMI instrumental problem had an impact on approximately half of the rows undergoing unpredictable patterns in cross-track directions relying on latitudes and seasons and prevented obtaining convincing daily product with continuous coverage (Boersma et al., 2011;Boersma et al., 2016). For NH<sub>3</sub>, the satellite instruments were strongly dependent on the meteorological conditions such as cloud fractions or the availability of the temperature profiles (Van Damme et al., 2014b;Boersma et al., 2011), and we cannot retrieve the whole coverage based on daily data over China. It will be beneficial to analyze a very local region with enough numbers of observations, but not appropriate to analyze such large coverage over China.

Facing this big challenge, we used the monthly data for the trend analysis over China. The uncertainty of DOMINO v2.0 NO<sub>2</sub> columns has been well documented in Boersma et al. (2011), and the relative error is reported lower than 20-30% in East Asian by an improved altitude-dependent air mass factor look-up table, a more realistic atmospheric profile, an increased number of reference vertical layers and advanced surface albedos (Boersma et al., 2011). The reader is strongly suggested to refer to Boersma et al. (2011) for more details on the uncertainty analysis.

The potential uncertainty of IASI NH3 columns resulted from IASI observation instruments and

retrieval algorithms. In this paper, the NH<sub>3</sub> datasets were generated based on the recent-updated robust and flexible NH<sub>3</sub> retrieval algorithms, which were designed to overcome some shortcomings of the current algorithms (Whitburn et al., 2016a). The current algorithms were designed firstly to calculate the hyperspectral range index (HRI), a measure for the  $NH_3$  signature strength in the spectrum, and then converted to IASI NH<sub>3</sub> columns by using the thermal contrast (TC) and lookup tables (LUT) of (HRI, TC) pair corresponding to  $NH_3$  columns. The retrieval of HRIs is strongly dependent on the amount of NH<sub>3</sub> and the thermal state of the atmosphere (Whitburn et al., 2016a). The quality of the IASI NH<sub>3</sub> product has been validated by atmospheric chemistry transport models, ground-based and airborne measurements, and NH<sub>3</sub> total columns obtained with ground-based Fourier transform infrared spectroscopy (FTIR). A first validation of the IASI NH<sub>3</sub> using the LOTOS-EUROS model was conducted over Europe, indicating the respective consistency of IASI measurements and model simulations (Van Damme et al., 2014c). A first evaluation of IASI NH<sub>3</sub> dataset using ground-based measurements was made worldwide, presenting consistency with the available ground-based observations and denoting promising results for evaluation by using independent airborne data (Van Damme et al., 2014a). A first validation of of IASI NH<sub>3</sub> dataset using ground-based FTIR derived NH<sub>3</sub> total columns was evaluated, demonstrating a mean relative difference of  $-32.4\pm(56.3)\%$ , a correlation r of 0.8 with a slope of 0.73 (Dammers et al., 2016).

#### 3.3. Atmospheric chemistry transport model NO2 and NH3 columns since 2008

Satellite  $NO_2$  and  $NH_3$  columns were observed at overpass time as an instantaneous point in a day (at 9:30 A.M. for IASI  $NH_3$  and at 1:45 P.M. for OMI  $NO_2$  local time). These instantaneous satellite observations may not be representative for the temporal trend analysis over China. We further retrieved the monthly variations of  $NO_2$  and  $NH_3$  columns since 2008 from MOZART varying 6 hours every day

(00, 06, 12, 18 h). We compared the temporal trend analysis of  $NO_2$  from MOZART at 12 h with that gained from satellite at the overpass time (OMI 1:45 P.M. local time) as well as for  $NH_3$ .

Fig. 5 shows the NO<sub>2</sub> columns at 12:00 during warm and cold months between 2008 and 2015 from MOZART. The percent increase rate for NO<sub>2</sub> columns at 12:00 during warm months (April-September) between 2008 and 2011 was 4.02% y<sup>-1</sup> (Fig. 5), which was comparable with that (4.23% y<sup>-1</sup>) derived from OMI (Fig. 3). During 2011-2015, we found a slightly lower decrease rate (-2.93% y<sup>-1</sup>) in NO<sub>2</sub> columns during warm months at 12:00 from MOZART (Fig. 5) than that (-3.62% y<sup>-1</sup>) gained from OMI at 13:45 (Fig. 3). The temporal variations of NO<sub>2</sub> columns at 12:00 from MOZART were generally in accord with those from OMI at 13:45 P.M. local time. Fig. 5 also demonstrates the average NO<sub>2</sub> columns (averaged at 00, 06, 12 and 18 h) during warm and cold months between 2008 and 2015. We found a close increase rate at 12:00 (4.02%) with that averaged at 00, 06, 12 and 18 h (4.23%) before 2011, as well as a similar decrease rate at 12:00 (-2.93%) and the average (-3.07%), implying that the temporal trend analysis at 12:00 vs. that averaged at 00, 06, 12 and 18 h can be considered mostly consistent over China from MOZART.

