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## Worldwide spatiotemporal atmospheric ammonia (NH<sub>3</sub>) columns variability revealed by satellite

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## RESEARCH LETTER

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## Key Points:

- Six years of NH<sub>3</sub> morning and evening IASI measurements are analyzed
- Seasonal cycles of atmospheric NH<sub>3</sub> are characterized for subcontinental areas
- Source processes are attributed from a climatology of the month of NH<sub>3</sub> maximum

## Supporting Information:

- Figures S1–S3

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Worldwide spatiotemporal atmospheric ammonia (NH<sub>3</sub>) columns variability revealed by satellite

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**Abstract** We exploit 6 years of measurements from the Infrared Atmospheric Sounding Interferometer (IASI)/MetOp-A instrument to identify seasonal patterns and interannual variability of atmospheric NH<sub>3</sub>. This is achieved by analyzing the time evolution of the monthly mean NH<sub>3</sub> columns in 12 subcontinental areas around the world, simultaneously considering measurements from IASI morning and evening overpasses. For most regions, IASI has a sufficient sensitivity throughout the years to capture the seasonal patterns of NH<sub>3</sub> columns, and we show that each region is characterized by a well-marked and distinctive cycle, with maxima mainly related to underlying emission processes. The largest column abundances and seasonal amplitudes throughout the years are found in southwestern Asia, with maxima twice as large as what is observed in southeastern China. The relation between emission sources and retrieved NH<sub>3</sub> columns is emphasized at a smaller regional scale by inferring a climatology of the month of maximum columns.

## 1. Introduction

Ammonia (NH<sub>3</sub>) is mainly emitted over continents by agricultural activities and biomass burning [Emission Database for Global Atmospheric Research, 2014]. As the primary form of reactive nitrogen (Nr) in the environment [Sutton *et al.*, 2013], it is a key component for the nitrogen cycle and several biogeochemical processes. Increased atmospheric NH<sub>3</sub> emissions from anthropogenic sources and subsequent Nr deposition are responsible for acidification of terrestrial ecosystems and loss of biodiversity [Erisman *et al.*, 2007, 2013; Bobbink *et al.*, 2010]. It can also cause algae blooms and drive eutrophication of water bodies [Paerl *et al.*, 2014]. NH<sub>3</sub>, being the major basic compound in the atmosphere, plays a significant role in the formation of secondary aerosols, which affect human and ecosystem health [Pope *et al.*, 2009; Erisman *et al.*, 2013; Paulot and Jacob, 2014]. This is especially the case when suburban NH<sub>3</sub> emissions occur close to sources of urban oxidized compounds (such as nitrogen oxides (NO<sub>x</sub>) and sulfur dioxide (SO<sub>2</sub>)), allowing the pollutants to react rapidly and causing air quality deterioration [Gu *et al.*, 2014].

NH<sub>3</sub> sources, sinks, and transport at various scales are still poorly known despite these important environmental issues. This is due to the fact that NH<sub>3</sub> emissions in the atmosphere are highly dependent on agricultural practices and environmental conditions and that up to recently there were insufficient monitoring means to capture their large variability [Sutton *et al.*, 2007]. Indeed, while some parts of the world are equipped with in situ measuring systems, others such as tropical agroecosystems and a large part of the Southern Hemisphere are completely unmonitored [Bouwman *et al.*, 2002; Van Damme *et al.*, 2015]. Furthermore, the representativity of point measurements for larger areas (e.g., regional/global model cell size) is also problematic [Wichink Kruit *et al.*, 2012]. These limitations of the global NH<sub>3</sub> monitoring network in place prevent, on the one hand, the accurate determination of emission magnitudes and distributions, which is needed to define and control national-scale policy requirements [Sutton *et al.*, 2013], and, on the other hand, accurate air quality model predictions [Gilliland *et al.*, 2003, 2006; Heald *et al.*, 2012].

Satellite instruments have shown their abilities to probe atmospheric NH<sub>3</sub> and fill this observational gap by providing daily global distributions, albeit at a medium spatial resolution [Clarisse *et al.*, 2009, 2010; Pinder *et al.*, 2011; Shephard *et al.*, 2011; Van Damme *et al.*, 2014a]. First analyses of the NH<sub>3</sub> temporal evolution were made with bidaily measurements from the Infrared Atmospheric Sounding Interferometer (IASI) over

California [Clarisse *et al.*, 2010], while measurements from the Tropospheric Emission Spectrometer (TES) were used for a preliminary study of the global seasonality, looking at composite means (to compensate for the limited coverage of TES) of January, April, July, and October [Shephard *et al.*, 2011]. There are now multiyear data sets of global NH<sub>3</sub> columns available from satellite instruments and in particular from IASI, which can be used to determine the spatial and temporal patterns of NH<sub>3</sub>, as was recently done for Europe [Van Damme *et al.*, 2014b].

