10–year satellite–constrained fluxes of ammonia improve performance of chemistry transport models

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Abstract

In recent years, ammonia emissions have been continuously increasing being almost four times higher than in the 20th century. Although an important species as its use as a fertilized sustains human living, ammonia has major consequences both for humans and the environment, because of its reactive gas phase chemistry that makes it easily convertible to particles. Despite its pronounced importance, yet, ammonia emissions are highly uncertain in most emission inventories. However, the great development of satellite remote sensing nowadays provides the opportunity for more targeting research in constraining ammonia emissions. Here, we used satellite measurements to calculate global ammonia emissions over the period 2008–2017. Then, the calculated ammonia emissions were fed to a chemistry transport model and ammonia concentrations were simulated for the period 2008–2017.

The simulated concentrations of ammonia were compared with ground measurements from Europe, North America and Southeastern Asia, as well as with satellite measurements. The satellite-constrained ammonia emissions represent global concentrations more accurately than state-of-the-art emissions, which underestimate ammonia with a factor of two. Calculated fluxes in the North China Plain were increased after 2015, not due to emission changes, but due to changes in sulfate emissions that resulted in less ammonia neutralization and hence in larger atmospheric loads. Emissions over Europe were also twice as much as those in traditional datasets with dominant sources to be industrial and agricultural applications. Four hot-spot regions of high ammonia emissions were seen in North America characterized by large agricultural activity (Colorado), animal breeding (Iowa, northern Texas and Kansas), animal farms (Salt Lake, Cache, and Utah) and animal breeding and agricultural practices (California). South America is dominated by ammonia emissions from biomass burning, which cause a strong seasonality. In Southeastern Asia, ammonia emissions from fertilizer plants in China, Pakistan, India and Indonesia are the most important, while a strong seasonality was observed with a spring and late summer peak due to rice and wheat cultivation. Modelled concentrations from the satellite-constrained ammonia emissions are overestimated in Eastern Europe, where state-of-the-art emissions capture observations better. Measurements of ammonia concentrations in North America were better reproduced with satellite-constrained emissions, while all emissions generally underestimate station concentrations in Southeastern Asia. The calculated ammonia emissions also reproduce global CrIS (Cross-track Infrared Sounder) observations more effectively.
1 Introduction

Ammonia (NH$_3$) has received a lot of attention nowadays due to its major implications for the population and the environment (Erisman, 2004; Erisman et al., 2007). These include eutrophication of semi-natural ecosystems and acidification of soils (Stevens et al., 2010), secondary formation of particulate matter in the atmosphere (Anderson et al., 2003), and alteration of the global greenhouse balance (De Vries et al., 2011). More specifically in the troposphere, ammonia reacts with the abundant sulfuric and nitric acids (Malm, 2004) contributing 30% to 50% of the total aerosol mass of PM2.5 and PM10 (Anderson et al., 2003). Ammonium aerosols are therefore a very important component in regional and global aerosols processes (Xu and Penner, 2012) also having significant implications for human health (Aneja et al., 2009). Ammonia alters human health indirectly mainly through formation of PM2.5 (Gu et al., 2014) that penetrate the human respiratory systems and deposit in the lungs and alveolar regions (Pope III et al., 2002) causing premature mortality (Lelieveld et al., 2015). As regards to the climate impact, the same ammonium aerosol particles affect Earth’s radiative balance, both directly by scattering incoming radiation (Henze et al., 2012) and indirectly as cloud condensation nuclei (Abbatt et al., 2006). They may also cause visibility problems and contribute to haze effect due to secondary PM formation.

Sources of ammonia include wild animals (Sutton et al., 2000), ammonia-containing watersheds (Sørensen et al., 2003), traffic (Kean et al., 2009), sewage systems (Reche et al., 2012), humans (Sutton et al., 2000), biomass burning (Sutton et al., 2008) and domestic coal combustion (Fowler et al., 2004), volcanic eruptions (Sutton et al., 2008) and agriculture (Erisman et al., 2007). The latter is responsible for the majority of ammonia global atmospheric emissions. Specifically, in the United States and Europe about 80% of all emissions is related to agriculture (Leip et al., 2015). Emissions have increased considerably since pre-industrial times and are unlikely to decrease due to the growing demand for food and feed (Aneja et al., 2008).

The growing attention in ammonia levels has enabled many monitoring actions in Europe (European Monitoring and Evaluation Programme, EMEP), in Southeastern Asia (East Asia acid deposition NETwork) and in the North America (Ammonia Monitoring Network in the US, AMoN-US; National Air Pollution Surveillance Program (NAPS) sites in Canada) to record surface concentrations of ammonia continuously. Recently, several satellite products have been also developed in an effort to identify global levels of ammonia considering that the
relatively sparse existing monitoring network has an insufficient coverage for this purpose. These are derived from satellite sounders as the Infrared Atmospheric Sounding Interferometer (IASI) (Van Damme et al., 2017), the Atmospheric Infrared Sounder (AIRS) (Warner et al., 2017), the Cross-track Infrared Sounder (CrIS) (Shephard and Cady-Pereira, 2015) the Tropospheric Emission Spectrometer (TES) (Shephard et al., 2015), and Greenhouse Gases Observing Satellite (Someya et al., 2020). Both IASI and CrIS ammonia products are being continuously compared and evaluated against other observations and products. For example, against column-integrated levels measured by Fourier transform infrared spectroscopy (FTIR) (Dammers et al., 2016, 2017), ground-based measurements (Van Damme et al., 2015; Kharol et al., 2018), bottom-up emissions (Van Damme et al., 2018; Dammers et al., 2019) and atmospheric chemistry transport models (CTMs) (Shephard et al., 2020; Whitburn et al., 2016a).

Despite its importance, ammonia is a poorly quantified trace gas, with uncertainties over 50% on the global emission budget and even higher on temporal and local scales (Dentener and Crutzen, 1994; Faulkner and Shaw, 2008; Reis et al., 2009). In the present paper, we grid 10 years (2008–2017) of satellite measurements of ammonia retrieved from IASI to calculate monthly surface emissions (hereafter named NE) (see section 2). The same is done using the gridded IASI ammonia column concentrations from Van Damme et al. (2018) (named as VD0.5 and VDgrfl) (see section 2). The three different emission inventories together with a state–of–the–art one, which is more often used by models (named as EGG), are then imported in a CTM to simulate ammonia for the same 10–year period. Finally, an evaluation of simulated surface concentrations against ground–based measurements from different monitoring stations and satellite products allow to quantify the improvements in ammonia emissions.