For NH<sub>3</sub>, we found the percent increase rate at 12:00 during warm months between 2008 and 2015 was 1.30% y<sup>-1</sup> from MOZART (Fig. 5), which was lower than that (2.37% y<sup>-1</sup>) from IASI during 2008-2014. The percent increase rate by daily average (at 00, 06, 12 and 18 h) during warm months between 2008 and 2015 was 1.36% y<sup>-1</sup> from MOZART (Fig. 5). In MOZART-4, the alkaline gaseous NH<sub>3</sub> and the acidic gaseous NO<sub>2</sub> (the precursor for HNO<sub>3</sub>) and SO<sub>2</sub> are very important precursors for bulk NH<sub>4</sub>NO<sub>3</sub> and (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> particles, which form the primary system of gas-particle partitioning (NH<sub>3</sub>-NH<sub>4</sub><sup>+</sup>-NO<sub>x</sub>-NO<sub>3</sub><sup>-</sup>-SO<sub>2</sub>-SO<sub>4</sub><sup>2-</sup>). The chemical shifts between particulate NH<sub>4</sub>NO<sub>3</sub> and gaseous NH<sub>3</sub> and NO<sub>x</sub> are correlated with the abundance of NH<sub>3</sub> and NO<sub>x</sub> and meteorological factors. The decreased

abundance of  $NO_x$  between 2011 and 2015 may also contribute to an increase in the  $NH_3$  abundance in the gas stage resulting from decreased conversion to particulate  $NH_4NO_3$ .

# 3.4. Implications for estimating long-term Nr deposition datasets and recommendations for future work

We found both the  $NO_x$  and  $NH_3$  over China increased continuously from 1980 to 2010 based on emissions data from REAS and EDGAR. In recent years, based on satellite observations, we found an increase of 2.37% y<sup>-1</sup> in NH<sub>3</sub> columns during 2008-2014. We also found high-level NO<sub>2</sub> columns over China from 2005-2011 (4.07%  $y^{-1}$ ) but a decrease from 2011 to 2015 (-3.62%  $y^{-1}$ ). Despite the decline, the NO<sub>2</sub> columns during 2011-2015 were still in high level with an average of  $1.87 \times 10^{15}$  molec. cm<sup>-2</sup>  $y^{-1}$  compared with that  $(1.65 \times 10^{15} \text{ molec. cm}^2 y^{-1})$  during 2005-2010. Notably, these emissions certainly lead to the deposition of atmospheric Nr in form of dry and wet processes into aquatic ecosystems and terrestrial, with implications affecting ecosystem and human health, biological diversity and greenhouse gas balances (Lu et al., 2016). Hence, it is very crucial to estimate Nr deposition with high spatiotemporal resolutions in order to drive ecological models such as the Denitrification-Decomposition (DNDC) model and Integrated BIosphere Simulator (IBIS), to assess its impact on soil, forest, water and greenhouse balance. Here, we call for a long-term dataset of Nr depositions both regionally and globally to investigate how the N emissions affect the environment. Challenge still exits in estimating both the dry (NO<sub>2</sub>, HNO<sub>3</sub> particulate NO<sub>3</sub><sup>-</sup>, NH<sub>3</sub> and particulate NH<sub>4</sub><sup>+</sup>) and wet ( $NH_4^+$  and  $NO_3^-$  in precipitation) depositions for a long-term dataset such as since 1980 or earlier possibly due to the complex scheme of N transformations and transportation or limited available data both from emissions, satellites and a limited number of ground measurements.