In this paper we investigate the spatiotemporal variability of NH<sub>3</sub> over the entire globe using six full years of bidaily retrieved columns. We examine the temporal evolution on the basis of global composite seasonal distributions and provide a more detailed analysis of seasonality for several subcontinental areas using monthly means. We finally derive for the first time a climatology of the month of maximum columns on the global scale, discussed in first order (i.e., neglecting chemistry, transport, and deposition) on the basis of underlying local emission processes.

## 2. IASI-NH<sub>3</sub> Observations

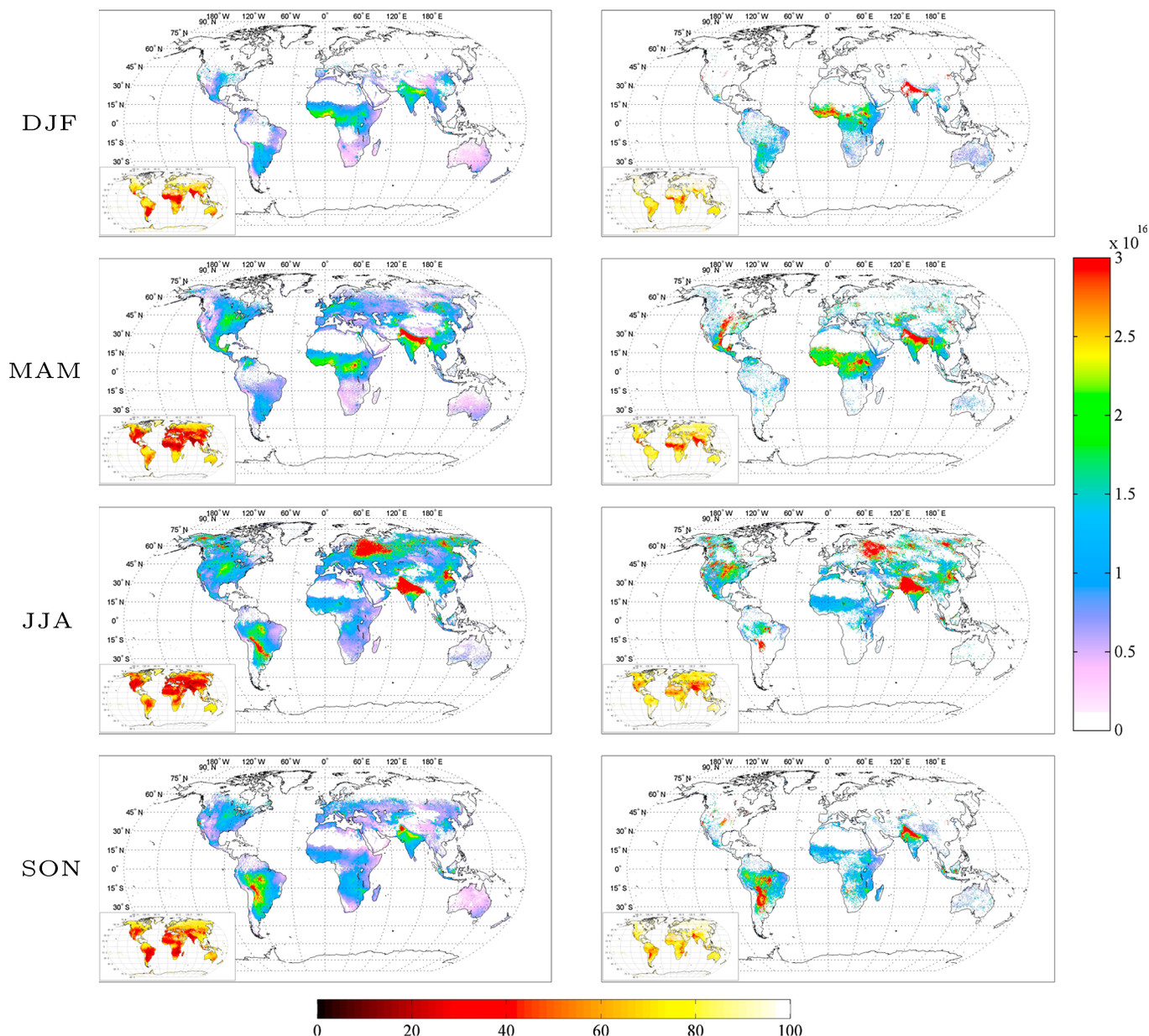
IASI is a polar Sun-synchronous infrared sounder on board the MetOp satellites. It covers the entire globe twice a day, crossing the equator at a mean solar local time of 9:30 A.M. and P.M. (hereafter referred to as the morning and evening overpasses) and using a scanning mode around the nadir [Clerbaux *et al.*, 2009]. A detailed description of the algorithm used to retrieve NH<sub>3</sub> is presented in Van Damme *et al.* [2014a], along with first global distributions and hemispheric time series from the IASI morning observations. In short, the algorithm consists of calculating a dimensionless spectral index from the IASI observations with a cloud fraction below 25%. This index is then converted into total columns (molecules/cm<sup>2</sup>) through look-up tables built from radiative transfer simulations performed under various atmospheric conditions. This method takes into account the dependency of the IASI sensitivity to NH<sub>3</sub> on the thermal contrast, which is provided for each IASI measurement by the IASI Level2 Product Processing Facility [August *et al.*, 2012]. It is worth noting that changes in Level2 version used in the retrieval procedure occurred in September 2010 [see Van Damme *et al.*, 2014b]. The NH<sub>3</sub> retrieved total columns are provided with an associated error, driven by the thermal contrast and the amount of NH<sub>3</sub>. The resulting NH<sub>3</sub> data set of columns has been evaluated against model simulations over Europe [Van Damme *et al.*, 2014b] and correlative ground-based and airborne measurements worldwide [Van Damme *et al.*, 2015]. In this study we consider for the first time NH<sub>3</sub> retrievals over land at a subcontinental scale, from both morning and evening overpasses of IASI/MetOp-A. A weighting procedure based on the relative error has been applied to average spatially and temporally IASI observations (columns and associated errors), as in Van Damme *et al.* [2014a, 2014b, 2015].