2 Methods

2.1 Satellite ammonia

2.1.1 IASI ammonia

The Infrared Atmospheric Sounding Interferometer (IASI) onboard the MetOp-A satellite measures Earth’s infrared radiation twice a day in a spectral range of 645–2,760 cm⁻¹ with an elliptical footprint (at nadir circular with a diameter of 12 km) at nadir (Clerbaux et al., 2009). Due to the larger thermal conditions that lead to smaller uncertainties, only morning data were used in the present assessment (Clarisse et al., 2010). The 10–year dataset used here is ANNI-NH3-v2.1R-I product (Van Damme et al., 2017). The Artificial Neural Network for IASI
(ANNI) algorithm converts the hyperspectral range index to an column–integrated NH$_3$ value (Whitburn et al., 2016a). The dataset also provides cloud coverage for each measurement (August et al., 2012). Only measurements with a cloud fraction below 10% were processed in consistency with Van Damme et al. (2018). Cloud coverage was not provided for all measurements until March 2010 resulting in smaller data availability before that date. Although the retrieval algorithm uses a fixed vertical profile, extended validation of the resulting dataset has verified small uncertainties (Van Damme et al., 2015, 2018; Dammers et al., 2016; Whitburn et al., 2016b). For instance, Van Damme et al. (2018) reported a difference of 2%±24% (global average) in column–integrated ammonia using different vertical profiles in the retrieval algorithm.

### 2.1.2 CrIS ammonia

The Cross-Track Infrared Sounder (CrIS) was first launched on the NASA Suomi National Polar-orbiting Partnership (S-NPP) satellite on 28 October 2011 in a sun-synchronous low Earth orbit. The CrIS sensor provides soundings of the atmosphere with a spectral resolution of 0.625 cm$^{-1}$ (Shephard et al., 2015). One of the main advantages of CrIS is its improved vertical sensitivity of ammonia closer to the surface due to the low spectral noise of ~0.04K at 280K in the NH$_3$ spectral region (Zavyalov et al., 2013) and the early afternoon overpass that typically coincides with high thermal contrast, which is optimal for thermal infrared sensitivity. The CrIS Fast Physical Retrieval (CFPR) (Shephard and Cady-Pereira, 2015) retrieves an ammonia profile (14 levels) using a physics-based optimal estimation retrieval, which also provides the vertical sensitivity (averaging kernels) and an estimate of the retrieval errors (error covariance matrices) for each measurement. As peak sensitivity is typically in the boundary layer between 900 and 700 hPa (~1 to 3 km) (Shephard et al., 2020), the surface and total column concentrations are both highly correlated with the retrieved levels in the boundary layer. Shephard et al. (2020) reports estimated total column random measurement errors of 10–15%, with estimated total random errors of ~30%. The individual profile retrieval levels have estimated random measurement errors of ~10 to 30%, with estimated total random errors increasing to 60 to 100% due to the limited vertical resolution. These vertical sensitivity and error output parameters are also useful for using CrIS observations in applications (e.g. data fusion, data assimilation; model-based emission inversions (e.g., Cao et al., 2020; Li et al., 2019) as a satellite observational operator can be generated in a robust manner. The detection limit of CrIS measurements has been calculated down to 0.3–0.5 ppbv (Shephard et al., 2020). CrIS ammonia has been evaluated against other observations over North America with the Ammonia Monitoring Network (AMoN) (Kharol et
al., 2018) and against ground-based Fourier transform infrared (FTIR) spectroscopy observations (Dammers et al., 2017) showing small differences and high correlations.

### 2.2 Inverse Distance Weighting (IDW) interpolation

To process large amounts of measurements in a 2-dimensional grid of high resolution, oversampling methods (Streets et al., 2013) can be used (Van Damme et al., 2018). However, considering that the resolution of the CTM is 2.5°×1.3° (see section 2.4), there is no need to process the measurements on such a high-resolution grid and therefore an interpolation method was used. The method has been extensively used after the Chernobyl accident in 1986 to process more than 500 thousand deposition measurements over Europe (De Cort et al., 1998; Evangeliou et al., 2016).

IASI ammonia was interpolated onto a grid of 2.5°×1.3° using a modified Inverse Distance Weighting (IDW) algorithm described by (Renka, 1988). This method is preferred due to its ease of use and to its high quality of interpolation. The IDW interpolation is defined by:

\[
\hat{v}(x, y) = \sum_{i=1}^{n} \frac{w_i v_i}{\sum_{i=1}^{n} w_i} \tag{Eq. 1}
\]

where \(\hat{v}(x, y)\) is the interpolated value at point \((x, y)\), \(w_1, ..., w_i\) are the relative weights and \(v_1, ..., v_n\) are the observation values. The weights are defined by the inverse distance functions:

\[
w_i = \left(\frac{r_i - d_i}{r_{iw}}\right)^2 \tag{Eq. 2}
\]

for \((r_{iw} - d_i) = \begin{cases} r_{iw} - d_i & \text{if } d_k < r_{iw}, \\ 0 & \text{if } d_k \geq r_{iw}. \end{cases}\)

where \(r_{iw}\) denotes the radius of influence of the point \((x_i, y_i)\), \(d_i\) the Euclidean distance between point \((x, y)\) and \((x_i, y_i)\), and \(d_k\) is the threshold distance. We used a threshold distance \(d_k\) of 50 km, which is similar to the size of each grid cell; different \(d_k\) values were included in a sensitivity study (see section 4.3). The Euclidean distance is calculated using Vincenty's formulae (Vincenty, 1975).

### 2.3 Emission flux calculation of ammonia

The emission fluxes of ammonia were calculated using a 1-dimensional box model that assumes first-order loss terms for ammonia and has been already used previously (Van Damme et al., 2018; Whitburn et al., 2016b). It takes into account the gridded column concentrations of ammonia that were calculated with the IDW interpolation method and all the potential removal
processes of ammonia occurring in a hypothetical atmospheric box according to the following equation:

\[ E_{NH3} = \frac{M_{NH3}}{\tau} \quad \text{Eq. 3} \]

where \( M_{NH3} \) is the mass of ammonia in each atmospheric box (grid-cell) in molecules cm\(^{-2} \) and \( \tau \) is the lifetime of ammonia in the box (given in seconds).

Van Damme et al. (2018) assumed a constant lifetime for ammonia, admitting that this is a limiting factor of their study on the basis that chemical loss and deposition are highly variable processes that can change the lifetime drastically. To tackle the large variability of the lifetime of ammonia, we used gridded lifetime calculated from a CTM. This gives robustness in the calculated emissions fluxes considering that at regions where sulfuric and nitric acids are abundant, the chemical loss will be more intense and, thus, lifetime will be much shorter affecting emissions dramatically.

The lifetime \( (\tau) \) of ammonia in each grid-box results from the three processes affecting ammonia concentrations: transport \( (t_{trans}) \) in and out of the grid-cell, chemical loss \( (t_{chem}) \) and deposition \( (t_{depo}) \):

\[ \frac{1}{\tau} = \frac{1}{t_{trans}} + \frac{1}{t_{chem}} + \frac{1}{t_{depo}} \quad \text{Eq. 4} \]

In a CTM, the lifetime can be easily calculated from the species mass balance equation (Croft et al., 2014):

\[ \frac{dC(t)}{dt} = S(t) - \frac{C(t)}{\tau(t)} \quad \text{Eq. 5} \]

where \( C(t) \) is the atmospheric burden of ammonia at time \( t \), \( S(t) \) is the time-dependent source emission fluxes and \( \tau(t) \) is the removal timescale. Assuming steady-state conditions and considering that emission fluxes of ammonia are continuous, there is a quasi-equilibrium between sources and removals of ammonia (Dentener and Crutzen, 1994), and the modeled lifetime of ammonia \( \tau_{mod} \) can be defined as:

\[ \tau_{mod} = \frac{C_{NH3}}{L_{NH3}^{\text{trans,chem,depo}}} \quad \text{Eq. 6} \]

where \( C_{NH3} \) is the atmospheric burden of ammonia and \( L_{NH3}^{\text{trans,chem,depo}} \) is the total loss due to any process affecting ammonia in the model (transport, chemical reactions, deposition).