Satellite observations provide a new perspective of estimating Nr depositions regionally, and have been

used to improve the estimation performance. For example, to improve the modeling performance in dry gaseous NO<sub>2</sub> depositions from GEOS-Chem (Goddard Earth Observing System chemical transport model), Nowlan et al. (2014) applied the OMI NO<sub>2</sub> columns to calibrate the simulated ground NO<sub>2</sub> concentrations, and then estimated the deposition between 2005 and 2007. Our previous work focusing on the dry particulate NO<sub>3</sub><sup>--</sup> deposition over China was also based on the OMI NO<sub>2</sub> columns, MOZART simulations and monitored-based sources (Liu et al., 2017b). Geddes et al. (2017) used the satellite NO<sub>2</sub> columns from GOME, GOME-2 and SCIAMACHY instruments to calibrate the NO<sub>x</sub> emissions in GEOS-Chem to estimate the NO<sub>x</sub> depositions since 1996. The simulations combining the satellite measurements and CTM models to derive Nr depositions (Geddes and Martin, 2017;Nowlan et al., 2014) in recent years will provide relatively accurate datasets (certainly need to be validated and modified by ground measurements).

Despite progress in satellite techniques in recent decades (for NO<sub>2</sub> since 1997 by GOME and for NH<sub>3</sub> since 2008 by IASI), we can hardly tracked studies concerning Nr depositions before 1997 based on satellite observations. Thus, with the help of emissions data such as REAS and EDGAR, we can derive long-term Nr depositions, especially before 1997. Long-term emissions data such as REAS and EDGAR will provide valuable dataset to expand the modeling Nr depositions in recent years. In order to derive the Nr depositions from the emission data, CTMs are frequently used through modeling the wet (simplified as the product of scavenging efficiency and precipitation amount) and dry process (simplified as the inferential method by multiplying the deposition velocity and gaseous or particulate concentrations). However, we still lack a comprehensive dataset of gridded long-term Nr depositions including both the dry (NO<sub>2</sub>, HNO<sub>3</sub> particulate NO<sub>3</sub><sup>-</sup>, NH<sub>3</sub> and particulate NH<sub>4</sub><sup>+</sup>) and wet (NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> in precipitation) processes over China, which will be addressed in future work.

Another gap is that, all the above mentioned studies focused on the NO<sub>x</sub> depositions and did not derive the NH<sub>y</sub> (NH<sub>3</sub> and NH<sub>4</sub><sup>+</sup>) depositions over China. Our recent work (Liu et al., 2017a) using IASI NH<sub>3</sub> columns combining the vertical profiles from MOZART benefits our understanding of the ground NH<sub>3</sub> concentrations over China, and the satellite-derived ground NH<sub>3</sub> concentrations were generally in accord with the national measurements from NNDMN. To date, there are still no reports of using the satellite NH<sub>3</sub> columns to derive the temporal and regional NH<sub>y</sub> depositions over China, which dominated the total Nr depositions (NO<sub>x</sub> plus NH<sub>y</sub>) (Liu et al., 2016b;Liu et al., 2013). The gaps of modeling NH<sub>y</sub> depositions by applying the satellite observations combining the CTMs simulations require more efforts and further research.

#### 4. Conclusion

Atmospheric ammonia (NH<sub>3</sub>) and nitrogen dioxide (NO<sub>2</sub>) play an important role in determining air quality, environmental degradation and climate change. The emission data, satellite observations and atmospheric transport modeling have great potential for understanding the temporal variations of atmospheric NH<sub>3</sub> and NO<sub>2</sub> on a regional scale, with high spatial and temporal resolutions. This study analyzed the characteristics of atmospheric NH<sub>3</sub> and NO<sub>2</sub> over China since 1980 based on the multiple datasets. The major findings were as follows:

1. Based on emission data, both significant continuous increasing trend of  $NH_3$  and  $NO_x$  were observed from REAS (for  $NH_3$  0.17 kg N ha<sup>-1</sup> y<sup>-2</sup> and for  $NO_x$  0.16 kg N ha<sup>-1</sup> y<sup>-2</sup>) and EDGAR (for  $NH_3$  0.24 kg N ha<sup>-1</sup> y<sup>-2</sup> and for  $NO_x$  0.17 kg N ha<sup>-1</sup> y<sup>-2</sup>) over China during 1980-2010.

2. Based on the satellite observations, we found high-level  $NH_3$  columns with the percent increase rate of 2.37% y<sup>-1</sup> from 2008 to 2014. For NO<sub>2</sub>, we found continuous high-level NO<sub>2</sub> columns over China from 2005-2011 but a decrease from 2011 to 2015 (still in high level). The decrease of NO<sub>2</sub> columns may result from more stringent strategies taken to control  $NO_x$  emissions during the 12th Five-Year-Plan, including successful application of SCR/SNCR systems in the power sector, tighter emission standards on vehicles and a series of standards with aggressive emission limits. Increasing trend of NH<sub>3</sub> columns may be due to continuous N fertilizer use for guaranteeing continuous increase of the crop productions. An increase in NH<sub>3</sub> columns may be due to decreased NH<sub>3</sub> removal leading to a larger fraction maintaining in gaseous state for a long time rather than changing to the condensed phase, which may be related with continuous decreased acidic gases including the NO<sub>2</sub> and SO<sub>2</sub> over China under strong control policy in 12-th FYP.