## 3. Spatiotemporal Variability

### 3.1. Seasonality and Monthly Variability

Composite seasonal distributions of NH<sub>3</sub> columns are shown separately for IASI morning (left column) and evening (right column) overpasses in Figure 1. The data correspond to weighted mean columns of the measurements performed during each season from 2008 to 2013 (0.25° × 0.5° cells); the resulting columns associated with a relative error above 75% have been filtered out, and the error distributions (not filtered) are shown as inset. These seasons are represented as 3 month periods: December-January-February (DJF, Figure 1, first row), March-April-May (MAM, second row), June-July-August (JJA, third row) and September-October-November (SON, fourth row). In Figure 2, we show the time evolution over the six years over six large areas, similar to the subcontinental boxes used by Shephard *et al.* [2011] (the six other regions are presented in the supporting information). Time series are shown separately for morning (red) and evening (blue) retrievals.

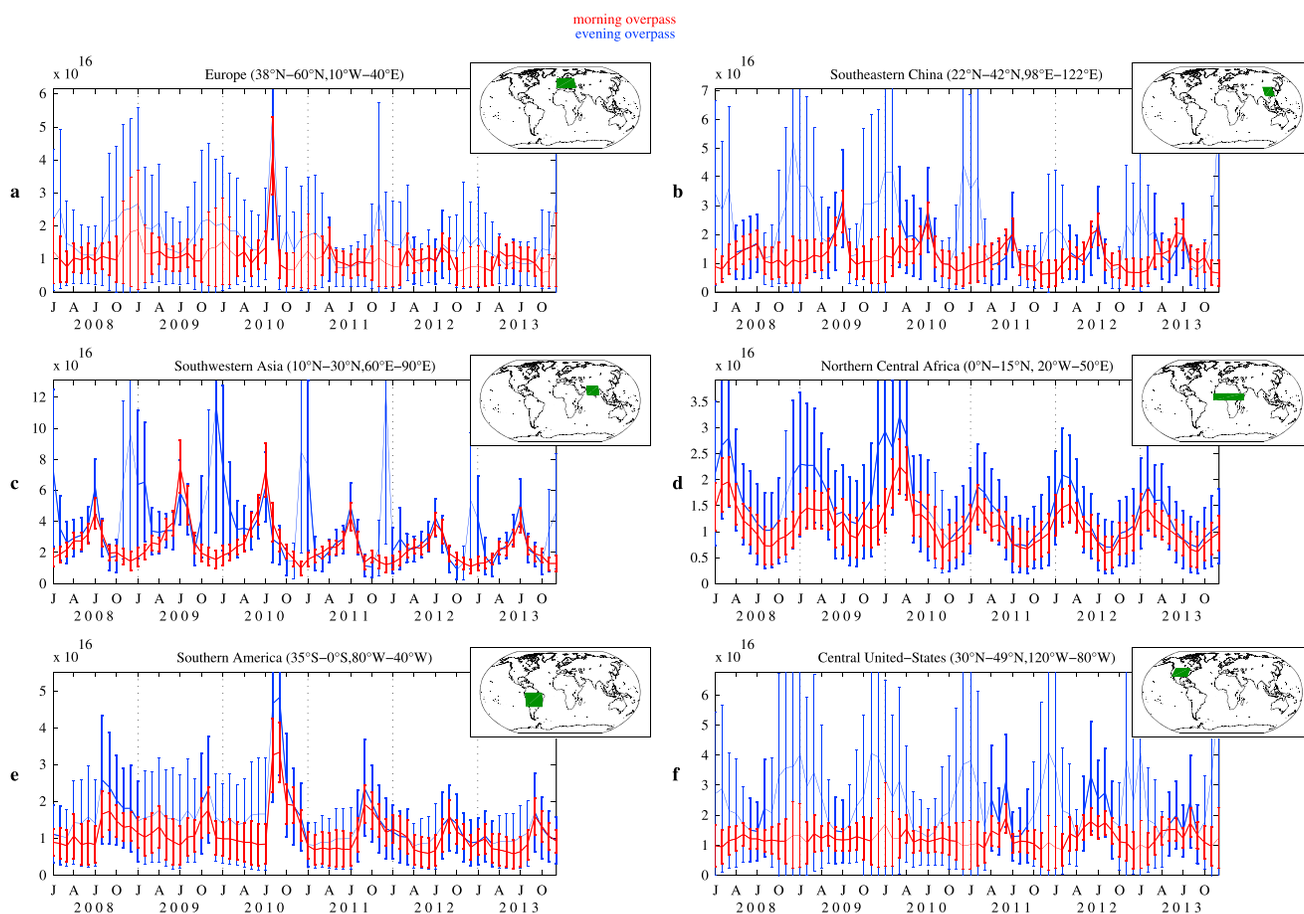
The seasonal distributions in Figure 1 together with the error maps highlight the varying sensitivity of IASI measurements for each region which makes the analyses of time series complex. It can be seen that the sensitivity is generally lower during the night (resulting in much larger error), due to less favorable thermal contrast, but that the main patterns are similar. On some occasions the measurements from the evening orbit concur in terms of sensitivity with those of the morning orbit (especially in case of temperature inversion [Clarisse *et al.*, 2010]), but this is not reflected in the seasonally aggregated distributions. Figure 2 shows further that IASI observes a clear seasonality, which is different from one area to another. It is very important to consider the error bars to account for the variation in sensitivity as a function of month (and between orbits); only the



**Figure 1.** Composite seasonal distributions (molecules/cm<sup>2</sup>, vertical color bar) averaged over 6 years (1 January 2008 to 31 December 2013) of IASI-NH<sub>3</sub> (left column) morning and (right column) evening measurements in 0.25° × 0.5° cells. (first to fourth rows) December-January-February (DJF), March-April-May (MAM), June-July-August (JJA), and September-October-November (SON) weighted mean distributions. A postfiltering consisting in excluding mean column values associated with a mean relative error above 75% has been applied. Relative weighted error distributions (% not filtered, horizontal color bar) are presented as an inset in the bottom left corner of each seasonal map.

main patterns, identified in Figure 2 by thicker red and blue line (when the error is below 75%), are therefore discussed here. The retrieved columns from both overpasses follow in most cases a similar evolution, with overall higher evening columns associated with larger errors. Northern central Africa (Figure 2d) shows in particular a seasonal cycle fully captured by IASI daytime and nighttime measurements, with the latest columns being consistently larger than the daytime ones. This could be explained by the building-up of NH<sub>3</sub> following emission during the day. Finally, in some regions, there are pronounced maxima in the evening measurements, which are absent in the morning ones. This is especially the case of southeastern China (Figure 2b) and southwestern Asia (Figure 2c), with the evening maxima observed in wintertime (see below).

In Europe (Figure 2a), the highest concentrations are measured in spring and summer (MAM and JJA, Figure 1) for the morning observations with maxima around 1.5 × 10<sup>16</sup> molecule/cm<sup>2</sup>. The stronger peak in August 2010,



**Figure 2.** Monthly weighted time series of the  $\text{NH}_3$  total columns ( $\text{molecules}/\text{cm}^2$ ) for the morning (red) and evening (blue) IASI overpasses covering the entire period from 1 January 2008 to 31 December 2013. Bold lines and error bars correspond to monthly means associated with a relative error below 75%. Each January is marked with a dashed vertical line, while the x axis labels are January (J), April (A), July (J), and October (O); the different years are also indicated on the time axis. The subcontinental areas selected are similar to the ones from Shephard et al. [2011] and are indicated by green patches in the upper right corner inset map.

also clearly seen in the JJA distributions, is linked with the exceptional fires that occurred around Moscow that summer [Konovalov et al., 2011]. It emitted large amounts of  $\text{NH}_3$ , which were transported to Central Europe [R'Honi et al., 2013; Van Damme et al., 2015].