We calculate ammonia emission fluxes using IASI satellite measurements that we interpolated (section 2.2) to the model resolution \((2.5^\circ \times 1.3^\circ)\) and applying a variable lifetime taken from a CTM (hereafter NE emissions). We also calculate ammonia emissions from the
oversampled IASI data of Van Damme et al. (2018), after bilinear re-gridding to the model resolution (2.5°×1.3°), applying a constant lifetime for ammonia of 12 hours (hereafter VD0.5 emissions) and a variable lifetime from a CTM (hereafter VDgrlf emissions).

### 2.4 LMDz-OR-INCA chemistry transport model

The Eulerian global CTM LMDz-OR-INCA was used to calculate ammonia lifetime, as well as to simulate ammonia concentrations from the emission fluxes calculated from IASI satellite products. The model couples the LMDz (Laboratoire de Météorologie Dynamique) General Circulation Model (GCM) (Hourdin et al., 2006) with the INCA (INteraction with Chemistry and Aerosols) model (Folberth et al., 2006; Hauglustaine et al., 2004) and with the land surface dynamical vegetation model ORCHIDEE (ORganizing Carbon and Hydrology In Dynamic Ecosystems) (Krinner et al., 2005). In the present configuration, the model has a horizontal resolution of 2.5°×1.3°, the vertical dimension is divided into 39 hybrid vertical levels extending to the stratosphere. Large-scale advection of tracers is calculated from a monotonic finite-volume second-order scheme (Hourdin and Armengaud, 1999), deep convection is parameterized according to the scheme of Emanuel, (1991), while turbulent mixing in the planetary boundary layer (PBL) is based on a local second-order closure formalism. More information and a detailed evaluation of the GCM can be found in Hourdin et al. (2006).

The model simulates atmospheric transport of natural and anthropogenic aerosols recording both the number and the mass of aerosols. The aerosol size distribution is represented using a modal approach that consists of the superposition of 5 log-normal modes that represent both the size spectrum and whether the aerosol is soluble or insoluble (Schulz, 2007). The aerosols are treated in three particle modes, sub-micronic (diameter < 1 μm) corresponding to the accumulation mode, micronic (diameter 1–10 μm) corresponding to coarse particles, and super-micronic or super coarse particles (diameter > 10 μm). LMDz-OR-INCA accounts for emissions, transport (resolved and sub-grid scale), and dry and wet (in-cloud/below-cloud scavenging) deposition of chemical species and aerosols interactively. LMDz-OR-INCA includes a full chemical scheme for the ammonia cycle and nitrate particle formation, as well as a state-of-the-art CH₄/NOₓ/CO/NMHC/O₃ tropospheric photochemistry. Further details about specific reactions, reaction rates and other information entering into the description of the ammonia cycle can be found in Hauglustaine et al. (2014).
The global transport of ammonia was simulated from 2007 to 2017 (2007 was the spin-up period) by nudging the winds of the 6-hourly ERA Interim Reanalysis data (Dee et al., 2011) with a relaxation time of 10 days (Hourdin et al., 2006). For the calculation of ammonia’s lifetime, the model ran with traditional emissions for anthropogenic, biomass burning and oceanic emission sources using emissions from ECLIPSEv5-GFED4-GEIA (hereafter called EGG) (Bouwman et al., 1997; Giglio et al., 2013; Klimont et al., 2017).

3 Results

In this section, the main results of the monthly emissions (NE) are presented for the 10-year period (2008–2017) of IASI observations. We first describe the simulated ammonia lifetimes (section 3.1). Then, we explain the main characteristics of the obtained emissions (section 3.2) and compare them with those calculated using the IASI gridded products from Van Damme et al. (2018) (VD0.5 and VDgrlf), as well as the ones from the state-of-the-art inventories of EGG and EDGARv4.3.1-GFED4 (Crippa et al., 2016; Giglio et al., 2013) that are often used in CTMs (section 3.3). We finally turn our focus to hot-spot regions and document their seasonal variation in emissions (section 3.4).

3.1 Modelled lifetime of ammonia

The lifetime of ammonia has been reported to range from a few hours to a few days (Behera et al., 2013; Pinder et al., 2008) so ammonia can only be transported over relatively short distances. This short spread of ammonia is also due to the fact that (a) the majority of its emissions are surface ones (major source is agricultural activity), and (b) its surface deposition velocities are high for most surfaces (Hov et al., 1994). The atmospheric lifetimes of ammonia were summarized in Van Damme et al. (2018). Specifically, Quinn et al. (1990) and more recently Norman and Leck (2005) reported lifetimes of a few hours in the West Pacific, South Atlantic and Indian Oceans, which is in agreement with Flechard and Fowler (1998), who reported a 2-hour lifetime in an area of Scotland where most sources are of agricultural origin. Similar to them, Dammers et al. (2019) recently reported a lifetime estimated from satellite measurements of 2.35±1.16 hours for large point sources based on satellite measurements. The majority of ammonia lifetimes reported regionally or globally fall within 10 and 24 hours independently of the different approaches (Hauglustaine et al., 2014; Hertel et al., 2012; Möller and Schieferdecker, 1985; Sutton et al., 1993; Whitburn et al., 2016b), while Dentener and Crutzen (1994) reported slightly higher lifetimes within a range between 0.9 and 2.1 days depending on ammonia emission fraction of natural origin. Monthly averaged atmospheric
ammonia lifetimes in the present study were derived using the version of the LMDz-OR-INCA that includes non-methane hydrocarbons (Hauglustaine et al., 2004).

Ammonia lifetime depends on numerous factors such as the presence of ammonia’s reactants (sulfuric and nitric acid), meteorological parameters (atmospheric water vapour, and temperature, atmospheric mixing and advection) and ammonia emissions. In ammonia–poor conditions, all ammonia is rapidly removed by neutralising sulfuric acid with an intermediate production of bisulfate. If ammonia increases further (ammonia–rich conditions), then reaction with nitric acid occurs forming nitric ammonium. At this point, the ammonia/sulfuric acid/nitric acid equilibrium becomes very fragile. If sulfate concentrations decrease, then free ammonia is produced, which gradually reacts with nitric acid resulting in production of aerosol phase nitric ammonium. But if particles are aqueous, then sulfate ions in solution increase the equilibrium vapour pressure of ammonia with nitric acid reversing the reaction towards gaseous phase reactants. So, sulfate reductions are linked with non-linear increases of aerosol nitrates and decreases of aerosol ammonium and water (Seinfeld and Pandis, 2000).