3. Based on MOZART simulations, the temporal variations of NO<sub>2</sub> columns at 12:00 from MOZART were generally in accord with those from OMI at 13:45 P.M. local time. We also found a close increase rate at 12:00 (4.02%) with that averaged at 00, 06, 12 and 18 h (4.23%) before 2011, as well as a similar decrease rate at 12:00 (-2.93%) and the average (-3.07%). For NH<sub>3</sub>, we found a lower percent increase rate from MOZART (1.30% y<sup>-1</sup>) than IASI (2.37% y<sup>-1</sup>) between 2008 and 2014.

4. The multiple datasets used in the current work have implications for estimating long-term Nr deposition datasets. The simulations combining the satellite measurements and CTM models to derive Nr depositions will provide relatively accurate datasets, and the REAS and EDGAR emissions have potential to expand the modeling Nr depositions to long-term datasets. In particular, modeling NH<sub>y</sub> depositions by applying the satellite observations combining the CTMs simulations require more efforts and further research.

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Figures



Fig. 1. The satellite-derived observation numbers for  $NO_2$  and  $NH_3$ . (a) denotes the percentages of observations in each month in 2010 for  $NO_2$  and in 2015 for  $NH_3$  and (b) represents the total observation numbers for  $NO_2$  and  $NH_3$  over China. Notably, the  $NO_2$  observation numbers were gained from DOMINO products with a cloud radiance fraction below 0.5, while the IASI observations with a relative error below 100% or an absolute error below  $5 \times 10^{15}$  molec. cm<sup>-2</sup> were processed for analysis over China.



Fig. 2. The NO<sub>2</sub> and NH<sub>3</sub> emissions over China. (a) denotes the NO<sub>2</sub> and NH<sub>3</sub> emissions over China from 1980 to 2010 from REAS, (b) represents the NO<sub>2</sub> and NH<sub>3</sub> emissions over China from 1980 to 2010 from EDGAR, (c) demonstrates the relationship of NO<sub>2</sub> emissions over China from REAS and EDGAR and (d) shows the relationship of NH<sub>3</sub> emissions over China from REAS and EDGAR and EDGAR.



**Fig. 3.** Time series of average OMI NO<sub>2</sub> and IASI NH<sub>3</sub> columns over China during warm months (April-September) and cold months (October-March). The time period of NO<sub>2</sub> columns was from 2005 to 2015, while the timespan of NH<sub>3</sub> columns was from 2008 to 2015 over China. The associated mean error for each period is presented here as error bars. The percent increase or decrease rate (%) was the long term mean calculated by  $100 \times (\frac{Y_2 - Y_1}{Y_1} + \frac{Y_3 - Y_2}{Y_2} + \dots + \frac{Y_{n+1} - Y_n}{Y_n}) \times \frac{1}{n}$ .



**Fig. 4.** Spatial distribution of the annual NH<sub>3</sub> (a) and NO<sub>2</sub> (b) columns (molecules cm<sup>-2</sup> year<sup>-1</sup>). The successfully full provincial names are Beijing (BJ), Tianjin (TJ), Hebei (HeB), Shandong (SD), Shanxi (SX), Henan (HeN), Shaanxi (SaX), Liaoning (LN), Jilin (JL), Heilongjiang (HLJ), Neimenggu (NMG), Gansu (GS), Ningxia (NX), Xinjiang (XJ), Shanghai (SH), Jiangsu (JS), Zhejiang (ZJ), Anhui (AH), Hubei (HuB), Hunan (HuN), Jiangxi (JX), Fujian (FJ), Guangdong (GD), Hainan (HaN), Yunnan (YN), Guizhou (GZ), Chongqing (CQ), Sichuan (SC), Guangxi (GX), Xizang (XZ) and Qinghai (QH).

(a) MOZART NO<sub>2</sub>



Fig. 5. Time series of MOZART  $NO_2$  and  $NH_3$  columns over China during average warm months (April-September) and cold months (October-March) from 2008 to 2015. The mean columns were calculated by averaging the columns at 00, 6, 12 and 18 h. The associated mean error for each period is presented here as error bars.