IASI measures large column values over several areas in Asia, with a maxima generally in July (well matching for morning and evening overpasses). These maxima are about twice as large for southwestern Asia (Figure 2c, around  $4 \times 10^{16}$   $\text{molecules}/\text{cm}^2$ ) as compared to southeastern China (Figure 2b, around  $2 \times 10^{16}$   $\text{molecules}/\text{cm}^2$ ). The amplitudes of the cycle for southwestern Asia are also the largest of all regions investigated, reaching  $6.3 \times 10^{16}$   $\text{molecules}/\text{cm}^2$  between the minimum (December) and the maximum (July) in 2010. For southeastern China, additional high monthly values ( $2 \times 10^{16}$   $\text{molecules}/\text{cm}^2$ ) are observed in June 2012 and 2013. These summer peaks are consistent with several in situ data sets [Ianniello et al., 2010; Meng et al., 2011; Shen et al., 2011]. Interestingly, for these two regions in Asia the measurements from the evening orbit are for many months also associated with low errors. In winter, very high values are measured (up to  $8 - 10 \times 10^{16}$   $\text{molecules}/\text{cm}^2$  in 2010 and 2011) possibly related to stable atmospheric conditions preventing the dispersion of pollutants [Boynard et al., 2014]. Southwestern Asia is an area where IASI is able to probe  $\text{NH}_3$  throughout the year with relatively low error, with persistent highest columns in July, from  $3.8 \times 10^{16}$  in 2012 up to  $7.5 \times 10^{16}$   $\text{molecules}/\text{cm}^2$  in 2009 during daytime and from  $4.2 \times 10^{16}$  in 2012 up to  $6.2 \times 10^{16}$   $\text{molecules}/\text{cm}^2$  in 2010 during nighttime, with additional secondary peaks in December and/or January, mainly in the evening (especially in December 2009:  $1.16 \times 10^{17}$   $\text{molecules}/\text{cm}^2$ ). Note that from ground-based measurements, the summer extrema for this part of the world have been reported around Delhi for 2011 by Singh and Kulshrestha [2012] and that high winter values have been also observed [Sharma et al., 2010].

In the central part of the African continent, high columns are retrieved in DJF and MAM for both morning and evening overpasses. The northern central Africa region (Figure 2d) presents maxima in the range  $1.5\text{--}2.5\times 10^{16}$  molecules/cm<sup>2</sup> (slightly higher in the evening) in February. The entire cycle is captured and is consistent with that observed by the IGAC/DEBITS/AFRICA (IDAF) ground-based network [Adon *et al.*, 2010]. The southern part of this continent is characterized by lower columns peaking later in the year, in SON. In southern America (Figure 2e), the entire cycle is captured using the retrieved columns from the morning overpass, while the nighttime measurements provide complementary information from August to January. The seasonality is characterized by enhanced NH<sub>3</sub> columns in JJA and SON with a persistent peak in September, however, generally below  $2\times 10^{16}$  molecules/cm<sup>2</sup>. Higher columns over the 6 year period are reported in August and September 2010, which is a year of exceptional burning [Chen *et al.*, 2013].

As for Europe, the sensitivity of IASI to NH<sub>3</sub> in the North American continent is not always sufficient and the total cycle cannot be perfectly characterized. From the morning-retrieved columns, we see that the values in Central U.S. (Figure 2f) are larger in MAM and JJA. This is similar in Europe, but we observe that the U.S. summer values are closer to the spring values than in Europe. These observed temporal patterns are consistent with the ones reported by Pouliot *et al.* [2012] based on emission inventories. The minima (around  $1\times 10^{16}$  molecule/cm<sup>2</sup>) and maxima (around  $1.5\times 10^{16}$  molecule/cm<sup>2</sup>) are also pretty similar for these two regions, possibly reflecting similarities in agricultural practices.

In comparison with the four (January, April, July, and October) monthly composite means provided by TES measurements [Shephard *et al.*, 2011], we have shown with the discussion above that IASI is able to provide quantitative monthly resolved seasonal patterns, fluctuating each year depending on environmental conditions. The general patterns of seasonality inferred from IASI are broadly consistent with those reported by TES. Similar summer peaks are indeed observed for the Asian subcontinental areas, as well as high values during SON period in South America, and large NH<sub>3</sub> columns from December to April in North Central Africa.