The calculated ammonia lifetime is shown in Figure 1a. The average lifetime was calculated to be 11.6±0.6 hours, which is in the range of the previously reported values. Lower values (~10 hours) were observed in clean remote areas characterized by low ammonia emissions (e.g., Amazon forest, Sahara and Australia), while in the rest of the globe the lifetime was closer to the average value. The highest lifetimes (~16 hours) occur over Southern Brazil and Venezuela, which are both areas with relatively high ammonia emissions and low sulfuric and nitric acid concentrations (Figure 1c). These conditions are characterized by a low atmospheric sulfuric and nitric acids availability to remove ammonia rapidly, hence causing an increase in lifetime.

3.2 Satellite-constrained emissions

The average ammonia emissions calculated from the 10-year IASI observations are shown in Figure 1b (also in Figure S 1a), the reactants’ atmospheric burden in Figure 1c and their seasonal variability in Figure 1d. The year-by-year total ammonia emissions are depicted in Figure S 1 with a monthly temporal resolution. Emissions decline from 242 Tg yr⁻¹ in 2008 to 212 Tg yr⁻¹ in 2011. In 2012 – 2014, emissions show little variation (194, 204 and 195 Tg yr⁻¹, respectively), before they increase steeply to 248 Tg yr⁻¹ in 2015. Finally, in 2016 and 2017 they remain at the same high level (197 and 227 Tg yr⁻¹, respectively).
The global average annual emission calculated from VD0.5 amounts to 189 Tg (9-year average), which is comparable to the average of the 10-year period that we have calculated in the present study (average±sd: 213±18.1 Tg yr⁻¹). The increase in the emissions we calculate during 2015 and 2017 stand out. The explanation for these increases could be twofold. If sulfur dioxide emissions decreased over time, less sulfates are available to neutralize ammonia, hence resulting in higher ammonia column concentrations seen by IASI that could be attributed to new emissions erroneously (see section 2.3). If sulfur dioxide and sulfates presented a constant year-by-year pattern or even increased, then the calculated ammonia emissions would be likely realistic.

To sort out between these two possibilities, we used sulfur dioxide measurements from NASA’s Ozone Monitoring Instrument (OMI, Yang et al., 2007) instrument, whereas sulfate column concentrations were taken from the Modern-Era Retrospective Analysis for Research and Applications, Version 2 (MERRA2, Gelaro et al., 2017) reanalysis data from NASA's Global Modeling and Assimilation Office (GMAO). Figure S 2 shows timeseries of column concentrations of sulfur dioxide and sulfates from OMI and MERRA2 averaged globally, for continental regions (Europe, North America, South America, Africa), as well as for regions where ammonia emissions are particularly high (India and Southeastern Asia, North China Plain). Although column concentrations of both sulfur dioxide and sulfates present strong interannual variability, they do not show significant changes on an annual basis. This indicates that sulfate amounts that neutralize ammonia and form ammonium sulfate are rather constant from year to year and, thus it is likely that the higher ammonia concentrations retrieved from IASI after 2015 are the result of an emission increase.

Another region of interest is the North China Plain, as it has been identified as an ammonia hotspot mainly due to extensive agricultural activities (Clarisse et al., 2009; Pan et al., 2018). Liu et al. (2018) reported a sulfur dioxide reduction of about 60% over the recent few years in the North China Plain, sulfates decreased by 50%, while ammonia emissions declined by only 7% due to change in agricultural practices. The suggested decrease in ammonia reactants over the North China Plain is illustrated by the calculated sulfur dioxide column concentration anomaly from OMI (Figure 2) and by the sulfate concentration anomaly from MERRA-2 after 2015 (the highest calculated one) (Figure S 3). However, the IASI-constrained ammonia emissions calculated here show only a tiny increase of 0.19±0.04 kt yr⁻¹ after 2015 in the North China Plain and of 10±3.1 Tg yr⁻¹ globally with respect to the 10-year
average (Figure 2). This is due to the change of SO2 and NOx emission regulations in China, which in turn led to reduced inorganic matter (sulfates, nitrates and ammonium) resulting in regional increases of gaseous ammonia (Lachatre et al., 2019).

### 3.3 Comparison with traditional emission datasets

In this section, we quantify the main differences of our IASI-constrained emission dataset with other state-of-the-art inventories used in global models and for different applications (air quality, climate change etc…). Aside from comparing our emissions with those calculated using Van Damme et al. (2018) data with a constant lifetime (hereafter called VD0.5), we extend our comparison to more traditional datasets such as those of ECLIPSEv5-GFED4-GEIA (EGG) for 2008–2017, and EDGARv4.3.1-GFED4 (Crippa et al., 2016; Giglio et al., 2013) for 2008–2012 period. Finally, the ammonia emissions presented in this study (NE emissions) are compared to emissions calculated from Van Damme et al. (2018) gridded IASI column data applying a variable (modelled) ammonia lifetime presented in Figure 1b (hereafter referred as VDgrlf).

The 10-year comparison of our calculated emissions with VD0.5 is shown in Figure 3. The 10-year average difference amounts to 29±15 Tg yr⁻¹ (average±sd). In all years, the largest differences could be seen over Latin America and over tropical Africa. Our emissions (NE) were lower in the Indo-Gangetic Plain, and situated slightly more northerly. Northern India has been previously identified as a hot-spot region for ammonia, mainly due the importance of agricultural activities in the region (Kuttipurath et al., 2020; Tanvir et al., 2019).

Figure S4 and Figure S5 present a comparison of our calculated emissions with the basic state-of-the-art datasets of EGG and EDGARv4.3.1-GFED4, respectively. In both datasets, ammonia emissions remain almost constant over time (average±sd: 65±2.8 Tg yr⁻¹ and 103±5.5 Tg yr⁻¹, respectively). The total calculated ammonia emissions are up to three times lower than those calculated from NE (average±sd: 213±18.1 Tg yr⁻¹) or from VD0.5 (9-year average: 189 Tg yr⁻¹). This results in 10-year annual differences that are very significant (average±sd: 150±19.3 Tg yr⁻¹ and 111±19.2 Tg yr⁻¹, respectively); the largest differences appear over South America, while European emissions are practically identical in all datasets. Emissions from South China Plain are much higher in the two traditional datasets that those calculated here. Based upon IASI retrievals, Liu et al. (2019) showed an increase of surface NH₃ concentrations trend of more than 0.2 µg N m⁻³ yr⁻¹ in eastern China and of around 0.1–0.2 µg N m⁻³ yr⁻¹ in northern Xinjiang during 2008–2016. Ammonia emissions derived over China in this work are
among the highest worldwide (Figure S 1), which agrees well with the 9-year average emissions calculated in VD0.5 inventory over China (see Figure 3). To assess to which extent emissions from EGG and EDGARv4.3.1-GFED4 are underestimated can only be done by comparing ammonia with ground or satellite observations.