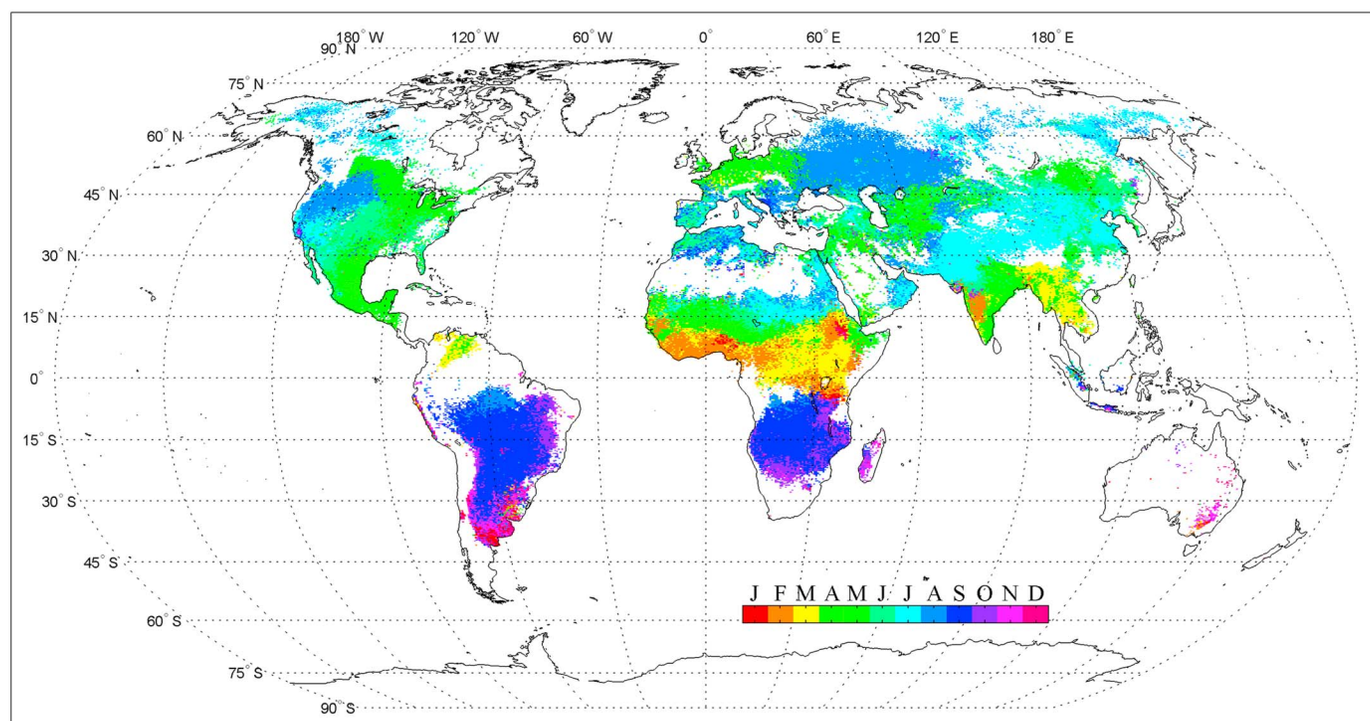
### 3.2. Interannual Variability

The yearly global distributions from 2008 to 2013 are presented in the supporting information (Figure S1). These distributions are weighted means of all the measurements (morning and evening overpasses) per year included in each  $0.25^\circ$  latitude  $\times$   $0.5^\circ$  longitude cell. The year 2010 stands out from the other years due to the large biomass burning events which occurred in Russia and in southern America. Large agricultural areas such as the North China Plain, the Indo-Gangetic Plain, and the Great Plains in U.S. and Canada, exhibit high columns each year. Other persistent high signals are observed by IASI above biomass burning areas in West Africa, South America, and the boreal regions. A significant decrease of the NH<sub>3</sub> columns over West Africa is observed in the years 2011–2013 (maximum column around  $1.5\text{--}2\times 10^{16}$  molecules/cm<sup>2</sup>), in comparison to the years 2008–2010 (maximum column well above  $2\times 10^{16}$  molecules/cm<sup>2</sup>). This could be due to the strong (2010–2011), moderate (2011–2012), and weak (2013) La Niña episodes [National Oceanic and Atmospheric Administration, 2014], which cool this part of Africa, implying reduced NH<sub>3</sub> volatilization [Sutton *et al.*, 2013]. An additional possible reason lies in the decrease of the savanna burning due to their conversion into cropland [Andela and van der Werf, 2014]. Another decrease easily identifiable from the yearly global distributions occurs in North India, especially in the region including the Hindus and the Ganges Valleys. While characterizing the trends (see Figure S2 in the supporting information), we found statistically significant values only for the African regions and Australia. However, a longer time period of IASI-NH<sub>3</sub> observations and consistent Level2 data are needed to confirm these first trend estimates.