The comparison of the annual ammonia emissions to the modified VDgrlf emissions is shown in Figure S 6. The latter showed a better agreement to the emissions presented in this study with mean annual different of 14±19 Tg yr\(^{-1}\) (average±sd). Previously observed emission differences in the two state-of-the-art inventories over South America and Africa have been now minimized, as well as the displacement north of the Indo-Gangetic Plain emissions remains important. Nevertheless, the smaller differences of our emissions from those of VDgrlf as compared with the respective difference from the VD0.5 emissions, show the large impact that a more realistic variable lifetime might have in emission calculations with this methodology.

3.4 Site-specific ammonia emissions and seasonal variation

Figure 4 illustrates specific regions that show the largest ammonia emissions (Europe, North America, South America and Southeastern Asia). These emissions correspond to the IASI-constrained emissions calculated in this study (NE) and are presented as total annual emissions averaged over the 10-year period of study. At the bottom panels of the same figure, the seasonal variation of the emissions is shown for each of the four hot-spot regions and each of the 10 years of the study.

European total ammonia emissions were estimated to be 15±2.2 Tg yr\(^{-1}\) (average±sd), more than double compared with those reported in EGG (6.9±1.1 Tg yr\(^{-1}\)) and similar to those in VD0.5 (11 Tg yr\(^{-1}\)) or those in VDgrlf (11±1.0 Tg yr\(^{-1}\)). The greatest emissions were calculated for Belgium, the Netherlands and the Po Valley in Italy (Figure 4). High emissions are also found in North and Northwestern Germany and over Denmark. In contrast, very low emissions are found in Norway, Sweden and parts of the Alps. It is not possible to quantitatively distinguish between different sources of ammonia. It has been reported that approximately 75% of ammonia emissions in Europe originate from livestock production (Webb et al., 2005), and 90% from agriculture in general (Leip et al., 2015). More specifically, ammonia is emitted from all stages of manure management, from livestock buildings during manure storage and application to land, as well as from livestock urine. These emissions are strong over most of Northwestern European countries, although sources like fertilization and non-agricultural
activities (traffic and urban emissions) can be also important. An example is Tange in Germany, which shows a late summer peak due to growing crops application. No obvious seasonality in the emissions can be seen for Europe as a whole, as the hot-spot regions are rather few compared to the overall surface of Europe. An exception to this stable emission situation over the year occurs during 2010 and during 2015, years for which a late summer peak. In 2010, large wildfires in Russia resulted in high ammonia emissions (R'Honi et al., 2013), while year 2015 has been also characterized as an intense fire year (though not like 2010), with fires occurring in Eurasia (Min Hao et al., 2016).

North America and in particular the US (Figure 4) has been characterized by four hot-spot regions. First, a small region in Colorado, Central US, which is the location of a large agricultural region that traditionally releases large ammonia emissions (Malm et al., 2013). Another example is the state of Iowa (home to more than 20 million swine, 54 million chickens, and 4 million cattle), northern Texas and Kansas (beef cattle), and southern Idaho (dairy cattle) (McQuilling, 2016). Furthermore, the three major valleys in Salt Lake, in Cache, and in Utah in the midwestern US show an evident, but lower intensity hot-spot, as they are occupied by massive pig farms associated to open waste pits. The largest emissions were calculated for the San Joaquin Valley in California (vegetables, dairy, beef cattle and chickens) and further to the South (Tulare and Bakersfield), an area characterized by feedlots (Van Damme et al., 2018; McQuilling, 2016). North American annual ammonia emissions over the 10-year period were averaged 1.1±0.1 Tg yr⁻¹ (average±sd). These values are over two orders of magnitude higher than those in EGG (0.062±0.0013 Tg yr⁻¹). Note that his estimate is three times lower than those reported in VD0.5 (3.1 Tg yr⁻¹) or in VDgrlf (3.4±0.5 Tg yr⁻¹). The 2008–2017 interannual variability (Figure 4) all show a minimum in winter. Maximum emissions were observed in late spring, due to the contribution from mineral fertilizer and manure application, in summer, due to influence of livestock housing emissions, and some years both in spring and summer (Makar et al., 2009; Zhu et al., 2013, 2015). A topographical dependence was also seen in midwest emissions that peaked in April, whereas over the rest of the US maximum emissions were appeared in summer (Paulot et al., 2014).

Ammonia emissions have different characteristics in South America and in Western Africa as both are fire-dominated regions. For simplicity we only present South America in Figure 4. This region is dominated by natural ammonia emissions mainly from forest, savanna and agricultural fires (Whitburn et al., 2014, 2016b) and volcanoes (Kajino et al., 2004;
Uematsu et al., 2004). This causes a strong seasonal variability in the ammonia emissions with the largest fluxes observed from August to October in all years (Figure 4). This strong dependence of South America from biomass burning emissions was first highlighted by Chen et al. (2013) and by van Marle et al. (2017). It also became particularly pronounced during the large wildfires in the Amazon rainforest in summer 2019 (Escobar, 2019). We estimated the 10-year average ammonia emissions to be 28±3.0 Tg yr\(^{-1}\) (average±sd) in agreement with VD0.5 (22 Tg yr\(^{-1}\)) and VDgrlf (24±1.3 Tg yr\(^{-1}\)). The respective emissions in EGG are four times lower than these estimates (7.1±0.3 Tg yr\(^{-1}\)).

The last column to the right of Figure 4 presents the 10-year average annual ammonia emissions and their respective interannual variability in Southeastern Asia. We define this region spanning from 70°E–130°E in longitude and from 0°N–45°N in latitude. Ammonia emissions were estimated to be 38±2.8 Tg yr\(^{-1}\) (average±sd) similar to VD0.5 (36 Tg yr\(^{-1}\)) and VDgrlf (39±1.8 Tg yr\(^{-1}\)) and slightly higher than those presented in EGG (25±1.2 Tg yr\(^{-1}\)). They comprise ammonia fertilizer plants, such as in Pingsongxiang, Shizuishan, Zezhou-Gaoping, Chaerhan Salt Lake, Delingha, Midong-Fukang and Wucaiwan (China), Indo-Gangetic Plain (Pakistan and India), Gresik (Indonesia). China and India contribute more than half of total global ammonia emissions since the 1980s with the majority of these emissions to originate from rice cultivation followed by corn and wheat (crop-specific emissions). More specifically, emissions from these crops due to synthetic fertilizer and livestock manure applications are concentrated in North China Plain (Xu et al., 2018). Considering that Southeastern Asia is the largest agricultural contributor in the global ammonia budget, a strong seasonality in the emissions was observed. Temporal ammonia emissions peak in late summer of most years, when emissions from rice cultivation, synthetic fertilizer application and livestock manure spreading (Xu et al., 2016) are important, and in spring when wheat cultivation dominates (Datta et al., 2012). Of course, the respective emissions from biomass burning should also be mentioned. However, these are difficult to be distinguish and are expected to be a relatively small source compared to agricultural emissions.

### 4 Discussion

In this section, we conduct simulations over the 10-year period (2008–2017, 1-year spin-up), with all the emissions derived and compare the NH\(_3\) concentrations with ground based observations over Europe, North America, Southeastern Asia (section 4.1), and observations from CrIS (section 4.2). These simulations consist of: (i) a simulation using traditional
emissions from ECLIPSev5 (Evaluating the CLimate and Air Quality ImPacts of Short-livEd Pollutants) for anthropogenic sources, GFED4 (Global Fire Emission Dataset) for biomass burning emissions and GEIA (Global Emissions InitiAtive) for oceanic sources from (Bouwman et al., 1997; Giglio et al., 2013; Klimont et al., 2017), where modelled lifetimes of ammonia were also calculated (EGG); (ii) a simulation using emissions calculated from IASI data from Van Damme et al. (2018) applying a constant lifetime of 12 hours for ammonia (VD0.5); (iii) a simulation using gridded emissions presented in the present paper (NE) calculated as described in section 2; and (iv) a simulation using emissions calculated from IASI data from Van Damme et al. (2018) applying a variable (modelled) lifetime (VDgrlf). Finally, we perform a sensitivity analysis in order to define the levels of uncertainty of our emissions in section 4.3 and discuss potential limitation of the present study in section 4.4.

4.1 Validation against ground-based observations

Figure 5 shows a comparison between modelled surface concentrations of ammonia with ground measurements from Europe (EMEP, https://emep.int/mscw/), North America (AMoN, http://nadp.slh.wisc.edu/data/AMoN/) and Southeastern Asia (EANET, https://www.eanet.asia). To avoid overplotting, the Gaussian kernel density estimation (KDE) was used, which is a non-parametric way to estimate the probability density function (PDF) of a random variable (Parzen, 1962):

$$ f(x) = \frac{1}{Nh} \sum_{i=1}^{N} K\left( \frac{x-x_i}{h} \right) $$  

Eq. 7

where $K$ is the kernel, $x_i$ the univariate independent and identically distributed point of the relationship between modelled and measured ammonia and $h$ is a smoothing parameter called the bandwidth. KDE is a fundamental data smoothing tool that attempts to infer characteristics of a population, based on a finite dataset. It weighs the distance of all points in each specific location along the distribution. If there are more points grouped locally, the estimation is higher as the probability of seeing a point at that location increases. The kernel function is the specific mechanism used to weigh the points across the data set and it uses the bandwidth to limit the scope of the function. The latter is computed using the Scott’s factor (Scott, 2015). We also provide the mean fractional bias (MFB) for modelled and measured concentrations of ammonia as follows:

$$ MFB = \frac{1}{N} \sum_{i=1}^{N} \left( \frac{C_m - C_o}{C_o} \right) \times 100\% $$  

Eq. 8

where $C_m$ and $C_o$ are the modelled and measured ammonia concentrations and $N$ is the total number of observations. MFB is a symmetric performance indicator that gives equal weights...
to under- or over-estimated concentrations (minimum to maximum values range from -200% to 200%). Furthermore, we assess the deviation of the data from the line of best fit using the root mean square error (RMSE) defined as:

\[ RMSE = \sqrt{\frac{\sum_{i=1}^{N} (C_m - C_o)^2}{N}} \quad \text{Eq. 9} \]

From 134 European stations, nearly 300,000 measurements made at a daily to weekly temporal resolution over the period of study (2007–2018) are presented on Figure 5. All emission datasets underestimate ammonia surface concentration over Europe. The most accurate prediction of concentrations was achieved using the traditional EGG emissions that underestimated observations by 67%, also being the least scattered from the best fit \((RMSE_{EGG} = 4.06 \mu g \ N \ m^{-3})\), followed by the emissions presented in this paper \((MFB_{NE} = -72\%), \ RMSE_{NE} = 4.65 \mu g \ N \ m^{-3})\), although they were more variable. VDO.5 or VDgrlf emissions further underestimated observations, though they were less sparse (Figure 5d). About 12% of the modelled concentrations using EGG were outside of the 10-fold limit from the observations, in contrast to only 17% and 15% in VD0.5 and VDgrlt, and 20% in NE. With regards to the spatial comparison with the observed concentrations, all datasets cause overestimations in the ammonia concentrations predicted in Eastern Europe (station AM0001R). EGG appears to be the most accurate in Central Europe (all stations with suffix DE00), NE emissions in all Spanish stations (suffix ES00) and VD0.5 and VDgrlf emissions in Italian stations (Figure S 7).

The comparison of simulated ammonia concentrations to observations over North America includes 119 stations, which represent nearly 27,000 observations (Figure 6) with a weekly, bi-weekly or monthly resolution. The only emission dataset that lead to an underestimation of ammonia concentrations was EGG \((MFB_{EGG} = -28\%)\). Two others, VD0.5 and VDgrlf caused ammonia observations to be strongly overestimated \((MFB_{VD0.5} = 52\% \text{ and } MFB_{VDgrlf} = 54\%)\), while NE slightly \((MFB_{NE} = 32\%)\). All inventories resulted in about the same variability in ammonia concentrations with RMSEs between 4.15 and 4.17 \(\mu g \ N \ m^{-3}\) (Figure 6). About 10% of the predicted concentrations using EGG emissions were at least 10 times off from the measured ones, more than twice the number of measurements compared to the other dataset. NE emissions better capture levels in the easternmost stations of the US (AL99, AR15, CT15, IL37, IN22, MI52, NY56, ON26) and in California (CA83) and Oklahoma (OK98), which are close to hot-spot regions (see section 3.4). EGG emissions perform better in Northwestern (ID03), Central (KS03) and several stations located over the
Eastern United States (KY03, KY98, OH09, AR03, IL46, KS03, GA41). The emission inventory VD0.5 leads to a very good agreement in ammonia concentrations over all stations of the North American continent (AL99, GA40, ID03, GA41, IL37, IL46, IN20, IN22, KS97, PA00, MD99, M152, TN04, NM99, NY96, OH99, OK98) (Figure S 8). In Southeastern Asia 62 stations from 13 countries were included in the comparison from the EANET monitoring network (Figure 7). These included about 8,000 surface measurements in monthly or 2-weekly resolution. All emission inventories underestimate station concentrations of EANET with MFBs beween -102% (EGG) and -61% (VD0.5 and VDgrlf ()). The least spread model concentrations were those simulated using VD0.5 and VDgrlf ($RMSE = 4.61 – 4.65 \mu g N m^{-3}$). Around 19% of model concentrations using EGG were outside the 10-fold limit of the 1×1 line with observations, 12% using NE emissions and only 5% and 6% using VD0.5 and VDgrlf, respectively. VD0.5 and VDgrlf emissions capture well the Japanese (suffix JPA) and Taiwanese stations (suffix THA). Given the short lifetime and the relatively coarse spatial scales, the model fails to capture the variability that exists within each gridbox (Figure S 9).