### 3.3. Climatology of the Month of Maximum Columns

To explore the global variability in the timing of NH<sub>3</sub> maxima at a higher spatial resolution, a climatology map of the monthly highest columns in  $0.25^\circ \times 0.5^\circ$  cells is presented in Figure 3, following an approach used earlier by van der A *et al.* [2008] for nitrogen dioxide (NO<sub>2</sub>). Monthly weighted means for each year based on all morning and evening columns over the 6 year period have been used to draw this map. The composite monthly means for the six years have been subsequently calculated to determine the month with the highest column value. Due to their associated lower error estimates, the daytime observations are driving the resulting distribution. The error on the timing of the month of maximum columns has been estimated at 1 month by comparing the second and third maximum to the maximum (see supporting information Figure S3).

Figure 3 highlights the timing diversity in NH<sub>3</sub> column maxima, which, if we neglect in first order the impact on the NH<sub>3</sub> columns of changing chemistry with season (e.g., enhanced nitrate aerosol formation with increased



**Figure 3.** Month of maximum  $\text{NH}_3$  column determined in  $0.25^\circ \times 0.5^\circ$  cells from  $\text{NH}_3$  measurements performed between 1 January 2008 and 31 December 2013. Monthly weighted columns associated with a weighted relative error above 35% have been set to zero.

emissions of  $\text{NO}_x$  and  $\text{SO}_2$ ), can be traced back to changes in emissions. It shows that a major part of northwestern Europe has an April maximum, consistent with agricultural temporal emission patterns for this area, which have a well-known associated spring peak due to fertilizer application [Friedrich and Reis, 2004; Paulot et al., 2014], and the fact that agriculture represents around 90% of  $\text{NH}_3$  reported emissions in Europe [European Environment Agency, 2014]. In contrast, southern Europe is characterized by June–July maxima and eastern Europe by August maxima. Those summer southern European emission peaks are likely due to the enhanced  $\text{NH}_3$  volatilization with increasing temperature [Sutton et al., 2013]. It is interesting to point out a well-marked autumn peak in Croatia and the Balkans, which could be related to vegetation fires. The August maxima for eastern Europe and part of Asia are highly consistent with the cycle of fire activity [Giglio et al., 2006], suggesting biomass burning as a major source of  $\text{NH}_3$  in these areas. These fires are generally caused by agricultural burning, whereas the wildfires in boreal areas are responsible for the July peaks [Korontzi et al., 2006; Magi et al., 2012].

In the northern part of southeastern Asia, IASI observes  $\text{NH}_3$  column maxima in March and April (light green and yellow in Figure 3), while for the Indian subcontinent the peak month varies from February (southwestern India) to July (Pakistan and northeastern India). The detailed interpretation of these varying seasonal patterns on a relatively small continental scale is not straightforward considering the numerous emission processes simultaneously at work (e.g., fertilization, livestock, industry, agricultural, and domestic fires). For instance, the Indus basin is well known for its intensive agriculture based on large-scale irrigated crops [Land Degradation Assessment in Drylands, 2013], and in that region fires do not seem to be responsible for the observed peak in July. In contrast, the  $\text{NH}_3$  maxima in southeastern Asia are well corresponding with the seasonality of fires, which indicates that biomass burning is contributing significantly to the  $\text{NH}_3$  columns [Giglio et al., 2006]—except for Thailand where the peak in March suggests the dominance of agricultural sources. The June–July peaks in the northern part and in particular in the North China Plain are likely linked with harvest season associated with the intense burning of agricultural crop residues [Huang et al., 2012a]. The Sichuan province, characterized by high livestock densities [Huang et al., 2012b], has its highest values sooner in the year, in May (green).

Africa presents a high diversity in the months of  $\text{NH}_3$  column maxima. The southern African continent is mainly driven by fire emissions, which are known to peak from September to November [Giglio et al., 2006].