### 4.2 Validation against satellite products

Here, we used surface ammonia concentrations from CrIS from 1st May 2012 to 31st December 2017 and we compared them with modelled ammonia concentrations using four emissions datasets (EGG, VD0.5, NE and VDgrlf), like in the previous section. The comparison is shown as PDF of surface modelled against CrIS concentrations of ammonia calculated with the Gaussian KDE in Figure 8. A total of 4.5 million surface measurements were used in the comparison with a global coverage. All datasets underestimated surface concentrations except NE emissions, which overestimate ammonia ($MFB = +0.48$). The best fit was achieved for the VDgrlf emissions, which slightly underestimate ammonia ($MFB = −0.37$), while 82% of the measurements were within one order of magnitude from the 1×1 line, which is also shown by the small $RMSE$. VD0.5 emissions produced similar concentrations, with respect to the $RMSE$ and $MFB$ values, whereas 79% of them were less than a 10-fold difference from the observations. NE emissions result in higher surface concentrations, also showing larger $RMSE$s. However, 90% of the modelled concentrations were within a factor of 10 from the CrIS observation. In general, a better agreement for the most recent years 2015 – 2017 was achieved. The baseline EGG emissions resulted in significantly larger deviations of modelled
surface concentrations of ammonia from the CrIS observations, as shown in Figure 8, comprising the largest RMSE and MFB values.

### 4.3 Uncertainty analysis

A sensitivity analysis in order to calculate the level of uncertainty that each of the parameter gives to the modelled surface concentrations of ammonia was also performed. The relative uncertainty was calculated as the standard deviation of ammonia’s surface concentrations from a model ensemble of 10 members (Table 1) divided by the average. The first six members are the surface concentrations that resulted from simulations of ammonia emissions after perturbation of the Euclidian distance \( d_k \) in the parameters of the IDW interpolation. The remaining four members are simulated concentrations using the previously reported emissions datasets (EGG, VD0.5, NE and VDgrf). The results are shown as a 10-year (2008–2017) annual average relative uncertainty in Figure 9 and as annual average relative uncertainty of surface concentrations for every year of the 10-year period in Figure S 10.

The surface concentrations resulting from the different calculated emissions mainly affects oceanic regions, with values reaching 100%. The reason for this could be threefold. First, the IDW interpolation shows to be affected by severe outlier values, which are found in several oceanic regions (Figure S 11); this creates high gridded column ammonia concentrations and, in turn, fluxes at regions that are not supported by previous findings or measurements. Second, the methodology with which ammonia concentrations are retrieved in IASI has certain limitation, with respect to (i) the use of constant vertical profiles for ammonia, (ii) potential dependencies of total column ammonia and temperature that are not taken into account, and (iii) instrumental noise that can cause a high bias of the measurements (Whitburn et al., 2016a). Third, there is much less ammonia over the Ocean, hence the relative error bars are much larger. Large uncertainties in surface ammonia concentrations were observed in regions characterized by large anthropogenic contribution, such as North India, North China Plain and Central USA. Smaller uncertainties were found in Central Africa and in Amazonia, regions that are linked with episodic biomass burning emissions (Figure 4).

### 4.4 Limitations of the present study

We discuss the importance of certain limitations in the methodology of the present study and in the validation of the results. These limitations will also be commented upon in the overall conclusion of the paper.
Regarding the methodology, emissions of short-lived species are determined, among other methods, using top-down approaches. When only satellite measurements are available, they are usually averaged over a particular location and surface emissions are calculated using a mass balance approach (Lin et al., 2010; Zhao and Wang, 2009). This is done by assuming a 1-dimensional box-model, where atmospheric transport between grids is assumed to be negligible and loss due to deposition or chemical reactions very fast. The solution to this problem is the use of Kernels (Boersma et al., 2008), which makes the computation of the emissions very intense. It has been reported that for resolutions, such as those used in the present paper (2.5°×1.3°), non-local contributions to the ammonia emissions are relatively small (Turner et al., 2012). Although, the use of Kernels is the proper way to account for non-local contributions, we believe that negligible transport here is a fair assumption, due to the small lifetimes of ammonia calculated from the CTM (11.6±0.6 hours); therefore, transportation from the adjacent grid-cells should be small. Note that already this method has been suggested for short lived climate pollutants, it is not suitable for species with lifetime from days to weeks (e.g. black carbon, Bond et al., 2013).

Another limitation of the present study is that the same model is used for the calculation of the modelled lifetimes and for the validation of the emissions that were calculated using these lifetimes (NE and VDgrlf). A more accurate validation would require an independent model for the simulations of surface concentrations using these emissions.

5 Conclusions

In the present paper, satellite measurements from IASI were used to constrain global ammonia emissions over the period 2008–2017. The data were firstly processed to monthly ammonia column concentrations with a spatial resolution of 2.5°×1.3°. Then, using gridded lifetime for ammonia calculated with a CTM, monthly fluxes were derived. This contrasts with previously reported methods that used a single constant lifetime. This enables a more accurate calculation in regions sensitive to the changing balance between nitrate and sulfate abundances.

The calculated ammonia emission fluxes were then used to simulate ammonia concentrations for the period 2008–2017 (referred to as NE). The same simulations were repeated using baseline emissions from ECLIPSev5-GFED4-GEIA (referred to as EGG), emissions constrained by Van Damme et al. (2018) IASI data using a constant lifetime for ammonia (named as VD0.5) and emissions based on Van Damme et al. (2018) retrievals using a modelled lifetime from a CTM (named as VDgrlf). The simulated surface concentrations of ammonia were compared with ground measurements over Europe (EMEP), North America (AMoN) and...
Southeastern Asia (EANET), as well as with global satellite measurements from CrIS. The main conclusions can be summarized as follows:

- The 10-year average annual ammonia emissions calculated here (NE) were estimated to be 213±18.1 Tg yr⁻¹, which is 15% higher than those in VD0.5 (189 Tg yr⁻¹), and 6% higher than those in VDgrlf (201±10.4 Tg yr⁻¹). These emission values amount to twice the published from datasets, such as EGG (65±2.8 Tg yr⁻¹) and EDGARv4.3.1-GFED4, (103±5.5 Tg yr⁻¹).

- In the North China Plain, a region characterized by intensive agricultural activities, a small increase of ammonia emissions is simulated after 2015. This is attributed to decreases in sulfur species, as revealed from OMI and MERRA-2 measurements. Less sulfates in the atmosphere leads to less ammonia neutralization and hence to larger loads in the atmospheric column as measured by IASI.

- In Europe, the 10-year average of ammonia emissions were estimated at 15±2.2 Tg yr⁻¹ (NE), twice as much as those in EGG (6.9±1.1 Tg yr⁻¹) and similar to those in VD0.5 (11 Tg yr⁻¹) or VDgrlf (11±1.0 Tg yr⁻¹). The strongest emission fluxes were calculated over Belgium, Netherlands, Italy (Po Valley), Northwestern Germany and Denmark. These regions are known for industrial and agricultural applications, animal breeding activities, manure/slurry storage facilities and manure/slurry application to soils.