The central and northern parts are much more variable with the peak month identified from January to August, pointing to mixed emission processes. In western Africa, and consistently with what is observed from ground-based measurement [Adon *et al.*, 2010],  $\text{NH}_3$  maxima are observed by IASI sooner above wet savannas (February to April) than above dry savannas (May to July). This is interesting and tends to confirm the hypothesis of Delon *et al.* [2012] that dry savanna emissions are dominated by  $\text{NH}_3$  volatilization (function of soil temperature among others) while wet savanna emissions occur also because of burning processes. A 1 month lag is observed compared to the fire seasonality (peaking in January) for a large part of the northern central Africa, which suggests agricultural emissions following biomass burning, as recently proposed in Whitburn *et al.* [2015].

The distribution of the month of maximum  $\text{NH}_3$  in southern America is, as for the southern parts of Africa, closely related with the fire seasonality (peaking from August to December), which indicates that this is the major source of  $\text{NH}_3$  in this continent. As an example, the area peaking in August in Brazil is part of the "Arc of Deforestation," well known for its fire activity [Chen *et al.*, 2013]. The maxima later in the year suggest  $\text{NH}_3$  emissions from agricultural areas, possibly linked with waste burning, especially in Argentina. The Llanos area (Venezuela) shows maxima in other months, March and April, which indicates that agricultural practices act there as a nonnegligible source, which add up to the vegetation burning that occurs primarily from January to March [Giglio *et al.*, 2006].

In central and northern America, the spatial patterns in month of maximum columns are more homogeneous. Western U.S. is characterized by peaks in summer months, more than likely due to enhanced  $\text{NH}_3$  volatilization in a drier environment at higher temperature. Consistent high July–August  $\text{NH}_3$  concentrations have, for instance, been reported recently above a rural gas production area in western Wyoming [Li *et al.*, 2014]. The San Joaquin Valley stands up in western U.S. showing its maximum of  $\text{NH}_3$  columns in October. Further studies are needed to identify the reason for this difference, which could be related to the large weight associated with the evening measurements in that regions, indeed peaking in October–November [Clarisse *et al.*, 2010]. The great plains in U.S. and Canada show an earlier maximum in April–May, suggesting agricultural spring emissions as a major source, similarly to Europe. The same can be said for Southern Ontario, in agreement with measurements from the ground [Zbieranowski and Aherne, 2013]. This timing is consistent with a study based on ammonium wet deposition data which have also shown this earlier peak of  $\text{NH}_3$  for the Midwest region, attributed to differences in agricultural activities [Paulot *et al.*, 2014].

#### 4. Conclusions

We have analyzed 6 years (2008–2013) of  $\text{NH}_3$  total columns retrieved globally from IASI measurements twice daily. Monthly mean total columns have been calculated with an error-weighting procedure for several regions, separately for morning and evening overpasses. In most situations, the monthly means from the IASI morning measurements have smaller errors. Despite this, the evening measurements allow capturing the seasonal cycle over northern central Africa as well; it highlights also additional winter peaks in southwestern Asia. In other regions, the evening measurements are not sufficiently accurate year-round, and for the months wherein measurements are available, a general consistency with the morning measurements is found. It is the first time that such an assessment of the added value of the evening IASI- $\text{NH}_3$  column observations is performed. It is also the first time that IASI- $\text{NH}_3$  observations are analyzed at subcontinental scale over a long period. It allows unprecedented investigations of the spatiotemporal variability of  $\text{NH}_3$  in the atmosphere.

The seasonal cycles are well marked in all regions, with the largest amplitude in Asia, where also the highest columns are measured (both at background and peak level). Europe's and northern America's seasonalities were categorized with a cycle of similar amplitude, characterized by high spring columns. The relations between atmospheric abundances and emission processes was also emphasized at smaller regional scale by extracting the global climatology of the month of maxima columns at high spatial resolution. In some regions, the predominance of a major source appears clearly (e.g., agriculture in Europe and north America and fires in central South Africa and south America), while in others mixed source processes on small scale are obvious (e.g., northern central Africa and southwestern Asia). We finally demonstrated that while the interannual variability does not allow to reveal clear trends, 2010 stands up as the year with largest  $\text{NH}_3$  concentrations, principally because of large emissions from fires in both hemispheres.

More generally, the results presented here underline the significance of the IASI satellite measurements of  $\text{NH}_3$ , which are performed at high temporal sampling and resolution, to characterize and understand the

variability of sources and atmospheric processes on the globe and thereby to help constraining models in the parameterization of these processes, as a function of environmental conditions and anthropogenic practices. In particular, the multiyear data set available from IASI satellites will allow, with the help of global chemistry-transport models, to assess bottom-up emission inventories through source inversion and to improve our knowledge of area-specific emission seasonality and interannual variability. It will also allow examining the effect of other changing geophysical or climatic parameters (temperature and NO<sub>2</sub> or SO<sub>2</sub> emissions) on the NH<sub>3</sub> global distributions.

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