- Some hot-spot regions with high ammonia emissions were distinguished in North America: (i) in Colorado, due to large agricultural activity, (ii) in Iowa, northern Texas and Kansas, due to animal breeding, (iii) in Salt Lake, Cache, and Utah, due to animal farms associated with open waste pits and (iv) in California, due to animal breeding and agricultural practices. Ammonia emissions in North America were 1.1±0.1 Tg yr⁻¹ or two orders of magnitude higher than in EGG (6.2±0.1 kt yr⁻¹) and three times lower than those in VD0.5 (3.1 Tg yr⁻¹) or in VDgrlf (3.4±0.5 Tg yr⁻¹), with maxima observed in late spring, due to fertilization and manure application and summer, due to livestock emissions.

- South America is dominated by natural ammonia emissions mainly from forest, savanna and agricultural fires causing a strong seasonality with the largest fluxes between August and October. The 10-year average ammonia emissions were as high as 28±3.0 Tg yr⁻¹ similar to VD0.5 (22 Tg yr⁻¹) and VDgrlf (24±1.3 Tg yr⁻¹) and four times higher than EGG (7.1±0.3 Tg yr⁻¹).

- In Southeastern Asia, the 10-year average ammonia emissions were 38±2.8 Tg yr⁻¹, in agreement with VD0.5 (36 Tg yr⁻¹) and VDgrlf (39±1.8 Tg yr⁻¹) and slightly higher than...
those in EGG (25±1.2 Tg yr⁻¹). The main sources were from fertilizer plants in China, Pakistan, India and Indonesia. China and India hold the largest share in the ammonia emissions mainly due to rice, corn and wheat cultivation. A strong seasonality in the emissions was observed with a late summer peak in most years, due to rice cultivation, synthetic fertilizer and livestock manure applications and in spring due to wheat cultivation.

- About 88% of the modelled concentrations over Europe using EGG were inside the 10-fold limit from the observations, higher than those with VD0.5 (83%), VDgrlf (85%) and NE (80%). All emission datasets overestimate of ammonia in Eastern Europe, EGG captures better Central Europe, NE emissions predict concentrations in Spain and VD0.5 with VDgrlf emissions in Italy.

- In North America, 90% of the modelled concentrations using EGG emissions were less than 10 times different from the measured ones; more than 95% of the modelled concentrations in North American stations were in the same range using NE, VD0.5 and VDgrlf emissions. NE emissions better capture levels in the easternmost stations of the US closer to the respective hot-spot regions, whereas EGG emissions perform better in Northwestern and Central USA. VD0.5 and VDgrlf emissions perform well in most of the North American stations.

- All emissions underestimate station concentrations in Southeastern Asia. The least spread model concentrations were those simulated using VD0.5 and VDgrlf. About 81% of modelled concentrations using EGG were in the 10-fold limit of the 1x1 line with observations, 88% using NE and only 95% and 94% using VD0.5 and VDgrlf, respectively. VD0.5 and VDgrlf emissions capture well the Japanese and Taiwanese stations.

- The comparison of the modelled ammonia with satellite observations from CrIS globally showed that the best agreement was achieved using the VDgrlf emissions in 2012–2014. After 2015, all satellite retrieved emissions show a better agreement with CrIS concentrations.

Overall, the satellite-constrained ammonia emissions calculated using a variable lifetime appear to give more realistic concentrations, with respect to station and satellite measurements. Accordingly, state-of-the-art emissions appear to underestimate ammonia significantly.
Data availability. All data and python scripts used for the present publication are open through the web address https://folk.niu.no/~nikolaos/AMMONIA/ or can be obtained from the corresponding author upon request.

Competing interests. The authors declare no competing interests.

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Author contributions. N.E. performed the simulations, analyses, wrote and coordinated the paper. S.E. contributed to the lifetime calculations. Y.B., D.H. and A.C. set up the CTM model. M.V.D., P.-F.C. and L.C. provided the IASI data, while M.W.S. and K.E.C.-P. provided the observations from CrIS. All authors contributed to the final version of the manuscript.

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Figure 1. (a) Average model lifetime of ammonia calculated from the LMDz-OR-INCA model, (b) total annual emissions averaged over the 10-year period, (c) atmospheric burden of the reactants sulfuric and nitric acid calculated in the model, and (d) monthly timeseries of lifetime (black), ammonia emissions (green), sulfate (red) and nitrate column concentrations (blue) for the whole 10-year period.
Figure 2. (a) Annual average total column sulfur dioxide anomaly after 2015 from OMI, and (b) annual average emission anomaly of ammonia calculated from IASI in the present study (NE).
Figure 3. Global differences of ammonia emissions calculated in the present study (NE) from those calculated using Van Damme et al. (2018) gridded concentrations applying a constant lifetime of 0.5 days (VD0.5). The results are given as 10-year average (2008–2017) and the number denotes the annual difference in the emissions.
Figure 4. Total annual emissions of ammonia averaged over the 10-year period (2008–2017) in Europe, North and South America, and Southeastern Asia, which are regions characterized by the largest contribution to the global ammonia budget. In the bottom panels, the monthly variation of the emissions is shown for each year of the study period.
Figure 5. Validation of modelled concentrations of ammonia for different emissions datasets (EGG, VD0.5, NE and VDgrlf) against ground-based measurements from EMEP for the 10-year (2008–2017) study period. Scatterplots of modelled against measured concentrations for the aforementioned emission inventories were plotted with the Kernel density estimation, which is a way to estimate the probability density function (PDF) of a random variable in a non-parametric way.
Figure 6. Validation of modelled concentrations of ammonia for different emissions datasets (EGG, VD0.5, NE and VDgrlf) against ground-based measurements from AMON for the 10-year (2008–2017) study period. Scatterplots of modelled against measured concentrations for the aforementioned emission inventories were plotted with the Kernel density estimation, which is a way to estimate the probability density function (PDF) of a random variable in a non-parametric way.
Figure 7. Validation of modelled concentrations of ammonia for different emissions datasets (EGG, VD0.5, NE and VDgrlf) against ground-based measurements from EANET for the 10-year (2008–2017) study period. Scatterplots of modelled against measured concentrations for the aforementioned emission inventories were plotted with the Kernel density estimation, which is a way to estimate the probability density function (PDF) of a random variable in a non-parametric way.
Figure 8. Kernel density estimation (KDE) of the probability density function (PDF) of modelled versus CrIS concentrations of ammonia in a non-parametric way. Modelled concentrations are results of simulations using different emissions datasets (EGG, VD0.5, NE and VDgrlf) for 2012–2017.
Figure 9. 10-year average relative uncertainty of modelled surface concentrations expressed as the standard deviation of surface concentrations from a model ensemble (Table 1) divided by the average.
Table 1. Model ensemble simulations using different emissions for ammonia that were used in the calculations of uncertainty. Uncertainties were calculated as the standard deviation of the surface concentrations of ammonia from the 10 ensemble members for the 10-year period (2008–2017).

<table>
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<th>Parameter perturbed</th>
<th>10-year average emissions (Tg yr⁻¹)</th>
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